1 Root water uptake patterns are controlled by tree species

2 interactions and soil water variability

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17 Abstract

18 Root water uptake depends on soil moisture which is primarily fed by throughfall in forests. Several biotic 19 and abiotic elements shape the spatial distribution of throughfall. It is well documented that throughfall 20 patterns result in reoccurring higher and lower water inputs at certain locations. However, how the spatial 21 distribution of throughfall affects root water uptake patterns remains unresolved. Therefore, we 22 investigate root water uptake patterns by considering spatial patterns of throughfall and soil water patterns 23 in addition to soil and neighboring tree characteristics. In a beech-dominated mixed deciduous forest in a 24 temperate climate, we conducted intensive throughfall sampling at locations paired with soil moisture 25 sensors during the 2019 growing season. We employed a linear mixed-effects model to understand 26 controlling factors for root water uptake patterns. Our results show that soil water patterns and interactions 27 among neighbouring trees are the most significant factors regulating root water uptake patterns. 28 Temporally stable throughfall patterns did not influence root water uptake patterns. Similarly, soil 29 properties were unimportant for spatial patterns of root water uptake. We found that wetter locations

30 (rarely associated with throughfall hotspots) promoted greater root water uptake. Root water uptake in 31 monitored soil layers also increased with neighbourhood species richness. Ultimately our findings suggest 32 that complementarity mechanisms within the forest stand, in addition to soil water variability and 33 availability, govern root water uptake patterns.

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35 Key words: root water uptake, throughfall, soil water, spatial patterns, beech

36 1) Introduction

37 Root water uptake depends on soil moisture, which is replenished by precipitation. At the same time, the 38 vegetation canopy intercepts and redirects precipitation into throughfall and stemflow, collectively 39 referred to as below-canopy precipitation. Thus, even before soil water can be taken up by roots, it has 40 already been influenced by the canopy.

41 Throughfall is typically the largest component of below canopy precipitation (Levia and Frost, 2006; 42 Sadeghi et al., 2020). For instance, in temperate forests about 70% of above canopy precipitation ends up 43 as throughfall (Levia and Frost, 2003; Sadeghi et al., 2020). Below-canopy precipitation is modified by 44 several biotic and abiotic factors (Levia and Frost, 2006; Levia et al., 2011), including vegetation type, 45 canopy architecture (Crockford and Richardson, 2000; Pypker et al., 2011; Levia et al., 2017), and forest 46 structure (Rodrigues et al., 2022), meteorological elements such as wind speed (Staelens et al., 2008; Van Stan et al., 2011; Fan et al., 2015), precipitation intensity and event size (Dunkerley, 2014; Magliano et 47 48 al., 2019; Zhang et al., 2016; Staelens et al., 2008). As a result, throughfall inherently varies across space 49 and time. However, previous studies showed that the spatial distribution of throughfall persists over time 50 (Keim et al., 2005; Staelens et al., 2006; Guswa and Spence, 2012; Carlyle-Moses et al., 2014; Metzger 51 et al., 2017; Van Stan et al., 2020).

52 Throughfall patterns have been hypothesized to affect the spatial variation in water uptake (Bouten et al., 53 1992; Coenders-Gerrits et al., 2013) and soil moisture distribution (Raat et al., 2002; Blume et al., 2009; 54 Zimmermann et al., 2009; Zehe et al., 2010; Bachmair et al., 2012; Rosenbaum et al., 2012; Zhang et al., 55 2016). Yet, empirical evidence is scarce. A decade ago Coenders-Gerrits et al., (2013) proposed that

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56 throughfall patterns are translated into soil wetting dynamics with a model based on combined hillslope 57 topographic and throughfall data collected in a beech-dominated catchment. However, in this model, the 58 effect of throughfall patterns on soil moisture patterns rapidly ceased, and became more similar to the 59 bedrock topography. Regarding the latter result, the model and reality differ, as the correlation between 60 measured bedrock topography and soil moisture is low (Tromp-van Meerveld and McDonnell, 2006). 61 which Coenders-Gerrits et al., (2013) attributed to root water uptake. Later, Metzger et al. (2017) showed 62 through field observations that although throughfall spatial variation strongly increases shortly after 63 rainfall it drops quickly again in the drained state, so the impact rapidly disappears. Later, Fischer-Bedtke 64 et al., (2023) confirmed in the same field site that recurring throughfall patterns left a notable imprint on 65 soil moisture response to rainfall yet the effect on absolute values of soil water content in drained state 66 was rather weak. More recently, Zhu et al. (2021) observed that stable throughfall patterns were weakly 67 related to the spatial distribution of soil moisture since this relationship was restricted only to relatively 68 wet soil locations and throughfall hotspots. They also showed that throughfall patterns had weaker 69 influence on the temporal dynamics of soil water content compared to soil bulk density and litter layer 70 properties.

71 Taken together, several studies have searched for patterns of throughfall in soil moisture spatial variation. 72 As comparatively weak relationships were found, some previous studies have suggested that root water 73 uptake (Bouten et al., 1992; Schwärzel et al., 2009) could be the cause. Specifically, based on a one-74 dimensional soil-water model, Bouten et al. (1992) proposed that throughfall patterns alter and localize 75 root water uptake as well as promote fast drainage. As a result, spatial variation in root water uptake could 76 diminish the effect of throughfall patterns into spatio-temporal variation of soil water. However, other 77 researchers suggested that other factors, such as soil properties (Metzger et al., 2017), preferential flow 78 (Jost et al., 2004; Blume et al., 2009; Molina et al., 2019; Fischer-Bedtke et al., 2023) and litter layer 79 processes (Raat et al., 2002) may be at the heart of the weak and short-term effects of throughfall patterns 80 on soil moisture variability.

81 However, to the best of our knowledge, the feedback mechanism of throughfall patterns on root water 82 uptake variation has not yet been investigated empirically. More common are studies related to soil water 83 distribution. Soil water availability, which could potentially be enhanced by throughfall, affects root water uptake patterns even more than root abundance (Kühnhammer et al., 2020; Guderle et al., 2018). On the
flip side, root water uptake can amplify or homogenize soil water variability (Hupet and Vanclooster,
2005; Teuling and Troch, 2005; Ivanov et al., 2010; Baroni et al., 2013; Martínez García et al., 2014).
Moreover, variations in soil water content reflect on root water uptake (Hupet et al., 2002; Schume et al.,
2004; Schwärzel et al., 2009; Guderle and Hildebrandt, 2015; Jackisch et al., 2020).

Next to water input, soil properties can alter root water uptake patterns (Nadezhdina et al., 2007; Kirchen et al., 2017). Also they control soil water redistribution (Grayson et al., 1997; Cosh et al., 2008; Jarecke et al., 2021) and water availability for root structures (Vereecken et al., 2007; Cai et al., 2018). For a given evaporative demand, water uptake at a particular location is a function of water transport resistance between root and soil in addition to the soil-water potential (Cardon and Letey, 1992; Shani and Dudley, 1996; Lhomme, 1998). Both characteristics depend on local soil properties and soil water status, and the latter in turn is affected by the local water uptake rate.

Finally, plant individual and ecosystem processes affect uptake: Root networks can connect wetter and
drier locations in a variety of ecosystems (e.g., Emerman and Dawson, 1996; Katul and Siqueira, 2010;
Yu and D'Odorico, 2015; Priyadarshini et al., 2016; Hafner et al., 2017). In addition, tree size, age,
neighboring tree species, and ecosystem structure affect the spatio-temporal variation in root water uptake
(Volkmann et al., 2016; Spanner et al., 2022; Kostner et al., 2002; Dawson, 1996; Brinkmann et al., 2019;
Gaines et al., 2016; Silvertown et al., 2015; Guo et al., 2018; Brum et al., 2019; Krämer and Hölscher,
2010).

103 Taken together, throughfall and soil water variability, soil properties, and root water uptake patterns form 104 complex and intertwined interactions in the terrestrial hydrological cycle. It has not yet been shown 105 empirically how root water uptake patterns are affected by throughfall and spatial distribution of soil 106 water content. In line with previous modeling results (Bouten et al., 1992; Coenders-Gerrits et al., 2013) 107 we hypothesize that throughfall hotspots enhance water availability at certain locations that elevate root 108 water uptake. Further we investigate the role of soil water variation in combination with soil properties 109 and neighboring tree characteristics on root water uptake patterns. We pose the following questions to 110 test the main hypothesis and guide the investigation:

111 i) How do throughfall patterns influence root water uptake patterns?

- 112 ii) How does soil moisture and its variation, along with soil properties, control variation in root113 water uptake?
- 114 iii) What is the role of biotic factors, namely size, distance, number, and species richness of115 neighbouring trees on root water uptake patterns?

Here, we address these questions by employing a linear mixed effects model based on weekly throughfall 116 117 sampling at locations paired with intensive soil moisture measurements in a beech-dominated unmanaged 118 forest. We estimate root water uptake using a water balance method applied at soil moisture measurement 119 point. This method dissects soil water flow and water uptake by exploring the differences in soil water 120 content change per time between day and night (Guderle and Hildebrandt, 2015; Jackisch et al., 2020). 121 While other methods exist, such as using isotopic tracers (Rothfuss and Javaux, 2017, Zarebanadkouki et 122 al., 2013), daily fluctuations in soil water allow for estimating the spatial distribution of ecosystem 123 evapotranspiration using standard measurements of soil water content (Guderle and Hildebrandt, 2016) 124 without the need for additional infrastructure. In addition, we incorporate data on field capacity, bulk 125 density, and neighboring tree characteristics namely size and species.

126 **2) Materials and Methods**

127 **2.1) Research Site and Field Sampling**

128 **2.1.1) Research Site**

The research site is located in the forested upper hill region of the Hainich low mountain range in Thuringia, Germany, as a part of the Hainich Critical Zone Exploratory (CZE) (Küsel et al., 2016). The altitude in the research site ranges from 362 m to 368 m a.s.l. Mean annual air temperature varies between 7.5 and 9.5 °C, and the mean annual precipitation ranges from less than 600 to 1000 mm in the CZE (Küsel et al., 2016).

In the study area, thin-bedded alternations of limestones and marlstones of carbonate rock (Middle
 Triassic) form the bedrock overlain by a shallow Pleistocene loess layer with cambisols and luvisols as

136 dominant soil types (IUSS Working Group, 2006; Metzger et al., 2021). The median soil depth above the

- 137 weathered bedrock is 37 cm, with soil depths ranging from 15 cm to a maximum depth of 87 cm (Metzger138 et al., 2017).
- In 2019, the tree community in the research site consisted of 574 individuals of various ages (diameter at breast height \geq 5cm). The dominant species is European beech (*Fagus sylvatica* L.), which makes up 70% of the tree community, followed by sycamore maple (*Acer pseudoplatanus* L.) with 21 %, and European ash (*Fraxinus excelsior* L.) with 4%. These dominant species are accompanied by Large-leaved linden (*Tilia platyphyllos* Scop.), European hornbeam (*Carpinus betulus* L.), Norway maple (*Acer platanoides* L.), Scots elm (*Ulmus glabra* L.), and Wild service tree (*Sorbus torminalis* (L.) Crantz). The stand has a total basal area of 40 m² ha⁻¹ and has been unmanaged since 1997 (Kohlhepp et al., 2017).

146 **2.1.2**) Soil moisture monitoring and soil properties

147 The forest site (1 ha) was equipped with a soil moisture monitoring network (SoilNet; Bogena et al., 2010) 148 consisting of SMT100 frequency domain sensors (Treuebner GmbH, Neustadt, Germany). Metzger et al. 149 (2017) first described the soil moisture monitoring setup. Briefly, the observation platform (Figure 1) was 150 divided into 100 subplots (10 m \times 10 m), and 49 subplots were equipped with soil moisture sensors at 151 two random measuring points each, for a total of 98 locations. At each measuring point, sensors were placed at two different depths, 7.5 cm (top sensors) and 27.5 cm (bottom sensors). The soil moisture 152 153 network is maintained through a regular bi-weekly routine to avoid potential failures such as depleted 154 sensors batteries, hardware problems, etc.

Undisturbed soil samples were collected during the sensor installation in 2014 and 2015 to estimate bulk density and water content at field capacity. In addition, we collected additional disturbed soil samples (n = 40) near sensor locations in 2019. Bulk density was determined from oven-dried (24h, 105°C) soil mass weight and water content at field capacity by applying 60 hPa pressure to the saturated undisturbed sample for 72 h.

Soil properties vary slightly from top to subsoil at the research site. While silty loam is the dominant soil texture in both layers, the clay content is higher in the subsoil (Metzger et al., 2021). The median volumetric water content at field capacity is 44% in the topsoil and 42% in the subsoil. Moreover, the water content at field capacity varies from 27% to 60% and from 31% to 62% in the topsoil and subsoil,

- 164 respectively. The average bulk density (d_{bulk}) of the topsoil is 1.16 g cm⁻³, with a range of 0.73 to 1.5 g
- 165 cm⁻³. In the subsoil, the average bulk density (d_{bulk}) is slightly higher at 1.37 g cm⁻³ but has a similar range
- 166 $(0.7 1.6 \text{ g cm}^{-3})$ (See supplement for details).



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Figure 1 (above) The photo of the site. (below) the field monitoring setup of stratified randomly distributed throughfall collectors and soil moisture sensors together with the trees which are sized according to the diameter at breast height (dbh)

170 and coloured according to the species. Throughfall collectors are paired with soil moisture sensors at 98 locations (n=182) in 171 the grey shaded subplots. White coloured subplots are equipped with only throughfall collectors.

172 **2.1.3**) Gross precipitation and throughfall sampling

173 Five gross precipitation funnels were placed 1.5 m above ground level in an adjacent open grassland (ca.

174 250 m distance to the research site). As described in Metzger et al. (2017) and Demir et al. (2022), the

175 precipitation funnels were made of a circular plastic funnel (12 cm in diameter) and sampling bottle (2 L

176 in volume), and ping pong balls were placed in the funnel orifice to prevent evaporation losses.

During the early growing season of 2019, we placed throughfall collectors in soil moisture monitoring subplots at 98 locations. We paired these throughfall collectors with the soil moisture sensors by placing them within 1 m of each other. The paired collectors were placed down-slope to avoid interference with soil moisture measurements. For the rest of the research site, in 51 other subplots, we adopted a separate independent stratified random design from Metzger et al. (2017). Briefly, we placed two throughfall collectors in each subplot that was not equipped with soil moisture sensors. All throughfall collectors

183 were placed roughly 37 cm above the ground.

We conducted weekly manual measurement of throughfall and gross precipitation during the 2019 growing season (April to August). Sampling was conducted on rain free days only. Thus, the sampling interval ranged between six and eight days.

We used the paired throughfall collectors (n = 98) to identify the drivers of root water uptake patterns, as we derived root water uptake values based on soil water content measurements (see below). However, we used all randomly placed throughfall collectors (n = 200) to describe the spatio-temporal variation of throughfall within the research site.

191 **2.2) Estimation of potential evapotranspiration**

We calculated the daily potential evapotranspiration by applying the concept of thermodynamic limits ofconvection (Kleidon and Renner, 2013; Kleidon et al., 2014):

194
$$E_{\text{pot}} = \frac{1}{\lambda} \frac{s}{s+\gamma} \frac{R_{\text{sn}}}{2}$$
(1)

195 Where R_{sn} is absorbed solar radiation (W m⁻²), λ is the latent heat of vaporization (2.5×10⁶ Jkg⁻¹), γ is 196 the psychrometric constant (65 PaK⁻¹), and *s* is the slope of the saturation vapor pressure curve (PaK⁻¹). 197 Here, we acquired solar radiation, air temperature, and precipitation data for the throughfall sampling

198 period from a nearby weather station ("Reckenbuel") which is located approximately 1.4 km northeast of

the research site and provides data in 10 minutes intervals. The site-specific albedo for the summer periodwas adopted from Otto et al. (2014).

We used the precipitation data measured at the weather station to define rain events and dry periods, asdescribed below.

203 2.3) Data analysis

204 **2.3.1) Quality control of soil water content data**

205 We systematically reviewed the six-minute soil water content data for quality control in two steps: 1) 206 identification of problems (such as jumps to extremely low and high values, duplicated time stamps of 207 different values, long discontinuities in the measurements, and lack of temporal variation in the time series 208 despite rain events), 2) classification and removal of detected outliers and irregularities. We visually identified and removed unrealistic measurements such as extremely low (< 5 vol-%) and high values far 209 210 beyond the field capacity (> 75 vol-%) and long plateaus of repeated values despite rain events. We also 211 excluded any time series that exhibited long-term discontinuities that prevented us from calculating root 212 water uptake. During the visual inspection, we eliminated values with duplicated time stamps that violated 213 the actual temporal trend. Next, we scanned the data using the Hampel filter function of the 'pracma' R 214 package (Borchers, 2021) with customized moving window length and Pearson's rule threshold value 215 (Pearson, 1999) to flag possible outliers.

Despite regular maintenance, many sensors failed to provide data that met the quality criteria during the growing season (March-August) in 2019. Only 56 sensor locations (out of 98) provided data from both top and bottom sensors that met the qualification criteria described above with varying date intervals throughout the growing season. Of these, only 34 sensor locations were used to estimate root water uptake as they simultaneously provided data from both top and bottom sensors within the dry periods.

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221 **2.3.2**) Soil water calculation

We estimated soil water (S) at measurement locations for the monitored soil layer based on volumetric soil water content measured by top and bottom sensors.

224
$$S_{i,d} = \sum z_t \theta_{i,d}^t + z_b \theta_{i,d}^b$$
(2)

225 We similarly integrated the soil water at field capacity $(S_{FC,i})$

226
$$S_{FC,i} = \sum z_t \theta_{FC,i}^t + z_b \theta_{FC,i}^b$$
(3)

where z_t is the depth of the soil column monitored by the top sensor and z_b is the depth of soil represented by the bottom sensor, and $\theta_{i,d}$ is volumetric soil water content at location *i* on date *d*, and $\theta_{FC,i}$ the soil water content at the field capacity.

230 We calculated bulk density at the sensors' locations for the monitored soil layer.

231
$$\overline{d_{bulk,i}} = \frac{\sum z_t d_{bulk,i}^t + z_b d_{bulk,i}^b}{\sum z_t + z_b}$$
(4)

where $d_{bulk,i}^{t}$ and $d_{bulk,i}^{b}$ are the bulk density of the topsoil and subsoil, respectively, at location *i*.

233 **2.3.3**) Descriptive Statistics

We calculated the coefficient of quartile variation (CQV) and the interquartile range to describe spatial variation of throughfall, volumetric soil water content, and root water uptake. Also, we estimated octile skewness (OS₈) of throughfall based on the first and seventh octile.

237
$$CQV = \frac{Q_3 - Q_1}{Q_3 + Q_1}$$
 (5)

238
$$OS_8 = \frac{(Q_7 - median) - (median - Q_1)}{Q_7 - Q_1}$$
 (6)

We characterized spatial patterns of daily root water uptake (E_t) by calculating the spatial deviation from the mean ($\delta E_{t,i,d}$, Equation 7) (Vachaud et al., 1985).

241
$$\delta E_{t\,i,d} = \frac{E_{t,i,d} - \overline{E_{t,d}}}{\overline{E_{t,d}}}$$
(7)

where $E_{t, i, d}$ is daily root water uptake estimated at *i* sensor location on date *d* and $\overline{E}_{t, d}$ is spatial average of daily root water uptake on date *d*.

244 Similarly, we calculated the spatial deviation of soil water and throughfall to identify their spatial patterns.

245 **2.4) Root water uptake estimation**

We estimated root water uptake using the multi-step, multi-layer regression method (MSML), which is a water-balance method and derives evapotranspiration from diurnal differences in soil water content (Guderle and Hildebrandt, 2015; Guderle et al., 2018). This approach does not require prior information on root structure but relies on high temporal and spatial resolution data on multiple soil layers. Previous studies using additional measurements such as sap-flow and lysimeters demonstrated that the MSML method successfully estimates transpiration in both forest and grassland ecosystems (Guderle et al., 2018; Jackisch et al., 2020).

As described in Guderle and Hildebrandt (2015), the MSML derives root water uptake from distinct differences in the day and night portions of soil moisture time series. The main assumption is that, in the absence of rainfall-driven rapid vertical soil water flow, evapotranspiration occurs only during the day, while soil water flow occurs both during the day and at night. As a result, soil moisture time series reflect a distinct day/night signal under dry weather conditions.

In applying this method to our study, we first excluded potential periods of fast vertical flow periods from the time series due to previous rainfall events and identified periods for estimating daily root water uptake. We considered an 8 h buffer period to include canopy dripping and 48 h for the cessation of rainfall influence on soil water. Thus, a total of 56 h was the time interval used to define the start of the water uptake estimation period. The period when the root water uptake is estimated is hereafter referred to as the dry period.

264 Next, we split each soil moisture time series into a day (transpiration active period) and a night branch, 265 as explained by Guderle and Hildebrandt (2015). We defined the transpiration period (starts 2 h after 266 sunrise and ends 2 h before sunset) based on local sunrise and sunset time. Sunrise and sunset times were 267 obtained from the R package 'suncal' (Thieurmel and Elmarhraoui, 2022). We fit linear models to each 268 split branch of the time series and derived the slopes. The difference between the slope of the day branch 269 (m_{tot}) and the average slope of the antecedent and preceding night $(\overline{m_{flow,l}})$ gives the rate of water uptake. 270 Thus, we estimated daily evapotranspiration at each soil water content location i (Equation 8, 9) by 271 accounting for soil layer thickness and slope difference-

272

273
$$E_{t,msml,i}^{t,b} = (m_{tot,i}^{t,b} - \overline{m_{flow,i}^{t,b}}) d_{z,i_{z,i}}^{t,b}$$
 (8)
274 $E_{t,i} = \sum (E_{t,msml,i}^{t} + E_{t,msml,i}^{b})$ (9)

276 2.5) Linear Mixed Effects Model

277 We employed a linear mixed effects model to investigate the driving factors for root water uptake patterns. A linear mixed effects model is a multivariate statistical tool that describes the relationship between a 278 279 dependent variable and explanatory variables (fixed effects) while controlling for dependencies in the 280 data that may arise due to repeated sampling with certain designs (random effects). Fixed effects are 281 informative, repeatable levels of explanatory and quantified variables that can influence the mean of the 282 dependent variable, and they can be tested. In addition, in a linear mixed-effects model, how the 283 relationship between the dependent variable and one predictor depends on the level of another predictor 284 can be represented via interaction term.

Random factors are uninformative levels of predictor variables but can explain parts of the residual of the fixed effects model by calculating different intercepts for different category levels. They are included in mixed effects models to account for qualitative information from repeated sampling with respect to individuals, time stamps, or treatments. Here, sensor location and dry period, i.e. date, are taken as random effects.

For the model, we used only paired throughfall and soil moisture measurement locations where both top and bottom sensors provided data during the dry periods. All considered explanatory drivers, which are included as fixed factors in the model, are listed in Table 1. These factors include abiotic and biotic variables that possibly influence relative local root water uptake: They are daily spatial average soil water storage, the spatial deviation of soil water from the mean, soil water at field capacity and bulk density of the monitored soil layer.

To account for spatial variability in throughfall, we calculated the spatial deviation from the mean by using Equation 7. Here we considered this variable at a two-different time scales: the sampling week(s) prior to root water uptake estimation, and over the entire throughfall sampling period. Further, as biotic factors, we included number of trees, and number of species within a 5 m radius of each soil moisture location, and inverse-distance-weighted basal area (BA) within 5 m radius of each soil moisture location, calculated as follows:

$$302 \qquad BA_i = \frac{\sum_{R=1}^R W_R A_{tree}}{A} \tag{10}$$

303 with
$$W_R = \frac{(x_i - x_R)^2}{\sum_R (x_i - x_R)^2}$$
 (11)

where i is the soil moisture sensor located at x_i , R is the tree index located at x_R , and A_{tree} is the individual basal area of the corresponding tree, A is the area around the soil moisture sensor i with 5 m in radius.

Even though our research plot is a beech-dominated forest, in some spots, two to four species were present
within a 5 m radius of the soil moisture sensors.

308 We also included interaction terms (Table 1) as fixed factors in the model to capture complex and non-309 linear relationships among the biotic and abiotic factors.

We conducted all analyses with the R statistical software (R Core Team, 2022) and used the *lmer* function in the 'lme4' package (Bates et al., 2015) for the model development. We visually checked the model assumptions using the 'check_model' function of the 'performance' package (Lüdecke et al., 2021).

In addition, we calculated both conditional and marginal R^2 of the model with the 'MuMIn' package (Bartoń, 2020). While the conditional R^2 includes the variance of the entire model, the marginal R^2 subsumes only the fixed effects (Bartoń, 2020). Before fitting the linear mixed effects model, we tested

316 for co-linearity of the considered variables and scaled the data with a Z-transformation by using the 'scale'

- 317 function in base R (R Core Team, 2022), which allowed us to evaluate the individual effect of fixed effects
- 318 by comparing slopes and significance levels.

We developed the optimal model by applying a systematic model selection procedure based on Akaike's Information Criterion (AIC) comparison in combination with the examination of the factors. Model selection began with the beyond-optimal model, which included all possible fixed and random effects. We stepwise evaluated each fixed effect based on its respective significance (*p* value comparison) by fitting the model the maximum likelihood (ML) to be able to compare AIC values (Zuur et al., 2009). In each step, starting with interaction terms, we identified the least significant effect and formulated a model without it. We compared the AIC values of the model before and after removing the effect, discarding it 326 in case the AIC was unaffected or decreased. We followed the procedure with the next equally detected

327 effect, and repeated it until only significant fixed effects remained, and the model with the lowest AIC

328 (the optimal model) was obtained.

As a final step, the best model was refitted with restricted maximum likelihood (REML) (Zuur et al.,2009).

331 Table 1 List of fixed and random factors considered for estimating the root water uptake patterns through linear mixed effects 332 model. Interaction is shown with 'x'.

Fixed Factors	
Single Factors	Interaction Factors
Spatial average of soil water storage in the monitored soil layer (\overline{S})	$\overline{S} \times S_{FC}$
Spatial deviation of soil water storage from the mean (δS)	$\delta S \times S_{FC}$
Field capacity of the monitored soil layer (S_{FC})	$\delta S \times BA$
Bulk density capacity of the monitored soil layer (d _{bulk})	$\overline{S} \times BA$
Spatial deviation of throughfall of events measured in sampling week previous to the corresponding dry period $(\delta P_{TF_{last ev.}})$	$\delta S \times n_{tree}$
The median of spatial deviation of throughfall measured within the whole sampling period $(\widetilde{\delta P_{TF}})$	$\overline{S} \times n_{tree}$
Number of trees (n _{tree})	$\delta P_{TF_{last ev.}} \times S_{FC}$
Basal area (BA)	$\delta P_{TF_{temp. stable.}} \times S_{FC}$
Number of species (n _{sp,tree})	$\delta P_{TF_{last ev.}} \times d_{bulk}$
	$\delta P_{TF_{temp. stable.}} \times d_{bulk}$
	$n_{sp,tree} \times WA_{int}$
Random factors	
Soil moisture sensor location	

Dry period

333 3) Results

334 **3.1)** Spatio-temporal distribution of throughfall and soil water content

335 In 12 out of the 16 sampling weeks, the weekly gross precipitation was more than half of the total potential 336 evapotranspiration. Table 2 shows the distribution of throughfall sampled in 2019 (April-August) at 200 337 collectors and the 98 collectors that were paired with soil moisture sensors. Weekly throughfall increased 338 with an increase in rain. The coefficient of quartile variation (CQV) of throughfall was generally lower 339 for larger cumulative weekly rains. On average, the collectors paired with soil-moisture sensors received 340 similar amounts of throughfall to all collectors (Table 2). The COV of data from the paired collectors 341 ranged from 0.27 to 0.6, which is similar to the CQV of throughfall sampled at all collectors. The octile 342 skew (OS_8) of paired and all collectors was also similar.

As the growing season progressed in 2019, the average soil water content decreased in both the topsoil and subsoil. In April and early May, the average volumetric soil water content in the topsoil was above 30%, and dropped to below 10% by the end of August. In the subsoil, the volumetric soil water content similarly declined from above 40 % to below 20 % over the sampling period (Figure 2). On average, soil water changed from 52.5mm to 17.5 mm in the topsoil and from 80 mm to 40mm in the subsoil.

348 We derived root water uptake for four periods (a total of 19 days) under different soil wetness conditions 349 that captured the seasonal variation of soil water content, including late spring when the soil water content 350 was higher and drier periods during the summer following re-wetted soil conditions with late summer 351 rains. As listed in Table 3 and shown in Figure 2, two periods were in late May and early June, and each 352 lasted two days. The third period began in late June and lasted 11 days; the last was four days in late July. 353 From the start of the first dry period to the end of the last, the average soil water content declined from 354 33 to 15 % in the topsoil and from 43 to 27% in the subsoil. Table 3 shows that within the dry periods, 355 the coefficient of quartile variation (CQV) of soil water content was between 0.09 -0.14 and 0.08 to 0.16 356 in the topsoil and subsoil, respectively. During the dry periods, the spatial heterogeneity of soil water 357 content in the subsoil increased systematically. In contrast, the spatial variation of soil water content in 358 the topsoil was not correlated with soil dryness.





Figure 2 Soil moisture temporal variation in top and subsoil together with the daily precipitation measured at the nearby Reckenbühl station (approximately 1.4 km to the Northeast). The solid and dashed lines are spatial mean of soil water content estimated based on top (7.5 cm) and bottom (27.5 cm) sensors, and grey shaded areas show first and third quartiles. The reddish shaded areas show defined dry periods within the throughfall sampling when root water uptake could be estimated.

P_{a}/F_{aab} $\overline{P_{aab}}$ $1\overline{OR}$ \overline{COV} \overline{OS}_8
a g/ - pot a TF P _{TF} P _{TF} P _{TF}
0.06 0.35 0.18 0.25 0.46
0.08 0.97 0.44 0.24 0.16
0.43 1.72 0.27 0.08 0.2
0.42 2.58 0.62 0.12 -0
0.62 3.77 1.24 0.17 -0
0.58 4.61 1.06 0.12 -(
0.96 5.19 1.06 0.10 -(
0.85 7.81 2.25 0.15
1.89 9.21 1.33 0.07 -
1.62 11.19 2.65 0.12 -
1.72 16.60 2.65 0.08 -
4.13 18.44 3.09 0.08
3.54 24.22 3.54 0.07
3.98 22.10 3.54 0.08
6.20 30.94 3.54 0.06
4.96 57.77 8.51 0.07

Table 3 The spatial average of daily volumetric soil water content ($\overline{\theta}_{top-soil}$, vol-%) in topsoil (0-17.5 cm), and ($\overline{\theta}_{subsoil}$, vol-%) in subsoil (17.5 – 37.5 cm) during the defined dry periods. The inter quartile range (IQR), and coefficient of quartile variation (COV) of daily volumetric soil water content in both layers during the dry periods.

Date	$\overline{oldsymbol{ heta}_{top-soil}}$ (vol-%)	IQR θ _{top-soil} (vol-%)	$\begin{array}{c} CQV \hspace{0.1 cm} \theta_{top-soil} \\ (vol-\%) \end{array}$	$\overline{oldsymbol{ heta}_{sub-soul}}$ (vol-%)	$\begin{array}{c} IQR \theta_{subsoil} \\ (vol-\%) \end{array}$	$\begin{array}{c} CQV \theta_{subsoil} \\ (vol-\%) \end{array}$	Dry Period
25 -05-2019	33.17	5.72	0.09	42.82	6.72	0.08	1
26-05-2019	32.12	6.62	0.10	42.46	6.67	0.08	1
01-06-2019	30.23	6.87	0.12	40.61	6.9	0.09	2
02-06-2019	29.22	7.23	0.13	40.11	6.85	0.09	2
23-06-2019	25.01	6.69	0.14	37.80	6.38	0.08	3
24-06-2019	24.04	6.45	0.14	36.94	6.22	0.08	3
25-06-2019	22.52	5.43	0.12	36.13	6.54	0.09	3
26-06-2019	21.48	5.07	0.12	35.24	6.71	0.10	3
27-06-2019	20.20	4.25	0.11	33.98	7.75	0.12	3
28-06-2019	19.45	3.85	0.10	33.31	8.08	0.12	3
29-06-2019	18.98	3.83	0.10	32.36	8.05	0.12	3
30-06-2019	18.44	3.52	0.09	31.37	8.15	0.13	3
01-07-2019	17.67	3.62	0.10	30.45	8.18	0.13	3
02-07-2019	17.29	4.18	0.12	29.84	8.87	0.15	3
03-07-2019	16.89	3.72	0.11	29.26	8.98	0.15	3
24-07-2019	16.15	3.48	0.11	28.56	8.7	0.16	4
25-07-2019	15.51	3.47	0.11	27.85	8.67	0.16	4
26-07-2019	14.98	3.57	0.12	27.21	8.49	0.16	4
27-07-2019	14.57	3.65	0.13	26.65	8.63	0.16	4

368

369 3.2) Soil water storage, potential evapotranspiration, and root water uptake

370 The integrated field capacity of the monitored soil depth was 160 mm on average at the research site. 371 Table 4 shows that soil water was much lower than the field capacity during the dry periods, and the mean 372 soil water storage dropped below 42 mm in late July. In addition, Table 4 demonstrates that the average root water uptake (\overline{E}_t) ranged from 0.94 mm d⁻¹ to 3 mm d⁻¹ while potential evapotranspiration (E_{pot}) 373 ranged from 1.75 mm d⁻¹ to 3.12 mm d⁻¹. The discrepancy between average root water uptake and the 374 375 potential evapotranspiration increased as soil water decreased, especially during the longest dry period 376 (Table 4). Root water uptake showed greater spatial variation than water input and soil wetness. The 377 coefficient of quartile variation (CQV) of root water uptake ranged from 0.15 to 0.28, which was higher 378 than the CQV of throughfall and volumetric soil water content in both soil layers.

Table 4 The daily average air temperature (T_{air} , °C), potential evapotranspiration (\underline{E}_{pot} , mm), mean soil water storage (\overline{S} , mm)

in monitored soil layer (0 - 37.5 cm), and spatial mean of daily root water uptake ($\overline{E_t}$, mm) based on all soil moisture sensors, and the ratio of the root water uptake to the potential evapotranspiration together with and standard deviation (SD) and are experimented as a standard deviation (SD) and the ratio of the daily root water uptake during the defined dry particle.

382 coefficient of quartile variation (CQV) of the daily root water uptake during the defined dry periods

Date	T _{air} (°C)	E _{pot} (mm)	<u> </u>	$\overline{E_t}$ (mm)	$\overline{E_t} / E_{\text{pot}}$	SD $\overline{E_t}$	CQV $\overline{E_t}$	Dry Period
25-05-2019	12.74	1.80	71.94	1.09	60.56	0.38	0.28	1
26-05-2019	14.43	1.90	70.57	1.30	68.42	0.48	0.25	1
01-06-2019	18.42	2.59	67.16	2.26	87.26	0.98	0.27	2
02-06-2019	21.38	2.77	65.79	2.50	90.25	1.12	0.18	2
23-06-2019	19.45	2.79	59.81	2.83	101.43	0.90	0.19	3
24-06-2019	20.22	2.82	58.16	2.62	92.91	0.76	0.17	3
25-06-2019	22.52	2.89	55.96	2.67	92.39	0.78	0.16	3
26-06-2019	25.73	2.96	54.13	3.00	101.35	0.88	0.15	3
27-06-2019	18.83	2.75	51.91	2.28	82.91	0.55	0.16	3
28-06-2019	16.07	2.58	50.55	1.53	59.30	0.40	0.20	3
29-06-2019	19.59	2.85	49.55	2.11	74.04	0.60	0.20	3
30-06-2019	25.54	3.12	48.26	2.57	82.37	0.86	0.18	3
01-07-2019	20.63	2.30	46.69	1.59	69.13	0.53	0.18	3
02-07-2019	14.88	1.75	45.81	1.08	61.71	0.42	0.24	3
03-07-2019	13.77	1.91	44.95	0.94	49.21	0.30	0.23	3
24-07-2019	24.39	2.76	43.61	1.88	68.12	0.64	0.19	4
25-07-2019	25.33	2.82	42.31	1.77	62.77	0.60	0.24	4
2019-07-26	23.27	2.64	41.18	1.40	53.03	0.55	0.18	4
2019-07-27	21.29	2.68	40.23	1.21	45.15	0.47	0.19	4

383 **3.3)** Soil water, throughfall, and root water uptake patterns

384 At soil moisture measurement points where daily root water uptake was determined (n = 34), we 385 calculated the spatial deviation from the median of throughfall, soil water storage, and root water uptake 386 to illustrate the spatial patterns. Figure 3 shows that some locations received repeatedly less (or more) 387 throughfall than average ($\delta P_{TF} < 0$), some locations were repeatedly wetter or drier ($\delta S < 0$), and some 388 places regularly had lower or higher root water uptake (δE_t) throughout the sampling period. However, 389 these locations were not related to each other. In fact, Figure 3 demonstrates that neither throughfall nor 390 soil water patterns are directly correlated with the root water uptake patterns. For example, the locations 391 with higher water uptake were not coupled with elevated throughfall input (locations coloured dark) or 392 higher soil water storage. In addition, soil water storage patterns were not correlated with throughfall 393 patterns.



394

Figure 3 Temporal stability of throughfall patterns which is estimated by the spatial deviation from the mean (δP_{TF}) throughout the sampling period in 2019 (April-August), soil water (δS) and root water uptake (δE_t) based on the spatial deviation from the mean during the defined dry periods. Soil moisture sensor locations colored according to throughfall input. Soil moisture sensor locations are colored from lighter to darker in the throughout figure according to throughfall input.

399 3.4) Fixed factors regulating root water uptake patterns

400 We used a linear mixed effects model to disentangle the effects of throughfall, soil water, soil properties,

401 and the neighbouring tree characteristics on root water uptake patterns. The fixed and random effects

- 402 contributed almost equally to the model. The R^2 of the model was 0.77, and the contribution of the fixed
- 403 effect to the R^2 was 0.39 (See the supplement for more details on the optimal model).
- 404 Figure 4 shows only the significant fixed effects for root water uptake patterns. Spatial deviation of soil
- 405 water from the mean (i.e., soil water patterns) was the only single and the most significant factor positively

related to the spatial deviation of root water uptake. Thus, water uptake was elevated at locations wherethe most water was retained in the soil at the given time, i.e., greater soil water storage.



408

Figure 4 The significant fixed factors of the best model to estimate root water uptake patterns (δE_t). Values on the x-axis indicate the slope of the relations. All variables were scaled by Z-transformation. Interaction is shown with 'x'. Here δS is the spatial deviation of soil water, S_{FC} is the field capacity, $n_{sp,tree}$ is the number of species, BA is the basal area, and \overline{S} is soil water storage. Significance codes are *** $\cong 0, ** \cong 0.001$. (the details on the model can be found in the supplement)

Field capacity by itself was not a significant factor affecting local root water uptake. However, it strongly influenced how local soil water controlled root water uptake as a part of the significant interaction term.Figure 5a illustrates how to root water uptake was more dependent on local soil water when field capacity was low (i.e., higher macroporosity). In contrast, soil bulk density and therefore total porosity was not part of the final model.

Although the spatial average of soil water storage, e.g., the state of wetness, was not an important factor for local root water uptake by itself, it moderated the impact of basal area (BA) on the spatial distribution of water uptake. We found that as the plot dries, uptake shifts from places with higher to places with lower basal area (Figure 5b). Furthermore, the statistical model revealed that water uptake increased with the higher basal area at locations where multiple species co-existed (Figure 5c). However, the number of species and the basal area were individually not significant fixed effects. Lastly, throughfall patterns were not significant predictors of local root water uptake. Only the median of the spatial deviation of 425 throughfall, which represents temporally stable patterns within the sampling period (δP_{TF}), marginally 426 improved the final model.



427

Figure 5 Visualisation of the significant relations shown in Figure 4, representing the significant drivers of root water uptake patterns during the defined dry periods. Relation to (a) interactive relation of the spatial deviation of soil water storage and field capacity (S_{FC}), (b) the interactive relation of basal area (BA) and the spatial average of soil water storage (\overline{S}), (c) the interactive relation of number of species ($n_{sp,tree}$) and basal area (BA).

432 **4) Discussion**

We investigated the role of throughfall, soil water patterns, and soil and tree characteristics on the spatial variation of root water uptake. In the following sections we discuss three main findings, which are: (1) Contrary to our hypothesis, throughfall patterns do not play a role not in root water uptake patterns despite the recurrence of distinctly localized greater and lesser throughfall inputs. (2) How and where water is stored in the soil, which is strongly determined by soil hydraulic properties, dominates water uptake patterns. (3) The size and species of neighbouring trees regulate relative local water uptake such that locations surrounded by more diverse neighbourhoods are subject to greater water uptake.

440 **4.1)** Spatial variation in throughfall does not affect root water uptake patterns

441 We adequately captured the spatial distribution and temporal stability of throughfall at locations where 442 local root water uptake was derived. Consistent with previous observations in temperate forests (e.g., 443 Whelan and Anderson, 1996; Staelens et al., 2006; Metzger et al., 2017), the amount of weekly rainfall 444 significantly altered the spatial distribution of throughfall such that more rainfall, and thus more 445 throughfall, resulted in less spatial variability. Previous studies repeatedly showed that throughfall 446 patterns exhibit temporal stability in forest ecosystems (e.g., Keim et al., 2005; Staelens et al., 2006; 447 Wullaert et al., 2009; Rodrigues et al., 2022). At our research site, using event-based sampling, Metzger 448 et al., (2017) and Fischer-Bedtke et al., (2023) demonstrated that throughfall patterns persist over time, 449 which was also true for our weekly sampling in 2019. With canopy cover being the key driver of 450 throughfall (Fischer-Bedtke et al., 2023), it is not surprising that weekly cumulative events resulted in a 451 localized high and low throughfall input.

452 Contrary to expectations (Bouten et al., 1992; Guswa and Spence, 2012; Coenders-Gerrits et al., 2013; 453 Fischer-Bedtke et al., 2023), our results showed that throughfall hotspots do not increase or facilitate 454 greater root water uptake. In addition, the linear mixed effects model results confirmed that throughfall 455 patterns do not drive the variation in root water uptake. We attributed the absence of this to two reasons: 456 (1) decoupled soil water and throughfall patterns, (2) non-water limited conditions.

457 Regarding (1), we confirmed that the temporally stable throughfall patterns do not correspond to the post-458 event soil water and root water uptake patterns. We paired the measurements of throughfall and soil water 459 content measurements – and thus the estimates of root water uptake- within a distance of 1 m. The spatial 460 correlation length of soil water content and throughfall is on the order of 6-10 m in natural temperate 461 forests (Keim et al., 2005; Gerrits et al., 2010; Zehe et al., 2010). In the same study site with the spatially 462 extended throughfall sampling, Fischer-Bedtke et al., (2023) found that the throughfall correlation length 463 increased with decreasing event size, varying from 6.2 m to 9.5 m depending on the size of the rain events. 464 Thus, the paired sampling design in our study likely provided co-located throughfall and soil moisture 465 measurements. However, variation in soil water storage was not related to throughfall patterns despite 466 temporally persistent local high and low throughfall inputs.

467 Some studies, mostly conducted in the arid regions and coniferous forests, reported that soil wetting 468 patterns were not or only partly linked to throughfall variation, despite recurrent throughfall patterns (Raat 469 et al., 2002; Shachnovich et al., 2008; Zhu et al., 2021). Forest floor thickness, horizontal water flow, and 470 soil properties were suggested as reasons for the decoupled patterns. Other modelling and field studies 471 conducted in temperate deciduous forests found that throughfall patterns influenced soil moisture 472 response to rain event rather than post-event soil water storage variability (Coenders-Gerrits et al., 2013; 473 Metzger et al., 2017; Fischer et al., 2023). These studies attributed possible reasons to local processes 474 such as preferential flow due to soil water repellency, the soil pore structure, or elevated root water uptake. 475 Our results support that it is not root water uptake but preferential flow paths that are likely to decouple 476 the throughfall and soil water patterns. In fact, Fischer-Bedtke et al., (2023) using independent throughfall 477 and soil water content sampling designs, demonstrated that the signature of throughfall patterns dissipated 478 in the post-event soil water variation. However, they detected the stronger influence of throughfall 479 patterns in the soil moisture response to rainfall in the 2015 and 2016 growing seasons. The temporal 480 variation in soil water content in the 2019 growing season was similar to the seasonal decline in soil water 481 content in 2015 (Metzger et al., 2017). Dry soil conditions can lead to rapid drainage due to reduced water 482 holding capability (Jost et al., 2004; Blume et al., 2009; Wiekenkamp et al., 2016; Demand et al., 2019; 483 Molina et al., 2019) regardless of throughfall amount and its variation. Therefore, our findings support 484 that the localized throughfall input potentially enhances preferential flow because of low soil retention 485 (Fischer-Bedtke et al., 2023) rather than local root water uptake. As a result, the fast flow processes likely 486 dominate how water is stored and transported at our site, erasing the throughfall distribution signature in 487 soil water and root water uptake patterns. Moreover, any short-term response of uptake to throughfall 488 could not be captured as water uptake was calculated only after 56 hours had elapsed since the last rain 489 event, yet we showed that temporally stable hotspots are not associated with elevated water uptake. 490 Hence, our results are consistent with previous propositions stating that the spatial variation of throughfall 491 affects drainage and subsurface flow (Keim et al., 2006; Blume et al., 2009; Guswa and Spence, 2012), 492 while root activities such as water uptake and hydraulic redistribution do not alter canopy-attributed 493 heterogeneity in drainage pathways (Guswa, 2012).

494 The second reason (2) is related to water-limitation conditions. In central Europe, 2019 was the second 495 consecutive extremely dry summer (Boergens et al., 2020), which damaged beech forests (Obladen et al., 496 2021). On average, however, the potential evapotranspiration demand was met at the study site despite 497 the low soil water storage. The ratio of root water uptake to potential evapotranspiration was mostly above 498 65%, which is within the expected range even in the absence of shallow groundwater storage (Nie et al., 499 2021). Hence, local biotic and soil tied abiotic factors determined the spatial variation of root water uptake 500 during growing season rather than throughfall -water input- patterns. However, the discrepancy between 501 daily potential evapotranspiration and root water uptake only increased as the soil in the sampled layers 502 dried out, due to a potential shift in the water uptake depth (see below).

503 4. 2) Relative and average soil wetness shapes root water uptake patterns

We found that spatial variation in soil water storage strongly regulates local water uptake such that wetter locations enhance root water uptake. This finding is consistent with expectations as transpiration rate relies on soil water availability and distribution (Couvreur et al., 2014; Klein et al., 2014; Hildebrandt et al., 2016). Here, we provide further support that root water uptake is likely to reduce the spatial variability in soil water storage as has been previously suggested (Hopmans and Bristow, 2002; Ivanov et al., 2010; Neumann and Cardon, 2012).

510 Trees take up more water in locations where water is not subject to throughfall-driven rapid drainage (see 511 above), as a result root water uptake patterns are determined by where water is retained longer in the soil. 512 Our results support previous studies suggesting that tree transpiration demand is met by water with longer 513 residence time in the soil matrix - passive storage - while groundwater recharge is fed by rapid flow -514 active storage (e.g. Evaristo et al., 2019; Sprenger et al., 2019). In our statistical analyses, we investigated 515 the soil properties of bulk density and field capacity, which are strongly dependent on other soil properties 516 that control aggregation and soil structure. Although bulk density is strongly related to texture, porosity, 517 soil organic carbon content, all of which also affect water retention (Zacharias and Wessolek, 2007; 518 Looy et al., 2017), surprisingly soil bulk density was not retained as a predictive variable in the optimal 519 model. In contrast, the interaction term including field capacity and local soil water storage was significant 520 in the model with a negative relationship with relative water uptake, showing that the combination of higher field capacity (fewer macropores) and low soil water hinders water uptake because water more is more strongly bound in the soil. Differences in local soil properties regulate the matric potential at a certain soil wetness. Thus, wetter locations do not necessarily correspond to those of easier root water uptake due to differences in the soil water retention characteristics (Vereecken et al., 2007; Cai et al., 2018) for which field capacity serves as a proxy. However soil properties alone were less important (smaller effects size of the interaction term including field capacity) than other factors despite their control on the spatial distribution of soil moisture (Vereecken et al., 2022).

In addition, the spatial mean of soil water - a measure of overall wetness of the stand - influenced root water uptake patterns, yet the effect depended on the basal area of neighboring trees. We found that as the study site dries out, local water uptake increased in locations with smaller basal areas. Conversely, wetter site conditions facilitate greater water uptake at locations with higher basal areas, i.e., dense clusters of large trees. We interpret this as a sign that larger trees are likely to shift their water uptake to deeper soil layers to meet transpiration demands, beyond the monitored soil depth (37 cm), as follows:

Higher basal area is likely to increase transpiration demand and enhance water uptake as long as water is available. Moreover, locations with higher basal area exhaust the water storage more rapidly as these locations host larger root structure and root biomass (Le Goff and Ottorini, 2001). At the same time, larger sized trees can shift uptake to deeper layers (Gaines et al., 2016).

538 Beech trees have extensive root systems at shallower depths similar to other temperate tree species, such 539 as European ash and sycamore maple (Kreuzwieser and Gessler, 2010; Brinkmann et al., 2019) Despite 540 their shallower root system (Leuschner, 2020) in response to declining soil water content in the topsoil, 541 temperate tree species can tap water from the deeper soil layers (Brinkmann et al., 2019; Agee et al., 542 2021; Seeger and Weiler, 2021). Recently, Agee et al. (2021) used a three-dimensional water uptake 543 model based on observations in temperate mixed-deciduous forest to show that water uptake is shifted to 544 the deeper soil layers as soil moisture depletes, which is consistent with the field observations. Moreover, 545 Krämer and Hölscher (2010) observed in beech and mixed deciduous stands that roots can extract water 546 at depths down to 70 cm soil depth. Similar to our site, theirs had a shallow soil layer underlain by 547 weathered limestone, but the soil depth varied between 50 and 120 cm. Brinkmann et al., (2019) also 548 observed similar depth range for beech-trees in a mixed forest by tracing stable water isotopes of soil and

549 xylem water.

550 Further tree age and size can affect both individual and stand level transpiration because of the different 551 physiological characteristics and biometrics of trees associated with them (Kostner et al., 2002; Tsuruta 552 et al., 2023). Within the same species, the larger -presumably older- trees have an advantage in accessing 553 the deeper water storages because of their larger root biomass (Le Goff and Ottorini, 2001) and root 554 plasticity may be able to shift the depth of water uptake while younger trees rely on shallower soil water 555 storages (Dawson, 1996). Our results can be interpreted as tree size, which can be attributed to tree age, 556 affecting root water uptake patterns through differential root biomass development. Furthermore, in the 557 Hainich the coexisting species most likely represent highly coherent rooting depth distribution among 558 trees (Gebauer et al., 2012; Meinen et al., 2009) yet adopt different water uptake strategies (see below). 559 Hence consistent with previous studies focusing on temperate tree species, the linear mixed effect model 560 results indicate that trees of different sizes response to declining soil water content by shifting water 561 uptake depth.

562 **4.3)** Tree species richness regulates root water uptake patterns

563 In addition to the basal area, we included the number of species and number of tree individuals in the 564 linear mixed effects analysis to further explore the biotic drivers of root water uptake patterns. While the 565 number of trees was unimportant, the number of species and the basal area showed a significant 566 interaction effect on the local water uptake. The result indicates that an increase in species richness leads 567 to greater root water uptake, depending on the size and/or density of the neighboring trees: Higher basal 568 area, combined with more species, elevates water uptake. In other words, the interactions among 569 neighboring tree species strongly determine root water uptake patterns, and for the same basal area, more 570 water can be taken up in a diverse neighborhood than in a less diverse locations.

In temperate forests, transpiration has been observed to change with tree species richness at the stand
level (Krämer and Hölscher, 2010; Gebauer et al., 2012; Kunert et al., 2012; Meißner et al., 2012;
Forrester, 2014). Although some studies indicate a positive relationship between tree diversity and water
uptake rate (Forrester et al., 2010; Krämer and Hölscher, 2010; Kunert et al., 2012), tree species diversity

575 is not always positively related to water uptake. While Krämer and Hölscher (2010) observed a positive 576 correlation between water uptake and species richness of the plots in the upper soil layers during soil 577 drying in 2006 at the same study site, Meißner et al. (2012) found no relationship between tree diversity 578 and root water uptake in 2009. They attributed this finding to wetter soil conditions. In contrast, Lübbe et 579 al. (2016) observed a weak effect of diversity on transpiration in wetter soil conditions but not in drier 580 conditions compared to previous studies (e.g., Pretzsch et al., 2013; del Río et al., 2014). Shortage of 581 water can inflate competition mechanisms for water among tree species (González de Andrés et al., 2018; Vitali et al., 2018; Magh et al., 2020). Our results indicate that competition between neighboring tree 582 583 species increases water uptake capacity at more diverse spots (Wambsganss et al., 2021).

584 In addition, different co-existing tree species can facilitate resource uptake or reduce competition, 585 depending on the temporal and spatial availability of the sources, which is often defined as 586 complementarity (Forrester and Bauhus, 2016). As reviewed and listed by Silvertown et al. (2015), 587 several studies suggest that co-existing tree species reduce competition for subsurface water sources by 588 adopting different vertical root water uptake strategies, referred to as hydrological niche partitioning. In 589 addition, trees can transport water from wet to dry parts of the soil layers through their roots (Neumann 590 and Cardon, 2012). The mechanism is called hydraulic redistribution or hydraulic lift, which can provide 591 water availability to the shallow roots in drier layers (Burgess et al., 1998; Jonard et al., 2011; Hafner et 592 al., 2017; Lee et al., 2018; Rodríguez-Robles et al., 2020; Hafner et al., 2021). In an experiment with six 593 temperate tree species, including the European beech, Hafner et al. (2021) found that the neighboring tree 594 species diversity may not be important for exploiting water uptake through hydraulic redistribution. Both 595 hydraulic niche partitioning and redistribution have been observed vertically, whereas horizontal patterns 596 are largely unexplored the context of niche partitioning (Hildebrandt, 2020). Our results do not provide 597 direct evidence for either hydraulic redistribution or horizontal niche partitioning. However, they indicate 598 that horizontal root water uptake patterns are regulated by species richness and interactions among 599 neighbouring trees. Thus, we emphasize here the complex interplay between tree species diversity, 600 complementary mechanisms, and water uptake patterns, which is consistent not only with the above-601 mentioned plot-scale studies, but also with larger-scale studies. For instance Knighton et al., (2019) using 602 the Budyko framework across more than one hundred catchments found that transpiration losses in 603 catchments with deep rooted and mixed species forests differed from those in monoculture catchments.
604 In other words, both plot and catchment scale studies support our results showing that interactions among
605 different coexisting species play a significant role in the spatio-temporal variation of root water uptake.

606 **5) Conclusion**

607 We investigated the factors that influence the spatial patterns of root water uptake by considering 608 heterogeneity in throughfall and soil water. To that end, we acquired a comprehensive data set based on 609 throughfall measurements paired with soil moisture sensors in a mixed deciduous forest. Soil and 610 neighboring tree characteristics were also included in the linear mixed effects model. We found that 611 variation in root water uptake did not correspond to throughfall consequently rejecting our hypothesis 612 that variation in throughfall is imprinted in water uptake patterns. Wetter soil locations, also poorly 613 associated with higher throughfall, increased local root water uptake. In contrast, how average soil water 614 conditions modified root water uptake depended on the neighborhood basal area. As the site dried out, 615 large trees likely took up water in deeper layers to meet transpiration demands. Furthermore, an increase 616 in species diversity promoted root water uptake, similarly depending on the size of neighboring trees, 617 suggesting active complementarity mechanisms in the forest stand. In conclusion, our results manifest 618 that soil water distribution and neighboring tree characteristics regulate root water uptake patterns more 619 than soil properties and throughfall variation.

620

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629 sampling.

630

631 Data availability

632 The dataset is currently being prepared for publication in an official repository. The DOI will be published633 with the data at the latest when the data are published.

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635 Author contributions

GD and AH designed the throughfall measurement setup, AH and JCM designed soil moisture measurement. GD conducted the field sampling with assistance from JF and the students listed in the Acknowledgments. GD analyzed the data, developed the linear mixed effects model, and analyzed the results with AH and AG. GD prepared the first version of the manuscript, and all authors contributed to

640 discussions and the final version of the manuscript.

641 **Competing interests**

Anke Hildebrandt is part of the editorial board of HESS. The peer-review process was guided by an independent editor, and the authors have also no other competing interests to declare.

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