REVIEW ARTICLE



Root hydraulic properties: An exploration of their variability across scales

Juan C. Baca Cabrera¹ | Jan Vanderborght¹ | Valentin Couvreur² | Dominik Behrend³ | Thomas Gaiser³ | Thuy Huu Nguyen³ | Guillaume Lobet¹

¹Institute of Bio- and Geoscience, Agrosphere (IBG-3), Forschungszentrum Jülich GmbH, Jülich, Germany

²Earth and Life Institute, Université catholique de Louvain, Louvain-la-Neuve, Belgium

³Institute of Crop Science and Resources Conservation, University of Bonn, Bonn, Germany

Correspondence

Juan C. Baca Cabrera and Guillaume Lobet, Institute of Bio- and Geoscience, Agrosphere (IBG-3), Forschungszentrum Jülich GmbH, Wilhelm-Johnen-Str., 52428 Jülich, Germany. Email: j.baca.cabrera@fz-juelich.de and g. Iobet@fz-juelich.de

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Abstract

Root hydraulic properties are key physiological traits that determine the capacity of root systems to take up water, at a specific evaporative demand. They can strongly vary among species, cultivars or even within the same genotype, but a systematic analysis of their variation across plant functional types (PFTs) is still missing. Here, we reviewed published empirical studies on root hydraulic properties at the segment-, individual root-, or root system scale and determined its variability and the main factors contributing to it. This corresponded to a total of 241 published studies, comprising 213 species, including woody and herbaceous vegetation.

We observed an extremely large range of variation (of orders of magnitude) in root hydraulic properties, but this was not caused by systematic differences among PFTs. Rather, the (combined) effect of factors such as root system age, driving force used for measurement, or stress treatments shaped the results. We found a significant decrease in root hydraulic properties under stress conditions (drought and aquaporin inhibition, p < .001) and a significant effect of the driving force used for measurement (hydrostatic or osmotic gradients, p < .001). Furthermore, whole root system conductance increased significantly with root system age across several crop species (p < .01), causing very large variation in the data (>2 orders of magnitude). Interestingly, this relationship showed an asymptotic shape, with a steep increase during the first days of growth and a flattening out at later stages of development. We confirmed this dynamic through simulations using a state-of-the-art computational model of water flow in the root system for a variety of crop species, suggesting common patterns across studies and species.

These findings provide better understanding of the main causes of root hydraulic properties variations observed across empirical studies. They also open the door to better representation of hydraulic processes across multiple plant functional types and at large scales. All data collected in our analysis has been aggregated into an open access database (https://roothydraulic-properties.shinyapps.io/database/), fostering scientific exchange.

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KEYWORDS

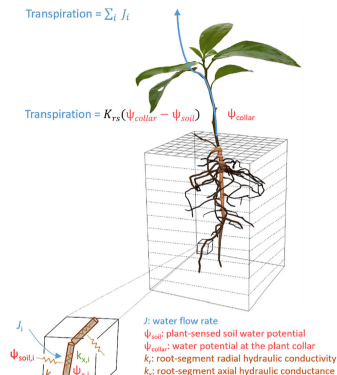
open access database, plant functional types, plant modeling, review, root hydraulic properties variability, whole root system conductance

1 | INTRODUCTION

Root water uptake is a fundamental mechanism essential for the survival of plants. The ability of plants to absorb water through their roots and transport it to the plant's above-ground tissues is crucial for enabling key physiological processes such as photosynthesis, nutrient absorption, and cell expansion (Lambers & Oliveira, 2019). The effectiveness of root systems in absorbing water allows plants to regulate their water balance, postpone or avoid water stress, regulate canopy temperature, and sustain physiological functions at their optimum (Abdalla et al., 2022; Lynch et al., 2014; Steudle, 2000a).

Water uptake is a passive process driven by the water potential gradients in the soil-plant-atmosphere continuum (catenary process, Cowan, 1965), where water is pulled up from the soil into the root xylem and up to the leaf following the cohesion-tension principle (Steudle, 2001). Water flow through the root system can be described analogously to electric current through a network of resistances (Landsberg & Fowkes, 1978). The water flow rate (J, m³ s⁻¹) between any two points is dependent on the water potential difference (ψ , MPa) and the hydraulic conductance (K, m³ MPa⁻¹ s⁻¹, the inverse of a resistance) between these points. In that, root water uptake from the root-soil interface to the above ground organs is affected by root hydraulic properties (the individual resistances) and the root system architecture (the way resistances are connected to form a network) (Doussan et al., 1998; Leitner et al., 2014; Lobet et al., 2014) (Figure 1).

Root hydraulic properties can be expressed at different tissue scales, from root segments up to the whole root system (Figure 1, Table 1). The radial conductivity (k_r) represents the capacity of roots to transport water from the root-soil interface to the root-xylem across their radial pathways, and depends on several anatomical features (North & Peterson, 2005; Steudle, 2000a) and aquaporin expression (Gambetta et al., 2017). The axial conductance k_x refers to the ability of roots to transport water longitudinally, which is a function of the number and diameter of xylem vessels (Hacke & Jansen, 2009). The resulting total conductivity of individual roots or root segments (k_{root}) can be limited by its radial (Bramley et al., 2009) or axial components (Bouda et al., 2018; Boursiac, Pradal, et al., 2022; Sanderson et al., 1988). The whole root system conductance (K_{rs}) integrates the contribution of all individual conductances along the root system, that is, it depends on k_r and k_x (Bouda et al., 2018; Meunier et al., 2019) but also on the root system architecture (Doussan et al., 2006), and reflects the overall hydraulic efficiency of the root system in transporting water from the soil to the above-ground tissues (see Table 1 for details). Understanding the variability in these key hydraulic properties among and within plant species and in response to changing environmental conditions and environmental stresses is essential for



 \hat{K}_{re} : root system hydraulic conductance

FIGURE 1 Root hydraulic properties and water flow in the soilplant-atmosphere continuum. Figure adapted from Vanderborght et al. (2021)

the study of plant water relations (Gallardo et al., 1996; Lambers & Oliveira, 2019).

A large range of empirical methods has been developed for the determination of root hydraulic properties, from the cell and tissue level (Steudle, 1990) up to the whole root system (Tyree et al., 1995), with the pressure chamber, the High Pressure Flow Meter (HPFM) and root exudation being the most common ones (Boursiac, Protto, et al., 2022). While these methods rely on the direct measurement of water flow across root tissues, also more indirect methods based on observations of soil water content and transpiration changes in combination with modeling have been applied (Abdalla & Ahmed, 2021; Abdalla et al., 2022). However, different measurement methods may produce different results, especially when comparing methods that rely on a hydrostatic driving force for water flow against those using an osmotic one (Kim et al., 2018). Additionally, empirical studies have shown that root hydraulic properties can strongly vary (up to orders of magnitude) among species (Bramley et al., 2009; Pratt et al., 2010; Steudle, 2000a), but also among genotypes of one species (Rishmawi et al., 2023) or even among individuals of the same genotype (Steudle, 2000a). This large variability can be explained, at least

TABLE 1 Root hydraulic properties definitions.



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Symbol	Definition	Tissue level	Units	Alternative symbols used in the literature	Specification			
k _r	Radial hydraulic conductivity	Individual roots or root segments	m MPa ⁻¹ s ⁻¹	L _r (Doussan et al., 2006; Huang & Nobel, 1994; North & Peterson, 2005)	Usually not directly measured, but calculated using k_{root} and k_x measurements, based on the model of Landsberg and Fowkes (1978).			
k _x	Specific axial hydraulic conductance	Individual roots or root segments	m ⁴ MPa ⁻¹ s ⁻¹	K_h (Doussan et al., 2006; Huang & Nobel, 1994; North & Peterson, 2005); K_x (Ahmed et al., 2018); L_x (Frensch & Steudle, 1989; Melchior & Steudle, 1993)	The ability of roots to transport water longitudinally			
k _{x_cs}	k _x normalized by cross sectional area	Individual roots or root segments	$m^2 MPa^{-1} s^{-1}$	K _s (Choat et al., 2012; Pratt et al., 2007)	 k_x data for woody species is very commonly reported on a cross sectional area basis (sapwood, stele, total root cross section) 			
k _{root}	(Total) root hydraulic conductivity	Individual roots or root segments	m MPa ⁻¹ s ⁻¹	L _{pr} (Boursiac, Protto, et al., 2022; Kim et al., 2018; Steudle, 2000a); L _p (Gambetta et al., 2017; Huang & Nobel, 1994; Lambers & Oliveira, 2019; North & Peterson, 2005)	The total water transport capacity of an individual root or a root segment. It can be separated into its radial and axial components. Often assumed to be an approximation of k_r in the literature (i.e., water transport only limited by k_r , not by k_x			
K _{rs}	Whole root system conductance	Entire root system	$m^3 MPa^{-1} s^{-1}$	$ \begin{aligned} & K_{\rm root} ({\rm Cai \ et \ al., 2022}); L_{\rm p} \\ & ({\rm Lambers \ \& \ Oliveira, 2019}); \\ & L_{\rm pr} ({\rm Kim \ et \ al., 2018}; \\ & {\rm Steudle, 2000a}), L_0 \\ & ({\rm Boursiac, \ Protto,} \\ & {\rm et \ al., 2022}; {\rm Maurel} \\ & {\rm et \ al., 2010}; {\rm Tyerman} \\ & {\rm et \ al., 2017}) \end{aligned} $	The water transport capacity of the entire root system.			
K _{rs_norm}	K _{rs} normalized by a measure of the root system size	Entire root system	Depends on normalization		Most common normalizations found in the literature include: Root surface area: K_{rs_area} (m MPa ⁻¹ s ⁻¹); Root fresh or dry weight: K_{rs_weight} (m ³ MPa ⁻¹ s ⁻¹ g ⁻¹); Root length: K_{rs_length} (m ³ MPa ⁻¹ s ⁻¹ m ⁻¹); Root volume: K_{rs_vol} (m ³ MPa ⁻¹ s ⁻¹ m ⁻³)			

partially, by the function of roots as hydraulic rheostats, that is, the dynamic changes that root hydraulic properties undergo during development and in response to environmental stimuli (Maurel et al., 2010). Interestingly, though, a systematic study of the range of variability of root hydraulic properties across multiple plant functional types (PFTs), experimental treatments and measurement techniques is still missing. PFTs provide a simplified description of plant diversity, facilitating the representation of ecosystem processes and vegetation dynamics (Wullschleger et al., 2014). Understanding the variability of root hydraulic properties among and within PFTs is therefore key for a better modeling representation of root water uptake processes across scales (Sulis et al., 2019; Nguyen et al., 2022, 2020).

In this context, the present study focused on improving the understanding of the variability of root hydraulic properties observed

across species and PFTs. For this, we systematically reviewed published empirical studies and addressed the following questions: (i) what is the total range of variation in root hydraulic properties observed in the literature?; (ii) are there systematic differences in root hydraulic properties among PFTs and which other factors affect root hydraulic properties variability?; (iii) are the responses of root hydraulic properties to environmental stresses consistent across PFTs?; and (iv) how are root hydraulic properties affected by root development (root age)?

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Given the large amount of data obtained in the review and its complexity (see Section 2.2 for a detailed data description), the results presented in this study have a stronger focus on K_{rs} , a key trait that might determine the water use of plants under changing environmental conditions (Vadez, 2014) and integrates the variability of k_r , k_x and root architecture. But, all original data that was collected in the review has been aggregated to an open access database (Baca Cabrera, 2023), which can be easily accessed through a web application (https://roothydraulic-properties.shinyapps.io/database/), facilitating data access and further use. Furthermore, we complemented our review by using functional-structural modeling, to improve our understanding of the mechanisms behind the emerging patterns in the empirical data.

2 | METHODS

2.1 | Literature review selection criteria

The main goal of this study was to obtain an overview about the range of variation in root hydraulic properties observed experimentally, and the main factors contributing to it. For this, we reviewed scientific articles in which whole root system hydraulic conductance, root hydraulic conductivity, radial conductivity and/or axial conductance were determined experimentally. The Web of Science search engine was used for the review, and following search terms and keywords were included: "root hydraulic conduct*"AND measur* or "root axial hydraulic conduct*" AND measur* or "root radial hydraulic conduct*"AND measur*. The boolean operator AND was used to limit the search to studies in which root hydraulic properties were directly measured and not indirectly modeled from soil water content and/or plant transpiration or theoretically derived. All papers resulting from the search were revised in detail and only those that met the selection criteria were retained in the database.

In a second step, we checked the citations included in the selected papers to look for additional publications that may meet the selection criteria. Additionally, we looked at previous metaanalyses (Bouda et al., 2018; Meunier et al., 2018), reviews (Aroca et al., 2011; Gambetta et al., 2017; Huang & Nobel, 1994; Kim et al., 2018; Maurel et al., 2010; Nobel & Cui, 1992; North & Peterson, 2005; Steudle, 2000a) and the Xylem Functional Traits Database (Choat et al., 2012) to check for missing publications that should be included in our review. In total, we reviewed 241 papers, which comprise the vast majority of experimental studies on root hydraulic properties published between 1973–2023. A complete list of references included in the database is presented in Table S1.

2.2 | Root hydraulic properties database

As part of the review process, we created an open access root hydraulic properties database, which aggregates all extracted data. Root hydraulic properties data were extracted manually and the software WebPlotDigitizer (Rohatgi, 2023) was used for digitalizing figures. The database contains detailed references to the original studies and provides easy, systematized access to the following data: root hydraulic properties (K_{rs} , k_{root} , k_r and/or k_x), plant functional type (PFT, Table 2), growth form (a coarser classification than PFT, that is, tree, shrub, succulent, graminoid and forb), tissue measured (whole root system, individual roots or root segments), root section (whole root or distal, mid-root or basal segments) measurement method, driving force for measurement, and experimental treatment(s) applied. When reported, plant age and morphological data were also included. The root hydraulic properties data corresponds to measurements in living root tissues, unless explicitly stated otherwise in the original publications. The values stored in the database correspond to average values per study, species, factor (with factor being one or many among experimental treatment, tissue, root section, measurement method and driving force) and age. This means, for example, that a study reporting on K_{rs} of maize, based on two different measurement methods, with two treatments at three developmental stages generated a total of $1 \times 2 \times 2 \times 3 = 12$ data points. Therefore, the number of data points aggregated to the database from each study varied greatly. All digitized data is available for download in the database repository.

Based on the digitalized data, we developed a web application (https://roothydraulic-properties.shinyapps.io/database/) that facilitates data selection, manipulation, visualization, and download. The main results presented in this study can be reproduced using the dynamic tools included there, and interested users are also encouraged to use these tools for their own research. The root hydraulic properties database, together with the web application, is conceived as a dynamic tool that will be updated continuously with newly reviewed studies. Readers are encouraged to share in the repository their new work or previously published work that may have been overlooked in our review process, by using the data sharing template available in the web application. The data included in the database is provided with free and unrestricted access for scientific (non-commercial) use (ODC-BY 1.0 license). Data users are requested to acknowledge the original data source and reference this review in resulting publications.

2.3 | Data analysis and statistics

The data stored in the database was used for a comprehensive analysis on root hydraulic properties variability, excluding data that could **TABLE 2** Plant functional type (PFT) classification. Selected PFTs and corresponding number of species, genera and studies for which root hydraulic properties were investigated. PFTs were defined based on commonly used classifications in land surface models (Poulter et al., 2015), and additional features such as growth form, differentiation between woody and herbaceous vegetation and agronomical importance.

PFT	Description	Species examples	Nr. Species	Nr. Genera	Nr. Studies
Crop herbaceous	Herbaceous crop species (legumes and non- legumes), excluding all C ₃ and C ₄ grasses	Tomato, soybean, lupin	23	17	50
Crop woody	Woody crop species	Cotton, grapevine	2	2	11
C ₃ grass	Grass species with a C ₃ photosynthetic pathway. Most species investigated corresponded to grasses used as crops	Barley, rice, wheat	9	7	50
C ₄ grass	Grass species with a C ₄ photosynthetic pathway. All species investigated corresponded to grasses used as crops	Maize, sorghum, pearl millet	4	4	40
Broadleaf tree	Decidious and evergreen broadleaf tree species, including fruit trees	Quercus spp., Populus spp., apple	64	30	54
Needle tree	Decidious and evergreen needle tree species	Pinus spp., Picea spp., Abies spp.	39	12	28
Tropical tree	Broadleaf tree species from tropical ecosystems	Piper spp., Shorea spp.	37	31	9
Shrub	Decidious and evergreen shrub species	Juniperus spp., Rhamnus spp.	29	17	10
Succulent	Succulent species from arid ecosystems	Agave spp., Opuntia spp.	6	3	10
Other	All species that could not be assigned to any of the defined PFTs	Arabidopsis thaliana., dendrobium, Iris germanica	3	3	6

not be classified into any PFT (defined as "Other", see Table 2). The data was highly imbalanced, and there were large differences in the number of studies and species investigated for the different PFTs and root hydraulic properties. Accordingly, appropriate data analysis methods had to be selected. Although applying a strict meta-analysis (Hedges et al., 1999) could have been reasonable for this purpose, we discarded this approach because of two reasons: too few articles reported all the information needed for performing a meta-analysis (i.e., sample size and standard deviations for each experimental factor); and the experimental factors varied extremely among studies (Table S1), which hampered an evaluation of their individual effects and interactions. Instead, we followed an ad-hoc step-wise approach, and performed a series of independent analyses that quantified the variability in root hydraulic properties observed across studies and evaluated some of the (most important) factors causing it (see Table 3 for factor description). This analysis was performed for all individual root hydraulic properties except for k_r , for which a very limited number of species and studies (n = 12, in both cases) was available. The original data showed a very large positive skewness in all root hydraulic properties ($K_{rs} = 3.6$; $K_{rs_area} = 9.9$; $k_{root} = 7.4$; $k_x = 10.2$; $k_{\rm x \ cs} = 5.8$). Therefore, reported values were log transformed before data analysis, and then back transformed. Thus, the presented results correspond to geometric averages. Approximate standard deviations and standard errors were calculated using the Delta Method (Cramér, 1999).

In a first step, we calculated the range of variation (i.e., minimum, mean and maximum values) for each of the PFTs described in Table 2. For this, we first calculated the geometric means for the different studies and of each species investigated. These values were considered independent and suited for the analysis and were used for the calculation of the range of variation. The results corresponded to geometric means and range of variation for each PFT and root hydraulic property investigated (3.1).

Secondly, Random Forest (RF) models were run and the drop in accuracy of the model -a permutation feature importance metric (Altmann et al., 2010)- was calculated to rank the importance of several factors on the variability of root hydraulic properties. Next, linear mixed models were fitted to test for significant differences in root hydraulic properties among PFTs. PFT and two other highest ranked factors according to the RF model (excluding taxonomical features) were defined as the fixed effects, and study and experimental treatment were defined as the random effects. Given the extremely large dissimilarity in experimental designs among publications (see Table S1 for treatment list), we simplified the factor experimental treatment to four levels: control (defined as such in the publications), stress (any treatment that causes stress, e.g., drought, salt stress, nutrient limitation), other (any treatment that cannot be strictly defined as control or stress, e.g., different soil types, genotypes, season) and no treatment (studies where no treatments were applied). Type III ANOVA with the Satterthwaite's method (Luke, 2017) was used for evaluating factor significance. The R-packages randomForest (Liaw & Wiener, 2002) and Ime4 (Bates et al., 2015) were used for fitting the models.

Finally, we evaluated in more detail three factors that have been repeatedly reported to affect root hydraulic properties: driving force used for measurement, drought stress, and aquaporin (AQP) inhibition (see e.g., Aroca et al., 2011; Gambetta et al., 2017; Kim et al., 2018). For this, the natural log response ratio ($\ln(r) = \ln$ (treatment) - ln (control)) (Hedges et al., 1999) was calculated for each individual study and species in which root hydraulic properties were measured under

TABLE 3 Factors affecting root hydraulic properties variability. Factors analyzed and their ranges (or factor levels) observed in the database.

Factor	Description	Factor levels or range		
PFT	Plant functional types, according to the classification in Table 2	Nine different PFTs		
Age	Root system age.Data principally corresponds to dicot crops and grasses. Root system age of trees and shrubs scarcely reported, mainly restricted to studies with seedlings	3–150 days (herbaceous crops and grasses); 12–485 days (woody crops)		
Driving force	Driving force used for measurement of root hydraulic properties	Hydrostatic or osmotic driving force		
Genus	Taxonomic genus	124 distinct genera		
Growth form	A coarser classification than PFT	Tree, shrub, succulent, graminoid or dicot crops		
Root section	Section of the root (segment) for which root hydraulic properties were determined. Several investigations measured whole roots instead of specific segments	Whole root or distal, mid-root or basal segments		
Root type	Type of root investigated	Primary, tap, seminal, lateral, adventitious, whole root system		
Species	Species investigated	214 distinct species		
Treatment	Simplified classification of the experimental treatments applied in the studies	Control, stress, other or no treatment		

both treatment and control conditions. The results were reported as the mean percentage change ([r - 1] * 100) (Ainsworth & Long, 2005) and response significance was tested with one-sample t-tests (on the log transformed data). Differences in the responses among PFTs were evaluated with one-way ANOVA tests. All data and statistical analyses were conducted in R v.4.3.1 (R Core Team, 2023).

2.4 | Modeling the relationship between K_{rs} and root system age

The results of the RF and linear mixed models (see Section 3.2) indicated a significant and (probably) non-linear relationship between root system age and K_{rs} (and K_{rs_area}). To investigate this relationship in more detail, we modeled the response of K_{rs} to the increase in root system age (and size) over time, using the functional-structural plant models CPlantBox (Schnepf et al., 2018) and MARSHAL (Meunier et al., 2019). Because data on root age was extremely scarce for trees and shrubs (see Table 3), this analysis was restricted to crop species (herbaceous crops and grasses).

CPlantBox was used to simulate the root system development of four different crops over a 120-day period: a C₃ grass (wheat), a C₄ grass (maize), a forb (cauliflower) and a legume (soybean). The species were selected based on plant-functional diversity and data availability. The XML-input parameters were obtained from the literature (de Moraes et al., 2020; Leitner et al., 2010; Morandage et al., 2021; Vansteenkiste et al., 2014). CPlantBox outputs (i.e., the root architecture at each time step) were coupled to MARSHAL to simulate water flow from the soil-root interfaces to xylem vessels at the plant collar, using the analytical solution of water flow within infinitesimal subsegments (Meunier, Draye, et al., 2017), and to calculate the macroscopic parameter K_{rs} (Couvreur et al., 2012). Segment-scale k_r and k_x values were extracted from the database and from modeling (Doussan et al., 1998) and used to parametrize MARSHAL. k_r and k_x are agedependent and vary among root types (Figure S1). To account for the uncertainty in their parameterization, a sensitivity analysis was performed by varying k_r , k_x or the k_r/k_x within the range of variation and the spatial heterogeneity observed in the literature (Figure S1). Modeled K_{rs} corresponds to the mean \pm standard error of all simulations, for each individual crop. Modeling results were contrasted with data gathered from the review, specifically for crop species (dicot crops and C₃ and C₄ grasses) measured using a hydrostatic driving force.

3 | RESULTS AND DISCUSSION

3.1 | Range of variability of root hydraulic properties

In this work, we reviewed a total of 241 root hydraulic properties publications, comprising 215 species from 124 genera (original data available in Baca Cabrera, 2023; complete list of references and species in Table S1). From this total, 165 studies focused on K_{rs} , 60 on k_{root} (including k_r) and 46 on k_x (some studies measured multiple hydraulic properties, simultaneously). We observed a large positive skewness in the reported data ($K_{rs} = 3.6$; $K_{rs_area} = 9.9$; $k_{root} = 7.4$; $k_x = 10.2$; $k_{x_cs} = 5.8$) and an extremely large range of variation (of orders of magnitude) in all root hydraulic properties, whereby this was especially pronounced for K_{rs} (Figure 2).

Reported K_{rs} values varied extremely across studies, species, and plant functional types (PFTs, see Table 2 for classification), ranging between 3.1×10^{-12} (measured in barley) to 9.4×10^{-8} m³ MPa⁻¹ s⁻¹ (measured in common bean). A very large range of variation was also observed within PFTs, with K_{rs} showing a range of variation of $\approx 2-3$ orders of magnitude in all PFTs, except for shrubs (for which only two studies were available). Large ranges of

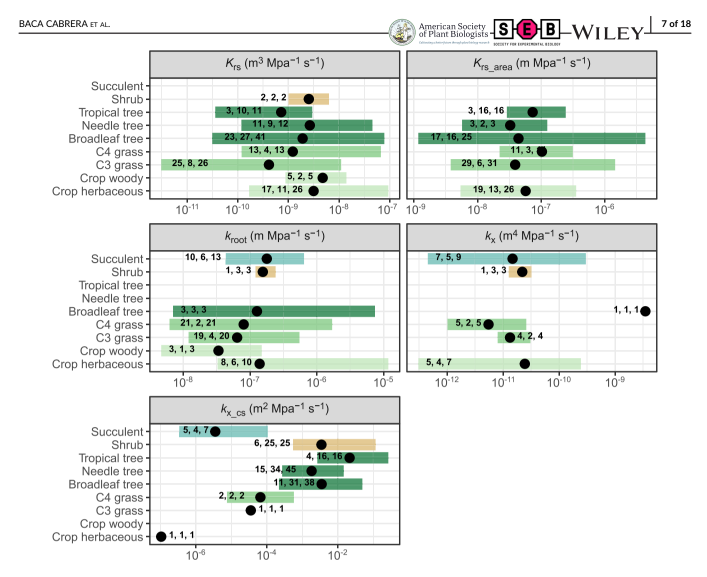


FIGURE 2 Range of variation in root hydraulic properties. Geometric means (filled circles) and range of variation (bars) of root hydraulic properties (see Table 1 for detailed definitions) for different plant functional types. The total number of studies, species, and individual data points for each PFT are indicated in bold (see Section 2.3 for details on the calculation).

variation (>1 order of magnitude) were also present in several species such as maize, barley, tomato or oak trees, which agrees with a previous meta-analysis that reported a total range of K_{rs} variation in maize of \approx 2 orders of magnitude (Meunier et al., 2019).

Clearly, K_{rs} variability within PFTs (and even within some species) was considerably larger than the differences in the geometric means among PFTs, which varied between 4.1×10^{-10} (C₃ grasses) and 4.8×10^{-9} m³ MPa⁻¹ s⁻¹ (woody crops). Because of this very large intra-PFT variability, possible systematic differences among PFTs could have been obscured (but see Section 3.2.1).

 $K_{\rm rs}$ is often reported in the literature on the basis of a measure of root size, to facilitate the comparison among plants of different age, with root surface area ($K_{\rm rs_area}$) being the normalization most widely used (see Table 1 for other common normalizations). Our results indicated that the range of variation of $K_{\rm rs_area}$ was indeed factors of magnitude smaller than that of $K_{\rm rs}$, but it was still extremely large (1.2×10^{-9} – 4.3×10^{-6} m Mpa⁻¹ s⁻¹) (Figure 2). A very large range of variation was also observed within each PFT (≈ 1 –3 orders of magnitude), indicating large intrinsic differences among species and/or experimental design of the studies. Surprisingly, even, both the lowest and the highest K_{rs_area} values found in the literature corresponded to broadleaf tree species (*Quercus petraea* and *P. tremula* × *tremuloides*). On the contrary, the geometric mean of K_{rs_area} varied comparatively slightly among PFTs (3.3×10^{-8} – 1.0×10^{-7} m Mpa⁻¹ s⁻¹).

Published root hydraulic properties data of individual roots and/or root segments (total, radial, and axial) also showed very large variability. The total conductance k_{root} (which is often reported as a proxy of k_r in the literature) varied extremely across studies (range = 4.7×10^{-9} – 1.2×10^{-5} m Mpa⁻¹ s⁻¹, Figure 2), but also within individual PFTs (ranges ≈ 1 –3 orders of magnitude). This large variation was observed despite the few species that have been investigated (2–6 species for the different PFTs). Additionally, the geometric means of k_{root} showed small variation among PFTs (3.4×10^{-8} – 1.8×10^{-7} m Mpa⁻¹ s⁻¹), and this range was almost identical to that of $K_{rs area}$.

Axial conductance also showed a very large variability, both for published data reported as k_x (range = 3.1×10^{-13} – 3.5×10^{-9} m⁴ Mpa⁻¹ s⁻¹) and on a cross sectional area basis ($k_{x,cs}$, range = 1.1×10^{-7} - 2.7×10^{-1} m² Mpa⁻¹ s⁻¹). However, we found very few studies on k_x (20 publications), and they were unevenly distributed across PFTs. While succulent species were the most frequently reported (seven studies, five species), only one tree species was available and showed by far the largest k_x (1–3 order of magnitudes larger than any other value). Excluding that species, k_x ranged between 3.1×10^{-13} - 3.0×10^{-10} m⁴ Mpa⁻¹ s⁻¹, with C₄ grasses showing the lowest (5.5 \times 10⁻¹² m⁴ Mpa⁻¹ s⁻¹) and dicot crops the highest (2.4 \times 10⁻¹¹ m⁴ Mpa⁻¹ s⁻¹) geometric means among PFTs. At the same time, $k_{x cs}$ has been widely reported for woody vegetation (26 publications, 105 species) and showed a range of variation between 2.2×10^{-4} - 2.7×10^{-1} m² Mpa⁻¹ s⁻¹, with tropical trees showing the highest values. These values were systematically higher than $k_{x,cs}$ of the very few non-woody species for which data was available (8 species, range = 1.1×10^{-7} - 1.1×10^{-4} m² Mpa⁻¹ s⁻¹) and confirm the results from previous meta-analyses (Bouda et al., 2018). However, our review also highlights the difficulty of comparing axial conductance of woody and non-woody vegetation, with the former almost entirely being reported as $k_{x cs}$ and the latter as k_x.

3.2 | Understanding root hydraulic properties variability

The results in Section 3.1 showed an extremely large range of variation in root hydraulic properties across published studies. Here, we further investigated to which degree the observed variability could be explained by the response of root hydraulic properties to the following factors: systematic differences among PFTs, driving force used for measurement (hydrostatic or osmotic), effect of environmental stresses, and root system age.

3.2.1 | Main factors affecting root hydraulic properties and differences among PFTs

One central question we addressed in this study was whether the observed variability in root hydraulic properties could be attributed to systematic differences among PFTs. For this, we first used Random Forest (RF) regressions to compare the importance of PFT with other variables that have been reported to affect root hydraulic properties. This included factors such as root system age, the driving force used for measurement (hydrostatic or osmotic), root section and root type, experimental treatment, or variation within species. According to the "drop in accuracy" metric (a permutation feature importance metric, more details in Section 2.3), root system age had the highest importance to explain the variability in K_{rs} (Table 4), which agrees with the general positive relationship between K_{rs} and root system size observed in the literature (Tyree, 2003). This is the case, as with

increasing age the root system grows, adding conductances (new root segments) in parallel in a hydraulic network, which increases the total conductance of that network. Interestingly, root system age also showed the highest importance for K_{rs_area} , suggesting complex interactions between root system growth and K_{rs} development (see Section 3.2.4 for further discussion). The importance of PFT for K_{rs} was 27.4% smaller (and 26.9% smaller for K_{rs_area}) than that of root system age and was similar to the importance of driving force or species and only clearly larger than that of experimental treatment (Table 4). These results indicate that the large variability of K_{rs} observed in the literature cannot be explained by systematic differences among PFTs, alone, but rather by the added effect of multiple factors.

We also analyzed the importance of PFT for k_{root} (Table 4) and observed that it was lower than the importance of driving force (-4.2%) and slightly higher to that of species, root type (seminal, adventitious, lateral) or root section (distal, mid-root, basal or entire root). This suggests that the observed variability of k_{root} is caused by the added effect of multiple factors and their interactions, rather than by systematic differences among PFTs. However, care must be taken in the interpretation of these results, because of the rather small number of species investigated (26) and the extremely low number of studies (5) in which species belonging to different PFTs were investigated simultaneously. On the contrary, the importance of PFT for $k_{\rm x \ cs}$ variability was much larger (at least more than twice) than that of any other factor, except for growth form, confirming the clear, systematic difference between woody and non-woody species depicted in Figure 2 and the observations of Bouda et al. (2018). These results are probably associated with large increases in axial conductance (2-3 orders of magnitude) following secondary growth in woody roots (Vercambre et al., 2002) and with large differences in xylem cross sections between woody and non-woody vegetation.

To confirm the results of the RF models and further investigate systematic differences in root hydraulic properties among PFTs, individual linear mixed models for K_{rs} , K_{rs_area} , k_{root} and k_{x_cs} were run, with PFT and additional non-taxonomical features (i.e., root system age, driving force, root section or root type, detailed factor and model description in Sections 2.2–2.3) as fixed effects, and study and treatment as random effects.

We found no significant effect of PFT on K_{rs} (p = .20), K_{rs_area} (p = .84) and k_{root} (p = .92), but k_{x_cs} varied highly significantly (p < .001) among PFTs (Table 4), which agrees with the results of the RF analysis and its conclusions. On the contrary, a highly significant effect of driving force (p < .001) on K_{rs} , K_{rs_area} and k_{root} was found, indicating systematic difference in root hydraulic properties measured using a hydrostatic driving force, against those using an osmotic driving force (see Section 3.2.2 for a detailed analysis). Additionally, root system age showed a highly significant positive effect on K_{rs} (p < .01), probably associated with an increase of K_{rs} with increasing root system size. Conversely, root system age had no effect on K_{rs_area} (p = .38), contradicting the high importance that root age had for K_{rs_area} prediction, according to the RF model. Interestingly, though, the linear mixed model showed a negative (albeit non-significant)

TABLE 4 Statistics of Random Forest and linear mixed models. Importance of several factors (as described in Table 3) for root hydraulic properties variability, according to the drop in accuracy metric (random Forest); *p*-value of the same factors, using type III ANOVA tests (linear-mixed models); and total variance explained by the fitted random Forest models. Data in bold indicate the 3 highest ranked factors (random Forest models) and effect significance (p < .05, ANOVA tests).

	Drop in mean square error			p-value (Satterthwaite)				
Factor	K _{rs}	K _{rs_area}	k _{root}	k _{x_cs}	K _{rs}	K _{rs_area}	k _{root}	k _{x_cs}
PFT	2.07	.98	.91	7.39	.20	.84	.92	<.001
Age	2.85	1.34	-	-	<.001	.38		
Driving force	1.63	1.1	.95	-	<.001	<.001	<.001	
Genus	1.84	1.05	.89	1.81	-	-	-	-
Growth form	1.14	.41	.68	6.72	-	-	-	-
Root section	-	-	.64	2.47			<.01	<.001
Root type	-	-	.75	1.79	-	-	-	-
Species	1.99	1.12	.76	1.8	-	-	-	-
Treatment	.67	.33	.31	.41	-	-	-	-
	-	-	-	-	-	-	-	-
Total variance explained (%)	76.9	65.9	64.3	83.6	-	-	-	-

relationship between K_{rs_area} and root age and this negative relationship became significant (p < .05) when a negative exponential function was fitted to the data, instead of a linear relationship. This implies a decrease in K_{rs} per unit root surface over time, a phenomenon that could be associated with the decrease in segment-scale radial conductivity with age, but also with axial transport limitation with increasing root length (Meunier, Draye, et al., 2017; Bouda et al., 2018, see also discussion in Section 3.2.4). Clearly, the relationship between root age and K_{rs} (and K_{rs_area}) observed in our review is complex and was therefore explored in more detail in Section 3.2.4.

The linear mixed models also showed a highly significant (p < .001) effect of root section – a factor describing whether root hydraulic properties were measured on basal, mid-root or distal root segments or on entire roots – on k_{root} and $k_{x cs}$, suggesting the presence of spatial gradients in roots across species and PFTs. Spatial variation alongside roots in k_r and k_x (and consequently in k_{root}) has been reported for the grass species maize (Doussan et al., 1998; Frensch & Steudle, 1989; Meunier et al., 2018) and barley (Knipfer & Fricke, 2011) and for A. deserti (Huang & Nobel, 1992), with radial conductivity decreasing from root tip to root base, while the opposite was the case for axial conductance (see also Figure S1). Variation can be caused by changes in root anatomy and function (e.g., formation of apoplastic barriers, increase in xylem diameter and density, differences in aquaporin expression) with increasing age. However, similar gradients were not evident (particularly in the case of k_r and k_{root}) in onion (Melchior & Steudle, 1993) or lupin (Doussan et al., 2006; Meunier et al., 2018), questioning the idea that they are ubiquitous across species and PFTs. Our review cannot answer this, because most of the studies reported data for one root section only, hampering systematic comparison among sections. For instance, the two largest $k_{\rm root}$ values in our review (1.2 \times 10⁻⁵ in Vicia faba and 7.4 \times 10⁻⁶ in Populus trichocarpa x deltoides) corresponded to measurements in distal segments, but unfortunately no other root section was

investigated in those studies. Nevertheless, the statistical results underscore the significance of spatial gradients as a factor of variability in root hydraulic properties and stress the need for further investigations on this topic, focusing on the differences (or lack thereof) among species from different PFTs.

In general, the statistical analyses did not reveal systematic differences in root hydraulic properties among PFTs, apart from the highly significant effect of PFT on axial conductance, a feature that has been reported previously. Rather, the results imply that the variation in multiple factors such as age, driving force, or root section analyzed (and probably their interactions) determined the extremely large variability observed here. This would also explain why root hydraulic properties varied so much within PFTs (Figure 2) or even within species. Accordingly, a detailed analysis on the influence of several factors on root hydraulic properties variability (with the main focus on K_{rs}) was also performed in this review, and the results are presented in the following Sections 3.2.2–3.2.4.

To our knowledge, this is the first systematic review on the topic of root hydraulic properties and their variability across PFTs, leaving little room for the comparison of our results with previous investigations. However, we cannot discard the possibility that systematic differences among PFTs - which we did not find - were obscured by the dissimilarity in experimental design among the publications. Actually, less than 10% of the reviewed studies included species corresponding to more than one PFT, and the hydraulic properties investigated there were unevenly distributed: while K_{rs} and k_{root} studies mostly focused on dicot and monocot crop species (e.g., Gallardo et al., 1996; Bramley et al., 2007; Hess et al., 2015), broadleaf and needle trees were predominant in k_x (or rather $k_{x cs}$) studies (e.g., Domec et al., 2010; Maherali et al., 2006). In fact, we only found one study in which root hydraulic properties of trees and herbaceous vegetation were measured simultaneously (Rieger & Litvin, 1999). Thus, more studies comparing root hydraulic properties across

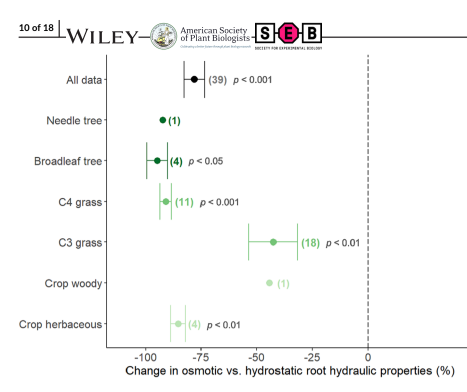


FIGURE 3 Difference between osmotic vs. hydrostatic root hydraulic properties. Data points and error bars represent the mean ± the standard error for each PFT (sample size *n* reported on the side). The mean value for all samples is represented with a black circle. Individual values were calculated based on the log response ratio. The *p*-values correspond to one-sample t-tests for all data and the individual PFTs.

species and PFTs are needed to confirm (or reject) the results in this review.

3.2.2 | The driving force matters

According to the results from the previous section, the driving force used for measurement was a key factor for explaining the very large variability observed in this review. Here, we quantified in more detail the differences in root hydraulic properties (specifically K_{rs} and k_{root} ; k_x data is not relevant for this analysis) estimated under osmotic gradients (hereafter osmotic root hydraulic properties), compared with those estimated under hydrostatic gradients (hereafter hydrostatic root hydraulic properties), based on the log response ratio of pairwise comparisons (methodological details in Section 2.3).

A total of 39 data pairs, corresponding to 29 studies and 16 species were investigated, whereby only four species (maize, barley, rice, and wheat) accounted for >60% of all values (see Table S2 for all studies and species included). On average, osmotic root hydraulic properties were 78.1% smaller than hydrostatic ones, and this effect was highly significant (p < .001). More interestingly, the observed response varied significantly among PFTs (p < .001), showing average decreases ranging from 42.6% (C₃ grasses) to 94.9% (broadleaf trees). In that, C₃ grasses showed a much lower decrease compared with the remaining PFTs, which varied very slightly among each other (range = 94.9–85.4%; woody crops were not included in this comparison, because only one value was available). For all PFTs, the reported decrease in osmotic root hydraulic properties was significantly different from zero (p < .05, Figure 3).

Clearly, the driving force affects the measurements of root hydraulic properties. Across all studies, the largest difference was observed in K_{rs} of oak trees and reached almost two orders of

magnitude (Steudle & Meshcheryakov, 1996). On average, a decrease of \approx 78% of osmotic compared with hydrostatic root hydraulic properties was observed, and in four PFTs (broadleaf and needle trees, C₄ grasses and dicot crops) a decrease of \approx 90% (i.e., 1 order of magnitude) was reached. Considering that the total range of variation within PFTs was \approx 1–3 orders of magnitude (Figure 2), the driving force can be described as one of the most important factors for explaining the variability in root hydraulic properties reported in this review.

That osmotic root hydraulic properties are systemically lower than hydrostatic ones has been reported before (Kim et al., 2018; Steudle, 2000a). In line with the principles of the composite transport model (Steudle, 2000a), the comparison between osmotic and hydrostatic root hydraulic properties has been widely used to differentiate the cell-to-cell path (obtained from osmotic measurements) from the overall path for water flow (i.e., cell-to-cell + apoplastic paths, obtained from hydrostatic measurements) and how the contribution of the former might change under conditions of environmental stress (see e.g., Barrios-Masias et al., 2015; Garthwaite et al., 2006; Kreszies et al., 2020). According to this approach, our results would imply that the cell-to-cell path had a (much) smaller contribution than the aploplastic path to the total water flow across PFTs, with the cell-to-cell contribution to total water flow being the lowest in broadleaf trees (4.9%) and the highest in C₃ grasses (36.5%). However, the accuracy of this approach has been questioned (Chaumont & Tyerman, 2014), as multiscale studies do not support this common assumption and rather indicate that the differences between osmotic and hydrostatic root hydraulic properties may stem from an erroneous estimation of the osmotic driving pressure and therefore of hydraulic properties (Bramley et al., 2007; Couvreur et al., 2018). Cell-scale simulations of the advection-diffusion of osmolytes suggest that their accumulation at apoplastic barriers (e.g., Casprian strip) may alone generate a fivefold overestimation of the effective water potential gradient across the endodermis (Couvreur et al., 2018; Steudle, 2000a; Knipfer & Fricke, 2011), while apoplastic, symplastic and transmembrane modes of water transport would vary radially regardless of whether the water potential difference between root surface and xylem is because of pressure or osmolytes. Nevertheless, the data clearly showed a differentiation between C₃ grasses and the remaining PFTs, and also very large discrepancies within the C₃ grasses: while osmotic and hydrostatic root hydraulic properties were almost equal in barley (\approx 6% higher osmotic root hydraulic properties, in average), osmotic root hydraulic properties were much smaller than hydrostatic ones in wheat and rice (\approx 55% and \approx 63% in average, respectively). To which degree these differences indicate functional heterogeneity in water transport patterns among species lies beyond the scope of this review, but the data presented here could be used to identify species or PFTs of interest for future studies.

3.2.3 | Responses to drought and AQP inhibition

Environmental stress has been widely reported as a factor affecting root hydraulic properties (Aroca et al., 2011; Gambetta et al., 2017; Maurel et al., 2010; Steudle, 2000b). Interestingly, though, our analysis showed that experimental treatment had the lowest importance of all variables in explaining the range of variation in K_{rs} , K_{rs_area} , k_{root} and $k_{\rm x \ cs}$ observed in the literature (Table 4). Two aspects could explain these results: (1) the variation across studies and PFTs was so large, that it obscured the effects of experimental treatments observed in individual studies; and (2) experimental treatments differed extremely among studies (Table S1), hindering a systematic analysis of the effect of environmental stress on root hydraulic properties variability. Thus, for the purpose of this review, the response of root hydraulic properties to stress was narrowed to two factors: drought stress and aquaporin (AQP) inhibition. For this, 28 studies on the effect of drought stress and 19 studies on the effect of AQP inhibition on K_{rs} (or its normalized values) were analyzed.

There was a significant decrease in K_{rs} under both drought stress and AQP inhibition (p < .001 in both cases). On average, K_{rs} decreased 61% under drought conditions and the decrease under AQP inhibition was very similar (59%). However, the K_{rs} response to drought showed more variation across PFTs, studies or species than that to AQP inhibition. The average K_{rs} decrease under drought varied among PFTs in a range between 80.8% (in dicot crops) and 38.3% (in C₃ grasses) and was only significant for broadleaf trees (p < .01), C₃ grasses (p < .05) and herbaceous crops (p < .01) (Figure 4). Accordingly, there was a marginally significant variation among PFTs (p = .07) in their response to drought stress. Meanwhile, K_{rs} decreased under AQP inhibition in a smaller range between 50.9% (in tropical trees) to 77.4% in (C₄ grasses), with the decrease being significant for herbaceous crops (p < .01), C₃ grasses (p < .001) and tropical trees (p < .001). But, the difference among PFTs in their individual responses to AQP inhibition was not significant (p = .16). Also, across all studies and species (n = 30), the K_{rs} response to drought varied greatly, between $\approx 98\%$ decrease (i.e., a decline of almost two orders of magnitude) and $\approx 35\%$

increase. On the contrary, K_{rs} responded negatively to AQP inhibition, without exception (n = 25), with the decrease ranging between $\approx 22\%$ -86%.

The average decline in K_{rs} under drought agrees with the conclusions of previous reviews (Aroca et al., 2011). This response corresponds to a water saving strategy under condition of limited water availability, which can be induced by short-term responses (e.g., changes in the aquaporin gating), but also on long-term droughtdriven anatomical changes (e.g., formation of apoplastic barriers, aerenchyma, changes in xylem vessel size) or changes in root size (Aroca et al., 2011; Bauget et al., 2023; Vadez, 2014). Furthermore, our review revealed differences among PFTs (albeit non-significant, probably because of a small sample size), with grasses (both C_3 and C_4) showing a weaker response to drought than trees or dicot crops. In fact, the only three studies in which an increase in K_{rs} under drought was reported, were conducted with rice (Ding et al., 2015; Lian et al., 2004) and maize (Zhang et al., 1995). Also, the K_{rs} decrease of maize (C₄ grass, \approx 44%) under drought was considerably weaker than that of tomato (dicot crop, \approx 63%), in the only study where grass and non-grass species were directly compared (Bárzana et al., 2012), supporting the overall trends reported here. However, the shown differences among PFT might be conditioned by the low number of species investigated within each PFT. For example, in the case of C_3 grasses seven out of nine studies were conducted with rice, and a similar behavior was observed for C_4 grasses (all four studies with maize) or dicot crops (four out of seven studies with tomato). But, regardless of these limitations, our results contribute to a better understanding of the expected root hydraulic properties variability under drought conditions across species and PFTs.

On the other hand, a negative response of K_{rs} to AQP inhibition was observed across all PFTs and species investigated. This effect is driven by a decrease in the cell-to-cell radial water flow (Aroca et al., 2011; Chaumont & Tyerman, 2014), such that the large range in K_{rs} responses to AQP inhibition (\approx 22%–86% decrease across studies) could be associated with differences in aquaporin activity of root cells among the investigated species and PFTs. However, we did not observe systematic differences among PFTs in our analysis. In a previous review on aquaporins and root water uptake, Gambetta et al. (2017) also identified a very large range in the response of root hydraulic properties to AQP inhibition, and mainly attributed this to variability in the experimental approach across studies. As such, further examinations of the responses exhibited by distinct tissues, species, and/or plant functional types (PFTs) are essential to enhance our understanding of water flow dynamics under stress conditions, and how this might impact the overall variability of root hydraulic properties.

3.2.4 | Non-linear K_{rs} increase with increasing root system age in crops and grasses

Root system age is a key factor for explaining the large variability in K_{rs} observed in this review (see Section 3.2.1). Here, we investigated

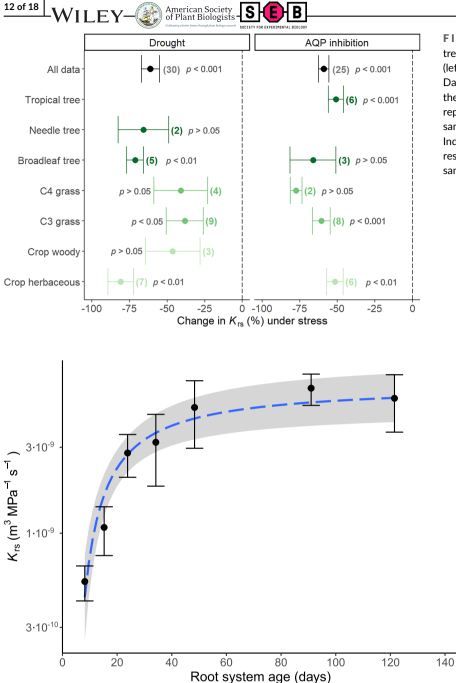


FIGURE 4 Response of K_{rs} to stress treatments. Changes in K_{rs} under drought stress (left panel) and aquaporin inhibition (right panel). Data points and error bars represent the mean ± the standard error for each PFT (sample size *n* reported on the side). The mean value for all samples is represented with a black circle. Individual values were calculated based on the log response ratio. The *p*-values correspond to onesample t-tests for all data and the individual PFTs.

FIGURE 5 Relationship between root system age and K_{rs} in crop species. Data points and error bars represent K_{rs} (mean \pm standard error) of crop species (C₃ and C₄ grasses and dicot crops) grouped according to age (0–10, 10–20, 20–30, 30–40, 40–60, 60–100, >100 days). The dashed blue line and the shaded area represent a fitted exponential model (\pm standard error).

this relationship in more detail, specifically for the hydrostatic K_{rs} of dicot crops and grass species (woody vegetation was excluded from the analysis, as precise data on root system age was very scarce for that type of vegetation; more details on selection criteria in Section 2.4). Across studies and species, there was a significant increase in K_{rs} with increasing age of the root system (p < .01, $R^2 = .32$, n = 25), with the relationship exhibiting a non-linear pattern (Figure 5). K_{rs} increased abruptly during the first 20–30 days of root development, and then slowly flattened out, with a total range of variation between $\approx 6 \times 10^{-11}$ – 2×10^{-8} m³ MPa⁻¹ s⁻¹. The steep increase in K_{rs} during the first days of development is probably caused by the growth of the root system adding new conductances (new roots) to the root hydraulic network, thus increasing the total

conductance of the network. However, the asymptotic behavior after days 30–40 suggests a partial decoupling between root size and K_{rs} at later stages of development. Unfortunately, root size data (e.g., root surface area or total root length) was not reported ubiquitously across studies, impeding the analysis of the interactions between K_{rs} , root age and root size. Interestingly, though, an analogous asymptotic relationship between root length and K_{rs} has been previously reported in a modeling study (Meunier, Couvreur, et al., 2017).

To explore the K_{rs} development with age in more detail, we modeled this relationship for four selected crop species, using CPlantBox coupled with MARSHAL (see Section 2.4 for details on data selection and model parametrization). Despite large differences in root size and root architecture (Figure S2), all species exhibited a very similar non-

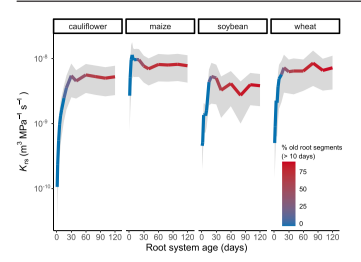
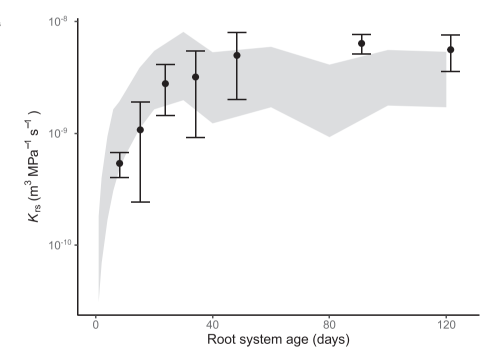


FIGURE 6 Modeled K_{rs} development with age in crop species. Colored lines and shaded areas represent K_{rs} (mean ± standard error) of simulations using CPlantBox coupled with MARSHAL, for four different crops: cauliflower (forb), maize (C_4 grass), soybean (legume) and wheat (C_3 grass). The species were selected based on plant-functional diversity. The color scale indicates the proportion of old (>10 days) root segments in the total root system.

FIGURE 7 Modeled and observed K_{rs} development with age in crop species. Data points and error bars represent K_{rs} (mean ± standard error) of crop species (C_3 and C_4 grasses and dicot crops) from the review and the shadowed area represents the total range of variation in K_{rs} according to simulations (CPlantBox coupled with MARSHAL using four crop species). - American Society of Plant Biologists - SEB-WILEY 13 of 18

are age dependent (Doussan et al., 1998). Specifically, k_r strongly decreases with age (Figure S1), and the radial pathway is commonly considered to be the more limiting one for water transport (Frensch & Steudle, 1989; Lynch et al., 2014). Thus, the counteracting effect of an increase in less conductive tissues (i.e., older root segments) proportionally to total root growth would explain the constancy in K_{rs} at later stages of development. Additionally, it has been shown that even under constant k_r and k_x , K_{rs} can display an asymptotic behavior for roots because of axial flow limitations with increasing root length (Meunier, Couvreur, et al., 2017). Furthermore, the modeled K_{rs} response to age strongly resembled the one observed in the empirical data. In fact, average K_{rs} values at different ages obtained from the review lay within (or very near) the range of variation of the models (Figure 7), indicating that the modeling results were representative of common patterns across studies and species. Whether the mechanisms observed in the models also explain the patterns evidenced in the review remains to be investigated.

The non-linear relationship between K_{rs} and root system age presented here has been reported previously. For instance, a similar pattern was observed in a modeling study with 10,000 virtual maize root systems (Meunier et al., 2019). However, our work is the first – at



linear pattern, that is, a pronounced increase in $K_{\rm rs}$ with age during the first 20 days, followed by rather constant values from day 20 onwards (Figure 6). This behavior was not related to cessation in root growth, as total root length showed a continuous increase during the 120 days of simulation (Figure S2). But, with increasing root age the proportion of "old" root segments (>10-day old segments) also increased (Figure 6). This could have impacted the development of $K_{\rm rs}$, as the radial (k_r) and axial (k_x) hydraulic properties of root segments – which, together with the root architecture, determine $K_{\rm rs}$ – least to our knowledge – to demonstrate a common pattern across studies and species in both experimental data and modeling and to quantify the associated range of variation in K_{rs} over time. Also, the combination of literature data and modeling gave insights about the (possible) causes for the emerging patterns. These results are therefore of relevance and can be a valuable input for the description of root water uptake processes at plant, field or regional scales (Couvreur et al., 2014; Jorda et al., 2022; Nguyen et al., 2022, 2020; Sulis et al., 2019; Vanderborght et al., 2021).

4 | CONCLUSIONS AND OUTLOOK

Here, we presented an extensive review on root hydraulic properties, their variability and some of the factors affecting them. A very large range of variation (orders of magnitude) in K_{rs} , k_{root} , k_r and k_x reported in the literature was identified, but this was not caused by systematic differences among plant functional types (with the only exception of significant differences between axial conductance of woody vs. non-woody species), but rather by the (combined) effect of several factors. We observed that the driving force used for measurement (osmotic vs. hydrostatic), root system age, drought stress and aquaporin inhibition caused changes in root hydraulic properties of 1-2 orders of magnitude, partially explaining the total range of variation in the data. However, because of the inherent constrains of a broad review, we did not quantify the magnitude of other factors that have been reported to affect root hydraulic properties across species, such as soil salinity, low temperature, nutrient availability, or mycorrhizal colonization (see e.g., Aroca et al., 2011; Gambetta et al., 2017; Lehto & Zwiazek, 2011). But, the data compiled here could serve as a valuable input for future studies exploring this topic.

Also, the scope of our study, as well as the interpretation of the results, was partially bounded by data availability. For instance, we did not analyze the effect of anatomical development on root hydraulic properties variability, as only a very low number of the studies included in this review investigated this relationship in detail. Anatomical factors such as root cortical aerenchyma formation (Fan et al., 2007), metaxylem formation (Passioura, 1983; Strock et al., 2021) or suberization (Henry et al., 2012; Ranathunge et al., 2017) affect root hydraulic properties, are involved in their responses to environmental stresses and could also explain the presence of spatial gradients that has been reported in the literature (Frensch & Steudle, 1989; Huang & Nobel, 1992; Knipfer & Fricke, 2011). Additionally, the exploration of the relationship between root system age and whole root system conductance described in Section 3.2.4 was restricted to crop species (grasses and dicot crops), as reliable data on this topic (i.e., detailed datasets on the evolution of K_{rs} with time) was extremely scarce for woody vegetation (shrubs and trees). Finally, we regarded all data included in this review as reflective of the root hydraulic properties of living tissues, unless explicitly mentioned otherwise in the original publications. It is important to note that this approach might potentially lead to misinterpretation of the observed range of variation and the influencing factors, given that the absorption capacity of roots diminishes with tissue decay (Ruark et al., 1983).

Despite these shortcomings, the present study yielded relevant new insights on the variability of root hydraulic properties variability across species and PFTs. Building on that, future studies should specifically target some of the open questions discussed here. The following topics are of special interest: (1) the difference between osmotic and hydrostatic root hydraulic properties was much lower in C_3 grasses (particularly in barley) than in other PFTs; how is this reflected in the water transport patterns of these species?; (2) a large range of

variation was observed in the response of root hydraulic properties to drought, with some indications of differences among PFTs, but clear conclusions were hindered by the extremely low number of studies comparing multiple species and PFTs. Hence, do species corresponding to different PFTs (e.g., dicot crops vs. grasses) respond differently to drought under the same environmental conditions?; (3) a common non-linear relationship between root system age and K_{rs} was identified for several crop species, according to both literature data and modeling. Is such a pattern also present in species from other PFTs (e.g., shrubs or young trees) and how is it reflected in the seasonality of perennial species?; and (4) how is the evolution of K_{rs} affected by processes of root senescence and death? To address these questions. a combination of empirical and modeling approaches could be applied, and we encourage interested researchers to make use of this review as an input for planning such future studies and identifying additional research gaps.

In summary, the present study represents an overview of root hydraulic properties variability across plant functional types, species and experimental conditions and their associated responses. The novel insights obtained here, together with the accompanying data (stored in a database and easily accessible through the web application, https://roothydraulic-properties.shinyapps.io/database/) and additional tools like modeling – as we applied in this study – should be a valuable input for future studies on the role of root hydraulics and root water uptake processes under changing environmental conditions. Moreover, the acquired knowledge holds potential applications in addressing environmental and agricultural challenges.

AUTHOR CONTRIBUTIONS

Juan C. Baca Cabrera: Conceptualization; software; formal analysis; investigation; data curation; visualization; writing—original draft; writing—review and editing. Jan Vanderborght: Formal analysis; funding acquisition; writing—review and editing. Valentin Couvreur: Formal analysis; writing—review and editing. Dominik Behrend: Writing review and editing. Thomas Gaiser: Funding acquisition; writing—review and editing. Writing—review and editing. Guillaume Lobet: Formal analysis; funding acquisition; writing—review and editing.

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CONFLICT OF INTEREST STATEMENT

The Authors did not report any conflict of interest.

PEER REVIEW

The peer review history for this article is available in the Supporting Information for this article.

DATA AVAILABILITY STATEMENT

Not applicable.

ORCID

Juan C. Baca Cabrera b https://orcid.org/0000-0001-8159-3837 Jan Vanderborght b https://orcid.org/0000-0001-7381-3211 Valentin Couvreur b https://orcid.org/0000-0002-1087-3978 Guillaume Lobet b https://orcid.org/0000-0002-5883-4572

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SUPPORTING INFORMATION

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