

Species-area relationships in continuous vegetation

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DOI:

[10.1111/jbi.13697](https://doi.org/10.1111/jbi.13697)

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Document Version

Publisher's PDF, also known as Version of record

Citation for published version (Harvard):

Dengler, J & Matthews, T 2019, 'Species-area relationships in continuous vegetation: evidence from Palaearctic grasslands', *Journal of Biogeography*. <https://doi.org/10.1111/jbi.13697>

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RESEARCH PAPER



Species–area relationships in continuous vegetation: Evidence from Palaearctic grasslands

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Funding information

State Fund for Fundamental Research of Ukraine, Grant/Award Number: Ø83/53427; Eusko Jaurlaritza, Grant/Award Number: IT936-16; Slovenská Akadémia Vied, Grant/Award Number: VEGA 02/0095/19; Narodowe Centrum Nauki, Grant/Award Number: 2017/27/B/NZ8/00316 and DEC-2013/09/N/NZ8/03234; Center for International Scientific Studies and Collaboration (CISSC), Grant/Award Number: NA; Eurasian Dry Grassland Group (EDGG) and the International Association for Vegetation Science (IAVS), Grant/Award Number: NA; Grantová Agentura České Republiky, Grant/Award Number: GA 17-19376S; MIUR initiative "Department of excellence", Grant/Award Number: Law 232/2016; Bavarian Research Alliance, Grant/Award Number: BayIntAn_UBT_2017_58; Bayreuth Center of Ecology and Environmental Research (BayCEER), Grant/Award Number: NA

Abstract

Aim: Species–area relationships (SARs) are fundamental scaling laws in ecology although their shape is still disputed. At larger areas, power laws best represent SARs. Yet, it remains unclear whether SARs follow other shapes at finer spatial grains in continuous vegetation. We asked which function describes SARs best at small grains and explored how sampling methodology or the environment influence SAR shape.

Location: Palaearctic grasslands and other non-forested habitats.

Taxa: Vascular plants, bryophytes and lichens.

Methods: We used the GrassPlot database, containing standardized vegetation-plot data from vascular plants, bryophytes and lichens spanning a wide range of grassland types throughout the Palaearctic and including 2,057 nested-plot series with at least seven grain sizes ranging from 1 cm² to 1,024 m². Using nonlinear regression, we assessed the appropriateness of different SAR functions (power, power quadratic, power breakpoint, logarithmic, Michaelis–Menten). Based on AICc, we tested whether the ranking of functions differed among taxonomic groups, methodological settings, biomes or vegetation types.

Results: The power function was the most suitable function across the studied taxonomic groups. The superiority of this function increased from lichens to bryophytes to vascular plants to all three taxonomic groups together. The sampling method was highly influential as rooted presence sampling decreased the performance of the power function. By contrast, biome and vegetation type had practically no influence on the superiority of the power law.

Main conclusions: We conclude that SARs of sessile organisms at smaller spatial grains are best approximated by a power function. This coincides with several other



comprehensive studies of SARs at different grain sizes and for different taxa, thus supporting the general appropriateness of the power function for modelling species diversity over a wide range of grain sizes. The poor performance of the Michaelis–Menten function demonstrates that richness within plant communities generally does not approach any saturation, thus calling into question the concept of minimal area.

KEYWORDS

logarithmic function, Michaelis–Menten function, minimal area, nested-plot sampling, nonlinear regression, Palaearctic grassland, plant biodiversity, power law, scaling law, species–area relationship (SAR)

1 | INTRODUCTION

Species–area relationships (SARs) represent one of the most fundamental laws in ecology (Lawton, 1999; Lomolino, 2000). Since the early studies by Arrhenius (1921) and Gleason (1922) they have attracted considerable attention (e.g. Connor & McCoy, 1979; Drakare, Lennon, & Hillebrand, 2006; Lomolino, 2001; Rosenzweig, 1995; Triantis, Guilhaumon, & Whittaker, 2012). SARs are of great theoretical interest as different theories of island biogeography (e.g. MacArthur & Wilson, 1967), species abundance distributions (e.g. Pueyo, 2006; Šizling & Storch, 2004) and neutral models (e.g. Hubbell, 2001) predict different shapes of SARs, with the implication that observed SARs can be deployed to test such theories. Furthermore, it is interesting to test how taxonomic group, scale, methodological settings and ecosystem or geographic context influence the relative performance of SAR functions and their parameters (e.g. Chiarucci, Viciani, Winter, & Diekmann, 2006; Crawley & Harral, 2001; Drakare et al., 2006; Storch, Evans, & Gaston, 2005). In addition, SARs allow the area effect to be controlled in assessments of ecological drivers of biodiversity (e.g. Price, 2004; Whittaker, Willis, & Field, 2001). SARs also allow extrapolation of species richness to larger areas that cannot be surveyed with reasonable effort (e.g. Kunin et al., 2018; Plotkin et al., 2000; Ulrich, 2005; Wilson, Peet, Dengler, & Pärtel, 2012). In addition, SARs allow standardization of richness records from several differently sized units to a common grain size, thereby facilitating scale-independent diversity comparisons and visualizations (e.g. Kier et al., 2005) and the identification of biodiversity hotspots (e.g. Fattorini, 2007). Finally, the slope parameters of certain types of SARs are suitable measures of beta-diversity (DeMalach, Saiz, Zaady, & Maestre, 2019; Jurasinski, Retzer, & Beierkuhnlein, 2009).

Historically, studies of SARs have largely been restricted to two functions, (a) the power function (often called the power law; Arrhenius, 1921; Preston, 1962) and (b) the logarithmic function (sometimes erroneously termed the 'exponential' function; Gleason, 1922). This was mainly because the fit of these two functions was easily explored using least squares linear regression techniques. A comparison of a broader set of functions became possible with

the advent of nonlinear regression techniques (e.g. Dengler, 2009; Flather, 1996; Guilhaumon, Gimenez, Gaston, & Mouillot, 2008; Stiles & Scheiner, 2007). In recent years, a wide array of different function types has been proposed and tested (Dengler, 2009, 2010; Tjørve, 2003, 2009). Consequently, several comprehensive studies have been conducted on the fit of different functions and parameters for island SARs as well as other broadscale SARs. Triantis et al. (2012) compared 20 different models for 601 true island datasets around the world and found strong support for the power function overall. Matthews, Guilhaumon, Triantis, Borregaard, and Whittaker (2016) extended this study to 182 habitat islands, with a similar finding. In a further step, they tested how ecological context affects the slope parameter of the power function, and they found systematic differences between island types and spatial scales, but not between major taxa.

While knowledge of functions and parameters of island SARs has been broadly consolidated during the last decade, comparable empirical evidence on small-grain SARs in continuous habitats is still lacking (for theory see Storch, 2016; Williamson, 2003). With continuous habitat or vegetation, we refer to situations where the sampling units do not have a natural border such as islands or habitat islands, but are delimited by the researcher. The influential study of Crawley and Harral (2001) on how biodiversity depends on scale in continuous vegetation, for example, *a priori* only considered the power function. Some regional studies have found a prevalence of the power function using multimodel inference, but were restricted to less than 20 datasets (e.g. Dengler, 2009; Dengler & Boch, 2008). Furthermore, Rosindell and Cornell (2007) obtained power function SARs from a spatially explicit ecological drift model (Hubbell, 2001) within a homogeneous grid model assuming skewed dispersal kernels. By contrast, there is a belief that the logarithmic function should be more suitable at small spatial scales (Gleason, 1922; van der Maarel, 1997). Saturated functions (i.e. functions with a horizontal upper asymptote) are also often assumed to represent SARs in continuous vegetation *weö*, inspired by the still widespread, but flawed (see Barkman, 1989) concept of so-called 'minimal areas' (e.g. Mueller-Dombois & Ellenberg, 1974), which was assumed to be the scale at which species richness is saturated for a given community. Additional confusion

around small-grain SARs was caused when confounding different sampling schemes with SARs in the strict sense (i.e. those originally considered by Arrhenius, 1921, or Preston, 1962). For example, Stiles and Scheiner (2007) and DeMalach et al. (2019) reported that the logistic function (a saturated function) and the logarithmic function, respectively, performed much better than the power function. However, they had analysed species accumulation curves, merging non-contiguous sample units (also called species-sampling relationships, SSRs; see Dengler, 2009; Fridley, Peet, van der Maarel, & Willems, 2006), and not SARs in the strict sense. In conclusion, this situation calls for a comprehensive, multimodel inference analysis of small-grain SARs in continuous vegetation, comparable to the analyses of Triantis et al. (2012) and Matthews et al. (2016) for island SARs.

As the Palaearctic biogeographic realm comprises more than one third of the world's ice-free terrestrial surface and spans a wide range of climatic and topographic gradients, it harbours a high number of vegetation types and considerable biodiversity (Rounsevell, Fischer, Torre-Marín, & Mader, 2018). Around 22% of the Palaearctic is composed of various grassland types (Török & Dengler, 2018), some of them being the world record holders of small-grain vascular plant diversity (Wilson et al., 2012). A large proportion of the Palaearctic grasslands are primary grasslands such as steppes and arctic-alpine grasslands. Even in regions where the potential vegetation is forest, natural grasslands occur in azonal and extrazonal conditions. Moreover, agricultural activities and pastoralism long present in the Palaearctic has resulted in the creation of secondary grasslands dependent on human land use that prevents succession towards shrublands or forests (Török & Dengler, 2018). The coverage of major ecological gradients and the high diversity of vegetation types across several biogeographic regions highlight Palaearctic grasslands as an excellent model system to study small-grain SARs and how they are affected by different factors.

Here, we used more than 2,000 nested-plot series from the GrassPlot database (Dengler et al., 2018), from a wide range of grassland types across six biomes, to perform a comprehensive analysis of small-grain (1 cm^2 – $1,024\text{ m}^2$) SARs in continuous vegetation for vascular plants, bryophytes and lichens. Specifically, we aimed to address the following questions using the Palaearctic grasslands as an example:

1. Which function is most appropriate to describe small-grain SARs?
2. Does the performance of the different functions depend on factors such as sampling method, taxonomic group, biogeographic setting or vegetation type?

2 | MATERIALS AND METHODS

2.1 | Vegetation-plot data

We used plot data from the collaborative vegetation-plot database GrassPlot (Dengler et al., 2018; <http://b.link/grassplot>), which is registered in the Global Index of Vegetation-Plot Databases (GIVD;

Dengler et al., 2011) as EU-00-003. GrassPlot collects vegetation-plot data (both richness and composition) together with methodological, environmental and structural information from grasslands as well as other plant communities dominated by herbs, dwarf-shrubs or cryptogams from the Palaearctic biogeographic realm (for delimitation see Figure S1.1). Requirements for inclusion are that the plots (sampling units) were precisely delimited in the field and carefully sampled with the aim of achieving complete species lists. One strength of GrassPlot is the numerous multi-scale datasets derived from a diversity of nested-plot sampling schemes (e.g. Dengler et al., 2016) of areas from 1 cm^2 to $1,024\text{ m}^2$ (schemes of the three main types of sampling designs in Figure S2.1).

We retrieved all nested-plot series contained in GrassPlot (v.1.27 on 4 January 2019) that comprised at least seven different grain sizes (see overview of the 69 datasets with these data in Table S1.4). In total, there were 2,057 series with vascular plant information (Figure 1), of which 757 also contained bryophyte data and 780 lichen data (Figure S1.2). The plots were distributed over 26 different countries from 34.9° to 68.9°N , from 9.1°W to 161.8°E and covered an altitudinal gradient from 0 to $4,387\text{ m a.s.l.}$ (Figure 1, Figure S1.2). In total, the nested-plot series consisted of 139,265 individual subplots with richness data, often with several replicates per grain size. Further characteristics of the used datasets are provided in Appendix S1.

For those nested-plots series with more than one subplot for a certain grain size, we averaged richness values across subplots and stored the information on how many subplots the average was based on. Thus, we obtained one single richness value per each grain size within each nested-plot series, if possible, for four different taxonomic groups (1 – complete terricolous macroscopic vegetation; 2 – vascular plants; 3 – terricolous bryophytes; 4 – terricolous lichens). We also recorded whether plots were sampled with the shoot presence or with the rooted presence method (for terminology, see Dengler, 2008).

2.2 | SAR modelling

From the numerous different functions proposed for modelling SARs (Dengler, 2009; Tjørve, 2003), we selected three main functions that have specifically been suggested and used for SAR modelling in continuous vegetation (DeMalach et al., 2019; Dengler & Boch, 2008): the power function, the logarithmic function (often erroneously termed the exponential function) and finally the Michaelis–Menten function as a simple two-parameter example of a SAR with saturation (i.e. an upper threshold of richness). To account for possible 'scale dependence' of the SAR, we added two variants of the power function that allow for exponents to change with area: the 'quadratic power function' with a continuous change of the exponent, and the 'breakpoint power function' with an abrupt change of the exponent at a certain grain (e.g. Dengler, 2010). The five functions were selected to represent fundamentally different shapes (Dengler, 2008; see also Figure S3.1) as well as different complexities (number of fitted parameters; Table 1).

We fitted all five functions for both species richness S (S -space; 'linear space') and for $\log S$ ($\log S$ -space; 'logarithmic space') as

FIGURE 1 Density and spatial distribution of the 2,057 nested-plot series in the Palearctic biogeographic realm that were analysed in this study. The colour scale indicates the number of available series per 10,000-km² grid cell. The map uses the Europe Lambert Conformal Conic projection

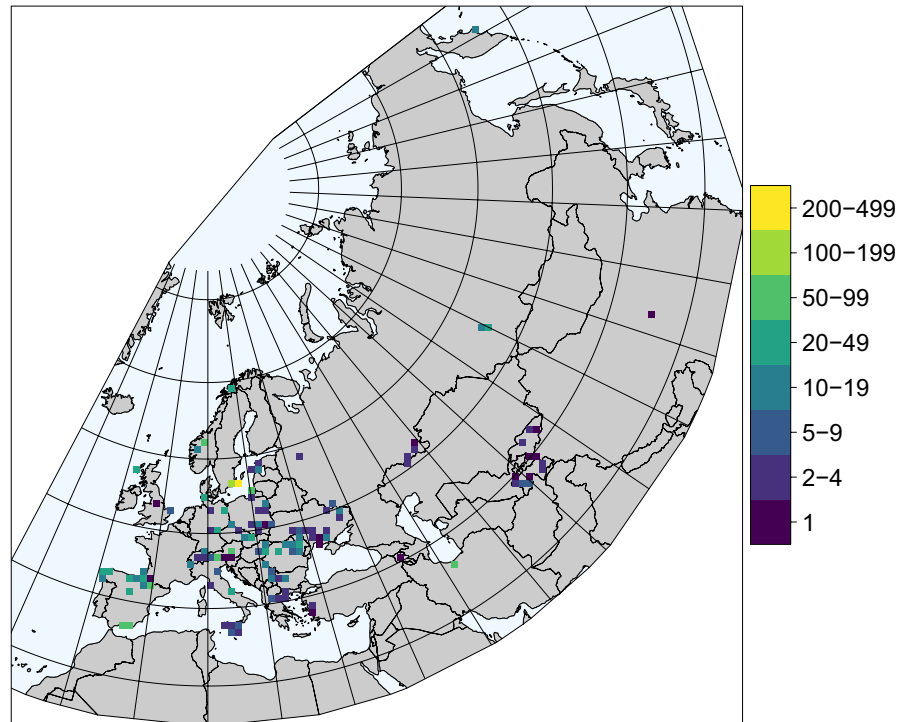


TABLE 1 The five function types used in this study to model species–area relationships (SARs). All functions were fitted both in *S*-space and in log *S*-space. The following notations are used: *S* = mean species richness; *A* = area/m²; log = log₁₀. The *k* fitted parameters (except the variance) are termed *c*, *z*, *z*₁, *z*₂, *b*₀, *b*₁ and *T*

Function name	Abbreviation	<i>k</i>	Formula in <i>S</i> -space	Formula in log <i>S</i> -space	Meaning of parameters
Power	powSAR	2	$S = c A^z$	$\log S = \log c + z \log A$	<i>c</i> = richness at unit area (1 m ²); <i>z</i> = steepness parameter (exponent in <i>S</i> -space or slope in log <i>S</i> -space)
Power quadratic	powQSAR	3	$S = 10^{(\log c + z_1 \log A + z_2 (\log A)^2)}$	$\log S = \log c + z_1 \log A + z_2 (\log A)^2$	<i>c</i> = richness at unit area (1 m ²); <i>z</i> ₁ = steepness parameter; <i>z</i> ₂ = change of steepness with increasing area
Power breakpoint	breakSAR2	4	$S = 10^{[\log c + (\log A < \log T) (z_1 \log A) + (\log A \geq \log T) (z_1 \log T + z_2 (\log A - \log T))]}$	$\log S = \log c + (\log A < \log T) (z_1 \log A) + (\log A \geq \log T) (z_1 \log T + z_2 (\log A - \log T))$	<i>c</i> = richness at unit area (1 m ²); <i>T</i> = breakpoint (area at which the steepness changes); <i>z</i> ₁ = steepness parameter for <i>A</i> > <i>T</i> ; <i>z</i> ₂ = steepness parameter for <i>A</i> ≥ <i>T</i>
Logarithmic	logSAR	2	$S = b_0 + b_1 \log A$	$\log S = \log (b_0 + b_1 \log A)$	<i>b</i> ₀ = intercept (in <i>S</i> -space); <i>b</i> ₁ = steepness parameter
Michaelis–Menten	mmSAR	2	$S = b_0 A / (b_1 + A)$	$\log S = \log (b_0 A / (b_1 + A))$	<i>b</i> ₀ = saturation value (modelled maximum richness); <i>b</i> ₁ = steepness parameter

Note: The logical expressions (log *A* < log *T*) and (log *A* ≥ log *T*) return 1 if they are true and 0 if they are false.

dependent variables using nonlinear regression (Table 1). Both approaches are valid, have been used in the literature, and have different strengths and limitations (see Dengler, 2009). Due to the different treatment of the error structure, the parameter estimates in the two spaces usually slightly deviate. Generally, fitting in *S*-space gives more weight to good fit at large grain sizes, whereas fitting in log *S*-space gives more weight to good fit at small-grain sizes. Moreover, fitting in log *S* typically reduces heteroscedasticity in the residuals.

As fitting in log *S*-space is not possible if some subplots have *S* = 0 (excluding such cases is not recommended; Dengler, 2010; Williams, 1996), we addressed this issue as follows. Fitting nested-plot series in the optimal case means that the richness value for the smaller grain sizes is representative for the whole area of the largest plot, which could be achieved by full tessellation of its area and averaging the richness values of all resulting subplots. In such optimal sampling, evidently the mean richness value of any smaller

grain size would be >0 if there was at least one species in the largest plot. However, in most cases, the nested-plot sampling schemes in GrassPlot recorded only one or few replicates of smaller plots. In such cases, the recorded (average) richness value may be $S = 0$, while the true average (calculated from a large number of plots) would be positive. As richness can take only positive values, an observed richness of 0 based on a single plot is a biased estimate as it represents the range of $[0, 0.5]$, while an observed richness based on a single plot of 1 is an unbiased value for the range of $[0.5, 1.5]$. Therefore, we replaced 0 with 0.25, that is, the mean of the lower and upper border of the range of values for which it stands. Likewise, if an observed mean richness of 0 was based on n replicates, we assigned a mean richness of $0.25/n$.

The five models were fitted in R (Version 3.5.1; R Core Team, 2018) using a combination of linear and nonlinear regression. Fitting in S -space always employed nonlinear regression, and optimization used the 'mle2' function in the 'bbmle' R package (Bolker & R Core Team, 2017). As the optimization algorithm was sensitive to the starting parameter values provided, a brute-force approach was used to find parameter values that maximized the likelihood, for a given model. For each model (e.g. power, breakpoint), a grid of multiple different starting parameter values was created. The size of this grid depended on the model, with the more complex models having a larger number of potential starting parameter values. Model optimization was then undertaken multiple times using the 'mle2' function and the starting parameter values from each row in the grid. Finally, the starting parameter values that resulted in the model fit with largest maximum likelihood were chosen. The AICc and $R^2_{adj.}$ values for the model fit optimized using these starting parameter values were then calculated. We consider AICc and $R^2_{adj.}$ as adequate measures for the relative appropriateness/superiority of the compared SAR functions despite the non-independence of the data points in our nested-plot data. In Appendix S4 (R codes in Appendix S5 and S6), we sampled from virtual landscapes where the shape of the SARs is known to test whether nested-plot sampling introduces biases in the model selection using AICc and $R^2_{adj.}$. The 'true shape' of the SARs in these virtual landscapes was determined by averaging the results of several random non-nested plot series of different grain sizes. The results show that the model ranking obtained by nested-plot sampling is close to the true pattern and actually depicts, on average, the true pattern better than SARs constructed from a series of single non-nested plots in the same landscape would do. Accordingly, we consider our approach as valid.

In a small number of cases, there were multiple optimized model fits (i.e. with different parameter estimates) with identical (maximum) likelihood values; here, we simply selected one set of parameter values at random. Following standard statistical convention, the variance was always considered as an additional parameter when calculating AICc. Thus, for example, the power model was considered to have three parameters when calculating AICc. For the power breakpoint model, a further model-fitting step was implemented. In certain cases, the best-selected power breakpoint model using the aforementioned approach contained a z -value that was greater than

1 or less than 0. This z -value was then fixed at either 0 or 1 (depending on which of these values it was initially closest to) and the model fitting process repeated. If both original z -values were out of bounds, this additional step was not undertaken. For a given model and plot series, the above model fitting process was repeated across all four taxonomic groups.

For the log S -space analyses, the logarithmic, Michaelis–Menten and breakpoint power functions were fitted using the nonlinear fitting procedure outlined above, whereas the power model and the quadratic power function were fitted using linear regression and the standard 'lm' function in R. The overall model fitting process was relatively computationally demanding and took approximately 48 hr on a 24-core computer cluster (100 GB RAM). Due to the brute-force approach, we achieved convergence of all models for all taxa in all datasets in the log S -space, and a negligible amount of non-convergence in the S -space (maximum 4% for lichens, but 0% for complete vegetation). The R code used to run the analyses is available as Appendix S7.

2.3 | Ranking and comparison of the SAR functions

We ranked model performance in five ways. First, we counted for how many nested-plot series a certain function performed best among all compared functions, using model selection based on AICc (Burnham & Anderson, 2002). Second, for each function we calculated the Akaike weights based on AICc in each nested-plot series. Akaike weights can be interpreted as the probability that the function i is the best model for the observed data, given the set of five candidate models (Johnson & Omland, 2004). Third, for each function by nested-plot series combination we calculated Δ_i , that is, the difference in AICc of the particular function compared to the respective best performing function ('delta AICc'). Fourth, we ranked models using $R^2_{adj.}$, which was calculated using the formula: $1 - (1 - R^2)(n - 1)/k$, where R^2 is the standard R^2 , n is the number of data points and k is the residual degrees of freedom. Fifth, we determined the best performing function based on the Bayesian Information Criterion (BIC) as there is no clear agreement on the superiority of AIC/AICc versus BIC (Burnham & Anderson, 2002; Johnson & Omland, 2004). The five comparisons were undertaken only in cases where our fitting procedure yielded a result for all five models. Note that model comparisons are restricted to each 'space', that is, measures of goodness of fit or information content (e.g. $R^2_{adj.}$, AICc) cannot be compared between S -space and log S -space (Dengler, 2009).

As sampling methodology has been repeatedly suggested to influence the shape of SARs (Dengler, 2008; Williamson, 2003), we tested for an effect of some key sampling method aspects using ANOVAs and linear regressions: (a) shoot versus rooted sampling of plants (both methods are widespread; see Dengler, 2008); (b) whether the richness of smaller grain sizes was averaged from several replicated subplots or not; and (c) number of grain sizes in a series (distribution of the different methodological choices and other data in Table S1.1 and Figure S1.4). Furthermore, we tested whether

the performance of the functions depended on (d) taxonomic group (vascular plants, bryophytes, lichens), (e) biome (Bruehlheide et al., 2019; based on Schultz, 2005), (f) vegetation type or (g) richness in the largest plot of the series (see Figure S1.1, Tables S1.2 and S1.3).

3 | RESULTS

3.1 | General suitability of the compared functions

Given the wide range of vegetation types studied, the species–area curves also varied widely (Figures S8.1 and S8.2). For all taxonomic groups and irrespective of S -space versus log S -space, the power function was by far the best model when using AICc as a model selection criterion (Figure 2). For the richness of the complete vegetation, it was the best model in 68.1% of all cases in S -space, with values slightly dropping from vascular plants (57.8%) to bryophytes (56.0%) to lichens (49.5%). The superiority of the model was even slightly higher in the log S -space than in the S -space. For the complete vegetation, the second best model, though clearly inferior, was the quadratic power function (in both spaces), while the logarithmic function was second best for vascular plants in both spaces as well as for bryophytes and lichens in log S -space. The Michaelis–Menten saturation function generally performed poorly, but was the second best model for bryophytes and lichens in S -space. When considering BIC instead of AICc (Figure S8.3), the ranking of functions changed. The breakpoint power function performed best followed by the ‘normal’ power function and the quadratic power function, while the logarithmic function and the Michaelis–Menten function had negligible support. The overall support for the three variants of the power function combined increased from c. 60%–90% in case of AICc to c. 90%–95% in case of BIC.

When considering not only the best model, but also the relative performance of all five models via Akaike weights (Figures S8.4 and S8.5) or delta AICc (Figure S8.6), the results remained qualitatively similar, but the superiority of the power function was even clearer, with the mean Akaike weight reaching as high as 71.1% in the case of the complete vegetation in log S -space. Based on R^2_{adj} , that is, considering only the fit, not the complexity, the three power functions performed very well (mostly above 0.95 for all taxa and vascular plants, and mostly above 0.85 for bryophytes and lichens), while the performance of the logarithmic and Michaelis–Menten functions was substantially worse (Figure S8.7). Owing to the one or two additional parameters, the quadratic and breakpoint power function, respectively, had a slightly better fit than the normal power function.

The resulting parameter estimates of all five models and their descriptive statistics are provided in Appendix S9. Here, we summarize only the results of the power function parameter estimates, as it was clearly the overall best model. In particular, we focus on a few parameters that are particularly relevant for interpretation. The slope parameter (z -value) of the overall best performing function (power function) in S -space was 0.20 ± 0.05 (mean \pm standard deviation) for all taxa, with slight variation among the three taxonomic groups (vascular plants: 0.26 ± 0.11 ; bryophytes: 0.19 ± 0.12 ; lichens: 0.28 ± 0.14). In log

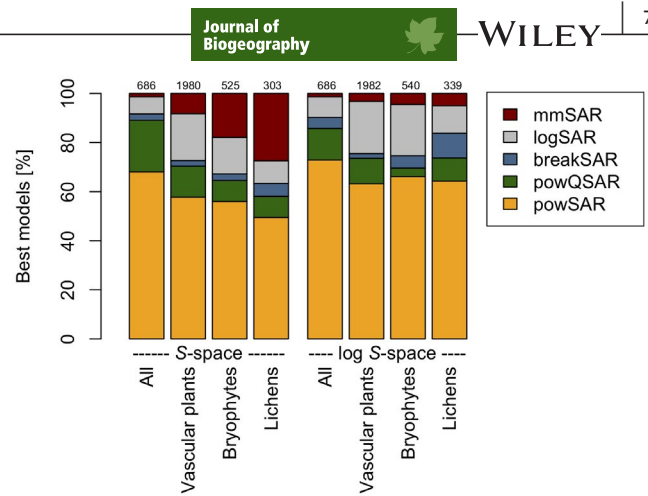


FIGURE 2 Model performance in comparison of the five function types: power (powSAR), power quadratic (powQSAR), power breakpoint (breakSAR), logarithmic (logSAR) and Michaelis–Menten (mmSAR), expressed as fraction of cases where a given model performed best based on AICc. The comparisons were run for the complete terricolous macroscopic vegetation (all species), vascular plants, terricolous bryophytes and terricolous lichens, and both in S -space and log S -space

S -space, the values showed a similar pattern with little deviation in absolute values from S -space (Table S9.1). The z_2 estimate of the quadratic power function was significantly negative (t -test with 0 mean as null model) for all taxa ($p < .001$; mean: -0.017 ± 0.047 , median: -0.012), with similar trends for vascular plants ($p = .09$; mean: -0.061 ± 1.634 , median: -0.019), bryophytes ($p < .001$; mean: -0.105 ± 0.548 , median: -0.009) and lichens ($p = .07$; mean: -0.844 ± 8.204 , median: -0.041) (Table S9.2).

3.2 | Factors influencing function performance

The relative performance of the five models was strongly influenced by several methodological factors: (a) rooted sampling drastically decreased the relative performance of the power function compared to shoot sampling (Figure 3), while the quadratic power and breakpoint power models performed relatively better (Figure S8.8). (b) Likewise, in nested-plot series where the smallest plots were not replicated and averaged, the relative performance of the power function was much worse than when an averaging had taken place (Figure 4, Figure S8.9). (c) The number of included grain sizes (not necessarily correlating with the grain size range) also decreased the superiority of the normal power function, while the two other variants of the power function increased in relative performance, and together all three variants of the power function were even more superior when more grain sizes were sampled (Figure S8.10).

Biome had hardly any influence on the superiority of the power function irrespective of taxonomic group (Figure 5). Only for vascular plants the relative performance of the power function slightly was worse in the ‘Dry tropics and subtropics’ and in the ‘Subtropics with winter rain’ than in the other four biomes.

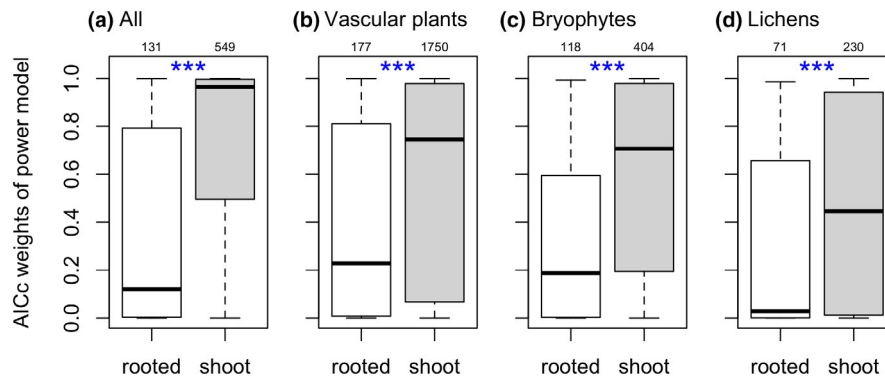


FIGURE 3 Differences in model performance of the power function expressed as AICc weights between the two fundamental ways of recording plants, rooted presence and shoot presence. 'Rooted presence' counts species in the point where they are attached to the ground irrespective whether they have roots in the anatomic sense or not. The displayed values are for the S-space (results in log S-space were consistent)

Likewise, the 18 different major vegetation types hardly showed any difference in the superiority of the power function; the few significant differences in the ANOVA were mostly related to types with only very few replicates (indicating that this might just be a random deviation and not a property of the respective type) (Figure S8.11). However, one vegetation characteristic had a significant influence on the relative performance of functions, at least in vascular plants: the relative performance of the power function strongly increased with the number of species in the biggest plot (Figure S8.12).

4 | DISCUSSION

4.1 | The nature of the species–area relationship

We found strong support for the power function SAR at small-grain sizes in continuous vegetation using more than 2,000 nested-plots over large ecological, geographical and diversity gradients for three major taxa and when focusing on the complete vegetation. Using AICc and R^2 as measures, the 'normal' power function was on average the best model. Using BIC, the breakpoint power function prevailed, and the quadratic power function had a similar level of

support to the normal power function. This difference is not astonishing as BIC penalizes complexity of a function differently than AICc, but actually less strongly for small sample sizes, which might lead to overfitting. If basing the conclusions on BIC, there might be some scale dependence of the SAR, that is, a minor change of the exponent z with grain size (see also Crawley & Harral, 2001). If all three variants of the power function are considered jointly, their prevalence as the best model increased from c. 60%–90% based on AICc to c. 90%–95% based on BIC. With our simulation (Appendix S4), we could further demonstrate that this result was not caused by the non-independence of the nested plots, but that this sampling approach, if at all, might even slightly underestimate the superiority of the normal power function.

The general superiority of the power function was largely unaffected by taxonomic group, biome or vegetation type. This finding is in line with previous regional studies analysing small subsets of the current database (Dengler, 2009; Dengler & Boch, 2008; Fridley et al., 2006). Although we restricted our comparisons for pragmatic reasons to a smaller set of functions, which still provides a good representation of the overall range of possible SARs, our findings are consistent with those of Triantis et al. (2012) and Matthews et al. (2016) for true islands and habitat

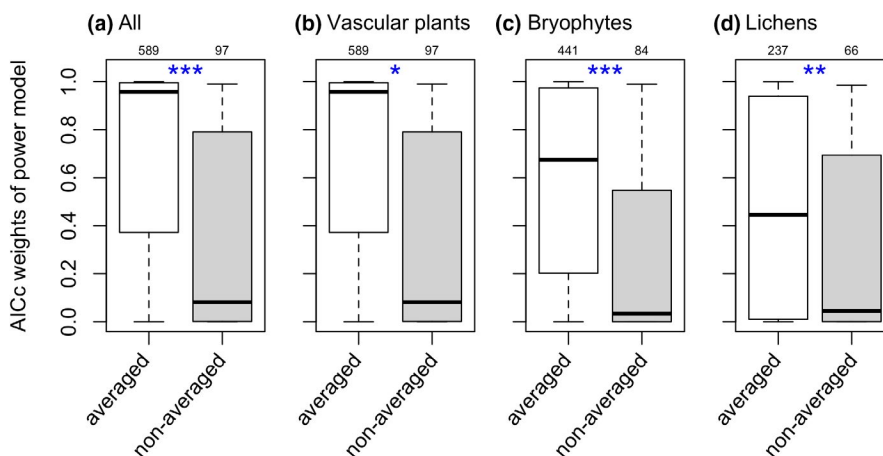


FIGURE 4 Differences in model performance of the power function expressed as AICc weights between sampling schemes where smaller grain sizes were replicated and their richness averaged and cases with only one subplot per grain size (non-averaged). The displayed values are for the S-space (results in log S-space were consistent)

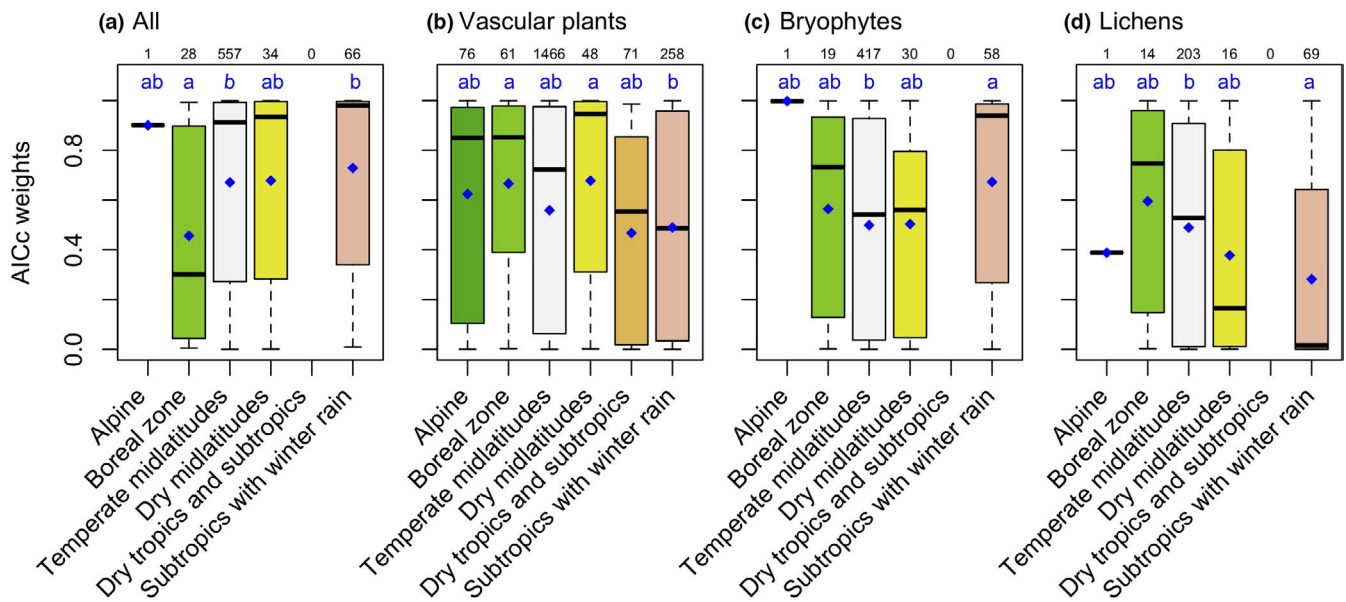


FIGURE 5 Comparison of model performance of the power function expressed as AICc weights between the six biomes represented in the study. The displayed values are for the *S*-space (results in log *S*-space were consistent)

islands, despite the very different study systems and scales. This suggests that, in spite of the commonly accepted notion that contrasting factors influence species diversity at different spatial scales (Brown & Peet, 2003; Field et al., 2009; Shmida & Wilson, 1985; Siefert et al., 2012), the resulting SARs are astonishingly similar over many orders of magnitude (see also Wilson et al., 2012) and across taxa and ecological conditions. Although the power function has been repeatedly criticized (e.g. Pan, Zhang, Wang, & Zhu, 2016; Stiles & Scheiner, 2007), our study supports the idea that it is indeed the one model that, at least for plant communities, can be universally applied (note that even in those cases where it was not the best model, it performed very well; see Figure S8.7). In contrast, other models are suitable, at best, in only a few specific cases.

This poses the question of why a single function (but with varying parameters, see next subsection) can be suitable across so many different situations. In fact, power law SAR-like relationships are far from restricted to species diversity versus area, but can likewise be found in other natural phenomena, such as species frequency versus body size, or body size versus area (Southwood, May, & Sugihara, 2006), or even in completely different realms of science and everyday life (Nekola & Brown, 2007; but see Stumpf & Porter, 2012, for a critical view). A general finding from these different disciplines is that power functions most often result from non-equilibrium conditions (Mitzenmacher, 2012) or skewed underlying distributions (e.g. Rosindell & Cornell, 2007). Power law relationships are likely the consequence of complex dynamical systems, not necessarily of specific ecological mechanistic processes (Nekola & Brown, 2007), even if the slopes of the power law SARs might well be effected by such processes. In this respect, it is interesting to compare the relative performance of the power function across

taxonomic groups. Performance was highest for all species groups combined, followed by vascular plants, bryophytes and lichens, which corresponds to the mean species richness of each group. Moreover, in vascular plants (the groups with the biggest dataset), we found a strong increase in the superiority of the power function with the richness in the biggest plot. It seems that the more elements (here: species) with slightly different properties (e.g. frequencies, habitat preferences, sizes) are involved, the more closely power functions are approached.

In addition, our findings suggest that there is likely no saturation in SARs in continuous vegetation as our saturation function (Michaelis-Menten) performed on average much worse than the functions without saturation across a wide array of different ecological conditions. We believe that even the few individual datasets where the Michaelis-Menten function appeared to be superior are likely artefacts of insufficient replication at the smaller grain sizes. As Dengler and Boch (2008) have shown, the relative performance of the power function versus saturation functions improves when the replication of smaller subplots is increased and thus the calculated average richness is closer to the true mean richness. This is in line with our finding of best fits for the Michaelis-Menten function for bryophytes and lichens in *S*-space. As these groups often have few species in grasslands, in many cases none of the smaller subplots (across several grain sizes) contained any species, resulting in a recorded richness of 0, despite the fact that the true average must be higher and increase with grain size (see Methods). We thus recommend that the concept of 'minimal area' (which only has a meaning if saturation exists), that has been presented in numerous textbooks of vegetation science (e.g. Barbour, Burk, & Pitts, 1999; Kent, 2012; Mueller-Dombois & Ellenberg, 1974) for over a century, should be completely abandoned, as has already

clearly been stated by previous studies (e.g. Barkman, 1989; van der Maarel, 1996).

The same holds for the logarithmic function (in the literature also termed 'exponential' or 'semi-log'). There is a widespread belief in vegetation science that this function is particularly appropriate at small-grain sizes (Gleason, 1922; He & Legendre, 1996; van der Maarel, 1997; Stohlgren, Falkner, & Schell, 1995), but the origin of this impression is unclear. For example, van der Maarel (1997) claims this despite the fact that the curvature of the SARs in his figure clearly suggests the better fit of a power law SAR. In addition, the logarithmic function cannot serve as an appropriate model for the SAR as it necessarily predicts negative richness values for small positive areas, often even within the fitted range (Dengler, 2008; see also Figure S3.1). The very poor performance of the logarithmic function across more than 2,000 grassland plots series throughout the Palaearctic matches the findings for multiple habitat types on the Curonian Spit, Russia (Dengler, 2009), and the south-eastern United States (Fridley, Peet, Wentworth, & White, 2005).

While we could rule out the logarithmic function and saturated functions as suitable models, at closer inspection, we found very small but consistent deviations from a power function with a uniform z -value. Increasing the number of replicates strongly increased the relative support of the power function variants with a varying exponent. Moreover, we found that the z_2 -value of the quadratic power function was significantly negative across all studied nested-plot series (e.g. -0.017 for all taxa), meaning that the actual slope is slightly decreasing towards larger grain sizes. While this pattern was to be expected for rooted sampling (Williamson, 2003), we found it also for shoot sampling (not shown), which would support the steep-flat-steep triphasic theory of SARs by Storch, Keil, and Jetz (2012). Our results show that such minor deviations could conveniently be accounted for in the power model by allowing z -values to change with grain size in a systematic manner (with the quadratic or the breakpoint variants of the power function).

4.2 | Methodological aspects

A few studies have found a much better performance of saturated and/or logarithmic functions compared to power functions at small spatial scales (DeMalach et al., 2019; Stiles & Scheiner, 2007). However, these authors analysed species accumulation curves and species-sampling relationships (SSRs; Dengler, 2009) rather than SARs in the strict sense (see the typology of Dengler, 2009), and thus these findings are not surprising. Even though their SSRs were also based on 'areas' (and thus many researchers continue calling them SARs in agreement with Scheiner, 2003), they have fundamentally different mathematical properties (Dengler, 2009). We illustrate this with our conceptual Figure 6 and Table 2. SSRs (whether based on individuals, samples or areas) increase sampling intensity within the same pre-defined focal area, while SARs in the strict sense actually increase the focal area. SSRs thus must be a saturation function by definition,

as also shown by the simulations of Dengler & Oldeland (2010). The fact that in such situations the logarithmic function also performs well (or even better than the rather inflexible saturated Michaelis-Menten function) as well as (or even better than) the rather inflexible saturated Michaelis-Menten function) has to do with the similar shapes of the two functions – at smaller grain sizes both become steeper or, in other words, shows an increasing negative deviation from the power function (Figure S3.1). Irrespective of whether area-based SSRs are called SARs or not, results (model superiority and parameter estimates) from analysing this type of curves are not comparable with those of SARs in the strict sense (whether these are nested-plot SARs in continuous habitats or island SARs).

Uncertain richness estimates, particularly underestimations might also mask fits of the power function and increase the relative performance of other models. For instance, Guilhaumon et al. (2008) reported relatively poor performance of power functions and large uncertainties in predictions of global hotspot species richness due to low or uncertain sample coverage. This coincides with our finding that the superiority of the power function was lowest for bryophytes and lichens, the two taxa with the lowest richness in most cases, because low absolute richness means that even a recording error of one species can be a substantial relative error. Likewise, the superiority of the power function increased when the mean richness values at small-grain sizes were based on averages and thus more reliable than when they were based on single counts. This was also found by Dengler & Boch (2008), who argued that adding random noise to the true relationship by chance will lead to higher superiority of other functions in some cases. We found that other methodological aspects can have pronounced effects on model superiority even when focusing on SARs in the strict sense. Specifically, we found that the power function performed much better for shoot presence sampling than for rooted presence sampling, which is in line with the predictions of Williamson (2003) and Dengler (2008). Theoretically, both of the widely applied ways to record plants in plots must theoretically lead to deviations from the shape of a perfect power function towards the smallest grain sizes, with z -values of the shoot presence method approaching 0 and those of the rooted presence method approaching 1. However, the deviation from a relatively constant z -value at larger grain sizes should appear at relatively larger grain sizes for rooted sampling and be more pronounced (Dengler, 2008), which evidently causes the much lower relative performance of the power function at the small-grain sizes studied here, in the case of rooted sampling in otherwise similar communities.

4.3 | Conclusions and outlook

While a perfect power function theoretically cannot hold across all grain sizes (Storch et al., 2012) from, for instance, 1 cm^2 to the terrestrial surface of the Earth (130 million km^2 , or 18 orders of magnitude), we found that it is a very good approximation for sessile organisms across the already large range of six orders of

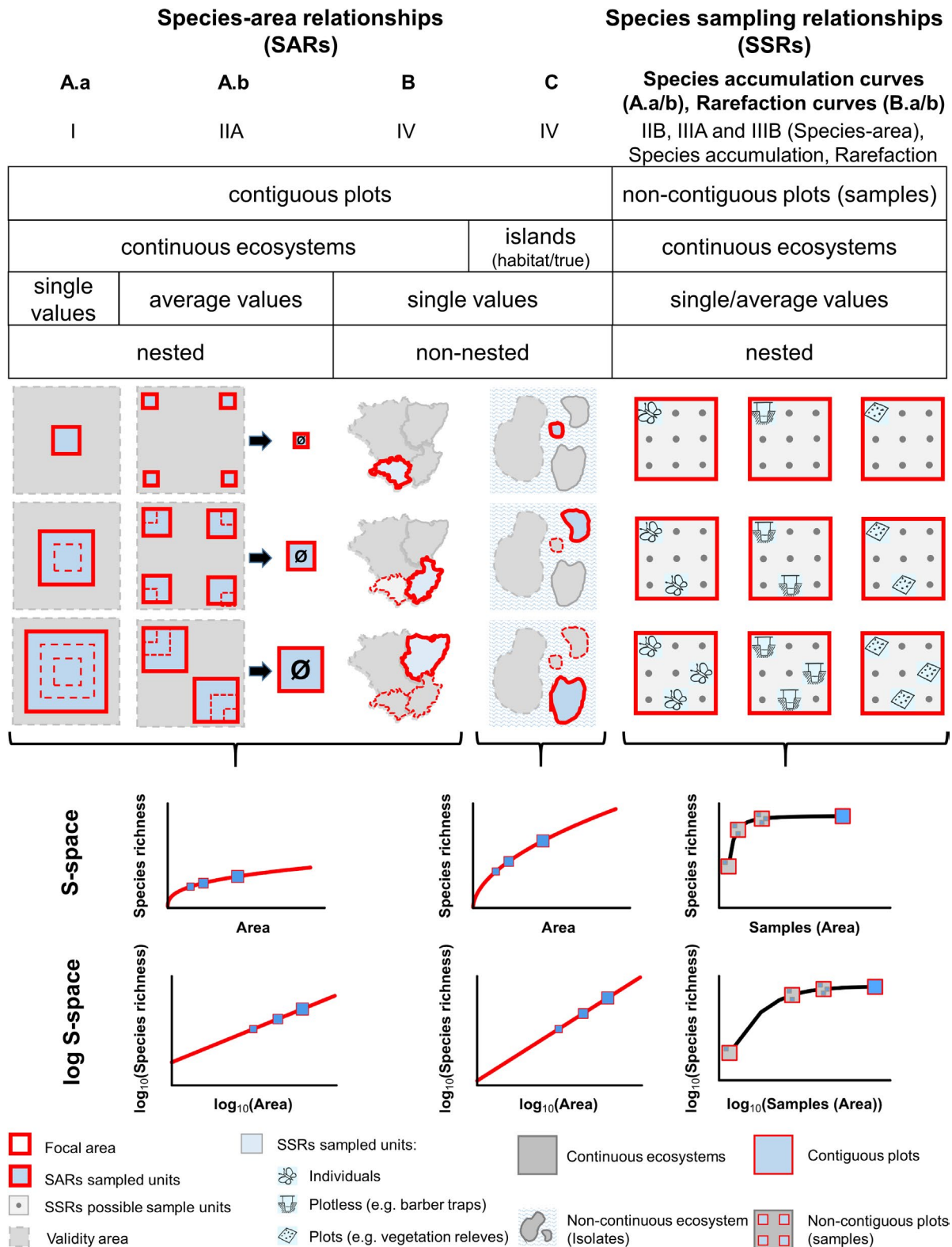


FIGURE 6 Overview of the main types of species richness curves in terms of spatial arrangement of sampling units and their combination to 'areas' as well as the resulting function shapes. It is evident that species-area relationships (SARs) in the strict sense differ fundamentally from area-based species-sampling relationships (SSRs = species accumulation curves). The species richness curves are assigned to the typologies of Dengler (2009, bold) and Scheiner (2003, normal font)

Criterion	SARs	SSRs
Sample coverage	Complete species list	Representative sample
Area on which one richness count is based	Contiguous	Non-contiguous (except for the finest grain size)
Sampling effort	Constant	Increasing
Focal area	Increasing	Constant
Function type	Unbound	Asymptotic

TABLE 2 Major differences between species–area relationships (SARs) in the strict sense and species–sampling relationships (species–sampling relationships = species accumulation curves)

magnitude in our study – despite the very wide ecological and floristical gradients included (e.g. 6 of the 10 global biomes, 18 major vegetation types). This is in line with the findings of the equally comprehensive studies of Triantis et al. (2012) and Matthews et al. (2016), who found a similar superiority of the power function across many orders of magnitude for multiple taxa in true and habitat islands at much larger grain sizes than in our study, but equally across many orders of magnitude. The superiority of the power function has also been shown at similar grain sizes and in continuous vegetation as well as habitats other than grasslands (e.g. forests and wetlands) (Dengler, 2009; Fridley et al., 2005). This leads us to conclude that the power function is a suitable (and mostly the best possible) model for SARs in nearly any situation, provided the areas from which the relationship is constructed are contiguous. For curves constructed from virtual areas consisting of non-contiguous sub-units (as in the case of area-based species accumulation curves), a saturated function, rather than a power function, is to be expected (Dengler & Oldeland, 2010). As a consequence, power functions are usually not suitable for estimating species loss due to habitat loss, as the remaining habitat is typically highly fragmented (Hanski, Zurita, Bellocq, & Rybicki, 2013). However, in all cases with contiguous areas, be it islands (of any type) or areas in continuous habitats delimited by the researcher, power function SARs are suitable tools for interpolation and extrapolation of species richness, or for removing the area effect if other drivers of biodiversity are the focus. Moreover, power function SARs provide, with their exponent (z-value), a meaningful (and standardized) beta-diversity measure in continuous vegetation (Jurasinski et al., 2009; Polyakova et al., 2016), enabling the effective comparison of species turnover among taxa or between different ecological conditions.

ACKNOWLEDGEMENTS

We thank all vegetation scientists who carefully collected multi-scale plant diversity data from Palaearctic Grasslands available in GrassPlot. The Eurasian Dry Grassland Group (EDGG) and the International Association for Vegetation Science (IAVS) supported the EDGG Field Workshops, which generated a core part of the GrassPlot data. The Bavarian Research Alliance (grant

BayIntAn_UBT_2017_58) and the Bayreuth Center of Ecology and Environmental Research (BayCEER) funded the initial GrassPlot workshop during which the database was established and the current paper was initiated.

A.N. acknowledges support by the Center for International Scientific Studies and Collaboration (CISSC), Iran. C.M., I.B., I.G.-M and J.A.C. were funded by the Basque Government (IT936-16). D.V. carried out the research supported by a grant of the State Fund For Fundamental Research $\Phi 83/53427$. G.F. carried out the research in the frame of the MIUR initiative ‘Department of excellence’ (Law 232/2016). I.D. was supported by the Polish National Science Centre (grant DEC-2013/09/N/NZ8/03234). J.Do. was supported by the Czech Science Foundation (GA 17-19376S). M.J. was supported by grant by Slovak Academy of Sciences (VEGA 02/0095/19). W.U. acknowledges support from the Polish National Science Centre (grant 2017/27/B/NZ8/00316).

DATA AVAILABILITY STATEMENT

The GrassPlot data used in this study are collectively owned by the GrassPlot Consortium, consisting of several hundreds of contributors. The GrassPlot database (Dengler et al., 2018; <http://b.link/grassplot>) is continuously growing, while specific older versions used for published analyses, such as version 1.27 analysed in this study, are permanently stored. They can be requested from the first author conditional on agreement with the GrassPlot Bylaws (<https://bit.ly/2lX2Svu>), which ensure a fair involvement of data originators in emerging products. The data that support the findings of this study will become openly available in Dryad at <http://doi.org/doi:10.5061/dryad.4j75656> after a 3-yr embargo period. The R code used in this study is provided in Appendix S5–S7.

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BIOSKETCH

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GrassPlot (<http://b.link/grassplot>) is an international consortium, organized by the EDGG, including 260 members. Much of the GrassPlot database consists of data from annual EDGG Field Workshops. Currently, GrassPlot includes 186,749 plots from 181 databases.

Author contributions: J.D. conceived the idea, initiated data collection and led the writing. Most authors contributed data, while J.D. served as custodian and I.B. as database manager of the GrassPlot database. T.J.M. and M.J.S. conducted the analyses, T.C. and C.M. prepared the maps, I.D. and S.W. the conceptual figures. S.W. also conducted the simulation study to demonstrate the appropriateness of our statistical approach. All authors checked, improved and approved the manuscript.

SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section at the end of the article.

How to cite this article: Dengler J, Matthews TJ, Steinbauer MJ, et al. Species–area relationships in continuous vegetation: Evidence from Palaeoarctic grasslands. *J Biogeogr.* 2019;00:1–15. <https://doi.org/10.1111/jbi.13697>