# Trade-offs among plant reproductive traits determine interactions with floral visitors

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# 25 ABSTRACT

Plant life-history strategies are constrained by cost-benefit trade-offs that determine plant form and function. However, despite recent advances in the understanding of plant trait variation, little is known about plant reproductive trade-offs and how these 28 constrain life-history strategies and shape interactions with floral visitors. Here, we 29 investigate plant reproductive trade-offs and how these drive interactions with floral visitors using a dataset of 16 reproductive traits for 1,506 plant species. We found that 31 over half of all plant reproductive trait variation was explained by two independent 32 axes. Specifically, the first axis indicated the presence of a trade-off between flower 33 number and flower size, while the second axis indicated a pollinator dependency trade-off. In addition, plant reproductive trade-offs determined important differences in the interaction level among floral visitor guilds. Our study shows the main plant 36 reproductive trade-offs and their relevance to understand plant-pollinator interactions in a global context.

# 39 INTRODUCTION

Despite the astonishing diversity of floral structures among flowering plants (Barrett 2002; Schiestl & Johnson 2013) and their importance in shaping plant-pollinator interactions (Fenster et al. 2004; Dellinger 2020), a unified framework that describes the 42 major plant reproductive trade-offs is currently lacking (Roddy et al. 2021). In addition, 43 macroecological studies that investigate plant reproductive traits are scarce (Baude et al. 2016; Munoz et al. 2016; Grossenbacher et al. 2017; Moeller et al. 2017) and consequently, there is poor understanding of how reproductive traits drive interactions with floral 46 visitors at large scales (Sargent & Ackerly 2008; Rech et al. 2016; Salguero-Gómez et al. 47 2016; Rüger et al. 2018). Linking the plant's position in trait-space with the different pollinator groups could help to improve our understanding of plant-pollinator associations (Dehling et al. 2016). Further, there is increasing interest in understanding 50 drivers of plant-pollinator interactions using trait-based approaches (Fenster et al. 2004; Rosas-Guerrero et al. 2014) and trait-matching analyses (Stang et al. 2009; Bartomeus et 52 al. 2016). However, plant reproductive traits have been overlooked beyond highly specialised pollination systems (Dellinger 2020) and remains unclear how specific plant reproductive biology traits (e.g., mating or compatibility system) influence plant-pollinator associations (Tur et al. 2013; Devaux et al. 2014). Species can optimise their fitness through various life-history traits, yet trade-offs 57 among those traits constrain the range of potential strategies that a species can use. With the recent availability of large trait databases (e.g., TRY, Kattge et al. 2011; and COMPADRE, Salguero-Gómez et al. 2015), plant ecological strategies are being increasingly examined, and are facilitating the identification of global patterns and constraints 61 in plant form and function (Díaz et al. 2016; Salguero-Gómez et al. 2016; Bruelheide et al. 2018; Carmona et al. 2021). However, most studies have focused on vegetative 63 traits such as leaf (Wright et al. 2004), wood (Chave et al. 2009), or root (Laughlin et al. 2021) trade-offs with little or no attention given to reproductive traits (E-Vojtkó et al. 2020; Roddy et al. 2021) which are critical to plant life strategies that shape

interactions with pollinators and ultimately determine plant reproductive success. For instance, short lived versus perennial species tend to have low versus high levels of outcrossing, respectively, (Barrett 2003; Moeller *et al.* 2017) and outcrossing levels are positively correlated with flower size (Goodwillie *et al.* 2010). In addition, the presence of costly rewards (e.g., pollen or nectar) and showy flowers or floral displays can only be understood through consideration of plant species' reliance upon animal pollination (pollinator dependence) and its role in attracting pollinators (Ollerton *et al.* 2011; Rodger *et al.* 2021). However, it is still unknown to what extent these different reproductive compromises determine plant-pollinator interactions.

Several studies have identified links between plant traits and plant-pollinator network 76 properties (Bartomeus 2013; Olito & Fox 2015; Rowe et al. 2020). Moreover, plant traits can define species' network roles (e.g., specialists vs generalists; Lázaro et al. 2013; Tur 78 et al. 2013). For example, plant species that occupy reproductive trait space extremes are more likely to exhibit higher levels of specialisation and be more reliant on the trait-80 matching with pollinators (Junker et al. 2013; Coux et al. 2016). Morphological matching between plant and floral visitors often determines plant-pollinator interactions, and can thus strongly influence interaction network structure (Stang et al. 2009; Ibanez 2012). Remarkably, the combination of traits has shown to increase the predictive power of the network interactions (Eklöf et al. 2013). Moreover, the position in the trait space can determine plant and pollinator species' functional roles (Dehling et al. 2016). 86 Therefore, by considering the multidimensional reproductive trait space we could 87 progress our understanding of plant-pollinator associations. Importantly, we know little if those patterns generally studied at the community level are representative of wider macroecological scales.

Here, we aim to explore the potential trade-offs among reproductive traits and how these influence plant-pollinator interactions. First, we identify the major axes of reproductive trait variation and trade-offs that determine plant form and function. Second, we investigate how plant species' position in trait-space influence interactions with floral visitors. Finally, we investigate how both the main axes of trait variation, and

individual traits, influence plant species' roles within networks using a set of complementary interaction network metrics (i.e., interaction strength, normalized degree and specialization).

# **MATERIALS AND METHODS**

## 100 Plant-pollinator network studies

We selected 28 studies from 18 different countries that constituted a total of 64 plant-101 pollinator networks. These studies recorded plant-pollinator interactions in natural systems and were selected so that we had broad geographical representation. Although 103 these studies differ in sampling effort and methodology, all studies provided infor-104 mation about plant-pollinator interactions (weighted and non-weighted), which we 105 used to build a database of plant species that are likely to be animal pollinated. Many of these networks are freely available either as published studies (Olesen et al. 2007; 107 Fortuna et al. 2010; Carvalheiro et al. 2014) or available in online archives (e.g., The Web of Life, Fortuna et al. 2010; and Mangal, Poisot et al. 2016). In total, our network 109 dataset (see Table S1) constituted 60 weighted (interaction frequency) and 4 unweighted (presence/absence of the interaction) networks, each sampled at a unique location and 111 year, as well as eight meta-webs where interactions were pooled across several locations and multiple years. 113

#### 114 Taxonomy of plants and pollinators

All species names, genera, families and orders were retrieved and standardized from the taxonomy data sources NCBI (https://www.ncbi.nlm.nih.gov/taxonomy) for plants and ITIS (https://www.itis.gov/) for pollinators, using the R package *taxize* (Chamberlain *et al.* 2020) version 0.9.99. We filled the 'not found' searches manually using http://www.theplantlist.org/ and http://www.mobot.org/ for plants and http://www.catalogueoflife.org/ for floral visitors.

#### 121 Functional traits

We selected a total of 19 different functional traits that included both reproductive and 122 vegetative traits (see Table 1 and Supplementary Information). From these, 16 were 123 reproductive traits (13 floral and 3 reproductive biology traits) and were selected based 124 on their relevance to plant reproduction and data availability. Note that autonomous 125 selfing was recorded quantitatively (fruit set) and qualitatively because this trait was 126 rarely available in its numerical form. The 3 remaining traits, were vegetative traits that are commonly used to characterize the global spectrum of plant form and function and 128 represent the fast-slow continuum of trait variation (e.g., short-lived versus perennial species). For each plant species, we undertook an extensive literature and online search 130 across a wide range of resources (plant databases, online floras, books, journals and images). From a total of 30,120 cells (20 columns x 1506 species) we were able to fill 132 24,341 cells (80.8% of the dataset, see Fig. S1 for missing values information for each 133 trait). 134

## 135 Phylogenetic Distance

We calculated the phylogenetic distance between different plant species using the function *get\_tree* from the package *rtrees* (https://github.com/daijiang/rtrees), which downloads phylogenetic distances from the extended R implementation of the Open Tree of Life (Smith & Brown 2018; Jin & Qian 2019).

#### 140 Data Imputation

Trait missing values were imputed with the function *missForest* (Stekhoven & Bühlmann 2012) which allows imputation of data sets with continuous and categorical variables. We accounted for the phylogenetic distance among species on the imputation process by including the eigenvectors of a principal component analysis of the phylogenetic distance (PCoA) which has been shown to improve the performance of *missForest* (Penone *et al.* 2014). To extract the eigenvectors, we used the function *PVRdecomp* from the package *PVR* (Chamberlain *et al.* 2018) based on a previous conceptual

Table 1 Plant functional traits used in this study.

Quantitative traits		Categorical traits		
Type	Traits	Type	Traits	Categories
Vegetative	Plant height (m)	Vegetative	Lifepan	Short-lived Perennial
Floral	Flower width (mm)	Vegetative	Life form	Herb Shrub Tree
Floral	Flower length (mm)	Floral	Flower shape	Brush Campanulate Capitulum Open Papilionaceous Tube
Floral	Inflorescence width (mm)	Floral	Flower symmetry	Actinomorphic Zygomorphic
Floral	Style length (mm)	Floral	Nectar	Presence Absence
Floral	Ovules per flower	Reproductive biology	Compatibility system	Self-incompatible Part. self-compatible Self-compatible
Floral	Flowers per plant	Reproductive biology	Breeding system	Hermaphrodite Monoecious Dioecious
Floral	Nectar (μl)	Reproductive biology	Autonomous selfing	None Low Medium High
Floral	Nectar (mg)			
Floral	Nectar concentration (%)			
Floral	Pollen grains per flower			
Reproductive biology	Autonomous selfing (fruit set)			

framework that considers phylogenetic eigenvectors (Diniz-Filho et al. 2012). Although the variable of autonomous selfing had a high percentage of missing values (68%), 149 we were able to solve this by back transforming a complementary qualitative column 150 of autonomous selfing to numerical. The categories of 'none,' 'low,' 'medium' and 151 'high' were converted to representative percentages of each category 0%, 13%, 50.5% 152 and 88%, respectively. This reduced the percentage of missing values for this column 153 from 68% to 35% and allowed the imputation of this variable. However, we were 154 unable to include nectar and pollen traits on the imputation process because of the high 155 percentage of missing values (Fig. S1). Hence, the imputed dataset had 1,506 species, 156 seven categorical and eight numerical variables and 5.79% of missing values. Further, we conducted an additional imputation process on the subset of species with data for 158 pollen per flower and microliters of nectar. This subset comprised 755 species, 8.01% missing values and all traits but milligrams of nectar (~50% of missing values) were 160 included in the imputation process.

## Plant strategies

We explored the trade-offs between different quantitative plant functional traits with a 163 phylogenetically informed Principal Component Analysis (pPCA). We did not include 164 the quantitative variables of flower length and inflorescence width because they were 165 highly and moderately correlated to flower width respectively (Pearson's correlation 166 = 0.72, P < 0.01 and Pearson's correlation = 0.36, P < 0.01), and thus we avoided 167 overemphasizing flower size on the reproductive spectrum of trait variation. Although 168 qualitative traits were not included in the dimensionality reduction analysis, we also 169 investigated the association of the different qualitative traits with the main axes of trait variation. Prior to the analyses, we excluded outliers and standardized the data. 171 Due to the high sensitivity of dimensionality reduction to outliers, we excluded values 172 within the 2.5th–97.5th percentile range (Legendre & Legendre 2012), and thus our 173 final dataset had 1,236 species. Then, we log transformed the variables to reduce the influence of outliers and z-transformed (X=0, SD=1) so that all variables were within 175 the same numerical range. We performed the pPCA using the function phyl.pca from

the package *phytools* version 0.7-70 (Revell 2012) with the method lambda ( $\lambda$ ) that 177 calculates the phylogenetic correlation between 0 (phylogenetic independence) and 178 1 (shared evolutionary history) and we implemented the mode covariance because 179 values for each variables were on the same scale following transformation (Abdi & 180 Williams 2010). Moreover, to corroborate that our imputation of missing values did not 181 affect our results, we conducted a pPCA on the full dataset without missing values (Fig. 182 S2). We found little difference between the explained variance with the imputed dataset 183 (51.08%) and the dataset without missing values (52.87%). In addition, the loadings 184 on each principal component had a similar contribution and correlation patterns, with 185 the exception of plant height which showed slight variations between the imputed 186 and non-imputed dataset. Finally, we conducted an additional phylogenetic informed 187 principal component analysis for the subset of species with pollen and nectar quantity. For this, we included all quantitative traits considered in the main pPCA plus pollen 189 grains and microlitres of nectar per flower.

# Phylogenetic signal of traits

We calculated the phylogenetic signal of the different quantitative traits on the imputed dataset with the full set of species (N = 1,506) with the package *phytools* (Revell 2012) version 0.7-70 and we used Pagel's  $\lambda$  as a measurement of the phylogenetic signal. However, for pollen and nectar traits, phylogenetic signal was calculated only on the subset of species that had quantitative information for these traits (N = 755).

#### 197 Network analyses

Analyses were conducted on the subset of 60 weighted networks sampled in a unique flowering season and site, which included 556 plant and 1,126 pollinator species. These networks were analysed in their qualitative (presence-absence) and quantitative (interaction frequency) form. First, we analysed the binary version of these weighted networks with presence-absence information that assumes equal weight across interactions. Second, we analysed the untransformed weighted networks with interaction frequency that accounts for the intensity of the interaction. Although floral visitors

are not always pollinators and interaction frequency does not consider each pollinator 205 species efficiency (Ballantyne et al. 2015), interaction frequency can provide valuable 206 information of the contribution of floral visitors to pollination (Vázquez et al. 2005, 207 2012). In total, our network dataset (excluding meta-webs and non-weighted networks) 208 included 2,256 interactions of bees with plants, 1,768 non-syrphid-Diptera interactions, 209 845 syrphids interactions, 437 Lepidoptera interactions, 432 Coleoptera interactions 210 and 362 non-bee-Hymenoptera interactions. Sampling methods varied across networks 211 but this was accounted for in analyses by considering them in the random effects of the 212 modelling process. All analyses were conducted in R version 4.0.3. 213

#### 214 Visitation patterns

We used Bayesian modelling (see below for details) to explore the effect of floral 215 visitor groups and the main axes of trait variation (pPCA with imputed dataset) on both qualitative (presence/absence) and quantitative (visitation rate) floral interac-217 tions per plant species. For this, we divided floral visitors into six main guilds 218 that differ in life form, behaviour and are likely to play a similar ecological role: 219 (i) bees (Hymenoptera-Anthophila), (ii) non-bee-Hymenoptera (Hymenoptera-non-220 Anthophila), (iii) syrphids (Diptera-Syrphidae), (iv) non-syrphid-Diptera (Diptera-non-221 Syrphidae), (v) Lepidoptera and (vi) Coleoptera. Moreover, because the guild of bees 222 was the most represented group with 2,256 records and had the highest frequency of vis-223 its of all groups, we also explored the presence-absence of interaction and visitation rate 224 of the main bee families (Andrenidae, Apidae, Colletidae, Halictidae and Megachilidae) 225 on the trait space. In addition, we found that *Apis mellifera* was the floral visitor with 226 the largest proportion of records counted (7.55% of the total). This finding is consistent 227 with previous research showing that A. mellifera was the most frequent floral visitor 228 in a similar dataset of 80 plant-pollinator networks in natural ecosystems (Hung et al. 229 2018). Hence, to control for the effect of *A. mellifera* on the observed visitation patterns 230 of bees, we conducted an analogous analysis with presence-absence of interaction and 231 visitation rate excluding A. mellifera. We found that A. mellifera, was partly driving 232 some of the observed trends on PC1 (Supplementary Fig. S3). However, we did not 233

detect major differences on PC2 and PC3.

We implemented Bayesian generalized linear mixed models using the R package brms 235 (Bürkner 2017) (version 2.14.6). We modelled the frequency of visits as a function of the 236 main axes of plant trait variation and their interactions with floral visitor functional 237 groups (Visits ~ PC1 x FGs + PC2 x FGs + PC3 x FGs). Because we were interested in 238 possible differences in the visitation patterns among floral visitors groups to plants with 239 different strategies, we included interactions between the main axes of trait variation (PC1, PC2 and PC3) and the floral visitor guilds. In this model, we added a nested 241 random effect of networks nested within the study system to capture the variation in 242 networks among studies and within networks. Moreover, we included the phylogenetic 243 covariance matrix as a random factor due to the possible shared evolutionary histories of species and therefore lack of independence across them. We specified this model 245 with a zero inflated negative binomial distribution and weakly informative priors from the brms function. We run this model for 3,000 iterations and with previous 1,000 warm 247 up iterations. We set delta ( $\Delta$ ) to 0.99 to avoid divergent transitions and visualized the posterior predictive checks with the function *pp\_check* using the *bayesplot* package 249 (Gabry et al. 2019) (version 1.7.2).

#### Plant species functional roles

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We investigated whether different quantitative traits determined plant species func-252 tional roles using Bayesian modelling and regression trees. For this, we selected simple 253 and complementary species-level network metrics commonly applied in bipartite net-254 work studies (Dormann et al. 2008) with a straightforward ecological interpretation 255 relevant to our research goals. The different plant species-level metrics were: (i) sum of 256 visits per plant species; (ii) normalized degree, calculated as the number of links per 257 plant species divided by the total possible number of partners; and (iii) specialization 258 (d') (Blüthgen et al. 2006), which measures the deviation of an expected random choice 259 of the available interaction partners and ranges between 0 (maximum generalization) 260 and 1 (maximum specialization). Normalized degree and specialization were calculated

with the *specieslevel* function from the R package *bipartite* (Dormann *et al.* 2008) (version 2.15).

First, we modelled the distinct plant species metrics (sum of visits, normalized degree and plant specialization) as a function of the three main axes of trait variation (plant species level metric ~ PC1 + PC2 + PC3). For each response variable (i.e., each plant species level metric), we used different distribution families (zero inflated negative binomial for the sum of visits, weibull for normalized degree and zero one inflated beta for specialization). Finally, we used the same random factors, model settings and conducted the same posterior predictive checks for each model as detailed above in the 'visitation patterns section.'

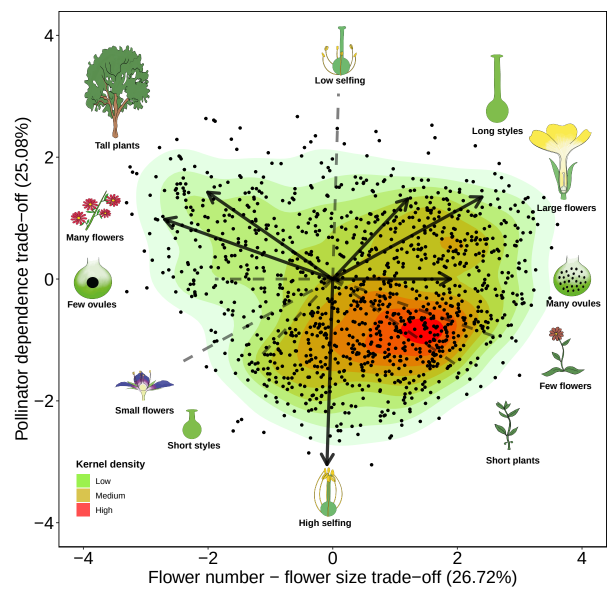
Second, to better understand these complex trait relationships, we used regression 272 trees. Regression trees are recursive algorithms which can detect complex relationships among predictors and allow identification of the relevance of specific trait combinations 274 on species functional roles. We focused exclusively on quantitative traits because 275 almost all categorical traits were statistically associated with the first two axes of 276 trait variation (Supplementary Table S2). We conducted this analysis using the rpart package (Therneau et al. n.d.) version 4.1-15 with method 'anova' with a minimum of 278 50 observations per terminal node and we used the *rpart.plot* package (Milborrow n.d.) 279 version 3.0.9 to plot the regression trees. We considered the species level indices as 280 response variables (interaction frequency, normalized degree and specialization) and 281 we performed one regression tree per metric using the different quantitative traits as 282 predictors. We calculated two regression trees per plant species-level metric, one for 283 the full set of species and another for the subset of species for which we had pollen 284 and nectar traits. We focused on regression trees that included floral rewards because 285 they consistently showed pollen and nectar traits as being the best for explaining the 286 different species-level metrics (see Supplementary Fig. S4). 287

# **RESULTS**

#### 89 Plant strategies

The phylogenetically informed principal component analysis (pPCA) captured by 290 the first two and three axes 51.8% and 70.97% of trait variation, respectively (Fig. 1 291 and Supplementary Fig. S5) and had a phylogenetic correlation ( $\lambda$ ) of 0.76. The first 292 principal component (PC1) represented 26.72% of the trait variation and indicated a 293 trade-off between flower number and flower size. We refer to this axis as the 'flower 294 number - flower size trade-off,' as already described in previous studies (Sargent et 295 al. 2007; Kettle et al. 2011). Hence, one end of the spectrum comprised species with 296 high investment in flower number and plant height but small flower size, short style 297 length and low ovule number. The other end of this spectrum comprised species that 298 were short in height and invested in large flowers, long styles, many ovules, but few 299 flowers. The main contributing traits to PC1 were plant height, flower number, ovule 300 number and flower size (loadings > | 0.5|; Supplementary Table S3) but style length also contributed moderately to PC1 (loading = -0.33). The second principal component (PC2) 302 represented 25.05% of the trait variation and indicated a trade-off between low and high 303 pollinator dependence. We refer to this axis as the 'pollinator dependence trade-off.' 304 The main driver of trait variation on PC2 was autonomous selfing (loading = 0.85) but the other traits (except ovule number) also made moderate contributions (loadings 306 from 0.27 to 0.4; Supplementary Table S3). We found that high pollinator dependence 307 was associated with larger and a higher number of flowers, greater plant height and 308 longer styles. In contrast, species with high levels of autonomous selfing tended to have 309 fewer and smaller flowers, had shorter styles and were shorter in height. Further, PC3 310 explained a considerable amount of trait variability (19.17%) and the main contributors to this axis were style length (loading = -0.66) and the degree of autonomous selfing 312 (loading = -0.51). The remaining traits, apart from ovule number, were moderately 313 correlated to changes on PC3 (loadings from -0.23 to -0.46; Supplementary Table S3). 314 Thus, because style length was correlated with all traits on PC3 and was the main driver

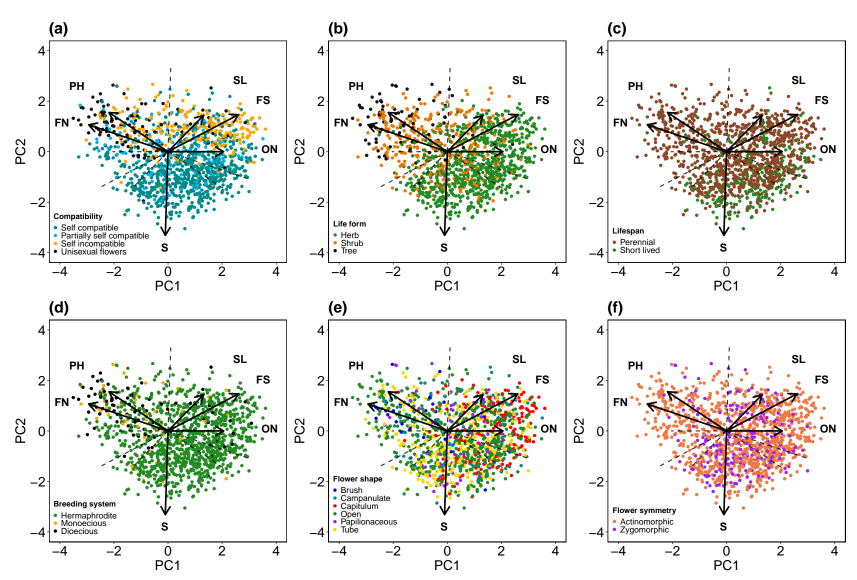
of trait variation, we refer to this axis as the 'style length trade-off.' Further, the pPCA 316 with the subset of species that had nectar and pollen quantity data showed that nectar 317 quantity (microlitres of nectar per flower) was positively associated with flower size, 318 style length and ovule number (PC1, 23.40%); and pollen quantity (pollen grains per 319 flower) was positively correlated with flower number and plant height and negatively 320 associated with autonomous selfing (PC2, 21.67%; Supplementary Fig. S6). This pPCA 321 explained similar variance with the first two principal components (45.07%) and similar 322 associations of traits despite some variability in the loadings (Supplementary Table S4). 323



**Figure 1** Phylogenetically informed principal component analysis (pPCA) of 1,236 plant species from 28 plant-pollinator network studies. The solid arrows indicate the direction of the different quantitative traits (flower number, plant height, style length, flower size, ovule number and level of autonomous selfing) across the two main axes of trait variation. The length of the arrows indicate the weight of the variables on each principal component and the dashed lines show the opposed direction of trait variation. The icons at both ends of arrows and dashed lines illustrate the extreme form of the trait continuum.

We found that most categorical traits were statistically associated with the first two axes of trait variation (Fig. 2 and Supplementary Table S2). Flower symmetry, which was only associated with PC2 (Sum of squares = 8.51, F-value = 14.72, P < 0.01), and nectar provision, which was independent of PC1 and PC2 (PC1: Sum of squares = 0.37, F-value

= 0.29, P = 0.59; PC2: Sum of squares = 0.83, F-value = 1.43, P = 0.23) showed lack of 328 statistical association. In addition, we found (with a Tukey test) statistical differences 329 between the different levels of categorical traits in the trait space (Supplementary Fig. 330 S7). Regarding self compatibility, we found larger differences on PC2 (i.e., species 331 with unisexual flowers that were self incompatible were statistically differentiated from 332 species with partial or full self compatibility; Supplementary Fig. S7a and Fig. S7b). Life 333 forms differed statistically across both axes of trait variation and followed a gradient 334 of larger life forms (trees and shrubs) with higher pollinator dependence to smaller 335 ones (herbs) with lower pollinator dependence (Supplementary Fig. S7c and Fig. S7d). 336 Consequently, lifespan also followed this gradient but perennial and short lived species 337 only differed statistically on PC2 (Supplementary Fig. S7e and Fig. S7f). Species with 338 unisexual flowers (monoecious and dioecious) were clustered on both extremes of the first two principal components and had the highest pollinator dependence and 340 highest number of flowers (Supplementary Fig. S7g and Fig. S7h). Moreover, we found that the campanulate and capitulum flower shapes were differentiated from tube, 342 papilionaceous, open and brush shapes in the trait space. The former morphologies 343 had larger flowers and greater pollinator dependence, while the latter had higher 344 flower number and greater autonomous selfing (Supplementary Fig. S7i and Fig. S7j). Regarding flower symmetry, zygomorphic flowers were associated with lower levels of 346 pollinator dependence, whereas actinomorphic flowers had higher levels of pollinator dependence (Supplementary Fig. S7k and Fig. S7l).



**Figure 2** Location of the different qualitative traits on the trait space. The panel is composed by the traits that showed statistical association with the first two axes of trait variation: compatibility system (a), life form (b), lifespan (c), breeding system (d), flower shape (e) and flower symmetry (f). The solid arrows indicate the direction of variation of the different quantitative traits showed in figure 1: flower number (FN), plant height (PH), style length (SL), flower size (FS), ovule number (ON) and the level of autonomous selfing (S).

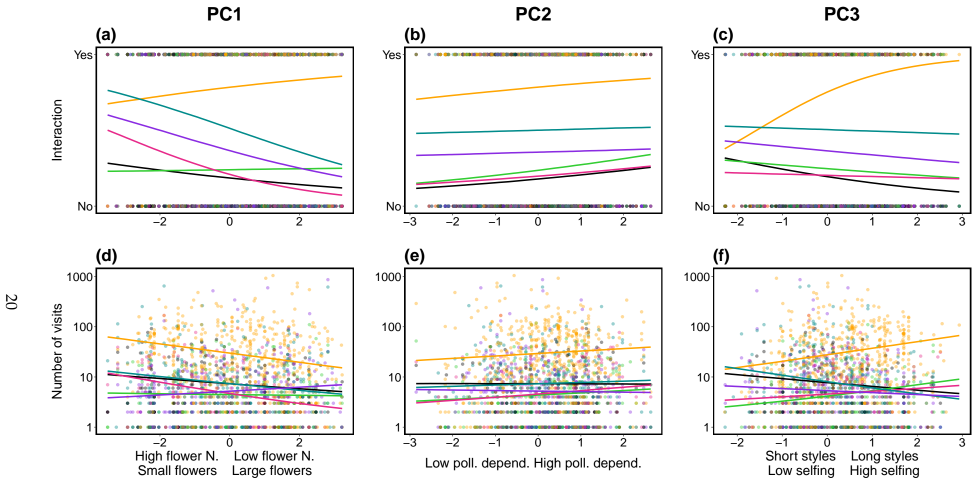
### Phylogenetic signal of traits

We found a strong phylogenetic signal (P < 0.01) in all quantitative traits (Supplementary Table S5). The traits that showed the highest phylogenetic signal were ovule number ( $\lambda = 1$ ), pollen grains per flower ( $\lambda = 1$ ) and plant height ( $\lambda = 0.96$ ), followed by flower length ( $\lambda = 0.75$ ), flower width ( $\lambda = 0.73$ ), number of flowers per plant ( $\lambda = 0.69$ ) and nectar concentration ( $\lambda = 0.65$ ). The traits that showed a moderate phylogenetic signal were inflorescence width ( $\lambda = 0.57$ ), style length ( $\lambda = 0.49$ ) and autonomous selfing ( $\lambda = 0.34$ ). Finally, microliters of nectar per flower showed the lowest phylogenetic signal of all traits ( $\lambda = 0.14$ ).

#### 358 Visitation patterns

The main axes of trait variation explained partly presence-absence interactions between 359 plant and floral visitors (conditional  $R^2 = 0.26$ ; marginal  $R^2 = 0.20$ ) but little of the 360 overall visitation rates (conditional  $R^2 = 0.31$ ; marginal  $R^2 = 0.06$ ). However, we 361 found relevant trends across the different floral visitor guilds on both presence-absence 362 and visitation interactions (Fig. 3). On the pollinator dependence trade-off, all floral 363 visitor guilds interacted more frequently with plant species with higher pollinator dependence (PC2; Fig. 3b and Fig. 3e). For presence-absence interactions we found that 365 all Diptera, Coleoptera and non-bee-Hymenoptera guilds interacted more frequently 366 with plants with high flower number and small flowers (flower number - flower size 367 trade-off, PC1; Fig. 3a) but bees and Lepidoptera interacted slightly more frequently 368 with plant species with low flower number but large flowers. For presence-absence 369 interactions on PC3 (style length trade-off; Fig. 3c), we found that bees interacted clearly more with plant species with long styles and high selfing and the rest of the 371 guilds interacted slightly more with plant species with short styles and low selfing. 372 In addition, all guilds other than Syrphids and Lepidoptera (i.e., all Hymenoptera, 373 non-syrphid-Diptera and Coleoptera) showed greater visitation rates on species with small numerous flowers (PC1; Fig. 3d). On the style length trade-off, bees, Lepidoptera 375 and non-bee-Hymenoptera showed greater visitation rates on plant species with larger

styles and higher levels of selfing; while syrphids, non-syrphid-Diptera and Coleoptera 377 showed higher visitation rates on species with shorter styles and lower selfing (Fig. 3f). 378 The additional model for both presence-absence of interactions (marginal  $R^2 = 0.29$ ; 379 conditional  $R^2 = 0.19$ ) and visitation rate (marginal  $R^2 = 0.30$ ; conditional  $R^2 = 0.03$ ) 380 for the most represented families of bees showed that the family Apidae was the main 381 driver of the observed patterns. The contrasting differences between presence-absence 382 and visitation rate for bees on PC1 (Fig. 3a and Fig. 3d) were driven by the family 383 Andrenidae, which interacted more frequently on presence-absence interactions with 384 plant species with low number of large flowers (Supplementary Fig. S8).



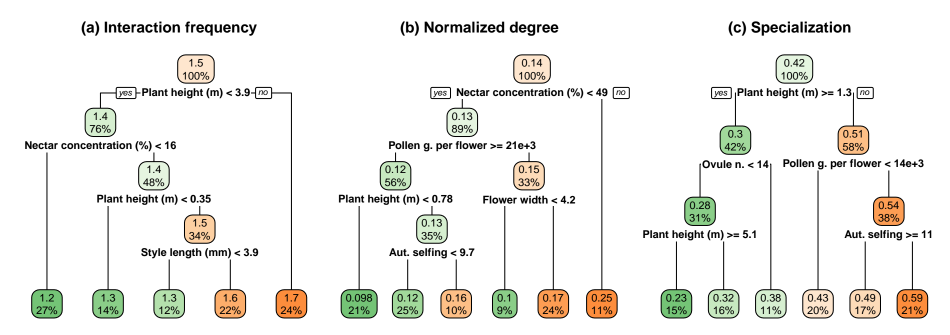
**Figure 3** Interaction (yes/no) and visitation rates across the three main axes of trait variation per floral visitor guild. Fitted posterior estimates of the presence/absence of interaction (a, b and c) and number of visits (d, e and f) made by the different floral visitors guilds in relation to PC1, PC2 and PC3. PC1 represents the flower number - flower size trade-off, PC2 represents the pollinator dependence trade-off and PC3, the style length trade-off. For visualization purposes, due to large differences between the visitation rates of bees and the rest of guilds, the number of visits was log-transformed (Y-axis of lower panel).

#### 86 Plant species functional roles

The variance of the different plant species-level network metrics was poorly explained 387 by the three main axes of trait variation (Supplementary Fig. S9; interaction frequency ~ 388 PCs, conditional  $R^2 = 0.11$ , marginal  $R^2 = 0.02$ ; normalized degree ~ PCs, conditional  $R^2$ 389 = 0.24, marginal  $R^2$  = 0.02; and, specialization ~ PCs, conditional  $R^2$  = 0.37, marginal  $R^2$ 390 = 0.03). Overall, the most notable trends were found on PC1 and PC3 for interaction 391 frequency and specialization. On the flower number - flower size trade-off (PC1), 392 interaction frequency was higher for plant species with more flowers but was lower 393 for plant species with larger flowers (Supplementary Fig. S9a). On PC1, specialization showed the opposite trend (Supplementary Fig. S9g). On the style length trade-395 off (PC3), interaction frequency was lower for plants with shorter styles and lower autonomous selfing and higher for species with longer styles and higher autonomous 397 selfing (Supplementary Fig. S9c). Again, specialization showed the opposite trend to 398 interaction frequency (Supplementary Fig. S9i). 399

When we further investigated the combination of traits that drive plant network roles, 400 we found that the regression tree for visitation frequency was best explained by plant 401 height, nectar concentration and style length (Fig. 4a). Specifically, species taller than 402 3.9m had the highest interaction frequency, while species that were shorter than 3.9m 403 and had a nectar concentration lower than 16% had the lowest interaction frequency. 404 Normalized degree was best explained by nectar concentration, pollen grains per 405 flower, plant height, flower width and autonomous selfing (Fig. 4b). Species with a 406 nectar concentration over 49% had the highest levels of normalized degree, whereas 407 species with nectar concentration lower than 49%, more than 21,000 pollen grains 408 per flower and height less than 0.78m had the lowest normalized degree. Finally, 409 specialization was best explained by plant height, ovule number, pollen grains per flower and autonomous selfing (Fig. 4c). Overall, plant species with the highest 411 specialization were shorter than 1.3m, had more than 14,000 pollen grains per flower and autonomously self-pollinated less than 11% of their fruits. In contrast, species 413 taller or equal than 5.1m and with lower than 14 ovules per flower had the lowest

specialization values.



**Figure 4** Contribution of traits in plant's network roles. Regression tree analysis of interaction frequency (log-transformed), normalized degree and specialization for the subset of species with quantitative data for pollen and nectar traits. The superior value inside the node indicates the mean value of the different species-level metric and the lower value, the percentage of species that are considered in each node. Thus, the top node has the mean value of the named trait for the 100% of species. Each node has a yes/no question and when the condition is fulfilled, the branch turns to the 'yes' direction and when not, to the 'no' direction. This rationale is followed in all the regression trees as indicated in the first branch division of the topmost node of each tree.

# DISCUSSION

This study demonstrates that plant species exhibit clear trade-offs among their vegetative and reproductive traits and that these trade-offs determine interactions with floral 418 visitors. These trade-offs are differentiated along three axes of trait variation: (i) flower 419 number - flower size, (ii) pollinator dependence and (iii) style length. These reproduc-420 tive trade-offs helped partly explain the presence of floral visitor interactions, but not 421 their visitation rates. However, floral visitor guilds formed distinct relationships with 422 the main axes of trait variation. Moreover, we found that the plant species functional 423 roles within pollination networks were best explained by plant size and floral reward 424 related traits.

Over half of all plant trait variation was captured by the flower number - flower size 426 and pollinator dependence trade-offs. Trait variation on these two axes was associated with the 'fast-slow continuum' in plant (Salguero-Gómez et al. 2016) and animal (Healy 428 et al. 2019) life-history strategies, as indicated by the different floral and reproductive biology traits associated with plant height, life form and lifespan. The 'slow' part of this 430 continuum (i.e., tall trees and shrubs) included plant species with many flowers, few ovules, higher pollinator dependence, frequent occurrence of self-incompatibility and 432 more complex breeding systems (e.g., monoecious and dioecious species). In contrast, plant species that employed the 'fast' strategy (i.e., short herbs), had fewer flowers, 434 more ovules, frequent occurrence of self-compatibility and lower pollinator dependence. Further, on the first two axes of trait variation, we found additional support for the 436 previously described positive association between higher outcrossing rate and larger 437 floral display (Goodwillie et al. 2010). The positive correlation between larger floral 438 display and higher pollinator dependence in our dataset further confirmed this trend (see Supplementary Fig. S10).

Despite the low predictive power of the main trait variation axes for broad-level interaction patterns (presence-absence of interactions and visitation rate), we found

changes in the interaction patterns among and within floral visitor guilds across these 443 axes that suggest plant life-history strategies influence plant-pollinator interactions. For 444 example, all floral visitor guilds visited plant species with higher pollinator dependence 445 more frequently, and high pollinator dependence was associated with large floral 446 displays and greater pollen quantities (Fig. 1 and Supplementary Fig. S6). This trend 447 is consistent with previous studies that show plant species with higher reproductive investment tend to be visited by pollinators more frequently (Hegland & Totland 2005; 449 Lázaro et al. 2013; Kaiser-Bunbury et al. 2014). In regard to the flower number -450 flower size and style length trade-offs, different pollinator guilds showed contrasting 451 visitation rates across the continuum of trait variation, which could be associated with 452 different pollination syndromes at a macroecological scale. For instance, bees and 453 syrphid flies were clearly associated with opposing life-strategies on PC1 and PC3 (Fig. 3) suggesting possible niche partitioning (Palmer et al. 2003; Phillips et al. 2020) 455 between these two guilds. However, despite floral rewards not being included in the 456 main analysis because there was insufficient data available, floral reward related traits 457 were among the best at characterising species functional roles (Fig. 4). More detailed 458 exploration of reproductive trade-offs in conjunction with floral rewards is needed to 459 help elucidate plant-pollinator associations. In any case, it is worth noting that other local factors such as species relative abundances, surely explain part of the observed 461 variability (Vázquez et al. 2007; Encinas-Viso et al. 2012; Bartomeus et al. 2016) that 462 reproductive trade-offs do not. 463

To conclude, we provide the first description of plant reproductive trade-offs using a large global dataset of plant traits. We identified the major reproductive strategies of flowering plants and how these strategies influence interactions with different floral visitor guilds. Although the explained variation that we found in the first two axes is lower than previous studies of vegetative traits (Díaz et al. 2016; Carmona et al. 2021) it is consistent with the largest and most recent study that has characterised plant life strategies with vegetative and reproductive traits (Salguero-Gómez et al. 2016). Future work needs to integrate the reproductive compromises that we have identified with

vegetative and physiological trade-offs to create a more comprehensive spectrum of
plant trait variation. Further, the varying level of phylogenetic signal among traits
deserves further attention to understand evolutionary changes on mating and flower
morphology in response to pollinators (Gervasi & Schiestl 2017; Mackin *et al.* 2021).
Finally, including plant-pollinator networks from unrepresented areas of the world and
a more complete description of plant reproductive trade-offs is essential for a better
understanding of the global patterns in plant-pollinator interactions.

# References

- Abdi, H. & Williams, L.J. (2010). Principal component analysis. *WIREs Comp. Stats.*, 2, 433–459.
- Ballantyne, G., Baldock, K.C.R. & Willmer, P.G. (2015). Constructing more informative plant–pollinator networks: Visitation and pollen deposition networks in a heathland plant community. *Proc. Royal Soc. B*, 282, 20151130.
- Barrett, S.C.H. (2002). The evolution of plant sexual diversity. *Nat. Rev. Genet.*, 3, 274–284.
- Barrett, S.C.H. (2003). Mating strategies in flowering plants: The outcrossing-selfing paradigm and beyond. *Philos. Trans. R. Soc. Lond. B Biol. Sci.*, 358, 991–1004.
- Bartomeus, I. (2013). Understanding Linkage Rules in Plant-Pollinator Networks by
   Using Hierarchical Models That Incorporate Pollinator Detectability and Plant Traits.
   PLoS One, 8, e69200.
- Bartomeus, I., Gravel, D., Tylianakis, J.M., Aizen, M.A., Dickie, I.A. & Bernard-Verdier,
   M. (2016). A common framework for identifying linkage rules across different types
   of interactions. *Funct. Ecol.*, 30, 1894–1903.
- Baude, M., Kunin, W.E., Boatman, N.D., Conyers, S., Davies, N., Gillespie, M.A., *et al.*(2016). Historical nectar assessment reveals the fall and rise of floral resources in
  britain. *Nature*, 530, 85–88.
- Blüthgen, N., Menzel, F. & Blüthgen, N. (2006). Measuring specialization in species interaction networks. *BMC Ecol.*, 6, 9.

- Bruelheide, H., Dengler, J., Purschke, O., Lenoir, J., Jiménez-Alfaro, B., Hennekens, S.M.,
- et al. (2018). Global trait–environment relationships of plant communities. Nat. Ecol.
- 502 Evol., 2, 1906–1917.
- <sup>503</sup> Bürkner, P.-C. (2017). Brms: An R package for Bayesian multilevel models using Stan. J.
- 504 Stat. Softw., 80, 1–28.
- Carmona, C.P., Tamme, R., Pärtel, M., Bello, F. de, Brosse, S., Capdevila, P., et al. (2021).
- Erosion of global functional diversity across the tree of life. *Sci. Adv.*, 7, eabf2675.
- <sup>507</sup> Carvalheiro, L.G., Biesmeijer, J.C., Benadi, G., Fründ, J., Stang, M., Bartomeus, I., et
- al. (2014). The potential for indirect effects between co-flowering plants via shared
- pollinators depends on resource abundance, accessibility and relatedness. *Ecol. Lett.*,
- 17, 1389–1399.
- 511 Chamberlain, S., Szoecs, E., Foster, Z., Arendsee, Z., Boettiger, C., Ram, K.,
- et al. (2018). Package 'PVR.' R package version 0.3. At https://CRAN.r-
- project.org/package=PVR.
- <sup>514</sup> Chamberlain, S., Szoecs, E., Foster, Z., Arendsee, Z., Boettiger, C., Ram, K., et al. (2020).
- Taxize: Taxonomic information from around the web. R package version 0.9.99. At
- https://CRAN.r-project.org/package=taxize.
- Chave, J., Coomes, D., Jansen, S., Lewis, S.L., Swenson, N.G. & Zanne, A.E. (2009).
- Towards a worldwide wood economics spectrum. *Ecol. Lett.*, 12, 351–366.
- coux, C., Rader, R., Bartomeus, I. & Tylianakis, J.M. (2016). Linking species functional
- roles to their network roles. *Ecol. Lett.*, 19, 762–770.
- Dehling, D.M., Jordano, P., Schaefer, H.M., Böhning-Gaese, K. & Schleuning, M. (2016).
- Morphology predicts species' functional roles and their degree of specialization in
- plant–frugivore interactions. *Proc. Royal Soc. B*, 283, 20152444.
- Dellinger, A.S. (2020). Pollination syndromes in the 21st century: Where do we stand
- and where may we go? *New Phytol.*, 228, 1193–1213.
- Devaux, C., Lepers, C. & Porcher, E. (2014). Constraints imposed by pollinator be-
- haviour on the ecology and evolution of plant mating systems. J. Evol. Biol., 27,
- <sub>528</sub> 1413–1430.

- Díaz, S., Kattge, J., Cornelissen, J.H.C., Wright, I.J., Lavorel, S., Dray, S., *et al.* (2016). The global spectrum of plant form and function. *Nature*, 529, 167–171.
- Diniz-Filho, J.A.F., Bini, L.M., Rangel, T.F., Morales-Castilla, I., Olalla-Tárraga, M.Á.,
- Rodríguez, M.Á., et al. (2012). On the selection of phylogenetic eigenvectors for
- ecological analyses. *Ecography*, 35, 239–249.
- Dormann, C.F., Gruber, B. & Fründ, J. (2008). Introducing the bipartite package:
- Analysing ecological networks. *R News*, 8/2.
- Eklöf, A., Jacob, U., Kopp, J., Bosch, J., Castro-Urgal, R., Chacoff, N.P., et al. (2013). The
- dimensionality of ecological networks. *Ecol. Lett.*, 16, 577–583.
- Encinas-Viso, F., Revilla, T.A. & Etienne, R.S. (2012). Phenology drives mutualistic
- network structure and diversity. *Ecol. Lett.*, 15, 198–208.
- E-Vojtkó, A., Bello, F. de, Durka, W., Kühn, I. & Götzenberger, L. (2020). The neglected
- importance of floral traits in trait-based plant community assembly. J. Veg. Sci., 31,
- 529–539.
- Fenster, C.B., Armbruster, W.S., Wilson, P., Dudash, M.R. & Thomson, J.D. (2004).
- Pollination Syndromes and Floral Specialization. Annu. Rev. Ecol. Evol. Syst., 35,
- 545 375–403.
- Fortuna, M.A., Stouffer, D.B., Olesen, J.M., Jordano, P., Mouillot, D., Krasnov, B.R., et
- al. (2010). Nestedness versus modularity in ecological networks: Two sides of the
- same coin? *J. Anim. Ecol.*, 79, 811–817.
- Gabry, J., Simpson, D., Vehtari, A., Betancourt, M. & Gelman, A. (2019). Visualization in
- Bayesian workflow. J. R. Stat. Soc. Ser. A Stat. Soc., 182, 389–402.
- Gervasi, D.D.L. & Schiestl, F.P. (2017). Real-time divergent evolution in plants driven
- by pollinators. *Nat. Commun.*, 8, 14691.
- Goodwillie, C., Sargent, R.D., Eckert, C.G., Elle, E., Geber, M.A., Johnston, M.O., et al.
- 554 (2010). Correlated evolution of mating system and floral display traits in flowering
- plants and its implications for the distribution of mating system variation. New
- 556 *Phytol.*, 185, 311–321.
- Grossenbacher, D.L., Brandvain, Y., Auld, J.R., Burd, M., Cheptou, P.-O., Conner, J.K.,

- et al. (2017). Self-compatibility is over-represented on islands. New Phytol., 215, 469–478.
- Healy, K., Ezard, T.H.G., Jones, O.R., Salguero-Gómez, R. & Buckley, Y.M. (2019).
- Animal life history is shaped by the pace of life and the distribution of age-specific
- mortality and reproduction. *Nat. Ecol. Evol.*, 3, 1217–1224.
- Hegland, S.J. & Totland, Ø. (2005). Relationships between species' floral traits and pollinator visitation in a temperate grassland. *Oecologia*, 145, 586–594.
- <sup>565</sup> Hung, K.-L.J., Kingston, J.M., Albrecht, M., Holway, D.A. & Kohn, J.R. (2018). The
- worldwide importance of honey bees as pollinators in natural habitats. *Proc. Royal*
- *Soc. B*, 285, 20172140.
- <sup>568</sup> Ibanez, S. (2012). Optimizing size thresholds in a plant-pollinator interaction web:
- Towards a mechanistic understanding of ecological networks. *Oecologia*, 170, 233–
- 570 242.
- Jin, Y. & Qian, H. (2019). V.PhyloMaker: An R package that can generate very large phylogenies for vascular plants. *Ecography*, 42, 1353–1359.
- Junker, R.R., Blüthgen, N., Brehm, T., Binkenstein, J., Paulus, J., Schaefer, H.M., et al.
- 574 (2013). Specialization on traits as basis for the niche-breadth of flower visitors and
- as structuring mechanism of ecological networks. *Funct. Ecol.*, 27, 329–341.
- Kaiser-Bunbury, C.N., Vázquez, D.P., Stang, M. & Ghazoul, J. (2014). Determinants of
- the microstructure of plant–pollinator networks. *Ecology*, 95, 3314–3324.
- Kattge, J., Díaz, S., Lavorel, S., Prentice, I.C., Leadley, P., Bönisch, G., et al. (2011). TRY –
- a global database of plant traits. *Glob. Chang. Biol.*, 17, 2905–2935.
- Kettle, C.J., Maycock, C.R., Ghazoul, J., Hollingsworth, P.M., Khoo, E., Sukri, R.S.H., et
- al. (2011). Ecological Implications of a Flower Size/Number Trade-Off in Tropical
- Forest Trees. *PLoS One*, 6, e16111.
- Laughlin, D.C., Mommer, L., Sabatini, F.M., Bruelheide, H., Kuyper, T.W., McCormack,
- M.L., et al. (2021). Root traits explain plant species distributions along climatic
- gradients yet challenge the nature of ecological trade-offs. *Nat. Ecol. Evol.*, 1–12.
- Lázaro, A., Jakobsson, A. & Totland, Ø. (2013). How do pollinator visitation rate

- and seed set relate to species' floral traits and community context? *Oecologia*, 173, 881–893.
- Legendre, P. & Legendre, L. (2012). Numerical ecology. 2nd edn. Elsevier, Amsterdam.
- Mackin, C.R., Peña, J.F., Blanco, M.A., Balfour, N.J. & Castellanos, M.C. (2021). Rapid
- evolution of a floral trait following acquisition of novel pollinators. *J. Ecol.*, 109,
- 2234–2246.
- Milborrow, S. (n.d.). Rpart. Plot: Plot rpart Models. An Enhanced Version of plot. Rpart.
- R package version 3.0.9 (2015). At https://CRAN.r-project.org/package=rpart.plot.
- Moeller, D.A., Runquist, R.D.B., Moe, A.M., Geber, M.A., Goodwillie, C., Cheptou,
- P.-O., et al. (2017). Global biogeography of mating system variation in seed plants.
- 597 Ecol. Lett., 20, 375–384.
- Munoz, F., Violle, C. & Cheptou, P.-O. (2016). CSR ecological strategies and plant
- mating systems: Outcrossing increases with competitiveness but stress-tolerance is
- related to mixed mating. *Oikos*, 125, 1296–1303.
- Olesen, J.M., Bascompte, J., Dupont, Y.L. & Jordano, P. (2007). The modularity of
- pollination networks. *PNAS*, 104, 19891–19896.
- 603 Olito, C. & Fox, J.W. (2015). Species traits and abundances predict metrics of
- plant–pollinator network structure, but not pairwise interactions. Oikos, 124,
- 428–436.
- Ollerton, J., Winfree, R. & Tarrant, S. (2011). How many flowering plants are pollinated
- by animals? *Oikos*, 120, 321–326.
- Palmer, T.M., Stanton, M.L. & Young, T.P. (2003). Competition and coexistence: Explor-
- ing mechanisms that restrict and maintain diversity within mutualist guilds. *Am.*
- Nat., 162, S63–S79.
- Penone, C., Davidson, A.D., Shoemaker, K.T., Marco, M.D., Rondinini, C., Brooks,
- T.M., et al. (2014). Imputation of missing data in life-history trait datasets: Which
- approach performs the best? *Methods Ecol. Evol.*, 5, 961–970.
- Phillips, R.D., Peakall, R., van der Niet, T. & Johnson, S.D. (2020). Niche perspectives
- on plant–pollinator interactions. *Trends Plant Sci.*, 25, 779–793.

- Poisot, T., Baiser, B., Dunne, J.A., Kéfi, S., Massol, F., Mouquet, N., *et al.* (2016). Mangal

   making ecological network analysis simple. *Ecography*, 39, 384–390.
- Rech, A.R., Dalsgaard, B., Sandel, B., Sonne, J., Svenning, J.-C., Holmes, N., et al. (2016).
- The macroecology of animal versus wind pollination: Ecological factors are more
- important than historical climate stability. *Plant Ecol. Divers.*, 9, 253–262.
- Revell, L.J. (2012). Phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, 3, 217–223.
- Roddy, A.B., Martínez-Perez, C., Teixido, A.L., Cornelissen, T.G., Olson, M.E., Oliveira,
- R.S., et al. (2021). Towards the flower economics spectrum. New Phytol., 229, 665–672.
- Rodger, J.G., Bennett, J.M., Razanajatovo, M., Knight, T.M., Kleunen, M. van, Ashman,
- T.-L., et al. (2021). Widespread vulnerability of flowering plant seed production to
- pollinator declines. *Sci. Adv.*, 7, eabd3524.
- Rosas-Guerrero, V., Aguilar, R., Martén-Rodríguez, S., Ashworth, L., Lopezaraiza-Mikel,
- M., Bastida, J.M., et al. (2014). A quantitative review of pollination syndromes: Do
- floral traits predict effective pollinators? *Ecol. Lett.*, 17, 388–400.
- Rowe, L., Gibson, D., Bahlai, C.A., Gibbs, J., Landis, D.A. & Isaacs, R. (2020). Flower
- traits associated with the visitation patterns of bees. *Oecologia*, 193, 511–522.
- Rüger, N., Comita, L.S., Condit, R., Purves, D., Rosenbaum, B., Visser, M.D., et al. (2018).
- Beyond the fast–slow continuum: Demographic dimensions structuring a tropical
- tree community. *Ecol. Lett.*, 21, 1075–1084.
- 636 Salguero-Gómez, R., Jones, O.R., Archer, C.R., Buckley, Y.M., Che-Castaldo, J., Caswell,
- H., et al. (2015). The compadre Plant Matrix Database: An open online repository
- for plant demography. *J. Ecol.*, 103, 202–218.
- 639 Salguero-Gómez, R., Jones, O.R., Jongejans, E., Blomberg, S.P., Hodgson, D.J., Mbeau-
- Ache, C., et al. (2016). Fast-slow continuum and reproductive strategies structure
- plant life-history variation worldwide. *Proc. Natl. Acad. Sci. U.S.A.*, 113, 230–235.
- Sargent, R.D. & Ackerly, D.D. (2008). Plant-pollinator interactions and the assembly of
- plant communities. *Trends Ecol. Evol.*, 23, 123–130.
- Sargent, R.D., Goodwillie, C., Kalisz, S. & Ree, R.H. (2007). Phylogenetic evidence for a

- flower size and number trade-off. *Am. J. Bot.*, 94, 2059–2062.
- Schiestl, F.P. & Johnson, S.D. (2013). Pollinator-mediated evolution of floral signals.
- 647 Trends Ecol. Evol., 28, 307–315.
- <sup>648</sup> Smith, S.A. & Brown, J.W. (2018). Constructing a broadly inclusive seed plant phylogeny.
- *Am. J. Bot.*, 105, 302–314.
- 650 Stang, M., Klinkhamer, P.G.L., Waser, N.M., Stang, I. & van der Meijden, E. (2009).
- Size-specific interaction patterns and size matching in a plant–pollinator interaction
- web. Ann. Bot., 103, 1459–1469.
- 653 Stekhoven, D.J. & Bühlmann, P. (2012). MissForest—non-parametric missing value
- imputation for mixed-type data. *Bioinformatics*, 28, 112–118.
- Therneau, T., Atkinson, B., Ripley, B. & Ripley, M.B. (n.d.). Package 'rpart.' R package
- version 4.1-15 (2015). At https://CRAN.r-project.org/package=rpart.
- Tur, C., Castro-Urgal, R. & Traveset, A. (2013). Linking Plant Specialization to Depen-
- dence in Interactions for Seed Set in Pollination Networks. *PLoS One*, 8, e78294.
- Vázquez, D.P., Lomáscolo, S.B., Maldonado, M.B., Chacoff, N.P., Dorado, J., Stevani,
- 660 E.L., et al. (2012). The strength of plant–pollinator interactions. Ecology, 93, 719–725.
- Vázquez, D.P., Melián, C.J., Williams, N.M., Blüthgen, N., Krasnov, B.R. & Poulin,
- R. (2007). Species abundance and asymmetric interaction strength in ecological
- networks. Oikos, 116, 1120–1127.
- Vázquez, D.P., Morris, W.F. & Jordano, P. (2005). Interaction frequency as a surrogate
- for the total effect of animal mutualists on plants. *Ecol. Lett.*, 8, 1088–1094.
- Wright, I.J., Reich, P.B., Westoby, M., Ackerly, D.D., Baruch, Z., Bongers, F., et al. (2004).
- The worldwide leaf economics spectrum. *Nature*, 428, 821–827.

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