The trade-offs of sharing pollinators: pollination service is determined by the community context

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Abstract

A fundamental feature of pollination systems is the indirect facilitation and competition that arises when plants species share pollinators. When plants share pollinators, the pollination 17 service can be influenced. This depends not only on how many partners plant species share, 18 but also by multiple intertwined factors like the plant species' abundance, visitation, or traits. 19 These factors inherently operate at the community level. However, most of our understanding of how these factors may affect the pollination service is based on systems of up to a handful 21 of species. By examining comprehensive empirical data in eleven natural communities, we 22 show here that the pollination service is—surprisingly—only partially influenced by the 23 number of shared pollinators. Instead, the factors that most influence the pollination service (abundance and visit effectiveness) also introduce a trade-off between the absolute amount of 25 conspecific pollen received and the amount relative to heterospecific pollen. Importantly, the ways plants appear to balance these trade-offs depend strongly on the community context, as most species showed flexibility in the strategy they used to cope with competition for pollination.

- **Keywords:** apparent competition, apparent facilitation, competition for pollinators, interspecific pollen transfer, pollen deposition, pollination costs and benefits, pollination network,
- pollination niche, and pollinator sharing

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33 Introduction

Animal pollination plays a disproportionally important role in food production and maintenance of global biodiversity (Klein et al. 2007, Bascompte and Jordano 2007, Ollerton et al. 35 2011). At a pairwise level, the mutually beneficial relationship between plants and pollinators underpins the pollination service. At a community level, sometimes involving hundreds of 37 species, both plant and pollinator species are connected in a myriad of indirect connections when pollination partners are shared. These indirect connections can dramatically alter the quality of the pollination service that plants receive because they determine how conspecific and heterospecific pollen is transferred across the community (Morales and Traveset 2008). Generally speaking, there is a trade-off between the benefits gained from a species maximising its number of partners and the costs of sharing them with other plant species (Waser 1978). However, due to the large number of factors that operate at the community level, we generally do not know how sharing pollinators affects the pollination service beyond systems with more than a handful of species. Here we investigate how pollinator sharing affects pollen transfer in natural communities and how it compares to other factors known to play a role in community dynamics like abundance, traits, and visitation patterns. There are two main mechanisms through which sharing pollinators can affect plant fertilisation (Morales and Traveset 2008). The first is by changes in intraspecific pollen transfer. Changes in intraspecific pollen transfer happen, for example, when plants with more attractive flowers might reduce the number of visits to those less attractive neighbouring plants, and hence reduce the amount of conspecific pollen deposited by animals (Yang et al. 2011). The second is via interspecific pollen transfer. In that case, even receiving a visit might not necessarily translate into fertilisation (Campbell and Motten 1985) because a focal plant might receive heterospecific pollen or because pollen from the focal plant might be lost to different species. Naturally, the precise effects on female or male plant fitness of conspecific and heterospecific pollen deposition depend on the species involved (Arceo-Gómez and Ashman 2016) and are

⁵⁹ unknown for many plant species.

Even for species well adapted to pollinator sharing, receiving foreign pollen on stigmas or losing pollen to foreign stigmas is neutral (at best). Indeed, there is substantial evidence supporting the idea that heterospecific pollen deposition can be detrimental to seed production and plant fitness (Ashman and Arceo-Gómez 2013, Arceo-Gómez and Ashman 2016). All else being equal, provided pollen is viable and compatible (de Jong et al. 1992, Dafni and Firmage 2000, Ramsey and Vaughton 2000), the higher the quantity of conspecific pollen and its purity (relative to heterospecific pollen), the better the pollination service received by the focal plant. As such, measuring conspecific and heterospecific pollen deposition provides a good indication of the potential levels of facilitation and competition a plant population might experience.

By definition, intra- and interspecific pollen transfer occur at the community scale. However, with few exceptions (Aizen and Rovere 2010, Tur et al. 2016), most of what we know about pollen transfer and its relationship with key ecological factors are based on studies with two plant species. That is partly so because the factors that determine the patterns of pollen deposition at the community scale are tightly intertwined, operate simultaneously, and may lead to emergent phenomena not observed at smaller scales (Flanagan et al. 2011). For instance, recent empirical evidence suggests that plants with flowering traits that are "original" relative to others in the community generally have fewer interaction partners (Coux et al. 2016).

This evidence aligns with the notion that a species that interacts with few species does so strongly with each of them whereas a species that interacts with a large number of species does so comparatively weakly (Bascompte et al. 2006, Vázquez et al. 2007, Thébault and Fontaine 2008). If evolutionary specialisation occurs by changing traits to focus on fewer but better partners (Caruso 2000), we should expect a reduction of competition for pollinators in species with "original" traits and an increase of competition in species with a large number of

interaction partners (Gibson et al. 2012, Carvalheiro et al. 2014). Alternatively, it might also
be the case that abundance (for example, in terms of flower or pollen counts) is the dominant
force driving pollen transfer (Seifan et al. 2014). Abundant plant species might experience a
dilution of available pollinators (Feinsinger 1987, Feldman et al. 2004) but might also receive
more effective visits by capitalising on a larger share of both visits and the pollen carried by
pollinators (Stavert et al. 2019). In this case, a potential reduction in the absolute amount of
conspecific pollen received could be compensated by an increase in the amount of conspecific
pollen relative to heterospecific pollen. Altogether, it is clear that these ecological factors can
indeed shape pollen deposition at the community level. However, we still do not understand
their relative importance and the trade-offs that might exist between them.

Here, we investigate pollen-deposition dynamics at the community scale using empirical data 95 from eleven plant-pollinator communities in the Argentinian Pampas. First, we investigate the relative contribution that four ecological factors make to the pollination service. Specifically, 97 we hypothesise that there are trade-offs on how these factors affect the quantity and purity of conspecific pollen deposition. While quantity and purity should decrease for plants that share many pollination partners, other factors like the plant's functional originality, its relative 100 floral abundance, and its visitation patterns should have the potential to compensate for 101 this decrease Second, we examine how these four factors that might affect pollen deposition 102 can change across communities where species are present. Because these factors may affect 103 the pollination service in contrasting ways, and a species role is relative to other species in 104 the community, we predict that species present in multiple communities should be flexible 105 enough to compete for pollinators under different community contexts.

$_{\scriptscriptstyle{107}}$ Methods

We collected data from eleven co-flowering plant communities and their pollinators in three locations in the Argentinian Pampas. In each location, we sampled two restored and two agricultural fragments, except in one located in the Flooding Pampas, where we were only able to sample one restored fragment due to the lack of available sites.

Factors affecting quantity and purity of pollination service

Our first objective was to investigate the relative contribution that different ecological factors 113 have on pollen deposition. Generally speaking, we expect that any factor that increases the 114 amount of conspecific pollen deposited in stigmas, both in quantity and purity relative to 115 heterospecific pollen, also has a positive effect on the pollination service. Specifically, we 116 investigated the effect of (i) a plant's number of shared pollinator species, (ii) a plant's 117 abundance relative to the rest of the community, (iii) the mean visit potential—a metric that 118 combines the amount and type of pollen carried by floral visitors and the number of visits it 119 receives from them, and (iv) the plant's functional originality (Laliberté and Legendre 2010). See Data Analysis section below for more details on these four factors. 121

122 Data collection

In each of the studied communities, we quantified pollen deposition in a subset of plant species between December 2010 and February 2011. This subset comprised between three and nine common insect-pollinated (entomophilous) plant species that were flowering during the sampling period. Based on data from previous years (Marrero et al. 2014), we chose plant species such that they cover a wide range on a specialization-generalization gradient as well as a wide range of abundances. In each of the selected plants, we removed all flowers leaving only buds that were expected to go into florescence on the next day. Two days after flowering,

we collected all remaining flowers and counted the number of conspecific and heterospecific pollen grains in their pistils. More details can be found in Marrero et al. (2016).

To obtain the number of shared pollinators for each species, we collected data to construct 132 qualitative and quantitative pollination networks. Qualitative networks were constructed 133 based on ten-hour observations of floral visits in each fragment. Quantitative networks were 134 constructed using two 50 m randomly located transects in each fragment. We counted and 135 collected all floral visitors found in a 2 m wide strip while walking at a pace of 10 m per 136 minute (Memmott 1999, Marrero et al. 2014). We visited the transects each month between 137 November 2010 and March 2011. To obtain floral abundance, we counted all units of floral 138 attraction found during an independent sampling of the same transects used to construct 139 the quantitative visitation networks. To estimate visit potential, we need to construct pollen 140 transfer networks in addition to the visitation networks. To do this, we examined the pollen 141 loads present on the floral visitors collected (Marrero et al. 2017). When the pollen count on 142 an individual animal was estimated to be less than 2,000 grains, we identified every grain 143 to the species level when possible and to pollen complexes when it was not. When the 144 pollen count was above 2,000 grains, we classified approximately 50% of pollen and total 145 pollen counts were extrapolated (Bosch et al. 2009). Finally, we also recorded morphological 146 traits that relate to plant type (herb, shrub, climber), life cycle (annual, perennial), flower 147 colouration, phenology, and whether the species is native in the study region. More details 148 can be found in Marrero et al. (2014 and 2017). 149

Data analysis

To investigate the impact of ecological factors on pollination services, we used two sets of linear mixed models (LMM) with bootstrap resampling. The response variables for these model sets were the number of conspecific and heterospecific pollen grains deposited per stigma in flowers open to animal-mediated pollination. We used LMMs in which pollen loads

were log-transformed because these models offered a better fit than equivalent GLMMs with Poisson (or quasi-Poisson) error structure. Models were fitted using the R package nlme 3.1-131 (Pinheiro et al. 2018).

Because the amount of deposited pollen can vary widely across species, and potentially also across communities, we evaluated two possible structures for the random effects: one that includes a random intercept for plant species, and one that treats measures from species across different communities independently. We selected the best random structure by comparing the median Akaike Information Criterion for small samples (AICc).

As fixed predictors in the models, we included the four ecological factors described above. Specifically, we calculated the number of shared pollinator species for each plant species by pooling data from the qualitative and quantitative pollination networks. To calculate the plants' relative floral abundance in their community, we aggregated floral counts for each species. We then calculated the mean visit potential of pollinator species i to plant species j as

$$o_{ij} = \frac{v_{ij}}{v_i} \frac{p_{ji}}{p_j},$$

where v_{ij} is the observed number of visits by i to j, p_{ji} is the number of pollen grains from j attached to i, v_i is the total number of visits performed by i, and p_j is the total number of grains carried by j. We log-transformed the number of shared pollinators, floral abundance, and visit potential before including them in the model.

Finally, functional originality is defined as the distance of a species from the community trait average—the centroid of functional space of the community (Laliberté and Legendre 2010, Coux et al. 2016). To include phenological variation, we treated floral abundance in each of the survey months (November to March) as a "trait" in this analysis. To account for the non-independence of floral counts and weight all traits equally, we assigned a weight of 1/5 to these abundances (one for each month). We scaled all traits before calculating the centroid of the functional space and calculated the species-specific functional coordinates using the R

package FD 1.0-12 (Laliberté et al. 2014). Finally, we scaled all four factors to have a zero mean and unit variance.

To estimate the coefficients, perform model selection, and quantify the associated uncertainty, 182 we used a combination of multi-model inference and bootstrap resampling with 99 replicates. 183 Using bootstrap replicates allow us to better understand the uncertainties associated with our 184 estimations. First, we performed model selection using AICc and determined the likelihood 185 of each candidate model (a particular combination of predictors) by calculating the median 186 \triangle AICc (relative to the most likely model) for each bootstrap sample. As we wanted model 187 coefficients from more likely candidate models to carry more weight in our results, we sampled 188 the coefficients for our factors proportionally to the likelihood of their candidate model. 189 Finally, we used these distributions of the model coefficients to estimate their mean impact 190 on the pollination service (in terms of quantity and purity of conspecific pollen deposition). 191

Flexibility of plant strategies

Our second objective was to tease apart whether and how these factors that might affect 193 pollen deposition might change across communities species are present. If community context 194 plays a relatively small role, or species are inflexible in regards to these factors, we would 195 expect plants of the same species to use similar "strategies" across different communities. 196 Alternatively, if the community plays a significant role and plant species are flexible, we 197 should be able to observe differences in the strategy a plant species uses across communities. 198 To test this, we first used a principal component analysis (PCA) of the four ecological 199 factors (number of shared pollinators, floral abundance, visit potential, and trait originality). 200 We scaled factors across the whole study to ensure that the PCA space does not change 201 according to the species present in each community. We define a species' strategy in a community as its coordinates in PCA space. For each species that was present in two or more 203 communities, we then calculated (i) the median distance between the points that correspond

to the strategy a species uses in different communities and (ii) the area of the convex hull
defined by these points in the first two principal components (only for species present in
three or more communities). We then compared the magnitude of these two metrics to those
obtained with 99 Monte Carlo randomizations in which we replaced the strategy of the focal
plant species by that of another randomly selected species in the dataset.

$_{\scriptscriptstyle{210}}$ Results

Factors affecting quantity and purity of pollination service

We first examined the potential roles played in pollen deposition by four ecological factors 212 (number of shared pollinators, abundance, mean visit potential, and functional originality). 213 We found that our models of pollen deposition had high explanatory power (the coefficient of 214 determination R² ranged between 0.76 and 0.93) although a large portion of the explanatory 215 power came from the random effects (Table S3). As determined by AICc, the random structure best supported by the data was the one that fit a separate intercept for each species in each community (as opposed to a common intercept for each species irrespective of the 218 community to which they belong). This structure was best for both the models of conspecific 219 and heterospecific pollen (Table S4). 220

Of the four factors we considered, we found that a plant's mean visit potential and relative floral abundance were the most important at predicting pollen deposition in plant stigmas (Fig. 1a). Surprisingly, the number of shared pollinators was comparatively unimportant, particularly for models of heterospecific pollen deposition, as it was only ever included in models with relatively large AICc values (Table S5).

[Figure 1 about here.]

We found that the relationship between each of the ecological factors and pollen deposition

was similar for both conspecific and heterospecific pollen. That is, strategies that were associated with an increase in conspecific pollen deposition were also associated with an 229 increase in heterospecific pollen deposition. Specifically, the plants' mean visit potential had 230 a positive effect on pollen deposition (Fig. 1b). However, the effect size was slightly larger for 231 heterospecific than for conspecific pollen. This larger effect indicates that, although there is 232 a positive association between visit potential and the quantity of pollen deposition, there is a 233 negative relationship with its purity (Fig. 1c). In contrast, a plants' relative floral abundance 234 negatively affected its deposition quantity, but the mean difference between the coefficients in 235 the models indicates a positive association with purity (Fig. 1c). The third most important 236 factor, functional originality, had a positive, although comparatively smaller, association 237 with both the quantity and purity. Finally, the number of shared pollinators had negative 238 and neutral associations with conspecific and heterospecific pollen, respectively, but these 230 impacts were small when compared to the other factors. Although the ecological factors were 240 positively correlated (Fig. S2), the collinearity between predictors did not qualitatively affect our findings (Fig. S3). 242

Flexibility of plant strategies

We used a PCA of the ecological factors—species matrix to investigate whether plants' strategies towards pollen deposition is similar across communities or whether they are flexible and therefore a reflection of the community context. The first two PCA components explained 75% of the total variance (Fig. 2a). The first component was dominated by visit potential and relative abundance while the second component was dominated by the number of shared pollinators and the plant's functional originality. When we locate the species that were sampled in more than one community in the first two PCA components (Fig. 2b), we observe that the positions of any given species do not tend to be close to each other. Indeed, when we measured the median distance between the plants' coordinates, we found that it was only

significantly smaller than that of randomisations for only two of the twelve species analysed (Fig. 2c).

[Figure 2 about here.]

Discussion

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Our results suggest that community context plays a central role in determining the pollen deposition dynamics and ultimately the net cost or benefit of sharing pollinators. First, we 258 found that multiple ecological factors can modulate the quality of the pollination service: 250 however, conspecific and heterospecific pollen deposition are tightly coupled and this creates 260 a clear trade-off between the quantity and purity of pollination (Thomson et al. 2019). 261 Second, we found that the way these factors shape pollen deposition for a species could be 262 dramatically different across communities. For instance, while a plant species in a particular 263 community could show high levels of pollinator sharing and relatively low trait differentiation, 264 the same species in another community can have relatively high trait differentiation and 265 low levels of pollinator sharing. Our findings highlight that trade-offs can at least partially explain the coexistence of facilitative and competitive effects of animal-mediated pollination 267 in the pollination service. 268

The trade-offs involved in attaining high-quality pollination service (and more broadly between facilitation and competition) are likely to arise when plants simultaneously maximise the deposition of conspecific pollen and minimise that of heterospecific pollen. In the short term, being a specialist and sharing no pollinators might reduce competition (Muchhala et al. 2010) and hence be preferable. This may be due to both costs to male fitness (Morales and Traveset 2008, Muchhala and Thomson 2012), and also, as we show here, because sharing pollinators reduces both the quantity and purity of the conspecific pollen deposited. However, over long periods of time, there could be a risk associated with a specialist plant having few pollinators

(Ricketts 2004). To ensure long-term survival, it is thus likely that plants also need to balance this risk with the costs of sharing pollinators (Aizen et al. 2012). One possible solution is to share pollinators and have original traits—as we show that trait originality is generally 279 beneficial to pollen deposition and it is commonly thought that species that are further from 280 others in trait space benefit from reduced competition. Yet, there are two possible caveats to 281 this strategy that highlight the interrelatedness of the ecological factors. First, in a mutualism 282 context, it is also possible that trait originality could come at the cost of being less 'apparent' 283 to pollinators (Reverté et al. 2016). Second, the negative relationship between originality and 284 generalism (Carvalheiro et al. 2014) has been shown to depend on plant abundance (Coux et 285 al. 2016), with generalist species being able to have original traits only when they are also 286 abundant enough to provide a valuable reward to make visiting worthwhile to pollinators. 287 Visit potential (high pollen and visits) and floral abundance, which were the most important 288 predictors of pollen deposition here, introduced an even more explicit trade-off between 289 gaining conspecific pollen and avoiding heterospecific pollen. Receiving high visitation 290 increases conspecific pollen deposition but increases heterospecific pollen deposition to a 291 greater extent—even when the visitors are likely to carry a high proportion of conspecific 292 pollen (Fang and Huang 2016). Contrastingly, being abundant reduces the amount of 293 conspecific pollen deposited and simultaneously reduces heterospecific pollen at a faster rate. 294 Our results corroborate the importance that two-species studies have ascribed to visitation 295 and abundance (Feldman et al. 2004, Muñoz and Cavieres 2008, Morales and Traveset 2008), 296 but they also suggest that (because visitation, pollen production and abundance are usually 297 correlated; Sargent and Otto 2006) balancing the pros and cons of sharing pollinators at the community level is not trivial. The fact that no species can easily outcompete others for pollination might be partially responsible for the diversity of plant-pollinator communities (Benadi and Pauw 2018). 301

We observed, as expected, that the effects of pollen deposition can vary widely among

species. For instance, the fitness of some plant species can be hurt even by low amounts
of heterospecific pollen, while the fitness of others can instead be limited by the amount of
conspecific pollen (Campbell and Motten 1985, Arceo-Gómez et al. 2019). Alternatively,
plant species can also differ substantially in the extent to which self- vs. outcross-pollen differ
in their value for fertilization. The difference can be particularly relevant for species that
are not self-fertile or those in which self-fertilization is rarely effective due to a temporary
separation in the maturation of the sexes (dichogamy).

Importantly, we show here that the balances between costs and benefits are determined not 310 only by species identity but also by the community to which plants belong. Specifically, most 311 plant species appear to be flexible enough to adopt markedly different "strategies" in different 312 communities. From an evolutionary perspective, our results suggest that selection for a 313 particular strategy might say something about the community in which a species has typically 314 inhabited during its evolutionary history. Furthermore, from a more applied perspective, 315 flowering plants are sometimes introduced to attract pollinators on other nearby plants. On 316 the one hand, our results suggest that plants that increase the relative originality of natives 317 (e.g. through distinct phenology) might have positive effects (Gibson et al. 2012). On the 318 other, because different strategies can lead to different outcomes across communities, our 319 results also highlight the difficulties involved in predicting whether the introduced plant 320 species will facilitate or compete with neighbours (Bartomeus et al. 2008). Other factors that 321 we were unable to measure (e.g. pollinator behaviour and densities or the spatial context) 322 have also been shown to play a role in the outcome of animal-mediated pollination (Cariveau 323 and Norton 2009, Flanagan et al. 2011, Ye et al. 2014, Thomson et al. 2019). Nevertheless, our results indicate that the strategies a plant might use to successfully minimise competition for pollination (or maximise facilitation) must be determined relative to other species in the community, rather than an absolute property of the species itself. 327

Overall, using empirical data on pollen deposition, we show at the community level that

sharing pollinators has a smaller effect on pollen deposition than what we expected based on experimental studies with a handful species. Other factors that underpin community dynamics 330 (abundance, traits, visitation) also influence patterns of pollination quantity and purity. The 331 interrelatedness of these factors, and the flexibility of species to position themselves across 332 them, means that their contributions to the quality of the pollination service cannot be 333 understood in isolation. All of the factors we analysed involve substantial trade-offs in pollen 334 deposition in the short and likely also in the long term. These trade-offs emphasise the 335 inherently competitive nature of pollination. However, many of the widely used theoretical 336 models of plant-pollinator communities do not account for the adverse effects of sharing 337 pollinators (but see Rohr et al. 2014 and similar). We therefore propose that achieving a 338 better understanding of species coexistence and how pollination supports plant biodiversity 339 will require seeing them as both mutualistic and competitive communities (Johnson and 340 Bronstein 2019).

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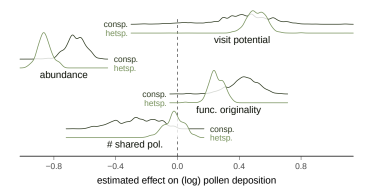
List of Figures

1	Effect of ecological factors on the pollination convice (a) The plant's visit	
1	Effect of ecological factors on the pollination service. (a) The plant's visit	
	potential and relative floral abundance are the most important factors deter-	
	mining the deposition of conspecific and heterospecific pollen. Meanwhile,	
	the number of shared pollinators was generally less important. The graph	
	shows the relative importance calculated as the sum of the Akaike weights of	
	the candidate models that included the selected factor. (b) The association	
	between ecological factors and heterospecific pollen (lighter line) tended to	
	align with their association with conspecific pollen (darker line). Visit potential	
	and functional originality had a positive association with pollen deposition,	
	while floral abundance and the number of shared pollinators had a negative	
	association. The plot shows the distribution of the effects (across 99 bootstrap	
	replicates) of the four ecological factors for conspecific and heterospecific pollen.	
	(c) The end result of these associations is that only the plants' functional orig-	
	inality has a positive impact on both the quantity and purity of conspecific	
	pollen deposition (relative to heterospecific pollen). The plot shows the model	
	averaged mean effect (± SE of 99 bootstrap replicates)	23
2	The flexibility of plant strategies. (a) The two first components explain a large	20
	proportion of the total variance. (b) When plants that were sampled in more	
	than one community are plotted in terms of these two components, we observe	
	that their points—which represent the strategy (the particular combination of	
	ecological factors) of that species in its community—do not seem to be grouped	
	by plant species. (c) This was confirmed using Monte Carlo randomizations	
	of the median distance between strategies of a plant species. Only two of the	
	examined species had strategies that were less flexible than would be expected	
	at random	24

(a) relative variable importance



(b) distribution of effects based on 100 bootstrap repllicates



(c) mean effect on pollination service

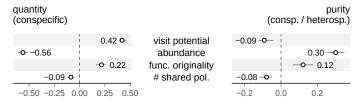
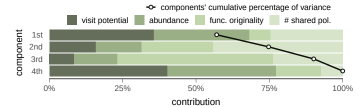


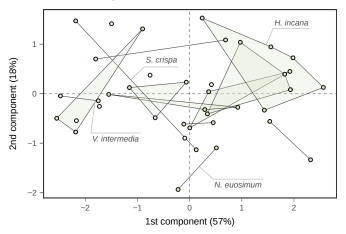
Figure 1

(a) components' variance and variable contributions principal component analysis of ecological variables



(b) plant realised niches in PCA space

convex hulls of species niches across communities



(c) flexibility of plant's strategies median distance between plant niches vs. randomisations

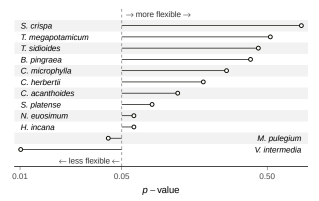


Figure 2