# The pollination trade-off

E. Fernando Cagua<sup>1</sup>, Hugo J. Marrero<sup>2</sup>, Jason M. Tylianakis<sup>1</sup>, Daniel B. Stouffer<sup>1</sup>

- $^{-1}$  Centre for Integrative Ecology, School of Biological Sciences, University of Canterbury, Private Bag 4800.
- : Christchurch 8041, New Zealand
- $^{\circ}$  Centro de Recursos Naturales Renovables de las Zonas Semiáridas, CONICET, Camino de la Carrindanga
- : Km. 7, 8000 Bahía Blanca, Argentina
- ${\it s-Author~for~correspondence:~E.~Fernando~Cagua~(efc 29\, {\it quelive.ac.nz}) Centre~for~Integrative~Ecology,}\\$
- School of Biological Sciences, University of Canterbury, Private Bag 4800, Christchurch 8140, New Zealand
- $\tau$  -Manuscript length: 35,176 characters, approx. 5,160 words (excludes front-matter, abstract, awknoledge-
- $z_{\parallel}$  ments, references, and figure/table legends); 26 min, reading time.
- Abstract length: 1 words.

Of course the figures are Methods + Results are very dense. Can anything be moved to supplemental?

in the annoying words of Nate 1. Michelle

■ Abstract

n TBC.

E Keywords: X. Y. Z

### . Introduction

 $_{\rm H}$  Animal pollination systems play a disproportionally important role in food production and maintenance of global biodiversity (Bascompte & Jordano 2007; Klein et al. 2007; Ollerton et al. 2011). Alongside the direct  $_{16}$  mutualisms between plants and their pollinators, biodiversity can be supported as a result of inter-species  $^{17}$  "facilitation"—the positive feedback loops that exist between plant species that share pollinators or pollinators that share plants (Moeller 2004; Ghazoul 2006; Molina-Montenegro et al. 2008). This inter-species facilitation <sub>19</sub> is able to promote species coexistence by offsetting the effects of direct resource competition (Stachowicz 2001).  $_{\odot}$  . Indeed, studies at the core of our current theoretical knowledge of mutualisms predict that the maximum  $_{11}$  number of coexisting species is achieved when the number of shared mutualistic partners is maximised in a = community (Bastolla et al. 2009).

23 However, in natural communities, plants rarely share all available pollinators. One explanation is that plants  $_{24}$  are effectively competing for mutualistic partners because there is a trade-off between the benefits gained 25 from maximising the number of instandistic partners and the costs of sharing them with other plant species zs. (Waser 1978). It has been clear for a long time (both experimentally and theoretically: Robertson 1895; Lewis 2 1961; Levin & Anderson 1970) that when competition for pollination is strong enough, it can be detrimental  $\kappa$  to plants fitness. In fact, it can be sufficient to drive ecological differentiation (Caruso 2000; for example by driving the evolution of both traits and reproductive strategies; Mitchell et al. 2009). It is less clear, however, 10 how exactly ecological differentiation, in turn, affect the trade-off between facilitation and competition that is involved in the pollimation service.

22 There are two main mechanisms through which competition for pollinators can affect plant fertilization Morales & Traveset 2008). The first is intraspecific pollen transfer. This happens, for example, when plants  $^{22}$  with distinctively attractive flowers might reduce the number of visits—and hence the amount of conspecific $_{35}$  -pollcu deposited by animals—to those less attractive neighbouring plants (Yang et al. 2011). The second is  $_{\infty}$  –intraspecific pollen transfer and occurs when plants share pollination partners. In that case, even receiving  $_{\scriptscriptstyle \mathcal{P}}$  a visit might not necessarily translate into fertilization (Campbell & Motten 1985) because a focal plant  $_{8}$  -might receive heterospecific pollen or pollen from the focal plant might be lost to different species. Generally aking, the higher the amount of conspecific pollen (both in absolute terms and relative to heterospecific 45 pollen) the better the pollination service received by the focal plant. These two mechanisms of competition.  $_{41}$  by definition, occur at the community scale. However, with few exceptions (Rathcke 1988; Lopezaraiza-Mikel 4: et al. 2007; Hegland et al. 2009; Aizen & Rovere 2010; Tur et al. 2016), most of what we know about the z deposition of conspecific and heterospecific pollen—and its relationship to competition vs. facilitation in

3

 $\omega$  pollination systems—is based on studies of two plant species.

z At the community scale, the factors that determine the patterns of conspecific and heterospecific pollen 45 deposition are tightly intertwined, operate simultaneously, and may lead to emergent phenomena not observed v at smaller scales (Flanagan cl 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 is aligned 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  $\alpha$  does so comparatively weakly (TODO; ref needed). If evolutionary specialisation occurs by changing traits to 52 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 in species with a large number of interaction partners (Gibson  $_{50}$  et al. 2012). Alternatively, it might also be the case that density (for example in terms of flower or pollen s counts) is the dominant force driving pollen transfer (Seilan et al. 2014). Abundant plant species might s experience a dilution of available pollinators (Feinsinger 1987) but might also receive more effective visits by <sup>57</sup> capitalising on a larger share of both visits and pollinator "real state". In this case, a potential reduction  $\kappa$  in the absolute amount of conspecific pollen received could be compensated by an increase in the amount verlative to heterospecific pollen. Altogether, it is clear that some ecological factors can influence pollen  $\infty$  deposition and that these factors, in the specific context of pollination, could be used to roughly define a species realised niche in their community. However, the intertwined nature of these factors makes it hard

large process.

| Process to determine whether they favour or hinder the pollination service at the community level Moreover, their  $_{60}$  relative importance has not yet been empirically evaluated in natural plant communities.

 $_{8}$  Here, we investigate competition for pollination at the community scale using empirical data from eleven plant-pollinator communities in the Argentinian Pampas. First, we estimate the extent to which the net outcome of plants sharing pollinators is facilitation or competition. If facilitation is the primary outcome we could expect two things. On the one hand, we could expect an overall positive relationship between the amount of conspecific pollen deposited in stigmas and the amount of heterospecific pollen (Tur  $et\ al.$ 2016). The higher the deposition of conspecific pollen relative to heterospecific, the larger the slope of that relationship. On the other, we could also expect a larger amount of conspecific pollen deposited in stigmas when flowers are open to animal pollinators than is deposited due to self-pollination. Second, we investigate the relative contribution that four ecological factors have on the pollination service. Specifically we hypothesize that competition (measured as absolute and relative conspecific pollen deposition) should increase for plants that share many pollination partners. However, we also hypothesize that other factors like the plant's functional originality, its relative abundance, and the visit effectiveness should have the potential

neleco

In first obj ratio of conspecific heterosp is response but in object it is predictor (competition)



to compensate for this increase in competition. Third, we examine how community context influences the
way plants use these factors to minimise competition. Although some plant species might be widespread
across multiple locations, competition for pollinators occurs at the community level and hence we predict
the competition for pollination niche to be influenced by the local context. If plant strategy is not flexible
across communities (TODO; ref needed), we could expect niches to be more similar within species than across
species in the study. On the contrary, if a species' strategy is flexible, we should expect plants to be able
to occupy different niches in each community such that competition could be minimised regardless of the
context.

#### 4 Methods

We collected data from eleven co-flowering plant communities and their pollinators in three locations in the
Argentinian Pampas. These locations are distributed across a precipitation gradient with a humid temperate
climate in the eastern Flooding Pampas and becoming drier as one moves inland and westward towards
the Pampean Grasslands. This precipitation gradient generated a natural gradient of network architecture
which allowed us to examine the generality of our findings. In each location, we sampled two restored and
two agricultural fragments, except in the Flooding Pampas, where we were only able to sample one restored
fragment due to the lack of available sites

### 22 Competition vs. facilitation

Our first objective was to identify whether the outcome of plants sharing pollinators at the community level is facilitation or competition. The first line of evidence can be found via the relationship between heterospecific and conspecific pollen deposited in plant stigmas (Tur et al. 2016). This is because heterospecific pollen can be seen as an unavoidable cost that plants must pay for receiving conspecific pollen from shared pollinators. A negative relationship (when shared pollinators increase heterospecific pollen loads and decrease conspecific loads) may indicate that competition for pollination is strong. This negative relationship could arise, for example, when another, more dominant species, monopolizes a limited number of pollinators (Lopezaraiza-Mikel et al. 2007; Mitchell et al. 2009), when heterospecific pollen deposited in the stigma mereferes with deposition of conspecific pollen (Runquist 2012), or when conspecific pollen is lost to heterospecific stigmas (Mitchell et al. 2009; Muchhala & Thomson 2012). In contrast, a positive relationship is what would be observed when facilitation is the dominant outcome. For example, when the abundance of shared pollinators

is positively influenced by floral resources, these positive effects ripple through the mutualistic network in a chain of positive feedback loops (Liao et al. 2011).

The second line of evidence can be obtained by examining the difference between the amount of conspecific pollen deposited in flower stigmas with and without animal-mediated pollination. Self-pollination can be favourable to plant reproduction when competition for pollination is strong because it provides insurance against poor pollination service (Kalisz & Vogler 2003). If competition for pollination is sufficiently strong, then the amount of conspecific pollen deposited in flower stigmas when animal pollination is prevented could be higher than when havel visits are allowed. In contrast, the opposite pattern may indicate that plants benefit from having pollen delivered directly to the stigma by animals. We did not specifically test the effect of pollen viability or the extent to which self-pollen vs. pollen from a conspecific may differ in their value for fertilization which can be important factors (TODO: ref needed). However, it is reasonable to expect that the difference between outcross and self-pollen may be generally relevant and particularly so 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) (TODO: ref needed).

#### 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 (see Figure 1a). We chose
plant species such that they cover a wide range on a specialization-generalization gradient as well as a wide
range of abundances. Briefler in each of the selected plants, we removed all flowers leaving only buds that
were expected to go into florescence on the next day. Approximately a quarter of these buds were bagged
to prevent animal pollination. Two days after inflorescence, we counted the pollen grains in the remaining
flowers' pistils, classified them between conspecific and heterospecific pollen, and calculated the number of
pollen grains per stigma in each flower. More details about the study sites, data collection, and laboratory
protocols can be found in Marrero et al. (2016).

#### Data analysis

To evaluate the relationship between heterospecific and conspecific pollen, we used a generalised linear mixed model (GLMM) with a Poisson error distribution (Tur *et al.* 2016). In this model, the number conspecific pollen grains per stigma was the response variable and the number of heterospecific grains the predictor (the

sowiest you test this favidure

number of stigmas sampled from the flower was added as an offset term). The slope and intercept of this 22 relationship were allowed to differ between plant species within each community, which were modelled as  $_{13}$  a random effect. Models were fitted using the function glmer from the R package lme4 1.1-19 (Bates  $c\bar{c}$ 2015). To compare the conspecific pollen deposited with and without animal-mediated pollination, we 107 followed the same approach as above. In this case, however, the predictor was the treatment (whether the flower was bagged or unbagged) instead of the number of heterospecific grains

### Ecological constraints the pollination service

Our second objective was to investigate the relative contribution of different factors that describe the plant  $_{131}$  competition for pollination niche to the pollination service. Specifically, we investigated the effect of (i) a plant's number of shared pollinator species. (ii) a plants abundance relative to the rest of the community. iii) the mean visit effectiveness—a metric that combines the share of pollen that a plant species is able to place on each of their floral visitors and the number of visits it receives from them (see Data Analysis section below for more details), and (iv) the plants functional originality (add ref). I don't really get

#### 146 Data collection

To obtain the number of shared pollinators for each species, we collected data to construct qualitative and quantitative pollination networks. Qualitative networks were constructed based on two-hour observations of floral visits in each fragment. Quantitative networks were constructed using two 50 m randomly located transects in each fragment. We counted and collected all floral visitors found in a 2 m wide strip while walking at a pace of 10 m per minute (Memmott 1999; Marrero et al. 2014). We visited the transects each onth between November 2010 and March 2011. To obtain plant abundance, we counted all units of floral attraction found during an independent sampling of the transects used to construct the quantitative visitation networks. To estimate visit effectiveness we need to construct pollen transfer networks in addition to the visitation networks. To construct the transfer networks, we examined the pollen loads present on the floral visitors collected (Marrero et al. 2017). When the pollen count on an individual animal was estimated to be less than 2,000 grains, we identified every grain to the species level when possible and to pollen complexes when it was not. When the pollen count was above 2.000 grains, we classified approx. 50% of pollen and total pollen counts were extrapolated (Bosch et al. 2009). Finally, we also recorded morphological traits that relate to plant type (herb, shrub, climber), life cycle (annual, perennial), flower colouration, phenology and whether the species is native in the study region. We used those traits because they were the only ones

his

available... (TODO: Add a better justification for the traits used) More details about the study sites. data

collection. and laboratory protocols can be found in Marrero et al. (2014 and 2017).

Data analysis

conspecific pollen deposited in stigmas, both absolutely and relative to heterospecific pollen, the better the pollination service. Specifically, we used two sets of linear mixed models (LMM) in which the response variables are the number of conspecific and heterospecific pollen grains deposited per stigma in flowers open to animal-mediated pollination. In general, using LMMs in which pollen loads were log-transformed (Figure ??) offered a better fit than a GLMM with Poisson (or quasipoisson) error structure. Models were fitted using the R package nlme 3.1-131 (Pinheiro et al. 2018). m 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 as independent. We selected 183 the best random structure by comparing the median Akaike Information Criterion for small samples (AICc) 38 As predictors in the models, we included the four ecological variables described above. Specifically, we

m -calculated the number of shared pollinators for each plant species by pooling data from the qualitative and

effectiveness of a visit by pollinator species i to plant species j as

quantitative pollination networks. To calculate the plants' relative abundance in their community, we simply

aggregated floral counts for each species. We then calculated the mean visit effectiveness. We define the

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

where  $v_{ij}$  is the observed number of visits by i to j.  $p_{ij}$  is the number of pollen grains from j attached to i.  $v_i$  is the total number of visits performed by i. and  $p_i$  is the total number of grains carried by j. Because the number of shared pollinators, floral abundance, visitation, and pollen production roughly followed a power-law distribution, we log-transformed these three predictors before including them in the model.

15 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 & Legendre 2010; Coux ct al. 2016). To include phenology we treated plant abundance in each of the survey months (November to March) as a "trait" in our analysis. Then we assigned the abundances a weight of 1/5 compared to other functional traits, this accounted for the non-independence of floral counts across months and effectively weights phenology equally
to other traits. We scaled all traits prior to 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, to
facilitate comparison across the four continuous explanatory variables in our models, we rescaled them all to
have a zero mean and unitary variance.

To estimate the coefficients and to perform model selection and quantify the associated uncertainty, we used a combination of multi-model inference and bootstrap resampling with one hundred replicates. First, we performed model selection using the AICc and determined the likelihood of each candidate model (a particular combination of predicting variables) by calculating the median  $\Delta$ AICc (relative to the most likely model) for each bootstrap sample. As we wanted model coefficients from more likely candidate models to carry more weight in our results, we sampled the coefficients for our variables proportional to the likelihood of its candidate model. We "shrank" the estimated coefficient of potentially unimportant variables by setting their estimate to 0 when the variable was not included in the model. Finally, we used these distributions of the model coefficients to estimate their mean impact on the pollination service (in terms of absolute and relative conspecific pollen deposition). Again, here we used bootstrap resampling to estimate the uncertainty around these means.

#### Flexibility of plant strategies

Our last objective was to tease apart the plant strategies that might reduce competition and, importantly, how the community context influences these strategies. If community context plays a relatively small role or the species strategy is inflexible we would expect plants of the same species to fill similar competition for pollination niches across different communities. Alternatively, if the community plays a large role and plants strategies are flexible, we should be able to observe differences in the niche a plant species occupies across communities. First, we used a principal component analysis (PCA) of the four ecological variables. We scaled variables across the whole study to ensure that the PCA space does not change according to the species present in each community. We define a species 'niche in a community as its coordinates in PCA space. For each species that was present in two or more communities, we then calculated (i) the median distance between the strategies that the species uses in different communities and (ii) the area of the convex laul 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.

#### Results

#### Competition vs. facilitation

We first examined the degree of facilitation/competition in natural communities. Specifically, we examined the slope of the relationship between the heterospecific and conspecific pollen deposited on stigmas of each species in a community. We found that overall neither facilitation nor competition dominates (Table S1). Indeed, we found that the proportion of species that experienced a significantly positive or neggative relationship between heterospecific and conspecific pollen than negative was very similar (35% and 37%, respectively; Figure 1a; Table S2). Most of the variation between plants in a facilitation-competition gradient was driven by differences between species; however, we also observed some important differences within species. For instance, six of the species sampled across multiple communities (Carduns acanthoides, Cirsium vulgare, Cypella herbertii, Diplotaxis tennifolia, Hirschfeldia incana, and Verbena intermedia) showed evidence that the same species, can experience facilitation (positive relationships) in some communities and competition (negative relationships) in others. Interestingly, although there was a wide variation in the slope of the relationship between heterospecific and conspecific pollen almost all plants had more conspecific than heterospecific pollen deposited on their stigmas (Figure 1b).

Second, we compared the amount of conspecific pollen deposited on flowers that were open to animal-mediated

Second, we compared the amount of conspecific pollen deposited on flowers that were open to animal-mediated pollinators with that of those that were closed. Here, we again found large differences between plants but with a clear tendency to find more pollen in open flowers (Table S3). Specifically, 66% of plants had significantly more pollen when open and 15% had significantly less pollen when open (the difference was non-significant for the remainding 19%; Figure 1c; Table S4). Variation within species was slightly smaller and only four species (C. acanthoides, C. valgare, H. incana, and Sisyrinchium platense) had both significantly positive and negative differences.

#### Ecological constraints in the pollination service

We then examined the potential roles played by four ecological variables (number of shared pollinators and abundance, mean visit effectiveness, and functional originality) play in pollen deposition. We found that our models of pollen deposition had high explanatory power (the coefficient of determination R<sup>2</sup> ranged between

regative

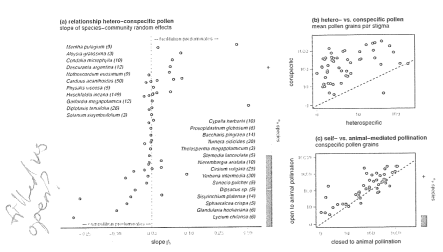


Figure 1: Competition vs. facilitation at the community level. (a) Some species experience more competition for pollination than others. Points indicate the estimated slopes for the relationship between the amount of heterospecific and conspecific pollen deposited per stigma across species in their communities. Significantly negative slopes (darkest shade) indicate that plants experience competition for animal-mediated pollination, while significantly positive slopes (lightest shade) indicate that facilitation is the predominant outcome. The number of plants analysed for each species is shown within brackets next to the species name. (b) Despite the variety of slopes, overall plants had more conspecific than heterospecific pollen deposited in their stigmas. (c) Generally, plants also had more pollen deposited in their stigmas when they were open to animal-mediated pollination than when they were closed. In all plots, error bars correspond to  $\pm$  the standard error (SE).

11

25 0.76 and 0.93) although a large portion of the explanatory power came from the random effects (Table S5).
26 As determined by AICc, the random structure best supported by the data was the one that fit an intercept
27 for each species in each community (as opposed to a common intercept for each species irrespective of the
28 community they belong to). This structure was best for both the models of conspecific and heterospecific
29 pollen (Table S6).

of the four variables, we considered, we found that a plant's mean visit effectiveness and abundance were the most important ecological variables predicting pollen deposition in plant stigmas (Figure 2a). 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 S7).

We found that the relationship between each of the ecological factors and pollen deposition was similar 55 for both conspecific and heterospecific pollen. That is, strategies that were associated with an increase in 237 conspecific pollen were also associated with an increase in heterospecific pollen deposition. Specifically, the plants' mean visit effectiveness had a positive effect on pollen deposition (Figure 2b). However, the effect size was slightly larger for heterospecific than for conspecific pollen. This indicates that, although there is a positive association between visit effectiveness and the absolute amount of conspecific pollen, there is a negative relationship with the relative amount of conspecific relative to heterospecific pollen (Figure 2c). In 🐹 contrast, a plants' relative abundance had a negative effect on the absolute amount of conspecific pollen but the mean difference between the coefficients in the models indicates a positive association with the relative amount of conspecific pollen (Figure 2c). The third most important variable, functional originality, had a positive, although comparatively smaller, association with both the absolute and relative amount of 264 conspecific pollen deposited. Finally, the number of shared pollinators had a negative and neutral association with conspecific and heterospecific pollen, but these impacts were small when compared to the other variables 262 Overall, collinearity did not influence our findings qualitatively. Although relative abundance, the number of shared pollinators, and the visit effectiveness were all positively correlated (Figure S1), the effect each had on conspecific pollen was similar among models that included all or just some of these three explanatory 22 variables (Figure S2). One exception was visit-effectiveness, which exhibits a positive association with the 22 relative amount of conspecific pollen under some variable combinations (all with relatively low AICc support).

#### Flexibility of plant strategies

We used a PCA of the analysed species to investigate whether plants' niche of competition for pollination is similar across communities or whether they are flexible, and therefore a reflection of the community

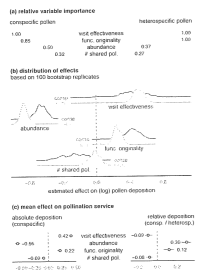


Figure 2: Effect of strategies to minimise competition in the pollination service. (a) The plant's visit effectiveness and relative abundance are the most important variables determining the deposition of conspecific and heterospecific pollen. Meanwhile, the number of shared pollinators was relatively unimportant. The graph shows the relative variable importance calculated as the sum of the Akaike weights of the candidate models that included the variable. (b) The association between ecological variables and heterospecific pollen tended to align with their association with conspecific pollen. Visit effectiveness and functional originality had a positive association with pollen deposition, while 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 variables for conspecific and heterospecific pollen. (c) The end result of these associations is that only the plants' functional originality has a positive impact on both the absolute and relative amount of conspecific pollen deposition (relative to heterospecific pollen). The plot shows the model averaged mean effect (± SE).

## (a) components' variance and variable contributions principal component analysis of ecological variables -o- components' cumulative percentage of variance func originality # shared polabundance contribution (b) plant realised niches in PCA space vex hulls of species niches across communities 1st component (57%) (c) flexibility of plant's strategies median distance between plant niches vs. randomisations S crispa T. megapotamicur B. pingraea C. microphylla C. herbertii C acanthoydes S platense N. euosimum H. incana M. pulegiun V. intermedia

Figure 3: The flexibility of plant strategies. (a) The two first components explain a large proportion of the total variance. (b) When plants that were sampled in more than one community are plotted with these two components, we observe that points, which represent the strategy (the particular combination of ecological variables) of a species in a 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 one of the examined species had strategies that were more tightly connected than what would be expected at random.

context. The first two PCA components explained 75% of the total variance (Figure 3a). The first component was dominated by changes in visit effectiveness 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 (Figure 3b), we observe that the niches 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 analysed species (Figure 3c).

#### 223 Discussion

All else being equal, our results suggest that community context plays a central role in determining net
cost or benefit of sharing pollinators. First, we found that the costs of pollinator sharing are pervasive.
In particular, we found that a significant proportion of species experiences overall neutral or competitive
effects from animal-mediated pollination. Second, we found that multiple ecological factors can modulate the
intensity of this competition; however, conspecific and heterospecific pollen deposition are rightly coupled and
therefore these factors are likely to involve a trade-off between the quantity and quality of pollination. Third,
we found that the way these factors influence competition is strongly shaped by the community context, this
is, if collectively these factors are used to define a niche of competition for pollination, the same plant species
can occupy dramatically different niches depending on the community it belongs to.

Both of the tests we used to identify the effect of animal-mediated pollination suggested the same outcomes
that net facilitation is only experienced by a fraction of species in a community. A recent study also suggested
as wide range of outcomes in a multi-species context (Tur ct al. 2016). However, the factors explaining why
some species experience more competition than others were not understood. Here we show that the prevalence
of neutral and competitive effects of animal-mediated pollination can be at least partially explained by the
trade-offs that minimising competition can involve.

These trade-offs were most evident in precisely the factors that most strongly modulate pollen deposition.

Specifically, we found that the factors related to the relative dominance of the species (in terms of pollen and visits, or abundance) also tended to be the most important predictors of pollen deposition. At the same time, receiving high visitation increases the absolute levels of conspecific pollen deposition but increases heterospecific deposition to a greater extent—even when the visitors are likely to carry a high proportion of conspecific pollen. Similarly, being abundant reduces the amount of heterospecific pollen deposited but at the expense of a noticeable reduction in the amount of conspecific pollen. Multiple studies recognise the

15

importance of visitation and abundance to competition for pollination (Feldman et al. 2004; Morales & Traveset 2008; Muñoz & Cavieres 2008). Our results strongly corroborate the importance of these two factors but also suggest that (because visitation, pollen production and abundance are correlated; Sargent & Otto 2006), plants must reach a fine balance between increasing conspecific pollen and reduce heterospecific pollen deposition to minimise competition for pollination.

Further evidence of the intricacies of balancing costs and benefits can be found when we examine the other two factors analysed in our models. We show that being a specialist and sharing no pollinators might be preferable in the short term (as it is generally detrimental to pollen deposition). However, over long periods of time, there could be a risk associated with a specialist plant having no pollinators. Hence, is likely that to ensure long term survival, plants also need to balance this risk with the costs of sharing pollinators. One possible solution is to share pollinators and have original traits—as we show that trait originality is generally beneficial to pollen deposition and it's commonly thought that species that are further from others in trait space suffer reduced competition. Yet again, there are two possible caveats to this strategy that highlight the interrelatedness of the ecological factors. First, in a mutualism context, it is also possible that trait originality could come at the cost of being less 'apparent' to pollinators. Second, the usually negative, relationship between originality and generalism has been shown depend on the plant's abundance (Coux et al. 2016), with generalist species being able to also have original traits only when they are abundant enough to provide a valuable reward to make it worth to pollinators.

How precisely plants balance the costs and benefits in the short and the long term likely depends on the species sensitivity to different risks but also, as we show here, on the community it belongs to. For instance, a plant species that experience a large reduction in seed set due to heterospecific pollen deposition might be better off by sharing few pollinators, while sharing pollinators makes more sense for those mainly affected by low levels of conspecific pollen deposition (Campbell & Motten 1985). Nevertheless, the diversity of the facilitation/competition outcomes across communities we observed and the relevance of random effects in our models suggests that the community context has a very important effect on the way plant species respond to competition for pollination beyond species identity. Perhaps overgeneralising, some plant families (Asteraceae for example) are often generalists in their communities, others (Orchidaceae) are known to be more commonly specialised. While it is possible, from an evolutionary perspective, that selection for a particular strategy might say something about the community in which a species has typically inhabited during its evolutionary history, we show that most plant species are still flexible enough to adopt markedly different niches in different communities. From a more applied perspective, flowering plants are sometimes introduced to attract pollinators on other nearby plants. Our results highlight the difficulties involved in

16

MINT

nen. Atmopie statues recognise the

predicting whether the introduced plant species will facilitate or compete with neighbours. This is so because
the outcome might depend not only on the particular niche each species occupies in a particular community
but also on other factors that we were unable to measure (Cariveau & Norton 2009; Flanagan et al. 2011;
predicting whether the introduced plant species will facilitate or compete with neighbours. This is so because
the outcome might depend not only on the particular niche each species occupies in a particular community
to but also on other factors that we were unable to measure (Cariveau & Norton 2009; Flanagan et al. 2011;
the outcome might depend not only on the particular niche each species occupies in a particular community
to but also on other factors that we were unable to measure (Cariveau & Norton 2009; Flanagan et al. 2011;
the outcome might depend not only on the particular niche each species occupies in a particular community
to but also on other factors that we were unable to measure (Cariveau & Norton 2009; Flanagan et al. 2011;
the outcome might depend not only on the particular niche each species occupies in a particular community
to but also on other factors that we were unable to measure (Cariveau & Norton 2009; Flanagan et al. 2011;
the outcome of the outcome

that introduced species are more likely to be beneficial if they have different traits to local plants.

Overall, our results suggest that pollination communities might not necessarily be the paradigmatic mutualistic communities we often believe. Instead, we confirm that plant competition is pervasive in pollination communities and that potential strategies to minimise competition are likely to involve strong trade-offs in the pollination service either in the short or the long term. As expected, not sharing pollinators and being functionally unique reduces competition for pollination by increasing conspecific pollen deposition (both in absolute numbers and relative to heterospecific pollen). However, this increase is surprisingly small when compared to the effect of relative abundance and visitation. Interestingly, these two factors also involve the largest trade-offs between increasing deposition of conspecific pollen and reducing that of heterospecific pollen. Despite large intra-species differences among species our results provide solid evidence of the role that competition for pollination plays in natural communities. Species coexistence is not solely determined by direct and indirect mutualistic relationships, but also strongly shaped by the multiple intertwined ecological factors that shape competition for pollination at the community level. Achieving a better understanding of how biodiversity is supported by pollination communities will require seeing them as both mutualistic and competitive communities.

#### 357 Acknowledgements

We thank Cátedra de Botánica General. Facultad de Agronomía. Universidad de Buenos Aires. The Agrasar
and Bordeu families, and the University of Buenos Aires, for logistical support and permission to conduct
this study at estancias Anquilóo. Las Chilcas and San Claudio, respectively. Fieldwork was supported
by grants PICT 08-12504 and 0851. EFC acknowledges the support from the University of Canterbury
Doctoral Scholarship, the University of Canterbury Meadow Mushrooms Postgraduate Scholarship, and
a New Zealand International Doctoral Research Scholarship, DBS and JMT acknowledge the support of a
Rutherford Discovery Fellowship. DBS also acknowledges the Marsden Fund Council from New Zealand
Government funding.

gens blue

Nice.

Table S1: Summary of the model used to analyse the relationship between heterospecific and conspecific redlen

predictor	estimate	S.E.	z-value
fixed component			
(Intercept)	1.976	0.279	17.862
heterospecific	0,008	0.017	0.474
random component (s	pecies:com	munity	)
S.D. random intercept	1.964	-	
S.D. random slope	0.120	-	

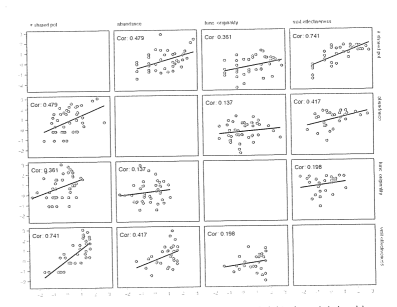


Figure S1: Correlation between the explanatory variables included in the statistical models.

:3

species name			
	community	slope	S.E.
Alogsia gratissima	Anquilóo - reservo - 2	0.0746	0.0144
Bacchavis pingraea	San Claudio - reserve - 1	-0.0012	0.0359
Cardaus acanthoides Cardaus acanthoides	Anquilóo - agricultural - 2	0.0116	0.0147
	San Claudio - agricultural - 1	-0.0106	0,0040
Carduus acanthoides Carduus acanthoides	San Claudio - agricultural - 2	0.0518	0.0044
	San Claudio - reserve - 1	0.0781	0.0710
Carduus ucunthoides	San Claudio - reserve - 2	-0.0008	0.0359
Cirsiam vulgare	Auquilóo - agricultural - 2	-0.0401	0.0025
Cirsium vulgare	Las Chilcas - reserve - 1	0.0007	0.0012
Cirsium valgare	San Claudio - agricultural - 2	0.0197	0.0158
Cirsium valgare	San Claudio - reserve - 1	-0.0149	0,0076
Condalia microphylla	Anquilóo - reserve - 1	0.0487	0.0200
Cypella herbertii	Las Chileas - agricultural - 2	0.0037	0.0002
Cypella herbertii	Las Chileas - reserve - 1	-0.0052	0.0001
Descurania argentina	Anquilóo - agricultural - 2	0.0429	0,0048
Diplotaxis tenuifolia	Auquilóo - reserve - 1	0.0008	0.0004
Diplotaxis tenuifolia	Anquilóo - reserve - 2	0.5173	0.0270
Diplotavis tennifolia	San Claudio - reserve - 2	-0.0045	0.0001
Dipsacus sp.	San Claudio - reserve - 2	-0.0368	0.0648
Gaillardia megapotamica	Auquilóo - reserve - 2	0,0016	0,0004
Glandularia hookeriana	Auquilóo - reserve - 2	-0.0942	0.0244
Hirschfeldia incana	Auquilóo - agricultural - 1	-0.0045	0.0013
Hirschfeldia incana	Anquilóo - agricultural - 2	~0.0148	0.0057
Hirschfeldio incana	San Claudio - agricultural - 1	0.0110	0.0020
Hirschfeldia incana	San Claudio - agricultural - 2	0.0031	0.0023
Hirschfeldia incana	San Claudio - reserve - 1	0.0022	0.0002
Hirschfeldia incana	San Claudio - reserve - 2	0.0432	0.0020
Lycium chilense	Auquilóo - reserve - 2	-0.3355	0,0087
Mentha palegium	Las Chileas - agricultural - 2	0.0136	0.0866
Mentho pulcyium	Las Chilcas - reserve - 1	0.3973	0.0388
Nierembergia aristata	Auquilóo - agricultural - 1	0.0197	0.0217
Nicrembergia avistata	Auquilóo - reserve - 1	-0,0065	0.0016
Nicrembergia aristata	Anquilóo - reserve - 2	-0.0048	0.0011
Nothoscordam cuosimum	Las Chilcas - agricultural - 1	0.0405	0.0034
Nothoscordam cuosimam	Las Chikas - agricultural - 2	-0,0045	0.1162
Physalis viscosa	Anquilóo - agricultural - 1	0.0041	0.0005
Prosopidastrum ylohosum	Auquilóo - reserve - 2	-0.0012	0.0194
Senecio pulcher	Las Chileas - agricultural - 1	-0.0104	0.0007
Sisyvinchium platense	Las Chileas - agricultural - 1	-0.2850	0.0203
Sisyvinchium platense	Las Chileas - agricultural - 2	-0.0487	0.0324
Sisyrinchium platense	Las Chilcas - reserve - 1	0.0206	0.1143
Solanum sisymbriifolium	San Claudio - agricultural - 1	0.0002	0.0004
Sphaeraleea crispa	Anquilóo - reserve - 1	-0.0601	0.0133
Stemodia lanceolata	Las Chilcas - agricultural - 1	-0,0044	0.0001
Thelesperma megapotamicum	Auquilóo - agricultural - 1	-0.0022	0.0025
Turnera sidioides Turnera sidioides	Anquilóo - agricultural - 1	-0.0002	1,000,0
	Anquilóo - agricultural - 2	-0.0140	0.0170
Turnera sidioides Verbena intermedia	Anquilóo - reserve - 2	-0,0014	0.0002
	Auquilóo - reserve - 2	-0.0643	0.0327
Verbena intermedia	San Claudio - agricultural - 2	0.0932	0.0071
Verbena intermedia	San Claudio - reserve - 2	-0.0073	0.0101

Table S3: Summary of the model used to analyse the relationship between conspecific pollen deposited in bagged and unbagged flowers (open to animal pollination).

predictor	estimate	S.E.	z-value
fixed component			
(Intercept)	4.215	0.318	13.235
treatment (unbagged flower)	0.845	0.205	4.128
random component (species	:communi	ty)	
S.D. random intercept	2.240	-	_
S.D. random slope	1.377	-	_



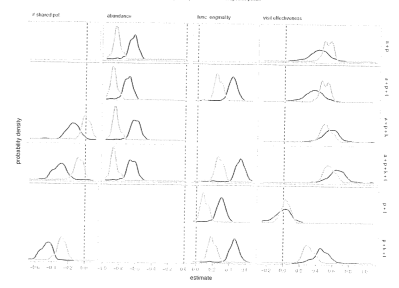


Figure S2: Distribution of effect estimates for models of conspecific and heterospecific pollen density gain. Model formulas have been abbreviated: a for abundance, k for the number of shared pollinators, p for the visit effectiveness, and t for trait originality. Only candidate formulas with a  $\Delta AICc < 4$  for either conspecific or heterospecific pollen are shown. Models candidates are arranged in decreasing order of support.

Table S4: Amount of conspecific pollen deposited in bagged and unbagged flowers (open to animal pollination) for each species in their community. Community names are constructed by location - agricultural/restored - fragment number.

		bagged		unbagged	
species name	community	mean	C.L.	mean	C.L
Mogsia gratissima	Anquilóo - reserve - 2	1	[0, 1]	39	[26, 59]
Baccharis pingraca	San Claudio - reserve - 1	()	[0, 0]	4	[2, 8]
Carduus acanthoides	Anquilóo - agricultural - 2	25	[23, 27]	56	[52, 61]
Carduns acanthoides	San Claudio - agricultural - 1	34	[32, 35]	42	[40, 44]
Carduus avanthoides	San Claudio - agricultural - 2	41	[39, 43]	16	[16, 17]
Carduus acanthoides	San Claudio - reserve - 1	12	[11, 14]	10	[9, 11]
Cardaus avanthoides	San Claudio - reserve - 2	10	9, 11]	1.5	[14, 17]
Cirsiam valgare	Auquilóo - agricultural - 2	211	[205, 217]	222	[215, 228]
Cirsium valgare	Las Chileas - reserve - 1	284	[277, 291]	320	313, 328
Cirsium onlgare	San Claudio - agricultural - 2	33	[8, 132]	76	[19, 302]
Cirsium vulgare	San Claudio - reserve - 1	218	[210, 226]	81	[78, 84]
Condulia microphylla	Auquilóo - reserve - 1	38	[36, 41]	35	[33, 37]
Cypella herbertii	Las Chileas - agricultural - 2	1095	[276, 4336]	2738	[691, 108]
Sypella herbertii	Las Chileas - reserve - 1	478	[462, 494]	2743	[2652, 28
Descurania argentina	Anquilóo - agricultural - 2	90	[86, 93]	117	[113, 121]
Diplotaxis tenuifolia	Auquilóo - reserve - 1	362	[92, 1435]	881	[222, 348]
Diplotaxis tenuifolia	Auquilóo - reserve - 2	177	[45, 700]	422	[107, 167
Diplotaxis tenuifolia	San Claudio - reserve - 2	769	[762, 776]	1153	[1143, 11]
Dipsacus sp.	San Claudio - reserve - 2	-4	[3, 5]	1.4	[12, 17]
Gaillardia megapotamica	Auquilóo - reserve - 2	590	[580, 601]	179	[175, 182]
Handularia hookeriana	Anquilóo - reserve - 2	185	[178, 192]	131	[127, 136]
Hirschfeldia incana	Anquilóo - agricultural - 1	432	[427, 437]	412	[408, 417
Hirschfeldia incana	Anquilóo - agricultural - 2	246	[240, 252]	75×	[740, 778]
Hirschfeldia incana	San Claudio - agricultural - 1	107	[403, 412]	271	268, 274
Tirschfeldia incana	San Claudio - agricultural - 2	291	[288, 294]	305	302, 308
Hirschfeldin incana	San Claudio - reserve - 1	384	[380, 389]	355	351, 359
Hirschfeldia incana	San Claudio - reserve - 2	340	[337, 344]	465	[460, 470]
Lycium chilense	Anquilóo - reserve - 2	998	[987, 1009]	1339	[1325, 13]
Mentha pulegium	Las Chileas - agricultural - 2	1	[1, 2]	3	[2, 4]
Mentha pulcyium	Las Chilcas - reserve - 1	7	[6, 8]	15	[12, 18]
Vicrembergia aristata	Anquilóo - agricultural - 1	116	[105, 128]	835	[756, 922]
Vicrembergia aristata	Anquilóo - reserve - 1	179	[171, 187]	1072	[1024, 11]
Vicrembergia aristata	Auquilóo - reserve - 2	71	[67, 76]	1054	984, 112
Nothoscordum cuosimum	Las Chileas - agricultural - 1	92	[88, 97]	408	388, 428
Nothoscordum cuosimum	Las Chileas - agricultural - 2	324	[315, 334]	352	343, 362
Oxalis violeta	San Claudio - reserve - 2	371	[122, 1131]	467	[153, 142]
Physalis viscosa	Anquilóo - agricultural - I	1227	[1211, 1244]	2732	2696, 27
Prosopidastrum globosum	Auquilóo - reserve - 2	10	[8, 11]	13	[12, 15]
šenecio pulcher	Las Chileas - agricultural - 1	358	[348, 367]	406	395, 117
Sisyrinchium platense	Las Chileas - agricultural - 1	91	[88, 95]	159	152, 165
Sisyrinchium platense	Las Chileas - agricultural - 2	35	[9, 139]	×1	[20, 319]
Sisyvinchium platense	Las Chileas - reserve - 1	193	[179, 208]	73	[67, 79]
Solanum sisymbriifolium	San Claudio - agricultural - 1	57	[50, 66]	2194	[1923, 25]
Sphaeraleca erispa	Anquilóo - reserve - 1	2	[2, 2]	9	[8, 10]
stemodia lanecolata	Las Chileas - agricultural - 1	387	[380, 394]	1919	[1884, 19
Phelesperma megapotamicum	Anquilóo - agricultural - 1	314	[306, 322]	327	[319, 330
Farnera sidioides	Anquilóo - agricultural - 1	53	51. 55	198	[189, 200
Farnera sidioides	Anquilóo - agricultural - 2	1	[0, 1]	1	[2, 8]
Furnera sidioides	Anquilóo - reserve - 2	1	[1, 2]	189	[113, 31]
Serbena intermedia	Anquilóo - reserve - 2	67	[64, 70]	125	[119, 13]
Verbena intermedia	San Claudio - agricultural - 2	34	[32, 36]	66	[63, 70]
Verbena intermedia	San Claudio - reserve - 2	136	[133, 139]	356	348, 367

Table S5: The coefficient of determination  $\mathbb{R}^2$  of the most parsimonious pollen deposition models (those with the lowest AICc). The marginal coefficient of determination describes the proportion of variance explained

conditional $R^2_{(\cdot\cdot)}$		marginal $R_{(m)}^2$			
mean	min	max	mean	min	max
conspe	eific p	ollen			
0.91	0.87	0.93	0.09	0.06	0.14
heteros	pecific	polle	1		
0.80	0.76	0.87	0.27	0.21	0.35

Table S6: Comparison of the two random structures we considered for the models of conspecific and heterospecific pollen deposition. The table shows median  $\Delta AIC$  values of 99 bootstrap resamples of the data. The 5th and 95th percentile are shown inside square brackets. Communities are defined by individual fragments but ignore the hierarchical arrangement of sampling sites.

	$\Delta AIC$		
random structure	median	C.L.	
conspecific pollen			
I   plant sp. * community	0,0	[0, 0]	
1 plant sp.	30.7	[8.2, 58.1]	
heterospecific pollen			
1   plant sp. * community	0.0	[0, 0]	
1   plant sp.	14.6	19.3, 88.1	

.

by just the fixed effects.

.

Table S7: Comparison of the different fixed structures we considered for the models of conspecific and heterospecific pollen deposition. The table shows median  $\Delta AIC$  values of 99 bootstrap resamples of the data. The 5th and 95th percentile are shown inside square brackets.

		$\Delta AIC$
fixed structure	median	C.L.
conspecific pollen		
- abundance + share pollen	0.0	(0, 0)
<ul> <li>abundance + share pollen + func, originality</li> </ul>	0.9	[0.4, 1.3]
- abundance + share pollen + degree	1.9	[1.6].2.1]
<ul> <li>abundance + share pollen + degree + func, originality</li> </ul>	2.2	1.6 .2.8
- share pollen + func. originality	2.8	2.1 ,3.8
<ul> <li>share pollen + degree + func, originality</li> </ul>	3.6	[2.3 ,4.6]
- share pollen	118.3	75.3 .178.7
- share pollen + degree	119.0	[76], [79,9]
- abundance	189.7	[150.1],239,
<ul> <li>abundance + func. originality</li> </ul>	191.6	151.7 .241.
~ abundance + degree	191.7	151.9 .241.
- func. originality	192.5	152.9 .242.
<ul> <li>abundance + degree + func, originality</li> </ul>	193.7	153.6 ,243.
- degree + func. originality	193.7	154.6 .243.
- degree	351.8	293.5 ,419.
heterospecific pollen		
- abundance + share pollen	0.0	[0], 0]
<ul> <li>abundance + share pollen + func. originality</li> </ul>	1.1	[0.5],1.5]
- abundance + share pollen + degree	2.1	[1.9 .2.1]
- abundance + share pollen + degree + func, originality	3.1	[2.6 ,3.5]
- share pollen + func, originality	11.9	[10],13.9
<ul> <li>share pollen + degree + func, originality</li> </ul>	13.2	[11.2],15.2]
share pollen	67.5	53.4 .87.5
- share pollen + degree	68.4	54.2 .88.7
- abundance + degree	206,9	160.6 ,251.
- abundance	207.6	162.8 .251.
<ul> <li>abundance + func, originality</li> </ul>	208.6	[163.2],252.6
- abundance + degree + func, originality	208.6	162.2 .253.:
- func. originality	214.3	168.3 ,258,
<ul> <li>degree + func, originality</li> </ul>	216.3	1170.3 .260.0
- degree	336.0	282.6 ,391,3