working title Compatibility system and stigma size are the main predictors of heterospecific pollen effect

Jose B. Lanuza ^{* 1} Ignasi Bartomeus ² Tia-Lynn Ashman ³ Romina Rader ¹

¹University of New England (Australia) 5 6 ²Estacion Biologica de Donana (EBD-CSIC), E-41092 Sevilla, Spain 7 8 ³ Department of Biological Sciences, University of Pittsburgh 4249 Fifth Avenue, Pittsburgh, Pennsylvania 15260-3929 USA 9 10 corresponding author: barragansljose@qmail.com

Pollinator sharing can have negative consequences for plant fitness with the arrival of foreign pollen. However, the costs of heterospecific pollen are not yet well understood. We conducted a glasshouse 13 experiment to understand how phylogenetic relatedness and plant traits mediate the impacts of heterospecific pollen transfer. We conducted 1800 crosses by experimentally transferring pollen (50%) 15 and 100% ratio) with reciprocal crosses between 10 species belonging to three different families: 16 Brassicaceae, Solanaceae and Convolvulaceae. Seed set was used as proxy of plant fitness. We found 17 that for 65% of the treatments with 50% mix reduced seed set. Moreover, the reduction in seed set was dependent on the degree of relatedness and reproductive traits of the pollen recipient and not the 19 pollen donor. Our results show that certain traits, particularly compatibility system, are critical in 20 understanding the costs of heterospecific pollen.

Keywords: heterospecific pollen, plant reproduction, fitness, interspecific competition, phylogenetic distance.

INTRODUCTION

- In most ecosystems, plant species normally coexist and share their floral visitors with other species Waser et al. (1996). From the plants' perspective, pollinator sharing can be positive for some plants
- Carvalheiro et al. (2014) or negative for others Pauw (2013), depending on the facilitation gradient. An
- increasing number of visits often correlates with higher chances of fertilization Engel and Irwin (2003).
- However this is not always the case, among these possible flower visitors there are also nectar robbers
- and pollen thieves Inouye (1980); Magrach et al. (2017). Receiving both sufficient quantity and quality
- deposited on the stigma is thus highly relevant to the pollination success of the plant Aizen and Harder

(2007).

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By visiting many plant species, many pollinators are responsible for conspecific pollen loss and the transport of foreign pollen, both of which can have important detrimental effects on species fitness Morales and Traveset (2008); Ashman and Arceo-Gómez (2013); Arceo-Gómez and Ashman (2016). Foreign pollen arrival can play an important role in plant species fitness but outcomes are variable and appear to be context dependent as there is not always a decrease in fitness Morales and Traveset (2008). 37 Some of this variation is likely due to the enormous variability of foreign pollen transferred across systems ranging from 0 to 75 percent. However, most studies report ranges of heterospecific pollen between 0 and 20 percent of the total pollen load Bartomeus et al. (2008) Montgomery and Rathcke (2012); Ashman and Arceo-Gómez (2013); Fang and Huang (2013), yet even these relatively low 41 amounts of heterospecific pollen transferred can decrease fitness greatly Thomson et al. (1982). While 42 we now have some understanding of the impacts of heterospecific pollen quantity, we have less 43 understanding of other factors that could be driving the variation in impacts upon fitness. Ashman and Arceo-Gómez (2013) postulated the first predictive framework that identifies a need to understand how plant traits might mediate heterospecific pollen effect, whereby mating system and pollen size were predicted to potentially mediate the impact of foreign pollen transfer on plant fitness. This concept is supported by specific case studies, such as Tong and Huang (2016) that demonstrate an asymmetrical effect in 6 species of *Pedicularis* whereby the pollen of long styled species was able to grow the full length of the style on short styled species but not vice versa. While this suggests that the impacts of heterospecific pollen may differ among pollen donor and recipient, few studies have been conducted to ascertain whether this pattern is in fact a general trend or to identify the extent to which other plant traits are critical to heterospecific pollen impacts. Plant traits are crucial to understand heterospecific pollen effect but the multifactorial nature of the traits that are involve in the pollen-pistil interaction make difficult to unravel what are the main traits in driving the effect. These traits can be seen from a male perspective of both donor and recipient where pollen size, pollen aperture number and pollen allelopathy are key components to understand the outcome of foreign pollen arrival Murphy and Aarssen (1995); Ashman and Arceo-Gómez (2013). In Ashman and Arceo-Gómez (2013) small pollen is predicted to cause a greater fitness decrease, although this can be true there are also other possibilities to consider which can obscure a predictive framework like big pollen can clogg small stigmas with fewer pollen grains, bigger stigmas are less likely to be

clogged by small pollen grains and bigger pollen can outcompete smaller pollen grains due to faster pollen tube growth rate Williams and Rouse (1990). Moreover, to understand the different mechanical or chemical effects of pollen also the female traits of the pollen recipient have to be considered, from the literature these main traits are: stigma size, style length, number of ovules, incompatibility system and also a structural trait such as flower morphology Montgomery and Rathcke (2012); Ashman and Arceo-Gómez (2013); Tong and Huang (2016). For example, greater stigmatic area is positively 67 correlated with greater heterospecific pollen deposition Montgomery and Rathcke (2012) and therefore possibly with an increase in negative effect. For species that are self-incompatible the barriers towards heterospecific pollen are stronger than self-compatible species Ashman and Arceo-Gómez (2013). Nonetheless, an effect of foreign pollen is a bit obscured by the variability within species, however species that are strong selfers or strong outcrossers have less variablity in mating systems and predictions of effect could be more realistic (see figure 1 from Whitehead et al. (2018)). Although past research has progress in the understanding of what traits can mediate the effect as we have shown here, there are multiple traits involved and multiple possible scenarios still to be explored empirically for a full understanding of the importance of heterospecific pollen effect in nature. For the understanding at what level or intensity the interference of pollen can occur is important to consider the relatedness of the interacting species. Closely related species are more likely to have similar traits Letten and Cornwell (2015). The similarity in traits between closely related species can lead to higher chances of ovule usurpation/abortion Arceo-Gómez and Ashman (2011) and therefore studies predict and show a greater negative effect of closely related species Ashman and Arceo-Gómez (2013); Arceo-Gómez and Ashman (2016). Few studies however, have focused on the impacts of heterospecific pollen on fitness of distantly related species Galen and Gregory (1989); Neiland and Wilcock (1999). Despite the fact that far related species are also able to decrease species fitness (REFS). Yet, most insects and most stigmas have been found to carry multiple species of foreign pollen with little attention to degree of relatedness (Arceo-Gómez and Ashman (2016); Fang and Huang (2013). Understanding the role of foreign pollen from distantly related species thus deserves greater attention. The relatedness of foreign pollen gives a first snapshoot of where the pollen competition can occur and therefore could be a proxy of effect. With that said, Arceo-Gómez and Ashman (2016) is the only work which has proven a greater effect of close related species through a meta-analysis but with

low sample sizes and lack of significance. Therefore, there is a need to study the effect of heterospecific pollen of far and close related species at community level beyond single pairwise interactions.

Heterospecific pollen studies in nature have the complexity added of great environmental variability which can lead to confounding interpretations. Moreover, the great diversity of floral structures and small sizes of some of them such as Asteraceae species make tedious or almost impossible the studies of heterospecific pollen on the field. For this reason, we investigated how floral reproductive traits and relatedness mediate the impact of heterospecific pollen by creating an artificial co-flowering community in a glasshouse with 10 species belonging to three different families with different traits. Our study tries to answer the following questions: (i) Do heterospecific pollen reduce seed set? To what extent do (ii) floral reproductive traits and (iii) relatedness, mediate the impacts of heterospecific pollen on seed set.

01 METHODS

The study was conducted in a glasshouse at University of New England (Armidale, Australia) from November 2017 to March 2018. Rooms were temperature controlled depending on the requirements of 103 the species with day and night temperature differences. The experimental design had species from three 104 different families: Solanaceae, Brassicaceae and Convolvulaceae (Table 1). The species of the study 105 had different reproductive traits and different degree of relatedness (Figure 1) where the reciprocal 106 crosses between species allowed us to have multiple different scenarios of both traits and relatedness. 107 Moreover, the species selected had fast life cycle and low structural flower complexity in order to 108 perform the pollination treatments and grow the different species from seeds. For the purpose of the 109 experiment all the species where considered as pollen recipient and as pollen donor (see interaction 110 matrix, fig 2). Species were watered once or twice per day and fertilized weekly (NPK 23: 3.95: 14).

112 **Table 1**

Family	Genus	Species
Brassicaceae	Brassica	Brassica rapa
Brassicaceae	Brassica	Brassica oleracea
Brassicaceae	Eruca	Eruca versicaria
Brassicaceae	Sinapis	Sinapis alba
Convolvulaceae	Ipomoea	Ipomoea aquatica
Convolvulaceae	Ipomoea	Ipomoea purpurea
Solanaceae	Capsicum	Capsicum annuum
Solanaceae	Petunia	Petunia integrifolia
Solanaceae	Solanum	Solanum lycopersicum
Solanaceae	Solanum	Solanum melongena

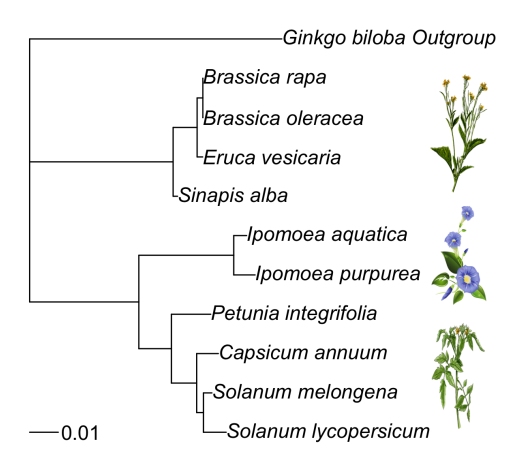


Figure 1: Phylogenetic tree of the ten species used in the experiment from three different families from top to bottom: Brassicaceae, Convolvulaceae and Solanaceae

113 Hand pollination

Foreign pollen effect was studied through two different treatments, one with 50% conspecific pollen and 114 50% heterospecific pollen and a second one with 100% foreign pollen in order to see if foreign pollen 115 can trigger fruit production by itself or even seeds through ovule usurpation. Therefore, we performed 116 180 different heterospecific treatments (N=10). Seed set was the proxy of effect for all our treatments. 117 Moreover, hand cross-pollination (between individuals of the same species), hand self-pollination, apomixis (bagged emasculated flowers) and natural selfing were tested for each species (N=10). For the treatments with foreign pollen and hand cross-pollination, flowers were emasculated the day prior 120 anthesis and hand pollinated next day with a toothpick. Hand-pollination was conducted with 3-4 121 gentle touches on the stigma surface. For each species 20 anthers were collected and their pollen 122 counted with a hemocytometer, each anther was counted 4 times and then an average of these counts 123 was performed. Once, the average number of pollen grains per anther was known, the proportion of anthers per mix was calculated in order to achieve a 50-50% mix. To confirm that the treatments 125 applied were the desire proportions, the total stigmatic load of pollen was counted and the proportions 126 calculated between the two species of the mix. Because pollen from the same family was difficult to 127 distinguish, and we expected similar properties in mixing, we counted pollen from just one randomly 128 selected species within each donor family different to the focal's family (N=3). 129

130 Traits and evolutive distance

The traits measured for each species were pollen per anther, number of ovules and stigma, style, ovary 131 width and length. For the stigma, the stigmatic area was also measured and moreover the stigmas were 132 divided in wet/dry type. All the morphometrical measurements were performed with a 133 stereophotomicroscope. Pollen was counted for 20 anthers of each species with 4 replicates per sample with an hemocytometer. Previously, anthers were squashed on a known solution with the pippete tip 135 and homogeneize with a vortex for 30 seconds. Ovule number was counted with the help of a 136 stereomicroscope and a small grid over a petri dish from 15 randomly selected flowers. Fruits per 137 number of flowers treated were counted for just Solanaceae species, we counted for all the species the 138 number of seeds produced per average number of ovules. Levels of self-incompatibility were estimated by dividing the fruit set of hand self-pollination by hand cross pollination Lloyd and Schoen (1992). 140

141 Analysis

To evaluate heterospecific pollen effect on seed production we performed linear mixed models. The
distinct heterospecific pollination treatments were compared through relevelling each variable with the
cross pollination treatment which was our control for optimum seed production for all the species. The
different replicates of each treatment were considered as random effects. Seed production was scaled for
all the species with mean 0 and standard deviation of 1 prior to the analysis. All the analysis were
conducted with the statistical language R (R Core Team 2018).

In order to test the relative effect of traits on seed production with foreign pollen we performed Mantel 148 test in R (Vegan package, Euclidean distance) between the assymetrical matrix of heterospecific pollen 149 effect (10 by 10 matrix) with the different distance matrices of traits. Heterospecific pollen effect was 150 obtained through the subtraction of seed production by hand cross-pollination minus seed production 151 of the different heterospecific pollen treatments. To find a model with the best explanatory traits we 152 used the function **bioenv** from R. We also conducted Mantel test between the matrix of heterospecific 153 pollen effect and the distance matrix from all the traits. Moreover, we explored also the correlation 154 between traits and heterospecific pollen effect through generalized mixed models where the response 155 variable was heterospecific pollen effect and the explanatory variable the different traits. In addition, 156 we tested the correlation between the total amount of pollen deposited on the stigma with heterospecific pollinations and the stigma size through Pearson's correlation. 158

We conducted mantel test to check for correlations between heterospecific pollen effect and phylogenetic distance. Due to improvents in statistical power we used the square root of the phylogenetic distance (Letten & Cornwell 2014). Two different phylogenetic distances were used from two kinds of markers: 1)

Internal transcribed spacer (ITS) and 2) ribulose-bisphosphate carboxylase (RBCL). The sequences were obtained from GenBank (https://www.ncbi.nlm.nih.gov/genbank/, accessed 20 Oct. 2018). The sequences were aligned with clustal omega and the pairwise evolutive distances calculated with MEGA7.

Phylogenetic signal of traits?

Add pollen on stigma counts to methods

Next bit to results, also divided it with

- Total pollen deposited on stigmas was significantly correlated with the stigmatic area, pearsons correlation=0.57 and p-value=0.008. Proportions of pollen talk about it to 50-50 mix? Talk about this in methods too.
- Add also plot with fruit production and pot it in appendix. Also the plots of ratios to appendices.
- 172 Add effect sizes to analysis!!
- 173 Mention that SI index is calculated (Schoen & Lloyd 1984)

74 RESULTS

Results of hand cross-pollination, hand self-pollination, natural selfing and apomixis are presented in Figure 1 (see appendix 1 for table with values). Heterospecific pollen reduced seed set significatively 176 with the 50-50% heterospecific pollen treatments for 65% of the pairwise interactions p<0.05. Moreover, 177 average effect sizes differed across species and across families, see Figure 2. Despite some variability in the effect of the different pollen donors per species, in general terms the effect of heterospecific pollen 179 from the distinct nine treatments per species was homogeneous (see Figure 3), just for four species out 180 of ten, just one donor did not overlap the confidence intervals with other donors. Therefore, none of the 181 donors had a clear stronger or weaker effect across species. When the donors were groped by family not 182 big differences were seen, just for S. lycopersicum the confidence intervals of Brassicaceae and 183 Convolvulaceae did not overlap (see **Figure 4**). In addition, for the 100% hetrospecific pollen 184 treatments we did not find almost seed production. However, for just one species (S. alba) the control 185 pollination and the heterospecific pollination with pollen from a confamilial had similar seed 186 production. For two Solanaceae species S. melongena and C. annuum 100% pollen treatments 187 produced few seedles fruits (3% and 9% respectively) and they did not for the apomictic treatments. Results of Mantel test between heterospecific pollen effect and phylogenetic distance gave a positive 189 statistically clear correlation for both markers (p<0.05). The correlations with ITS and RBCL markers 190 was respectively of 0.29 and 0.25. We found a significant phylogenetic signal of traits for pollen size, 191 stigma measurements and style length (p<0.05). Although with a lack of a significant correlation 192

pagel's lambda values were also relatively high (>0.45) for incompatibility index, ovary length and levels of selfing. Moreover, Mantel test between heterospecific pollen effect and traits gave also a positive significant correlation with a r value of 0.4. When the effect was look trait by trait with Mantel test, stigma type and stigma measurements (length, width and area) gave a significant positive correlation with heterospecific pollen effect.

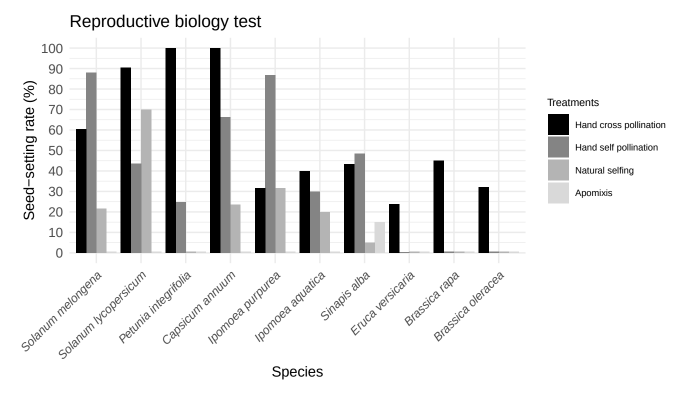


Figure 2: Barplot with the different treatments that provide information of the reproductive biology of the ten species. The y axis is the proportion of ovules converted to seed in percentage. The different treatments (N=10) which are presented in the legend are, hand cross-pollination, hand self-pollination, natural selfing and apomixis. More information about these treatments can be found in Methods and Appendices.

Maybe for this plot I have to do hedges'g instead of cohean because of different sample sizes when I

199 group the treatments per family. Check

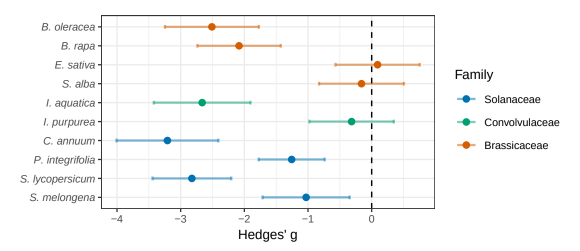


Figure 3: Average effect size and confidence intervals (95%) of the 9 heterospecific treatments of 50% pollen to the different 10 focal species coloured by family.

Mantel test indicates that a possible?? It exists! correlation exist between heterospecific pollen effect and the evolutive relative distances, for ITS and RBCL markers we had r coefficients of 0.29 and 0.25 respectively p<0.05 think on a figure - maybe using NMDS. Moreover, Mantel test indicates that also a possible?? correlation between stigma width and stigma type exist (stats??). Trait correlations were also explored with GLMM

I have done it at the moment just for Compatibility system Also I have to fix from mixed linear model to GLMM, just realize that

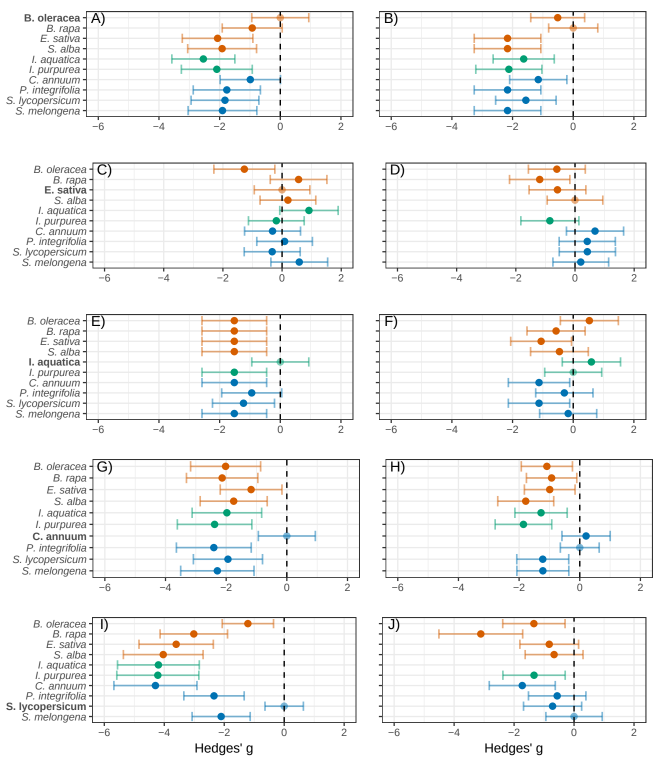


Figure 4: Effect sizes for the 10 different species. The different families appear with different colours, when a species was focal was coloured differently from its family.

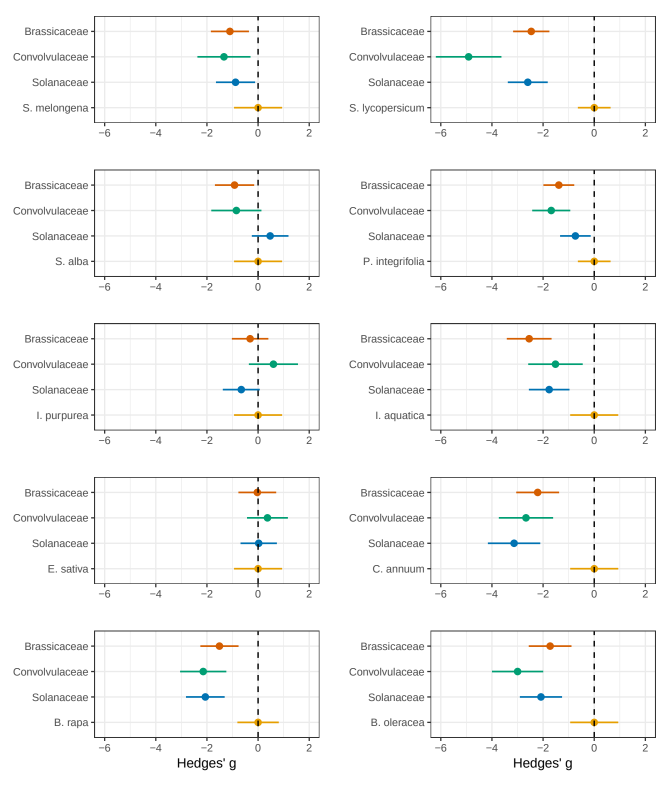


Figure 5: A

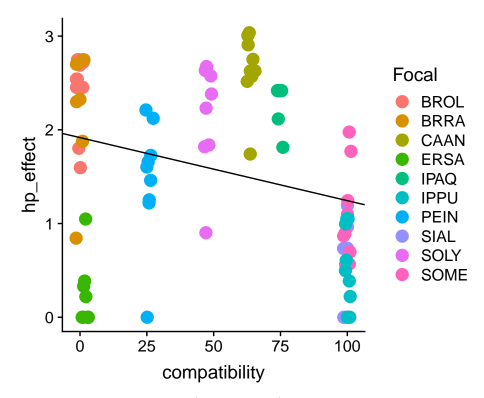


Figure 6: The effect of heterospecific pollen (scaled see set) is represented in function of the compatibility system (self/cross*100) for the different species. Each coulored dot represents the interaction of a focal species with a different pollen donor.

Compatibility index don't multiply per 100 from Lloyd

208 DISCUSSION

Discussion Herbs vs tress, annual vs perennial... Many flowers vs few flowered species; structural composition on 210 a system 211 What are the implications of the findings? Ideas about pollen size in heterospecific pollen effect. (still have to develop it more...) Let's classify pollen size in three groups in order to understand the interaction between pollen donor 214 and recipient: 1) Donor pollen size < Recipient pollen size 2) Donor pollen size = Recipient pollen size 215 3) Donor pollen size > Recipient pollen size Now I try to develop each part 217 1) Donor pollen size < Recipient pollen size 218 Effect: • Donor's pollen could clogg the stigma 220 • Chemical inhibition 221 Traits associated with bigger pollen of the recipient: 222 • Recipient's pollen have faster pollen tube growth (example with my data) 223 • Reduction in number of ovules (Also with my species) 224 • Big differences in pollen size can be traduced in low relatednes therefore less likely of pollen 225 germination on a far related stigma. 226

16 March 11, 2019

2) Donor pollen size = Recipient pollen size

• Very relatedness dependant this point

227

228

- Similar probabilities of taken space on the stigma
- 3) Donor pollen size > Recipient pollen size
- 231 Effect:

229

- 232 -In small stigmas big pollen grains can occupy great part of the stigmatic area.
- -small pollen grains can get embeded
- ²³⁴ IB: Think also on using tree analysis to test if hp effect depends on complex trait combinations. Tree
- 235 analysis are great when two different strategies lead to the same outcome. This would never been pick
- 236 up by GLMs. The r package is party{} . You can see an example applied to birds is Sol et al 2010
- 237 Science. Ask me if you want more details or code examples.

238 CONCLUSIONS

239 ACKNOWLEDGEMENTS

240 REFERENCES

- Aizen, M. A., and L. D. Harder. 2007. Expanding the limits of the pollen-limitation concept: Effects of
- pollen quantity and quality. Ecology 88:271–281.
- ²⁴³ Arceo-Gómez, G., and T.-L. Ashman. 2011. Heterospecific pollen deposition: Does diversity alter the
- consequences? New Phytologist 192:738–746.
- ²⁴⁵ Arceo-Gómez, G., and T.-L. Ashman. 2016. Invasion status and phylogenetic relatedness predict cost
- of heterospecific pollen receipt: Implications for native biodiversity decline. Journal of Ecology
- 247 104:1003-1008.
- Ashman, T.-L., and G. Arceo-Gómez. 2013. Toward a predictive understanding of the fitness costs of

- ²⁴⁹ heterospecific pollen receipt and its importance in co-flowering communities. American Journal of
- 250 Botany 100:1061–1070.
- Bartomeus, I., J. Bosch, and M. Vilà. 2008. High invasive pollen transfer, yet low deposition on native
- 252 stigmas in a carpobrotus-invaded community. Annals of Botany 102:417–424.
- ²⁵³ Carvalheiro, L. G., J. C. Biesmeijer, G. Benadi, J. Fründ, M. Stang, I. Bartomeus, C. N.
- ²⁵⁴ Kaiser-Bunbury, M. Baude, S. I. Gomes, V. Merckx, and others. 2014. The potential for indirect effects
- between co-flowering plants via shared pollinators depends on resource abundance, accessibility and
- relatedness. Ecology letters 17:1389–1399.
- Engel, E. C., and R. E. Irwin. 2003. Linking pollinator visitation rate and pollen receipt. American
- 258 Journal of Botany 90:1612–1618.
- ²⁵⁹ Fang, Q., and S.-Q. Huang. 2013. A directed network analysis of heterospecific pollen transfer in a
- biodiverse community. Ecology 94:1176–1185.
- Galen, C., and T. Gregory. 1989. Interspecific pollen transfer as a mechanism of competition:
- 262 Consequences of foreign pollen contamination for seed set in the alpine wildflower, polemonium
- viscosum. Oecologia 81:120–123.
- Inouye, D. W. 1980. The terminology of floral larceny. Ecology 61:1251–1253.
- Letten, A. D., and W. K. Cornwell. 2015. Trees, branches and (square) roots: Why evolutionary
- relatedness is not linearly related to functional distance. Methods in Ecology and Evolution 6:439–444.
- Lloyd, D. G., and D. J. Schoen. 1992. Self-and cross-fertilization in plants. i. functional dimensions.
- ²⁶⁸ International Journal of Plant Sciences 153:358–369.
- Magrach, A., J. P. González-Varo, M. Boiffier, M. Vilà, and I. Bartomeus. 2017. Honeybee spillover
- 270 reshuffles pollinator diets and affects plant reproductive success. Nature ecology & evolution 1:1299.
- Montgomery, B. R., and B. J. Rathcke. 2012. Effects of floral restrictiveness and stigma size on

- 272 heterospecific pollen receipt in a prairie community. Oecologia 168:449–458.
- Morales, C. L., and A. Traveset. 2008. Interspecific pollen transfer: Magnitude, prevalence and
- 274 consequences for plant fitness. Critical Reviews in Plant Sciences 27:221–238.
- Murphy, S. D., and L. W. Aarssen. 1995. Reduced seed set in elytrigia repens caused by allelopathic
- pollen from phleum pratense. Canadian Journal of Botany 73:1417–1422.
- Neiland, M., and C. Wilcock. 1999. The presence of heterospecific pollen on stigmas of nectariferous
- ²⁷⁸ and nectarless orchids and its consequences for their reproductive success. Protoplasma 208:65–75.
- Pauw, A. 2013. Can pollination niches facilitate plant coexistence? Trends in ecology & evolution
- 280 28:30-37.
- ²⁸¹ R Core Team. 2018. R: A language and environment for statistical computing. R Foundation for
- 282 Statistical Computing, Vienna, Austria.
- Thomson, J. D., B. J. Andrews, and R. Plowright. 1982. The effect of a foreign pollen on ovule
- development in diervilla lonicera (caprifoliaceae). New Phytologist 90:777–783.
- ²⁸⁵ Tong, Z.-Y., and S.-Q. Huang. 2016. Pre-and post-pollination interaction between six co-flowering
- pedicularis species via heterospecific pollen transfer. New Phytologist 211:1452–1461.
- Waser, N. M., L. Chittka, M. V. Price, N. M. Williams, and J. Ollerton. 1996. Generalization in
- pollination systems, and why it matters. Ecology 77:1043–1060.
- ²⁸⁹ Whitehead, M. R., R. Lanfear, R. J. Mitchell, and J. D. Karron. 2018. Plant mating systems often vary
- ²⁹⁰ widely among populations. Frontiers in Ecology and Evolution 6:38.
- Williams, E., and J. Rouse. 1990. Relationships of pollen size, pistil length and pollen tube growth
- rates in rhododendron and their influence on hybridization. Sexual Plant Reproduction 3:7–17.

293 APPENDIX

294 1.

Table S1. Perecentage of seeds produced per ovule for the ten species used in the experiment. The treatments presented are hand cross-pollination, hand self-pollination, natural selfing and apomixis (emasculated flowers).

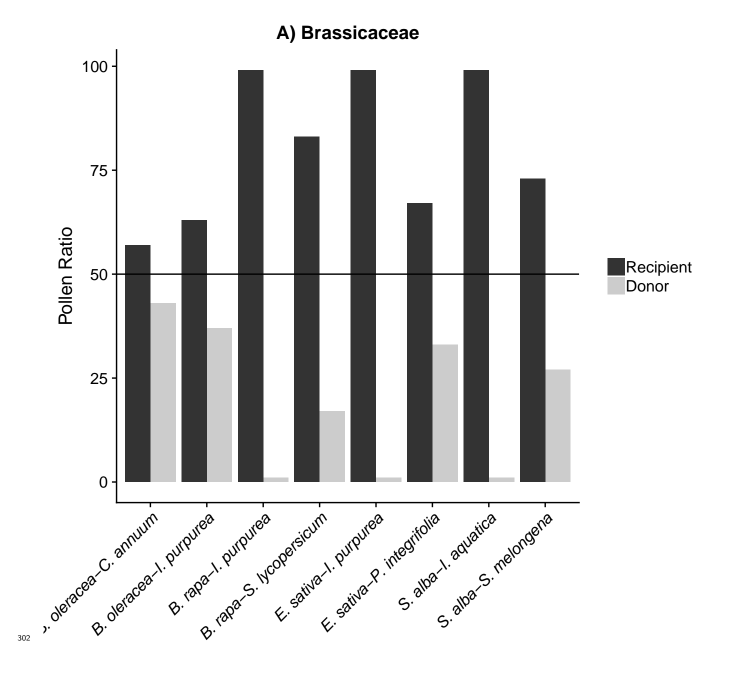
Species	Cross	Self	Natural_selfing	Apomixis
Brassica oleracea	32.06897	0.0000000	0.00000	0
Brassica rapa	44.97041	0.0000000	0.00000	0
Eruca versicaria	23.75000	0.4166667	0.00000	0
Sinapis alba	43.33333	48.3333333	5.00000	15
Ipomoea aquatica	40.00000	30.0000000	20.00000	0
Ipomoea purpurea	31.66667	86.6666667	31.66667	0
Capsicum annuum	100.00000	66.2240664	23.48548	0
Petunia integrifolia	100.00000	24.7727273	0.00000	0
Solanum lycopersicum	90.38043	43.4782609	70.00000	0
Solanum melongena	60.47525	87.9702970	21.56436	0

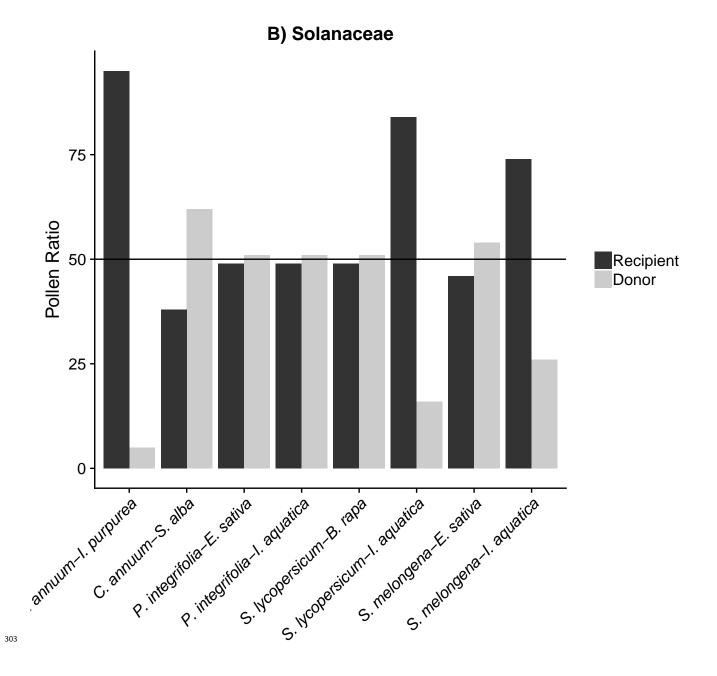
298 2.

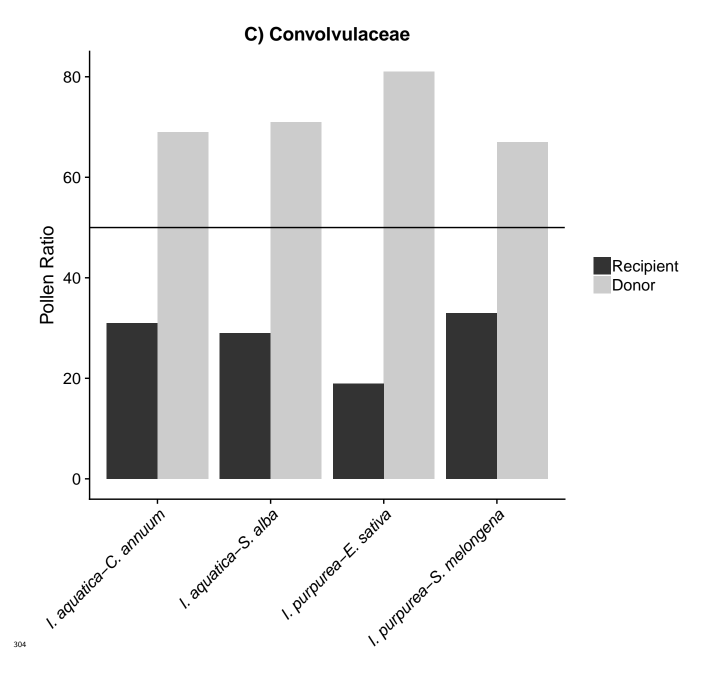
Figure S1. Pollen ratios separated by family A) Brassicaceae, B) Solanaceae, C) Convolvulaceae.

 $^{300}\,$ Pollen was counted for crosses with species of the other two families (N=3). To prepare these ratios all

the pollen grains on the stigma were counted.







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