Effect of heterospecific pollen deposition on pollen tube growth depends on the phylogenetic relatedness between donor and recipient

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Abstract

Co-flowering plant species may interact via pollinators leading to heterospecific pollen transfer with consequences for plant reproduction. What determines the severity of heterospecific pollen effect on conspecific pollen performance is unclear, but it may depend on the phylogenetic relatedness of the interactors (pollen donors and recipient). The heterospecific pollen effect might also depend on the extent to which plants are exposed to heterospecific pollen over ecological or evolutionary timescales. For instance, generalist-pollinated plant species might tolerate heterospecific pollen more than specialists. Here, we tested whether heterospecific pollen effects are stronger between closely related species than phylogenetically distant ones in a tropical highland community. Then, based on these results, we determined whether responses to heterospecific pollen were stronger in generalized vs. specialized plant species. We applied heterospecific pollen from close (congeneric) or distant (different families) donors alone or with conspecific pollen on stigmas of three recipient species (one generalist, Sisyrinchium wettsteinii; and two specialists, Fuchsia campos-portoi and Fuchsia regia) and scored pollen tube performance in styles. In all species, pollen from closely related donors grew pollen tubes to the base of the style indicating a high potential to interfere with seed set. Conversely, distantly related heterospecific pollen had no effect on either specialist Fuchsia species, whereas enhanced performance of conspecific pollen was observed in generalist S. wettsteinii. The strong effect of phylogenetic relatedness of donor and recipient might have obscured the role of pollination specialization, at least for the three species examined here. Therefore, phylogenetic relatedness mediated the effect of heterospecific pollen on post-pollination success, with possible consequences for reproductive trait evolution and community assembly for further studies to explore.

Keywords: Competition; facilitation; interspecific pollen transfer; pollen germination; pollinator sharing.
Introduction

Most flowering plants rely on animal pollinators to transfer their pollen grains (Ollerton et al. 2011) to conspecific stigmas and set seeds. Plants sharing pollinators may compete or facilitate pollinator visits (i.e. pre-pollination interactions; Rathcke 1983; Moeller 2004; Mitchell et al. 2009). If two species exploit the same pollinator, interspecific pollinator movements can result in conspecific pollen (hereafter CP) loss and heterospecific pollen (hereafter HP) deposition on stigmas (i.e. post-pollination interactions) with a potential impact on the reproductive success of both (Morales and Travesset 2008; Ashman and Arceo-Gómez 2013; Moreira-Hernández and Muchhala 2019). Hence, plant–plant interactions via pollinators are traditionally interpreted as negative, at least for the donor’s perspective (male component), because loosing CP is always a waste of gametes that could otherwise affect conspecific reproduction (Waser 1978; Bell et al. 2005). Historically, HP receipt (female component) was also mainly interpreted as negative because loss of seed fitness is even more costly than loss of pollen grains (Rathcke 1983). However, once pollen grains are on heterospecific stigmas they can encounter diverse barriers that can lead pollen tubes to different fates (Swanson et al. 2004). So even though there is considerable evidence that HP can reduce recipient fitness (e.g. Da Silva and Sargent 2013; Briggs et al. 2015; Fonseca et al. 2016; Arceo-Gómez et al. 2018), there is also evidence that HP has no effect on recipient reproduction (e.g. Kohn and Waser 1985; Caruso and Alfaro 2000; Montgomery 2009). The lack of a consistent pattern may be related to the fact that the outcome of the interaction does not depend solely on the recipient, but also on interactive effects of recipient with HP donors (Arceo-Gómez et al. 2019). Assessing what underlies the prevalence and strength of HP effects on post-pollination success is essential to understand its role in shaping flowering communities.

Heterospecific pollen can impact recipient plants by physically blocking the stigma (Waser and Fugate 1986; Galen and Gregory 1985) and/or interfering with CP performance (Arceo-Gómez and Ashman 2011). The interference can be through allelopathic effects that retard CP tube or ovule growth (Sukhada and Chandra 1980; Thomson et al. 1981) or through HP fertilizing conspecific ovules (Harder et al. 1993; Burgess et al. 2008). The latter is most common among closely related species and can, ultimately, result in hybridization (Wendt et al. 2001; Wendt et al. 2008; Arceo-Gómez and Ashman 2011). As pollen–pistil interactions may be compatible between species with recent evolutionary history (Moreira-Hernández and Muchhala 2019), those with distant history face considerable morphological and genetical incongruities that preclude much HP development after deposited on stigmas (Hogenboom 1975). Thus, we predict HP effects will increase with decreasing recipient-donor relatedness (Ashman and Arceo-Gómez 2013; Arceo-Gómez and Ashman 2015). Moreover, intrinsic traits of recipients can mediate HP effects, such as its mating system since self-incompatible species possess stronger barriers than self-compatible ones to improper pollen growth (Harder et al. 1993; Ashman and Arceo-Gómez 2015). The degree to which recipients restrict pollinator accessibility to flowers could also correlate with tolerance to HP receipt. In this sense, generalist-pollinated species (i.e. that exploit a wide variety of pollinators; Ollerton et al. 2007) might not be impaired as much by HP as specialist-pollinated ones, because the former were presumably exposed more often and to higher and more diverse loads of HP over generations (Fang and Huang 2013; Arceo-Gómez et al. 2016; Fang et al. 2019). Donor traits like HP size and apertures (Ashman and Arceo-Gómez 2013), as well as extrinsic factors such as HP arrival time on stigmas (Suárez-Mariño et al. 2019), abiotic conditions (Celaya et al. 2015) and HP load diversity and identity (Arceo-Gómez and Ashman 2011) also are known to contribute on recipient’s post-pollination outcomes.

Species floral traits influence how plants exploit pollinators and consequently also affect the likelihood of HP transfer (Minnaar et al. 2019). For instance, species with specialized floral morphology constrain accessibility to only few pollinators that provide the best pollen transfer (Stebbins 1970; Ollerton et al. 2007), reducing the chance of HP deposition. Nevertheless, generalist-pollinated flowers are the majority in flowering communities (Waser et al. 1996; Memmott 1999; Olesen and Jordano 2002), making HP deposition a frequent phenomenon, even in spite of all mechanisms to control HP transfer (McLernon et al. 1996; Montgomery and Rathcke 2012; Fang and Huang 2013). But the effects vary between species and thus selection to avoid HP is expected to vary. Hence, species might possess mechanisms that act filtering only HP that is indeed detrimental to recipients (Ashman and Arceo-Gómez 2013) or even compensate for recipient losses by maximizing pollen dispersal to conspecifics (Muchhala et al. 2010). Moreover, the benefits of sharing pollinators with heterospecifics via facilitation may outweighs HP costs (Tur et al. 2016) indicating that species with overlapping pre-pollination mechanisms may be unaffected by receipt of each other’s pollen (Gross et al. 2000). Thus, to fully understand the magnitude of HP as an evolutionary force driving floral trait divergence or community assembly, we need to consider its costs on both female and male reproductive components as well as in a community context (Muchhala et al. 2010).

Here, we used an experimental approach to assess the post-pollination HP effects between sets of species with similar flowers that share pollinators in a tropical highland community. The species studied here were categorized as specialist- or generalist-pollinated depending on whether they were visited by one or more than one group of pollinators, respectively (see Bergamo et al. 2020a). The taxonomic classification of these species (genus and family) was used to establish pairs of closely related taxa (species in the same genus) vs. more distantly related (species in different families). Our main goal was to test whether the effects of HP differ with phylogenetic distance between donor and recipient. Because there are fewer post-pollination barriers between closely related species, we hypothesize that HP effects are stronger between closely related species than between phylogenetically distant ones (Ashman and Arceo-Gómez 2013; Arceo-Gómez and Ashman 2016). As we choose species with distinct pollination systems to address this question, we used the results from experiments to establish whether recipiente respond differently depending on their likelihood of receiving HP. Since species pollinated by numerous animals can receive more HP (Fang and Huang 2013; Arceo-Gómez et al. 2016), we hypothesize that generalist-pollinated species are more able to tolerate the presence of HP than specialist-pollinated species, and thus suffer less impact by HP receipt.

Materials and Methods

Study site

The study was conducted in the plateau of the Itatiaia National Park (22°21′S, 44°40′W) that is in the Atlantic forest domain,
Selection of studied species

We conducted hand pollination experiments with three trios of sympatric co-flowering species (Fig. 1; Table 1). Each trio was composed of (i) one pollen recipient species, (ii) one HP donor phylogenetically close to the recipient species and (iii) one HP donor phylogenetically distant but with phenotypically similar flowers. We considered congeners as phylogenetically close and species from different families as phylogenetically distant. Based on this assumptions, the recipients chosen included two *Fuchsia* hummingbird-pollinated species, considered as specialists (previously classified in Bergamo et al. 2020a): *Fuchsia campos-portoi*, that has 113.7 ± 20.83 (mean ± standard deviation) ovules per ovary and a stigmatic surface area of 0.01 cm² and *F. regia*, that has 122.38 ± 25.23 (mean ± standard deviation) ovules per ovary and a stigmatic surface area of 0.05 cm². The other recipient was *Sisyrinchium wettsteinii* that has 60.5 ± 16.8 (standard deviation) ovules per ovary, a stigmatic surface area of 0.001 cm² and is visited by various insects (i.e. bees, flies and beetles), considered a generalist-pollinated species (previously classified in Bergamo et al. 2020a). Besides fitting the assumptions described before, these species were chosen to be recipients also due to their high flower abundance in field, their pollination system and the fact that they experience HP deposition naturally (e.g. 52 % of *F. campos-portoi*, 50 % of *F. regia* and 53 % of *S. wettsteinii* stigmas received HP in the field; N. S. Streher et al., unpubl. data). Species used as pollen donors were *Barbacenia gounelleana*, *Oxalis confertissima*, *Sisyrinchium glaziovii* and *F. campos-portoi* and *F. regia*. *Barbacenia gounelleana*, *S. glaziovii* and *O. confertissima* were used as pollen donors but not as recipients because they did not fit our requirements to be a recipient model in this study.

Experimental treatments

Five types of hand pollinations were conducted in the field: (i) outcross conspecific pollen (CP); (ii) heterospecific pollen from the phylogenetically distant species (HP distant); (iii) heterospecific pollen from the phylogenetically close species (HP close); (iv) mixture of outcross (CP + HP distant); (v) mixture of outcross (CP + HP close). Pollination by pure loads of HP was conducted to assess post-pollination barriers and to aid in the interpretation of the results of mixtures as pollen grains from congeners are usually hard to distinguish and once pollen tubes grow into the style species identity is unknown. Pistils were fixed in 50 % FAA (formalin-acetic acid-alcohol) solution (Johansen 1940) 24 h after hand pollinations since pilot experiment demonstrated that this was enough time for pollen tubes to reach style base in all species. Fixed materials were cleared with NaOH 9 N, heated at 60 °C for 20 min, stained with blue aniline and observed in a fluorescent microscope (Martin 1959). For each pistil, we counted the number of pollen grains deposited on stigmas, the number of pollen grains germinated in stigmas, pollen tubes at

Figure 1. Flowers of species used in experiments of heterospecific pollen effects. (A) *Fuchsia campos-portoi*. (B) *Fuchsia regia*. (C) *Barbacenia gounelleana*. (D) *Sisyrinchium wettsteinii*. (E) *Sisyrinchium glaziovii*. (F) *Oxalis confertissima*. Bar = 0.5 cm.
the tip and base of styles. Based on what we scored, we refer to post-pollination success as how many of the CP grains adhered to stigmas successfully develop further along the style.

Buds were bagged to avoid pollen contamination from visitors. Because anthers and stigmas in S. wettsteinii flowers are positioned close to each other we emasculated them in bud prior to experiments to avoid self-pollen contamination. Emasculation was not necessary for Fuchsia species because their flowers are protogynous (stigmas are receptive before anthers dehiscence preventing self-pollination). Pollen applied was fresh which means that recipient and donor species were flowering at the same time during experiments in the field. To standardize our method of pollen transfer we touched one anther per individual donor in each recipient stigma. We used three individuals as CP donors and only one as HP donor so mixed treatments had ca. 75 % CP and 25 % HP mix of pollen grains. Heterospecific pollen receipt in natural communities varies extensively, with most species receiving on average 20 % of HP (see Ashman and Arceo-Gomez 2015; Fang and Huang 2013). Because of that, we checked the HP naturally deposited in flowers of the species studied here during one flowering season. The HP deposition varied from 0–16 % in F. campos-portoi, 0–2 % in F. regia and 0–100 % in S. wettsteinii and the mean percentage of HP loads in flowers was 4 %, 0.8 % and 27 % for each species, respectively (N. S. Streher et al., unpubl. data). Hence, the usual ratio 50:50 of CP:HP used in experiments seemed too high in general for these species (even for the generalist-pollinated S. wettsteinii), leading us to use the ratio 75:25. In these treatments, CP was applied first and HP immediately after. Differences in pollen size and pollen adherence capability among species may have influenced the final pollen load (see Table 2 for pollen load). To avoid self-incompatibility reactions, the individuals that were used as CP donors were always separated by at least 200 m from the recipient.

**Statistical analysis**

To evaluate the effect of pollination treatments on recipient post-pollination success, we fitted generalized linear mixed models (GLMMs) using binomial distributions in the glmmTMB package (Brooks et al. 2017) in R v. 3.5.1 (R Core Team 2018). We fitted models for each of the three response variables resulting in three models for each recipient species. The response variable of each model was a matrix containing the total pollen deposited in stigma and the pollen response resulting from this deposition in the different portions of the same pistil (pollen grains germinated in the stigma, pollen tubes at the tip of the style and pollen tubes at the base of the style). This model accounted for variation in the amount of pollen deposited on the stigmas. For all models, pollination treatments were the fixed effect and individuals were included as a random effect. Model assumptions were checked graphically. To verify the significance of each model, we compared the built models with a null model that included the respective response variable and only the intercept. Since we were specifically interested in comparing treatment effects relative to outcross CP, we performed a post hoc test using the package emmeans (Lenth et al. 2020).

**Results**

Pollination treatments influenced recipient post-pollination success revealing different outcomes depending whether HP was present and its source. Pollen performance was explained by differences among treatments since all models performed better than the null models (Table 3). Model comparisons within each pistil portion, described below, were interpreted relative to CP treatment as this corresponds to the ideal situation for reproduction (i.e. only conspecific and no foreign pollen grains on stigmas; see Table 4 for more).

**Recipient F. campos-portoi (specialist-pollination)**

At the stigma, treatments containing HP from phylogenetically distant species showed greater probabilities of pollen germination (>78 %) compared to treatment containing only CP (HP distant, \( t = -7.460, df = 99, P < 0.001 \); CP + HP distant, \( t = -10.064, df = 99, P < 0.001 \); Fig. 2). On the other hand, treatments with HP

| Table 1. Species used as pollen recipient in the experiments, their floral phenotypes and the identity of heterospecific pollen donors. |
|-----------------|------------------|-----------------|-----------------|
| Pollen recipient species | Floral phenotype | HP donor species |
|-----------------|------------------|-----------------|
| Fuchsia campos-portoi (Onagraceae) | Specialist (hummingbird-pollinated) | Fuchsia regia (Onagraceae) | Barbacenia gounelleana (Velloziaceae) |
| Fuchsia regia (Onagraceae) | Specialist (hummingbird-pollinated) | Fuchsia campos-portoi (Onagraceae) | Barbacenia gounelleana (Velloziaceae) |
| Sisyrinchium wettsteinii (Iridaceae) | Generalist (various insects) | Sisyrinchium glaziovii (Iridaceae) | Oxalis conspersa (Oxalidaceae) |

| Table 2. Number of pollen grains applied in each treatment for each recipient species (mean ± standard deviation). The number of recipients used in each treatment are in parenthesis. |
|-----------------|-----------------|-----------------|-----------------|
| Treatment | Fuchsia campos-portoi | Fuchsia regia | Sisyrinchium wettsteinii |
| CP | 128.05 ± 45.55 (20) | 293.81 ± 105.62 (22) | 81.09 ± 51.21 (21) |
| HP close | 135.913 ± 109.76 (23) | 280.61 ± 97.20 (13) | 52.34 ± 28.84 (41) |
| CP + HP close | 164.58 ± 77.48 (24) | 340.5 ± 147.48 (14) | 80.67 ± 37.37 (28) |
| HP distant | 83.15 ± 62.48 (19) | 119.166 ± 176.58 (8) | 8.68 ± 11.64 (32) |
| CP + HP distant | 106.7 ± 45.63 (20) | 200.76 ± 124.20 (14) | 45.64 ± 34.59 (17) |
from congeners lead to lower probabilities of pollen germination (<45%), being worse than CP alone (HP close, \( t = 9.954, df = 99, P < 0.001 \); CP + HP close, \( t = 13.199, df = 99, P < 0.001 \); Fig. 2). At the tip of the style, the only treatment that had a similar probability of pollen tubes as CP was the one containing the mix of CP + HP (\( t = -1.375, df = 99, P = 0.645 \)), while all other treatments had lower probabilities (Fig. 2). This pattern remained when we looked at pollen tubes in the base of the style (CP + HP distant, \( t = -2.453, df = 99, P = 0.110 \); Fig. 2).

**Recipient Fuchsia regia (specialist-pollination)**

For this species, it is worth noticing that CP grains had the lowest germination probability on the stigma of all treatments (27%), being different from both treatments containing HP from congeners (HP close, \( t = -5.371, df = 64, P = 0.013 \) and CP + HP close, \( t = -15.625, df = 64, P < 0.001 \)), and also from CP + HP distant (\( t = -14.837, df = 64, P < 0.001 \); Fig. 2). Once pollen tubes entered the style, all treatments had similar probabilities as CP alone (20%), except HP distant (with only 2%, \( t = 5.333, P < 0.001 \); Fig. 2). All treatments ended up showing similar probabilities than CP (that itself had a small probability, only 7%) of having pollen tubes in the end of the style (CP close, \( t = 0.543, df = 63, P = 0.982 \); CP + HP close, \( t = 2.526, df = 63, P = 0.098 \); HP distant, \( t = 1.895, df = 63, P = 0.331 \) and CP + HP distant, \( t = -0.891, df = 63, P = 0.899 \); Fig. 2).

**Recipient S. wettsteini (generalist-pollination)**

All treatments had high probabilities of pollen germination on the stigma (75–99%; Fig. 2), but only CP + HP distant was high as CP (\( t = -0.738, df = 132, P = 0.947 \)). Within the tip of the style, most treatments had lower probabilities than CP, except CP + HP distant that had a greater probability (\( t = -5.822, df = 132, P < 0.001 \); Fig. 2). Once pollen tubes reached the base of the style, the treatment containing only HP distant continued to show a lower probability relative to CP (\( t = 6.396, df = 132, P < 0.001 \)) and both treatments containing CP close had similar probabilities to CP (HP close, \( t = 0.879, df = 132, P = 0.904 \); CP + HP close, \( t = 0.022, df = 132, P = 0.899 \); Fig. 2).

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**Table 3. Comparisons between outcross/pollinator (CP) and the other treatments. EMM = estimated marginal means; SE = standard error. Bold values indicate significant effects at \( P < 0.05 \).**

| Treatment | EMM | SE  | t     | P   |
|-----------|-----|-----|-------|-----|
| CP close  | 0.724 | 0.073 | 9.954 | <0.0001 |
| CP + HP close | 0.881 | 0.067 | 13.199 | <0.0001 |
| HP distant | -0.806 | 0.108 | -7.460 | <0.0001 |
| CP + HP distant | -1.028 | 0.102 | -10.064 | <0.0001 |
| CP close  | 0.647 | 0.074 | 8.688 | <0.0001 |
| CP + HP close | 0.768 | 0.066 | 11.596 | <0.0001 |
| HP distant | 2.603 | 0.294 | 8.864 | <0.0001 |
| CP + HP distant | -0.126 | 0.091 | -1.375 | 0.645 |

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**Table 4. Contrasts between outcross conspecific pollen (CP) and the other treatments. EMM = estimated marginal means; SE = standard error. Bold values indicate significant effects at \( P < 0.05 \).**

| Treatment | EMM | SE  | t     | P   |
|-----------|-----|-----|-------|-----|
| CP close  | 0.441 | 0.120 | 3.678 | 0.003 |
| CP + HP close | 0.886 | 0.118 | 7.340 | <0.0001 |
| HP distant | 4.103 | 0.551 | 7.443 | <0.0001 |
| CP + HP distant | -0.284 | 0.116 | -2.453 | 0.022 |

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**Recipient Fuchsia regia**

**Table 5. Comparisons between outcross conspecific pollen (CP) and the other treatments. EMM = estimated marginal means; SE = standard error. Bold values indicate significant effects at \( P < 0.05 \).**

| Treatment | EMM | SE  | t     | P   |
|-----------|-----|-----|-------|-----|
| CP close  | 0.557 | 0.104 | 5.371 | <0.0001 |
| CP + HP close | -1.546 | 0.099 | -15.625 | <0.0001 |
| HP distant | -0.319 | 0.161 | -1.988 | 0.284 |
| CP + HP distant | -1.774 | 0.12 | -14.837 | <0.0001 |
| HP close  | 0.266 | 0.124 | 2.144 | 0.151 |
| CP + HP close | 0.24 | 0.114 | 2.098 | 0.323 |
| HP distant | 2.453 | 0.460 | 5.333 | <0.0001 |
| CP + HP distant | -0.284 | 0.116 | -2.453 | 0.022 |

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**Recipient S. wettsteini**

**Table 6. Comparisons between outcross conspecific pollen (CP) and the other treatments. EMM = estimated marginal means; SE = standard error. Bold values indicate significant effects at \( P < 0.05 \).**

| Treatment | EMM | SE  | t     | P   |
|-----------|-----|-----|-------|-----|
| CP close  | 0.633 | 1.167 | 0.543 | 0.582 |
| CP + HP close | 2.579 | 1.021 | 2.526 | 0.098 |
| HP distant | 2.769 | 1.461 | 1.895 | 0.331 |
| CP + HP distant | -0.911 | 1.023 | -0.891 | 0.389 |
Figure 2. Probability of pollen performance of each treatment in the three portions of each recipient’s pistil (stigma, style tip and style base). In each block, different letters indicate significant differences at P < 0.05.

P = 1.0). At this point, CP + HP distant continued to perform better than CP alone (t = −5.252, df = 132, P < 0.001), with ca. 23 % greater chance of pollen tubes reaching the base of the style (Fig. 2).

Discussion

Our results reinforce the idea that when species from the same genus are interacting via pollination, HP has a stronger negative impact on recipient (Ashman and Arceo-Gómez 2013; Arceo-Gómez and Ashman 2016). However, when the interaction is between distantly related species, recipients can tolerate or even benefit at the pollen tube stage from sharing pollinators with heterospecifics. As phylogenetic identity of HP source affects responses, the contribution of pollination systems was conditioned to that. Even though, our results indicate that both generalist- and specialist-pollinated species can tolerate HP in some level. The magnitude of HP response depends on the interactive effects between donor and recipient (Arceo-Gómez et al. 2019) and here we provide evidence that species phylogenetic relatedness is one of the factors involved in the complex equation of plant–plant post-pollination interactions.

The effects of HP considering phylogenetic distance between donor and recipient

Even though we cannot differentiate which pollen tubes are from HP close and which ones are CP in the same style, our experiments using only HP close reveal that there are chances of hybridization between congers. The lack of strong post-pollination barriers for HP close, as the ones that we present here, does not necessarily mean that these pollen tubes will fertilize the ovules. However, if the ovules are usurped by HP tubes this can be extremely costly to recipient plants since these ovules will no longer be available for CP tubes (Levin et al. 1996; Burgess et al. 2008). HP close tubes reach the base of the style in all species, but performances were different (greater or lesser probability of pollen-tube growth) depending on the specific treatment. We can check these differences by comparing the CP + HP close treatment with the treatments CP alone and HP close alone within each portion of the style. These comparisons allowed us to interpret which interactions modulated the final costs in each recipient species. In S. wettsteinii, the interaction between CP and HP close is competitive in stigmas, relaxing along the style since pollen tubes had similar probabilities of reaching style base in the three treatments. This possibly indicates a lack of incompatibility between these two species (S. wettsteinii and S. glaziouii), which might be a by-product of the recent speciation process of the genus (Chauveau et al. 2011). The competition is also strong during pollen germination in F. campos-portoi, but in this species it continues intense until the style base where both CP + HP close and HP close alone performed worse than CP alone. In F. regia stigmas, there is no apparent competition between CP and HP close germination; however, most pollen tubes of CP + HP close treatment are blocked prior to the first portion of style. Hence, their probabilities of success decrease from 64 % in stigmas to less than 1 % in style base, indicating that style strongly sieves pollen tubes. Considering how Fuchsia species impact each other reproduction, it is possible to see that when pollen tubes get in the style base, F. regia pollen plainly decreases F. campos-portoi post-pollination success. The reverse cross (F. campos-portoi donating pollen to F. regia) is more complex to interpret since all treatments (including the ones with HP distant) had slightly the same way, but based on the already mentioned substantial decrease of CP + HP close performance since arriving in stigma till style base, it also seems to be negative. Hence, it is likely that these responses are driven by an active mechanism (de Nettancourt 1977) rather than by incongruity since CP tubes are also being blocked. Nevertheless, further studies should consider applying genetic markers to check the strength of HP close in siring seeds in recipients, because some species might show a conspecific advantage over heterospecific in fertilizing ovules (Arnold 1997; Campbell et al. 2008) which then changes the signal of the interaction between closely related species to positive.
Heterospecific pollen distantly related (HP distant) germinated in all species when applied alone in stigmas, against general expectations (Martin 1970; Moreira-Hernández and Muchhala 2019). In fact, HP distant treatment had high probabilities of germination (especially in F. campos-portoi), which may suggest that in these species the stigmas themselves do not function to select pollen. This role seems to be played by the tip of the style where most of these pollen tubes are arrested, probably due to the lack of recognition resulted from the genetic distance (Hogenboom 1975). As they rarely got into the style (<4 %), this means that we observed mainly CP tubes in the final portion of styles in the treatment that combined CP + HP distant. Hence, HP can germinate even in phylogenetically distant species and have neutral (Fuchsia species) or even positive (S. wettsteinii) effects on recipient post-pollination success (pollen tubes in style end). A similar result was reported for Canikle edentula that when received HP of Bidens pilosa also increased CP tube growth, which was suggested to be stimulated by the release of biochemical compounds and result in a herd effect (Suárez-Martíno et al. 2019).

In the context of pollination, the herd effect can be interpreted as the greater CP growth when in the presence of foreign pollen (HP in our case); however, this is an idea that has yet to be formally tested (Ashman et al. 2020). In this sense, when greater CP tubes number reach style base they are the result of recipient interaction with heterospecifics early in the stigma.

One important factor that might influence the extension of HP effect on distantly related species is how well pollen grains can adhere to recipients' stigmatic surface. In the case of S. wettsteinii as recipient, only a small percentage of O. confertissima pollen adhered to its stigmas suggesting a weak attachment between the two species, which can be essential to avoid stigma blocking and its detrimental effects. Plus, the positive effect from their interaction could be due to their long history of coexistence. For instance, it has been shown that the opposite (new interactions between distantly related species) can lead to negative effects on recipient, like the ones provoked by only a few Zea mays pollen grains on Mimulus guttatus female fitness (Arceo-Gómez et al. 2018). Nevertheless, a previous study showed that other Sisyrinchium species (S. campestre) was not affected by a distantly related pollen donor (Euphorbia esula) invasive to the community (Montgomery 2009). These results taking together could indicate that species that encounter HP often in the evolutionary time (i.e. unrestricted generalist-pollinated flowers) may have evolved mechanisms for tolerating it regardless of which species they are interacting with. However, whether the response is due to floral exposure to HP and detached from the history of coexistence with donors still needs to be formally tested.

The effects of HP considering recipient's pollination system

The role of pollination systems (i.e. specialized or generalized) determining the degree of plant response to HP remains an area in need of more study. Indeed, in here, recipients responded differently which might be associated with their historical exposure to HP but there was no consistent pattern within each category. Sisyrinchium wettsteinii, that has unrestricted flowers visited by various groups of insects characterizing a generalist-pollination system (functional pollination sensu Ollerton et al. 2007), not only tolerate, as our hypothesis predicted, but can also respond positively to HP. On the other hand, both Fuchsia species, that show restrictive flowers pollinated by hummingbirds, being more specialized in the spectrum of plant-pollinator interactions (functional pollination sensu Ollerton et al. 2007), can also tolerate HP depending on the phylogenetic relatedness of donor. Hence, as our experiments were designed to assess the role of phylogenetic distance of HP relative to recipient, they might have hidden the real contribution of pollination systems.

Additional traits potentially influencing recipient–donor interactions

Several floral traits besides phylogenetic distance are hypothesized to influence recipient–donor interactions affecting recipients' responses. In the case of stigmatic surface, large stigmas capture more HP in nature (Montgomery and Rathcke 2012) but since there is still enough space for CP adherence, HP post-pollination effects seem to be minimum. Hence, we might expect that small stigmas are more negatively impacted by HP than larger ones. Interestingly, by our experiments it is possible to see the opposite when HP distant was in the arena. Sisyrinchium wettsteinii is the species with smallest stigmatic area in our set of recipients and was the only one that had its post-pollination success improved by the presence of HP distant. On the other hand, HP close seems to always have a negative effect regardless of recipient stigmatic area.

Pollen features like its size and apertures are also potential influencers of plant responses to HP (Ashman and Arceo-Gómez 2013). Our experiments indirectly suggest that other pollen feature that might affect the interaction as well is its water content. This was noticed due to F. regia low probabilities of pollen germination in CP treatment, which could be due to the application of a non-intentional amount of non-viable pollen. Fuchsia pollen grains were described as partially hydrated (high water content; Franchi et al. 2002) which means that they are fast germinators and, hence, strong competitors (Nepi et al. 2001). However, this condition makes them highly vulnerable to water loss, decreasing their viability rapidly after removed from anthers (Franchi et al. 2002), which possibly occurred in this specific treatment. Even happening in the treatment on which donor and recipient are the same species, this can be extended to heterospecific interactions. That is, we can hypothesize that partially hydrated pollen may not impair recipient post-pollination success as much as dry pollen due to its faster viability loss.

The distance between recipients' stigmas and donors' anthers may also influence HP effect in recipient since it represents interactors' pollen flow. For instance, the different size of flowers of the two Fuchsia species may indicate that pollen flow between them is asymmetrical. Fuchsia campos-portoi, that has a smaller stigma height, has more chances of receiving HP from F. regia than the other way around. The former has some small chances of picking up B. gounelleana pollen from pollinator's body while for the latter this is very unlikely to occur. Therefore, the long styles of F. regia are more effective as an avoidance mechanism to secure that few HP close pollen will reach stigmas. In the set of interactions with S. wettsteinii as recipient, the stigma–anther distance between this species and pollen donors is negligible, regardless of phylogenetic distance. Such absence of a mechanical barrier is likely because generalist-pollinated flowers usually do not show a mechanical fit with pollen vectors resulting in a pollen placed diffusely in pollinator’s bodies (Minnaar et al. 2019). The lack of specificity in pollen deposition and picking up could denote that conspecific pollination assurance is more relevant than HP costs, or basically that HP is not costly and can be even advantageous to recipient plants with generalist-pollinated systems, as we demonstrated in this case.
Concluding remarks

Pollen performance was worst when HP close was applied to the stigmas. This reinforces that interactions via HP between congeners leads to more detrimental effects to recipients than when they get pollen from distantly related species (Arceo-Gómez and Ashman 2016). Nevertheless, our fine-scale study was able to demonstrate that plants can possibly also have positive reinforcements from receiving pollen of distantly related species. The evidence of tolerance and even benefits of HP in our experiments could be related to the community context that species are inserted. Tropical mountaintop communities, as the one here studied, are highly vulnerable to climatic variations, which makes the pollination environment very unpredictable (Freitas and Sazima 2006). For this plant community, it has been shown that species that flower nearby heterospecifics get more CP and grow more pollen tubes suggesting that the joint attraction of pollinators is advantageous under low pollinator availability circumstances (Bergamo et al. 2020a), but the role of HP receipt in such outcomes was not considered. Therefore, we hypothesize that the positive effect observed when S. wettsteinii received HP distantly related in our study could be a reflex of the pollinator scarcity context, especially because this interspecific facilitation mentioned before was a trend among the generalist-pollinated species of this community (Bergamo et al. 2020a, b). Our results together with what is known from the studied community give new insights on how plant–plant post-pollination interactions may influence community assembly for further studies to explore.

Data

All data and code are available at https://doi.org/10.6084/m9.figshare.c.4950777.v1

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Contributions by the Authors

N.S.S., M.W. and M.S. conceived and designed the study. N.S.S. and P.J.B. performed the experiments and collected the data. N.S.S. and T.-L.A. designed data analyses. N.S.S. wrote the first draft and all authors provided input to the final manuscript.

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Conflicts of Interest

None declared.

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