

VIEWPOINT

Pollen on stigmas as proxies of pollinator competition and facilitation: complexities, caveats and future directions

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• **Background** Pollen transfer via animals is necessary for reproduction by ~80 % of flowering plants, and most of these plants live in multispecies communities where they can share pollinators. While diffuse plant–pollinator interactions are increasingly recognized as the rule rather than the exception, their fitness consequences cannot be deduced from flower visitation alone, so other proxies, functionally closer to seed production and amenable for use in a broad variety of diverse communities, are necessary.

• **Scope** We conceptually summarize how the study of pollen on stigmas of spent flowers can reflect key drivers and functional aspects of the plant–pollinator interaction (e.g. competition, facilitation or commensalism). We critically evaluate how variable visitation rates and other factors (pollinator pool and floral avoidance) can give rise to different relationships between heterospecific pollen and (1) conspecific pollen on the stigma and (2) conspecific tubes/grain in the style, revealing the complexity of potential interpretations. We advise on best practices for using these proxies, noting the assumptions and caveats involved in their use, and explicate what additional data are required to verify interpretation of given patterns.

• **Conclusions** We conclude that characterizing pollen on stigmas of spent flowers provides an attainable indirect measure of pollination interactions, but given the complex processes of pollen transfer that generate patterns of conspecific–heterospecific pollen on stigmas these cannot alone determine whether competition or facilitation are the underlying drivers. Thus, functional tests are also needed to validate these hypotheses.

Key words: Conspecific pollen, heterospecific pollen, plant–plant interactions, plant–pollinator network, pollination, pollinator sharing, pollen tube, pollen transfer, stigmatic pollen load, visitation network.

INTRODUCTION

Approximately 80 % of flowering plant species depend on animal pollinators for sexual reproduction (Ollerton *et al.*, 2011). The sufficiency of pollination for a given species, however, can be strongly affected by the abundance and diversity of other plant species in the community (co-flowering effects; Pauw, 2013; Thomson *et al.*, 2019). Thus, understanding how pollinator-mediated interactions between plants affect plant reproduction has been an important pursuit for many decades (reviewed in Knight *et al.*, 2018). It is even more urgent now, given the accelerated rate of anthropogenic threats to this crucial ecosystem service (e.g. land use and climate change; IPBES, 2016).

Pollinator-mediated plant-plant interactions not only affect plant fitness and population growth (e.g. Ashman *et al.*, 2004) but can have implications for the evolution of floral traits (e.g. Caruso, 2000) and for the structure and assembly of plant communities (e.g. Sargent and Ackerly, 2008). Interactions between plant species for pollination services have been described as positive (facilitation), negative (competition) or neutral (commensalism) for one or both of the interacting plant species (reviewed in Braun and Lortie, 2019). While these interactions have been broadly demonstrated in pairs or in a few interacting plant species (e.g. Moeller, 2004; Ghazoul, 2006), the complexity of interactions represented in multispecies communities has necessitated a shift in emphasis away from focusing on pairwise interactions to the more diffuse ones found in most natural communities (e.g. Bascompte *et al.*, 2003).

One approach has been to characterize plant–pollinator networks based on visitation patterns, and these have revealed, among other things, extensive pollinator sharing among plants and high levels of plant and pollinator generalism (e.g. Bascompte *et al.*, 2003). However, visitation networks are limited in that they do not characterize the direction (positive or negative) of plant–plant interactions occurring, nor can they evaluate the full fitness consequences of these interactions (King *et al.*, 2013; Bennett *et al.*, 2018). Thus, attention has shifted away from solely observing pollinator visitation to recording

© The Author(s) 2020. Published by Oxford University Press on behalf of the Annals of Botany Company. All rights reserved. For permissions, please e-mail: journals.permissions@oup.com. community-level outcomes of the pollination process that take place between flower visitation and seed production (Fig. 1A). Specifically, researchers have begun to record the number and identity of conspecific pollen (CP) and heterospecific pollen (HP) grains transferred to stigmas for multiple species in a community (e.g. Fang and Huang, 2013; King et al., 2013; Emer et al., 2015; Tur et al., 2016; Johnson and Ashman, 2019). Patterns of pollen deposition on stigmas can be powerful in that they convey information on the strength of plant-plant interactions that result from pollinator sharing (e.g. Tur et al., 2016; Thomson et al., 2019), as well as serving as reliable proxies of the fitness consequences of these interactions (CP tube number and seed production; Fig. 1A, B). This is particularly true when the relationship between CP and HP grains and conspecific pollen tubes in the style is further evaluated as it affects the probability of ovule fertilization (Fig. 1B; Aizen and Harder, 2007; Alonso et al., 2012; Arceo-Gómez and Ashman 2014b; Tur et al., 2016). This shift in focus reflects the growing recognition that interacting communities of pollen grains on stigmas can affect post-pollination processes and overall plant fitness (Ashman and Arceo-Gómez, 2013).

Recently, post-pollination patterns of CP and HP receipt have been used as proxies of the strength and direction of pollinatormediated competitive and facilitative plant–plant interactions in an entire community (Fig. 1B; Tur *et al.*, 2016). This was motivated by the challenges presented by experimental studies of multispecies interactions (e.g. Ghazoul, 2006) and the complex processes influencing pollen transfer (e.g. pollinator diversity, efficiency, visitation rate, floral density and morphology) in multispecies communities (e.g. Minnaar *et al.*, 2018; Albor *et al.*, 2019; Thomson *et al.*, 2019). In particular, the patterns

of covariation between CP and HP on stigmas have been used as proxies for pollinator-mediated competition (negative relationship) and facilitation (positive relationship; Fig. 1B) among plant species at the pre-pollination stage (i.e. before pollen deposition on stigmas; Tur et al., 2016). Furthermore, the likelihood of conspecific pollen tube growth as a function of CP and HP receipt (Fig. 1B) has been used in a similar manner to describe plant-plant interactions at the post-pollination stage (i.e. after pollen deposition on stigmas; Alonso *et al.*, 2013; Tur et al., 2016). Evaluating the relationship between HP and CP tubes/grain is particularly important as it more directly informs the fitness effects of HP on the recipient species (reviewed in Morales and Traveset, 2008). This, in turn, better informs the potential for overall facilitative or competitive plant-plant interactions, as the final outcome will strongly depend on the extent of the cumulative effects of HP transfer (i.e. effects via floral visitation and HP interference on the stigma), which have been shown to be highly variable (Ashman and Arceo-Gómez, 2013). Overall, these studies have revealed that there is a wealth of information to be gained from analysis of stigmatic pollen loads collected from natural communities.

The difficulty of conducting manipulative studies to evaluate the strength and direction of community-level plant–plant interactions (e.g. Ghazoul, 2006; Thomson *et al.*, 2019) further suggests that patterns of pollen receipt gathered from stigmas will continue to be an important tool to evaluate ecological interactions among multiple plant species within contemporary communities and as references for understanding how these change over time (Johnson *et al.*, 2019). However, we argue that the lack of acknowledgement of the myriad factors (e.g. relative flower abundance, flower morphology, pollinator



FIG. 1. Schematic representation of the pre- and post-pollination processes in the pathway from pollen transport to seed production (A) and how these processes translate into patterns of conspecific pollen (CP) versus heterospecific pollen (HP) on the stigma or patterns of HP and CP/tubes per grain in the style (B). (A) Pre-pollination processes affect CP and HP received by flowers, whereas post-pollination interactions between CP and HP on the stigma affect the number of CP tubes or seeds produced. (B) Patterns between CP and HP grains received, and between CP tubes/grain and HP received per stigma, can reflect pollinator-mediated interactions and their fitness consequences, respectively. Such patterns have been used to infer competitive or facilitative plant–plant interactions. The focus of this paper is to clarify the diversity of drivers that can underlie these patterns (Fig. 2).



FIG. 2. Patterns of conspecific pollen (CP) versus heterospecific pollen (HP) receipt that can be observed on stigmas of spent flowers and potential drivers of variation in these patterns. Potential drivers include variation in pollinator visitation (A, B) or variation in the pollinator pool and floral morphology (C–E). (A, B) Patterns that result from variation in quantity of pollinator visits. In (A), when visitation rate for the focal species is higher (>) or lower (<) in the presence of coflowering species, the pattern can reflect facilitation (green) or competition (red), respectively. However, in (B) lower visitation rate is accompanied by a pollen-transport trade-off leading to a negative CP–HP relationship. (C–E) CP–HP relationships when variation in pollinator pool and floral avoidance mechanisms exist regardless of visitation differences (blue for emphasis of mechanism). (C) Uneven pollinator pools combined with or without floral avoidance can lead to non-linear positive CP–HP relationships. (D) Uneven pollinator pools or perfect floral avoidance of HP represent forms of plant–pollinator specialization and lead to the lack of a CP–HP relationship.

visitation rate and diversity) that influence these proxies limits the conclusions that can be made regarding the types of diffuse plant–plant interactions (e.g. competition and facilitation) occurring pre-pollination in natural communities. Thus, a clear framework to study and interpret patterns of covariation between HP receipt, CP receipt, and CP tube growth is urgently needed if we aim to continue using these as proxies of ecological interactions within communities.

Our aim with this viewpoint is to stimulate awareness by discussing caveats, limitations and assumptions of using patterns of pollen receipt and pollen tubes in styles to interpret pollinator-mediated facilitative and competitive processes. We do so by explicating a few examples of how multiple processes can produce similar CP-HP patterns and vice versa and how multiple CP-HP patterns can result from the same prepollination processes (e.g. competition). We hope to shed light on the complexity, but also provide some guidelines on how to better evaluate and interpret data on stigmatic pollen loads in a way that helps move this growing field forward. It is also worth acknowledging for clarity that here we exclusively focus on interpreting these patterns from a female fitness perspective because recent studies have only considered this aspect (e.g. Tur et al., 2016). However, male fitness consequences of interspecific pollen transfer (i.e. CP loss) are important contributors to total plant fitness (Muchhala and Thomson, 2012; Moreira-Hernandez and Muchhala, 2019) and these would need to be eventually integrated at the community level to gain a full understanding of plant-plant interactions from pollen loads on stigmas.

Here we focus on key drivers of the associations (1) between CP and HP receipt, and (2) between the proportion of CP tubes/ grain and HP receipt (Fig. 1B). To illustrate our arguments we

conceptually explore how flower visitation rate (as affected by flower density), floral morphology and features of local pollinator pools could combine to generate the varied linear and non-linear relationships observed in natural communities (e.g. Arceo-Gómez *et al.*, 2016; Tur *et al.*, 2016), although we recognize that other factors may also be important (Minnaar *et al.*, 2018). This heuristic approach reveals that a more careful examination of the variation in pollen loads on stigmas is required to distinguish between pollinator-mediated competition and facilitation. In addition, our analysis indicates that supplementary information will be required to tease apart dominant underlying drivers of variation in pollen loads on stigmas. Nevertheless, stigmatic proxies are an important component of a research programme to understand plant–plant–pollinator interactions.

PROCESSES THAT UNDERLIE CP-HP RELATIONSHIPS

We consider how pollinator visitation rate, pollinator pool diversity/efficiency and floral HP avoidance morphology can contribute to the variation in transfer of pollen to stigmas. We then evaluate the ways these could combine to create a variety of observed patterns among flowers in CP and HP received (Fig. 2). In doing so we illustrate the complexity in interpretation of these patterns and reveal the features that need to be confirmed independently to confidently ascribe underlying processes.

The distribution of shared pollinator visits among co-flowering plants depends on their density and their attractiveness to pollinators (Rathcke, 1983). If plants are equally attractive, pollinators are expected to be recruited only as a function of flower densities in a patch, such that pollinator visitation reflects species' proportional flower abundance (i.e. neutral interactions). However, when pollinators are a limited resource, differences in relative flower density can disproportionally increase visitation rate to species with high flower densities at the expense of species with lower densities (i.e. competition; Rathcke, 1983; Braun and Lortie, 2019), as pollinators seek to exploit the most abundant resource. On the other hand, rare species (those at low densities) could benefit from increased overall flower abundance if pollinators are most attracted to highly rewarding flower patches, such that pollinator visitation to rare species increases in the presence of other co-flowering species (i.e. facilitation; Ghazoul, 2006). These different types of plant-plant pre-pollination interactions are in turn expected to lead to decreased (e.g. competition) or increased (e.g. facilitation) patterns of CP receipt. However, relative flower abundances cannot only affect CP receipt, but also affect HP delivery (Thomson et al., 2019). For instance, rare species that are facilitated are expected not only to receive more CP but also to receive larger HP loads due to increased pollinator visits to heterospecific flowers, potentially creating a trade-off between conspecific pollen load size and purity (Thomson et al., 2019). Nonetheless, if we only consider relative flower abundance and its effects on visitation rate, then facilitative interactions may be identified by a linear increase in CP (increased pollinator visitation) with increasing HP receipt (increasing visits to heterospecific flowers), as shown by Tur et al. (2016). However, the sign of the relationship (competition versus facilitation) may depend on the accrual rate of HP versus CP (see below).

Pollen transfer, however, depends on more than just visitation rate (reviewed in Minnaar et al., 2018). In particular, pollen transfer can be significantly influenced by inequalities in the efficacies of pollinators visiting flowers and by plant manipulation of pollen acquisition and deposition via floral morphological adaptations (e.g. Herrera, 1987; Conner, 1995; Moreira-Hernandez and Muchhala, 2019). For instance, pollinator pool diversity can be even (all pollinator taxa are equivalently represented) or uneven (some taxa are frequent while others are rare) and when pollinators differ in their effectiveness the degree of evenness can affect pollen receipt (Arceo-Gómez et al., 2016; Koski et al., 2018). Pollinator efficacy, i.e. the capacity of pollinators to transfer pollen from stamens to the conspecific stigma (Freitas, 2013; King et al., 2013), may depend on their behaviour at a flower as well as their constancy during single bouts (Minnaar et al., 2018). On the other hand, floral morphology can mediate the placement of pollen on a pollinator's body (Herrera, 1987; Huang et al., 2015; Tong and Huang, 2016; Fantinato et al., 2018; Minnaar et al., 2018), and thus the likelihood of pollen deposition on a stigma. For instance, relative positioning of anthers and stigmas can increase CP transfer and reduce the receipt of HP while absence of such floral avoidance strategies is expected to lead to higher HP transfer (Armbruster et al., 2009; Ashman and Arceo-Gómez, 2013; Minnaar et al., 2018; Moreira-Hernández and Muchhala, 2019). Other plant traits, such as pollen morphology, may also affect pollen adherence to, or resistance to grooming from, pollinators (Konzmann et al., 2019) and can be viewed similarly to HP floral avoidance morphology in their influence on pollen transport (Minnaar et al., 2018). In the following paragraphs we discuss specifically how the pollinator pool and floral avoidance can modify the effect of visitation rate and in turn affect the CP-HP relationship and thus influence our

interpretation of facilitative versus competitive plant-plant interactions.

We predict that the above processes combine to create different accrual patterns of CP and HP on stigmas resulting in different relationships between CP and HP when flowers are sampled at the end of their lifetimes. So far, however, their influences on CP and HP receipt have only been considered independently from each other (Arceo-Gómez et al., 2016; Tur et al., 2016). As explained above, linear increasing relationships can arise entirely from quantitative differences in the relative availability of pollinators and flowers (Fig. 2A; for examples see Arceo-Gómez et al., 2016; Tur et al., 2016). However, this would only occur if both CP and HP are deposited with each pollinator visit - as pollinator visits increase, the stigma accumulates HP at the same rate as CP. This pattern would be expected when the assemblage of pollinators is even and of similar effectiveness (depositing similar amounts of CP and HP; Arceo-Gómez et al., 2016), but only when the recipient species' flowers do not display floral adaptations for avoidance of HP. Under this scenario, differences in the effect of co-flowering species presence on the visitation rate to a focal species, i.e. neutral (visits proportional to focal species abundance), facilitative (visits greater than proportional abundance) or competitive (visits less than proportional abundance), are expected to affect the steepness but not the sign of the CP-HP accrual relationship (Fig. 2A). For instance, if the effect of co-flowering species presence is to reduce visitation rate of the focal species, i.e. a competitive interaction, then the slope of the CP-HP relationship is expected to be low (Fig. 2A), as visitors deposit more HP than CP with every visit owing to spending more time on the competitors' flowers than those of the focal species. In this case, both CP and HP will still increase as more visits are received but HP will accumulate faster than CP (Fig. 2A). In an artificial array of two flower types for instance, Thomson et al. (2019) observed that unattractive flowers receive larger pollen loads but with much higher amounts of HP compared with CP when intermingled with a competitor compared with when they are not. This result indicates that HP can accumulate faster than CP in the presence of competition. Wide among-species variation in the value of CP-HP positive slopes has also been observed within a single community (Tur et al., 2016). Hence, we propose that not only the direction (Tur et al., 2016) but also the strength of the slope of the CP-HP relationship is an important indicator of the potential type of plant-plant interaction taking place (competitive versus facilitative). Furthermore, under extreme asymmetrical competition for pollinators, not only will stigmas of the competitive loser accrue more HP and less CP, but limited space on the pollinators' bodies may result in a trade-off in transport between CP and HP because higher HP loads will prevent transport of CP. This trade-off can lead to an overall negative relationship between CP and HP receipt (Fig. 2B; for an example see Tur et al., 2016). Thus, we propose that when there is no floral HP avoidance and the effectiveness of the pollinator pool is even, the type of pollinator-mediated plant-plant interactions (competitive versus facilitative) reflected by CP-HP relationships can be differentiated by both the steepness of the positive slope of the CP–HP linear relationship when positive (Fig. 2A) and the negative sign of the slope in the presence of pollen placement trade-offs (Fig. 2A versus 2B). Some observational evidence of

these CP-HP relationships exists (Arceo-Gómez et al., 2016; Tur *et al.*, 2016) but these predictions are vet to be confirmed. While we recognize the difficulty of testing these predictions using stigmas collected in the field alone, they could be corroborated with other field data, such as characterizing pollen on pollinator bodies to evaluate pollen transport trade-offs, or they could be tested in experimental settings where floral morphology (e.g. artificial flowers), pollinators and relative flower densities are manipulated (e.g. Thomson et al., 2019). Nonetheless, this set of outcomes illustrates a salient point that a single ecological process, in this case pollinator competition, can yield two different patterns of CP-HP covariation (positive and negative; Fig. 2A, B) and that the same CP–HP pattern (positive) can be reflective of facilitation (Fig. 2A; visit rate greater) or competition (Fig. 2A; visit rate lower) depending on the magnitude of the slope. Thus, interpretation of the functional drivers must be cautious and well informed with respect to floral features and other aspects of patterns of pollen deposition on stigmas.

Quantitative relationships between CP and HP delivery are modified not only by competition and facilitation for visits but also by differences in quality aspects of pollination (amount and purity of CP loads). Specifically, the relationship between CP and HP can be non-linear when the pollinator pool is uneven, i.e. composed of one or few frequent, high-quality pollinators that deliver mostly CP plus several less frequent, low-quality pollinators that deliver CP and HP (or vice versa). Such patterns have been observed in several communities (Arceo-Gómez et al., 2016). Thus, we propose that unequal distributions of CP relative to HP on stigmas will occur as some stigmas receive highly effective pollinators (depositing abundant CP and rare HP) whereas others receive visits by less effective pollinators (carrying small amounts of CP and varying amounts of HP). As most HP would be delivered along with some small amounts of CP during visits by these inefficient pollinators, this scenario leads to an exponentially decreasing relationship of CP with HP receipt ('uneven' in Fig. 2C; for an example see Arceo-Gómez et al., 2016). In contrast, if numerous high-quality pollinators deliver small but pure CP loads and infrequent low-quality pollinators deliver large mixed (CP and HP) loads, then CP will show a negative exponential increase with HP ('uneven' in Fig. 2D; for an example see Arceo-Gómez et al., 2016). Thus, the quality of pollinators, not just the quantity of visits, affects CP-HP relationships. And specifically, the type of non-linear relationship that occurs with unevenness of pollinators depends on the amount of CP delivered by the most effective pollinators (also see Arceo-Gómez et al., 2016).

Interestingly, when floral avoidance morphology is not perfect at reducing HP deposition by all pollinators in the pool, it differentially impacts their transfer of CP relative to HP, and thus can also lead to non-linear relationships. For instance, when floral avoidance is more effective at reducing HP carried by pollinators bringing small loads than those carrying large loads of pollen a negative exponential CP–HP relationship would result. Large amounts of HP would only be delivered along with large CP loads, even under conditions of pollinator pool evenness and quantitative facilitation ('present' in Fig. 2D). Likewise, floral HP avoidance could exacerbate the variance in effectiveness of pollinators in the pool contributing to the non-linearities between CP and HP represented in

Fig. 2C, D (e.g. 'uneven' and 'present' in Fig. 2C and 'even' and 'present' in Fig. 2D), among other forms. It is worth noting that the most extreme case of pollinator pool unevenness is equivalent to absence of pollinator sharing (i.e. a single specialist, highly effective pollinator) and for these plants we predict there would be a non-significant CP-HP relationship with low variance in HP among flowers (Fig. 2E; for an example see Arceo-Gómez et al., 2016). A similar pattern would be expected when floral avoidance is perfectly effective and excludes all HP transfer, or when flowers exhibit early deposition of self CP ('prior selfing'; Lloyd and Schoen, 1992), preventing HP deposition by late-arriving visitors (e.g. Randle et al., 2018). This discussion makes clear that CP-HP relationships could arise from forces unrelated to the degree of competition and facilitation for visits. The described cases are examples of how the CP-HP relationships are strictly influenced by the single or combined action of flower morphology, pollinator diversity and autonomous selfing mechanisms (also see Arceo-Gómez et al., 2016; Minnaar et al., 2018). Thus, future studies should test for the presence of non-linear (in addition to linear) CP-HP relationships to determine whether these factors are at play in pollinator-mediated plant-plant interactions.

In conclusion, while variation in pollinator visitation rates alone can lead to linear CP-HP relationships that could reflect competition or facilitation for pollinator visits (e.g. Tur et al., 2016), we argue against this simplistic perspective on two grounds. First, CP-HP relationships are influenced by numerous factors other than the nature of the pre-pollination interaction - of which we have focused on two (pollinator pool and floral morphology) - that can result in complex (and non-linear) relationships (Fig. 2C-E). Second, both positive and negative slopes could be outcomes of competitive interactions, and positive slopes could arise from neutral, facilitative or competitive interactions (Fig. 2A, B). Thus, additional experiments are needed to avoid misinterpretation of the role of pollinator-mediated facilitative and competitive interactions in nature. We argue that to tease apart these pollinator-mediated plant-plant interactions, one will require not only knowledge of CP-HP patterns but of aspects of the reproductive biology (e.g. floral HP avoidance strategies, potential for autogamous self-pollen deposition, level of pollinator specialization) of the species being studied as well as manipulations to verify the predicted interpretations. For instance, when a positive CP-HP relationship is observed (Fig. 2A), manipulations of the whole floral community and/or key interacting species (i.e. those identified as sharing pollinators from the identity of pollen of the stigma; see below) can be used to test whether the pre-pollination interactions are competitive or not. If removal of heterospecifics leads to a higher CP-HP slope by the focal species, then pre-pollination interactions were competitive. However, if removal leads to a lower slope of the CP-HP relationship for the focal species, then pre-pollination interactions were facilitative. These community-level experiments can be logistically daunting, but by identifying key facilitators/competitors', effects can be simultaneously evaluated on multiple focal species. Focused manipulative experiments will also avoid the time-consuming task of observing pollinator visits on a wide range of species sometimes leading to insufficient sampling. Likewise, manipulations of the pollinator pool evenness can be used to confirm its role in creating non-linear relationships (Fig. 2C, D). For instance, selective removals of pollinators (e.g. Brosi and Briggs, 2013) could be used to increase evenness in the pollinator pool, an effect predicted to reduce non-linearities under competitive pre-pollination interactions (Fig. 2C), potentially revealing a negative relationship resulting from CP-HP trade-offs (Fig. 2B). Finally, manipulation of focal species floral traits, for instance style length (e.g. Creswell, 2000), can be used to test whether floral avoidance contributes to non-linearities in CP-HP relationships (Fig. 2C, D). If changing the height of the stigma alters the CP–HP relationship, then manipulations of flower abundance (see above) can be used to determine the nature of pre-pollination interactions (i.e. via visitation). Pairing CP-HP patterns with a judicious combination of experimental approaches can provide more rigorous information about the nature of plant-plant interactions than either alone.

PROCESSES THAT UNDERLIE CP TUBES/GRAIN-HP RELATIONSHIPS

A second, often overlooked, relationship that can be observed from flowers at the end of their lifetime is that between CP tubes in the style and the load of CP and HP on the stigma (e.g. Briggs et al., 2016). The relationship between the number of CP tubes and the number of CP grains alone can provide information on the CP quality aspect of pollination (Kalla and Ashman, 2002; Parra-Tabla and Bullock, 2005; Alonso et al., 2012). The relationship between the proportion of CP grains that produce a tube (i.e. CP tubes/grain) and HP (Fig. 1B) is particularly relevant because it informs on the consequences of HP receipt for effective fertilization and is thus central for assessing the final outcome of pollinator-mediated competitive or facilitative interactions. Thus, consideration of both the quality of CP delivered to the stigma and the potential CP-HP interactions that occur after deposition is essential for understanding the contribution of post-pollination interactions to plant evolution and community structure (Morales and Traveset, 2008; Ashman and Arceo-Gómez, 2013; Briggs et al., 2016). Below we briefly describe the factors that can influence CP-HP interactions on the stigma and thus our interpretation of the HP-CP tubes/grain patterns observed.

The quality of CP that is delivered to a stigma is affected by genetic and environmental factors. Environmental conditions during pollen development and dehiscence, such as temperature, drought and UV irradiance, can reduce pollen grain viability and tube vigour (e.g. Kakani et al., 2002; Koski and Ashman, 2015). Pollen genotype can directly determine whether CP grows a tube on a given stigma, e.g. genetic self-incompatibility (Rea and Nasrallah, 2008, Vaughton et al., 2010), and can interact with the environment to determine CP potency (e.g. Kakani et al., 2002). The genetic relatedness (e.g. outcross versus self) of CP on a recipient stigma can be dictated by plant floral traits (herkogamy or dichogamy), population genetic structure/density, or the behaviour of pollinators within or among plants (e.g. Mazer et al., 2010). Even changes in pollinator behaviour in response to the presence of heterospecific plants can alter the genetic composition of CP delivered to stigmas (e.g. increased selfing in the presence of competitors; Flanagan et al., 2009). Because outcross pollen can be more

vigorous, and produces tubes at higher rates than self pollen, the genetic relatedness of CP alone can affect the relationship between the number of CP tubes and grains (e.g. Parra-Tabla and Bullock, 2005, Arceo-Gómez and Ashman, 2014*a*).

Interactions between HP and CP grains on the stigma, or tubes within the recipient style, can also influence the quality of pollination. Heterospecific pollen can reduce CP tubes as a result of allelochemical or physical interference (Morales and Traveset, 2008: Ashman and Arceo-Gómez, 2013: Wipf et al., 2016; Takemori et al., 2019). Heterospecific pollen can sometimes lead to increased seed production (Arceo-Gómez et al., 2019; Suárez-Mariño et al., 2019), possibly reflecting increased CP tubes/grain as a result of the 'herd' or 'mentor'-like effect (as in CP; Cruzan, 1990; Niesenbaum, 1999). Seed production may also increase via release of biochemical compounds that influence pollen tube growth (i.e. spermidine; Palmer-Young et al., 2019) when HP germinates or bursts on the stigma (e.g. Prieu et al., 2016; Wipf et al., 2016). In multispecies loads of HP, antagonistic effects between HP grains can lead to non-additive effects on CP performance (Arceo-Gómez and Ashman, 2011). And finally, because HP can interact more strongly with self CP than outcross CP (Arceo-Gómez and Ashman, 2014a), the quality of CP deposited with HP can affect the relationship between CP tubes/grain and HP.

Once again, pollen-pollen interactions on the stigma could combine to create different relationships between CP tubes/ grain and HP under natural pollination conditions, but their actions have often been evaluated independently from each other. For instance, a decline in CP tubes/grain with increasing HP could result from HP interfering with CP germination or tube growth ('negative' in Fig. 3A), whereas an increase could occur if HP has positive effects on CP germination or growth ('positive' in Fig. 3B; see Tur et al., 2016 for examples). However, these same types of relationship could also arise when there is no effect of HP directly on CP but rather the presence of heterospecifics in the community changes the quality of CP deposited by pollinators. For instance, when increased presence of heterospecific plants leads to increased selfing and thus lower-quality CP deposited with HP ('decrease' in Fig. 3A), or alternatively when pollinators fly longer distances between rare species and more outcrossed pollen is brought along with increased deposition of HP ('increase' in Fig. 3B). Opposing impacts of CP quality and CP-HP interactions (e.g. increase in CP quality along with an increase in HP interference) can create the lack of a significant relationship between CP tubes/grain and HP ('increase' and 'negative' in Fig. 3C; see Tur et al., 2016 for examples), so it is important not to disregard such relations entirely by assuming neutral effects ('no change' and 'none' in Fig. 3C) when a lack of significant slope is observed. Interestingly, when there are dose-dependent effects of HP on CP tube growth (e.g. allelopathic effects; Thomson et al., 1981; Arceo-Gómez et al., 2018b) or when herd effects only occur after surpassing a threshold pollen load size, non-linear responses to HP are possible (Fig. 3D-F, respectively). If HP only interferes with CP germination after a relatively large load size is reached, then non-linear declining threshold effects could result (Fig. 3D). But if HP interferes with CP intensely even in small amounts then pronounced declines in CP tubes/ grain with increasing HP could result (Fig. 3E; for an example see Thomson et al., 1981). If HP has positive effects, then the



FIG. 3. Patterns of conspecific (CP) tubes/grain versus heterospecific pollen (HP) received on stigmas of spent flowers and potential drivers of variation in these patterns. Potential drivers are CP quality and HP interference with CP, with positive interaction noted in green and negative ones in red for emphasis. In (A) increasing interference or decreasing quality of CP can lead to a negative linear CP tubes/grain–HP relationship, which can be interpreted as forms of competitive interaction. In (B) positive linear content on the store of the s

opposite trend would be expected (Fig. 3F). Explicating these possibilities makes clear that the CP tubes/grain versus HP relationship can be complex. Teasing apart the different factors that drive these relationships will require careful experimentation where the origin, identity and quality of pollen deposited on stigmas is manipulated (for examples see Arceo-Gómez and Ashman, 2011, 2014a). Nevertheless, data on the relationship between CP tubes/grain and HP provide crucial insight into the consequences of pollinator sharing that occur after pollen deposition (e.g. effects of CP quality and HP interference, and mentor effects). Thus, CP tubes/grain remains an important proxy that deserves more empirical attention and careful interpretation.

ASSUMPTIONS AND ADDITIONAL CONSIDERATIONS

The insights gained from pollen composition on stigmas depend on how closely the underlying assumptions are met. Below we consider how several assumptions may affect the dynamics and interpretation of pollination processes.

Adequacy of sampling effort

The first assumption is that large sample sizes afforded by use of stigmatic pollen loads are adequate to fully characterize the multipartite relationships of interest. Lack of power, however, could lead to incorrect interpretations of the dominant pre-pollination processes, such as concluding that a linear relationship exists (Fig. 2A) when it does not, or concluding that a relationship does not exist ('no change' and 'none' in Fig. 3C) when a non-linear one does (Fig. 3D–F). The large flower-toflower intraspecific variance in stigmatic pollen loads that is often observed (Alonso *et al.*, 2013; Arceo-Gómez *et al.*, 2016; Thomson *et al.*, 2019) suggests that appropriate sample sizes need to be seriously considered, although this has yet to become commonplace for stigmatic load data sets. However, two studies provide some insight into adequate sampling for these purposes. Alonso *et al.* (2012) recommends 150 stigmas per species to assess non-linearities in CP tubes/grain relationships, whereas Arceo-Gómez *et al.* (2018*a*) recommend sampling a minimum of 50 stigmas per species to capture HP diversity. These sample size targets can be used as starting points for future studies, although rarefaction approaches will also be highly effective for assessing the sampling adequacy of a given study (e.g. Arceo-Gómez *et al.*, 2018*a*). Evaluating power will be a key component of any study using these proxies.

Influences of temporal and spatial variation

Another implicit assumption in the analysis of stigmatic pollen loads from spent flowers is that CP and HP arrive somewhat simultaneously, or at least have the same possibility of arriving throughout flower life, as might be the case for pollinators with diffuse pollen placement (Minnaar et al., 2018). The arrival time of HP and CP cannot, however, be inferred from collections made at flower senescence, but this timing could affect pollen load composition as well as pollen tube growth. For instance, flowers with pronounced autonomous autogamy (prior or delayed self-pollination relative to outcross pollination; Dole, 1990; Lloyd and Schoen, 1992; Randle et al., 2018) could have stigma loads that are not purely pollinatormediated, and thus differ dramatically in the time of self or outcross pollen arrival even in the absence of pollinator visits. Self and outcross CP grains can differ in their tube growth rates (e.g. Sorin et al., 2016) so different schedules of their delivery will affect interpretations of quality aspects of pollination and CP tubes/grain-HP relationships. Timing of HP delivery can affect the impact of HP on CP tubes/grain: simultaneous HP deposition has a greater negative effect on CP than staggered deposition (Bruckman and Campbell, 2014). Finally, autogamy will reduce flower-to-flower variance in CP receipt, and thus could obscure patterns of variation. For example, prior selfing could

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lead to high CP with low HP, as in Fig. 2E. As a consequence, it will be important to take the mode of delivery of self CP into consideration when evaluating CP–HP covariance as well as for the interpretation of the CP tubes/grain–HP relationship.

Finally, it is also implicitly assumed that the multispecies communities under study lack pronounced spatial structuring of the interacting plants and/or that pollinators disregard such structure if it exists. The evidence for impact of these on pollen transfer, however, is mixed. Strong spatial structure in simple communities (e.g. illustrated with arrays of artificial flowers; Thomson *et al.*, 2019) can impact HP and CP receipt, but local communities (Table 3, 'conspecific density effect' in Arceo-Gómez *et al.*, 2016). The degree to which spatial structure affects the proxies described herein can be tested with observational data of local plant distributions (e.g. Thomson *et al.*, 2016).

Influences of pollen identity and diversity

The proxies as described thus far implicitly assume that HP behaves as a uniform population of interspecific pollen. The specific identities and phylogenetic relationships of the HP donors (to each other and to the recipient), however, can result in varied individual and non-additive combinatorial effects on recipient CP tube growth and seed production (Arceo-Gómez and Ashman, 2011; Arceo-Gómez et al., 2016, 2019). Moreover, if HP diversity increases with load size, as seen in Arceo-Gómez et al. (2016), and HP diversity affects CP tubes/ grain (Arceo-Gómez and Ashman, 2011), then non-linear relationships between them may be more common in nature than not (e.g. Figs 2D, E and 3D-F). Thus, a full understanding of pollination interactions from the community perspective will require characterization of the diversity and composition of HP in addition to abundance on the stigma, as well as testing whether the composition of the HP stigmatic community affects CP tubes/grain versus HP relationships (e.g. Arceo-Gómez and Ashman, 2011).

CONCLUSIONS

Even in the light of the complex relationships described here, stigmatic pollen loads can add important information to our understanding of pollination processes at the community scale and in the wild. They directly address post-pollination interactions that cannot be captured by visitation metrics alone. For instance, facilitation for pollinator visits that does not lead to higher pollen deposition represents only 'apparent' facilitation because it cannot lead to higher reproduction. Here CP and HP on stigmas would be more reliable than visits as an indicator of the net interaction. In addition, evaluating the CP tubes/grain and HP relationship provides relevant information on whether a positive relationship between HP and CP receipt ultimately leads to a negative fitness effect or not. Thus, stigmatic proxies are valuable for understanding pollinator-mediated plant-plant interactions in multispecies communities, but we urge researchers to be clear about what can and cannot be inferred from these data alone. In particular, we highlight that competition and

facilitation can lead to the same CP–HP outcomes. To this end, we suggest that validation of hypothesized mechanisms (with knowledge of the natural history and reproductive biology of species and via experimentation) are required for a true understanding of the ecological process underlying the patterns observed on stigmas. We also argue, however, that stigmatic pollen loads can provide insight into contemporary pollinator-mediated plant–plant interactions, as well as those that occurred in the past. For instance, Johnson *et al.* (2019) used pollen on stigmas sampled from herbarium sheets to infer historical pollination interactions. Thus, stigmatic pollen loads represent a crucial tool for pollination ecologists faced with describing communitywide pollination interactions that are rapidly being transformed globally by human activities (IPBES, 2016).

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LITERATURE CITED

- Aizen MA, Harder LD. 2007. Expanding the limits of the pollen-limitation concept: effects of pollen quantity and quality. *Ecology* 88: 271–281.
- Albor C, García-Franco J, Parra-Tabla V, Díaz-Castelazo C, Arceo-Gómez G. 2019. Taxonomic and functional diversity of the co-flowering community differentially affect *Cakile edentula* pollination at different spatial scales. *Journal of Ecology* 107: 2167–2181.
- Alonso C, Herrera CM, Ashman T-L. 2012. A piece of the puzzle: a method for comparing pollination quality and quantity across multiple species and reproductive events. *New Phytologist* 193: 532–542.
- Alonso C, Navarro-Fernandez CM, Arceo-Gómez G, Meindl GA, Parra-Tabla V, Ashman T-L. 2013. Among-species differences in pollen quality and quantity limitation: implications for endemics in biodiverse hotspots. *Annals of Botany* 112: 1461–1469.
- Arceo-Gómez G, Ashman T-L. 2011. Heterospecific pollen deposition: does diversity alter the consequences? New Phytologist 192: 738–746.
- Arceo-Gómez G, Ashman T-L. 2014a. Heterospecific pollen receipt affects self pollen more than outcross pollen: implications for mixed-mating plants. *Ecology* 95: 2946–2952.
- Arceo-Gómez G, Ashman T-L. 2014b. Coflowering community context influences female fitness and alters the adaptive value of flower longevity in *Mimulus guttatus. American Naturalist* 183: E50–E63.
- Arceo-Gómez G, Ashman T-L. 2016. Invasion status and phylogenetic relatedness predict cost of heterospecific pollen receipt: implications for native biodiversity decline. *Journal of Ecology* 104: 1003–1008.
- Arceo-Gómez G, Abdala-Roberts L, Jankowiak A, et al. 2016. Patterns of among- and within-species variation in heterospecific pollen receipt: the importance of ecological generalization. *American Journal of Botany* 103: 396–407.
- Arceo-Gómez G, Alonso C, Ashman T-L, Parra-Tabla V. 2018a. Variation in sampling effort affects the observed richness of plant-plant interactions via heterospecific pollen transfer: implications for interpretation of pollen transfer networks. *American Journal of Botany* 105: 1601–1608.
- Arceo-Gómez G, Jameel MI, Ashman T-L. 2018b. Effects of heterospecific pollen from a wind-pollinated and pesticide-treated plant on reproductive success of an insect-pollinated species. *American Journal of Botany* 105: 836–841.
- Arceo-Gómez G, Kaczorowski RL, Patel C, Ashman TL. 2019. Interactive effects between donor and recipient species mediate fitness costs of

heterospecific pollen receipt in a co-flowering community. *Oecologia* **189**: 1041–1047.

- Armbruster WS, Hansen TF, Pelabon C, Perez-Barrales R, Maad J. 2009. The adaptive accuracy of flowers: measurement and microevolutionary patterns. Annals of Botany 103: 1529–1545.
- Ashman TL, Arceo-Gómez G. 2013. Toward a predictive understanding of the fitness costs of heterospecific pollen receipt and its importance in co-flowering communities. *American Journal of Botany* 100: 1061–1070.
- Ashman T-L, Knight TMS, Amarasekare JAP, et al. 2004. Pollen limitation of plant reproduction: ecological and evolutionary causes and consequences. Ecology 85: 2408–2421.
- Bascompte J, Jordano P, Melián CJ, Olesen JM. 2003. The nested assembly of plant–animal mutualistic networks. *Proceedings of the National Academy of Sciences of the USA* 100: 9383–9387.
- Bennett JM, Thompson A, Goia I, et al. 2018. A review of European studies on pollination networks and pollen limitation, and a case study designed to fill in a gap. AoB Plants 10: ply068.
- Braun J, Lortie CJ. 2019. Finding the bees knees: a conceptual framework and systematic review of the mechanisms of pollinator-mediated facilitation. *Perspectives in Plant Ecology, Evolution and Systematics* 36: 33–40.
- Briggs HM, Anderson LM, Atalla LM, Delva AM, Dobbs EK, Brosi BJ. 2016. Heterospecific pollen deposition in *Delphinium barbeyi*: linking stigmatic pollen loads to reproductive output in the field. *Annals of Botany* 117: 341–347.
- **Brosi BJ, Briggs HM. 2013.** Single pollinator species losses reduce floral fidelity and plant reproductive function. *Proceedings of the National Academy of Sciences of the USA* **110**: 13044–13048.
- Bruckman D, Campbell DR. 2014. Floral neighborhood influences pollinator assemblages and effective pollination in a native plant. *Oecologia* 176: 465–476.
- Caruso CM. 2000. Competition for pollination influences selection on floral traits of *Ipomopsis aggregata*. *Evolution* **54**: 1546–1557.
- Conner J. 1995. The effect of wild radish floral morphology on pollination efficiency by four taxa of pollinators. *Oecologia* 104: 234–245.
- **Creswell JE. 2000.** Manipulation of female architecture in flowers reveals a narrow optimum for pollen deposition. *Ecology* **81**: 3244–3249.
- Cruzan MB. 1990. Pollen-pollen and pollen-style interactions during pollen tube growth in *Erythronium grandiflorum* (Liliaceae). *American Journal* of Botany 77: 116–122.
- Dole JA. 1990. Role of corolla abscission in delayed self-pollination of Mimulus guttatus (Scrophulariaceae). American Journal of Botany 77: 1505–1507.
- Emer C, Vaughan IP, Hiscock S, Memmott J. 2015. The impact of the invasive alien plant, *Impatiens glandulifera*, on pollen transfer networks. *PLoS ONE* 10: e0143532.
- Fang Q, Huang SQ. 2013. A directed network analysis of heterospecific pollen transfer in a biodiverse community. *Ecology* 94: 1176–1185.
- Fantinato E, Del Vecchio S, Giovanetti M, Acosta ATR, Buffa G, Collins B. 2018. New insights into plants co-existence in species-rich communities: the pollination interaction perspective. *Journal of Vegetation Science* 29: 6–14.
- Flanagan RJ, Mitchell RJ, Knutowski D, Karron JD. 2009. Interspecific pollinator movements reduce pollen deposition and seed production in *Mimulus ringens* (Phrymaceae). *American Journal of Botany* 96: 809–815.
- Freitas L. 2013. Concepts of pollinator performance: is a simple approach necessary to achieve a standardized terminology? *Brazilian Journal of Botany* 36: 3–8.
- Ghazoul J. 2006. Floral diversity and the facilitation of pollination. *Journal of Ecology* 94: 295–304.
- Herrera CM. 1987. Components of pollinator "quality": comparative analysis of a diverse insect assemblage. *Oikos* 50: 79–90.
- Huang ZH, Liu HL, Huang SQ. 2015. Interspecific pollen transfer between two coflowering species was minimized by bumblebee fidelity and differential pollen placement on the bumblebee body. *Journal of Plant Ecology* 8: 109–115.
- IPBES. 2016. Summary for policymakers of the assessment report of the Intergovernmental Science-Policy Platform on Biodiversity and Ecosystem Services (IPBES) on pollinators, pollination and food production. https:// ipbes.net/sites/default/files/spm_deliverable_3a_pollination_20170222.pdf.
- Johnson AL, Ashman T-L. 2019. Interspecific pollen transfer network structure shifts along spatial invasion gradients in a diverse island ecosystem. *New Phytologist* 221: 142–154.
- Johnson AL, Rebolleda-Gómez M, Ashman T-L. 2019. Pollen on stigmas of herbarium specimens: a window into the impacts of a century of

environmental disturbance on pollen transfer. American Naturalist 194: 405-413.

- Kakani VG, Prasad PVV, Ceaufurd PQ, Wheeler TR. 2002. Response of in vitro pollen germination and pollen tube growth of groundnut (*Arachis hypogaea* L.) genotypes to temperature. *Plant, Cell and Environment* 25: 1651–1661.
- Kalla SE, Ashman T-L. 2002. The effects of pollen competition on progeny vigor in *Fragaria virginiana* (Rosaceae) depend on progeny growth environment. *International Journal of Plant Sciences* 163: 335–340.
- King C, Ballantyne G, Willmer PG, Freckleton R. 2013. Why flower visitation is a poor proxy for pollination: measuring single-visit pollen deposition, with implications for pollination networks and conservation. *Methods in Ecology and Evolution* 4: 811–818.
- Knight TM, Ashman T-L, Bennett JM, Burns JH, Passonneau S, Steets JA. 2018. Reflections on, and visions for, the changing field of pollination ecology. *Ecology Letters* 21: 1282–1295.
- Konzmann S, Koethe S, Lunau K. 2019. Pollen grain morphology is not exclusively responsible for pollen collectability in bumble bees. *Scientific Reports* 9: 4705.
- Koski MH, Ashman T-L. 2015. Floral pigmentation patterns provide first example of Gloger's rule in plants. *Nature Plants* 1: 4013.
- Koski MH, Ison JL, Padilla A, Pham AQ, Galloway LF. 2018. Linking pollinator efficiency to patterns of pollen limitation: small bees exploit the plantpollinator mutualism. *Proceedings of the Royal Society B* 285: 20180635.
- Lloyd DG, Schoen DJ. 1992. Self- and cross-fertilization in plants. I. Functional dimensions. International Journal of Plant Sciences 153: 358–369.
- Mazer SJ, Hove AA, Miller BS, Barbet-Massin M. 2010. The joint evolution of mating system and pollen performance: predictions regarding male gametophytic evolution in selfers vs. outcrossers. *Perspectives in Plant Ecology, Evolution and Systematics* 12: 31–41.
- Minnaar C, Anderson B, de Jager ML, Karron JD. 2018. Plant-pollinator interactions along the pathway to paternity. *Annals of Botany* 123: 225–245.
- Moeller DA. 2004. Facilitative interactions among plants via shared pollinators. *Ecology* 85: 3289–3301.
- Morales CL, Traveset A. 2008. Interspecific pollen transfer: magnitude, prevalence and consequences for plant fitness. *Critical Reviews in Plant Sciences* 27: 221–238.
- Moreira-Hernández JI, Muchhala N. 2019. Importance of pollinatormediated interspecific pollen transfer for angiosperm evolution. *Annual Review of Ecology, Evolution, and Systematics* **50**: 8.1–8.27.
- Muchhala N, Thomson JD. 2012. Interspecific competition in pollination systems: costs to male fitness via pollen misplacement. *Functional Ecology* 26: 476–482.
- Niesenbaum RA. 1999. The effects of pollen load size and donor diversity on pollen performance, selective abortion and progeny vigor in *Mirabilis jalapa* (Nyctaginaceae). *American Journal of Botany* 86: 261–268.
- Ollerton J, Winfree R, Tarrant S. 2011. How many flowering plants are pollinated by animals? *Oikos* 120: 321–326.
- Palmer-Young EC, Farrell IW, Adler LS, et al. 2019. Chemistry of floral rewards: intra and interspecific variability of nectar and pollen secondary metabolites across taxa. *Ecological Monographs* 89: e01335.
- Parra-Tabla V, Bullock SH. 2005. Ecological and selective effects of stigmaanther separation in the self-incompatible tropical tree *Ipomoea wolcottiana* (Convolvulaceae). *Plant Systematics and Evolution* 252: 85–95.
- Pauw A. 2013. Can pollination niches facilitate plant coexistence? Trends in Ecology and Evolution 28: 30–37.
- Prieu C, Matamoro-Vidal A, Raquin C, et al. 2016. Aperture number influences pollen survival in Arabidopsis mutants. American Journal of Botany 103: 452–459.
- Randle AM, Spigler RB, Kalisz S. 2018. Shifts to earlier selfing in sympatry may reduce costs of pollinator sharing. *Evolution* 72: 1587–1599.
- Rathcke B. 1983. Competition and facilitation among plants for pollination. In: Real L, ed. *Pollination biology*. London: Academic Press, 305–329.
- Rea AC, Nasrallah JB. 2008. Self-incompatibility systems: barriers to self-fertilization in flowering plants. *International Journal of Developmental Biology* 52: 627–636.
- Sargent RD, Ackerly DD. 2008. Plant–pollinator interactions and the assembly of plant communities. *Trends in Ecology and Evolution* 23: 123–130.
- Suárez-Mariño A, Arceo-Gómez G, Sosenski P, Parra-Tabla V. 2019. Patterns and effects of heterospecific pollen transfer between an invasive and two native plant species: the importance of pollen arrival time to the stigma. *American Journal of Botany* 106: 1308–1315.

- Sorin YB, Mitchell RJ, Trapnell DW, Karron JD. 2016. Effects of pollination and postpollination processes on selfing rate in *Mimulus ringens*. *American Journal of Botany* 103: 1524–1528.
- Takemori A, Naiki A, Takakura KI, Kanaoka MM, Nishida S. 2019. Comparison of mechanisms of reproductive interference in *Taraxacum*. *Annals of Botany* **123**: 1017–1027.
- Thomson JD, Andrews BJ, Plowright RC. 1981. The effect of a foreign pollen on ovule development in *Diervilla lonicera* (Caprifoliaceae). *New Phytologist* 90: 777–783.
- Thomson JD, Fung HF, Ogilvie JE. 2019. Effects of spatial patterning of co-flowering plant species on pollination quantity and purity. *Annals of Botany* 123: 303–310.
- Tong ZY, Huang SQ. 2016. Pre- and post-pollination interaction between six co-flowering *Pedicularis* species via heterospecific pollen transfer. *New Phytologist* 211: 1452–1461.
- Tur C, Saez A, Traveset A, Aizen MA. 2016. Evaluating the effects of pollinator-mediated interactions using pollen transfer networks: evidence of widespread facilitation in south Andean plant communities. *Ecology Letters* 19: 576–586.
- Vaughton G, Ramsey M, Johnson SD. 2010. Pollination and late-acting self-incompatibility in *Cyrtanthus breviflorus* (Amaryllidaceae): implications for seed production. *Annals of Botany* 106: 547–555.
- Wipf HM, Meindl GA, Ashman TL. 2016. A first test of elemental allelopathy via heterospecific pollen receipt. American Journal of Botany 103: 514–521.