Diverse pollinator communities enhance plant reproductive success

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Understanding the functional consequences of biodiversity loss is a major goal of ecology. Animal-mediated pollination is an essential ecosystem function and service provided to mankind. However, little is known how pollinator diversity could affect pollination services. Using a substitutive design, we experimentally manipulated functional group (FG) and species richness of pollinator communities to investigate their consequences on the reproductive success of an obligate out-crossing model plant species, Raphanus sativus. Both fruit and seed set increased with pollinator FG richness. Furthermore, seed set increased with species richness in pollinator communities composed of a single FG. However, in multiple-FG communities, highest species richness resulted in slightly reduced pollination services compared with intermediate species richness. Our analysis indicates that the presence of social bees, which showed roughly four times higher visitation rates than solitary bees or hoverflies, was an important factor contributing to the positive pollinator diversity–pollination service relationship, in particular, for fruit set. Visitation rate at different daytimes, and less so among flower heights, varied among social bees, solitary bees and hoverflies, indicating a niche complementarity among these pollinator groups. Our study demonstrates enhanced pollination services of diverse pollinator communities at the plant population level and suggests that both the niche complementarity and the presence of specific taxa in a pollinator community drive this positive relationship.

Keywords: biodiversity–ecosystem function relationship; complementary resource use; ecosystem services; plant–pollinator interactions; positive selection effect; social bees

1. INTRODUCTION

Understanding the consequences of biodiversity loss on ecosystem functioning and services has developed into a central theme in ecology [1–3]. Animal pollination is a critical ecosystem service as most angiosperms are pollen-limited [4] and rely on animals for sexual reproduction [5]. A large proportion of the human diet depends directly or indirectly on animal pollination [6]. However, there is evidence that pollinators are declining in many parts of the world as a consequence of environmental degradation [7–9]. Recent research has linked reductions in the pollination of crops and wild plants in intensively managed agro-ecosystems to declines in density and diversity of pollinators [10–18]. However, in these observational studies, aggregate abundance and diversity of pollinators are correlated. Thus, an experimental approach controlling for the confounding effect of aggregate abundance is needed in order to gain a better and more mechanistic understanding of the role of pollinator diversity in the provisioning of pollination services ([19–21], see also [22]).

Mainly three mechanisms have been proposed to explain positive pollinator diversity–pollination service relationships [20]: first, such relationships may result from a positive sampling [23,24] or selection effect [25], by which diverse communities are more likely to include functionally highly effective species or species groups. Pollination effectiveness among pollinator groups can vary in the quality of the pollen transfer as a consequence of morphological and behavioural differences, but primarily owing to variation in flower visitation rates [26]. Secondly, under functional facilitation some community members enhance the functional performance of others [27]. For example, honeybees switched more often between plant individuals of sunflower hybrids in the presence of foraging wild bees, thereby facilitating cross-pollination [28]. Third, functional complementarity through niche partitioning [29] in the flower-visitation patterns of pollinators may lead to enhanced pollination services provided by diverse pollinator communities [16,19,21]. Such niche partitioning may occur at multiple temporal and spatial scales [21]. At a large scale, inter-annual [18] or regional complementarity may enhance pollination services and their stability. Furthermore, diversity effects in pollination may be owing to a diverse plant community being pollinated (corresponding to resource heterogeneity) [19] or to differences among pollinators visiting a single-plant species [21,30]. The latter can occur, for example, when different species in a pollinator community partition their foraging activities during different daytimes [31] or among flowers at different positions within plant individuals [16]. However, experimental knowledge of the functional consequences of pollinator niche partitioning in diverse pollinator communities for the pollination success of single-plant species is currently lacking [but see 22].

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Received 13 July 2012
Accepted 13 September 2012

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In the present study, we experimentally manipulated functional group (FG) richness (one versus three FGs) and species richness (one, three and nine species) of caged pollinator communities in a substitutive design and analysed patterns of pollinator visitation to the single plant *Raphanus sativus* L. as a model species to address the following questions: (i) How does the presence and richness of pollinator species and FGs affect fruit and seed set of a self-incompatible, insect-pollinated plant species? (ii) Do single FGs and communities of pollinators differ in their temporal (diurnal) and spatial (flower visitation height) niches? While functional facilitation was not a focus of this study, we had the following expectations regarding positive sampling and complementarity effects: under a positive sampling effect, we would expect increased visitation rates and plant reproductive success in the presence of a given pollinator species or FG. If complementarity contributes to a positive pollinator diversity–pollination service relationship, we would expect complementary flower visitation patterns of different pollinator groups during the day or between different flower heights, and a significant contribution of these differential visitation patterns among pollinator groups to increased visitation rates at the pollinator community level in diverse communities.

## 2. MATERIAL AND METHODS

### (a) Study plant species

*Radish, R. sativus* ssp. *oleiformes* L. (Brassicaceae), is a hermaphroditic, annual herb native to Europe. It produces up to several hundred flowers per plant and is visited by a wide variety of flower visitor, including solitary and social bees, hoverflies and butterflies [13,26,32]. *Raphanus sativus* has a sporophytic self-incompatibility system and relies on animal pollination for reproduction [33]. Thus, *R. sativus* is an ideal model species to study the effects of diverse pollinator communities on life-time plant reproductive success [13,26].

### (b) Experimental design

For the pollination treatments, 12 cubic cages (side length, 2 m, mesh width, 0.8 mm [19]) were set up in spring 2007 in the experimental garden of the University of Zurich (Switzerland). The cages prevented natural pollination of the experimental plant populations and a ground-covering plastic foil prevented natural plant establishment inside cages. During a pollination treatment a cage contained nine potted, abundantly flowering plants of *R. sativus* arranged in a grid with a width of 50 cm between plants. During spring 2007, *R. sativus* had been sown directly into these pots containing 5 l of standardized, nutrient-rich garden soil and grown in a pollinator-free glasshouse. To ensure that flowering plants of roughly the same age and size were available for the pollination treatments, they were sown at four different dates with a time-lag of several hours [37]. Most pollinator species were captured at first flowers of *R. sativus* at the age of 7 days when flowers are open and receptive for pollination [10]. They represent the main FGs pollinating radish [13,32,34], although butterflies can also be functionally important in some regions and ecosystems [26]. Each FG comprised three species, thus pollinator communities of up to nine species were used in the experiment: the social bee species *Bombus terrestris* L. (A1), *Bombus pascuorum* Scopoli (A2) and *Apis mellifera* L. (A3); the solitary bee species *Halictus rubicundus* Christ (B1), *Andrena flavipes* Panzer (B2), *Lasioglossum* sp. (B3); and the hoverfly species *Eristalis tenax* Lateville (C1), *Episyrphus balteatus* De Geer (C2) and *Sphaerophoria* sp. (C3) (figure 1). The *Lasioglossum* bees were most likely all *L. morio* Fabricius and the *Sphaerophoria* hoverflies most likely all *S. scripta* L., but we cannot totally rule out the possibility that also some individuals of morphologically very similar congenic species were collected as identification of these species can be difficult. All species are highly generalized, polylectic flower visitors [35,36].

### (c) Pollination rounds

The experiment was conducted during four days (four time blocks, hereafter ‘pollination rounds’) on 17 July, 25 July, 1 August and 9 August 2007 under sunny weather conditions. A 1-day period for each pollination round was used because flowers of *R. sativus* are open and receptive for roughly 1 day, with most ovules available for fertilization for a few hours [37]. Most pollinator species were captured at...
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different locations in the northeast of Switzerland. Individuals of the hoverfly species *E. balteatus* were purchased as pupae from Katz Biotech AG, Germany. Two large colonies of *B. terrestris* were purchased from Leu & Gygax AG, Switzerland. For each community comprising *B. terrestris* half of the individuals required for a certain pollinator treatment were taken from colony 1 and the other half from colony 2. Pollinators (both purchased and captured individuals) were kept in boxes (acryl-glass and fine-meshed fabric; 50 × 50 × 150 cm) in a climate chamber (20°C, 60% humidity) and were fed with sugar- and honey-water and finely ground pollen (Leu & Gygax AG) until the day before using them for a pollination round. For each pollination round, pollinators were introduced into cages at 8 h and removed at 19 h on the same day. Immediately after a pollination round, stalks of open, not wilted flowers of each *R. sativus* plant were marked with a permanent marker. The next day, potted plants were brought back to the pollinator-free glasshouse until fruit collection. Seed set and fruit set was determined as the number of seeds and number of fruits set, respectively, per marked flower.

(d) Pollinator observations

To investigate possible mechanisms of pollinator-mediated consequences on plant reproduction, the number of visits— and the identity of the pollinator species performing the visit—was recorded for a randomly selected focal plant for each pollinator community treatment. Observations were made during each of the four pollination rounds. No observations were made before 9 h to ensure that pollinators had enough time to calm down after introduction into cages. Bees needed approximately 30 min to calm down before starting to visit flowers, while hoverflies usually started to visit flowers immediately (M. Albrecht 2007, personal observation; in agreement with observation by Fontaine et al. [19]). For each visit to a randomly selected focal plant the height of the flower visited by a pollinator was estimated and assigned to one of three flower height classes (basal, less than 40 cm; medium, between 40 and 80 cm; apical, more than 80 cm). Observations of pollinator communities were done during 30-min observation periods during each of four different daytime periods: 9.00–11.30, 11.30–14.00, 14.00–16.30 and 16.30–19.00. Despite extensive observations (3660 min of total observation time) and several people observing simultaneously during pollination rounds, it was not possible to observe all replicates of the different pollinator communities, and the three-species community A2B2C2 was not observed.

(c) Data analysis

Linear mixed-effects models were fitted using the lme-function of the nlme package supplied in the R-system of statistical computing [38]. A model selection procedure based on Akaike’s information criterion (AIC) was used to select the most adequate model, using maximum likelihood for model comparisons and backward selection starting from the full model [39]. Most adequate models were fitted with restricted maximum likelihood and model fit was assessed by testing the residuals for normality and homoscedasticity and by plotting the residuals against the predicted values. Means ± 1 s.e. are reported. The relative importance of the predictor variables of the full model was calculated as the proportion of the total variance explained by each variable using increments of multiple $R^2$ squared (i.e. percentage typ 1 sum of squares) of the fixed model versions of the fitted linear mixed models [40]. The calculated percentages can be used as measures of effect sizes [40].

To test the effect of pollinator FG richness, species richness (and their interaction) and the presence of FGs on the response variables fruit and seed set, they were included as fixed effects in the full model. Species richness was log-transformed because this gave a better fit than linear species richness. However, we also calculated models in which species richness was fitted with a second degree polynomial (i.e. [linear species richness + [linear species richness]$^2$] to test for a hump-shaped relationship (results not shown). Cage (nested within pollination round) and community composition were included as random effects. Because this analysis indicated that species richness did not explain much variation when fitted after FG richness (see §3), we focused on FG identity and richness as explanatory variables in the subsequent analysis of niche complementarity. The full model, fitted to test whether spatio-temporal resource use (square-root transformed number of visited flowers) differed among pollinator FGs, contained the fixed factors pollinator FG identity, daytime (four daytime periods) and flower height (three height classes), and all their possible interactions, and pollinator species identity, cage and pollination round as random effects. To analyse visitation-rate patterns of whole pollinator communities, the same model but with the fixed effect FG richness instead of FG identity and the random effect pollinator community identity instead of pollinator species identity was fitted. Data have been deposited in the Dryad repository [41].

3. RESULTS

(a) What are the functional consequences of pollinator richness?

Functional group richness of pollinators increased plant seed set ($F_{1,27} = 9.60, p = 0.005$; 1 FG: 2.91 ± 0.17; 3 FGs: 3.48 ± 0.28; figure 2a), explaining 55 per cent of the total variation owing to the fixed effects of the full model. However, log(species richness) only explained an additional non-significant amount of 9 per cent (fitted after FG richness in the full model; $F_{1,27} = 1.45, p = 0.239$; figure 2a). The presence or absence of social bees explained an additional marginally significant amount of 19 per cent (fitted after FG richness and log(species richness); $F_{1,27} = 3.19, p = 0.085$). Furthermore, there was no significant interaction between FG richness and log(species richness) (fitted after FG richness, log(species richness) and presence of social bees; $F_{1,27} = 1.44, p = 0.241$).

Functional group richness of pollinators also increased plant fruit set ($F_{1,26} = 10.86, p = 0.003$; 1 FG: 0.61 ± 0.04; 3 FGs: 0.78 ± 0.04; figure 2b), explaining 37 per cent of the total variation owing to the fixed effects of the full model. Similar to seed set, the positive effect of log (species richness) on fruit set only explained an additional non-significant amount of 2 per cent (fitted after FG richness in the full model; $F_{1,26} = 0.94, p = 0.342$; figure 2b). Most of the total variation in fruit set owing to fixed effects, namely 60 per cent, was explained by the presence or absence of social bees (fitted after FG richness and log(species richness); $F_{1,26} = 19.66, p < 0.001$). The interaction between FG richness and log(species richness) was very small and not significant (fitted after FG richness, log(species richness) and
The presence of social bees; $F_{1,27} = 1.44$, $p = 0.241$. The most adequate model for fruit set contained social bee presence (present: $0.78 \pm 0.04$, absent: $0.52 \pm 0.04$) as the only fixed explanatory variable.

**Do pollinator functional groups differ in spatio-temporal flower visitation?**

Pollinator FGs tended to differ in the number of flowers visited ($F_{2,6} = 4.86$, $p = 0.055$), which was primarily a result of the roughly four times higher visitation rate of social bees compared with solitary bees or hoverflies (figure 3). The three FGs differed in their flower-visitation patterns during the day, irrespective of whether only communities consisting of a single FG were analysed ($F_{6,87} = 4.81$, $p < 0.001$) or also three-FG communities were included in the analysis ($F_{6,229} = 3.21$, $p = 0.005$; figure 3). However, when only the communities consisting of three FGs were analysed, the three FGs did not significantly differ in flower visitation among the four different daytime periods ($F_{6,108} = 0.92$, $p = 0.484$), but social bees still tended to differ from solitary bees and hoverflies in visitation rate early (morning and noon) compared with later during the day (afternoon and evening; $F_{1,115} = 3.82$, $p = 0.053$). Social bees visited most flowers between 14.00 and 16.30 h, in contrast to solitary bees, visiting most flowers in the morning between 9.00 and 11.30 h, while hover flies visited most flowers in the morning and noon, with similar numbers recorded between 9.00 and 11.30 h and between 11.30 and 14.00 h, respectively (figure 3). The three FGs did not differ in the relative number of visits to flowers at lower parts of plants compared with more upper parts when single and three FG communities were analysed separately, although there was a trend for social bees to visit a relatively higher number of basal flowers when all communities were analysed (interaction FG × flower height contrast (basal versus medium and apical): $F_{2,69} = 2.56$, $p = 0.085$). While including the FG × daytime interaction (indicating diurnal visitation differences among pollinator groups) substantially enhanced the model fit ($\Delta$AIC $7.18$; all communities included), including the FG × flower height interaction only slightly increased model performance ($\Delta$AIC $< 2$), confirming that diurnal differences in visitation rates among FGs played a more important role than spatial differences (visits to flowers at different heights).

**Do pollinator communities differ in spatio-temporal flower visitation?**

Analysing visitation rates of whole communities, the average number of visited flowers increased with FG richness from $14.4 \pm 3.4$ to $22.7 \pm 6.9$ visits per plant during 30 min, but this increase was statistically not significant ($F_{1,13} = 1.17$, $p = 0.299$). However, visitation rate was significantly increased in communities including the FG social bees (present: $25.4 \pm 3.1$, absent: $7.9 \pm 2.8$;
group assemblages, owing to morphological constraints in short-tongued hoverflies to pollinate plants with tubular flowers, while more efficiently pollinating open flowers. In this case, similar to experiments with plant or bacterial communities, the functional consequences of diversity were stronger in more heterogeneous resource environments [47,48]. In contrast, our study is among the first to experimentally demonstrate positive effects of pollinator richness in a more homogeneous pollinator resource environment, namely that of a single-plant species. From the plant’s point of view this indicates that even at the population level a diverse pollinator community may increase plant reproductive success.

Apart from the study of Fontaine et al. [19] and the present study, existing evidence for positive effects of pollinator species diversity on pollination services comes mainly from correlational studies of animal-pollinated crop [10,12,16] and wild-plant species [13,42]. Some of these correlations between pollinator diversity and pollination service have been attributed to temporal complementarity among years [10] or combined spatio-temporal complementarity [16]. However, collinearity among aggregate abundance and diversity of pollinators in these studies makes it difficult to assess the importance of different components of pollinator diversity and the mechanisms driving the observed patterns [20,21]. Indeed, a recent simulation study suggested that similar functional patterns may arise from the relationship between relative abundance and the effectiveness of the pollinator species present in diverse communities [30]. Our results demonstrate positive pollinator diversity effects on pollination services and plant reproductive success that are independent of aggregate pollinator abundance.

Despite the clear need for controlled experiments to address some of the important aspects of the functional consequences of pollinator diversity [19–21], they come at the cost of simplifying some of the real-world complexity. Cages represent an artificial environment to pollinators, hindering them in the performance of some types of natural behaviour, such as the provisioning of nests in the case of bees. However, in agreement with observations of Fontaine et al. [19], visitation rates and duration of flower visits of foraging pollinators were in the range of those observed under natural conditions for the model plant species (M. Albrecht 2004, unpublished data). Pollinator communities of a generalized plant species such as R. sativus are likely to be more species-rich, at least those of plant populations in relatively un-degraded habitats [13,42]. From our study using a relatively low maximum pollinator number, we cannot rule out the possibility that at considerably higher levels of pollinator richness the richness–pollination service relationship becomes humped-shaped [42] if negative selection effects owing to many highly inefficient pollinators or nectar-robbing flower visitors [49] play a more important role. Furthermore, it is conceivable that at high richness antagonistic pollinator interactions in the simple one-resource plant environment, e.g. through disturbance or even competitive exclusion of functionally superior pollination service providers by inferior ones from some pollination niches, could lead to negative complementarity effects. Such increases in antagonistic interactions in simple resource environments leading to negative biodiversity effects have recently been found in bacterial biodiversity–ecosystem functioning experiments.
More complex resource environments including larger temporal and spatial scales are expected to considerably broaden the scope for complementarity effects and associated increased ecosystem functioning [18, 51–53].

(b) What are the drivers of the positive pollinator richness effects?

Our results indicate that the presence of social bees in a community was an important factor explaining positive pollinator richness effects on seed set, and in particular fruit set, of *R. sativus*, suggesting that a positive selection effect [25] played an important role in the observed diversity effects. Indeed, social bees visited roughly four times more flowers than solitary bees or hoverflies, and the three social bee species showed the highest pollination service, measured as fruit and seed set, in the single-species treatments. The pollinator species providing the highest pollination service was the bumblebee *B. pascuorum*.

However, we found strong evidence that—in addition to the higher overall visitation rates of social bees—niche complementarity was a key mechanism driving the positive pollinator richness–pollination service relationship: the three FGs differed in their relative foraging activity at different times of the day—social bees showing particularly distinct diurnal visitation patterns compared with solitary bees and hoverflies—and they tended to differ in the relative number of flowers visited at different heights within plants.

By exploiting different spatio-temporal niches, pollinators can maximize their resource-use efficiency, while simultaneously increasing pollination efficiency at the community level [20, 21]. Diurnal foraging activity is determined by intrinsic factors, such as physiological attributes and environmental tolerances, and behavioural responses in relation to the daily course of extrinsic factors [54] that may result in pollinator species-specific or pollinator group-specific ‘daily activity windows’ [31, 51, 55, 56]. In our study, solitary bees foraged most in the morning, hoverflies in the morning and noon and social bees in the afternoon. Similarly, peak visitation rates of bumblebees and honeybees in the afternoon have been observed for other wild plant and crop species in temperate climates, which have mostly been attributed to positive temperature–foraging activity relationships [57]. Our findings are also in agreement with the scarce existing data on diurnal foraging activity of hoverflies, suggesting that pollen feeding of most species is highest in the morning hours [58]. Previous observations of foraging activity patterns of solitary bees indicate that they can be highly variable, with some solitary bee species visiting more flowers in the morning than in the afternoon [59], while others show bimodal patterns with peaks in the morning and afternoon [31]. In our study, social bumblebees and honeybees, in contrast to the other pollinator groups, also continued to forage after 18 h (M. Albrecht, 2007, personal observation). In many plant species, including *R. sativus*, stigma receptivity is rather short (usually a few hours) and can show some variation among plant individuals during the day [37], which could have contributed to the importance of diurnal complementarity in pollinator foraging activity.

Spatio-temporal niche partitioning in pollinator communities is likely to be greatest in heterogeneous or plant species-rich landscapes that offer a broad array of niches to be partitioned [20, 53]. This is in accordance with the general finding that biodiversity effects increase with biotope space [48]. Indeed, our results indicate that even within a simple environment with a single flowering plant as a resource, such spatio-temporal niche partitioning was effective, with the temporal component of complementarity in diurnal visitation times being clearly more important than the spatial component of complementarity in flower visitation heights.

5. Conclusions

Our study demonstrates enhanced pollination-mediated reproductive success in a single-plant species owing to higher pollinator functional-group richness—and if only one FG is present also at higher species richness—indeed independent of aggregate pollinator abundance. Such fine-scale functional effects of pollinator richness are likely to be important for the population dynamics of local populations of natural plant species, and have economic implications for the many animal-pollinated plant crops worldwide, which are typically grown as monocultures of single-plant species [6]. In agreement with predictions of biodiversity–ecosystem functioning relationships for simple resource environments, highest species richness resulted in slightly reduced pollination services compared with intermediate levels of species richness. Our results suggest that both complementarity effects, primarily resulting from different realized daytime niches among pollinator FGs, and the presence of particular taxa in a pollinator community, in our case social bees, contributed to the positive pollinator richness–pollination service relationship. These findings provide an important step towards a more mechanistic understanding of the effect of pollinator diversity on pollination services. Our results emphasize the importance of the conservation and restoration of diverse pollinator communities for the provisioning of pollination services to animal-pollinated plants.

We thank Jochen Fründ and Colin Fontaine for valuable discussions and advice, and two anonymous reviewers for excellent suggestions that considerably improved an earlier version of the manuscript. We further thank Dominique Bühler, Lauren Cottle, Anna Kopps, Angelica Lopez, Kevin Richards, Daniel Trujillo and Tobias Züst for assistance at various stages of the experiment and the analysis of fruit and seed set. Thanks for funding the project go to the Swiss National Science Foundation (grant to CBM; 631-065950).

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