Foraging efficiency and size matching in a plant–pollinator community: the importance of sugar content and tongue length

Saskia G.T. Klumpers,1,2,3* Martina Stang1,2 and Peter G.L. Klinkhamer1

1Plant Ecology and Phytochemistry, Institute of Biology Leiden, Leiden University, Sylviusweg 72, 2333 BE Leiden, The Netherlands
2Rocky Mountain Biological Laboratory, Crested Butte, CO 81224, USA
3School of Life Sciences, University of KwaZulu-Natal, Private Bag X01, Scottsville, Pietermaritzburg 3209, South Africa

*Correspondence: E-mail: s.g.t.klumpers@biology.leidenuniv.nl

INTRODUCTION

The description and understanding of the structure of species interaction networks is growing into a major research field in ecology. Especially mutualistic interactions between plants and pollinators provide an important model system in this field. Studies of these interaction networks have revealed general interaction patterns such as nestandedness, connectance and asymmetry in species dependencies (e.g. Bascompte et al. 2003; Vázquez & Aizen 2004; Bascompte & Jordano 2007). These patterns can be explained by a number of non-mutually exclusive factors, including species abundance, phylogenetic relationships, overlap in spatio-temporal distributions and matching of phenotypic traits of interacting species (e.g. Vázquez et al. 2009a and references therein; Olesen et al. 2011). In spite of a relatively high ability to predict these aggregate network characteristics, only a few studies have been able to predict pairwise interactions (Stang et al. 2009; Olesen et al. 2011; Verdu & Valiente-Banuet 2011; Kaiser-Bunbury et al. 2014; Maglianesi et al. 2014; but see Vázquez et al. 2009b). Nevertheless, insight into the mechanisms that determine pairwise species interactions is important as it can help to understand and better predict commonly observed structural features of ecological communities.

Size matching between floral nectar tube depth and pollinator proboscis length has been shown to be an important determinant of plant–pollinator interactions. In general, flowers with deep nectar tubes restrict access of nectar to visitors with long mouthparts (Haber & Frankie 1989; Corbet 2000; Stang et al. 2007). For specialised and co-dependent species pairs, evolutionary theory predicts that (1) there should be strong natural selection for pollinator proboscis to exceed floral nectar tubes in order to access all the nectar within the flowers; and (2) that selection on flowers should favour nectar tubes that are longer than their pollinator’s proboscis because this ensures contact with the reproductive parts of the flowers and thus maximises pollen transfer (e.g. Alexandersson & Johnson 2002; Anderson & Johnson 2008; Muchhala & Thomson 2009). This co-evolutionary arms race should lead to ever-increasing nectar tube depths and proboscis lengths and a close matching between these traits of strongly interacting species (Darwin 1862; Nilsson 1988; Anderson et al. 2010). However, such an arms race cannot explain size matching in multispecies assemblages with a variety of nectar tube depths and proboscis lengths and asymmetrically specialised species pairs. In these systems, flowers with a long nectar tube can only be visited for nectar by pollinators with a long proboscis, but these pollinators can potentially visit plants species with both long- and short-tubed flowers (Harder 1985; Haber & Frankie 1989). Although positive correlations between nectar tube depth of flowers and the average proboscis length of their pollinators have been found (e.g. Brian 1957; Gilbert 1981; Harder 1985; Shmida & Dukas 1990; Plowright & Plowright 1997; Borrell 2005; Stang et al. 2009), such a correlation does not necessarily imply that there is close size matching. For close size matching to occur, pollinators with a long proboscis need to forage preferentially from flowers with a long nectar tube. Only a few community studies have shown...
close size matching or that size matching can affect interaction strength or interaction probability of pairwise interactions (Stang et al. 2009; Maglianesi et al. 2014 for hummingbirds).

Although insects with a long proboscis can potentially visit a wide variety of flowers, they often seem to avoid relative shallow flowers (Stang et al. 2009) and preferentially forage on plant species with flowers of which the nectar tube matches their proboscis length (Haverkamp et al. 2016). One would expect such a preference if pollinators forage more efficiently on matching flowers. Flower foraging efficiency depends on both handling time and nectar reward. Concerning handling time, it has been shown that long-tongued bumblebees are generally faster foragers than short-tongued bumblebees when foraging on a single plant species (Harder 1983), irrespective of whether floral nectar tubes are longer or shorter than their proboscis. In contrast, several studies suggest that it takes pollinators with a long proboscis longer to handle short-tubed flowers than pollinators with a short proboscis; in other words, these studies suggest that tongues can be too long (e.g. Inouye 1980; Plowright & Plowright 1997; Kunte 2007; Karolyi et al. 2013; Bauder et al. 2015). For hummingbirds, it was found that the handling time of nectar resources was related to the degree of matching between bill length and nectar tube depth (Maglianesi et al. 2014). A morphological mismatch (longer bill than nectar tube) led in this study system to longer handling times when accounting for differences in nectar content across plant species. However, other than Maglianesi et al. (2014), we know of no studies that have revealed that handling time and the degree of trait matching of interacting species are related. What is more, the question whether foraging efficiency (defined as sugar extraction rate) of insect pollinators is higher on plant species with matching flowers remains largely unanswered. Insect pollinators seem to prefer more rewarding plants of the same species (Anderson et al. 2016). However, the specific roles of nectar reward and handling time for determining foraging efficiency and its effect on size matching in multispecies assemblages in which both a negative mismatch (proboscis is shorter than nectar tube depth) and a positive mismatch (proboscis is longer than nectar tube depth) occur has not been examined.

In this study, we investigated whether the degree of size matching between floral nectar tube depth and pollinator proboscis length can predict the interaction probability among sixteen co-occurring subalpine Asteraceae species and their pollinators, and whether foraging efficiency is an important driver of this size matching. As an indicator for foraging efficiency, we used sugar extraction rate. We included flower head display size and flower head density as covariates in this analysis as interaction probability is expected to be related to these floral characteristics (Hegland & Totland 2005; Verdu & Valiente-Banuet 2011; Kaiser-Bunbury et al. 2014). In addition, we determined the relationship between degree of size matching and both handling time and sugar extraction rate and examined how sugar extraction rate, handling time and nectar reward are related. Harder (1983) has tested the effect of proboscis length and nectar tube depth on probing time of bumblebees foraging on artificial flowers. The result of this study show that probing time increased gradually with increasing depth for flowers shallower than bees’ glossa. However, beyond that depth, probing time increased much more rapidly. Based on these findings, we hypothesize that the highest interaction probability and sugar extraction rate occur when proboscis length and nectar tube depth match. We expect that interaction probability and sugar extraction rate decrease rapidly when the nectar tube depth exceeds the proboscis length. This is because pollinators cannot reach the nectar in these flowers easily and therefore it will take pollinators longer to handle them. When pollinator proboscis exceeds the nectar tube depth, we expect interaction probability and sugar extraction rate to decrease more gradually, either because pollinators may need longer time to handle these flowers since a longer proboscis may be of hindrance, or because relative shallow flowers produce less nectar than deeper, closely matching, flowers. We choose species of the Asteraceae as a study system, because they are functionally generalised due to their relatively shallow tubular flowers. They have been shown to be visited by a high diversity of insect species, mostly belonging to the insect orders of Hymenoptera, Lepidoptera and Diptera (Lane 1996). This high level of functional generalisation makes these plants an ideal system to analyse the degree of size matching and foraging efficiency and its effect on flower choice in multispecies assemblages.

MATERIAL AND METHODS

Study system

Data were collected in the Colorado Rocky Mountains, USA, in the vicinity of the Rocky Mountain Biological Laboratory (RMBL). We included sixteen co-flowering Asteraceae species which were flowering most abundantly (see Table S1 in Supporting Information). Plant–pollinator interactions were observed from mid-June to mid-August 2011 in five different plots of approximately 50 by 50 m. These plots were 100–200 m apart. Pollinators could move freely between these plots. The elevation of the plots was between 2900 and 3200 m above sea level, exposed mainly to the south.

Interaction probability

Each plant species was observed for 4 hours in total by randomly walking through the plots where the focal plant species was flowering abundantly. Total sampling time was divided into eight sampling intervals of approximately half an hour. These sampling intervals were spread over several days, and during different times of day (morning and afternoon). All observations took place between 09:00 and 16:00 h on days without strong wind and rain.

Insect species were not caught to avoid changing their densities, distributions and behaviour. Instead, all insects encountered were filmed with a digital video camera for later identification. Individuals were identified up to the species- or morpho-species level based on video material and a reference collection. In total we observed 610 individuals belonging to 64 (morpho-)species, mostly Hymenoptera (28 species accounting for 74% of all the individuals), Lepidoptera (18 species, accounting for 16% of all individuals) and Diptera.
(12 species, accounting for 7% of all individuals). Videos were used to determine whether an insect visited a flower for nectar and/or pollen and whether it touched anthers or stigmas. Only those interactions where insects were foraging for nectar were included in the analyses. All insects touched anthers or stigmas, thus potentially acting as pollinators.

Handling time
Handling time was determined using the video recordings. Video recordings were analysed frame-by-frame (50 frames/s) with the software Adobe Premiere Pro CC 2014. We measured handling time as the time that an insect has its proboscis in the flower. Thus, our measurement of handling time includes the time an insect needs to enter a flower with its proboscis and extract the nectar. Not all individuals were filmed long enough to be able to analyse their handling time. In total, we recorded the handling time of 117 unique pairs of interacting plant-pollinator species. Handling time per insect species–plant species combination are averages of several individuals with an average 31 probes per species pair (range between 3 and 260 probes).

Trait matching: degree of (relative) mismatch
To calculate the degree of mismatch, we measured pollinator proboscis length and floral nectar tube depth. Proboscis length measurements of butterflies, flies, solitary bees and wasps were based on specimens from a reference collection. The specimens in this collection were collected in 2008 and 2010, at the same study site as where our study was conducted. The proboscis lengths of these specimens were measured according to the methods described by Stang et al. (2006). Data on the average proboscis lengths for bumblebees were obtained from Macior (1974) (for queens and workers) and from Inouye (1976) (for males), who also collected their specimens in the vicinity of our study sites.

For all plant species, we measured the corolla tube of disk flowers. This corolla tube consists of two parts: a (very) narrow lower part and an upper wider bell-shaped part. The wider part roughly begins where the stamens insert and ends at the beginning of the corolla lobes, at the base of the deepest cleft in the corolla. Nectar is produced at the base of the narrow part of the corolla tube and migrates up into the wider upper part of this tube. The lower narrow part of the corolla tube is completely filled by the style (see Table S1). Consequently, it can only contain traces of nectar and it is unlikely that insects can access this part of the corolla tube. Therefore, for each plant species, we only measured the depth of the upper wider part of the corolla tube, hereafter referred to as nectar tube (see Fig. S1). We measured the nectar tube depth for at least twenty freshly picked flowers (of at least ten different individuals), to the nearest 0.01 mm, under a dissecting microscope.

The degree of mismatch between pollinator proboscis length and floral nectar tube depth was calculated by subtracting the length of the nectar tube from the length of the proboscis. Consequently, positive values imply that pollinator proboscis is longer than the nectar tube of the flowers they visit. In turn, negative values imply that pollinator proboscis is shorter than the nectar tube of the flowers they visit. In addition, we then calculated the degree of relative mismatch by dividing the degree of mismatch by pollinator proboscis length. The degree of relative mismatch rather than the degree of mismatch might determine whether a long proboscis is of hindrance as e.g. a 0.5 mm longer proboscis would be of more hindrance for an insect with a proboscis of 1 mm than for an insect with a proboscis of 10 mm.

Sugar content and sugar extraction rate
To estimate nectar reward, we measured the sugar content of flowers that were bagged for 24 h. We used nectar production rate (NPR) as an estimate for nectar standing crop (NSC) because we were unable to measure the NSC of eight of the sixteen species. NPR is an appropriate estimate of NSC because these measurements were linearly correlated (Klummers, unpublished data). NPR and NSC measurements were performed on individuals flowering at the same location as where plant–pollinator interactions were observed. Flowers were bagged between 8 and 9 AM and collected 24 h later. We used bags made of fine gauze that do not change the temperature within the bags and thus do not influence the nectar solute concentration (Corbet 2003). As ray flowers of our species are sterile and do not produce nectar (Mani & Saravanan 1999), we only measured the nectar production of disk flowers. Our video footage also showed that only disk flowers were probed by insects. Nectar volume was measured in the laboratory under a dissecting microscope, within approximately an hour after collection. Nectar was extracted with micropipettes, ranging in size between 0.2, 0.25, 0.5 and 1.0 µL. Capillary size depended on the quantity of the nectar produced by the flowers and the nectar tube width. With the micropipette tubes we probed the base of the nectar tube until no more nectar could be removed and then used digital callipers to measure the distance the nectar had migrated up the microcapillary tubes. Nectar volume was determined by converting the distance measured to volumes (µL). Nectar sugar concentration (expressed in % w/w in a solution) was measured with a hand-held refractometer (Bellingham and Stanley, Tunbridge Wells, UK). Sugar content (mg) was quantified as the product of the nectar sugar concentration, which was first converted from wt/wt to wt/vol, x nectar volume (µL), as suggested by Bolten et al. (1979). When nectar volumes of single flowers were too small, nectar samples of flowers from the same flower head were pooled to obtain an estimate of the average sugar concentration of these flowers. Consequently, sugar content for individual flowers was estimated based on the nectar volume of a single flower and the average sugar concentration of the flowers in a particular flower head.

After the visit of a pollinator, there was no nectar left in flowers that could be extracted with micropipette tubes. Therefore, we obtained an approximation of sugar extraction rate by dividing the mean sugar content per flower by the handling time for each plant-pollinator species pair. Thus, sugar extraction rate is the amount of sugar a pollinator ingests per time unit, while handling a single flower.
Flower head display size

For each plant species, we measured the diameter of the flower head (including the rays for the species that have ray flowers) for at least twenty flower heads and calculated the surface area, which we refer to as flower head display size.

Flower head density

For each plant species, we counted the number of open flower heads per square meter. For most species, this number was calculated based on ten quadrats of one square meter which were randomly placed within the plots at places where at least one individual of a species was flowering. For Cirsium sp. 2 and Arnica purpurea, we measured flower head density based on one plot of ten square meters. This is because these species were flowering at only one location in our plots.

Statistical analyses

Statistical analyses were performed using R version 3.4.1 (R Development Core Team 2014). To test whether the probability that an interaction between a plant and pollinator species occurred was related to the degree of mismatch, we performed a generalised additive mixed model (GAMM), with presence/absence as response variable and degree of mismatch as explanatory variable, assuming a Binomial error structure. To account for the interspecific differences in flower head display size and flower head density and differences between insect orders, we included these variables as additional explanatory variables. Furthermore, to account for random variation among species, we included insect species as a random effect. As Hymenoptera, Lepidoptera and Diptera made up most of the interactions (97%), we only included these insect orders in the analyses.

To test whether both handling time and sugar extraction rate were related to the degree of mismatch and the degree of relative mismatch, we also used GAMMs, with either handling time or sugar extraction rate as response variable and either degree of mismatch or degree of relative mismatch as explanatory variable, assuming a Gaussian error structure. We calculated the mean handling time and mean sugar extraction rate per plant–pollinator species pair. To account for the differences in handling time among insect orders (Herrera 1989), we included insect order as an additional explanatory variable. For the analyses of handling time, we also included the mean sugar content per flower as an additional explanatory variable. This is to account for the influence of nectar production per flower on handling time. We did not include sugar content in the analyses of sugar extraction rate because the latter was calculated as sugar content divided by handling time and thus would lead to spurious correlations. Finally, to account for the random variation among species, we included insect species as a random factor in these models. We performed GAMMs because this allows for fitting nonlinear and asymmetric relationships between explanatory and response variables. All GAMMs were performed, using the gamm function within the package “mgcv” (Wood 2018).

To analyse whether handling time was affected by sugar content, we used linear regression models with handling time as response variable and sugar content and insect order as explanatory variables. We analysed the effect of sugar content on handling time of negative and positive degree of mismatch separately.

Finally, we analysed the data of those insect species which visited more than three different plant species separately to illustrate the species–specific relationships between handling time or sugar extraction rate and degree of mismatch or relative mismatch. We could only include positive mismatches because of the lack of sufficient data for species with a negative mismatch. We performed linear regression models with handling time or sugar extraction rate as the response variable and either degree of mismatch or degree of relative mismatch and insect species as explanatory variables. In addition, to account for the effect of nectar reward on handling time, we fitted multivariate linear regression models with handling time as response variable and either degree of mismatch or degree of relative mismatch and insect species as explanatory variables. For all analyses, we log10-transformed handling time and sugar extraction rate to normalise residuals.

RESULTS

Interaction probability

The traits of the plant and insect species that were observed are summarised in Tables S1 and S2 respectively. Plants were on average visited by eleven different insect species (range 2–31 species), while insects visited on average three different plant species (range 1–11 species). Thus, plants were more generalised than insects.

Among Hymenoptera and Diptera, the highest interaction probability occurred when their proboscis was slightly longer than the nectar tube (Fig. 1a; Table 1). Among Hymenoptera, we found a rapid decrease in interaction probability when proboscides were increasingly shorter than nectar tubes. When proboscides were increasingly longer than nectar tubes, interaction probability decreased more gradually (Fig. 1a). Diptera experienced the highest interaction probability when their proboscides were approximately four mm longer than nectar tubes. When proboscides were shorter, interaction probability decreased. A mismatch greater than four mm did not occur among Diptera. Lepidoptera did not show a significant relationship between size matching and interaction probability. They did not visit flowers that were much longer than their proboscis (Fig. 1a; Table 1). Interaction probability also increased with flower head display size and flower head density (Figs 1b and 1c; Table 1). Together, the degree of mismatch, flower head display size and flower head density explained 15% of the variance in interaction probability.

Handling time

Handling time when proboscis was shorter than nectar tube

Handling time data are summarised in Table S3. Handling time was exponentially related to the degree of (relative)
mismatch as indicated by the linear relationship between degree of (relative) mismatch and $\log_{10}$ handling time (Figs 2a and 2b). Handling time strongly increased when proboscides were increasingly shorter than the nectar tube (negative degree of mismatch) (Figs 2a and 2b; Table 2). This was not because deeper flowers contained more sugar, since handling time was not related to sugar content when the proboscis was shorter than the nectar tube (Fig. 3a). This indicates that it takes insects much longer to extract nectar when they visit flowers of which the nectar tube is longer than their proboscis, not because these flowers contain more nectar but because their proboscis is too short to reach and extract the nectar efficiently.

Handling time when proboscis was longer than nectar tube
In contrast to the results when the degree of (relative) mismatch was negative, among Hymenoptera and Lepidoptera, handling time decreased when proboscides were increasingly longer than the nectar tube (Fig. 2a and b; Table 2). This indicates that when a proboscis is longer than the nectar tube, it is not of hindrance to the pollinator. When insects foraged on flowers that were shorter than their proboscis (positive mismatch), handling time increased with increasing sugar content (Fig. 3b). Analysing the data of only those insect species that visited at least three different plant species revealed that for positive mismatches, the degree of (relative) mismatch did not affect handling time when accounted for sugar content (Table S4). When accounting for the degree of relative mismatch, Lepidoptera and Diptera had significantly longer handling times than Hymenoptera, as indicated by a greater intercept in the GAMMs (Table 2).

Sugar extraction rate
Insects experienced the highest sugar extraction rate when their proboscis was slightly longer than the nectar tube, although this relationship was only significant among Hymenoptera (Fig. 2c and d). These were also the flowers that Hymenoptera and Diptera seemed to prefer, as their interaction probability was the highest on these flowers (Fig. 1a). Among Hymenoptera sugar extraction rate rapidly decreased when their proboscides were increasingly shorter than the nectar tubes. Sugar extraction rate decreased more gradually when proboscides were increasing longer than nectar tubes (Fig. 2c and d).

Figure 4 shows that most insect species experienced the highest handling time and sugar extraction rate when their proboscis length matched the nectar tube depth. Mainly the intercept in the regression models of the different insect species, and thus their average handling time and sugar extraction rate, rather than the slopes, differed.

DISCUSSION
Our results show that the probability that plant- and pollinator species interact can be predicted by the degree of size matching between floral nectar tube depth and pollinator proboscis length. Among Hymenoptera and Diptera, the highest interaction probability occurred when their proboscis was slightly longer than the nectar tube. On these flowers, insects experiencing the highest sugar extraction rate, and thus foraged most efficiently. This suggests that foraging efficiency is an important driver of pairwise plant–pollinator interactions. For Lepidoptera we did not find such a
relationship. This finding supports the idea that, in contrast to Hymenoptera, Lepidoptera are mainly opportunistic nectar feeders, using nectar resources as they become available (Shreeve 1992). Other floral traits, such as display or blossom size might also be more important for butterfly flower choice (Corbet 2000).
The role of handling time and nectar reward as determinants of pollinator foraging efficiency depended on the type of mismatch (positive or negative) between floral nectar tube depth and pollinator proboscis length. When proboscides were shorter than the nectar tube (negative mismatch), sugar content did not determine sugar extraction rate while handling time did. For these interactions, a decrease in sugar extraction rate was caused by the fact that it becomes more difficult to handle the flowers efficiently and not because these flowers contain less nectar. The reverse is true when proboscides were longer than the nectar tube (positive mismatch). When pollinator proboscis was longer than the nectar tube, a decrease in sugar extraction rate was not caused by increasing handling times resulting from a morphological mismatch, but from a smaller nectar content in flowers with shorter nectar tubes. Thus, pollinators experienced lower foraging efficiency when foraging on relatively shallow flowers, not because long tongues are of hindrance, as often proposed (Inouye 1980; Plowright & Plowright 1997; Stout 2000; Peat et al. 2005) and shown by Maglianesi et al. (2014) for hummingbirds. It is because those shallow flowers contain less sugar. Furthermore, Hymenoptera visited on average flowers that were more rewarding and handled these flowers faster than Lepidoptera.

Instead of exact matching, interaction probability was the highest in plant species of which flowers were slightly shorter than pollinator proboscides, an average mismatch which was also shown to occur in a Spanish pollination web (Stang et al. 2009). As our results indicate, pollinators may preferentially visit these plants because they forage most efficiently on their flowers. However, as there is intraspecific variability in nectar tube depth, as well as in pollinator proboscis length, by choosing to forage from flowers which are on average slightly shorter than their proboscis, there is a much lower probability that pollinators encounter flowers with nectar tubes that are

| Parametric coefficients | Smooth terms |
|-------------------------|--------------|
| Estimate | Std.error | T-value | P-value | Edf | F-value | P-value | R²_adj |
| (a) Handling time – degree of mismatch | | | | | | | |
| Intercept (Hym) | −0.26 | 0.03 | −8.77 | <0.001 | Degree of mismatch Hym | 2.35 | 17.45 | <0.001 | 0.48 |
| Dip | 0.13 | 0.13 | 1.00 | 0.319 | Degree of mismatch Dip | 1.00 | 0.28 | 0.598 |
| Lep | 0.62 | 0.08 | 7.65 | <0.001 | Degree of mismatch Lep | 1.00 | 5.34 | 0.023 |
| Sugar content | | | | | | | |
| (b) Handling time – degree of relative mismatch | | | | | | | |
| Intercept (Hym) | −0.27 | 0.03 | −0.01 | <0.001 | Degree of relative mismatch Hym | 1.70 | 35.18 | <0.001 | 0.54 |
| Dip | 0.17 | 0.08 | 2.13 | 0.036 | Degree of relative mismatch Dip | 1.00 | 1.64 | 0.203 |
| Lep | 0.42 | 0.20 | 2.04 | 0.044 | Degree of relative mismatch Lep | 2.27 | 5.07 | 0.019 |
| Sugar content | | | | | | | |
| (c) Sugar extraction rate – degree of mismatch | | | | | | | |
| Intercept (Hym) | −0.92 | 0.06 | −14.54 | <0.001 | Degree of mismatch Hym | 2.14 | 3.14 | 0.059 | 0.28 |
| Dip | −0.24 | 0.26 | −0.94 | 0.350 | Degree of mismatch Dip | 1.00 | 0.55 | 0.461 |
| Lep | −0.50 | 0.16 | −3.08 | 0.003 | Degree of mismatch Lep | 1.00 | 0.56 | 0.455 |
| (d) Sugar extraction rate – degree of relative mismatch | | | | | | | |
| Intercept (Hym) | −0.93 | 0.06 | −15.25 | <0.001 | Degree of relative mismatch Hym | 2.67 | 9.04 | <0.001 | 0.37 |
| Dip | −0.36 | 0.16 | −2.22 | 0.028 | Degree of relative mismatch Dip | 1.00 | 0.41 | 0.521 |
| Lep | −0.31 | 0.20 | −1.57 | 0.118 | Degree of relative mismatch Lep | 1.00 | 2.76 | 0.099 |

**Table 2** Results of the generalised additive mixed models (GAMMs) predicting the effect of degree of mismatch or degree of relative mismatch on handling time or sugar extraction rate, fitting a Gaussian distribution. Handling time and sugar extraction rate were log10 transformed. Sugar content was included as an additional explanatory variable in de models predicting handling time. In all models, insect species was included as a random factor.
Figure 4 Handling time as a function of (a) degree of mismatch (\(R^2_{\text{adj}} = 0.02, P = 0.109, N = 67\)), (b) species-specific degree of mismatch (\(R^2_{\text{adj}} = 0.19, P = 0.030, N = 67\)), (c) degree of relative mismatch (\(R^2_{\text{adj}} = 0.06, P = 0.028, N = 67\)), (d) species-specific degree of relative mismatch (\(R^2_{\text{adj}} = 0.18, P = 0.037, N = 67\)), and sugar extraction rate as a function of (e) degree of mismatch (\(R^2_{\text{adj}} = 0.11, P = 0.004, N = 67\)), (f) species-specific degree of mismatch (\(R^2_{\text{adj}} = 0.44, P < 0.001, N = 67\)), (g) degree of relative mismatch (\(R^2_{\text{adj}} = 0.25, P < 0.001, N = 67\)), and (h) species-specific degree of relative mismatch (\(R^2_{\text{adj}} = 0.47, P < 0.001, N = 67\)). Figures only show insect species that visit at least three plant species and only interactions of which the nectar tube depth < pollinator proboscis length (positive mismatch). In Figs a, c, e and g, colours represent different insect orders (blue: Lepidoptera, green: Hymenoptera, red: Diptera). In Figs b, d, f and h, colours represent different insect species and shapes represent different insect orders (open triangles: Lepidoptera, closed circles: Hymenoptera, and open squares: Diptera).
longer than their proboscis and consequently are not able to feed on the nectar.

The type of mismatch that we found appears to be in contrast to the general trend in co-evolutionary studies for pollinator tongues to be shorter than the nectar tube of the flowers they pollinate (Anderson et al. 2010; Paudel et al. 2016). This trend is generally explained by asymmetries in specialisation and thus dependence (Anderson et al. 2010). Selection may act more strongly on the interacting partner that is more specialised, as it has more at stake in the interaction (Dawkins & Krebs 1979; Brodie 1999). In coevolved systems, this is usually the plant as they have extremely long nectar tubes. In contrast, in our study system, pollinators were on average more specialised than plants. Thus our results still support the idea that asymmetries in specialisation might drive the degree of mismatch between interacting partners. Moreover, in more generalised pollination systems, of which the flowers are relatively shallow (such as the Asteraceae), flowers produce minute amounts of nectar that can only be found at the bottom of the nectar tube (Pleasants 1981). Therefore, the actual nectar tube depth equals the functional depth and consequently nectar is only accessible to visitors with a proboscis length equal to, or longer than, the nectar tube depth. This is in contrast to extremely long-tubed flowers in which the nectar wells up in the flowers and becomes accessible to pollinators with shorter mouthparts (Johnson et al. 2017). Deviation from 1:1 phenotypic matching does not necessarily imply a functional mismatch. For example, Toju & Sota (2006) showed experimentally that a functionally perfect phenotypical match (i.e. fitness of the two interacting species is balanced) occurs when weevil rostrums are 1.74 times longer than fruit pericarps. For the Asteraceae, anthers and styles stick out of the flowers. Therefore, functionally perfect phenotypical matching, i.e. optimal pollination and pollen transfer, could actually occur when pollinators’ proboscis is slightly longer than the nectar tube so that the head touches the anthers.

Overall, this study demonstrates that size matching between pollinator proboscis length and floral nectar tube depth can predict plant–pollinator pairwise interactions, and that foraging efficiency is an important driver for such size matching in plant–pollinator communities. As deeper tubed flowers generally produce more nectar (e.g. Galetto & Bernardello 2004; Petanidou 2007; Johnson et al. 2017), it is likely that this is a universal mechanism shaping plant–pollinator interactions. Being able to predict plant–pollinator interactions and understand the underlying mechanisms is of great importance for understanding (the structure of) interaction networks. Generally, short-tongued insects are not able to visit deep-tubed flowers (Haber & Frankie 1989; Corbet 2000; Stang et al. 2007) and the absence of these ‘forbidden’ interactions are often assumed to results in a nested network structure (Stang et al. 2006; Olesen et al. 2011; Ibanez 2012). However, our results indicate that interactions between long-tongued pollinators and short-tubed flowers do not occur often. Consequently, in quantitative networks, species morphology may lead to a compartmentalised, modular structure, rather than nestedness. Furthermore, most long-tongued and generalised pollinators are expected to be relatively tolerant (less at risk) to plant species loss because they are not dependent on a single species (Memmott et al. 2004). However, our results indicate that the loss of their ‘matching’ interaction partners may not only increase interspecific resource competition but also decrease their foraging efficiency, which may have negative fitness consequences (Hainsworth et al. 1991). Finally, foraging on the deepest possible flowers (given the length of the proboscis) leads to fitness benefits because these flowers produce more nectar. Therefore, higher pairwise specialisation and a co-evolutionary arms race between plants and pollinators leading to deeper flowers and longer proboscides is only likely to occur if this is accompanied by larger nectar rewards and consequently increased foraging efficiencies. We are only starting to unravel the role of nectar reward, handling time and foraging efficiency as determinants of pollinator flower choice and thus the structure of plant–pollinator interaction networks. We hope that this paper will stimulate more research on the role of nectar production and handling time as important determinants for pairwise interactions and community wide interaction patterns, especially in more generalised pollination systems.

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AUTHORSHIP

SGTK and MS designed the project and collected data, SGTK conducted statistical analyses and wrote the manuscript, all authors provided feedback on analyses, interpretation of results, and contributed substantially to revisions of the manuscript.

DATA ACCESSIBILITY STATEMENT

Data available from the Dryad Digital Repository: https://doi.org/10.5061/dryad.r2s74q4.

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