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Effect of expanded variation in anther position on pollinator visitation to wild radish, *Raphanus raphanistrum*

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● **Background and Aims** Plant–pollinator interactions shape the evolution of flowers. Floral attraction and reward traits have often been shown to affect pollinator behaviour, but the possible effect of efficiency traits on visitation behaviour has rarely been addressed. Anther position, usually considered a trait that influences efficiency of pollen deposition on pollinators, was tested here for its effect on pollinator visitation rates and visit duration in flowers of wild radish, *Raphanus raphanistrum*.

● **Methods** Artificial selection lines from two experiments that expanded the naturally occurring phenotypic variation in anther position were used. In one experiment, plant lines were selected either to increase or to decrease anther exertion. The other experiment decreased anther dimorphism, which resulted in increased short stamen exertion. The hypothesis was that increased exertion would increase visitation of pollen foragers due to increased visual attraction. Another hypothesis was that exertion of anthers above the corolla would interfere with nectar foragers and increase the duration of visit per flower.

● **Key Results** In the exertion selection experiment, increased exertion of both short and long stamens resulted in an increased number of fly visits per plant, and in the dimorphism experiment bee visits increased with increased short stamen exertion. The duration of visits of nectar feeders declined significantly with increasing long stamen exertion, which was opposite to the hypothesis.

● **Conclusions** Until now, anther position was considered to be an efficiency trait to enhance pollen uptake and deposition. Anther position in wild radish is shown here also to have an ecological significance in attracting pollen foragers. This study suggests an additional adaptive role for anther position beyond efficiency, and highlights the multiple ecological functions of floral traits in plant–pollinator interactions.

Key words: Anther position, artificial selection, Brassicaceae, pollen-foraging insects, pollinator-mediated selection, *Raphanus raphanistrum*, visitation rate.

INTRODUCTION

Plant–pollinator interactions play a major role in plant speciation (Grant, 1949; van der Pijl, 1961; Levin and Kerster, 1967; Grant, 1994) and are a classic example of evolution mediated through biotic interactions (Grant, 1949; van der Pijl, 1961; Fægri and van der Pijl, 1979; Proctor *et al.*, 1996). A useful way to understand the role of floral traits in pollination is to group traits by their specific functions. First, advertisement traits attract the attention of animal pollinators and provide cues for associative learning (e.g. Laverty, 1994; Fenster *et al.*, 2006; Higginson *et al.*, 2006; Ushimaru *et al.*, 2007). Secondly, reward traits are the ultimate reason for pollinator visits (e.g. Fægri and van der Pijl, 1979; Burd, 1995; Fenster *et al.*, 2006). Last are floral traits that influence pollination efficiency via the fit of flower and pollinator, and include flower and flower-part size, shape and position (e.g. Mitchell and Shaw, 1993;

Campbell *et al.*, 1994; Hansen *et al.*, 2003; Gómez *et al.*, 2009). For example, the position of the anthers relative to the corolla tube opening (anther exertion) affects pollen removal in wild radish (Conner *et al.*, 2003).

Floral attraction and reward traits have been shown to affect visitation behaviour of pollinators in numerous studies (e.g. Conner and Rush, 1996; Melendez-Ackerman *et al.*, 1997; Jones and Reithel, 2001; Kunze and Gumbert, 2001; Ne'eman and Kevan, 2001; Biernaskie and Cartar, 2004; Harder *et al.*, 2004; Armbruster *et al.*, 2005; Huber *et al.*, 2005; Irwin and Strauss, 2005; Wolfe *et al.*, 2005; Buide, 2006; Fenster *et al.*, 2006; Hoballah *et al.*, 2007). Less obvious and less well studied are the effects of efficiency traits on visitation behaviour. We propose two possible mechanisms for how anther position might affect pollinator visitation. First, prominent anthers can be part of the visual display of the flower (Nakanishi, 1982; Langanger *et al.*, 2000; Lunau, 2000; Andersson and Jorgensen,

2005), providing a direct signal of the reward for pollen foragers. On the other hand, prominent or exerted anthers may interfere with visitors foraging for nectar and may result in longer duration visits, which in turn can increase pollen removal or deposition (Young and Stanton, 1990; Conner *et al.*, 1995; Kudo, 2003). These hypotheses are not mutually exclusive, but both predict that anther position in the flower has a role beyond efficiency of pollen deposition.

Dimorphic or polymorphic positions of anthers, where anthers are at different heights or in whorls, is common in some angiosperm groups, such as in *Narcissus*, *Linum* and most Polemoniaceae and Brassicaceae species. Flowers of most of the approx. 4000 species in the mustard family (Brassicaceae) exhibit tetradynamy, in which there are four long and two short stamens in each flower. This trait is used as a diagnostic trait for the family (Zomlefer, 1994), but the reasons for the evolutionary stasis of this trait are unclear. There is significant genetic variation for the difference in anther heights in *Brassica rapa* and *Raphanus raphanistrum* (Karoly and Conner, 2000), and in the latter species tetradynamy is under stabilizing selection and affects pollinator efficiency (Conner *et al.*, 2003).

Experimental studies of anther position effects on pollinator behaviour are usually categorical, i.e. they use experimental manipulation of the numbers of anthers at different heights to assess the effect of anther position on the duration of visits and pollen removal per visit (e.g. Golding *et al.*, 1999; Kudo, 2003; Ornelas *et al.*, 2004; Syafaruddin *et al.*, 2006). While these studies provide insight to the function of anthers in whorls, they do not address the effects of continuous natural variation in anther positions. Using continuous variation in anther position is a complementary approach that more closely mimics natural variation for most traits (Stanton *et al.*, 1986; Campbell *et al.*, 1991; Conner and Rush, 1996; Conner *et al.*, 2003). The weakness of this approach is that traits that are well adapted will show reduced variation, as the unfit variants have been removed by selection in the past. Therefore, it is difficult to determine the effect of variation in these traits on fitness. Artificial selection is a solution to this problem of a lack of variation in continuous traits (Conner, 2003, 2006; Fuller *et al.*, 2005; Lehtila and Brann, 2007). Artificial selection is the source of most domesticated species (Meyer *et al.*, 2012), and can be used in quantitative genetic studies for evolutionary inferences (Callahan, 2005; Fuller *et al.*, 2005). However, it has been rarely, if ever, applied to pollination studies.

Here we used artificially selected lines of wild radish (*Raphanus raphanistrum*) with expanded variation in anther position to test the hypothesis that anther position, usually considered an efficiency trait, can affect pollinator visitation behaviour. Specifically, we tested the effect of long and short stamen exertion (relative to the corolla opening) on number of visits to the plant, number of flowers probed in each such visit and the duration of these visits. We tested two predictions. (1) More exerted anthers may act as a visual attractant and increase the accessibility of reward for pollen foragers. Therefore, we predict that increased short and long stamen exertion will increase the number of visits by pollen foragers and the time spent foraging per flower. (2) More exerted anthers may interfere with

visitors foraging for nectar and may thus cause increased time spent per flower.

MATERIALS AND METHODS

Study organism

Wild radish (*Raphanus raphanistrum*) is an annual or short-lived perennial native to the Mediterranean region, growing mainly in disturbed areas, and was introduced to North America in the middle of the 19th century (based on herbarium specimens; Panetsos and Baker, 1967; J. Conner, unpubl. data). Flowers of wild radish exhibit tetradynamy, in which there are four long and two short stamens in each flower. Wild radish is self-incompatible and thus its reproduction is entirely dependent on insect pollination; therefore, floral adaptations for successful pollination are crucial for fitness. The most common pollinators of wild radish in Europe and North America are honey-bees, small bees, bumble-bees, syrphid flies and butterflies, particularly *Pieris rapae* (Conner *et al.*, 2009); 14 of the 15 genera of these taxa tested were effective pollinators of wild radish (Sahli and Conner, 2007).

Artificial selection

Artificial selection was carried out in separate experiments for stamen dimorphism and for anther exertion of the long stamen, with two replicates of all selected and control lines. Anther exertion is defined as filament length minus the length of the corolla tube (Conner, 1997). Before selection, mean exertion of the long stamen was near zero, meaning that the entire anther is exerted, whereas mean short stamen exertion is about -2 mm (Conner and Via, 1993), which means the top of the anther is at the corolla tube opening (mean short anther length is about 2 mm; J. Conner, unpubl. res.). The initial base population for the exertion lines was derived from a single natural population (Conner and Via, 1993) that had been maintained in large random-mating greenhouse populations ($N_e \approx 600$) for seven generations (Conner, 2002); the dimorphism lines were established directly from the seeds collected from the same natural population. Filament and corolla tube lengths were measured on one early flower on each plant, usually the third (the first two often develop abnormally; J. Conner, pers. obs.). For both traits, within-family selection was used, in which the most extreme individual within each full-sib family was chosen as a parent for the next generation; completely randomized crosses were performed, with each selected plant serving as a male and a female, with no reciprocal crosses. This technique maximizes N_e and thus minimizes genetic drift (Falconer, 1989). One member of each family was randomly chosen for crosses in the control lines.

The anther exertion experiment involved 12 families in each of six lines: two selected for increased exertion, two for decreased exertion and two randomly selected controls. After three generations of selection performed at Kellogg Biological Station (MI, USA; hereafter KBS), replicates of each line were split between KBS and Reed College (OR, USA; hereafter RC) for two (RC) or three (KBS) more generations of selection. For more details of this artificial selection experiment, see Conner *et al.* (2011).

Stamen dimorphism has a lower heritability than anther exertion in *R. raphanistrum* (0.25 vs. 0.47, respectively; Karoly and Conner, 2000); thus, 20 families were used in each of four lines for the stamen dimorphism experiment. Two lines were selected for decreased dimorphism, and the other two were randomly mated controls. Each pair of lines was initiated from a base population consisting of one plant from each of 200 field-collected sibships grown in either the KBS or RC greenhouse, and selection continued for five generations, one pair of selection and control lines at KBS and the other at RC. On average, 7.4 offspring were grown and measured from each of the 20 full-sib families in each generation. We selected for an increased ratio of short to long stamen lengths, ensuring that selection is not biased toward plants with smaller flowers.

There was no evidence for significant genetic correlations between short and long stamen exertion and reward traits: pollen production, nectar volume and nectar concentration did not differ between the selection and control lines. This is not surprising, because since exertion is the difference between two linear dimensions, it is not correlated with overall flower size. Phenotypic correlations between short and long stamen exertion and the attraction traits of floral display (number of flowers open when the plant was placed in the field) and petal area were all <0.3 and with one exception not significant (Supplementary Data Table S1). Still, these two advertisement traits were included in the visitation rate analyses to correct for these low correlations (see below).

Field experimental arrays

After the fifth generation of selection (sixth in the case of KBS exertion lines), four plants from each family (anther exertion, $n = 288$; stamen dimorphism, $n = 320$) were grown in a stratified random design in the KBS greenhouse and exertion or dimorphism was measured on each; this was done in spring 2001 for the exertion selection lines and spring 2002 for the dimorphism lines. One plant from each family was selected to produce the most uniform distribution of exertion and dimorphism possible across all lines, resulting in a group of 72 plants for the exertion lines and 80 for the dimorphism lines. For the pollinator observations, the exertion selection plants were divided into three arrays of 24 plants each in the summer of 2001, and the dimorphism selection plants into four arrays of 20 plants each in the summer of 2002. Field work was done at KBS, which consists of a mixture of old fields, woodlands and agricultural fields. The visitors to wild radish at this site are primarily small native bees and syrphid flies (Sahli and Conner, 2007).

Pollinator observations

On 20 d from 18 June to 30 August 2001 and 28 d from 13 June to 21 August 2002, one group of plants was transported to the field site; thus, each array was in the field seven times (six times for one array in 2001), spaced across the normal core flowering season of wild radish. On each day of observation, floral traits were measured from one recently opened flower on each plant, the height of the plant was measured with a metre stick from the base of the plant to the top of the central inflorescence, and the number of open flowers on the plant was counted.

The plants were placed randomly in a rectangular grid with 1 m spacing between plants (4×6 in 2001, 4×5 in 2002). Each plant was observed for 10 min, and all pollinators foraging on flowers were recorded. Total observation times were 76 h and 40 min in 2001 and 84 h and 10 min in 2002. For a sub-set of focal visitors (see the Results), we recorded the number of flowers probed by that visitor, the total duration of its visit and whether it was foraging for pollen or nectar. Flies and most bees feed only on pollen because their proboscides are too short to reach the nectar at the bottom of the 12 mm corolla tubes. The mouthparts of butterflies only allow them to feed on nectar. Honey-bees and long-tongued bees (families Anthophoridae and Megachilidae) forage for both pollen and nectar; nectar feeding is obvious because the tongues are <12 mm, so the bees struggle to insert their heads into the corolla tube. Mean time per flower was calculated as the total time a visitor spent on a plant divided by the number of flowers probed.

Statistical analysis

Individual plants were the unit of analysis; thus, for all analyses, we averaged the floral and pollinator visitation measurements across each of the six or seven field days for each plant. Exertion of long and short filaments was calculated as the length of the filament minus the length of the corolla tube. Analysis of variance was performed to test for the effect of the artificial selection on mean exertion compared with the control lines.

Insect visitors were lumped into three groups (bees, flies and butterflies) to ensure adequate sample sizes for analysis; other visitors (wasps, beetles and ants) totalled <3 % of visitors and were excluded from the analyses. Two response variables were used to assess the effect of floral morphology on pollinator behaviour: the average number of pollinators visiting the plant per 10 min observation period and the average time spent foraging per flower probed. The latter was obtained from the observations on the sub-sample of focal visitors (see above). The residuals were checked for normality using the Box–Cox transformation family (MASS package for R; Venables and Ripley, 2002); based on this test, time per flower was log-transformed for the significance testing.

To analyse the effect of anther position on attraction to pollinators, we used analysis of co-variance (ANCOVA) of the number of visits and time per flower against both long and short filament exertion separately for the 2001 and 2002 experiments. In order to control for the effect of advertisement other than anthers, we added both petal area and number of open flowers as predictor variables. Poisson link-function was implemented in the model to account for the nearly Poisson distribution of the number of visits. For all analyses, we used generalized linear models (GLMs) performed in R version 3.1.2 (R Development Core Team, 2014) using RStudio. All data are available from the Dryad Digital Repository: <http://dx.doi.org/10.5061/dryad.lgc03>.

RESULTS

Effects of artificial selection on anther exertion

Selection on long stamen anther exertion resulted in significant differences among treatment groups for this trait and for

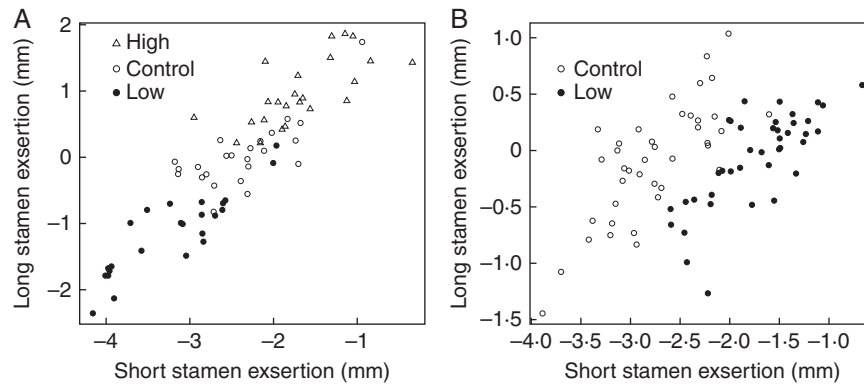


FIG. 1. Values of long and short stamen exertion in experimental populations of wild radish. Points represent mean values across flowers for each plant used in the field pollination experiment. Exsertion was measured as the difference between filament length and tube length, hence, negative values indicate insertion of the bottom of the anther into the corolla tube. Because the anthers are 2 mm long on average, an exertion of -2 means that the top of the anther is even with the tube opening, and values above this mean at least part of the anther is exerted. (A) Exsertion selection experiment ($n = 72$); plants were selected either to increase exertion ('High') or to decrease exertion ('Low'). (B) Dimorphism selection experiment ($n = 80$); plants were selected for decreased dimorphism ('Low').

TABLE 1. Mean short and long stamen anther exertion in the exertion selection experiment (2001) and dimorphism selection experiment (2002)

Stamen	2001 High	Control	Low	2002 Control	Low
Short filament exertion	-1.67 ± 0.07	-2.37 ± 0.06	-3.21 ± 0.08	-2.75 ± 0.06	-1.73 ± 0.05
Long filament exertion	0.98 ± 0.07	0.03 ± 0.06	-1.14 ± 0.08	-0.09 ± 0.06	-0.07 ± 0.05

Values are the mean \pm s.e. of exertion of the anthers, expressed as the difference between filament length and tube length (in mm).

In 2001 all means were significantly different among treatments for both short and long stamens (ANOVA: $F_{2,69} = 40.57$, $P < 0.001$ and $F_{2,69} = 94.17$, $P < 0.001$, respectively; Tukey post-hoc: $P < 0.001$ for all comparisons).

In 2002 mean length of the long filament was not different between treatments (Student's t -test: $t_{74.6} = 0.17$, $P = 0.866$), but short filament length was different between treatments ($t_{77.6} = 9.29$, $P < 0.001$).

short anther exertion through a correlated response (Fig. 1A; Table 1; see also Conner *et al.*, 2011). In the dimorphism selection experiment, where the target of selection was the relative lengths of the two filaments, no difference was found in mean long stamen exertion, but short stamen exertion was significantly less in the selection lines compared with the control (Fig. 1B; Table 2). These mean differences among lines caused increased variance in the traits when lines were combined for the field studies, increasing our power to test the effects of exertion on pollinators. Thus, variance in both short and long stamen exertion was increased relative to controls alone (representing the natural population) in the 2001 study, and only variance in short stamen exertion was increased in the 2002 experiment (Fig. 1; Table 2B).

In order to test for an increase in variance, we performed an F -test of equality of variances (Snedecor and Cochran, 1989). In the 2001 exertion selection experiment, the variance in exertion of short filaments in the full array, including both control and selection lines, was 0.74, more than 2.5 times higher than the variance measured in control lines (0.296; $F_{47,23} = 3.287$, $P = 0.003$); the latter represent the natural variation. Variance in long filament exertion was 1.447 in selection lines, about six times larger than in control lines (0.242, $F_{47,23} = 5.977$, $P < 0.001$; Fig. 1A). Nonetheless, the overall variance in short filaments across all the plants in this experiment was expanded, compared with variance in the control

plants alone. Variance was 0.497 for the full array of all the plants, 2.25 times higher than control lines alone ($F_{79,39} = 2.249$, $P = 0.006$). Variance of long filaments was not significantly increased by the selection treatment, where all the plants combined exhibited variance of 0.216, only 1.25 times larger than control lines ($F_{79,39} = 1.256$, $P = 0.436$; Fig. 1B).

Composition of pollinators

The composition of pollinators differed between years (Table 2): in the 2001 study on the exertion selection lines, flies were the majority of visitors, while in the 2002 study on the dimorphism lines bees were dominant. Bees in both years foraged mostly for pollen (69 and 88 % of bee visits in 2001 and 2002, respectively).

Effect of anther position on pollinator visitation rates

The results from both years supported the hypothesis that more exerted anthers increase attractiveness to pollen foragers, with the effect on the dominant pollinator and the traits with expanded variation in each year being statistically significant. In 2001, increased exertion of both short and long stamen anthers resulted in an increased number of fly visits per plant (Fig. 2A, B). In 2002, number of visits of bees per plant increased with

increased short stamen exertion (Fig. 2C), but long stamen exertion did not have a significant effect (Fig. 2D); recall that variance in long filament exertion was not increased significantly in this experiment.

Effect of anther position on duration of visits

In 2001, time per flower was higher for pollen feeders (mean visit duration 35.3 s, $n = 135$) compared with nectar feeders (mean visit duration 18.0 s, $n = 83$; $t = 3.68$, d.f. = 193.23, $P < 0.001$; data were log-transformed for significance testing).

TABLE 2. Number of visitors recorded in wild radish plants in each year, including the sub-set that were tracked and their behaviour recorded in detail (see the Materials and Methods)

	2001		2002	
	Recorded	Tracked	Recorded	Tracked
Flies	1249	488	811	317
Bees	437	269	3825	1229
Butterflies	137	84	13	5
Other taxa	21	5	18	5
Total	1844	846	4667	1556

2001: exertion selection experiment; total observation time 76 h and 40 min.

2002: dimorphism selection experiment; total observation time 84 h and 10 min.

Recall that individual bees were classified as either pollen or nectar feeders based on their behaviour during each visit. Visit time per flower was not significantly affected by anther exertion and there was no significant interaction between foraging mode and exertion of either of the stamens (ANCOVA: $P > 0.3$; Table 3; Fig. 3A, B). In 2002, the mean difference was opposite to 2001 – time per flower was significantly higher for nectar feeders (mean visit duration 33.8 s, $n = 65$) compared with pollen feeders (mean visit duration 26.1 s, $n = 160$; $t = -2.82$, d.f. = 131.56, $P = 0.005$). The long stamen exertion by foraging mode interaction was marginally significant (Table 3), and the visit duration of nectar feeders declined significantly with increasing long stamen exertion (Fig. 3D).

DISCUSSION

In this study, we used artificial selection to expand the variance of anther position within a species (Fig. 1) and found that pollinator visitation was affected by this expanded variance in anther exertion, which suggests that anther position functions in pollinator attraction as well as efficiency. The effect of floral traits on pollinator behaviour has been shown in many studies across many angiosperm families (Fægri and van der Pijl, 1979; Proctor *et al.*, 1996; Johnson and Steiner, 2000; Willmer, 2011). These studies usually rely on natural floral variation in wild populations, and only a few studies have experimentally expanded trait variation beyond the natural range of the traits, using inter- or intra-specific hybrids (e.g. Bradshaw and

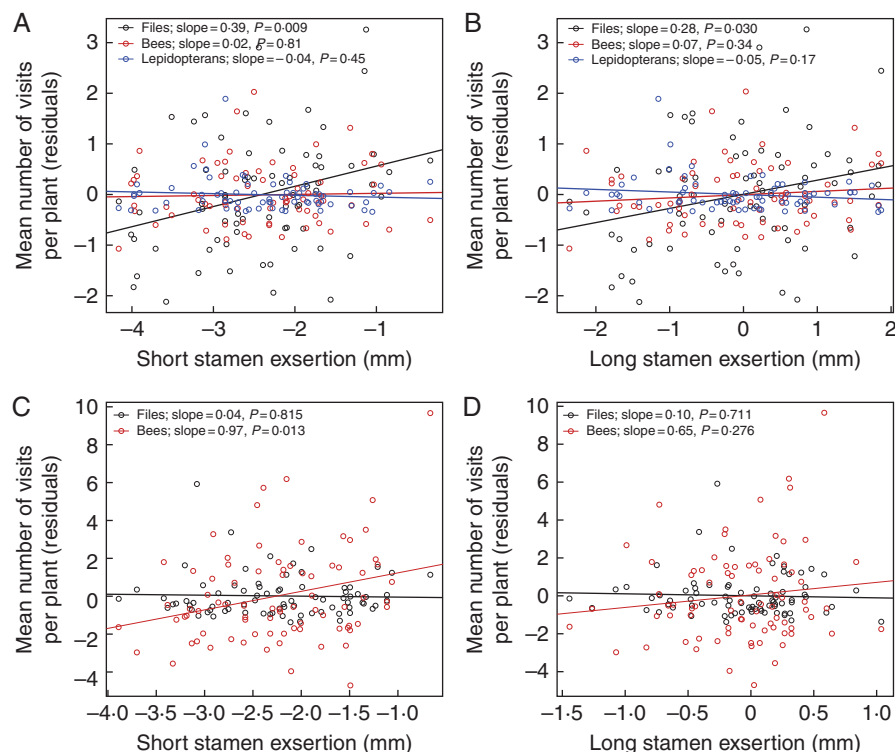


FIG. 2. Effect of short and long anther exertion on mean number of visits to plants by bees and flies. Data are residuals after controlling for petal area and number of open flowers per plant. In 2002, butterflies were not included because of the low number of visits (see Table 2). P -values for the significance of the difference from zero of the regression slopes are denoted for each visitor type in the key. (A and B) Effects of short and long stamen exertion in the 2001 exertion selection experiment. (C and D) Effects of short and long stamen exertion in the 2002 dimorphism selection experiment.

Schemske, 2003; Martin *et al.*, 2008; Sapir, 2009; Anton *et al.*, 2013). We used artificial selection within a species to achieve a similar increase in statistical power (Conner, 2003). Previous work on wild radish, including one study using the selection lines used here, found evidence for selection on stamen exertion (Morgan and Conner, 2001; Sahli and Conner, 2011), and here we provide evidence for one possible mechanism of this selection by examining the effects of anther exertion on the behaviour of the main pollinators of wild radish. This enables us to test hypotheses regarding the role of anthers as

TABLE 3. Analysis of covariance for the effect of foraging mode (pollen or nectar collection) and short and long stamen exertion on duration per flower

Experiment	Source	d.f.	Sum of squares	F	Significance
2001	Forage	1	14.7	12.46	$P < 0.001$
	Long exertion	1	0.27	0.23	$P = 0.630$
	Short exertion	1	0.98	0.83	$P = 0.364$
	Forage \times long exertion	1	0.51	0.44	$P = 0.510$
	Forage \times short exertion	1	1.03	0.03	$P = 0.868$
	Residuals	212	1302.5		
2002	Forage	1	5.1	7.29	$P < 0.001$
	Long exertion	1	1.1	1.50	$P = 0.222$
	Short exertion	1	0.5	0.71	$P = 0.401$
	Forage \times long exertion	1	1.95	2.79	$P = 0.096$
	Forage \times short exertion	1	0.1	0.17	$P = 0.680$
	Residuals	219	2147.3		

Duration per flower data were log-transformed for significance testing.

advertisement, as well as their interference for nectar-foraging insects.

Our first hypothesis was that anthers are used as a direct signal of the reward to pollen foragers and thus are a visual attraction trait, as suggested by studies on the UV absorbance and reflectance by anthers (Nakanishi, 1982; Langanger *et al.*, 2000). Support for this hypothesis in *R. raphanistrum* was found here: increased exertion of short filaments in both years and of long filaments in 2001 significantly increased the number of visits by the dominant pollinator (Fig. 2A–C). Recall that the main group of insects that visited flowers each year was either exclusively or mainly foraging for pollen (flies in 2001 and bees in 2002). To our knowledge, this is the first report that anther position, largely considered as an efficiency trait (Kudo, 2003; Castellanos *et al.*, 2004; Armbruster *et al.*, 2009; Conner *et al.*, 2009), is also used as an advertisement.

As with any study, it is possible that the increased attraction is not caused by anther exertion, but rather by unmeasured traits that are genetically correlated to anther exertion, and thus responded to our artificial selection. However, by focusing on exertion, which is a difference between two dimensions, we remove inter-trait correlations that are caused by floral size; indeed, there were no significant responses to selection for reward and display traits (see the Materials and Methods). Further, because we included display traits in the models, we are correcting for unmeasured traits correlated with these as well. Given these facts, it is likely that the effect of anthers on pollinator behaviour is direct rather than through correlation with other traits.

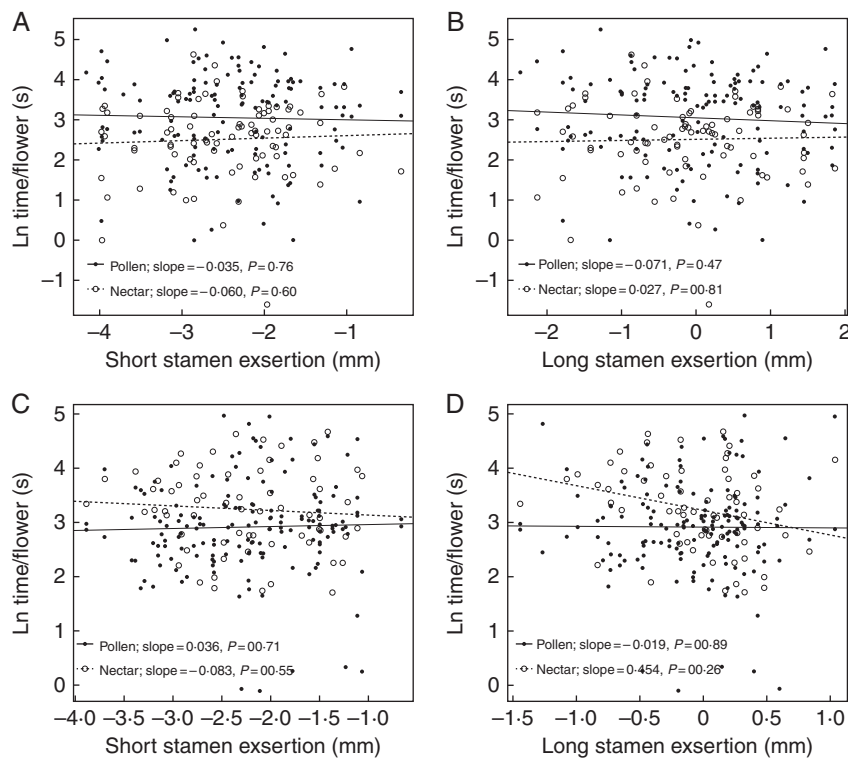


FIG. 3. Mean duration of insect visits to single flowers as a function of stamen exertion and foraging mode (pollen or nectar). (A and B) Effects of short and long stamen exertion in the 2001 exertion selection experiment. (C and D) Effects of short and long stamen exertion in the 2002 dimorphism selection experiment. Values of time per flower are log-transformed.

Our second hypothesis was that increased anther exertion increases the duration of nectar foraging visits through interference. This hypothesis was not supported; the only significant effect of exertion was a decrease in time per flower of nectar feeders with increasing long stamen exertion in 2002 (Fig. 3D). In 2001, we did not find evidence for the effect of anther exertion on duration of visit per flower (Fig. 3A, B), while in 2002 we did observe a decrease in the duration per flower of nectar foragers with increase of exertion of the long stamens (Fig. 3D). This evidence is opposite to our expectation and requires further detailed observation to understand this effect.

A few studies have tried to assess the adaptive role of anther position in Brassicaceae, usually proposing pollination efficiency as an adaptive explanation (Conner and Via, 1993; Conner and Sterling, 1995; Conner et al., 1995; Morgan and Conner, 2001). Here we show that anther position can act as a visual attraction, providing an alternative adaptive role for anther position, other than efficient pollen delivery. Pollen presentation as a visual attraction, as well as interference of anthers with visiting pollinators, had been proposed in the past (Lunau, 2000; Cocucci et al., 2014; Ren and Bu, 2014). However, our study is the first to test the combination of both attraction and interference and to partition it between two foraging modes, which suggest a possible mechanism of selection on anther position. Moreover, while past studies examined the natural variation of anther position, this is the first study that we are aware of that expanded the variation beyond the natural limits of the traits by using a few generations of artificial selection. These effects found here might not have been uncovered without the expanded variation in anther position traits introduced experimentally through artificial selection.

SUPPLEMENTARY DATA

Supplementary data are available online at <https://academic.oup.com/aob> and consist of Table S1: correlations between anther position traits and advertisement traits.

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