Effects of pollination limitation and seed predation on female reproductive success of a deceptive orchid

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Abstract. For many species of conservation significance, multiple factors limit reproduction. This research examines the contributions of plant height, number of flowers, number of stems, pollen limitation and seed predation to female reproductive success in the deceit-pollinated orchid, *Cypripedium candidum*. The deceptive pollination strategy employed by many orchids often results in high levels of pollen limitation. While increased floral display size may attract pollinators, *C. candidum*'s multiple, synchronously flowering stems could promote selfing and also increase attack by weevil seed predators. To understand the joint impacts of mutualists and antagonists, we examined pollen limitation, seed predation and the effects of pollen source over two flowering seasons (2009 and 2011) in Ohio. In 2009, 36 pairs of plants size-matched by flower number, receiving either supplemental hand or open pollination, were scored for fruit maturation, mass of seeds and seed predation. Pollen supplementation increased proportion of flowers maturing into fruit, with 87% fruit set when hand pollinated compared with 46% for naturally pollinated flowers. Inflorescence height had a strong effect, as taller inflorescences had higher initial fruit set, while shorter stems had higher predation. Seed predation was seen in 73% of all fruits. A parallel 2011 experiment that included a self-pollination treatment and excluded seed predators found initial and final fruit set were higher in the self and outcross pollination treatments than in the open-pollinated treatment. However, seed mass was higher in both open pollinated and outcross pollination treatments compared with hand self-pollinated. We found greater female reproductive success for taller flowering stems that simultaneously benefited from increased pollination and reduced seed predation. These studies suggest that this species is under strong reinforcing selection to increase allocation to flowering stem height. Our results may help explain the factors limiting seed production in other *Cypripedium* and further emphasize the importance of management in orchid conservation.

Keywords: Conservation; orchid; plant reproduction; plant–insect interactions; pollen limitation; pollination ecology; reproductive trade-offs; seed predation; supplemental pollination.

Introduction
The complex dynamics between plants, pollinators and seed predators, and how these interactions affect plant reproduction, are important in understanding the evolution of plant floral displays (Irwin and Brody 2011). While many have examined the roles of both mutualists and antagonists in plant reproduction (Gómez 2003; Strauss and Irwin 2004; Abdala-Roberts et al. 2009;
Burkhardt et al. 2009; Carlson and Holsinger 2010; Kolb and Ehrlén 2010; Ågren et al. 2013), relatively few have examined the effect of these interactions on the selection of floral traits (Cariveau et al. 2004; Parachnowitsch and Caruso 2008). Both mutualists and antagonists may forage on individuals based on floral display size, where larger floral displays provide concentrated resources for pollinators (Peakall 1989; Brody and Mitchell 1997; Mitchell et al. 2004) as well as seed predators (Stephens and Myers 2012). The goal of this study was to assess the relationships among mutualists, antagonists and floral display size, and how these biotic interactions influence reproductive success in a deceptive pollination system of conservation concern.

Maximizing pollen transfer efficiency has greatly shaped the evolution of a multitude of floral forms and functions in angiosperms (Barrett 2003). Increased floral display size is expected to increase pollinator attraction and visitation (Peakall 1989; Burd 1995; Aragón and Ackerman 2004; Grindeland et al. 2005; Li et al. 2011; Sletvold and Ågren 2011). In an experimental manipulation of floral display size in *Mimulus ringens*, Mitchell et al. (2004) found that bumblebee pollinators strongly respond to floral display size, probing more flowers in sequence on plants with large numbers of flowers. Larger floral displays also increase plant visibility, thereby increasing the attraction of pollinators from greater distances (Kindlmann and Jersákova 2006). Flowering stem height may similarly increase visibility and pollination efficiency. Synchronous flowering within a plant may also effectively increase floral display size, potentially increasing individual visibility, while synchronous flowering on the population level may increase conspecific competition for the limited pollinator resources (Crone and Lesica 2004; Crone et al. 2009). Measures of plant size such as numbers of stems and leaves, which may or may not correlate with floral display size, while not directly affecting the recruitment of pollinators, represent increased photosynthetic resources available for subsequent fruit maturation (Bazzaz et al. 1987).

Many orchids rely on a deceptive pollination strategy, a pollination strategy in which the flower provides floral cues indicating a food reward while not providing that reward (Faegri and van der Pijl 1971; Cozzolino and Widmer 2005). Deceptive pollination systems often show lower visitation and pollination relative to rewarding relatives (Nilsson 1980, 1984). To account for this reduced visitation and pollination, some have hypothesized that deception reduces geitonogamous pollination by causing pollinators to forage non-rewarding patches (Smithson 2002; Johnson et al. 2004; Schiestl 2005; Kindlmann and Jersákova 2006; Sun et al. 2009), referred to as the outcrossing hypothesis (Jersákova et al. 2006). Orchids relying on food deception often depend on newly emergent or otherwise inexperienced insects for pollinator services (Jersákova et al. 2006). In Dactylorhiza iaponica, Sletvold et al. (2010) demonstrated strong pollinator-mediated selection on spur length and plant height in the open lawn community of a Norwegian fen. This and other studies of deceptive species (O’Connell and Johnston 1998; Johnson and Nilsson 1999; Gigord et al. 2001) indicate that a variety of floral traits (plant height, flower number, petal colour) may be targets of selection for increasing female reproductive success through pollinator visitation.

Pollen limitation, defined as the difference in seed production between open pollinated (natural pollination with no supplementation) and supplemental pollination treatments, occurs when the average open-pollinated seed production is significantly less than the average seed production of individuals receiving supplemental pollen (Knight 2003). While increased floral display size may reduce pollinator limitation, it may also substantially increase geitonogamy (de Jong et al. 1993; Barrett and Harder 1996; Snow et al. 1996; Galloway et al. 2002). Geitonogamy reduces female function by reducing the number or quality of the offspring, but also impacts male function by reducing the quantity of pollen available for export to other plants, also known as pollen discounting (Harder and Barrett 1995; Barrett 2003; Johnson et al. 2004).

After pollination, female reproductive success may be eroded by seed predation. Pre-dispersal seed predation can play an important role in determining fecundity and long-term population persistence (Louda and Potvin 1995; Russell et al. 2010). Chronic seed predation rates can limit population growth by reducing fecundity, while stochastic predation rates can play a more diffuse, but equally important role in population dynamics (Kolb et al. 2007). Just as pollinators are often attracted to large floral displays, seed predators may be attracted to the accompanying large ovule resource (Brody and Mitchell 1997; Kudoh and Whigham 1998; Galen and Cuba 2001; Irwin et al. 2003; Adler and Bronstein 2004; Shimamura et al. 2005; Stephens and Myers 2012) as many seed predators rely on ovule development to feed their offspring (Cariveau et al. 2004). Therefore, although dense floral resources may attract mutualist pollinators, the accompanying dense floral and fruit resources may simultaneously attract antagonist herbivores and seed predators, creating conflicting selective pressures (Louda and Potvin 1995; Strauss and Irwin 2004; Ågren et al. 2008). Similar to the influence of co-flowering species on plant–pollinator interactions, rates of pre-dispersal seed predation also can be influenced by community context through their attraction to other species that share seed predators (Recart et al. 2013).
Cypripedium species are deceptive, deciduous, terrestrial orchids with growth emerging from a subterranean rhizome (Stoutamire 1967). Cypripedium candidum Muhlenberg ex. Willdenow, the Small White Lady's Slipper, with yellow-green lateral sepals and petals with a white, purple-spotted labellum (Stoutamire 1967), occurs in calcareous prairies as well as fens and limestone barrens (Cusick 1980). The plants occur as single plants or in large clumps (1–12 vegetative stems) containing a single large flower per vegetative stem. This floral architecture, combined with a short flowering period, severely limits the probability of reproduction by restricting opportunities for pollination. Cypripedium candidum flowers are pollinated by small (4–6 mm long) adrenid and halictid bees (Catling and Knerer 1980; Bowles 1983; Wake 2007). While pollen transport distances have not been specifically studied for adrenid and halictid bees, previous studies have found pollinia transport distances to vary widely (Nilsson et al. 1992; Kropf and Renner 2008). In the larger (8–17 mm long) Andrena bee species, Kropf and Renner (2008) found a maximum transport distance of 6.9 m. Data collected in a separate study (Walsh 2013) in Ohio C. candidum found pollen dispersal to be limited to 1 m within the focal plant.

The primary antagonist for C. candidum, Stethobaris ovata (Family: Curculionidae; subfamily: Baridinae), is a known weevil seed predator of Cypripedium spp. and other temperate orchid genera, with reports of adults in Canadian populations feeding on emerging shoots and flower buds (Light and MacConaill 2011). Adult weevils emerge in early spring along with Cypripedium shoots, and oviposit in developing fruits and, possibly stems, resulting in either fruit abortion or near total loss of the developing embryos (Light and MacConaill 2011). Predation rates on C. parviflorum in Canada vary from 32 to 53 % among plants in a population, depending on climate and availability of fruit resources (Light and MacConaill 2002). Little is known about the life history of the weevil, although they may complete two life cycles within a growing season (M. Light, pers. comm.).

The goal of this study was to assess the effects of plant and floral display sizes on both pollination and seed predation, and understand how these factors influence female reproductive success in the long-lived, highly specialized deceptive orchid, C. candidum. Cypripedium candidum is highly dependent on full sun in open areas and, as with many prairie species, populations begin to decline with the invasion of woody plants (Curtis 1946). In addition to the potential shading effects of encroaching woody vegetation, increased heterospecific stem density has been shown to reduce pollination and population recruitment (Wake 2007). Although the major challenges in orchid conservation research reside in understanding their symbiotic associations with fungi and improving survival following propagation (Krupnick et al. 2013), effective management and restoration will require a more mechanistic understanding of how habitat changes influence all biotic interactions limiting recruitment. We hypothesized that an increased floral display size would (i) increase pollinator visitation as indicated by increasing fruit set, (ii) increase geitonogamy, resulting in increased fruit abortion, decreasing fruit maturation and offspring fitness, and (iii) increase attraction of antagonists by providing an attractive resource concentration for seed predators. To address these hypotheses, we conducted two pollen limitation experiments over the course of the 2009 and 2011 field seasons in two separate sites, examining the effects of plant size on pollinator limitation and seed predation in 2009, and the effect of pollen quality in a seed predator exclusion experiment in 2011.

**Methods**

The primary field site for this study was located in Northern Ohio (GPS coordinates available upon request). Historically C. candidum existed in at least seven Ohio counties (herbarium records OSU and BGSU). However, the Northern Ohio site is now one of the only two locations in the state where C. candidum remains (ODNR, pers. comm.). This site has a large, actively managed prairie area (~900 ha) with wooded areas intertwined. Vegetative cover at this location is dominated by Andropogon gerardii, Sorghastrum nutans and Silphium terebinthinaceum, with Viola spp., Sisyrinchium montanum and Fragaria virginiana co-flowering with C. candidum. The prairie is maintained through controlled burns in early spring approximately every 3 years (J. Windus, ODNR, pers. comm.), producing large, thriving populations (total N ~ 6000) in the calcareous soil, although our access to the area was restricted to a subset of the total population by the Ohio Division of Natural Resources.

**2009 Study**

In early May 2009, we established three randomly selected 60-m line transects through a patch of C. candidum (n > 250). The population density of C. candidum at the prairie was previously estimated at 3.26 plants m⁻² (range = 1–9 plants m⁻²; SD = 2.68) (Walsh 2013). At 5-m intervals, the plant closest to a transect was selected and a second plant of equal size (number of stems and flowers) was chosen within 0.5 m of the transect on the opposite side (total N = 72). Any plants with flowers that had already opened, or had any pollinia removed or deposited prior to set up of the experiment were
excluded from the study, resulting in the exclusion of one sample point. The standard method of assessing pollen limitation compares the female fitness of open-pollinated plants with that of plants that have had all of their flowers hand pollinated (Wesselingh 2007) to avoid redirection of resources from non-pollinated flowers, which may bias information on fruit or seed set (Knight et al. 2006; Wesselingh 2007). One of the paired plants was randomly chosen to receive a hand pollination treatment on all its flowers (mean = 2.06, SE = 0.14, range 1–7) with pollinia from a different population from a site at least 100 m away, while the other member of the pair was open pollinated. The numbers of flowering stems, total stems, leaves per stem and the height of each flowering stem (to the nearest 0.1 cm) were also recorded. Because the site was burned in 2009 ca. 1 month prior to sampling, surrounding vegetation was sparse during the flowering period, precluding the collection of surrounding vegetation data.

Each flower received a single pollinium from a mixed batch of pollinia gathered earlier the same day from a different population at least 100 m away. After hand pollination, flowers were bagged with a mesh (mesh opening 3 mm × 3 mm) to prevent accidental removal of the pollinium. Mesh bags were removed after all flowers in the experiment had dehisced (ca. 2 weeks) to allow weevil predation. Capsule development was recorded 1 month after floral dehiscence (June), as well as at maturity in August when all capsules were collected and scored for insect damage and seed production. Fruit abortion was scored as the number of fruits initiated in May minus the number of fruits matured in August. We saw no evidence of browsing or herbivory other than weevil damage during the experimental period. Mature capsules were dried at 60 °C for 3 days prior to separate weighing of capsule and seed masses. Mature capsules were scored as predated when circular insect exit holes, ca. 1 mm in diameter, weevil body parts (possibly molts) and a lack of mature seeds were observed.

2011 Study
In May 2011 prior to flower opening, we set up a second pollen limitation study on a different nearby population of similar size to the 2009 study on the same property. In this experiment, a hand self-pollination treatment was added, stem and flower number were controlled and fruits were protected from weevil predation to obtain enough fruits for analysis of the effect of pollen quality on fruit set and seed mass. Five 50-m transects were randomly located across the population. For this study, only three-flowered, three-stemmed individual and wrapped unopened flowers in a mesh to prevent visitation and weevil predation. Each stem was randomly assigned to one of three treatments: hand self-pollinated, hand-outcross pollinated (from a population >100 m away) or open pollination. Plants were checked daily for open flowers and treatments were applied when the stigma became receptive. Open-pollinated stems had mesh bags removed as soon as flowers opened, while hand-outcross and hand-selfed flowers were re-bagged after receiving appropriate treatments. Following floral dehiscence, the pollinator exclusion bags were removed. Initial fruit set was scored 2 weeks after flower dehiscence, with green, enlarged fruits scored as pollinated and pale, shrunken or missing fruits scored as a failed pollination. All fruits at this time were covered in dialysis tubing and secured at both ends to exclude insect damage. Fruit abortion was scored 4 weeks after flower dehiscence as well as at the end of the study (August 2011). We collected fruits 3 months after flower dehiscence, dried them at 65 °C for 48 h and then weighed dissected seed mass to the nearest hundred-thousandth of a gram on a Mettler AE-240 scale (Mettler-Toledo Inc.). We estimated the effect of inbreeding depression ($\delta$) on female reproductive success by calculating the mean per-family seed production for each maternal family that matured capsules on both the self and outcross pollination treatments as $\delta = 1 - (o_s/o_x)$ where $o_s$ is the seed mass produced by selfing and $o_x$ is the seed mass produced by hand-outcross (Johnston and Schoen 1994).

Data analysis. Analyses were performed using JMP v.9.0.2 (SAS Institute, Cary, CA, USA, 1989–2013). A generalized linear model (GLM) with an identity link function was used to assess the effects of treatment, average flowering stem height, number of flowers, number of stems and the interaction between treatment and average flowering stem height on percentage of initial capsule development (number of capsules produced/number of flowers), proportion of capsules matured and proportion of capsules preyed upon. The number of flowers and number of stems were not significant variables in any initial models and therefore we used pooled error terms to test for other effects in the final models. The effects of number of flowers, number of stems, average height of flowering stems, treatment and percentage of capsules produced on the probability of abortion, final fruit and seed mass were analysed using a GLM. The proportions of fruit set and predation were arcsin square root transformed and all count data were log transformed. To test for proportionality of increase in
response to size variables we used an In–ln regression, testing for a slope of one (Klinkhamer and de Jong 2005; Karron and Mitchell 2012). For the 2011 study, a GLM with an identity link function examined the effect of pollination treatments on initial fruit set, final fruit set and seed mass, followed by one-way ANOVAs to test for differences among treatment groups on fruit set, abortion rates and final seed mass. Post hoc analyses on the one-way ANOVA’s were performed using a Tukey–Kramer HSD.

Results

2009 Study

Plants in the study had an average of 2.06 flowers (SE = 0.14, range 1–7) and 3.05 stems (SE = 0.23, range 1–9) per individual. Mature flowering stems had an average height of 22.6 cm (SE = 0.47), ranging between 15.1 and 40.9 cm. The average number of leaves on a plant varied little, with an average of 3.2 leaves per stem (SE = 0.01, range 3–4). Leaf length and width were not measured in this study based on prior work (Walsh 2008) that showed little between-individual variation.

In these experiments, C. candidum showed strong pollen limitation. Initial fruit set, measured 1 month after floral dehiscence, significantly increased in plants receiving supplemental pollen compared with open-pollinated flowers (pollination treatment \( P < 0.0001 \); whole model \( P < 0.0001 \); df = 5, AIC = 73.6, \( R^2 = 0.33 \); Fig. 1). Plants that received supplemental pollen had an initial fruit set of 87 % (SE = 0.037), while plants only serviced by pollinators had substantially lower initial fruit set (mean = 46 %, SE = 0.06, Fig. 1). Number of flowers and number of stems did not significantly predict the percentage of initial or final capsules produced and were omitted from further analysis. In the subsequent GLM analysis (whole model: \( P < 0.0001 \); df = 3; AIC = 71.2; \( R^2 = 0.29 \)), initial fruit set was similarly affected by pollination treatment \( (P < 0.0001) \), but was significantly influenced by flowering stem height only for open-pollinated plants (interaction effect of treatment × the average flowering stem height, \( P = 0.02 \); main overall effect of height \( P = 0.11 \)).

Final fruit set, measured 3 months post-floral dehiscence, also differed between pollination treatments (model \( P = 0.0004 \); df = 1; AIC = 121.14; \( R^2 = 0.16 \); Fig. 1). Study plants receiving supplemental pollen matured capsules 87 % (SE = 0.87) of the time, while only 46 % (SE = 0.057) of open-pollinated plants fully developed fruit (Fig. 1). However, no measured plant size traits (number of flowers, number of stems, height of flowering stems and number of leaves) or interaction effects explained variation in final fruit set (Fig. 2).

Of the flowering stems that set fruit, 73 % were preyed upon. Surprisingly, predation rates were not influenced by availability of food for the weevils (numbers of flowers or fruits) or by plant size (number of stems) (whole model \( P = 0.0024; \text{df} = 5; \text{AIC} = 93.6; R^2 = 0.23 \)). The average height of flowering stems was the only size variable to significantly explain the probability of predation \( (P = 0.002); \text{whole model: } P = 0.0020; \text{df} = 1; \text{AIC} = 98.5; R = 0.14 \), with taller stems less likely to be attacked \( (F_{1,61} = 10.1, P = 0.002, R^2 = 0.14); \text{Fig. 3} \). Fruits suffering predation had poor seed production, as the seed mass of predated capsules was 89 % lower than the seed mass of capsules without predation. A total of 22 plants in the study aborted at least one fruit between initial and final fruit set measurements. Seed mass, measured as the dried and extracted seeds from the capsules, increased

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**Figure 1.** Effect of hand and open pollination treatments on the mean per cent of initial and final fruit set. Plants receiving supplemental pollen produced higher fruit set for both the initial time period (Hand mean = 0.87, SE = 0.037; Open mean = 0.46, SE = 0.06; \( t = -4.11, n = 36, P < 0.001 \)) and final time period (Hand mean = 0.87, SE = 0.033; Open mean = 0.46, SE = 0.057, \( t = -4.30, n = 36, P < 0.001 \)).

**Figure 2.** Relationship between per cent fruit set (number of fruits/number of flowers) of open-pollinated plants and the mean height of flowering stems. Linear regression, % fruit set = \(-0.34 + 0.037 \times \text{avg. height flowering stems}, F_{1,35} = 7.0, P = 0.012, R^2 = 0.17 \).
with increased fruit production (numbers of capsules, \( P = 0.0161 \)) and decreased with the proportion of capsules predated (\( P < 0.0001 \)) (whole model \( P < 0.001; \) df = 6; AIC = 320.5; \( R^2 = 0.36 \)).

2011 Study

When fruits were protected from weevil predation and plant size and flower number were controlled, initial fruit set varied with pollen source. Plants receiving hand pollination had higher fruit set 2 weeks after floral dehiscence, with plants receiving self-pollen setting 63% of their capsules (SE = 0.08) and plants receiving outcrossed pollen setting 43% of their capsules (SE = 0.09). In contrast, fruit set on open-pollinated plants was considerably lower, with only 16% (SE = 0.06) of open-pollinated flowers initially setting fruit. Initial fruit set differed among pollination treatments (\( F_{2,89} = 7.73, P = 0.0008; \) Fig. 4).

Plants that received hand self-pollinations had a significantly higher initial fruit set than the open-pollinated plants (\( P < 0.05; \) Fig. 4).

Final fruit set, scored 12 weeks after floral dehiscence, was affected by the pollination treatments (\( F_{2,89} = 5.73, P = 0.0046; \) Fig. 4). As in the 2009 study, plants receiving natural pollinator service were pollen limited, as self and outcross hand-pollinations matured more fruits (self mean fruits per flower = 0.46, \( P = 0.0058 \); outcross = 0.4, \( P = 0.0292 \)) compared with open-pollinated stems (mean fruits per flower = 0.1; SE = 0.081). However, pollen quality did not influence the probability of fruit maturation, as the fruit set of selfed and outcrossed hand-pollination treatments were similar (\( P = 0.83; \) Fig. 4). However, pollen quality of the pollination treatment did significantly influence final seed mass (\( P < 0.0001, F_{2,89} = 18.72, \) Fig. 5). The open and outcross treatments produced significantly larger seed mass per capsule than the self-pollinated treatments (open = 0.027 g, SE = 0.002; outcross = 0.026 g, SE = 0.001; self = 0.017 g, SE = 0.001), while seed masses of the open pollinated and outcross hand-pollination treatments were similar (Fig. 5). There was no significant difference in abortion rates among treatments (\( F_{2,89} = 1.79, P = 0.172 \)). A power analysis indicated that 81 replicates would have been needed to reach a significance level of \( P < 0.05 \).

When the 12 individuals that matured fruit on both hand-pollination treatments were used to estimate the effect of inbreeding depression on seed production, most individuals showed increased seed mass when pollinated...
with outcrossed pollen (mean $\delta = 0.463$; SD = 0.31). However, the effect of selfing was variable across families. Nine families had an 11–67% reduction in seed mass when selfed, two families matured a fruit but failed to produce any selfed seed within the fruit, while one family produced similar masses of seed in both the outcross and self-hand-pollination treatments.

**Discussion**

Orchid species have been routinely shown to demonstrate consistent pollen limitation across multiple years (Snow and Whigham 1989; Ackerman and Montalvo 1990; Calvo 1990; Primack and Hall 1990; Dudash and Fenster 1997). Furthermore, deceptive orchids often produce only half as many fruits as their non-deceptive counterparts (Johnson and Bond 1997; Neiland and Wilcock 1998; Tremblay et al. 2004; Jersáková et al. 2006). This study provides strong evidence of pollen limitation in a deceptive orchid over two flowering seasons. In 2009, we observed a moderate fruit set from open pollination (46%), while fruit set was greatly reduced (16.6%) for *C. candidum* plants in 2011. This number closely parallels open-pollinated fruit set seen across multiple non-burn years at this site (16.5%; Walsh 2013), as well as levels reported in other relatives, such as 10.5% in *C. calceolus* (Kull 1998) and 5–13% in *C. acaule* (O’Connell and Johnston 1998). Hand-pollinated fruit set was consistent between study years and burn/non-burn years, with at least 40% of flowers setting fruit when supplemental pollen was provided. The mean temperature for the flowering month did not differ between 2009 and 2011 (15.5 °C); however, the 2011 study year received more than double the precipitation (18.5 cm) compared with 2009 (8.8 cm).

Pollen limitation, in principle, has two components, pollen quantity and pollen quality (Aizen and Harder 2007). In plant systems producing normal, dust-like pollen and large numbers of ovules, inadequate saturation of the stigmatic surface may result in only partial pollination. Orchids produce pollen aggregated into sac-like pollinia containing large amounts of pollen, although misplacement of the pollinia on the stigmatic surface by pollinators may result in incomplete pollination. Studies in the deceptive *Dactylorhiza* orchid have shown that flowers may need multiple visits in order to receive enough pollination for complete seed set (Sletvold and Ågren 2010; Sletvold et al. 2010). Although Aizen and Harder (2007) argue that pollen supplementation often involves high-quality outcross pollen that could inflate pollen deposition estimates, our 2011 study found that pollen quality manipulation in this system did not significantly increase fruit production, although it did significantly increase seed mass. Different experiments from both years show no significant change in seed mass/fruit between open-pollinated and hand-outcrossed flowers, suggesting that open-pollinated flowers are usually outcrossed.

Although numerous studies have shown that an increase in floral display and plant size increases pollen receipt and fruit maturation (Peakall 1989; Meléndez-Ackerman and Ackerman 2001; Aragón and Ackerman 2004; Mitchell et al. 2004; Li et al. 2011), we saw no effect of number of flowers, number of stems or number of leaves on pollen receipt or overall fruit maturation. The population studied was relatively dense for an orchid population, 3.26 plants m$^{-2}$ (range = 1–9 plants m$^{-2}$; SD = 2.68), and the large numbers of closely spaced individuals may have limited the ability to detect any effect of floral display size on fruiting success. However, we found strong evidence that greater flowering stem height increases initial fruit set, suggesting that taller plants were more likely to attract pollinators. Others have shown previously that the height and density of surrounding vegetation affects pollination and fruit production in deceptive orchids. Wake (2007) found increased seed set in *C. candidum* when surrounding vegetation was experimentally reduced, while the height of the flowering stem also increased pollination and fruit production in the closely related species *C. acaule* (O’Connell and Johnston 1998). Similarly, Sletvold et al. (2013) found strong pollinator-mediated selection for taller plants in the presence of taller vegetation in the deceptive *D. lapponica*. Given the tall grass prairie vegetation in which *C. candidum* occurs, a taller flower would be more visible to pollinators through the vegetation and therefore be more likely to receive pollinator servicing. Compared with other years (measured in a concurrent demographic survey; Walsh 2013), fruit set was unusually high in the 2009 study, which took place immediately after a controlled burn. This would suggest that both increased visibility to pollinators and increased nutrients from the burn may have contributed to this relatively high fruit set. Furthermore, taller flowering stems with increased sun exposure may offer a warmer microenvironment for the small bee pollinators in early spring, as well as greater opportunity for photosynthesis by developing fruits. Future studies on the effects of flowering stem height on seed dispersal in response to local and landscape variation in vegetation density and height may provide additional insights into the functional role of selection on plant traits affecting floral displays.

The presence of a deceptive pollination system may explain why our results with *C. candidum* are contrary to reports in the literature involving non-orchids. While other plants with larger floral displays attract pollinators from greater distances (Sih and Baltus 1987; Hessing...
1988), the absence of a reward may discourage further foraging on the same plant, limiting any increase in fitness that would otherwise occur in a large, multi-flowered rewarding plant. Jersáková et al. (2006) cite numerous examples of deceptive orchids with reduced geitonogamy, while nectar addition experiments in deceptive orchids have found dramatic increases in self-pollination when reward is added (Johnson et al. 2004; Jersáková et al. 2006; Walsh 2013). In a deceptive system, a taller stem may increase the probability of pollinator attraction, but the visitor is expected to quickly depart after receiving no compensation for its efforts. Although food deceptive systems may increase pollen limitation compared with rewarding ones, it may be more advantageous to produce fewer, but higher quality fruits than producing additional lower quality (selfed) offspring, as seen in our 2011 data (in which seed mass substantially decreased with hand self-pollination) and several others (e.g. Tremblay et al. 2004; Jersáková et al. 2006).

These data describe a potential mechanism driving the classical outcrossing hypothesis, which explains the benefits of deceptive orchid pollination via increased outcrossing (Dafni and Ivri 1979; Nilsson 1983; Ackerman 1986; Johnson and Nilsson 1999; Jersáková et al. 2006). In their review of published estimates of inbreeding depression in orchids, Sletvold et al. (2012) noted that mean inbreeding depression for seed production was 33%, regardless of mating system. While most individuals in this study produced greater seed mass on average when receiving outcrossed pollen (leading to a mean inbreeding depression of $\delta = 0.46$), this outcome was variable across all families, with one individual producing equal seed mass in both self and outcross treatments. The average number of flowers per plant within this population over a 4-year observation period was 1.75, with ~51% of flowers setting fruit (Walsh 2013), indicating that although inbreeding depression in C. candidum might be overcome by setting an additional fruit, the floral display architecture (single flower/stem) of C. candidum makes this highly unlikely. Furthermore, a true estimate of inbreeding depression would require data on the germination and future growth and reproduction of the offspring and is likely to depend on environmental conditions (Cheptou and Donohue 2011; Murren and Dudash 2012). Although seed packets have been previously used to quantify germination in the field in some orchids (Rasmussen and Whigham 1993; Sletvold et al. 2012), attempts to germinate C. candidum using this method produced no seedlings over a 2-year study period (Walsh 2013).

In this study seed predators preferentially preyed on fruit with shorter flowering stems, exerting a strong concordant selective pressure reinforcing that of the pollinators. In total, seed predation heavily reduced total reproductive output of the population, with 73% of all capsule-bearing stems attacked by a seed predator. All capsules appeared to be damaged by the same insect, most likely the weevil in the Stethobaris genus previously reported to prey upon Cypripedium fruit (Light and MacConaill 2011). Weevil predators in this genus are known to feed upon the leaves, flowers and developing capsules of many orchids, destroying most seeds by ovipositing in the maturing capsules. Contrary to our predictions, neither predation rates nor pollination success was related to other size variables such as numbers of flowers, stems and leaves. Our data suggest these weevil predators may prefer to forage on resources closer to the ground, where a lower predation risk may be associated with cover from litter and canopy vegetation, or where less energy is required to climb shorter stems to reach capsules for oviposition. Although our analyses do not indicate any response to food resource abundance such as the number of fruits on a plant, Recart et al. (2013) found a significant increase in abundance of another orchid weevil, Sethobaris polita, along with increased floral damage and reduced fruit set on a native Puerto Rican orchid, Bletia patula, in sites where an invasive orchid Spathoglottis plicata co-occurs. Complex plant–pollinator–seed predator interactions have been documented by others (Sethobaris polita 2004), although instances of conflicting pressures seem to greatly outnumber instances of concurrent pressure.

Conclusions

Our study quantifies a three-way interaction between plants, pollinators and seed predators in a deceptive orchid system in which mutualists and antagonists are exerting concordant, reinforcing selection on a plant trait, reproductive stem height. Pollinators visited plants with taller stems more often, while another trait often associated with increasing pollinator visitation, floral display size, had no effect on pollination. Furthermore, measures of plant size, such as numbers of stems and leaves that might influence resources available for fruit maturation, did not affect female reproductive success. Seed predators may be attracted to more easily reached resources that are sheltered by surrounding vegetation and less apparent to their invertebrate predators (Marquis 1992; Palo et al. 1993). Additionally, the greater seed mass from both outcrossed hand and open pollination events compared with hand self-pollinations suggests that deceit pollination effectively prevents geitonogamy, so that most pollinations in C. candidum arise predominately from outcrossing. Although variable in magnitude and based on a small sample, inbreeding depression reduced
seed mass by 11–67 % in all but one family, suggesting that conservation of small, at-risk populations that may be vulnerable to decreased pollination opportunities and increased geitonogamy should focus on facilitating outcrossing to increase recruitment.

In addition to the strong directional selective pressure on flowering stem height from both antagonists and mutualists, the height and density of the surrounding heterospecific vegetation matrix likely enhances this selection for taller flowering stems. The evolution of this complex interaction may hold an important lesson in the conservation of *Cypripedium* spp. and other deceptive plants. Management activity that controls surrounding heterospecific vegetation density and height during the flowering period may increase pollination success and fruit maturation, functioning as a cost-effective method to potentially increase rare plant offspring recruitment by modifying the pre-existing natural selective pressures on the biotic interactions of the system. However, while there have been several studies examining the effect of nectar addition on deceptive orchids (e.g. *Johnson and Nilsson* 1999; *Johnson et al.* 2004; *Jersakova and Johnson* 2006), there has been no study to date explicitly examining the demographic consequences of the deceptive pollination strategy in any plant. Although further research across multiple years and populations is needed to quantify the relative importance of pollination limitation and seed predation, orchid restoration efforts may benefit from research on whether management can be specifically targeted to ameliorate chronic pollination limitation or seed predation in orchids. As recently argued by *Sletvold and Aãgren* (2014), spatial and temporal variation in selection mediated by the biotic environment strongly affects the extent of pollinator-mediated selection. We propose that the surrounding vegetation context that provides the arena for plant–pollinator–predator interactions, as well as seed dispersal, is an important, relatively understudied component that should be considered for both managers of species of conservation concern as well as biologists seeking greater understanding of the evolution and functional significance of floral traits in these complex interactions.

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**Contributions by the Authors**

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**Conflicts of Interest Statement**

None declared.

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**Literature Cited**

*A. Abdala-Roberts*, L. *Parra-Tabla*, V. *Salinas-Peba*, L. *Herrera*. 2009. Noncorrelated effects of seed predation and pollination on the perennial herb *Ruellia nudiflora* remain spatially consistent. *Biological Journal of the Linnean Society* 96:800–807.

*Ackerman JD*. 1986. Mechanisms and evolution of food-deceptive pollination systems in Orchids. *Lindleyana* 1:108–113.

*Ackerman JD*, Montalvo AM. 1990. Short- and long-term limitations to fruit production in a tropical Orchid. *Ecology* 71:263–272.

*Adler LS*, Braunstein JL. 2004. Attracting antagonists: does floral nectar increase leaf herbivory? *Ecology* 85:1519–1526.

*Aãgren J*, *Ehrlén J*, Solbreek C. 2008. Spatio-temporal variation in fruit production and seed predation in a perennial herb influenced by habitat quality and population size. *Journal of Ecology* 96:334–345.

*Aãgren J*, *Hellström F*, *Toräng P*, *Ehrlén J*. 2013. Mutualists and antagonists drive among-population variation in selection and evolution of floral display in a perennial herb. *Proceedings of the National Academy of Sciences of the USA* 110:18202–18207.

*Aizen MA*, Harder LD. 2007. Expanding the limits of the pollen limitation concept: effects of pollen quantity and quality. *Ecology* 88:271–281.

*Aragón S*, *Ackerman JD*. 2004. Does flower color variation matter in deception pollinated *Psychilis monensis* (Orchidaceae)? *Oecologia* 138:405–413.

*Barrett SCH*. 2003. Mating strategies in flowering plants: the out-crossing–selfing paradigm and beyond. *Philosophical Transactions of the Royal Society B: Biological Sciences* 358:991–1004.

*Barrett SCH*, Harder LD. 1996. Ecology and evolution of plant mating. *Trends in Ecology and Evolution* 11:73–79.

*Bazzaz FA*, Chiariello NR, Coley PD, Petelka LF. 1987. Allocating resources to reproduction and defense. *BioScience* 37:58–67.

*Bowles ML*. 1983. The tallgrass prairie orchids *Platanthera leucophaea* (Nutt.) Lindl. and *Cypripedium candidum* Muhl. ex Willd.
Walsh et al. — Pollen limitation and seed predation in Cypripedium candidum

some aspects of their status, biology, and ecology, and implications toward management. Natural Areas Journal 3:14–37.

Brody AK, Mitchell RJ. 1997. Effects of experimental manipulation of inflorescence size on pollination and pre-dispersal seed predation in the hummingbird-pollinated plant Ipomopsis aggregata. Oecologia 110:86–93.

Burd M. 1995. Pollinator behavioural responses to reward size in Lobelia deckeni: no escape from pollen limitation of seed set. Journal of Ecology 83:865–872.

Burkhardt A, Delph LF, Bernasconi G. 2009. Benefits and costs to pollinating, seed-eating insects: the effect of flower size and fruit abortion on larval performance. Oecologia 161:87–98.

Calvo RN. 1990. Inflorescence size and fruit distribution among individuals in three Orchid species. American Journal of Botany 77:1378–1381.

Cariveau D, Irwin RE, Brody AK, Garcia-Mayeya LS, von der Ohe A. 2004. Direct and indirect effects of pollinators and seed predators to selection on plant and floral traits. Oikos 104:15–26.

Carlson JE, Holsinger KE. 2010. Natural selection on inflorescence color polymorphisms in wild Protea populations: the role of pollinators, seed predators, and intertrait correlations. American Journal of Botany 97:934–944.

Catling PM, Knerer G. 1980. Pollination of the small white lady’s-slipper (Cypripedium candidum) in Lambton County, southern Ontario. The Canadian Field-Naturalist 94:435–438.

Cheptou PO, Donohue K. 2011. Environment-dependent inbreeding depression: its ecological and evolutionary significance. New Phytologist 189:395–407.

Cozzolino S, Widmer A. 2005. Orchid diversity: an evolutionary consequence of deception? Trends in Ecology and Evolution 20:487–494.

Crone EE, Lesica P. 2004. Causes of synchronous flowering in Astragalus scaphoides, an iteroparous perennial plant. Ecology 85:1944–1954.

Crone EE, Miller E, Sala M. 2009. How do plants know when other plants are flowering? Resource depletion, pollen limitation and mast-seeding in a perennial wildflower. Ecology Letters 12:1119–1126.

Curtis JT. 1946. Use of mowing in management of white ladyslippers. Journal of Wildlife Management 10:303–306.

Cusick AW. 1980. Cypripedium candidum Muhl ex Willd. Ohio Heritage Database.

Dafni A, Ivri Y. 1979. Pollination ecology of, and hybridization between, Orchis coriophora L. and O. calluna Sol. ex Russ. (Orchidaceae) in Israel. New Phytologist 83:181–187.

De Jong TJ, Waser NM, Klinkhamer PGL. 1993. Geitonogamy: the neglected side of selfing. Trends in Ecology and Evolution 8:321–325.

Dudash MR, Fenster CB. 1997. Multiyear study of pollen limitation and cost of reproduction in the iteroparous Silene virginica. Ecology 78:484–493.

Faeigi K, van der Pijl L. 1971. The principles of pollination ecology. Oxford, NY: Pergamon Press.

Galen C, Cubo J. 2001. Down the tube: pollinators, predators, and the evolution of flower shape in the alpine skypilot, Polemonium viscosum. Evolution 55:1963–1971.

Galloway LF, Cirigliano T, Gremski K. 2002. The contribution of display size and dichogamy to potential geitonogamy in Campanula americana. International Journal of Plant Sciences 163:133–139.

Gigord LDB, Macnair MR, Smithsonian A. 2001. Negative frequency dependent selection maintains a dramatic flower color polymorphism in the rewardless orchid Dactylorhiza sambucina (L.) Soo. Proceedings of the National Academy of Sciences of the USA 98:6253–6255.

Gómez J. 2003. Herbivory reduces the strength of pollinator-mediated selection in the Mediterranean herb Erysimum mediohispanicum: consequences for plant specialization. The American Naturalist 162:242–256.

Grindel J, Otte T, Sletvold N, Imms RA. 2005. Effects of floral display size and plant density on pollinator visitation rate in a natural population of Digitalis purpurea. Functional Ecology 19:383–390.

Harder LD, Barrett SCH. 1995. Mating cost of large floral display in hermaphrodite plants. Nature 373:512–515.

Hessing MB. 1988. Geitonogamous pollination and its consequences in Gernium caespitosum. American Journal of Botany 75:1324–1333.

Irwin RE, Brody AK. 2011. Additive effects of herbivory, nectar robbing and seed predation on male and female fitness estimates of the host plant Ipomopsis aggregata. Oecologia 166:681–692.

Johnson SD, Bond WJ. 1997. Evidence for widespread pollen limitation of fruiting success in Cape wildflowers. Oecologia 109:530–534.

Irwin RE, Brody AK. 2011. Additive effects of herbivory, nectar robbing and seed predation on male and female fitness estimates of the host plant Ipomopsis aggregata. Oecologia 166:681–692.

Irwin RE, Strauss SY, Storz S, Emerson A, Guibert G. 2003. The role of herbivores in the maintenance of a flower color polymorphism in wild radish. Ecology 84:1733–1743.

Jersákova J, Johnson SD. 2006. Lack of floral nectar reduces self-pollination in a fly-pollinated orchid. Oecologia 147:60–68.

Jersákova J, Johnson SD, Kindlmann P. 2006. Mechanisms and evolution of deceptive pollination in orchids. Biological Reviews of the Cambridge Philosophical Society 81:219–235.

Johnson SD, Bond WJ. 1997. Evidence for widespread pollen limitation of fruiting success in Cape wildflowers. Oecologia 109:530–534.

Johnson SD, Nilsson LA. 1999. Pollen carryover, geitonogamy, and the evolution of deceptive pollination systems in orchids. Ecology 80:2607–2619.

Johnson SD, Peter CI, Ager J. 2004. The effects of nectar addition on pollen removal and geitonogamy in the non-rewarding orchid Anacamptis morio. Proceedings of the Royal Society B: Biological Sciences 271:803–809.

Johnston MO, Schoen DJ. 1994. On the measurement of inbreeding depression. Evolution 48:1735–1741.

Karron JD, Mitchell RJ. 2012. Effects of floral display size on male and female reproductive success in Mimulus ringens. Annals of Botany 109:563–570.

Kindlmann P, Jersákova J. 2006. Effect of floral display on reproductive success in terrestrial orchids. Folia Geobotanica 41:47–60.

Klinkhamer PGL, de Jong T. 2005. Evolutionary ecology of plant reproductive strategies. Cambridge, UK: Cambridge University Press.

Knight TM. 2003. Floral density, pollen limitation, and reproductive success in Trillium grandiflorum. Oecologia 137:557–563.

Knight TM, Steets JA, Ashman T-L. 2006. A quantitative synthesis of pollen supplementation experiments highlights the contribution of resource reallocation to estimates of pollen limitation. American Journal of Botany 93:271–277.

Kob A, Ehrlén J. 2010. Environmental context drives seed predation-mediated selection on a floral display trait. Evolutionary Ecology 24:433–445.

Kob A, Ehrlén J, Eriksson O. 2007. Ecological and evolutionary consequences of spatial and temporal variation in pre-dispersal seed
predation. Perspectives in Plant Ecology, Evolution and Systematics 9:79–100.
Kropf M, Renner SS. 2008. Pollinator-mediated selfing in two deceptive orchids and a review of pollinium tracking studies addressing geitonogamy. Oecologia 155:497–508.
Krupnick GA, McCormick MK, Mirenda T, Whigham DF. 2013. The status and future of orchid conservation in North America. Annals of the Missouri Botanical Garden 99:180–198.
Kudoh H, Whigham DF. 1998. The effect of petal size manipulation on pollinator/seed-predator mediated female reproductive success of Hibiscus moscheutos. Oecologia 117:70–79.
Kull T. 1998. Fruit-set and recruitment in populations of Cypripedium calceolus L. in Estonia. Botanical Journal of the Linnean Society 126:27–38.
Li P, Huang BO, Pemberton RW, Luo YB, Cheng J. 2011. Floral display influences male and female reproductive success of the deceptive orchid Phaius delavayi. Plant Systematics and Evolution 296:21–27.
Light MHS, MacConaill MC. 2011. Potential impact of insect herbivores on orchid conservation. European Journal of Environmental Sciences 1:115–124.
Louda SM, Potvin MA. 1995. Effect of inflorescence-feeding insects on the demography and lifetime of a native plant. Ecology 76:229.
Marquis RJ. 1992. Selective impact of herbivores. In: Frits RS, Simms EL, eds. Plant resistance to herbivores and pathogens, ecology, evolution and genetics. University of Chicago Press, 301–325.
Meléndez-Ackerman EJ, Ackerman JD. 2001. Density-dependent variation in reproductive success in a terrestrial orchid. Plant Systematics and Evolution 227:27–36.
Mitchell RJ, Karron JD, Holmgquist KG, Bell JM. 2004. The influence of Mimulus rings floral display size on pollinator visitation patterns. Functional Ecology 18:116–124.
Murren CJ, Dudash MR. 2012. Variation in inbreeding depression and plasticity across native and non-native field environments. Annals of Botany 109:621–632.
Neiland MRM, Wilcock C. 1998. Fruit set, nectar reward, and rarity in the Orchidaceae. American Journal of Botany 85:1657–1671.
Nilsson LA. 1980. The pollination ecology of Dactylorhiza sambucina (Orchidaceae). Botaniska Notiser 133:367–385.
Nilsson LA. 1983. Mimesis of bellflower (Campanula) by the red hel-leborine orchid Cephalanthera rubra. Nature 305:799–800.
Nilsson LA. 1984. Anthecology of Orchis morio (Orchidaceae) at its outpost in the north. Nova Acta Regiae Societatis Scientiarum Upsaliensis 3:167–179.
Nilsson LA, Rabakonandrianina E, Pettersson B. 1992. Exact tracking of pollen transfer and mating in plants. Nature 360:666–668.
O’Connell LM, Johnston MO. 1998. Male and female pollination success in a deceptive orchid, a selection study. Ecology 79:1246–1260.
Palo TR, Gowda J, Högberg P. 1993. Species height and root symbiosis, two factors influencing ant herbivore defense of woody plants in East African savanna. Oecologia 93:322–326.
Parachnowitsch AL, Caruso CM. 2008. Predisperse seed herbivores, not pollinators, exert selection on floral traits via female fitness. Ecology 89:1802–1810.
Peekall R. 1989. The unique pollination of Leporella fimbriata (Orchidaceae): pollination by pseudocopulating male ants (Myrmecia urens, Formicidae). Plant Systematics and Evolution 167:137–148.
Primack RB, Hall P. 1990. Costs of reproduction in the Pink Lady’s Slipper Orchid: a four-year experimental study. The American Naturalist 136:638–656.
Rasmussen HN, Whigham DF. 1993. Seed ecology of dust seeds in-situ; a new study technique and its application in terrestrial orchids. American Journal of Botany 80:1374–1378.
Recart W, Ackerman JD, Cuevas AA. 2013. There goes the neighborhood: apparent competition between invasive and native orchids mediated by a specialist florivorous weevil. Biological Invasions 15:283–293.
Russell FL, Rose KE, Louda SM. 2010. Seed availability and insect herbivory limit recruitment and adult density of native tall thistle. Ecology 91:1081–1093.
Schiestl FP. 2005. On the success of a swindle: pollination by deception in orchids. Naturwissenschaften 92:255–264.
Shimamura R, Kachi N, Kudoh H, Whigham DF. 2005. Visititation of a specialist pollen feeder Althaeus hisbici Olivier (Coleoptera: Bruchidae) to flowers of Hibiscus moscheutos L.(Malvaceae) 1. The Journal of the Torrey Botanical Society 132:197–203.
Sih A, Baltus MS. 1987. Patch size, pollinator behavior, and pollinator limitation in catnip. Ecology 68:1679–1690.
Sletvold N, Ågren J. 2010. Pollinator-mediated selection on floral display and spur length in the orchid Gymnadenia conopsea. International Journal of Plant Sciences 171:999–1009.
Sletvold N, Ågren J. 2011. Nonadditive effects of floral display and spur length on reproductive success in a deceptive orchid. Ecology 92:2167–2174.
Sletvold N, Ågren J. 2014. There is more to pollinator-mediated selection than pollen limitation. Evolution 68:1907–1918.
Sletvold N, Grindelund JM, Ågren J. 2010. Pollinator-mediated selection on floral display, spur length and flowering phenotype in the deceptive orchid Dactylorhiza tapponica. New Phytologist 188:385–392.
Sletvold N, Grindelund JM, Zu P, Ågren J. 2012. Strong inbreeding depression and local outbreeding depression in the rewarding orchid Gymnadenia conopsea. Conservation Genetics 13:1305–1315.
Sletvold N, Grindelund JM, Ågren J. 2013. Vegetation context influences the strength and targets of pollinator-mediated selection in a deceptive orchid. Ecology 94:1236–1242.
Smithson A. 2002. The consequences of rewardlessness in orchids: reward-supplementation experiments with Anacamptis morio (Orchidaceae). American Journal of Botany 89:1579–1587.
Snow AA, Whigham DF. 1989. Costs of flower and fruit production in Tipularia discolor (Orchidaceae). Ecology 70:1286–1293.
Snow AA, Spira TP, Simpson R, Klips RA. 1996. The ecology of geitonogamous pollination. In: Lloyd DG, Barrett SCH, eds. Floral biology. New York: Chapman & Hall, 191–216.
Stephens AEA, Myers JH. 2012. Resource concentration by insects not pollinators, exert selection on floral traits via female fitness. Ecology 89:1802–1810.
Strauss SY, Irwin RE. 2004. Ecological and evolutionary consequences of multispecies plant-animal interactions. *Annual Review of Ecology, Evolution, and Systematics* **35**:435–466.

Sun H-Q, Cheng J, Zhang F-M, Luo Y-B, Ge S. 2009. Reproductive success of non-rewarding *Cypripedium japonicum* benefits from low spatial dispersion pattern and asynchronous flowering. *Annals of Botany* **103**:1227–1237.

Tremblay RL, Ackerman JD, Zimmerman JK, Calvo RN. 2004. Variation in sexual reproduction in orchids and its evolutionary consequences: a spasmodic journey to diversification. *Biological Journal of the Linnean Society* **84**:1–54.

Woke CM. 2007. Micro-environment conditions, mycorrhizal symbiosis, and seed germination in *Cypripedium candidum*: strategies for conservation. *Lankesteriana* 7:423–426.

Walsh RP. 2008. *Cypripedium* (Orchidaceae) Hybridization along a Prairie/Woodland ecotone: evidence for hybridization using morphology and genetics. MS Thesis, Bowling Green State University, USA.

Walsh RP. 2013. Pollination ecology and demography of a deceptive orchid. PhD Dissertation, Bowling Green State University, USA.

Wesselingh RA. 2007. Pollen limitation meets resource allocation: towards a comprehensive methodology. *New Phytologist* **174**:26–37.