Pollination and Nectar Larceny by Birds and Bees in Novel Forests of the Hawaiian Islands

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Abstract—The extinction of native species and introduction of non-native species may lead to the disruption of biotic interactions. Pollination is a critical ecosystem process that often requires mutualisms between animals and plants. Non-native animals may interact with native flowering plants, with the potential to pollinate or steal nectar (larceny) from flowers without pollination. In the Hawaiian Islands, many native plants have lost their original pollinators. Birds and insects are known to visit native plant flowers, but it is unclear whether they pollinate or steal nectar, whether native and non-native species differ in their interactions with flowers, and what influences visitation to flowers. On O‘ahu, we deployed camera traps and conducted in-person observations on four at-risk species of Hawaiian lobelioids (Campanulaceae). We observed birds, mammals, and insects visiting flowers, with a native bird and native bee visiting most frequently. Regardless of native versus non-native status, bees made contact with reproductive structures during most visits (90.5% of visits), while birds stole nectar during most visits (99.3% of visits). Bee and bird visitation increased with the number of flowers on focal plants. Bird visitation also increased with canopy cover and the number of nearby conspecific flowers and decreased with the number of nearby heterospecific flowers. Our results indicate that bees may pollinate plants that were historically bird-pollinated, while native and non-native birds have neutral or negative impacts on these plants. Broadly, we contribute to an understanding of how native plant pollination can be altered in changing ecosystems.

Keywords—Pollination, nectar larceny, nectar robbing, novel ecosystems, extinction driven change, invasion impacts

INTRODUCTION

Rates of species extinction and invasion are accelerating worldwide, leading to a new suite of species co-existing and forming novel ecosystems (Hobbs et al. 2009). If and how species interact, and the subsequent impacts on biodiversity and ecosystem function are often uncertain in novel ecosystems (Morse et al. 2016). For example, mutualistic interactions, such as seed dispersal or pollination, can be lost, gained, or altered in response to shifts in the community, with differential impacts on animal and plant populations (Vaughton 1996; Ollerton et al. 2009; Anderson et al. 2011). Mutualisms are known to be particularly vulnerable to environmental changes and this may be even more true on island ecosystems, where populations tend to be small and evolution has occurred in isolation (Traveset & Richardson 2006; Tylianakis et al. 2008; Mitchell et al. 2009; Kiers et al. 2010). Therefore, examining these interactions in novel island ecosystems is critical for predicting species persistence and conserving biodiversity in response to global change.

Pollination is often needed for seed production and genetic connectivity between plant populations, affecting population persistence and community structure (Castilla et al. 2017). Pollination typically requires mutualistic interactions between animals and flowering plants (Bond 1994). Pollination mutualisms can be highly specialized, with plant and animal partners
exhibiting matched functional traits (e.g., matching of bird bill and floral tube shapes) to improve gene transfer between plants and to improve nectar or pollen acquisition for animals (Krauss et al. 2017). However, pollination mutualisms can be cheated through floral larceny. Here we use the term nectar larceny; this occurs when animals obtain nectar rewards while bypassing structures for pollen transfer, with (e.g., nectar robbers) or without (e.g., nectar thieves or base workers) damaging floral tissues (Inouye 1980; Irwin et al. 2001). Nectar larceny may reduce visitation of legitimate pollinators by decreasing potential rewards and shortening floral longevity (Inouye 1980; Irwin et al. 2001). The loss of pollinators, increase in nectar larceny, or shifts in the frequency of floral interactions may increase risk of pollination limitation for plant populations (Vaughton 1996; Pratt et al. 2010; Abe et al. 2011; Anderson et al. 2011). Alternatively, the introduction of pollinators may serve as valuable substitutes for lost pollinator species (Lord 1991; Ollerton et al. 2009; Pattemore & Wilcove 2012).

The frequency of interactions between animals and flowering plants may depend on traits of the focal plant and the surrounding floral neighbourhood, including the number of flowers on the focal plant, the number of flowers in the neighbourhood, and canopy cover (Ghazoul 2005; Nottebrock et al. 2017; Santos et al. 2018). For instance, a greater number of flowers on the focal plant may increase animal visitation by increasing detection of the plant and signalling higher resource availability (Eckhart 1991; Harder & Barrett 1995; Conner & Rush 1996). An increased number of flowers in the surrounding neighbourhood may either reduce focal flower visitation, owing to competition for visitors, or increase visitation, by attracting animals to an area (Rathcke 1983; Ghazoul 2005). These outcomes may vary with pollinator type and whether the neighbourhood flowers are from conspecific or heterospecific plants (Nottebrock et al. 2017). For example, for visitors that specialize on particular plants, the number of conspecific flowers, but not the number of heterospecific flowers in the neighbourhood, is predicted to influence visitation (Lázaro et al., 2009). In contrast, for generalist visitors, the number of flowers in the neighbourhood, whether conspecific or heterospecific, may impact visitation (Lázaro et al. 2009). Furthermore, increased canopy cover can either increase visitation to plants, by providing perching sites, or can decrease visitation, by concealing flowering plants (Kolb & Ehrlén 2010; Hopson et al. 2020).

In a novel forest ecosystem of O‘ahu, Hawai‘i, where over 65% of plant cover (Vizentin-Bugoni et al. 2019), 86.7% of forest bird species (Vizentin-Bugoni et al. 2019), and 30% of bee species are non-native (Snelling 2003), we examined visitation to flowers of four at-risk Hawaiian lobelioids (Campanulaceae) (Tab. 1). We documented the

| Species | Delissea waianaeensis | Cyanea grimesiana obatae | Cyanea longiflora | Clermontia persicifolia |
|---------|----------------------|--------------------------|------------------|------------------------|
| Growth form | Woody shrub (1-2.5 m)¹ | Woody shrub (1-3.2m)² | Woody shrub (1-3m)³ | Shrub/tree (2-6m)⁴ |
| Corolla colour | White, greenish-white⁵ | Greenish-white, yellowish-white, purplish-white, striped with magenta⁵ | Dark magenta⁵ | White, greenish-white⁵ |
| Corolla length (mm) | 44-60⁵ | 55-80⁵ | 60-90⁵ | 48-60⁵ |
| Conservation status | Critically endangered (ICUN)¹ | Critically endangered (ICUN)² | Critically endangered (ICUN)³ | Near threatened (ICUN)⁴ |
| Distribution (sq km) | 32 km¹ | 12 km² | 1 km³ | Unknown |
| Estimated population number | 169¹ | 16² | 79³ | Unknown |

¹(Keir, M., Portner, T., Caraway, V.L. & Kwon, J. 2015); ²(Keir 2015); ³(Keir, M., Kwon, J., Caraway, V.L. & Weisenberger, L 2015); ⁴(World Conservation Monitoring Centre 1998); ⁵(Wagner et al. 1999)
identity of species visiting lobelioid flowers and whether native and non-native animal species made contact with or bypassed reproductive structures (stigma or pollen brush at the tip of the staminal column) while feeding on pollen or nectar. We also examined if and how the characteristics of the focal plant and the floral neighbourhood influenced visitation. Since human arrival in the Hawaiian Islands (~800 years ago) (Wilmshurst et al. 2011), forest ecosystems have undergone major changes owing to high rates of species extinction and introduction (Boyer 2008; Elphick et al. 2010). Studies of pollination in Hawai‘i have observed both native and non-native birds and insects playing a role in pollination of native plants, as well as observations of nectar larceny by native and non-native birds (Cox 1983, Junker et al. 2010, 2011, Pratt et al. 2010, Koch & Sahli 2013, Aslan et al. 2014, 2016, Hanna et al. 2014, Freed et al. 2016, Shay et al. 2016). However, rates of pollination versus nectar larceny between birds and insects visiting native Hawaiian flora, in addition to factors affecting visitation, remain uncertain. Nectar analyses and morphological traits (e.g. floral tube shape) suggest that most genera of Hawaiian lobelioids (all except Brighamia) (Walsh et al. 2019) were originally pollinated by birds with long bills, including honeycreepers (Drepanidae) and ‘ō‘ōs (Mohoidae) (Pender et al. 2014, Walsh et al. 2019). Owing to the extinction or decline in populations of historic pollinators (Elphick et al. 2010; Magnacca & King 2013) in modern forests, present lobelioid pollinators may potentially include remaining native species of honeycreeper (original pollinating species are not likely present on O‘ahu) and yellow-faced bees (Hylaenus sp.), as well as non-native birds and western honey bees (Apis mellifera) (Lammers et al. 1988, Kevin M. Roddy 1997, Koch & Sahli 2013). There is limited evidence from O‘ahu that the non-native warbling white-eye (Zosterops japonicus) and the yellow-faced bee visit lobelioid flowers, that warbling white-eyes may rob nectar, and the flowers of some lobelioid species are never visited by animals (Gardener & Daehler 2006; Cory et al. 2015). Ecological communities are rapidly changing worldwide (Hobbs et al. 2009), and to conserve native biodiversity, it will be critical to evaluate how species interactions are changing and what the impacts of novel interactions are for native species.

**Materials and Methods**

**Study Species and Sites**

We examined four endemic species of Hawaiian lobelioids - *Delisea waianaeensis*, *Cyanea grimesiana obatae*, *Cyanea longiflora*, and *Clermontia persicifolia*. All of the species are endemic to the island of O‘ahu and three are critically endangered (Tab. 1). These forest understory plants all have hermaphroditic protandrous flowers, comprising either male or female floral stages, with plants exhibiting all male, all female, or a mix of male and female stage flowers at a given time (Lammers 2005, Lammers 1991). Flowers are all tubular, tetracyclic, fundamentally pentameric, perfect, complete, dichogamous via proterandry, and epigynous. The five petals are bisymmetrically and bilabiate arranged, with a pair of dorsal petals and a trio of ventral petals. Stamens form a hollow staminal column around the style, and pollen dehiscences longitudinally and introrsely, depositing pollen inside the column. A single style is present with a pair of stigmatic lobes at its apex. As the style extends, it pushes the pollen mass out of the staminal column, once extending past the column the lobes unfurl and become receptive to fertilization. When viewed from the side, the middle of the floral tube is higher in the middle then on the ends, leading to a “humpback” appearance (Lammers 2005).

We studied plants at three mesic forest sites in the Wai‘anae Range of O‘ahu, HI - Paohole Natural Area Reserve (21°32’11.30”N 158°10’47.64”W, hereafter PAH), Kahanāki Management Unit (21°32’12.55”N 158°11’35.40”W, hereafter KAH), and ‘Ekahanui Gulch (21°26’35.90”N 158°54’27”W, hereafter EKA) (details on sites provided in (Vizentin-Bugoni et al. 2019). Plant communities at each site are novel, with over 30% of plant species being non-native. In particular, 30.4% of plant species are non-native at PAH, 40.2% are non-native species at KAH, and 92.9% of plants are non-native species at EKA (Vizentin-Bugoni et al. 2019). Furthermore, bird assemblages at our study sites comprise mostly non-native bird species (Vizentin-Bugoni et al. 2019). All observed plants occurred in restored populations, being initially grown at an off-site facility and then planted in forested areas (hereafter called out-planting plot) by land managers. We examined plants within 5 out-plantings located across sites (3 at PAH, 1 at...
KAH, and 1 at EKA). Interactions with *D. waianaeensis* and *C. grimesiana obatae* were examined at all three study sites. Interactions with *C. persicifolia* were examined at PAH and KAH, and interactions with *C. longiflora* were examined at PAH. Out-planting plots are continuously managed by the Army Natural Resources Program on O‘ahu (ANRPO) and/or the State of Hawaii Department of Land and Natural Resources (DLNR).

**Detection of Interactions**

We used two methods to detect interactions: in-person observations and game camera deployment. In-person observations were used to detect interactions with invertebrates, while game camera deployment was used to detect interactions with vertebrates. Both methods were used because some insects in our system are quite small (e.g., yellow-faced bees) and may fail to trigger game cameras during every visit. However, game cameras are reliable for larger taxa and are capable of detecting night-time visitation (Rowcliffe et al. 2011).

We conducted in-person observations of flowering plants at all three sites from May – July 2019. During observations, a single observer with binoculars was positioned 3 meters from the focal plant. In each interaction, we recorded the animal species (to family or genus), the sex of flower visited, whether the animal made contact with reproductive structures, and whether the animal entered floral tubes. In-person observations were conducted for one-hour intervals during daylight hours (8:00 to 17:00), with a median observation time of 11:00. In total, observation times were 11 hours (N = 7 plants) for *D. waianaeensis*, 10 hours (N = 6 plants) for *C. grimesiana obatae*, and 7 hours (N = 2 plants) for *C. persicifolia*. No plants were observed more than once on a given day. Owing to logistical constraints and the rarity of these plants, we monitored few individuals, and we were not able to choose only focal plants spatially segregated from each other. Further, individual plants from different focal species were sometimes monitored within the same out-planting locations. From all species examined, focal plants ranged from 1-23,053 m apart (mean = 7,951 m, median = 9,883 m), and 2-9 plants (across all plant species examined) were observed per out-planting plot (median = 5). Owing to the timing of flowering, we were unable to conduct in-person observations of *C. longiflora* and only obtained data on this species from game cameras (see below). Even with small sample sizes of in-person observations, our results indicate high variation in the pollinator community, and this work should serve as a starting point for future studies.

Game cameras were deployed on flowering plants at all three sites from April – September 2019, and February – April 2017 and 2018. Specifically, we used the Bushnell Trophy Camera Brown Model 119836, programmed to record videos for 10-15 seconds following motion detection for 24 hours/day. Cameras were placed within 1 meter of the focal plant for a duration of 6 to 54 days per plant (median: 14 days). We deployed game cameras on *D. waianaeensis* (N = 14 plants for 5,688 hours), *C. grimesiana obatae* (N = 7 plants for 19,364 hours), *C. persicifolia* (N = 7 plants for 2,640 hours), and *C. longiflora* (N = 6 plants for 2,352 hours). Some individual plants were monitored simultaneously using both in-person observations and game camera deployment (N = 5 plants for *D. waianaeensis*, N = 6 plants for *C. grimesiana obatae*, and N = 2 plants for *C. persicifolia*).

**Classification of Interactions**

We defined three types of interactions - reproductive contact, nectar larceny, and visitation - based on the behaviour of the visitor at focal flowers. Events in which animals made contact with reproductive structures (stigma or pollen brush at the tip of the staminal column) were recorded as reproductive contact, whereas events in which animals took nectar or entered floral tubes without touching reproductive structures were recorded as nectar larceny (Inouye 1980). In particular, we defined bird interactions as reproductive contact when a bird inserted its bill into a flower in such a way that the feathers on its head or body came into contact with the reproductive structures of the flower. We defined bee reproductive contact as occurring when bees landed on the reproductive structures (stigma or pollen brush depending on stage of flower). We defined bird nectar larceny as occurring when a bird accessed nectar reserves without making contact with reproductive structures, occurring when a bird inserted its bill into the front or back of the flower, often through a hole in the base of the flower. We defined bee nectar larceny as
occurring when a bee entered the floral tube without making contact with reproductive structures. Visitation events occurred when there was physical contact between animals and flowers, irrespective of whether the animal was feeding on nectar, collecting pollen, or only making physical contact with flowers without collection of nectar or pollen. Among visitation events, feeding visits occurred when animals were observed feeding on or collecting pollen or nectar. Each instance of contact between an animal and flower was recorded as a separate visitation.

**Focal and Neighbourhood Characteristics**

For plants monitored between May-July 2019 (N = 38 plants), we recorded focal plant data and conducted vegetation surveys of the surrounding neighbourhood (N = 116 vegetation surveys). For plants with game camera deployment, surveys were conducted on each plant every 14–21 days (mean = 15.7, median = 14) owing to likely changes in the number of flowers on focal plants and in the floral community over time. If we conducted an in-person observation on a plant without game camera deployment, we conducted a vegetation survey immediately before the in-person observation. In each survey, we recorded the number of flowers and sex of each flower on focal plants. From all vegetation surveys (N = 46 vegetation surveys from 38 plant individuals), 20 plants had all male flowers, 8 had all female flowers, and 18 had a mix of female and male flowers. In total, 169 male flowers and 100 female flowers were observed on the 38 plants. We also identified the species of all flowering plants and counted the number of open flowers for each species in the surrounding neighbourhood. We defined the neighbourhood as a 7 m radius surrounding each focal plant. We chose a 7 m radius because we expected that plants within this range could be perceived by animals (both birds and bees) and a greater radius would introduce more potential for perceptual obstruction by the dense vegetation. Furthermore, a 7 m radius is similar in size to other neighbourhood studies (Sargent 1990; Saracco et al. 2005; Smith & McWilliams 2014; Hopson et al. 2020). For our metrics of the number of focal plant flowers, conspecific flowers, and heterospecific flowers, we took the average number of flowers observed between two consecutive vegetation surveys to account for changes in the number of flowers over time, providing a single value for each observation interval. Upon camera setup, we measured canopy cover over focal plants using a spherical crown densiometer, positioned directly over the focal plant during measurement.

**Statistical Analyses**

For all analyses we tested different random effects structures (out-planting plot within a given site, plant ID), and used the one that explained the greatest variance in the model. We did not perform model selection, other than removing correlated variables or ones due to sample size constraints, meaning our results are based on the global models listed below. All analyses were performed in Program R 3.6.1 (R Development Core Team, 2019).

We created animal group-specific models to examine how characteristics of the neighbourhood and traits of the focal plant influenced the probability of visitation or the number of visits. Fixed effects of interest included the number of flowers on the focal plant, the number of conspecific flowers in the neighbourhood, the number of heterospecific flowers in the neighbourhood, percent of non-native flowers in the neighbourhood, and percent canopy cover. The percent of non-native flowers and the number of heterospecific flowers were highly correlated (0.8); therefore, we removed the percent of non-native flowers from all models. We conducted two separate models, the bee visitation model which was based on our in-person observations and a bird visitation model which was based on our game cameras. We conducted two separate analyses because the game cameras could not detect the smaller, native bees, while birds did not visit during in-person observations.

For the bee visitation model, we used the in-person observation data. Owing to a small sample size, we were only able to examine the probability of visitation, rather than what influenced the number of visits. We used a GLMM with a binomial error structure, a logit link function, and visitation (yes, no) as the response variable. Only half of our plants had canopy cover measures taken; thus, we first examined whether canopy cover influenced the probability of bee visitation, and because we found that it did not, we removed canopy cover from subsequent analyses to increase
sample size (N = 28 hours of observation from 15 individual plants). Our global model included the number of focal flowers, number of heterospecific flowers, and number of conspecific flowers as fixed effects, and out-planting plot as the random effect.

For bird visitation models, we used data from game camera traps, and we conducted two separate analyses owing to the data being zero-inflated (e.g., 67% of observation intervals had no visitors, and few observations with more than one visit, Zuur et al. 2010). First, we examined the probability of visitation using a GLMM with a binomial error structure, a clog-log link function, and the response variable was visitation (yes, no). Second, we examined the number of visits, given at least one visit was observed, using a GLM with a negative binomial error structure, a log link function, and the response variable was number of visits. Unlike the in-person observations, game cameras were placed on focal plants for variable lengths of time and we accounted for this in both models (see below). For the probability of visitation model, we tested a separate model for canopy cover and removed it from subsequent analyses because it did not influence the probability of visitation. Therefore, our global model included the average number of flowers on the focal plant, average number of conspecific flowers, average number of heterospecific flowers, and number of days the plant was filmed as fixed effects, and out-planting plot as a random effect (N = 49 observation intervals) from 27 plants, with an average of 1.8 observation intervals per plant). For the number of visits model, we broke this down into separate models for each variable of interest owing to the small sample size for the number of plants visited one or more times (N = 20 observation intervals). Each model had an offset of the number of days a plant was filmed, and then a fixed effect of either average number of focal flowers, average number of conspecific flowers, average number of heterospecific flowers, or canopy cover. Owing to the small sample size, the potential random effects of plant ID and out-planting plot did not explain any variation, and therefore, were not retained.

**RESULTS**

**VISITING SPECIES**

Although flowers were visited by both native and non-native species, the native O‘ahu ‘amakihī (*Chlorodrepanis flava*) was the most frequent bird visitor and the yellow-faced bee was the most frequent bee visitor (Tab. 2). We observed floral

| Animal group | Species                        | Native status | Total visitations | Nectar larceny (%) | Contact with reproductive structures (%) | Non-feeding visits (%) |
|--------------|--------------------------------|---------------|-------------------|-------------------|------------------------------------------|-----------------------|
| Bird         | O‘ahu ‘amakihī (*Chlorodrepanis flava*) | Native        | 621               | 96.3              | 0.0                                      | 3.7                   |
|              | Warbling white-eye (*Zosterops japonicus*) | Non-Native    | 103               | 84.4              | 3.9                                      | 11.7                  |
|              | ‘Apapane (*Himatione sanguinea*) | Native        | 13                | 100.0             | 0.0                                      | 0.0                   |
|              | Red-billed leiothrix (*Leiothrix lutea*) | Non-Native    | 2                 | 0.0               | 50.0                                     | 50.0                  |
| Mammal       | Rat (*Rattus sp.*)               | Non-Native    | 16                | 0.0               | 75.0                                     | 25.0                  |
| Insect       | Yellow-faced bee (*Hylaeus sp.*)  | Native        | 88                | 2.3               | 71.6                                     | 26.1                  |
|              | Western honey bee (*Apis mellifera*) | Non-Native    | 55                | 14.5              | 50.9                                     | 34.6                  |
|              | Ant (*Formicidae sp.*)           | Non-Native    | 3                 | 100.0             | 0.0                                      | 0.0                   |
|              | Moth (larva) (*Lepidoptera sp.*)  | Unknown       | 1                 | 0.0               | 0.0                                      | 100.0                 |
|              | Hoverfly (*Syrphidae sp.*)       | Unknown       | 1                 | 0.0               | 0.0                                      | 100.0                 |
visitations by the O‘ahu ‘amakihi (*Chlorodrepanis flava*), warbling white-eye (*Zosterops japonicus*), ‘apapane (*Himatione sanguinea*), red-billed leiothrix (*Leiothrix lutea*), rat (*Rattus* sp.), yellow-faced bee (*Hylaeus* sp.), western honey bee (*Apis mellifera*), ant (*Formicidae* sp.), moth larvae (*Lepidoptera* sp.), and hoverfly (*Syrphidae* sp.) (Tab. 2). Rats could not be identified to species from video footage, but potential species include the black rat (*Rattus rattus*), Norway rat (*Rattus norvegicus*), and Pacific rat (*Rattus exulans*) (Pratt et al. 2009; Shiels and Drake 2011). In visitations captured by game cameras (N = 847), 73.3% were by the O‘ahu ‘amakihi, 12.1% by the warbling white-eye, 6.5% by the western honey bee, 4.1% by the yellow-faced bee, 1.9% by rats, 1.5% by the ‘apapane, 0.2% were unknown insects, and 0.002% by the red-billed leiothrix. In visitations observed in-person (N = 59), 91.5% were by yellow-faced bees, 5.08% by ants (*Formicidae* sp.), 1.6% by moth larvae (*Lepidoptera* sp.), and 1.6% by hoverflies (*Syrphidae* sp.). Of the flowers visited by bees during the in-person observations (N = 40), 73% of visits were on male flowers and 27% were on female flowers (55 male and 22 female flowers were observed in total). Overall, bees visited flowers far more frequently than birds, with bees visiting flowers on average 1.89 visits/hour and birds visiting flowers on average 0.05 visits/hour (Tab. 3).

Visitation varied amongst the different plant species, with visitations ranging from 0 - 2.6 visits/hr (Tab. 3). One plant species, *C. longiflora*, was never visited by any species, while *C. persicifolia*, the species that is of lower conservation concern out of the four plant species, received the highest visitation rate out of all plant species.

| Animal group | Plant species               | Mean visitation rate (visits/hr) | Plants sampled | Hours sampled |
|--------------|------------------------------|----------------------------------|----------------|--------------|
| (a) Bees     | *Clermontia persicifolia*    | 2.000                            | 2              | 7            |
|              | *Cyanea grimesiana obatae*   | 1.900                            | 6              | 10           |
|              | *Delissea waianaeensis*      | 1.818                            | 7              | 11           |
| (b) Birds    | *Clermontia persicifolia*    | 0.003                            | 5              | 1,365        |
|              | *Cyanea grimesiana obatae*   | 0.103                            | 10             | 6,253        |
|              | *Cyanea longiflora*          | 0.000                            | 5              | 1,547        |
|              | *Delissea waianaeensis*      | 0.020                            | 15             | 4,446        |

Bees contacted reproductive structures in 90.6% of their feeding interactions, whereas birds contacted reproductive structures in only 0.7% of feeding visits. In particular, from a total of 106 feeding visits by bees, 96 involved landing upon reproductive structures (Fig. 1), and 10 involved nectar larceny, whereby bees crawled into floral tubes without damaging tissues. Birds contacted reproductive structures in only 5 of the 703 feeding visits observed. Birds appeared to be primarily nectar robbers, damaging floral tissues. In particular, while feeding, birds would perch at the base of the flower, and they either created small holes or used previously created holes with their bills in the bottom of the corolla (as opposed to entering the floral tube opening). Birds also split the corolla using their bills while positioned from above, allowing them access to the nectar without touching reproductive structures (Fig. 1).

Focal and neighbourhood characteristics influencing visitation

Characteristics of the focal plant and/or the neighbourhood influenced visitation, with both bees and birds increasing their probability of visitation when the focal plant had more flowers (Tab. 4, Fig. 2A-B). For bees and birds, only the number of flowers on the focal plant influenced the probability of visitation, while the number of conspecific or heterospecific flowers in the surrounding neighbourhood did not (Tab. 4). For birds, we found that characteristics of the focal plant and the neighbourhood influenced the number of visits to a focal plant, given that a bird visited at least once (Tab. 4, Fig. 2C-F). In particular, we found increased visitation to focal plants with an increased number of flowers on focal plants and an increased number of...
Discussion

In the Hawaiian Islands, at least 134 plant species (Wood et al. 2019) and 79 bird species have gone extinct since human arrival (T. K. Pratt et al. 2009). Hawaiian lobelioids, which are renowned for exemplifying coevolutionary processes with avian nectarivores, have been disproportionately affected, with 25% of species extinct and many extant species presently at-risk of extinction (Smith et al. 1995). Although Hawaiian lobelioids exhibit floral traits adapted for historic bird pollinators (Givnish et al. 2009; Pender et al. 2014), in novel forest ecosystems of O'ahu, we found that birds of modern forests were unlikely to transfer much pollen between lobelioid plants, owing to low floral visitation and infrequent contact with reproductive structures. In contrast, bees were more likely to transfer pollen, owing to more frequent visitation and contact with flowers in both male and female stages of development. Thus, bees are likely to be more important than birds for maintaining pollination in the at-risk lobelioid species we examined, while the effects of nectar larceny by birds on bee pollination remains largely unexplored. Other animal groups had low visitation rates. For example, rats infrequently visited flowers \((N = 16\) visits), during which they consumed floral tissues, likely destroying entire flower structures. Nevertheless, further research is

| Model | Predictor | \(\beta\) | SE | P-value |
|-------|-----------|---------|----|---------|
| (a) Probability of bee visitation \((R^2 = 0.54)\) | Number of flowers on focal plant | 1.88 | 1.01 | 0.06 |
| | Number of conspecific flowers in neighborhood | -0.44 | 0.65 | 0.50 |
| | Number of heterospecific flowers in neighborhood | 0.03 | 1.08 | 0.98 |
| (b) Probability of bird visitation \((R^2 = 0.74)\) | Number of flowers on focal plant | 1.04 | 0.55 | 0.06 |
| | Number of conspecific flowers in neighborhood | -1.01 | 0.93 | 0.28 |
| | Number of heterospecific flowers in neighborhood | 0.62 | 0.59 | 0.30 |
| | Number of days filmed | 0.65 | 0.52 | 0.21 |
| (c) Univariate models* | Number of flowers on focal plant \((R^2 = 0.83)\) | 0.91 | 0.32 | <0.01 |
| | Number of conspecific flowers in neighborhood \((R^2 = 0.74)\) | 2.79 | 0.97 | <0.01 |
| | Number of heterospecific flowers in neighborhood \((R^2 = 0.90)\) | -0.41 | 0.23 | 0.08 |
| | Percentage of canopy cover above focal plants \((R^2 = 0.65)\) | 3.26 | 0.95 | <0.01 |

*All four models included an offset of the number of days the plant was filmed.
needed to assess the impacts of these infrequent visitors. Furthermore, native and non-native species appeared to have similar effects on the flowers (potential for either pollination or larceny). Thus, the more frequent visitation by native species, compared to non-native species, to lobelioid flowers did not necessarily lead to more pollination. We showed that characteristics of the focal plant and the surrounding neighborhood altered bird and bee visitation. Because Hawaiian lobelioids are often out-planted strategically by land-managers (James 2009), the manipulation of these characteristics could improve conservation strategies. More broadly, our results demonstrate that in novel ecosystems, both native and non-native species may help fill the functional role of extinct species.

**NECTAR LARCENY BY BIRDS**

We recorded visitation to lobelioids by two native bird species, the ʻapapane and Oʻahu ʻamakihi, and two non-native bird species, the warbling white-eye and red-billed leiothrix (Tab. 2). However, pollination was unlikely; instead, birds were mostly committing nectar larceny. Nectar larceny by birds generally occurs in specialized flowers with long floral tubes that are difficult for birds with small bills to reach (Rojas-Nossa et al. 2016). In our study, bird species had small bills (13.12 – 16.66 mm culmen length, Gleditsch & Sperry 2019) compared to the floral tube lengths of the lobelioid species we examined (45 – 90 mm corolla length) (Wagner et al. 1999). Previous studies have reported nectar larceny of Hawaiian lobelioids by warbling white-eye and Oʻahu ʻamakihi (Gardener & Daehler 2006, Aslan et al. 2014). To our knowledge, we are the first to describe nectar larceny by ʻapapane on lobelioids (Tab. 2). Oʻahu ʻamakihi and ʻapapane are known to feed upon nectar of other native plants (Pratt et al. 2010) and our work indicates nectar from lobelioid flowers is another food resource for the remaining native birds. Nevertheless, as found in other studies on the Hawaiian Islands, we found that bird visitation to lobelioids was infrequent (Lammers et al. 1988). Therefore, we would expect that the impact of nectar larceny by birds on legitimate pollination is low, but given the high abundance and widespread distribution of these birds in Hawaii (BirdLife International 2017), future work should examine whether nectar larceny by these species impacts pollination of Hawaiian plants.

**Figure 2.** Model results from bird and bee visitation models, with the number of flowers on the focal plant affecting (A) the probability of bee visitation, (B) the probability of bird visitation, and (C) the number of visits by birds. The number of visits by birds was influenced by (D) the number of conspecific flowers, (E) the number of heterospecific flowers, and (F) the percentage of canopy cover. The green shading is the 95% CIs from the model output.
Reproductive contact by bees

We found that both native yellow-faced bees and non-native western honey bees visited plants frequently and made contact with reproductive structures during visits to the three lobelioid species (D. waianaeensis, C. grimesiana obatae, and C. persicifolia) we observed in-person. The majority of observed bee visits (all in-person observations) were by yellow-faced bees. In Hawai’i, yellow-faced bees carry less pollen but exhibit higher pollinator fidelity compared to western honeybees, and western honeybees often forage on pollen of invasive plants (Miller et al. 2015). Thus, yellow-faced bees are expected to be more effective pollinators of native plants compared to western honeybees, but more research is needed to compare bee species (Miller et al. 2015). Future studies should, for instance, examine pollinator effectiveness on different plants and for different bee species, considering pollinator fidelity and movement patterns. Yellow-faced bees have been observed visiting flowers of two other lobelioid species on O‘ahu (Gardener & Daehler 2006). We found that bees visited both male and female flower phases, suggesting that pollination events are likely occurring. We also detected nectar larceny in 2.3% of visits by yellow-faced bees. Although nectar larceny by yellow-faced bees has not been previously reported, nectar larceny is common in other bee species (Roubik 1982; Hanna et al. 2013). At least 63 endemic yellow-faced bee species are known, 33 of which are species of conservation concern, and 7 species are listed as endangered by the United States Fish and Wildlife Service (Daly & Magnacca 2003; Fish and Wildlife Service 2016). Because they tend to specialize on few flowering plant species (Wilson et al. 2010), lobelioid pollen and nectar may be a critical food resource for some yellow-faced bee populations.

Visitation and influence of the focal plant and neighbourhood

Floral visitation is highly variable across ecosystems worldwide and may depend on factors ranging from site-level characteristics to features of entire biogeographic regions (Primack & Inouye 1993; Ollerton et al. 2009). Novel ecosystems may be particularly susceptible to changes in floral visitation from loss or gain of pollinator species (Ollerton et al. 2009). In our system, visitation was low for all plants examined, but bees visited lobelioids much more frequently than birds. Thus, the overall impact of bees on lobelioid populations is likely greater than impacts from birds. Because all lobelioids examined are of conservation concern, low visitation rates suggest potential for pollination limitation as a threat to remaining populations. For instance, one plant species, Cyanea longiflora, received zero visitation. Owing to low visitation rates in bees and nectar-robbing by birds, the functional roles of extinct lobelioid pollinators are unlikely being completely substituted by animals in modern forests.

Characteristics of the focal plant and the neighbourhood influenced visitation by birds and bees suggesting that these traits alter the probability of detecting and visiting a plant. In particular, bird and bee visitation increased when there were more flowers on the focal plant, likely owing to a larger visual signal to animals and the potential for higher rewards (Baude et al. 2011). Furthermore, bird visitation increased with increased canopy cover and conspecific flowers in the neighbourhood, while decreasing with increased heterospecific flowers in the neighbourhood. Increased canopy cover increases perching locations, reduces temperatures, and may improve visual cues by shading flowers, all of which may increase the amount of time birds spend in these areas (Primack & Inouye 1993, Altshuler 2003, Champlin et al. 2009, Hopson et al. 2020). Further, we expect that floral density of conspecific and heterospecific plants surrounding focal plants serves as a visual cue for birds, affecting selection of foraging patches and individual plants within foraging patches (Sargent 1990). Conspecific flowers in the neighbourhood may facilitate interactions with focal plants by cueing in nectarivores to lobelioid nectar (Elliott & Irvin 2009). In contrast, heterospecific flowers in the neighbourhood may decrease bird visitation due to interspecific competition between plants (Totland et al. 2006; Bruckman & Campbell 2014). Further work is needed to evaluate floral neighbourhood relationships in our system and the behaviours of different visiting species.

Island ecosystems and the introduction of non-native floral visitors

Island ecosystems are particularly vulnerable to species extinction and invasion, and the introduction of non-native pollinators can have
varying effects on pollination processes (Traveset & Richardson 2006). Pollination success generally declines for native flora following the extinction of historic pollinators and introduction of non-native pollinators, owing to differences in the roles of species lost and gained (Traveset & Richardson 2006; Mortensen et al. 2008). For instance, on both New Zealand and Guam, where many or all native pollinators have gone extinct, a suite of new species pollinate native plants, but seed set, fruit production, and juvenile recruitment are lower in these novel communities compared to intact communities nearby (Lord 1991; Mortensen et al. 2008; Kelly et al. 2010; Anderson et al. 2011; Pattemore & Wilcove 2012). Non-native bees, in particular, have the potential to substitute vacant pollinator roles or disrupt interactions between native pollinators and native flora. For instance, non-native honeybees substitute roles of extinct or declining native bee populations in the Ogasawara Islands and Bonin Islands (Kato et al. 1999; Abe et al. 2011). However, in New Caledonia and Australia, non-native bees have disrupted pollination by native fauna by removing the majority of pollen from native flora (Whelan et al. 2001; Kato & Kawakita 2004). Furthermore, following the extinction or population decline of avian pollinators in island ecosystems, bees often shift into dominant pollinator roles for historically bird-pollinated flora (Rosas-Guerrero et al. 2014). For example, in New Zealand, a mistletoe species that was historically bird-pollinated, now has two native bee species (Hylaeus spp. and Leioproctus spp.) that appear to pollinate the species (Robertson et al. 2005). Therefore, shifts in both the status of the pollinator (native vs. non-native) and the taxonomic group (e.g., birds vs. bees) have been observed on islands, with differential impacts on native plants.

The Hawaiian Islands have experienced high rates of pollinator extinction and introduction (Junker et al. 2010; Hanna et al. 2014; Aslan et al. 2016; Shay et al. 2016). Previous work has shown that, like other island systems, non-native birds and insects contribute to the maintenance of pollination processes for some native plant species, but they are unlikely to fully substitute the roles of lost pollinators (Walsh et al. 2019). Intact populations of native yellow-faced bees are currently maintaining pollination for many endemic plant communities (Wilson et al. 2010; Koch & Sahli 2013) and for certain plant populations, non-native western honeybees may play a critical role in pollination (Junker et al. 2010). In our study, we found that Hawaiian lobelioids that have lost their historic bird pollinators may now depend on native and non-native bees for pollination services, and the modern bird community, composed of native and non-native species, may either have no impact or negatively impact plants by stealing nectar. Thus, our findings suggest that, in relation to impacts on native flora, the taxonomic grouping (bird or bee) of the floral visitor is more important than whether or not the organism is native to the ecosystem.

Major shifts in plant-animal interactions following species extinction and introduction have likely contributed to observed population declines in lobelioids and may continue to impact the distribution and abundance of remaining populations into the future. Further, shifts in the visiting species may have differing effects on plant population fitness, which may impact the demography and evolutionary trajectories for lobelioids species over time (Zhang et al. 2007; Castro et al. 2008). Ultimately, increased understanding of how critical plant-animal interactions are responding to species extinctions and introductions will improve efforts to conserve native species in a rapidly changing world.

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