

Nectar robbing in the trainbearers (*Lesbia*, Trochilidae)

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ABSTRACT

Many flower visitors engage in floral larceny, a suite of so-called ‘illegitimate’ visits in which foragers take nectar without providing pollination services. The data on prevalence of illegitimate visits among hummingbirds, as well as the total proportion of foraging and diet that such visits comprise is broadly lacking. Here, we report the occurrence of nectar larceny in the two currently recognized species of trainbearers and analyze the proportion of plant visits categorized by mode of visitation as: robbing, theft, and/or pollination. We augment our original field observations using a trove of data from citizen science databases. Although it is difficult to distinguish primary vs. secondary robbing and theft vs. pollination, we conservatively estimate that ca. 40% of the recorded nectar foraging visits involve nectar robbing. Males appear to engage in robbing marginally more than females, but further studies are necessary to confidently examine the multi-way interactions among sex, species, mode of visitation, and other factors. Our results also indicate that the relationship between serration of bill tomia and traits such as nectar robbing or territorial defense may be more complicated than suggested in the literature. We discuss the significance of our findings in the context of recent developments in the study of nectar foraging, larceny, and pollination from both avian and plant perspectives.

INTRODUCTION

A growing list of bird species across several clades are known to forage for nectar on so-called ‘illegitimate’ flower visits, in which flower rewards are taken without the requisite provision of pollination services. Also termed ‘floral larceny,’ this mode of foraging is correspondingly gaining a broader appreciation as an important factor shaping the ecology and evolution of plant-animal interactions (Lara and Ornelas, 2001; Irwin et al., 2010; Rojas-Nossa et al., 2016; Boehm, 2018). Although species such as flower piercers (*Diglossa*, Passeriformes) are widely known to depend on nectar larceny, there are many reports of illegitimate visits by hummingbirds (Lara and Ornelas, 2001; Gonzalez and Loiselle, 2016). Some morphological characteristics of hummingbirds, including bill length and tomial serrations, are thought to be particularly closely associated with nectar larceny (Ornelas 1994, but see Rico-Guevara et al. 2019).

Nearly all plants that provide nectar and pollen rewards experience larceny, most often from insects and vertebrates, and many defend against it (Irwin et al., 2010). The remarkable frequency of illegitimate visits necessitated the development and adoption of a more precise lexicon of larceny (Inouye, 1980), which attempts to separate it into canonical modes, partly in service of conceptual clarification useful in pursuit of identifying its ecological and evolutionary causes and consequences. Thus, “primary nectar robbers” mechanically create a hole in flower tissue through which they remove nectar, bypassing the floral opening. By contrast, “secondary nectar robbers” remove nectar utilizing openings previously fashioned by primary robbers. Deceptively resembling pollinators, “nectar thieves” also access nectar through the floral opening. Due to a mismatch between flower and thief morphology or behavior, however, pollination does not take place. Finally, some plant species are vulnerable to larceny by “base workers,” visitors who probe along the base of the flower (between the free petals) and obtain nectar, while still

47 failing to effect pollination.

48 These modes of illegitimate nectaring visits may be difficult to distinguish from each other and/or
49 legitimate visits—pollination—without careful observations and manipulations. Moreover, they can
50 clearly quantitatively overlap so that, for example, thieving may merely reduce pollination efficiency
51 without eliminating it, or individual birds can engage in a mix of primary and secondary robbing. While
52 many studies report one or more modes of nectar foraging among hummingbirds (McDade and Kinsman,
53 1980; Roubik et al., 1985; Feinsinger et al., 1987; Ornelas, 1994), we lack assessments of their relative
54 importance (Snow and Snow, 1980), which are critical for a more complete understanding of the incentives
55 and behaviors that drive the observed patterns or plant-animal interactions.

56 Here, we present original observations and collect data from public photograph databases with a
57 focus on documenting the frequency of modes of nectar foraging by Green- and Black-tailed Trainbearers
58 (*Lesbia nuna* and *L. victoriae*). We discuss the significance of common occurrences of nectar robbing
59 in the context of plant-pollinator co-evolution and bill morphology. The original observations took
60 place in and around Ollantaytambo, Peru, on *Brugmansia sanguinea*, *Fuchsia boliviensis*, and *Passiflora*
61 *tripartita*. We coarsely document the proportion of pollination, theft, and primary- and secondary-robbing
62 visits to flowers, by leveraging images from citizen-science databases eBird, iNaturalist, and other pho-
63 tographed/reported occurrences in the literature. Our highly preliminary analyses find that floral larceny
64 is surprisingly common and dominated by robbing (at least ca. 40%). Specifically, if secondary rob-
65 bing is common among hummingbirds, as we suggest, assessment of the proposed associations between
66 tomial serrations and mode of nectar foraging (Ornelas, 1994; Rico-Guevara et al., 2019) could be difficult.

68 MATERIALS AND METHODS

69 Species and site description

70 The genus *Lesbia* is currently comprised of two short-billed hummingbird species, Green-tailed Train-
71 bearer, *Lesbia nuna* (Lesson, 1832), and Black-tailed Trainbearer, *Lesbia victoriae* (Bourcier & Mulsant,
72 1846). Their geographic distributions largely overlap, ranging from Colombia to Peru in montane scrub
73 and other semi-open habitats. The species are differentiated by color, specifically the extent of emerald
74 green on dorsal side and tail (more in *L. nuna*). Black-tailed Trainbearers are generally larger, as well.
75 Both species are strongly sexually dimorphic, with males in possession of a spectacularly long tail. The
76 species are difficult to differentiate and confidently assign because of a great amount of geographic
77 variation within each species (many subspecies names exist) and possible existence of undescribed species
78 (Weller and Schuchmann, 2004; Stiles, 2004). These difficulties are especially apparent in attempts to
79 identify individuals from single photographs, and we consequently conducted our analyses (see below) at
80 the generic level.

81 Description of original observation

82 On January 14th, 2019, we recorded and photographed a male Green-tailed Trainbearer, *Lesbia nuna*,
83 which nectar-robbed several flowers of *Fuchsia boliviensis* (Onagraceae) in Ollantaytambo, Peru (13°15'44"
84 S, 72°16'14" W), and then followed the same individual on a 50m foraging bout, while it nectar-robbed
85 flowers of *Brugmansia sanguinea* (Solanaceae)—a species with spectacular ca. 30cm-long tubular
86 crimson-to-yellow flowers—and then legitimately visiting *Salvia leucantha* (Lamiaceae). Both rusty and
87 black-throated flowerpiercer (*Diglossa sittoides* and *D. brunneiventris*) occur at relatively high densities
88 in the area (B.I. and P.B.F., unpub. data), are known to pierce both *Fuchsia* and *Brugmansia* species,
89 and may have been responsible for the previously made calyx and petal hole at the base of *B. sanguinea*
90 flower.

91 Literature searches

92 We examined the literature and internet resources to accumulate a database of additional instances of nectar
93 collecting visits by *Lesbia* species. First, we used internet and Google Scholar exact keyword searches
94 of the existing scientific literature, including: ‘*Datura*’ or ‘*Brugmansia*’ and ‘*Lesbia*’ or ‘trainbearer’;
95 ‘*Lesbia*’ and ‘diet’. We employed “advanced search” filtering, screening out a variety of ribald nouns, to
96 reduce the considerable volume of images with unrelated or offensive content. Documentation of any
97 details on foraging ecology of the trainbearers is sparse. We found three sources either documenting or
98 hinting at illegitimate flower visits by trainbearers. Gould (1861, p.15) cited in Ornelas (1994), contains

a passing reference to piercing of a *Brugmansia* sp. flower by a *Lesbia* sp., but the visit was originally inferred as insect gleaning, not nectar robbing, as later re-framed by Ornelas (1994), and neither the species nor locations were identified. Snow and Snow (1980) documented feeding behavior for a species identified as *Lesbia nuna* in the vicinity of Bogotá, Colombia. The observed individuals—nearly all females—visited six plant species, and were thought to either utilize holes in corolla or probe around the base of the corolla split in *Castilleja fissifolia*. In a paper documenting secondary robbing behavior in Cinereous Conebill, Vogt (2006) mentions unpublished observations of secondary nectar robbing by a trainbearer. Additionally, although we could find no papers detailing their insect foraging in any detail, 21.6% of *Lesbia nuna* feeding records comprised of insect hawking (Snow and Snow, 1980), and gut contents of a single individual included arthropods (Remsen et al., 1986). Therefore, as generally holds for hummingbirds, the trainbearers' diet at a minimum contains arthropods and nectar. The proportion of time spent on each mode of nectaring is not known.

Foraging mode collection and scoring procedures

Consequently, we examined all identified species records for the Genus *Lesbia* in two public databases—iNaturalist (iNaturalist, 2019) and eBird (Sullivan et al., 2009), and in online searches, primarily of the photo service flickr (<https://www.flickr.com>). In each case, we made last visits or downloaded database records on January 6th, 2020. For all resulting records, we applied the same methodology. Accessions, metadata, image locations, and scored data used in our analyses are available in Supplemental Materials. Those without any flowers were discarded, as were those in which birds were simply perched or flying near a flower. We then closely examined the remaining images, in which birds were hovering in close proximity (about a body length) to flowers, and facing them. For that subset of 180 records, we extracted the location, date, and comment metadata. We recorded individual sex ('m', 'f', or 'NA'), species identification (scoring specific epithets as 'nuna', 'victoriae', or 'sp.'), estimated mode of visitation (primary nectar-robbing, secondary nectar-robbing, thieving, pollination, or 'NA'), as well as our confidence in the estimate of the mode of interaction ('low' vs. 'high'; to help facilitate critical re-examinations of our results). To ensure consistent scoring of nectar foraging mode, both B.I. and P.B.F. examined all 180 records and jointly agreed on marked scoring. Finally, with the help of several colleagues, we identified and recorded many (but not all) of at least 50 visited plant species.

Often, modes of visitation are unclear and we combined terms for the most accurate characterization of visitation. For example, primary vs. secondary nectar robbing and pollination vs. thieving are generally hard to distinguish, so we simply combined the likely modes of visitation (there were only two instances of clear thieving). This enables us to conservatively summarize modes of interactions as approximate ranges, without making unnecessary and possibly incorrect assumptions. We should note here that nearly all of the combined pollination/thieving visits appear to have been legitimate visits, but establishing the extent of pollen transfer is prohibitively demanding for most studies of modes of nectar feeding. Similarly, we recorded the species identifications, but found them to be uncertain for a number of individuals, and we consequently highlight the jointly presented feeding habits for all of *Lesbia*. Published (uploaded) observations were not *a priori* assigned or randomized in any way. Due to these limitations we simply report a number of pooled summary statistics. Moreover, we find it unlikely that they suffer of a strong bias with respect to visit mode, because the mode of visit is often unclear to a photographer until after the photo is taken. Instead, we initially found it more likely that there would be a photography bias in favor of photographing males, with their outstanding long tails (68 out of 114 records with foraging mode; exact binomial test, $p < 0.03$). We examine associations and interactions with sex in part due to this possibility. As pooling over the ignored variable(s) can be misleading, so we also perform limited three-way visual analyses, but caution against their over-interpretation. We coarsely examine associations between species ID (nuna/victoriae), sex (f/m), and binarized mode of interaction with flowers (robbing/other), examining the null hypothesis that the three traits are mutually independent. For model selection, we used log-linear models implemented in R package MASS (R Core Team, 2019), and AIC scores from LRstats. The nature of our data collection and resulting exploratory analyses suggests that associations ought to be re-tested by obtaining a new set of data in advance of further testing.

Bill morphology

Short-billed hummingbirds and those with serrated tomia are thought to be more likely to nectar-rob flowers with long corolla tubes, co-adapted with long-billed hummingbirds (Lara and Ornelas, 2001). We did not find a reference to presence/absence of serrated tomia in trainbearers, so we scored this trait in 17

specimens of *Lesbia nuna* and four specimens of *Lesbia victoriae* at the Field Museum of Natural History. We used a dissecting microscope to visualize and score this trait on the available collections, as well as measure their bill lengths (tip to operculum; all performed by I.N.).

RESULTS

Field observation of robbing

In our field observations, we recorded a single adult male foraging bout, during which this individual visited flowers of three plant species. Approximately ten flowers of *Fuchsia boliviensis* were robbed (unclear if primary or secondary robbing took place) as well as two flowers of *Brugmansia sanguinea*, while five flowers of *Salvia boliviensis* were apparently legitimately visited (ruling out or confirming pollination success requires controlled experiments). Photographs collected in the course of observations were deposited in iNaturalist (<https://www.inaturalist.org/observations/37206000>; iNaturalist 2019).

Database collection

We collected and examined more than 1550 photographs of *Lesbia* and found floral visits for 180 foraging bouts (Figure 1, and Supplementary Data Table) on at least 42 flowering plant genera, identified thus far. It is unlikely that many of these instances are replicates, recording the same individual, although this is somewhat uncertain for the vicinity of Cerro de Monserrate in Bogotá, Colombia, with about a dozen documented foraging bouts.

We assigned the mode of visitation for 135 visits with some confidence, while the mode remains unknown for 45 visits, most often because pictured birds were hovering in the vicinity of a flower, immediately before or after nectaring (Table 1). The most common recorded mode was ‘pollination/thieving’ (46 visits), a category that we scored conservatively (vaguely) because it is clearly continuous, virtually impossible to confidently score separately without functional study. Ideally, one would examine pollen transfer or effected seed set, after bagging and marking flowers. Our impression, however, is that the majority of these visits at least plausibly resulted in pollen transfer. The second most common mode was combined ‘nectar robbing’ (primary and secondary robbing; 41 visits) indicating that an individual fed through a pierced hole in the side of the corolla. It was followed by ‘pollination’ (36 visits), which indicated a fairly clear legitimate interaction, ‘secondary nectar robbing’ (10 visits), and thieving (2 visits). Although we could not confidently assign any visits to the ‘primary robbing’ mode, this is entirely due to the one key limitation of our observational approach, reliance on scoring from photographs. If we conservatively lump all ‘pollination/thieving’ and ‘pollination’ observations as legitimate, such visits comprise approximately 60.7% of the total and illegitimate visits 39.3% (combining all robbing and thieving).

With a number of limitations in mind, we present a few preliminary analyses in an attempt to estimate the association of species ID, sex, and mode of visitation. When these factors are jointly considered in a three-way contingency table, we find that three variables are likely not mutually independent (AIC=56.43, LR χ^2 =13.55, d.f.=4, p <0.01). Non-robbing visits by *Lesbia nuna* females and robbing visits by *Lesbia victoriae* males have the highest residuals. The lowest AIC model posits conditional independence (mode is independent of species, given sex; AIC=48.78, LR χ^2 =1.91, d.f.=2, p =0.39), indicating that sex differences are greater than species differences. One broad interpretation is that the females generally rob less than the (territorial) males. We emphasize that this result is preliminary, and the associations may stem from a number of data limitations, especially confounding unmeasured variables, or those not considered *a priori*. For example, more *Lesbia victoriae* males were observed in urban parks where plant communities and robbing frequency may differ, compared to rural or natural areas. Moreover, a model with all two-way interactions, but without the three-way interaction of the saturated model, provides only slightly worse fit (Δ AIC=0.15). It is therefore presently unclear whether species or sex differences are more important in predicting nectar foraging modes.

Bill morphology

We examined 17 specimens of *Lesbia nuna* and found no serrated tomia. The imbalance of sex ratio (15 males) is concerning, but perhaps unimportant in this case, because serrations are more likely to be present in males (Rico-Guevara et al., 2019). The mean bill length for males was 15.9 ± 0.7 mm, and for females 14.2 ± 3.2 mm. Similarly, we found no tomial serrations in four specimens (three male, one female) of

205 *Lesbia victoriae*. Their mean bill length for males was 14.7 ± 0.4 mm, and one female measured 11.0
206 mm.

207 DISCUSSION

208 Hummingbirds are generally thought to serve as legitimate flower visitors, and effect pollination in
209 exchange for host plant nectar. There is, however, an increasing appreciation of the fact that nectar
210 obtained during illegitimate visits—including both primary and secondary robbing, as well as thieving—
211 may comprise a substantial portion of diets of many species (Lara and Ornelas, 2001; Irwin et al., 2010;
212 Boehm, 2018). Presently, the frequency distribution of nectar larceny within and among bird individuals
213 and species, and a variety of covariates (community composition, territoriality, seasonality, geography,
214 etc.) is unclear. Most reports of hummingbird nectar foraging document the presence (or imply absence)
215 of nectar robbing, and do not provide an estimate of the proportion of visits dedicated to illegitimate visits.
216 This is in large part due to the difficulty of accruing this kind of data and, when larceny occurs, separating
217 its modes.

218 Our study originated with a simple natural history observation of robbing by a trainbearer, which
219 we augmented using a trove of citizen-science data from crowd-sourced public databases. We believe
220 that such estimates are a first step in improving our understanding of the dynamics between plants and
221 pollinators, especially why and how (e.g. primary vs. secondary robbing) some individual birds or species
222 steal more than others, how larceny affects plants, and how they respond to larceny. Below, we discuss
223 the results and their immediate implications of this small-scale study and place them in the context of the
224 literature on ecology and evolution of plant-hummingbird interactions.

225 We find that both currently described species of *Lesbia* steal nectar, mostly by robbing. Unambiguous
226 illegitimate visitation constitutes a substantial proportion, around 40% of all nectar foraging visits.
227 Although Snow and Snow (1980) surveyed a relatively small area, as a part of a study with a far greater
228 scope, they found that robbing comprises 64% of all visits by *Lesbia nuna* in the northern Andes, and
229 all illegitimate visits occurred on one of five visited plant species (*Castilleja fissifolia*, Orobanchaceae).
230 Observations of nine species at this high-elevation site found that five species visited flowers illegitimately,
231 and robbing constituted 17% of all flower visits (range 0% to 64%). Plant-side studies, which estimate
232 the proportion of flowers robbed on a single plant species, uncover a wide range of robbery rates due to
233 visiting hummingbirds. For example, Maloof and Inouye (2000) reviewed reports of visitation to eight
234 plant species (by a variety of animal species), and found robbing ranged from 11% to 89%. Similarly,
235 Arizmendi (2001) found that total robbing in four highland Mexican species ranges from 9% to 58%,
236 and Navarro (2001) found that 75% of *Macleania bullata* (Ericaceae) flowers contained pierced holes
237 consistent with primary robbing. An unpublished report (D. Boose, cited in Maloof and Inouye 2000)
238 found that the Stripe-tailed Hummingbird (*Eupherusa exima*) robs 98% of *Razisea spicata* (Acanthaceae)
239 flowers, but the mode is not stated. In each of these plant-side studies, however, the proportion of
240 hummingbirds' visits comprised of each visit mode is unknown, and in some cases the primary robber is
241 not a hummingbird.

242 Despite its critical importance for understanding the animal-side foraging incentives and plant-side
243 pollination and defense strategies, primary and secondary robbing are difficult to disentangle (Vogt,
244 2006; Boehm, 2018). Snow and Snow (1980) is the only hummingbird study we found, which lists data
245 that enables a calculation of the relative frequencies of visit modes, including primary vs. secondary
246 robbing. While we may have missed existing records, it appears that such estimates are rare at best.
247 Confidently scoring the specific mode of robbing (primary vs. secondary) demands evidence bearing
248 on the question whether 'breaking-and-entering' took place, which in turn requires careful experimental
249 data collection. For example, capturing the act of piercing of flowers or documenting entry through a
250 pre-existing hole could enable unambiguous scoring of robbing mode. We find that secondary nectar
251 robbing is relatively common, comprising between 7.4% and 37.8% of the visits (Figure 1). This large
252 range estimate reflects uncertainty (not variability) due to the limitations of our approach. As secondary
253 robbing is critically dependent on primary robbing—an able robber must first make a hole—we examined
254 additional photographs of many robbed plant species, and we found extensive flower damage as well as
255 records of several species of flowerpiercers (*Diglossa*) commonly observed in the areas in close proximity
256 to trainbearer robbing locations (B.I. and P.F. unpub. data). It is therefore likely that secondary robbing
257 may represent the majority of illegitimate visits by trainbearers, especially because their range overlaps
258 with multiple species of flowerpiercers. Finally, it is possible that due to the difficulty of observation, this

259 mode of nectar foraging is underestimated in other hummingbird species.

260 From the hummingbird perspective, if primary robbing is more likely to allow access to previously
261 unvisited, relatively nectar-rich flowers, then perhaps secondary robbing ought to be less advantageous
262 (Irwin et al., 2010). Framed in this way, any secondary robbing may appear somewhat paradoxical. It is
263 possible that, instead, one or more of the common underlying assumptions are flawed. Specifically, we
264 can imagine that, for example, if they are faced with resource limitation (of flowers that are ordinarily
265 legitimately exploited), trainbearers respond by robbing or thieving from inferior resources, a situation
266 that may be exacerbated by territoriality. Although our study cannot strongly substantiate it, a trend of
267 higher robbing frequency in males than females (complicated by association with species), may be related
268 to such phenomena. On the other hand, the evolutionary game between hummingbirds, flowers, and all
269 other interactors (whole pollinator guilds, herbivores, etc.) is likely substantially more complex (e.g.
270 Maloof and Inouye 2000), broadening the set of conditions for stable non-zero frequencies of secondary
271 robbing. If we are to gain a more complete understanding of hummingbird foraging strategies, it will be
272 critical to record these patterns and more closely scrutinize the ecological context, including target plants
273 and other visitors (Richman et al., 2017).

274 Floral larceny is often targeted at flowers of plants that have close co-evolutionary relationships
275 with other legitimately pollinated animal species, such as the sword-billed hummingbird and its guild
276 of long-flowered species (Soteras et al., 2018). Some characteristics of hummingbirds, including short
277 length and tomial serrations are hypothesized to be particularly closely associated with primary nectar
278 robbing (Ornelas, 1994). The idea that the serration of bill tomia affects nectar robbing ability and
279 efficiency remains broadly untested. Recent studies suggest that the presence of serrations is sexually
280 dimorphic, with greater or exclusive expression in males, and that this feature could instead be shaped by
281 a role in plumage preening or fighting, especially male territorial defense (Rico-Guevara et al., 2019).
282 Although additional field studies of nectar robbing and examination of bill morphologies for the presence
283 of serrate tomia are needed to clarify their role, our data provide some insight. First, our data imply
284 that, if secondary nectar robbing is more common than previously thought, it may be difficult to detect
285 an association between serration and (combined) robbing observations. Serrations may well provide
286 flower-robbing utility, but they do not appear necessary for secondary robbing. Second, if males perform
287 primary robbing more frequently, then the existence of sexual dimorphism for tomial serrations is less
288 problematic for the Ornelas hypothesis. Moreover, neither sexual dimorphism nor the proposed role
289 in defense precludes simultaneous or exaptive function of serrated tomia in piercing. Consequently,
290 it will be difficult to accumulate a great weight of evidence that exclusively support or detracts from
291 this hypothesis. It is maybe instructive that many other specialized piercers, such as the Wedge-billed
292 Hummingbird (*Schistes geoffroyi* and Purple-crowned fairy *Heliothryx barroti*, fail to present sexually
293 dimorphic serrations (Rico-Guevara et al., 2019). A systematic recording of the nectar feeding mode,
294 along with a number of predictors, coupled with careful studies of feeding behavior and morphology are
295 likely to be most persuasive in examining the validity and generality of the Ornelas hypothesis.

296 Predominance of different modes of larceny is likely to be associated with differing evolutionary
297 responses from flowering plants, including variation in amount and/or time-dependence of nectar release,
298 mating system (increased reliance on self-fertilization), and defenses, both physical (spines, thicker
299 calyces or bracts) and chemical (secondary metabolites). A number of plant species visited by trainbearers
300 are members of the nightshade family (Solanaceae), known for common production of alkaloids, which
301 are broadly toxic to vertebrates. Perhaps not coincidentally, a great number and quantity of alkaloids are
302 produced by species of *Brugmansia* and *Nicotiana*, and tissue at the base of flowers (calyx) contains some
303 of the highest concentrations of such compounds (Saitoh et al., 1985; Alves et al., 2007). *Brugmansia*
304 species have not been studied in equal detail to tobacco, but at least some plants in this genus can
305 incapacitate juvenile humans, orders of magnitude larger than hummingbirds, following casual skin
306 contact (Andreola et al., 2008). Moreover, given that there is a strong association between hummingbird
307 pollination and self-fertility (Wolowski et al., 2013), stereotypically hummingbird-pollinated flowers may
308 be more commonly robbed by both avian and hymenopteran robbers. This may be especially true for
309 those flowers with relatively longer tubular corollas and large amounts of nectar (Lara and Ornelas, 2001;
310 Rojas-Nossa et al., 2016), precisely the syndrome commonly associated with hummingbird pollination.

311 Our observations and analyses point to additional needed work on systematics and ecology of *Lesbia*,
312 especially their unsettled taxonomy, feeding ecology and energy budgets. More broadly, we sorely
313 lack observational data to establish the relative frequencies of nectar feeding modes and manipulative

314 field study to obtain direct evidence for nectar feeding dynamics across all hummingbirds and other
315 nectar-feeding birds (Irwin et al., 2010). Our study documents the presence of larceny in trainbearers
316 and, more specifically, shows that floral larceny is surprisingly common. These findings indicate that
317 larceny, especially secondary robbing, may be more common among hummingbirds than is generally
318 appreciated. Furthermore, we demonstrate that citizen science databases can be fruitfully employed to
319 establish the presence of nectar robbing and coarsely examine its frequency. Such approaches can also
320 generate hypotheses and guide an improved understanding of plant-side effects and responses, including
321 pollination efficiency, the distribution of visitor interactions (Maloof and Inouye, 2000; Arizmendi, 2001),
322 plant incentives and defenses (as well as overall fitness) (Pelayo et al. 2011), and ultimately better explain
323 the dynamics in the co-evolutionary game between flowers and their visitors.

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Table 1. Cross tabulation data of nectar foraging mode, sex, and species identity for *Lesbia* spp.. ‘Other’ denotes lumped pollination and thieving, and ‘Robbing’ combines both modes of robbery. Sex identification relied on tail length and subtle color differences as a proxy, and species identification was in nearly all cases copied from the original source. Of 180 total foraging bouts captured, we assigned visit mode to 135, and each of visit (nectar foraging) mode, sex, and species to 114 shown in the table.

		Other	Robbing
<i>L. nuna</i>	Female	18	5
	Male	12	7
<i>L. victoriae</i>	Female	16	7
	Male	23	26

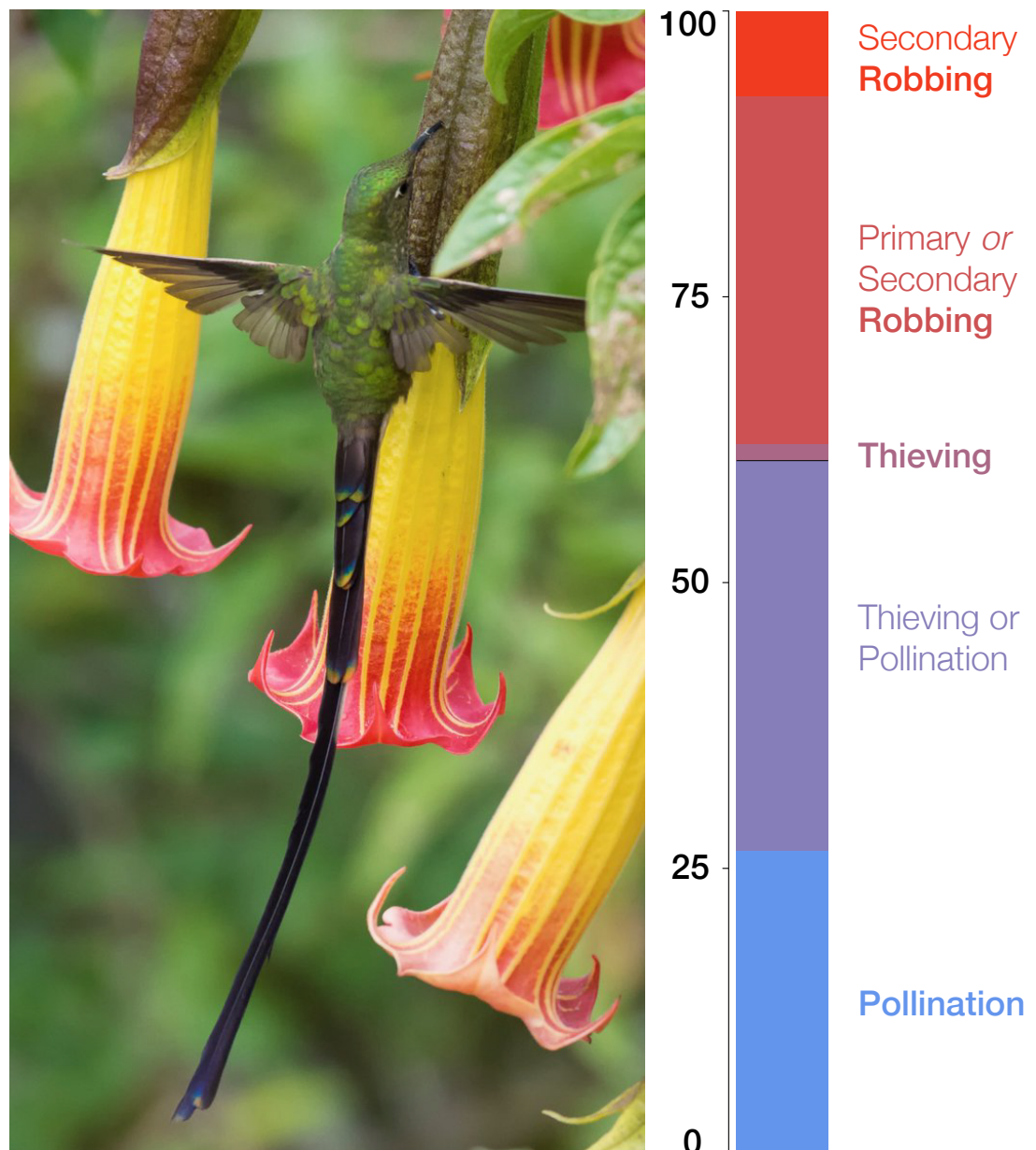


Figure 1. A nectar robbing visit by a Black-tailed Trainbearer *Lesbia victoriae* on *Brugmansia sanguinea*, whose flower was previously pierced by a species of *Diglossa* (Photo by Diego Emerson Torres; used with permission). Right panel: A bar plot illustrating the relative visit mode frequencies of *Lesbia* species (n=135). Of the 135 visits whose mode is known (could be assigned), 60.7% were legitimate (combining pollination and pollination/thieving) and 39.3% illegitimate (combining all robbing and thieving).