Male and female bees show large differences in floral preference

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14 Abstract

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1. Intraspecific variation in foraging niche can drive food web dynamics and 16 17 ecosystem processes. Field studies and theoretical analysis of plant-pollinator 18 interaction networks typically focus on the partitioning of the floral community 19 between pollinator species, with little attention paid to intraspecific variation 20 among plants or foraging bees. In other systems, male and female animals 21 exhibit different, cascading, impacts on interaction partners. Although the 22 foraging ecology of male bees is little known, we expect foraging preferences to 23 differ between male and female bees, which could strongly impact plant-24 pollinator interaction outcomes.

25 2. We designed an observational study to evaluate the strength and 26 prevalence of sexually dimorphic foraging preferences in bees.

We observed bees visiting flowers in semi-natural meadows in New
 Jersey, USA. To detect differences in flower use against a shared background
 resource availability, we maximized the number of interactions observed within
 narrow spatio-temporal windows. To distinguish observed differences in bee use
 of flower species, which can reflect abundance patterns and sampling effects,
 from underlying differences in bee preferences, we analyzed our data with both a
 permutation-based null model and random effects models.

34 4. We found that the diets of male and female bees of the same species 35 were often as dissimilar as the diets of bees of different species. Furthermore, we 36 demonstrate differences in preference between male and female bees, and 37 identify plant species that are particularly attractive to each sex. We show that 38 intraspecific differences in preference can be robustly identified within 39 interactions between hundreds of species, without precisely quantifying resource 40 availability, and despite high phenological turnover.

5. Given the large differences in flower use and preference between male
and female bees, ecological sex differences should be integrated into studies of
bee demography, plant pollination, and coevolutionary relationships between
flowers and insects.

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46 Key Words

dimorphism, dissimilarity, Morisita-Horn, phenology, plant-pollinator interaction,
pollination, pollinator habitat, preference

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50 Introduction

51 Intraspecific variation in traits and behavior, including foraging niche, has

52 important consequences for species interactions and conservation (Bolnick et al.,

53 2011; Durell, 2000). Sexual dimorphism is a large source of individual niche

54 variation, and an important factor in plant-animal interactions, such as seed

dispersal (Zwolak, 2018). Sexual dimorphism underlies adaptation, speciation,

and the way in which animals exploit their ecological niche (Butler, Sawyer, &

57 Losos, 2007; Temeles, Miller, & Rifkin, 2010). Morphological, behavioral, and life-

58 history dimorphisms can also drive the form and function of ecosystems, for

59 example when predator sex ratio drives the community composition of lower

60 trophic levels, shown experimentally (Start & De Lisle, 2018) and in nature,

- 61 where responses extended to water chemistry as well (Fryxell, Arnett, Apgar,
- 62 Kinnison, & Palkovacs, 2015).
- 63

64 Though ecological dimorphisms were first studied in vertebrates (Selander,

- 1966), they are common across taxa, including insects (Shine, 1989).
- 66 Surprisingly, in bees (Hymenoptera, Apoidea) for which both foraging (P.
- 67 Willmer, 2011) and sexual dimorphism (Alcock et al., 1978) have been well
- studied, sexually dimorphic foraging has rarely been documented. Intraspecific
- 69 variation in floral preference is well known in social (Heinrich, 1979) and to a
- 70 lesser extent, solitary bee species (Bruninga-Socolar, Crone, & Winfree, 2016;
- 71 Tur, Vigalondo, Trøjelsgaard, Olesen, & Traveset, 2014), yet most community-
- level studies focus on species-level interaction networks, and furthermore, on theforaging preferences of only female bees.
- 74

75 Male bees differ from their better-studied female counterparts in their life history 76 and ecology. Female bees construct, maintain, provision, and defend nests, 77 whereas male bees primarily seek mates (P. G. Willmer & Stone, 2004). Both 78 sexes drink floral nectar for their own caloric needs, but only females collect 79 pollen to provision young, and thus forage at greater rates. While the pollen from 80 each flower species (the term we use throughout for the flowers from a species 81 of plant) tends to be morphologically and nutritionally distinct, interspecific 82 variation in the chemical composition of nectar is comparatively subtle (Cane & 83 Sipes, 2006). Thus, we expect the foraging ecology, including floral preferences, 84 of male and female bees to differ as well.

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86 Male bees prove to be important pollinators when studied, both in specialized oil-87 or scent-collecting pollination systems (Eltz et al., 2007; Etl, Franschitz, Aquiar, 88 Schönenberger, & Dötterl, 2017; Janzen, 1971) and also when males are foraging for nectar and pollen (Cane, 2002; Cane, Sampson, & Miller, 2011; 89 90 Ogilvie & Thomson, 2015). Male bees may also be particularly relevant for bee 91 conservation. Males may be limiting in declining populations, either because 92 genetic diversity is necessary for the development of female offspring as a result 93 of complementary sex determination, or because mate or sperm limitation results 94 from poor male condition (Elias, Dorn, & Mazzi, 2010; Straub et al., 2016). As the 95 dispersing sex in most bee species, males may be crucial for gene flow and 96 metapopulation persistence even when they are not locally limiting (López-Uribe, 97 Morreale, Santiago, & Danforth, 2015; Ulrich, Perrin, & Chapuisat, 2009). 98 99 Foraging niche is only partly described by resource use. Indeed, resource 100 preferences may be more important than use alone in many contexts, including 101 conservation. Preference-the use of a resource in excess of its relative 102 availability—is challenging to measure, because both resource use and 103 availability must be known. Floral resource availability for pollinators is

- 104 particularly hard to quantify outside an experimental context because the
- 105 appropriate scale and units of floral resource availability are unclear. The

106 composition, amount, and supply rate of pollen and nectar per flower, the number 107 of flowers per inflorescence, of inflorescences per individual, and the number and 108 distribution of individual plants over the square kilometers of a bee's foraging 109 range are all important components of availability (Hicks et al., 2016). 110 Furthermore, floral availability can change rapidly over time. However, 111 differences in flower use between bees foraging at the same place and time 112 indicate differences in preference, which may occur between species, or between 113 individuals of the same species. 114 115 In this study, we assess differences between floral preferences of male and 116 female bees in the wild. We collected bees foraging on flowers in meadows in 117 New Jersey, USA. In order to observe preference differences, we collected as many individuals as possible during replicated, short (3-day) windows, during 118 119 which we assumed floral availability and bee abundance were constant at each 120 site. We compare the species composition of flowers visited by males and 121 females of the most common bee species across the entire study as a naïve 122 measure of differences in preference between the sexes. Then, using random 123 effects models, we assess when differential flower species use by male and 124 female bees likely arises from sex-specific floral *preference*, as opposed to 125 shifting overlap between foragers and floral resources (i.e. changes in availability 126 without differences in preference). Specifically, we ask

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1) How much do male and female bee diets overlap?

- 2) To what degree are particular flower species disproportionately visited by bees of one sex?
- 131 132
- 3) To what extent are differences in floral use driven by preference, rather than phenological differences between male and female bees?

134 Materials and Methods

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136 Study design and data collection137

138 Because absolute preference is nearly impossible to observe outside of an 139 experiment, we designed our study to reveal differences in preference between 140 groups of bees. In order to collect a large number of males and females from 141 many native bee species, we selected six meadows (sites) in New Jersey, USA 142 with a high abundance and diversity of flowers. These meadows were managed 143 for pollinator-attractive, summer-blooming forbs through seed addition, and a 144 combination of mowing, burning, and weed removal. Most flower species present 145 in the meadows are native to the eastern United States. We collected our data 146 during peak bloom and maximum day length (6 June to 20 August, 2016), and 147 during good weather (sunny enough for observers to see their own shadow, no 148 precipitation). We visited each site for three consecutive good weather days over 149 five evenly spaced sampling rounds in the 11-week period of our study. In all 150 analyses, we assume that bees and flowers detected at a site within one 3-day 151 sampling round co-occurred. In contrast, we assume that turnover of both plant

152 species in bloom and bee species activity can occur in the ~10 days between 153 sampling rounds.

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155 During each 3-day sampling round, an observer walked parallel transects 156 through the meadow (which ranged in size from 0.8–2.2 ha; mean=1.4 ha). 157 observing every open flower within a moving 1-m semicircle, and net-collecting 158 any bee seen actively foraging, which we defined as contacting anthers or 159 collecting nectar from a flower (Fig. S1). We collected all bee species except 160 Apis mellifera L., the domesticated western honey bee, because Apis males do 161 not forage. Observations began as soon as pollinator activity picked up in the 162 morning (7–9 am) and continued into the late afternoon or evening until pollinator 163 activity slowed substantially. Observers sampled nearly continuously, in 30-164 minute timed collection bouts with short breaks in between. If inclement weather 165 precluded a minimum of six 30-minute sampling bouts in a day, we added an 166 additional day to the sampling round as soon as weather permitted. 167

168 Flower species were identified in the field by the data collector. Bee species were 169 identified using a dissecting microscope and published keys; Jason Gibbs

170 (University of Manitoba), Joel Gardner (University of Manitoba), and Sam Droege

171 (USGS) assisted with identification for bees in the genera Andrena, Anthophora,

- 172 Coelioxys, Halictus, Heriades, Hoplitis, Hylaeus, Lasioglossum, Megachile,
- Melissodes, Nomada, Osmia, Pseudoanthidium, Ptilothrix, Sphecodes, Stelis,
 and Triepeolus, and at least one of them confirmed voucher specimens for every
- and *Triepeolus,* and at least one of them confirmed voucher specimens for every species. We determined every specimen to species except for the following four

176 complexes: Most bees in the genus *Nomada* with bidentate mandibles (*ruficornis*

group) were treated as one species. All specimens from the *Hylaeus* species

178 complex that includes *Hylaeus affinis*, *H. modestus*, and at least one additional

- species, informally dubbed "species A," were treated as a single species,
- denoted *Hylaeus affinis-modestus*, because females cannot be reliably
 distinguished. There is a cryptic species in the genus *Halictus* unlikely to occur in
 our area, *Halictus poeyi*, which is not morphologically distinct from *H. ligatus*; we
- 182 our area, *Halictus poeyl*, which is not morphologically distinct from *H. ligatus*; we 183 treat all specimens in this complex as *Halictus ligatus*. We could not confidently
- 183 separate all specimens of the two closely related *Lasioglossum* species

Lasioglossum hitchensi and *L. weemsi*. Thus, we treat all specimens from either species as one, denoted *Lasioglossum hitchensi-weemsi*. All bee specimens are

187 curated in the Winfree lab collection at Rutgers University, and the data used in

- this paper are available from the Dryad Digital Repository
- 189 http://dx.doi.org/XXXXXXX (Roswell et al.)
- 190

191 Analytical methods

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We performed all statistical analyses and simulations using R 3.5.1 (R CoreTeam, 2018).

- 195
- 196 **1)** How much do male and female bee diets overlap?
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198 To compare the diets of male and female bees, we used the Morisita-Horn index 199 of resource overlap (Horn, 1966; Morisita, 1959). This dissimilarity index 200 compares the proportion of all female bees found on each flower species to the 201 proportion of all male bees found on each flower species. In other words, it 202 compares the contribution of each flower species to female diets (where this term 203 includes the food that females collect for themselves and also to feed to young) 204 to the contribution of the same flower species to male diets. The Morisita-Horn 205 index ranges from zero (completely similar) to one (maximally dissimilar), and 206 has several good properties for our purposes. First, it uses proportions, placing 207 visits from male and female bees on the same scale, even though most visits 208 come from females. Second, it is much more sensitive to large proportions than 209 to small ones, thereby down-weighting the contribution of flower species for 210 which we have little information. Third, the Morisita-Horn estimates are resilient 211 to undersampling and uneven sample size between groups (Barwell, Isaac, & 212 Kunin, 2015).

213

214 To determine whether the male-female differences we observed exceeded those 215 expected by chance, we compared the observed compositional dissimilarity 216 between flower visits from male and from female bees to dissimilarity measures 217 from a null model that randomly permuted the bee sex associated with each flower-visit record. This permutation holds constant the total number of male and 218 219 of female visits, and the total number of visits to each flower species from both 220 sexes combined (Fig. S2). The range of dissimilarity values from this simulation 221 is the difference we would observe in our sample, if there were no true difference 222 in flower species use between males and females of the same bee species. We 223 evaluated the hypothesis that male and female diets overlap less than would be 224 expected by chance; thus we use a one-sided alpha of p<0.05. We iterated this 225 null simulation 9999 times, which was sufficient to stabilize p-values near our 226 chosen alpha (North, Curtis & Sham 2002). When the observed dissimilarity was 227 greater than 9500 of the 9999 simulated dissimilarities, we concluded that we 228 had detected a difference in the pattern of floral visitation between conspecific 229 male and female bees, given the observed diet breadth and abundance of each 230 sex.

231

232 To compare the diet overlap we observed between sexes to a meaningful 233 benchmark, interspecific diet overlap, we repeated the same null model analysis, 234 this time comparing females of the focal species to females of other species. We 235 performed one analysis for each bee species for which we collected at least 20 236 visitation records for each sex (19 species). This sample size threshold is 237 arbitrary, but null model variance shrinks with sample size, such that apparent 238 patterns for species with smaller sample sizes are rarely interpretable (Fig. S3). 239 Because we analyze 19 bee species, females of each species are compared to 240 18 others. We then compared the male-female difference (observed minus mean 241 null dissimilarity in flower communities visited) to the analogous species-species 242 difference (observed minus null dissimilarity).

244 For this analysis, which evaluates holistic differences between male and female 245 bees of the same species, we combined observations across the full season and 246 all sites. This allows us to observe foraging niche differences that are driven by 247 flower and/ or bee phenology, in addition to any sex-specific floral preference.

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249 2) To what degree are particular flower species disproportionately visited by bees 250 of one sex?

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252 This analysis uses our entire data set of 153 bee species to determine whether 253 particular flower species are disproportionately visited by male or female bees. 254 and whether the answer varies by bee species. We can infer a preference 255 difference between male and female bees for a flower species when predicted 256 odds of visitors to that flower species being male are especially high or low. To 257 do this, we use a random effects model in which bee sex is the response, and 258 flower species, bee species, site, and their interactions are random effects. We 259 statistically control for variation in the overall sex ratio across bee species 260 through a random intercept of bee species, and variation in sex ratios across 261 sites, through random intercepts for site, and the site-bee species and site-flower 262 species interactions. Because it is unlikely that, within bee species, sex ratios at 263 birth vary greatly across space, any variability attributed to site terms would likely 264 result from differential overlap of bee foraging activity and flower bloom across 265 space.

266

267 We call this model the "summed model" because we sum interactions observed 268 across the entire season (all five sampling rounds) at each site. In the summed 269 model, the relationship between phenological overlap and the odds of flower-270 visiting bees being male would be incorporated into the species effects. This 271 perspective is helpful for considering flower species' contributions to the overall 272 diets of male versus female bees. We fit the model with the R package Ime4 273 (Bates, Maechler, & Walker, 2016) with the following call: 274

275 Summed model

```
276
      Ime4::glmer(bee sex ~ (1|site)+ (1|flower species)+ (1|bee species)+
277
             (1)flower species:bee species)+ (1|site:bee species)+
278
```

```
(1|site:flower species), family="binomial", data=data)
```

279

280 We included bee species and site as random, rather than fixed, effects to directly 281 compare the variability in bee sex associated with each of these predictors to the 282 variability associated with flower species (preference). Comparing the overall 283 variability across these groups was more important to us than assessing 284 predictions on a per-site or per-bee-species basis. We fit flower species, the 285 primary covariate of interest, as a random effect to facilitate model fitting (fewer 286 degrees of freedom) as well as interpretation. In our summed model, we included 287 all two-way interactions, but omitted the three-way interaction, bee species by 288 flower species by site. Although the sort of context-dependent preference this 289 term could represent (e.g. males from bee species 1 prefer flower A at one site

290 (relative to females), but shun it at another) may exist in nature, it is unlikely we 291 would estimate it accurately in our model.

292

293 We confirmed model convergence by comparing several fitting methods using 294 the allFit function in Ime4 (Bates et al., 2016), which all showed similar parameter 295 estimates (Table S1). We tested whether residuals from our model fit were 296 overdispersed using Bolker's function "overdisp" (Bolker, 2017), and visually 297 assessed our additivity assumptions with binned residual plots (Gelman & Hill, 298 2007) (Fig. S4).

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3) To what extent are differences in floral use driven by preference, rather than 301 phenological differences between male and female bees?

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Over the 11 weeks of our study, we observed turnover in bee species. in flower 303 304 bloom, and within-bee species changes in sex ratio. Therefore, phenological 305 overlap between male versus female bees and the bloom period of particular 306 flower species, rather than preference of those bees for those flowers, may 307 explain much of the variation in sex ratio we observed across visitors. In question 308 3, we are explicitly interested in distinguishing sex-specific diet *preferences* from 309 variable use resulting from seasonal resource availability and male vs. female 310 abundance. We do this in the "seasonal model" by incorporating sampling round 311 (our measure of phenology) as an additional random intercept effect, along with 312 random intercepts for the interactions between sampling round and the other 313 covariates. We chose to include sampling round as a random effect because this 314 enables direct comparison to all other terms in both models. We ignored the 315 three- and four-way interactions between bee species, flower species, and other 316 covariates. We fit this model with the following call in the R package Ime4, with 317 new terms in bold: 318 319 Seasonal model

- 320 glmer(bee sex ~ (1|site)+(1|flower species)+(1|bee species)+
- 321 (1)flower species:bee species)+ (1)site:bee species)+
- 322 (1|site:flower species)+ (1|sampling round)+ (1|site:sampling round)+
- (1)flower species:sampling_round)+ 323
- 324 (1) bee species:sampling round)+
- 325 (1|site:bee species:sampling round)+
- 326 (1|site:flower_species:sampling_round), family="binomial", data=data)
- 327 328 Our index of preference for both the *summed model* and the *seasonal* 329 model is the change in odds that a bee is male when the flower species it visits is 330 given. To describe the importance of model terms, we calculated a bootstrapped 331 median odds ratio using code from Seth (Seth, 2017), which gives the expected 332 difference in odds that a flower-visiting bee is male between levels of a predictor 333 (Merlo et al., 2006). For example, a median odds ratio of five for the main effect 334 of sampling round would indicate that the odds of a flower-visiting bee being 335 male differ by about a factor of five between sampling rounds, while a median

336 odds ratio of one would indicate that the odds of a flower-visiting bee being male 337 do not change across rounds. If the median odds ratio is large for flower species 338 in both models, we could say that there are intrinsic (i.e. not simply phenological) 339 properties of flower species identity that male or female bees prefer. If flower 340 species is a strong predictor of bee sex in the summed model but not in the 341 seasonal one, we would still conclude that flower species often contribute more 342 strongly to the diet of one sex than the other, though these differences may not 343 arise due to differing preferences. If the sampling round terms have large median 344 odds ratios, then accounting for phenology is critical for identifying differences in 345 preference in addition to differences in use.

346

347 **Results**

348

In total we collected 18,698 bee specimens belonging to 152 bee species (table
S2) from a total of 109 flower species (table S3), which together comprised 1417
unique species-species interactions. Roughly 18% of specimens were male
(n=3372). Thus, the overall ratio of male to female bees we collected was 0.22,
although this ratio varied markedly between flower species (Fig. 1).

- 354
- 355 How much do male and female bee diets overlap?
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We found that male and female bee diets overlap significantly less than would be expected at random (Fig. 2), and that the differences in diet composition between male and female bees of several species were of similar magnitude to the differences in diet between species of bee (Fig. 3).

361

362 To what degree are particular flower species disproportionately visited by bees of363 one sex?

364

The sex ratio of flower-visiting bees varied across species of flower (Fig. 2). After controlling for bee species identity (the strongest predictor of sex in our models, Fig. 4), and site, we still found that some flower species received a

disproportionate number of male bee visitors (Figs 4–5). The median odds ratio
 for the main effect of flower species was 3.6 (bootstrapped CI 3.0–4.2) in our

370 summed model, indicating that, typically, the visitor sex ratio differs between two

371 flower species by more than a factor of 3. Furthermore, we observed sex-based

differences in flower use specific to particular bee species: the median odds ratiofor the flower species by bee species interaction in our summed model was

- nearly as large (median=3.1, bootstrapped CI 3.0–3.3) as the main effect of
- 375 flower species. By contrast, sex ratios are not expected to differ between sites
- 376 (median odds ratio for main effect of site=1).
- 377

378 To what extent are differences in floral use driven by preference, rather than 379 phenological differences between male and female bees?

- 380
- 381 The flower species blooming in our system turned over throughout our 11-week

sampling period, with several highly visited species blooming for only one of the 382 383 three months during which we sampled. This turnover, along with potential sex-384 specific bee flight seasons, means that differences in diet between male and 385 female bees could reflect seasonal availability and use, without also indicating 386 preference differences between the sexes. Indeed, phenology predicts bee sex 387 somewhat, with the odds of a flower-visiting bee being male expected to change 388 by a factor of 1.5 (bootstrapped CI 1.1–1.9) between sampling rounds (Fig. 4). 389 Phenological patterns of male vs. female flight seasons vary across bee species; 390 the median odds ratio for the bee species by sampling round interaction is 2.2 391 (bootstrapped CI 2.1–2.3) (Fig. 4). Even after accounting for these effects, 392 however, there remains a strong association between the species of flower a bee 393 visits and its sex (Figs 4-5). The relative effects of each flower species on the 394 sex of its visitors were changed very little by accounting for phenology; Pearson 395 and Spearman correlations between the random effect of flower species in the 396 seasonal model and the same random effect in the simpler summed model were 397 both 0.98. In addition to finding overall preference difference between male and 398 females, we found evidence for bee-species-specific difference in floral 399 preferences between the sexes (median odds ratios in both models for the bee 400 species by flower species interaction > 2.8).

401

402 **Discussion**

403

404 We found strong differences between the flower species preferences of male and 405 female bees. The difference in floral visits between male and female bees of the 406 same species was similar in magnitude to differences between females of 407 different species. The partitioning of the floral community among bee species is a 408 primary focus of pollination ecology and ecological network analysis (Bascompte 409 & Jordano, 2014), but male bees are typically disregarded or lumped together 410 with their female counterparts. Our study suggests this may represent an 411 important oversight. Further, our study provides strong confirmation of the few 412 studies that investigate the foraging behavior of male bees, which found that 413 males play a unique role in plant pollination (Cane, 2002; Cane et al., 2011; 414 Ogilvie & Thomson, 2015; Pascarella, 2010). Our result also implies that male 415 bees contribute substantially to the complexity of plant-pollinator networks in 416 nature, and that network analyses might benefit from separating males and 417 females into different nodes (Bolnick et al., 2011; Zwolak, 2018). 418 419 Phenology, a previously reported mechanism for distinct use of floral resources 420 by male and female bees (Ogilvie & Thomson, 2015; Robertson, 1925), 421 explained some variation in the sex ratio of flower-visiting bees, but was less 422 important than flower species identity over the period of our study. We expected 423 to find an effect of phenology because both the identity of the flower species 424 blooming within sites, and also the sex ratio of foragers within bee species, vary 425 across the season. Males emerge first in most solitary bees; for social species, 426 initial broods usually consist primarily of female workers, then males and

427 reproductive females emerge at the end of the colony cycle (P. G. Willmer &

428 Stone, 2004). Surprisingly, however, phenology only weakly predicted the sex of 429 flower-visiting bees. This is despite the fact that, as predicted by natural history, 430 the sampling round(s) in which males were relatively more prevalent depended 431 on bee species (the bee species by sampling round interaction was much bigger 432 than the sampling round main effect; Fig. 4). This indicates that our evidence for 433 floral preference differences between male and female bees was robust to 434 accounting for seasonal turnover in flower species bloom, bee species flight 435 seasons, and the sex ratios within bee species.

436

437 Whereas female bees collect both nectar and pollen, male bees forage primarily 438 for nectar to fuel flight. Thus, we predicted that male bees would avoid flowers that produce no nectar. Indeed, in both our models, the predicted odds of a bee 439 visiting a nectar-less flower species being male were approximately half that of a 440 441 bee visiting a flower species that produces nectar (Fig. S5). A second biological 442 difference between male and female bees is that adult male bee activities orient 443 around mate seeking (Alcock et al., 1978). These behaviors, such as patrolling 444 routes (Barrows, 1976) or seeking flowers visited by conspecific females (Rossi, 445 Nonacs, & Pitts-Singer, 2010) could generate differences from females via 446 complementarity (males visiting flower species not visited by females), or 447 nestedness (one sex primarily visiting a subset of species visited by the other). 448 We found evidence for both (Fig. S6). Divergent floral preferences between 449 sexes may reflect nutritional needs or mating behavior, or simply biases resulting 450 from previous flower encounters, or visual or olfactory sensitivities that differ 451 between the sexes (Robert, Frasnelli, Collett, & de Ibarra, 2016; Somanathan, 452 Borges, Warrant, & Kelber, 2017; Streinzer, Kelber, Pfabigan, Kleineidam, & 453 Spaethe, 2013).

454

455 While natural and semi-natural habitats are critical landscape elements for many 456 bee species (Senapathi, Goddard, Kunin, & Baldock, 2016), what constitute 457 suitable and/ or limiting resources within these habitats remains less clear (De 458 Palma et al., 2015). Flowers, which provide food for adult and larval bees, are 459 likely among them (Roulston & Goodell, 2011). Whether floral diversity per se 460 tends to benefit individual pollinator taxa remains unclear (Spiesman, Bennett, 461 Isaacs, & Gratton, 2017: Sutter, Jeanneret, Bartual, Bocci, & Albrecht, 2017). 462 However, complementary flower species use between the sexes implies a 463 mechanism by which a bee species could benefit from a diversity of flower 464 choices. In addition, it is currently unknown how the distinct foraging niches of 465 male bees mediate either the robustness of pollinator communities to species 466 loss and environmental perturbations (Brosi & Briggs, 2013; Ramos-Jiliberto, Valdovinos, Moisset de Espanés, & Flores, 2012; Tur et al., 2014), or the 467 468 effectiveness of different habitat ameliorations (Rundlöf, Persson, Smith, & 469 Bommarco, 2014; Rusterholtz & Erhardt, 2000; Williams & Lonsdorf, 2018). This 470 study suggests that both questions warrant further investigation. 471

472 Patterns in bee-flower interaction data can arise from the sampling process itself473 (Blüthgen, 2010; Fründ, McCann, & Williams, 2016). Our analyses control for

474 these patterns. To evaluate diet overlap, we used a dissimilarity index that 475 downweights rare diet items, and implemented a null model that accounts for 476 differences that could arise from sampling and abundance effects. To evaluate 477 preference, we used random effects models that incorporated all (nearly 19,000) 478 observations, and shrank extreme values for rarely observed species-species 479 interactions towards the global mean for each effect. Thus, our estimates for sex-480 specific preferences should be robust to the inevitable under-sampling of rarer 481 taxa. Establishing differences in preference between categories of bees such as 482 males and females, even when resource availability is seasonal and difficult to 483 quantify, is possible using methods such as these, though absolute preference 484 remains elusive.

485

Pollination ecology and pollinator conservation still face the question of how
important sexually dimorphic foraging is. Does it enhance or reduce the stability
of bee populations? Should pollinator restorations explicitly include "male bee"
flowers and "female bee" flowers? Are floral traits under selection to favor female
versus male visitors? While our study does not answer these questions, by
showing that the diets and preferences of male bees commonly differ from those

- 492 of their female conspecifics, we suggest they are worthy of future study.
- 493

494 Author contributions

495

496 MR and RW conceived the ideas and designed field methodology; MR, JD, and
497 RW designed statistical methodology. MR collected the data; MR, RW, and JD
498 analyzed the data; MR and RW led manuscript drafting. All authors contributed
499 critically to the drafts and gave final approval for publication.

500

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502

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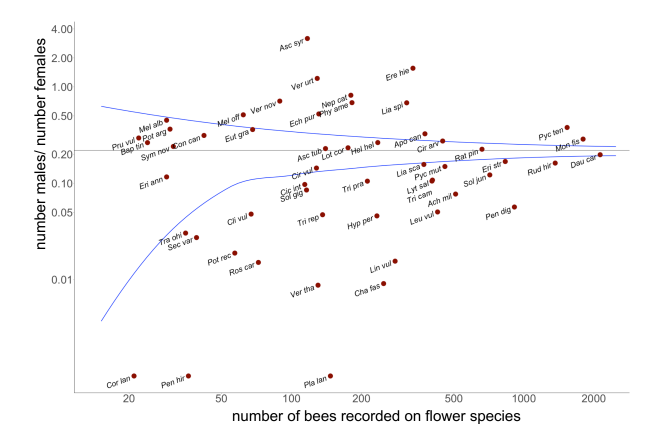
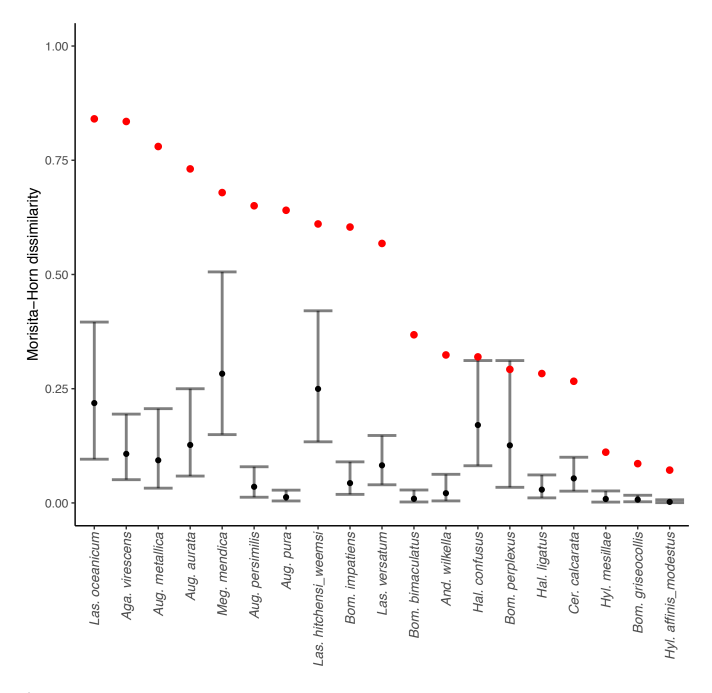
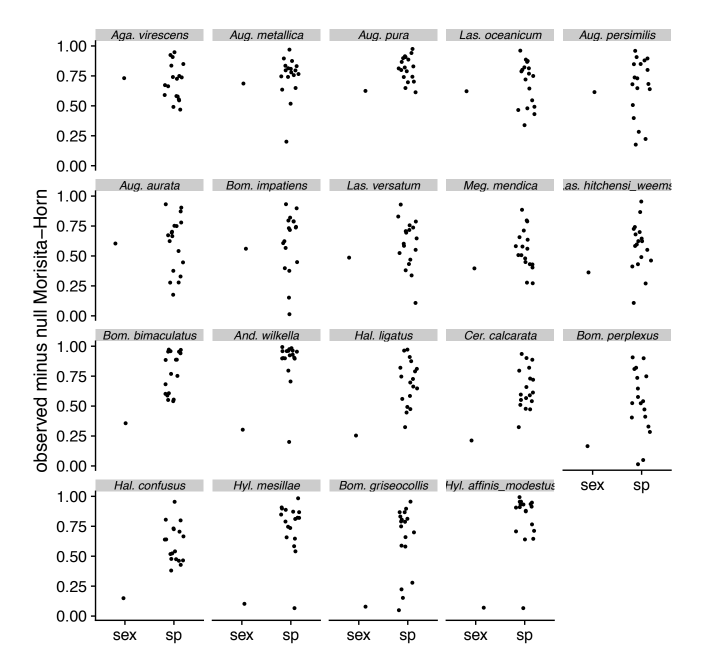


Figure 1. The sex ratio (M:F) of flower-visiting bees varies across flower species. 701 702 Each red point represents a flower species, the first three letters of the Latin 703 genus and species names for the flower species label each point. The x-axis is 704 the number of bees collected from that species, the y-axis is the ratio of male to 705 female bees collected from the flower. Flower species that received >19 visits are plotted (n=54). Blue lines are smooth fits to the 97.5th and 2.5th percentiles of the 706 707 binomial distribution given by the observed ratio of males to females in our 708 overall dataset (M/F=0.22; i.e. M/(M+F)=0.18). This distribution represents our 709 expectation for random variation in sex ratio across flower species, if the sex ratio of flower-visiting bees is independent of flower species identity (male and 710 711 female bees exhibit the same floral preferences), and remains nearly constant 712 across time and space.



- 715 **Figure 2:** Flower visit patterns of male and female bees of the same species
- 716 differed significantly. Red points are observed Morisita-Horn dissimilarities
- 517 between flower communities visited by all male and all female bees of a
- 718 particular species across all sites and sampling rounds. Black points are the
- mean dissimilarity (gray bars, 95% CI) from a permutation-based null model that
- randomly shuffles the sex associated with each visit record, maintaining the total
- number of males, females, and overall combined visits to each floral species.
- 722



723 724 Figure 3. The diets of male and female bees of the same species can be as 725 dissimilar as the diets of females of two different bee species. Dissimilarities in this figure are the observed statistic minus, for each pairwise comparison, the 726 727 mean dissimilarity in the null model. Each panel focuses on a bee species (panel name) and shows: above the label "sex", observed diet dissimilarity between 728 729 male and female bees of the focal species, minus the average null dissimilarity 730 resulting from randomly permuting the sex identity of each visit record; above the 731 label "sp", observed diet dissimilarity between female bees of the focal species 732 and each other bee species, minus the average null dissimilarity resulting from 733 randomly permuting the species identity of each visit record.

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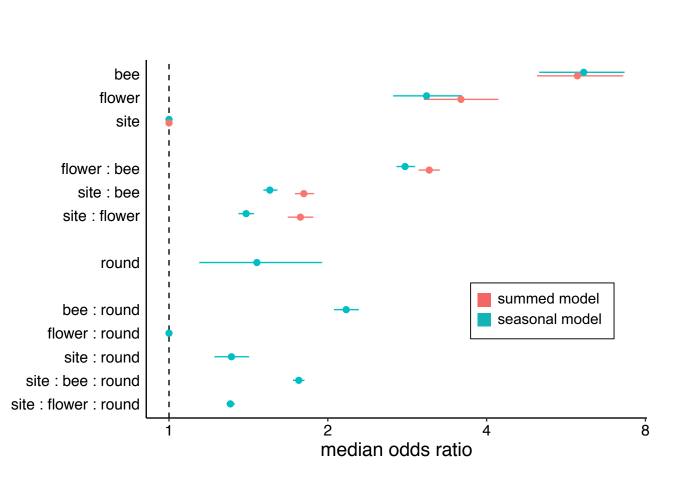


Figure 4. Flower species, along with bee species, predicts the sex of visiting 736 737 bees, which suggests floral preferences differ between male and female bees. 738 Flower species is an important predictor of bee sex even after accounting for 739 phenology (seasonal model). For each term ("bee"= bee species, "flower"=flower 740 species, "round"=sampling round) in each model, the median odds ratio (+/- 95% 741 bootstrapped credible interval) indicates the expected difference in odds that a flower-visiting bee is male between two levels. For example, a median odds ratio 742 743 of 3.7 for the flower species term means the odds of a visitor being male are 744 expected to differ by a factor of 3.7 between two randomly selected species of 745 flower.

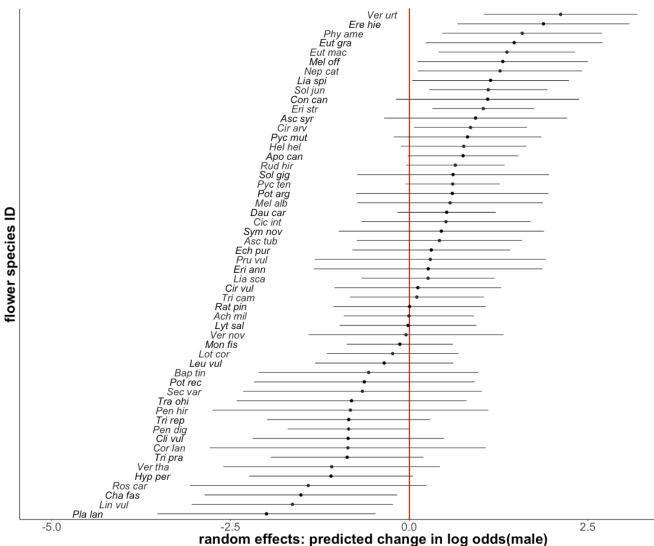
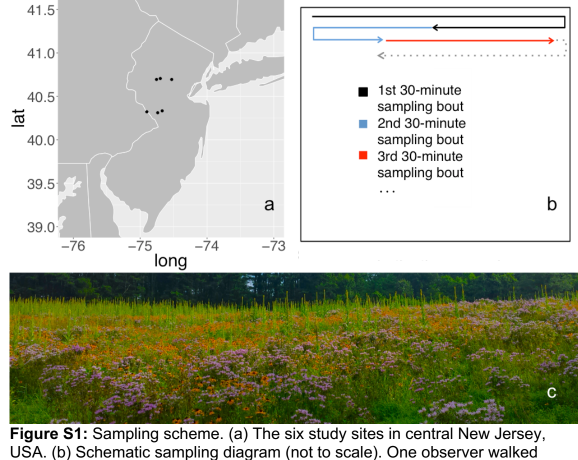


Figure 5. Male bee preferences for and against flower species vary across 747 748 flower species. Each point is the conditional mode of the random effects 749 prediction (the random-effects analog to an estimate), for a flower species that 750 received at least 20 visits, on the logit scale. Zero represents the odds of a visitor being male on a random flower, and -2 or 2 indicates a ~7 fold decrease or 751 752 increase in those odds, given flower species identity. Error bars are the square 753 root of the conditional variances on the conditional mode * 1.96, and can be 754 interpreted as the expected range in which the random effect for a particular 755 flower truly lies, analogous to 1.96 times the standard error of the mean for a 756 fixed effect. 757



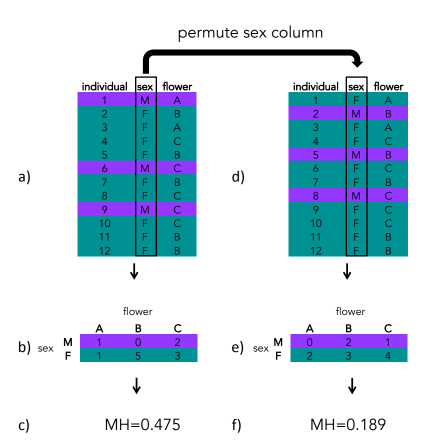
USA. (b) Schematic sampling diagram (not to scale). One observer walke
 parallel 2m transects covering the entire sampling area. Each 30-minute

sampling bout resumed where the previous one left off; observers typically

- covered the entire meadow once over a 3-day sampling round. (c) The
- southwestern-most site in peak bloom.

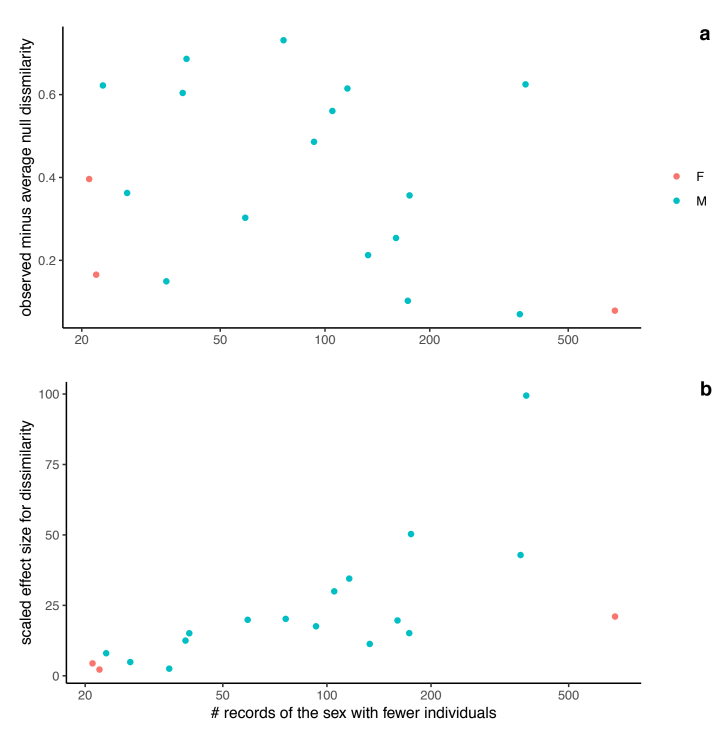
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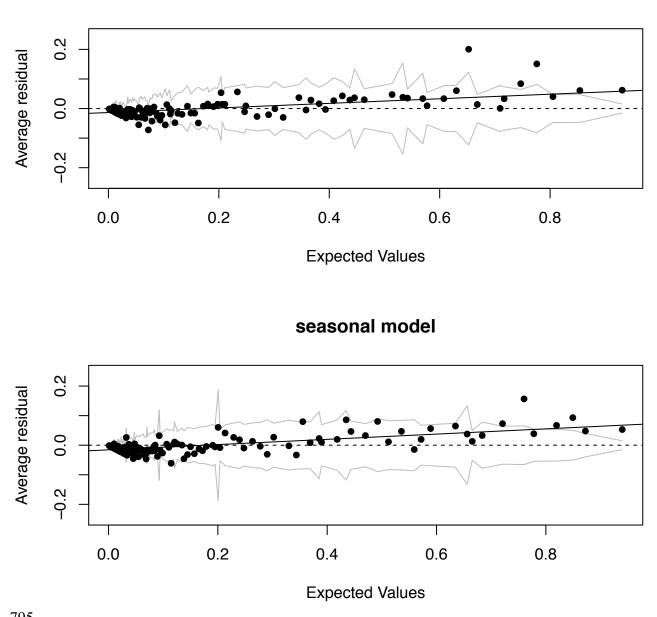
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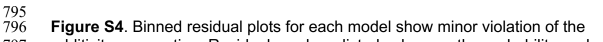
Figure S2. Schematic cartoon of our simulation for the dissimilarity values 768 769 associated with our null hypothesis that diets of male and female bees do not 770 differ. (a) Each collection record for each bee species associates the sex of an 771 individual bee to the flower species from which it was collected. (b) To compute 772 the dissimilarity between males and females, we compare all visits to each flower 773 species from males (purple vector) to all visits to each flower species from 774 females (green vector). (c) The Morisita-Horn index summarizes the differences 775 between the two vectors as a value between 0 (identical) and 1 (maximally 776 dissimilar). (d) For our null model, we shuffle the sex column from our observation table. (e) This produces two null vectors. The row and column sums 777 778 for the matrices in (b) and (c) are identical, but the elements can differ. (f) For our 779 null model, we compute the dissimilarity between the null vectors. We repeated 780 steps d-f 9999 times to generate confidence intervals for the null hypothesis that 781 the sex of a visiting bee is unrelated to the flower species it is collected from. 782 When comparing the flower species visited by different species of bee, we 783 conducted an analysis identical except that rather than comparing two sexes of 784 the same species, we compared two species of the same sex (i.e. exchanging 785 "sex" and "species" throughout figure S1).



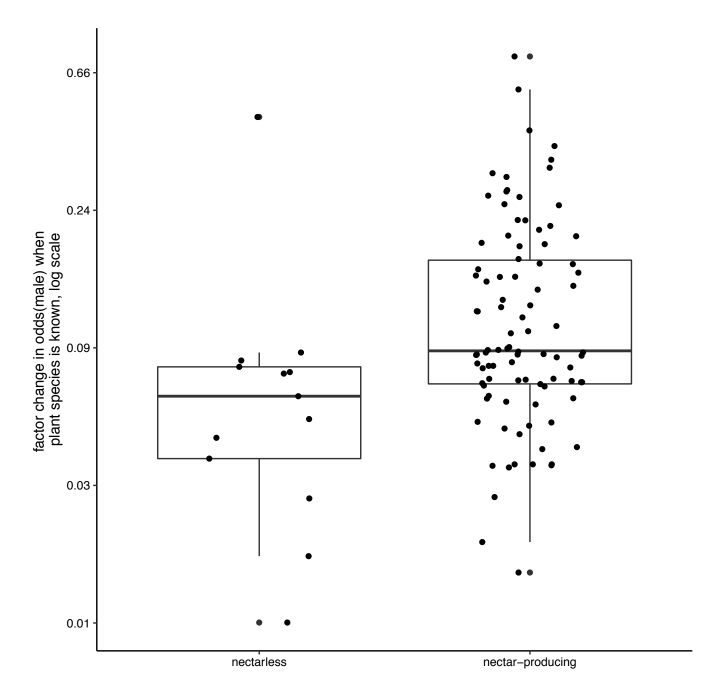
787 Figure S3. Effect size for diet dissimilarity is independent of sample size, while 788 standardized effect is strongly driven by the number of individuals of the sex with 789 the fewest records. a) Observed Morisita-Horn dissimilarity in flower communities 790 visited by male and female bees of a single species, minus average null 791 dissimilarity vs. the number of records for the less frequently observed sex. b) 792 Observed minus null dissimilarity in composition of flowers visited by male and 793 female bees of a single species, scaled by the variation in the null model, versus 794 the number of records for the less frequently observed sex.

summed model





additivity assumption. Residuals and predicted values on the probability scale.

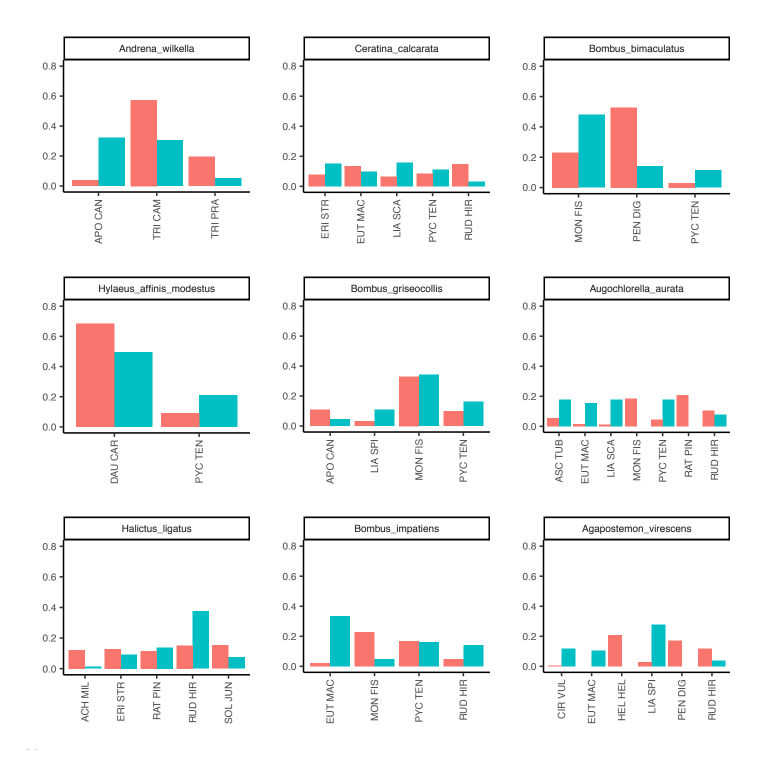


799 **Figure S5.** Seasonal model predictions are consistent with the hypothesis that

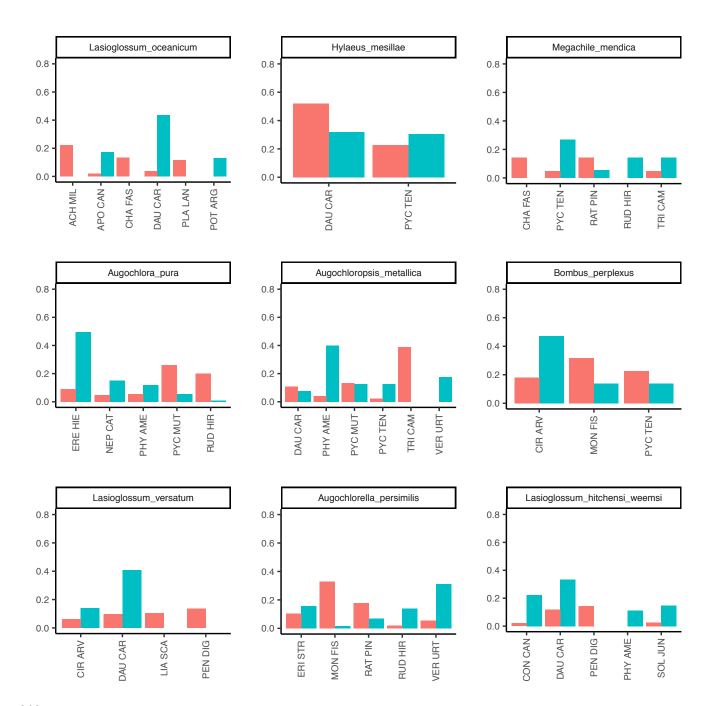
800 male bees avoid flower species that do not produce nectar, relative to females.

801 Each point is the random effect prediction (change in odds that a bee visiting that 802 flower is male) for a flower species.

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Figure S6. Dissimilarity in flower communities visited by male and female bees arise due to complementarity in addition to nestedness patterns. For each bee species, the proportion of male (blue) and female (red) visits to each flower species that received >10% of at least one sex's visits are pictured. Due to omitted flower species, bars may sum to <1.

813 **Table S1.** Model convergence confirmed based on similar parameter estimates across fitting routines. For each model,

814 the estimate for each term is given for each of 6 fitting algorithms in the R package Ime4. Subsequent analyses used

815 parameter estimates in yellow, in both cases tied for the highest estimated likelihood with other very similar fits.

term	model	bobyqa	Nelder Mead	nlminbw	optimx.L- BFGS-B	nloptwrap.NLOPT_ LN_NELDERMEAD	nloptwrap.NLOPT_ LN_BOBYQA
intercept	summed	-2.43	-2.43	-2.43	-2.43	-2.43	-2.43
· ·							
bee species	summed	2.04	2.04	2.04	2.04	2.04	2.04
flower species	summed	1.40	1.40	1.40	1.40	1.40	1.40
site	summed	0.00	0.00	0.00	0.00	0.00	0.00
bee species:flower species	summed	1.21	1.21	1.21	1.21	1.21	1.21
site:bee species	summed	0.62	0.62	0.62	0.62	0.62	0.62
site:flower species	summed	0.61	0.61	0.61	0.61	0.61	0.61
intercept	seasonal	-2.38	-2.45	-2.38	-2.38	-2.45	-2.45
bee species	seasonal	2.09	2.14	2.09	2.09	2.13	2.13
flower species	seasonal	1.25	1.27	1.25	1.25	1.27	1.27
site	seasonal	0.00	0.00	0.00	0.00	0.00	0.00
bee species:flower species	seasonal	1.09	1.10	1.09	1.09	1.10	1.10
site:bee species	seasonal	0.46	0.47	0.46	0.46	0.47	0.47
site:flower species	seasonal	0.35	0.35	0.35	0.35	0.35	0.35
sampling round	seasonal	0.38	0.36	0.38	0.38	0.36	0.36
sampling round:bee species	seasonal	0.83	0.84	0.83	0.83	0.84	0.84
sampling round:flower species	seasonal	0.00	0.00	0.00	0.00	0.00	0.00
sampling round:site	seasonal	0.29	0.29	0.29	0.29	0.29	0.29
sampling round:site:bee species	seasonal	0.60	0.60	0.60	0.60	0.60	0.60
sampling round:site:flower species	seasonal	0.28	0.29	0.28	0.28	0.28	0.28
316	•		-		-		

Table S2. Bee species with number of female and male specimens collected.

This table will be removed from the final submission when data are deposited on Dryad.

family	genus	species	females	males
Andrenidae	Andrena	brevipalpis	1	0
Andrenidae	Andrena	carlini	3	0
Andrenidae	Andrena	commoda	3	0
Andrenidae	Andrena	cressonii 16		0
Andrenidae	Andrena	fragilis	2	0
Andrenidae	Andrena	hippotes	4	0
Andrenidae	Andrena	imitatrix	6	0
Andrenidae	Andrena	krigiana	14	0
Andrenidae	Andrena	nasonii	13	0
Andrenidae	Andrena	nuda	2	0
Andrenidae	Andrena	pruni	6	0
Andrenidae	Andrena	robertsonii	8	0
Andrenidae	Andrena	rudbeckiae	8	11
Andrenidae	Andrena	rugosa	1	0
Andrenidae	Andrena	spiraeana	1	0
Andrenidae	Andrena	vicina		0
Andrenidae	Andrena	wilkella	277	59
Andrenidae	Andrena	wilmattae	2	0
Andrenidae	Calliopsis	andreniformis	4	1
Apidae	Anthophora	abrupta	4	0
Apidae	Anthophora	terminalis	3	2
Apidae	Bombus	auricomus	1	0
Apidae	Bombus	bimaculatus	577	175
Apidae	Bombus	citrinus	0	5
Apidae	Bombus	fervidus	18	0
Apidae	Bombus	griseocollis	681	815
Apidae	Bombus	impatiens	2358	105
Apidae	Bombus	perplexus 22		36
Apidae	Bombus	vagans		2
Apidae	Ceratina	calcarata	1417	133
Apidae	Ceratina	dupla	151	19
Apidae	Ceratina	mikmaqi	130	5
Apidae	Ceratina	strenua	285	13

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Apidae	Melissodes	agilis	0	7
Apidae	Melissodes	bimaculatus	9	1
Apidae	Melissodes	denticulatus	7	73
Apidae	Melissodes	desponsus	1	7
Apidae	Melissodes	subillatus	31	6
Apidae	Melissodes	trinodis	1	7
Apidae	Nomada	articulata	4	0
Apidae	Nomada	bidentate_gr	7	0
Apidae	Nomada	erigeronis	1	0
Apidae	Nomada	lehighensis	1	0
Apidae	Nomada	maculata	2	0
Apidae	Nomada	pygmaea	15	0
Apidae	Ptilothrix	bombiformis	0	1
Apidae	Triepeolus	cressonii	0	1
Apidae	Triepeolus	eliseae	1	0
Apidae	Triepeolus	remigatus	1	0
Apidae	Xylocopa	virginica	137	13
Colletidae	Hylaeus	affinis_modestus	1376	363
Colletidae	Hylaeus	fedorica	1	0
Colletidae	Hylaeus	leptocephalus	1	3
Colletidae	Hylaeus	mesillae	575	173
Halictidae	Agapostemon	sericeus	5	5
Halictidae	Agapostemon	virescens	203	76
Halictidae	Augochlora	pura	1036	377
Halictidae	Augochlorella	aurata	397	39
Halictidae	Augochlorella	persimilis	434	116
Halictidae	Augochloropsis	metallica	121	40
Halictidae	Dufourea	novaeangliae	0	1
Halictidae	Halictus	confusus	174	35
Halictidae	Halictus	ligatus	2432	160
Halictidae	Halictus	parallelus	6	18
Halictidae	Halictus	rubicundus	31	19
Halictidae	Lasioglossum	abanci	6	0
Halictidae	Lasioglossum	admirandum	15	0
Halictidae	Lasioglossum	anomalum	17	0
Halictidae	Lasioglossum	atwoodi	7	1
Halictidae	Lasioglossum	birkmanni	1	0
Halictidae	Lasioglossum	bruneri	6	4
Halictidae	Lasioglossum	callidum	54	0

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Halictidae	Lasioglossum	cattellae	14	4
Halictidae	Lasioglossum	coeruleum	2	0
Halictidae	Lasioglossum	coreopsis	1	0
Halictidae	Lasioglossum	coriaceum	14	0
Halictidae	Lasioglossum	cressonii	16	5
Halictidae	Lasioglossum	ellisiae	0	3
Halictidae	Lasioglossum	ephialtum	1	0
Halictidae	Lasioglossum	foxii	2	2
Halictidae	Lasioglossum	fuscipenne	9	0
Halictidae	Lasioglossum	gotham	74	2
Halictidae	Lasioglossum	hitchensi_weemsi	152	27
Halictidae	Lasioglossum	illinoense	70	7
Halictidae	Lasioglossum	imitatum	462	15
Halictidae	Lasioglossum	leucocomum	2	0
Halictidae	Lasioglossum	leucozonium	2	0
Halictidae	Lasioglossum	nigroviride	2	0
Halictidae	Lasioglossum	oblongum	4	2
Halictidae	Lasioglossum	obscurum	7	1
Halictidae	Lasioglossum	oceanicum	104	23
Halictidae	Lasioglossum	oenotherae	1	0
Halictidae	Lasioglossum	paradmirandum	50	0
Halictidae	Lasioglossum	pectorale	3	0
Halictidae	Lasioglossum	pilosum	2	0
Halictidae	Lasioglossum	platyparium	2	3
Halictidae	Lasioglossum	rozeni	15	11
Halictidae	Lasioglossum	smilacinae	4	0
Halictidae	Lasioglossum	subviridatum	5	1
Halictidae	Lasioglossum	tegulare	31	2
Halictidae	Lasioglossum	trigeminum	44	0
Halictidae	Lasioglossum	truncatum	2	0
Halictidae	Lasioglossum	versatum	681	93
Halictidae	Lasioglossum	viridatum	11	2
Halictidae	Lasioglossum	zephyrum	12	1
Halictidae	Sphecodes	atlantis	0	1
Halictidae	Sphecodes	dichrous	3	5
Halictidae	Sphecodes	heraclei	10	5
Megachilidae	Anthidiellum	notatum	4	1
Megachilidae	Anthidium	manicatum	7	8
Megachilidae	Anthidium	oblongatum	18	19

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Megachilidae	Coelioxys	alternatus	1	2
Megachilidae	Coelioxys	banksi	1	0
Megachilidae	Coelioxys	germanus	0	1
Megachilidae	Coelioxys	hunteri	0	1
Megachilidae	Coelioxys	modestus	0	1
Megachilidae	Coelioxys	obtusiventris	1	0
Megachilidae	Coelioxys	octodentatus	1	1
Megachilidae	Coelioxys	porterae	0	1
Megachilidae	Coelioxys	sayi	2	6
Megachilidae	Heriades	carinatus	31	2
Megachilidae	Heriades	leavitti	1	6
Megachilidae	Heriades	variolosus	10	0
Megachilidae	Hoplitis	pilosifrons	46	1
Megachilidae	Hoplitis	producta	8	0
Megachilidae	Hoplitis	spoliata	2	1
Megachilidae	Lithurgus	chrysurus	0	6
Megachilidae	Megachile	brevis	25	3
Megachilidae	Megachile	campanulae	6	18
Megachilidae	Megachile	exilis	11	29
Megachilidae	Megachile	frugalis	26	6
Megachilidae	Megachile	gemula	4	2
Megachilidae	Megachile	georgica	1	0
Megachilidae	Megachile	inimica	4	0
Megachilidae	Megachile	integra	1	0
Megachilidae	Megachile	melanophaea	0	1
Megachilidae	Megachile	mendica	21	56
Megachilidae	Megachile	montivaga	15	9
Megachilidae	Megachile	petulans	0	2
Megachilidae	Megachile	pugnata	2	3
Megachilidae	Megachile	rotundata	11	8
Megachilidae	Megachile	sculpturalis	17	32
Megachilidae	Megachile	xylocopoides	2	1
Megachilidae	Osmia	albiventris	3	0
Megachilidae	Osmia	atriventris	9	0
Megachilidae	Osmia	bucephala	21	0
Megachilidae	Osmia	distincta	7	0
Megachilidae	Osmia	georgica	5	0
Megachilidae	Osmia	pumila	30	0
Megachilidae	Pseudoanthidium	nanum	0	1

Megachilidae	Stelis	lateralis	1	0
Megachilidae	Stelis	louisae	1	2

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824 **Table S3.** Number of male and female visitors to each plant species, and bias 825 towards attracting male bee visitors. This bias is the random effect prediction 826 from the seasonal model, which indicates the change in log(odds) that a visiting 827 bee is male when the species of flower it visits is given; greater values indicate 828 male bias.

family	genus	species	female visits	male visits	random effect
Alliaceae	Allium	vineale	3	0	-0.224
Apiaceae	Daucus	carota	1778	350	0.526
Apiaceae	Eryngium	yuccifolium	2	0	-0.225
Apiaceae	Sanicula	canadensis	1	0	-0.049
Apocynaceae	Apocynum	cannabinum	283	92	0.754
Asclepidaceae	Asclepias	incarnata	7	0	-0.736
Asclepidaceae	Asclepias	syriaca	28	89	0.929
Asclepidaceae	Asclepias	tuberosa	114	26	0.423
Asteraceae	Achillea	millefolium	472	36	-0.004
Asteraceae	Bidens	trichosperma	1	0	-0.015
Asteraceae	Carduus	nutans	1	0	-0.366
Asteraceae	Centuarea	stoebe	321	50	0.265
Asteraceae	Cichorium	intybus	104	10	0.516
Asteraceae	Cirsium	arvense	351	96	0.858
Asteraceae	Cirsium	discolor	5	2	0.349
Asteraceae	Cirsium	vulgare	112	16	0.121
Asteraceae	Conyza	canadensis	32	10	1.097
Asteraceae	Coreopsis	lanceolata	21	0	-0.857
Asteraceae	Coreopsis	tinctoria	1	0	-0.069
Asteraceae	Crepis	capillaris	6	0	-0.236
Asteraceae	Doellingeria	umbellata	1	0	-0.052
Asteraceae	Echinacea	purpurea	86	45	0.309
Asteraceae	Erechtites	hieraciifolius	130	203	1.878
Asteraceae	Erigeron	annuus	26	3	0.266
Asteraceae	Erigeron	strigosus	712	119	1.037
Asteraceae	Euthamia	graminifolia	50	18	1.467
Asteraceae	Eutrochium	maculatum	461	166	1.368
Asteraceae	Gaillardia	aristata	4	0	-0.274
Asteraceae	Helianthus	angustifolius	11	0	-0.411

Actorecco	Holicathus	otrumosous		4	0 5 4 7
Asteraceae	Helianthus	strumosus	1	1	0.517
Asteraceae	Heliopsis	helianthoides	186	49	0.763
Asteraceae	Hieracium	pilosella	2	0	-0.148
Asteraceae	Krigia	biflora	19	0	-0.542
Asteraceae	Lactuca	serriola	4	0	-0.257
Asteraceae	Leucanthemum	vulgare	405	20	-0.349
Asteraceae	Liatris	spicata	186	127	1.137
Asteraceae	Ratibida	pinnata	539	121	0.005
Asteraceae	Rudbeckia	hirta	1174	189	0.645
Asteraceae	Solidago	altissima	8	0	-0.586
Asteraceae	Solidago	canadensis	8	2	0.816
Asteraceae	Solidago	gigantea	107	9	0.613
Asteraceae	Solidago	juncea	636	77	1.106
Asteraceae	Solidago	rugosa	1	0	-0.015
Asteraceae	Symphyotrichum	novae-angliae	25	6	0.451
Asteraceae	Vernonia	noveboracensis	52	37	-0.045
Brassicaceae	Barbarea	vulgaris	3	0	-0.114
Campanulaceae	Lobelia	inflata	12	3	1.147
Campanulaceae	Lobelia	siphilitica	2	0	-0.034
Caryophyllaceae	Dianthus	armeria	3	0	-0.280
Commelinaceae	Tradescantia	ohiensis	34	1	-0.805
Convulvulaceae	Calystegia	silvatica	5	0	-0.538
Cornaceae	Swida	amomum	4	0	-0.250
Cornaceae	Swida	racemosa	12	1	0.546
Fabaceae	Baptisia	tinctoria	19	5	-0.566
Fabaceae	Chamaecrista	fasciculata	246	2	-1.514
Fabaceae	Desmodium	paniculatum	6	1	-0.138
Fabaceae	Lotus	corniculatus	142	33	-0.231
Fabaceae	Melilotus	albus	20	9	0.571
Fabaceae	Melilotus	officinalis	41	21	1.309
Fabaceae	Securigera	varia	38	1	-0.654
Fabaceae	Senna	hebecarpa	5	5	1.678
Fabaceae	Trifolium	aureum	1	0	-0.033
Fabaceae	Trifolium	campestre	365	39	0.106
Fabaceae	Trifolium	hybridum	11	7	0.739
Fabaceae	Trifolium	pratense	192	20	-0.869
Fabaceae	Trifolium	repens	130	6	-0.847
Fabaceae	Vicia	tetrasperma	1	0	-0.048
Gentianaceae	Sabatia	angularis	1	0	-0.034

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Hypericaceae	Hypericum	perforatum	223	10	-1.095
Hypericaceae	Hypericum	punctatum	1	0	-0.176
Iridaceae	Sisyrinchium	angustifolium	9	0	-0.351
Lamiaceae	Clinopodium	vulgare	64	3	-0.854
Lamiaceae	Monarda	fistulosa	1397	400	-0.131
Lamiaceae	Monarda	punctata	0	1	1.242
Lamiaceae	Nepeta	cataria	99	81	1.270
Lamiaceae	Prunella	vulgaris	17	5	0.296
Lamiaceae	Pycnanthemum	muticum	398	59	0.811
Lamiaceae	Pycnanthemum	tenuifolium	1113	421	0.609
Lamiaceae	Pycnanthemum	verticillatum	5	8	0.886
Lamiaceae	Teucrium	canadense	3	0	-0.104
Loniceraceae	Lonicera	japonica	1	0	-0.131
Lythraceae	Lythrum	salicaria	364	38	-0.016
Onagraceae	Oenothera	biennis	2	0	-0.143
Onagraceae	Oenothera	fruticosa	2	1	0.221
Oxalidaceae	Oxalis	stricta	8	0	-0.249
Phytolaccaceae	Phytolacca	americana	108	74	1.580
Plantaginaceae	Plantago	lanceolata	147	0	-1.997
Poaceae	Glyceria	grandis	1	0	-0.187
Poaceae	Sorghastrum	nutans	1	0	-0.092
Polygonaceae	Fallopia	convolvulus	3	4	1.044
Polygonaceae	Persicaria	pensylvanica	4	0	-0.391
Polygonaceae	Persicaria	setacea	3	1	0.158
Ranunculaceae	Ranunculus	hispidus	1	0	-0.057
Rosaceae	Drymocallis	arguta	22	8	0.604
Rosaceae	Potentilla	recta	56	1	-0.628
Rosaceae	Rosa	carolina	71	1	-1.412
Rosaceae	Rosa	multiflora	5	0	-0.264
Rosaceae	Rubus	flagellaris	1	0	-0.027
Rosaceae	Rubus	pensilvanicus	7	0	-0.241
Rubiaceae	Galium	mollugo	4	0	-0.369
Scrophulariaceae	Linaria	vulgaris	275	4	-1.633
Scrophulariaceae	Penstemon	digitalis	862	48	-0.847
Scrophulariaceae	Penstemon	hirsutus	36	0	-0.822
Scrophulariaceae	Verbascum	blattaria	15	0	-0.721
Scrophulariaceae	Verbascum	thapsus	129	1	-1.085
Solanaceae	Solanum	carolinense	14	0	-0.518
Verbenaceae	Verbena	hastata	8	3	0.927

Verbenaceae	Verbena	simplex	11	3	0.482
Verbenaceae	Verbena	urticifolia	58	71	2.118