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# Floral abundance and resource quality influence pollinator choice

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22 Abstract

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Pollinator declines caused by forage habitat loss threaten insect pollination services. Pollinating insects
 depend on adequate floral resources, and their ability to track these resources. Variability of these resources
 and the effect on insect foraging choice is poorly understood.

27 2. We record patterns of visitation to six wildflower species and test the hypotheses that: pollinators
 28 preferentially visit the most rewarding flowers; nectar diurnal variations affect foraging preferences;
 29 pollinators respond most strongly to nectar rewards.

30 3. Nectar volume and sugar concentration were negatively correlated within plant species over time of day
31 where greater concentration and lower volume was evident in the afternoon, but this did not correspond to
32 pollinator visitation. Both floral abundance and nectar quality (total sugar per inflorescence) positively affect
33 insect visitation. For some foragers the positive effects of high quality rewards were only evident when floral
34 abundance was high (>50 inflorescences per patch), perhaps reflecting the low probability of pollinators
35 detecting scarce rewards. Pollen quality (total protein per inflorescence) was negatively related to visitation
36 of *Apis mellifera* and *Bombus pascuorum*.

Fewer pollinators visiting flowers of higher pollen quality could reflect plant allocation trade-offs or the
presence of secondary metabolites in pollen, meaning pollen foraging is likely affected by factors other than
protein concentration. Nectar rather than pollen appeared to be the main driver of floral choice by insects in
this system.

- 41 5. Conservation schemes for bees in farmland or gardens might benefit from ensuring that rewarding plant
  42 species are present at high density and/or are aggregated in space.
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52 Over the past 80 years, local and UK-wide changes in farming practice and agricultural intensification 53 have led to a reduction in diversity of crop and non-crop plants (Robinson & Sutherland, 2002; Öckinger et al., 54 2007; Geiger et al., 2010). This includes the loss of meadow plants (Goulson et al., 2005), arable weeds and 55 hedgerows (Hanley & Wilkins, 2015), which provide valuable foraging resources for flower visiting insects. This 56 habitat degradation has been identified as the primary reason for population declines in insects reliant on flowers 57 to provide nectar and/or pollen as a main food source, including adult butterflies (Brereton et al., 2011) and adult 58 and offspring honey bees (Potts et al., 2010), bumblebees (Goulson et al., 2008) and solitary bees (Ollerton et al., 59 2014). Nesting and hibernation resources aside, it is imperative that bees and other pollinating insects are able to 60 forage effectively for nutritional resources in increasingly fragmented landscapes in order to survive and 61 reproduce, particularly as they are facing other pressures such as diseases and pathogens (Cox-Foster *et al.*, 2007), 62 global environmental change (Tylianakis et al., 2008) and pesticide use (Goulson et al., 2015), all of which impact 63 on their survival. Population decline in flower visiting insects could jeopardise the pollination services provided 64 to entomophilous crops (Kremen et al., 2002; Klein et al., 2007).

65 Nectar provides sugars (mainly sucrose, glucose and fructose), which energise pollinators to continue 66 foraging (on nectar or other sources of nutrients), undertake nesting activities, find mates, and provide for 67 offspring. It also contains ions, water and small amounts of amino acids, which may contribute to nutrition (Kim 68 & Smith, 2000). Pollen comprises proteins, carbohydrates, lipids, vitamins and minerals (Roulston & Cane, 2000; 69 Nicolson, 2011). Although many flower visiting insects consume pollen for sustenance (e.g. beetles, flies), bees 70 (Hymenoptera: Apoidea) also collect pollen to feed larvae, and many simultaneously collect both nectar and pollen 71 from flowers, depending on what requirements they have and what flower species they are feeding on (Heinrich, 72 1979a; Heinrich, 1979b; Goulson et al., 2005).

Survival and reproductive success of pollinating insects is dependent on them successfully gathering adequate protein and sugar to provide for their energy and nutritional needs. For example, many Lepidoptera, Diptera and Hymenoptera rely on energy gained from nectar to undertake mating flights, for dispersal and/or migration, and to find suitable places to lay their eggs. Butterflies tend to exhibit lower fecundity when nectar limited (Boggs & Ross, 1993), and the hoverfly *Episyrphus balteatus* has greater longevity when fed a sugar and protein rich diet (Pinheiro *et al.*, 2015). Bees require both pollen and nectar to feed to their offspring. When fed on protein rich diets, bumblebee colonies are more reproductively successful (Génissel *et al.*, 2002; Kitaoka &
Nieh, 2009), and the solitary bee *Lasioglossum zephyrum* (Roulston & Cane, 2002) and honey bees (Basualdo *et al.*, 2012) produce larger adults. Larger bees generate and retain heat faster leading to earlier and more frequent
forage flights (Stone, 1993), they have greater diapause survival (Strohm and Linsenmair, 1997) and are better
able to cope with adverse conditions such as parasitism and disease (Alaux *et al.*, 2010; Di Pasquale *et al.*, 2013).

In order to forage effectively, pollinators use olfactory cues to enable detection of non-depleted nectar resources (Goulson *et al.*, 1998a; Howell & Alarcón, 2007) and greater nectar volume and sugar content (Pyke, 1978; Heinrich, 1979a; Wolff, 2006). Likewise, insect foragers tend to select pollen with greater protein and essential amino acid content (Levin & Bohart, 1955; Schmidt, 1982; Robertson *et al.*, 1999; Cook *et al.*, 2003; Arenas & Farina, 2012), and obligate insect pollinated plant species produce pollen that is richer in protein and amino acids compared to facultative species (Hanley *et al.*, 2008, but see Roulston *et al.*, 2000).

90 Few studies have investigated how eusocial or solitary bees integrate information on nectar and pollen 91 quantity and quality simultaneously, and those that have, indicate that nectar is the primary factor influencing 92 foraging preference for bees, and pollen a secondary consideration (Konzmann & Lunau, 2014; but see Somme 93 et al., 2015). In addition, studies focusing on flower selection by pollinators tend to look only at individual species 94 of bee and specific pollinators for individual plant species (e.g. Robertson et al., 1999), or solely bees as a group 95 (e.g. Heinrich, 1979b; Pernal & Currie, 2001). Floral choice tests are frequently undertaken in controlled 96 conditions (e.g. Konzmann & Lunau, 2014), but the limited number of field studies investigating the influence of 97 rewards on the foraging decision of pollinators makes it difficult for the conclusions gained by laboratory studies 98 to be applied, especially as there are external factors in the natural environment likely to influence the results, 99 such as the spatial distribution and reward phenology of flowers. For example, Hanley & Wilkins (2015) describe 100 greater food plant abundance in roadside, compared with field facing hedgerows, and noted a corresponding 101 increase in bumblebee abundance. In addition, several studies indicate that some flower species offer less nectar 102 in the middle of the day and afternoon compared to the morning and evening (Mačukanović-Jocić et al., 2004; 103 Silva et al., 2004; Mačukanović-Jocić & Đurđević, 2005), providing a further challenge to efficient foraging by 104 pollinators since the relative rewards provided by different flower species may alter hour by hour through the day.

105 In this study, we examine the foraging choices made by all flower visiting insects in relation to the 106 relative nectar and pollen quantity and quality of six common plant species under natural conditions. We tested 107 the hypothesis that nectar quantity or quality shows diurnal variation and that nectar or pollen quantity or quality, 108 or a combination of reward metrics, predicts insect visits. We record how nectar volume and sugar concentration 109 changes through the day, and also the pollen weight and protein concentration for each test plant species, and 110 relate these values to insect flower choices through the day. Specifically, this study's objectives were to; (i) record 111 nectar and pollen quantity and quality estimates for six test plant species, (ii) investigate the diurnal variation in 112 nectar quantity and quality for the test plant species, and (iii) assess how nectar and pollen quantity and quality 113 influence foraging choices by flower visiting insects.

- 114
- 115 Methods
- 116
- **117** *Test Plant Species*

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119 Six species of flowering plant that are common in southern UK were selected for nectar and pollen 120 quantity and quality estimation, and insect visitation surveys in the field. These were Lamium album L. and 121 Glechoma hederacea L. (Lamiaceae), Crataegus monogyna Jacq. and Rubus fruticosus L. agg. (Rosaceae), 122 Symphytum officinale L. (Boraginaceae) and Ranunculus repens L. (Ranunculaceae). These spring or early 123 summer flowering plants considered to be beneficial foraging plants for insects in previous studies (Lack, 1982; 124 Goulson et al., 1998b; Kipling & Warren, 2013) (further species details are listed in Table S1 in the Electronic 125 Supplementary Material). Previously, R. fruticosus and S. officinale were included in studies of pollen protein 126 quality and its effects on bee foraging preferences (Hanley et al., 2008), but to our knowledge the other test plant 127 species have not been studied in this context.

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129 Site Selection

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Sites were chosen to be included in this study if they contained at least three of the test plant species within 50 m of each other, were on chalk soil, were easy to access and were subject to intermediate levels of disturbance (e.g. mowed once a year, or grazed intermittently by sheep). Sites could be divided into two groups, road verges and semi-improved grassland. In terms of management, road verge sites were mowed once a year and semi-improved grassland were grazed several times throughout the year. However, none of the survey sites were either mown or grazed whilst this study was taking place. Once sites were identified, nectar and pollen samples were collected from test plant species and visitation sampling was conducted.

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#### 139 Nectar Volume and Sugar Concentration

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141 Nectar sampling was undertaken on 30 inflorescences of each test plant species, collected from at least 142 three different survey sites. Nectar was sampled from inflorescences of plant species that had recently come in to 143 bloom and were easily accessible. For each inflorescence nectar production was estimated from morning, 144 afternoon and evening periods for each plant species tested. For each species then flowers were emptied of nectar 145 and bagged by 09:00 (GMT) using a fine-mesh cotton fabric and masking tape. At 15:00, 21:00 and 09:00 the following morning flowers were emptied of nectar using 5 µl micropipettes (BLAUBRAND® intraMARK, 146 147 Wertheim, Germany). The nectar from each inflorescence was then expelled onto a field refractometer 148 (Bellingham and Stanley Ltd, Basingstoke, Hants, UK) to measure the proportion of sugar in each sample (Bolten 149 et al., 1979). This produced both mean nectar volume and mean sugar concentration for each species in each time 150 period, and over a 24 hour period when totalled.

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#### 152 Pollen Weight and Protein Concentration

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Pollen was collected from 10 inflorescences from at least three different sites for each plant species. Standing crop of pollen was taken just once for each replicate so variation in pollen quantity or quality was not measured over time of day. Flowers were collected in the field before anthers had dehisced, and placed in water in an unheated and well ventilated laboratory until anthers opened. Pollen was collected systematically until anthers were empty. It is conceivable that placing flowers in water affects pollen quality, but all plant species were treated in the same way. Pollen was either stored in a – 20 °C freezer for drying at a later date, or immediately dried in an oven for 24 hours at 40 °C. After drying, pollen was weighed to measure total pollen weight per 161 inflorescence. Protein extraction and detection was undertaken using the Bradford assay (Bradford, 1976). The 162 assay binds protein to Coomassie Brilliant Blue G-250 dye (4.7% [weight: volume] ethanol, and 8.5% [weight: 163 volume] phosphoric acid dissolved in water). Light absorbance is then measured against known protein 164 standards. From each of the 10 inflorescence pollen from each species, 1 mg pollen were dusted with aluminium 165 powder, wetted with 20 µl 0.1 mol/L NaOH and ground with a micro-pestle. Ground pollen was reanimated 166 with 480 µl 0.1 mol/L NaOH and placed in a refrigerator for 24 hours before analysis, but used within 1 week. 167 Prior to absorbance measurement samples were placed in boiling water for 5 minutes and centrifuged for 5 168 minutes. Then 10 µl of supernatant was slowly vortexed with 300 µl of dye reagent. This was repeated in 169 triplicate for each sample and left to incubate at room temperature for 15 minutes.

Protein standards were made up each time samples were run, using pre-mixed concentrations of Bovine
Serum Albimum (BSA) from the BIO-RAD (Hertfordshire, UK) Quick-Start<sup>TM</sup> Bradford Protein Assay kit. Once
samples and standards were created, they were measured for absorbance within an hour of mixing at 595 nm using
a Thermo Scientific (Paisley, UK) Nanodrop 2000 UV-Vis Spectrophotometer. This produced the standing crop
of both mean pollen weight and a crude mean protein concentration for each test plant species.

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176 Visitation Surveys

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178 Visitation surveys took place at seven sites across Sussex, UK between May and June 2014 (for site 179 details see Table S2). Surveys were undertaken in the morning (08:00-10:00), afternoon (13:00-15:00) and 180 evening (19:00—21:00). Each survey consisted of a 10 minute standing observation of a 4  $m^2$  area of each test 181 plant species at each site. The number and species of all visiting insects to that plant species were recorded. All 182 observed flower visiting insects appeared to collect nectar and, in most cases, pollen from plant species, therefore 183 we did not attempt to distinguish which resources insects were collecting. The number of inflorescences of the 184 plant species, ambient temperature, and wind speed were recorded for each survey. Surveys were only conducted when air temperature was >  $14^{\circ}$ C and average wind speeds was < 20 mph. 185

186 Common species of bumblebee (genus *Bombus*) and European honeybees (*Apis mellifera* L.) were
187 identified on the wing. Due to the unreliability of morphological characters for separating *Bombus terrestris* L.
188 and *B. lucorum* agg. workers in the field, these two species were grouped. Although others have recorded flower

visiting bumblebees into groups based on colour type (Haughton *et al.*, 2003), in this study the bumblebee species observed were easily separated to species in the field. However, faded or suspected cleptoparasitic bees (*Psithyrus* spp.) were caught and checked using a hand magnifying glass. Other visiting insects too small to be identified on the wing were collected and identified to species or genus. Although unlikely given the short period of each survey, the possibility of double counting insects was the same for all surveys due to equal sampling time and therefore assumed to be unbiased across the study.

195

196 Statistical Analysis

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Means tests of variance were used to assess differences in floral resources available from the plant species, as well as the number of insect visits, between periods of the day. Based on whether resource and insect visit metrics showed homogeneity of variance between species or periods of the day using Levene's test, either ANOVA (parametric) or Kruskal-Wallis (non-parametric) tests were applied to assess variance for all groups before either Tukey HSD tests or pairwise Wilcox tests were applied to assess differences between groups, respectively.

204 Generalized Linear Mixed Models (GLMMs) were used to test the effects of resource metrics on insect 205 visitation (i.e. counts per 10 minute survey). Prior to applying models, proposed explanatory variables were 206 checked for multicollinearity using Variance Inflation Factors (VIFs), if variables had VIFs greater than 3 or 207 correlation coefficients more than 0.6 with other variables, they were excluded from models (Zuur et al., 2010). 208 These analyses indicated nectar volume and pollen weight were correlated with sugar and protein concentration, 209 respectively. To enable modelling of resource quality and quantity without violating model assumptions, nectar volume and pollen weight were multiplied by sugar and protein concentration to create total sugar and protein in 210 211 milligrams per inflorescence as measurements of quality in the plant species. To account for the change in density 212 of the solution based on the amount of sugar recorded, percentage sugar was multiplied by the mass density (g/cm3) 213 of sugar at that concentration. These measurements also allowed the interpretation of the nutritional gain available 214 to foraging insect pollinators, and is referred to as total sugar and total protein from here on.

GLMMs were initially run with the environmental variables temperature and wind speed included asexplanatory variables, however these had no effect on visitation or the outcome of the models so were removed.

217 We modelled five visitation response variables (including: all insects; all bumblebees; the three most recorded 218 species, B. pascuorum Scopoli, B. pratorum L., and A. mellifera) in response to the number of inflorescences (log 219 transformed) for each test plant species surveyed in each observation area  $(4 \text{ m}^2)$  (floral abundance), and total 220 sugar and protein. We did not analyse data on other groups of flower visiting insects alone as they were recorded 221 in such small numbers that data analysis would have been unreliable. However, they were included in the data 222 analysis as part of 'all insects' in statistical models. We included interactions between floral abundance and total 223 sugar and protein explanatory variables. Sample site was fitted as a random factor because observations were 224 nested within sample sites and contained different combinations of test plant species. Visitation rates were count 225 data, therefore models were applied with Poisson errors (O'Hara & Kotze, 2010). Model residuals were assessed 226 for normality and heteroscedasticity. Model simplification was carried out using likelihood ratio tests, and 227 sequentially deleting terms that did not significantly decrease model deviance, beginning with higher order 228 interactions.

To further investigate how particular plant species affect insect visitation, linear mixed effects (LME) models were used to test the effects of floral abundance and species on the same response variables as the GLMMs, with site as a random factor. Models were tested for significance using likelihood ratio tests, with and without species as an explanatory variable.

Already published protein estimates for each plant species were compared to confirm the extraction results and accuracy of our values. We found that only *Lamium album* (4.48%) differed from the literature for *Lamium* sp. (22.8% in Roulston *et al.*, 2000), so to assess whether this estimate affected our findings, we substituted it in the two models that included total protein (for *A. mellifera* and *B. pascuorum* visitation).

Statistical analysis was conducted using R v3.1.2 (R Core Team, 2014) within RStudio (RStudio, 2012)
using packages 'lme4' (Bates *et al.*, 2015) and 'usdm' (Naimi, 2013), and plots were generated using 'ggplot2'
(Wickham, 2009).

240

241 Results

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243 *Nectar and Pollen Quantity and Quality* 

A large amount of between-inflorescence variation was found in nectar and pollen resources for each test plant species (Table 1). Nevertheless, nectar volume (Kruskal-Wallis H = 211.64, d.f. = 5, P < 0.001), nectar sugar concentration (H = 112.61, d.f. = 5, P < 0.001) and pollen protein concentration (ANOVA  $F_{5,54}$  = 20.6, P < 0.001) varied significantly between plant species, with only pollen weight ( $F_{5,54}$  = 1.25, P = 0.3) showing no significant differences.

Over 24 hours, *Rubus fruticosus* produced the greatest volume of nectar, which contained the lowest sugar concentration (Table 1). In contrast, *Ranunculus repens* produced the least nectar, whilst *Crataegus monogyna* offered the highest sugar concentration. *Lamium album* pollen offered the greatest mass per inflorescence but had the lowest protein concentration of all species. *C. monogyna* produced the least pollen mass whilst *G. hederacea* offered the highest protein concentration. We found no correlation between nectar and pollen quantity (Pearson R = -0.46, P = 0.26) or quality (Pearson R = -0.33, P = 0.52).

Nectar volume and pollen weight were multiplied by sugar and protein concentration, respectively, to create total sugar and protein in milligrams per inflorescence (Table 1). Over 24 hours, *Symphytum officinale* produced the greatest total sugar per inflorescence whilst *R. repens* produced the least. *C. monogyna* contained the most total protein per inflorescence and *S. officinale* the least (Table 1).

260

#### 261 Diurnal Variation in Nectar Quantity and Quality

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Mean nectar volume and sugar concentration averaged between plant species (regardless of time period) were not significantly correlated (Pearson R = -0.49, P = 0.32, Fig. 1A). When mean for each species at each time period was taken, there was a significant negative correlation between nectar volume and sugar concentration (Pearson R = -0.52, P = 0.02, Fig. 1B).

Volumes of nectar produced differed markedly between species, and between morning, afternoon and evening periods. *R. fruticosus* produced the greatest volume of nectar of all species in the evening and morning, yet recorded the lowest sugar concentration of all species in these periods (Table 1). The lowest volume of nectar was detected in *R. repens* in the evening, and the greatest sugar concentration was found in *C. monogyna* in the afternoon. When averaged across species, mean nectar volume did not differ significantly between periods of the day ( $F_{2, 15} = 1.18$ , P = 0.35). However, mean nectar sugar concentration across species increased significantly in the afternoon compared to the morning ( $F_{2, 15} = 4.01$ , P = 0.04) (Fig. 2).

Once sugar and protein concentrations were calculated between time periods, total sugar was significantly lower in the afternoon period compared to the morning or evening for *R. fruticosus*, *G. hederacea*, *L. album* and *R. repens* (Fig. 3). The exceptions were *C. monogyna* which produced comparable amounts of total sugar in each time period sampled, and *S. officinale* which had marginally more total sugar in the afternoon compared to morning and evening periods.

279

**280** *Foraging choices of flower visiting insects* 

281

282 In total, we made 112 ten-minute observations (38, 39 and 35 in morning, afternoon and evening periods, 283 respectively), between May and June 2014 at seven sites in Sussex, during which 574 insects were recorded 284 visiting test plant species. Proportionately, of all insect visits, 93% were by bees (Hymenoptera: Apidae), 6% by 285 Diptera (< 1% of which were hoverflies [Syrphidae]), < 1% by Lepidoptera and < 1% by Vespidae (Table S3). 286 Of all flower visiting insects, the three most numerous were the bee species Bombus pascuorum, B. pratorum, and 287 Apis mellifera, which represented 33%, 30% and 16% of all insect visits, respectively (Table S3). The majority 288 of insects were recorded visiting R. fruticosus (47%), S. officinale (24%) and L. album (19%) (Table S3). LME 289 models showed significant species-specific preferences for the most commonly recorded flower visiting insects 290 (Table S4). B. pascuorum mainly visited L. album and S. officinale (49% and 31% of this species' visits, 291 respectively), B. pratorum preferred R. fruticosus and S. officinale (57% and 36%), and A. mellifera mainly visited 292 R. fruticosus (87%) (Fig. S1 & Table S3).

Fewer insects were recorded in the evening (122) than the morning (221) and afternoon periods (231), and differences in abundance were also evident within and between recorded insect and test plant species (Table S3). However, these differences were not significant when tested between surveys for all insect abundance ( $F_{2, 109}$ = 2.59, P = 0.07). No significant difference was found between any response variable between different periods of the day when all test plant species were grouped or tested separately, except *G. hederacea* which had significantly more insects visiting during the afternoon compared to morning and evening periods ( $F_{2, 24}$  = 9.03, P = 0.001; Fig. 4). This however does not correspond with the timing of maximum sugar availability, which
appeared highest in the morning or evening for most species (Fig. 3). As standing crop of pollen was taken just
once for each replicate, we could not quantify variation in pollen quantity or quality over the day.

302 Overall, flower abundance had a positive effect on the total number of bumblebee (bees within the genera 303 Bombus) and all insect visitation (Table 2). Total sugar predicted insect visitation, with this relationship being 304 positive for all response categories apart from B. pascuorum which was negative (Table 2). There were also a 305 number of significant higher order interactions between floral abundance and total sugar and protein. For Apis 306 mellifera, all bumblebees and all insects, the relationship between insect visitation and total sugar was weak or absent at low floral abundance  $(10 - 50 \text{ inflorescences per 4 } \text{m}^2)$  but positive at high floral abundance  $(51 - 1000 \text{ m}^2)$ 307 inflorescences per 4 m<sup>2</sup>) (Fig. 5). Total protein had a significant negative effect on A. mellifera visitation (Table 308 309 2). In addition, a negative interaction between total protein and floral abundance was found for B. pascuorum 310 (Table 2), with the relationship weak or absent at low floral abundance  $(10 - 50 \text{ inflorescences per 4 m}^2)$  but 311 negative at high floral abundance  $(51 - 1000 \text{ inflorescences per 4 } m^2)$ . However, the three plant species most 312 visited by insects (R. fruticosus, S. officinale and L. album) were also recorded as producing the lowest total pollen 313 protein. When the protein value for L. album was replaced with the higher value reported in Roulston et al. (2000) 314 there was no longer a negative significant effect of total protein on B. pascuorum visitation, and we found a 315 positive significant effect of total sugar (Table S5).

316

#### 317 Discussion

318

319 Many insects forage on flowering plants to gain key nutritional resources, largely nectar and pollen. What 320 factors determine when plants secrete nectar is still largely unknown, as both internal and external factors can 321 affect the rate of secretion (Heil, 2011). In our study, floral rewards of nectar and pollen (apart from pollen weight) 322 significantly differed between test plant species. Similarly, nectar resources varied between periods of the day, 323 whilst the single measure of pollen for each replicate meant this resource could not be tested for diurnal differences. 324 Nectar volume and sugar concentration were negatively correlated in test plant species, with sugar concentration 325 greater in the afternoon, which corresponds with previous studies showing similar trends (Mačukanović-Jocić et 326 al., 2004; Silva et al., 2004; Mačukanović-Jocić & Đurđević, 2005). This may be due to decreased water

327 availability or flowers reabsorbing nectar in the afternoon (Silva et al., 2004) when humidity decreases and 328 temperatures rise (Silva et al., 2004; Mačukanović-Jocić & Đurđević, 2005). While our results suggests that nectar 329 resources vary in quantity and quality across the day, insect visitation did not track nectar availability (with the 330 exception of G. hederacea where the opposite was found: more insects were recorded visiting this species in the 331 afternoon when lower total sugar was recorded). This may be an indication that some pollinators are not capable 332 of accurately responding to changes in nectar production throughout the day, or it may be that their activity is 333 constrained by other factors such as temperature. It is important to note that our methods meant that afternoon and 334 evening nectar sampling occurred at 6-hour intervals, compared with morning sampling which was undertaken 335 12 hours after the flowers had been emptied and bagged the previous evening. We considered this as providing a 336 fair estimate of how much nectar is likely available to foraging insects by the morning, however, the effect of 337 early morning foraging between 05:00 and 09:00 GMT was not represented by our data and this should be taken 338 into account in interpretations.

339 Due to the high energy cost of foraging, successfully selecting the most rewarding flowers is predicted 340 to have a large impact on survival and reproductive success, especially when floral resources are fragmented. The 341 most documented resource offered to insects by plants as an attractant for pollination is nectar. Foraging insects 342 are capable of learning nectar rewards gained from visited flowers (Pyke, 1978), preferring to forage on flowers 343 with significantly more nectar (Heinrich, 1979a; Wolff, 2006) and with greater sugar concentration (Hendriksma 344 et al., 2014). After testing for relationships between nectar and pollen resources and pollinator visitations our 345 results support this, as typically, we found that greater nectar resources had a positive effect on insect and bee 346 visitation. Single species specialism in insect-plant mutualisms is rare (Waser et al., 1996), and the majority of 347 flower visiting insects have flexible foraging choices. Foraging bumblebees show flower constancy ('majoring' 348 on one particular species of known reward) and flower infidelity ('minoring' on other flowers checking reward 349 change over time) (Heinrich, 1979b). This behaviour allows foragers to track resources in multiple flower species 350 in a habitat. In our study, the positive effects of high floral rewards i.e. sugar were often only apparent when the 351 floral abundance of test plant species was high (> 50 inflorescences), with scarce flowers tending to be visited 352 less frequently even when comparatively highly rewarding. Our results appear to support Heinrich (1979b); if a 353 flower species is both abundant and rewarding then insects are very likely to have discovered its value and 354 preferentially visit it. In addition, where there are more flowers in a patch, there is a greater total quantity of nectar. 355 Hence, although quality has an influence on forager choice, the abundance of floral rewards in the local 356 environment is important in insect selection of resources.

357 There is evidence that bees show preferences for pollen with higher protein (Levin & Bohart, 1955; 358 Robertson et al., 1999) and essential amino acid content (Cook et al., 2003), and these preferences are supported 359 by obligate insect pollinated plant species producing pollen that is richer in protein and amino acids (Hanley et 360 al., 2008). However, other studies describe contrasting results where protein-rich pollen seems to be preferred in 361 some cases, but not in others (Wille et al., 1985 in Praz et al., 2008; Roulston & Cane, 2002). Our results suggest 362 greater total protein was negatively related to visitation by A. mellifera, and for B. pascuorum the negative 363 relationship between protein content and visitation was more apparent when the floral abundance of test plant 364 species was high (> 50 inflorescences). There are several possible explanations. Firstly, although not significant, 365 there was a negative relationship between nectar and pollen quantity per inflorescence (correlation coefficient -366 0.46), so if bees are basing decisions primarily on nectar rewards they will tend to visit flowers with less pollen. 367 We did not attempt to discern whether bees were collecting pollen only, nectar only, or both, but previous studies 368 suggest that the majority of visits are for nectar (e.g. Goulson et al., 2005). Second, bees may have been responding 369 to other nutritional compounds present in pollen. Plant species may be making trade-offs between protein and 370 other nutritional elements that drive foraging preferences such as sterols (Somme et al., 2014), lipids and starch 371 (Roulston & Cane, 2000) or pollen-specific odours (Dötterl & Vereecken, 2010) not addressed in this study. Third, 372 some plant species protect their pollen with defensive secondary compounds that may affect bee foraging choices 373 (Gosselin et al., 2013). For example, Echium vulgare has high protein content and also high concentrations of the 374 hepatotoxins 1,2-dehydropyrrolizidine alkaloids and their N-oxides (Boppré et al., 2005), which can be toxic to 375 insects (e.g. Boppré et al., 2005; Sedivy et al., 2012) and affect flower selection (Kessler & Halitschke, 2009). In 376 this study the species recorded with the greatest amount of pollen protein were Glechoma hederacea and 377 Ranunculus repens, and pollen of Ranunculus spp. is known to contain the toxic lactone protoanemonin 378 ranunculin with reportedly negative effects on honeybees (Sedivy et al., 2012; Jurgens & Dotterl, 2004); and G. 379 hederacea is toxic to some species of herbivorous insects as it produces a defensive insecticide protein (Hutchings 380 & Price, 1999; Van Damme, 2008). It is also important to note that the negative effect we found did not persist 381 for B. pascuorum once our values for Lamium album were replaced with higher values found in the literature. Our 382 methods for extracting and measuring protein content are crude, and results can be variable so should be treated 383 with caution. More detailed investigations are needed in which the full range of compounds present in pollen are 384 quantified if we are to fully understand how bees choose which flowers to visit when collecting pollen.

385 Most studies do not measure both nectar and pollen rewards in relation to insect visit frequency, and in
386 studies that do, conflicting results have been found. Konzmann & Lunau (2014) found that, in bumblebees, nectar

rewards appear more important than pollen quality, whereas Somme *et al.*, (2014) found when pollen loads are analysed in conjunction with nectar from forage plants, both nectar and pollen quality appear important. In this study, while pollen and nectar are not negatively correlated, total nectar production appears to influence the visitation of insects to a greater extent than pollen. This could mean that flower visiting insects are more concerned with the quality of nectar, with pollen as a secondary consideration.

392 Insect visitation to test plant species appears to be species-specific, which can go further to explain our 393 results. B. pascuorum mainly visited L. album and S. officinale, most likely due to its longer tongue allowing 394 access to their deeper corollas. Contrary to other visiting species, we detected a negative effect of total sugar on 395 B. pascuorum visitation. This may be because B. pascuorum had little competition from other insect foragers for 396 L. album (only 16 other individuals recorded foraging on this species, which produced comparatively low amounts 397 of sugar), or because this species was at the start of its life cycle when this study's sampling was undertaken and 398 newly emerged queens were focusing foraging efforts on pollen collection. Bombus pratorum, which as a short-399 tongued bee may have exhibited restricted foraging choices, tending to visit flowers with greater total sugar. 400 Although this species has a short tongue, it is a secondary nectar robber; 93% of recorded visits to S. officinale 401 were via robbing (behaviour previously reported in Goulson et al., 1998b). Apis mellifera, although also a short 402 tongued species, mainly visited R. fruticosus and was not recorded robbing in this study, though it has been 403 recorded acting as a secondary robber elsewhere (Darwin, 1872). Differences between foraging behaviour of 404 bumblebees and honeybees suggest bumblebees (B. terrestris and B. pascuorum) show less fidelity when 405 collecting pollen than honeybees, which have a highly flower-constant strategy (Leonhardt & Blüthgen, 2012).

406 Promoting and developing resources for pollinating insects is predominantly conducted through agri-407 environmental schemes promoting flower-rich field edges (Carvell et al., 2007), or through targeted planting in 408 urban spaces or private gardens (Hanley *et al.*, 2014). However, our understanding of the way in which pollinating 409 insects respond to differences in the quality of resources offered by managed planting is limited. Our results 410 suggest more consideration should be given when selecting plants for conservation management efforts, notably 411 in terms of differing insect species requirements for pollen and nectar quality. Differences in flower selection 412 between pollinator species may relate to the variation of life histories and may reduce competition for resources. 413 Nectar resource quality appears to be the main driver of flower selection by most insect foragers in this study but, 414 importantly, the benefits of greater resource quality in plants is dependent on local floral abundance. One practical 415 conclusion to be drawn from this is that bees may benefit more from plantings of flowers (be they in farmland,

416	parks or gardens	) where species are	presented in large clui	nps rather than in hetero	geneous mixtures. More broad	ly.
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417 it is clear that there is still much that we do not understand about the role of sugars, proteins and other compounds

418 in nectar and pollen in determining the foraging preferences of pollinators under field conditions.

419

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**Table 1** Measurements per inflorescence of nectar (30 replicates) and pollen (10 replicates) quality and quantity (Mean ± SD) for *Crataegus monogyna*, *Rubus fruticosus*,

*Glechoma hederacea, Lamium album, Ranunculus repens* and *Symphytum officinale* between times and totalled over 24 hours from seven sites in Sussex, UK.

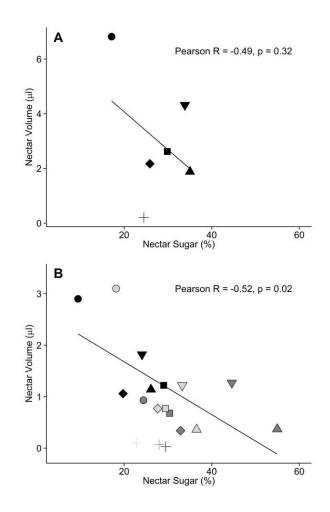
Nectar (µl)			1	Nectar (%)			Pollen (mg)	Pollen (%)	Sugar per Infl. (mg)				Protein per		
Test Plant Species	Morning	Afternoon	Evening	Over 24 Hours	Morning	Afternoon	Evening	Over 24 Hours	(8)		Morning	Afternoon	Evening	Over 24 Hours	Infl. (mg)
Crataegus monogyna	1.1 ± 1.7	$0.4 \pm 0.4$	$0.4 \pm 0.5$	1.9 ± 2.0	26.13 ± 2.26	54.95 ± 3.15	36.54 ± 1.81	35.03 ± 9.42	0.11 ± 0.03	15.39 ± 2.24	0.6 ± 0.9	0.6 ± 0.1	0.3 ± 0.2	1.2 ± 1.2	0.02 ± 0.01
Rubus fruticosus	2.9 ± 2.8	0.9 ± 1.2	3.1 ± 2.1	6.8 ± 3.7	9.48 ± 0.52	24.41 ± 1.87	18.16 ± 1.26	17.13 ± 4.72	0.48 ± 0.23	12.6 ± 2.28	$0.3 \pm 0.2$	$0.3 \pm 0.3$	0.6 ± 0.5	1.3 ± 0.7	0.06 ± 0.02
Glechoma hederacea	1.2 ± 0.6	0.7 ± 0.4	$0.8 \pm 0.4$	2.6 ± 0.8	29.03 ± 1.20	30.40 ± 1.57	29.43 ± 1.43	29.86 ± 4.56	0.89 ± 0.29	19.07 ± 3.46	$0.4 \pm 0.2$	$0.2 \pm 0.1$	0.2 ± 0.1	0.9 ± 0.2	0.17 ± 0.06
Lamium album	1.1 ± 0.4	$0.3 \pm 0.2$	$0.8 \pm 0.5$	2.2 ± 0.6	19.76 ± 0.98	32.88 ± 2.13	27.67 ± 1.38	25.88 ± 7.61	1.22 ± 0.33	4.48 ± 1.28 <sup>1</sup>	$0.2 \pm 0.1$	0.2 ± 0.1	$0.3 \pm 0.2$	0.6 ± 0.3	0.06 ± 0.02
Ranunculus repens	0.1 ± 0.1	0.1 ± 0.1	$\begin{array}{c} 0.0 \pm \\ 0.0 \end{array}$	0.2 ± 0.2	22.79 ± 2.25	28.00 ± NA	29.5 ± 2.60	24.50 ± 8.13	0.76 ± 0.3	12.06 ± 1.50	$0.0 \pm 0.0$	0.1 ± NA	$0.0 \pm 0.0$	0.1 ± 0.0	0.09 ± 0.04
Symphytum officinale	1.8 ± 0.8	1.3 ± 0.5	1.2 ± 0.6	4.3 ± 1.1	24.07 ± 1.06	44.59 ± 0.77	33.25 ± 1.61	33.88 ± 4.60	0.95 ± 0.31	9.71 ± 1.38	$0.5 \pm 0.2$	$0.7 \pm 0.2$	0.5 ± 0.2	1.7 ± 0.4	0.09 ± 0.03

<sup>1</sup>This value differed from the previously recorded value of 22.8% in Roulston *et al.* (2000).

Bold values indicate the total for nectar measurements over 24 hours, and the total for pollen measurements (for which there was only a single measure per replicate)

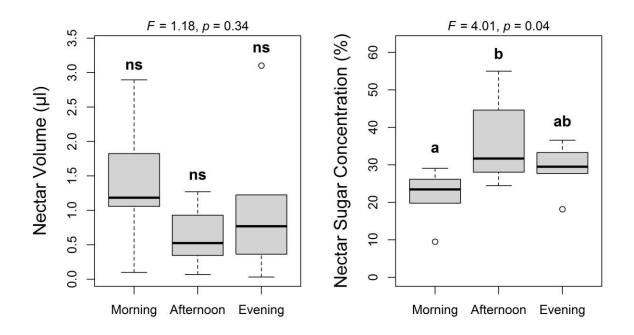
Table 2 Generalized Linear Mixed Models (GLMM) test results investigating the effects of floral abundance, total nectar and pollen quality and interactions between floral
 abundance and each quality metric on visitation of insect foragers from seven sites in Sussex, UK. Displayed are parameter estimates ± standard error, z-value and p-value for
 each explanatory variable in the final model after simplification, and model AIC and log-likelihood tests. P-values are in bold if < 0.05 significance.</li>

Visitation	Explanatory Variables left in model	Parameter Estimate ± SE	z- value	p-value	AIC	logLik
Apis mellifera	Log flower abundance	$-0.066 \pm 0.253$	-0.261	0.794	331.2	-159.6
	Total sugar	$7.845 \pm 3.288$	2.386	0.017		
	Total protein	$-2.491 \pm 0.927$	-2.687	0.007		
	Log flower abundance × Total sugar	$-1.405 \pm 0.639$	-2.201	0.028		
Bombus pratorum	Total sugar	$2.838 \pm 0.424$	6.687	< 0.001	423.2	-208.6
Bombus pascuorum	Log flower abundance	$0.281 \pm 0.249$	1.125	0.260	419.5	-203.7
	Total sugar	$-1.380 \pm 0.453$	-3.048	0.002		
	Total protein	-12.584 ± 4.158	-2.785	0.005		
	Log flower abundance × Total protein	$2.879 \pm 0.888$	3.241	0.001		
All Bombus Abundance	Log flower abundance	$0.849 \pm 0.168$	5.049	< 0.001	605.5	-297.7
	Total sugar	$6.275 \pm 1.889$	3.322	< 0.001		
	Log flower abundance × Total sugar	$-1.074 \pm 0.366$	-2.932	0.003		
All Insect Abundance	Log flower abundance	$0.425 \pm 0.124$	3.404	< 0.001	705.7	-347.9
	Total sugar	$4.741 \pm 1.522$	3.113	0.001		
	Log flower abundance × Total sugar	$-0.776 \pm 0.296$	-2.623	0.008		



613 **Fig. 1** Negative correlation between nectar volume (µl) and nectar sugar concentration (%) averaged between a) 614 test plant species (Pearson R = -0.49, p = 0.32) and b) test plant species and time period (Pearson R = -0.52, p = 615 0.02). ▼ = Symphytum officinale, ▲ = Crataegus monogyna, ■ = Glechoma hederacea, ◆ = Lamium album, • =

- 616 *Rubus fruticosus* and + = *Ranunculus repens*. Black symbols = Morning (09:00 GMT), dark grey symbols =
- 617 Afternoon (15:00) and light grey symbols = Evening (21:00) periods of nectar sampling.



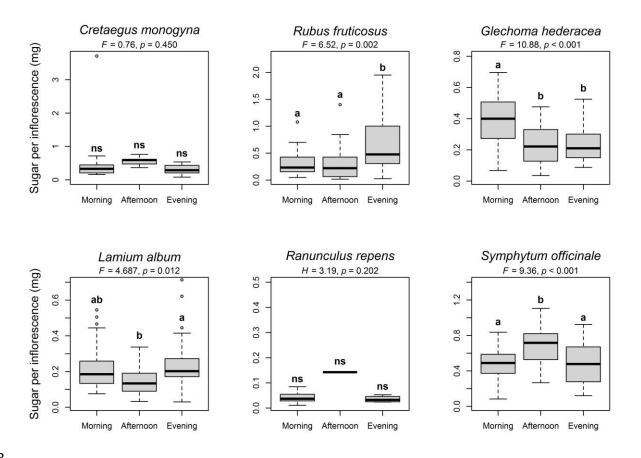


**Fig. 2** The mean nectar volume (μl) and mean nectar sugar concentration (%) measured from 30 inflorescences

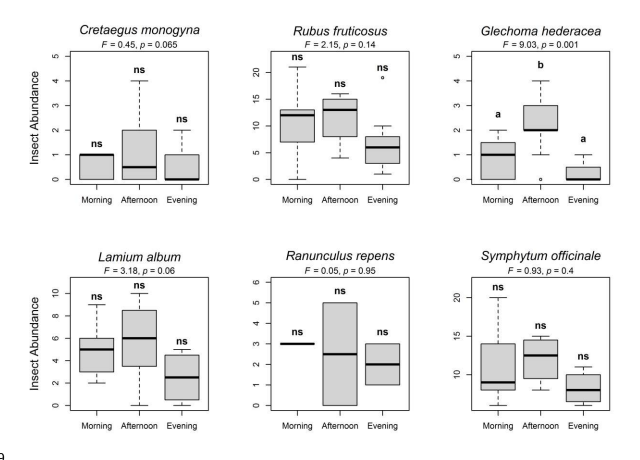
620 in Morning (09:00 GMT), Afternoon (15:00) and Evening (21:00) time periods averaged from six test plant

621 species. Significant differences between time periods were identified using post-hoc tests; time periods that do

622 not share a letter show significant variation (p < 0.05), ns = no significance.



**Fig. 3** Total sugar per inflorescence (mg) of test plant species at Morning (09:00 GMT), Afternoon (15:00) and Evening (21:00) time periods, with the test statistic ('F' for ANOVA and 'H' for Kruskall-Wallis, respectively) and significance levels for each analysis of variance between periods of the day for each species. Significant differences between time periods were identified using post-hoc tests; time periods that do not share a letter show significant variation (p < 0.05), ns = no significance.





**Fig. 4** Mean insect visits to test plant species in Morning (08:00–10:00 GMT), Afternoon (13:00–15:00) and

631 Evening (19:00—21:00) time periods, with the test statistic and significance levels of ANOVAs between

632 periods of the day for each species. Significant differences between time periods were identified using post-hoc

633 tests; time periods that do not share a letter showed significant variation (p < 0.05).

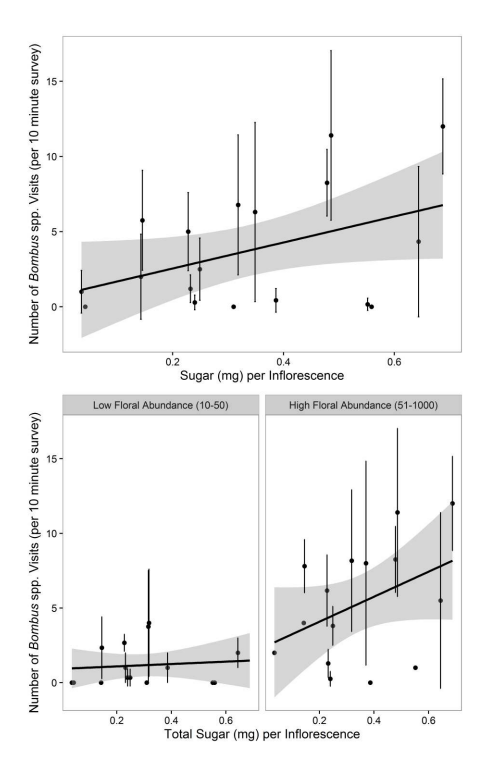


Fig. 5 Mean (± standard deviation) visits per test plant species to illustrate the positive effect of greater total
sugar on visitation in *Bombus* spp. (above), and total sugar on *Bombus* spp. visits (below) when separated
between sample plots with high or low floral abundance; low floral abundance (10—50 inflorescences per a 4
m<sup>2</sup>, left) shows little or no trend, whilst high floral abundance (51—1000 inflorescences, right) shows a positive
trend. Grey lines and polygons indicate model best fit and 95% Confidence Interval.