

1 **Quantifying exposure of wild bumblebees to mixtures of agrochemicals in agricultural and**  
2 **urban landscapes**

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9

10 **Abstract**

11 The increased use of pesticides has caused concern over the possible direct association of  
12 exposure to combinations of these compounds with bee health problems. There is growing  
13 proof that bees are regularly exposed to mixtures of agrochemicals, but most research has  
14 been focused on managed bees living in farmland, whereas little is known about exposure of  
15 wild bees, both in farmland and urban habitats. To determine exposure of wild bumblebees to  
16 pesticides in agricultural and urban environments through the season, specimens of five  
17 different species were collected from farms and ornamental urban gardens in three sampling  
18 periods. Five neonicotinoid insecticides, thirteen fungicides and a pesticide synergist were  
19 analysed in each of the specimens collected. In total, 61% of the 150 individuals tested had  
20 detectable levels of at least one of the compounds, with boscalid being the most frequently  
21 detected (35%), followed by tebuconazole (27%), spiroxamine (19%), carbendazim (11%),  
22 epoxiconazole (8%), imidacloprid (7%), metconazole (7%) and thiamethoxam (6%).  
23 Quantifiable concentrations ranged from 0.17 to 54.4 ng/g (bee body weight) for individual  
24 pesticides. From all the bees where pesticides were detected, the majority (71%) had more  
25 than one compound, with a maximum of seven pesticides detected in one specimen.  
26 Concentrations and detection frequencies were higher in bees collected from farmland  
27 compared to urban sites, and pesticide concentrations decreased through the season. Overall,  
28 our results show that wild bumblebees are exposed to multiple pesticides when foraging in  
29 agricultural and urban landscapes. Such mixtures are detected in bee tissues not just during  
30 the crop flowering period, but also later in the season. Therefore, contact with these  
31 combinations of active compounds might be more prolonged in time and widespread in the  
32 environment than previously assumed. These findings may help to direct future research and  
33 pesticide regulation strategies to promote the conservation of wild bee populations.

34 **Main findings:** There is a widespread presence of multiple pesticide residues in free-flying wild  
35 bumblebees, with different levels and frequencies of detection according to the species,  
36 sampling period and landscape context.

37 **Keywords:** neonicotinoids; fungicides; pesticide mixtures; wild bumblebees; pesticide  
38 exposure

## 39 **1. Introduction**

40 Bees are exposed to environmental pollutants via contaminated food resources such as pollen,  
41 nectar or water (Bonmatin et al., 2015), and through external contact with aerosols during  
42 spraying and contaminated dust emitted during the sowing of dressed seeds as their hairy  
43 bodies trap particulate residues (Greig-Smith et al., 1994; Pistorius et al., 2015). Many studies  
44 have used honeybees as relevant organisms to monitor environmental pollution (Celli and  
45 Maccagnani, 2003; Porrini et al., 2003). Bumblebees also forage in a great diversity of places  
46 and strongly interact with the environment, mainly the flora, surrounding their nests in a range  
47 of maximum foraging distances of 363 to 1650 m depending on the species (Walther-Hellwig  
48 and Frankl, 2000; Wood et al., 2015), and are thus also suitable organisms for monitoring  
49 landscape-based ecological pollution.

50 While most pesticide research has been focused on managed bees, there has been less work  
51 on wild bee populations. For instance, the only European bumblebee that has been studied in  
52 relation to pesticide exposure and toxicology is *Bombus terrestris*, simply because this species  
53 is easy to rear in captivity and commercially reared colonies are readily available (Baron et al.,  
54 2014; Gill et al., 2012; Rundlöf et al., 2015; Whitehorn et al., 2012).

55 There is increasing evidence that managed bees living in agricultural landscapes are routinely  
56 exposed to mixtures of agrochemicals (David et al., 2016; Lambert et al., 2013; Long and  
57 Krupke, 2016; Mullin et al., 2010; Pettis et al., 2013), but little is known about the exposure of  
58 wild bees in these environments (Hladik et al., 2016). Nevertheless, bees constitute a highly  
59 diverse group where different taxonomic groups differ widely in their vulnerability to pesticide  
60 exposure (Biddinger et al., 2013; Devillers et al., 2007; Piironen and Goulson, 2016; Thompson  
61 and Hunt, 1999). Furthermore, bee species exhibit pronounced differences in floral  
62 preferences and foraging habits, collecting pollen and nectar more frequently from particular  
63 plant species according to their morphological traits (e.g. tongue length, body shape and size)  
64 and nutritional needs (Goulson et al., 2008; Vanderplanck et al., 2014; Vaudo et al., 2016).  
65 Such foraging choices may profoundly influence the probability of bees to be more or less

66 exposed to some active compounds (Woodcock et al., 2016). Treated crop plants growing in  
67 agricultural landscapes have often been regarded as the only source of exposure to  
68 agrochemicals for pollinators, but recent research revealed their presence in wild plants  
69 growing near crops (Botías et al., 2015; David et al., 2016; Long and Krupke, 2016; Mogren and  
70 Lundgren, 2016). We would expect bees that visit flowering arable crops to have higher  
71 exposure than those that do not, but also those that visit wild plant species may have varying  
72 exposure depending on the ecology, physiology and morphology of their preferred flowers  
73 (Botías et al., 2016). Therefore, it is essential to understand the possible differences in levels of  
74 exposure among bee species, since this could reveal which are the most likely exposed and the  
75 most frequent mixtures of agrochemicals that they are exposed to.

76 The widespread occurrence of mixtures of agrochemicals in bee tissues (Hladik et al., 2016)  
77 increases concerns regarding the possible detrimental effects of simultaneous exposure to a  
78 cocktail of compounds. In general, only the effects of single active substances are studied in  
79 toxicity studies both for research and pesticide registration protocols, and exposure to  
80 mixtures are only evaluated in risks assessments when they are part of the same formulation.  
81 However, the application of two or more plant protection products during the same cropping  
82 season is a common practice in conventional farming (Botías et al., 2015; Garthwaite et al.,  
83 2013), and hence complex mixtures of agrochemicals which are not co-formulants of a single  
84 product can be simultaneously detected in bee forage and bee tissues (David et al., 2016;  
85 Hladik et al., 2016; Long and Krupke, 2016). This issue is worrisome given that exposure to  
86 mixtures might pose higher risks for animal health than the single impact of a specific class of  
87 compounds (Cedergreen, 2014; Rizzati et al., 2016). For example, some combinations of  
88 insecticides (*e.g.* pyrethroids with neonicotinoids) and of insecticides with fungicides can lead  
89 to additive and synergistic toxicity for bees at the individual and the colony level (Gill et al.,  
90 2012; Iwasa et al., 2004; Schmuck et al., 2003; Sgolastra et al., 2016). The scarcity of  
91 information on the field-relevant mixtures of agrochemicals and levels of exposure for bees  
92 could lead us to overlook the possible additive or synergistic effects of pesticide mixtures when  
93 risk assessment studies are performed, some of which have been designed to evaluate the  
94 hazards of such combinations (Sánchez-Bayo and Goka, 2014).

95 Another major gap in knowledge regarding exposure of bees to pesticides is the potential  
96 uptake and contact with these compounds in urban areas, where ornamental nursery plants  
97 can also be treated with pesticides (Brown et al., 2013; Fevery et al., 2016) and no information  
98 is available on their use in domestic gardens. The possible exposure of bees to harmful

99 pesticides through forage collected in gardens is of high ecological concern, since these  
100 habitats are of great value for bees, providing nectar, pollen and nest sites, and sustaining a  
101 remarkably high pollinator species richness and abundance (Baldock et al., 2015; Kaluza et al.,  
102 2016; Samnegård et al., 2011), including bumblebees (Fetridge et al., 2008; Goulson et al.,  
103 2010). If foraging resources and nesting sites in urban habitats are contaminated with  
104 pesticide residues, it is likely that exposure to certain active compounds could be more widely  
105 spread in the landscape and more prolonged in time than previously assumed.

106 The aim of this study was to evaluate and compare exposure in different wild bumblebee  
107 species. To do this, we analysed the levels of five neonicotinoid insecticides, thirteen currently-  
108 used fungicides and a pesticide synergist in tissues of five bumblebee species (*B. hortorum*, *B.*  
109 *pascuorum*, *B. terrestris*, *B. lapidarius* and *B. pratorum*). These wild bumblebee samples were  
110 collected in agricultural and urban habitats to compare levels of exposure in both  
111 environments and to study distribution of agrochemicals in the landscape. The bees were  
112 gathered in three different periods (late spring, early summer and midsummer) in order to  
113 monitor the length of exposure to agrochemicals through the season.

114 Our results show evidence that wild bumblebees are frequently exposed to mixtures of  
115 agrochemicals when they forage in arable and urban habitats, with peak concentrations  
116 decreasing through the season.

## 117 **2. Materials and Methods**

### 118 **2.1. Sampling sites and field collection**

119 Wild bumblebees were collected in five farms and five urban landscapes in East Sussex (South-  
120 East England, UK), all sites being at least 2 kilometers apart from each other (Figure 1) (Table  
121 S1). The sites selected to collect bees in agricultural land consisted of arable fields within  
122 mixed farms, where the predominant crops were oilseed rape, winter wheat and spring barley,  
123 and part of the land was pasture. The urban sampling sites consisted of ornamental public  
124 gardens and parks surrounded by houses that had private gardens in most cases. Foraging  
125 bumblebees were collected using insect nets and kept in individual labeled tubes and put on  
126 ice during transport back to the lab, and then kept at -80°C until pesticide analysis was  
127 performed. Specimens of five bumblebee species with different ranges of tongue length were  
128 sampled (Brodie, 1996; Prys-Jones and Corbet, 2011): short-tongued bumblebees were *B.*  
129 *pratorum* (6.4 - 7.1 mm), *B. lapidarius* (6 - 8.1 mm), *B. terrestris* (5.8 - 8.2 mm); medium-  
130 tongued was *B. pascuorum* (7.6 - 8.6 mm); and long-tongued was *B. hortorum* (12 - 13.5 mm)

131 (Table 1). The flowers where the bees were foraging at the time of capture were recorded  
132 (Tables S2a-S1c), since bumblebees exhibit a high degree of floral constancy (Wilson and Stine,  
133 1996), and this may help predict exposure.

134 Bumblebee individuals were gathered during three sampling periods, spring (27/04/14 –  
135 14/05/14), early summer (5/06/14 – 23/06/14) and midsummer (15/07/14 – 2/08/14), and 150  
136 bee individuals were collected in total. Oilseed rape crops were in bloom during the first  
137 sampling period (late spring), and 18 out of the 25 individuals gathered in arable sites during  
138 that period were foraging in oilseed rape crops when collected (Table S2a). The pesticide usage  
139 information of the crops where bees were foraging was not provided by the farmers. The EU  
140 moratorium on the use of neonicotinoid insecticides started on the 1<sup>st</sup> December 2013, but the  
141 oilseed rape crops that were in bloom in the 2014 spring were sown at the end of August-  
142 beginning of September 2013, so these crops were still allowed to be seed-treated with  
143 neonicotinoids.

## 144 **2.2. Pesticide analysis**

### 145 *Chemicals and reagents*

146 Eight classes of contaminants were chosen to be tested in the bumblebee samples, including  
147 the five neonicotinoid insecticides that are registered for use in the UK, based on the most  
148 used (by weight or area treated) in UK crops including oilseed rape, wheat, spring barley, field  
149 bean, strawberry and raspberry crops  
150 (<https://secure.fera.defra.gov.uk/pusstats/surveys/2014surveys.cfm>) (Table S3). Except for  
151 certified standards of carbendazim-d3 and tebuconazole-d6, which were purchased from LGC  
152 standards UK and carbamazepine-d10 and prochloraz-d7 from QMX Laboratories Limited UK,  
153 all the other certified standards as well as formic acid, magnesium sulphate, ammonium  
154 formate, sodium acetate and Supel<sup>TM</sup> QuE PSA/C18/GCB (ratio 1/1/1) were obtained from  
155 Sigma Aldrich UK. The compound purity of the pesticide standards was > 99%, apart from  
156 spiroxamine (98.5%), triticonazole (98.8%), piperonyl butoxide (97.9%). Isotopic purity of  
157 deuterated standards was > 97%. HPLC grade acetonitrile, toluene and water were obtained  
158 from Rathburns UK.

### 159 *Preparation of samples and residue analysis*

160 The extraction of pesticides from the bumblebee samples was performed as reported in David  
161 et al., 2015.

162 Briefly, individual whole bumblebee specimens (N = 150) were ground in liquid nitrogen and  
163 weighed (mean weight  $\pm$  S.D =  $108 \pm 46$  mg; range = 21 – 236 mg). Each sample was spiked  
164 with 10  $\mu$ l acetonitrile containing the mixture of deuterated internal standards (IS) at 40 ng/ml  
165 (400 pg of each IS). Subsequently, the extraction was performed by the addition of 400  $\mu$ l of  
166 water, 500  $\mu$ l of acetonitrile, 250 mg of magnesium sulphate: sodium acetate mix (4:1) and 50  
167 mg of SupelTM QuE PSA/C18/GCB for the dispersive solid phase extraction step (dSPE)  
168 (QuEChERS method). After the dSPE step, the sorbent was extracted with acetonitrile/toluene  
169 (3/1, 150  $\mu$ l) and the supernatant was combined with that of the previous acetonitrile extract  
170 and spin filtered (0.22  $\mu$ m). After evaporation, reconstitution was made with 120  $\mu$ l  
171 acetonitrile:water (30:70) and the extract was centrifuged (20 min). The supernatant obtained  
172 was stored in the dark at -20°C before analysis.

### 173 *UHPLC-MS/MS analyses*

174 Extracts of samples were analysed by ultra high-performance liquid chromatography tandem  
175 mass spectrometry (UHPLC-MS/MS) as described in David et al. (2015), using a Waters Acquity  
176 UHPLC system coupled to a Quattro Premier triple quadrupole mass spectrometer from  
177 Micromass (Waters, Manchester, UK). Acquisition of data was performed with the software  
178 MassLynx 4.1 and the quantification was established by the calculation of the response factor  
179 of each pesticide to its internal standard. Least-square linear regression analysis of the native  
180 analyte to deuterated internal standard (concentration ratio) versus the peak area was used to  
181 determine analyte concentrations. Further methodological details are described in David et al.  
182 (2015).

183 Table S4 reports the method detection limits (MDL) and the method quantification limits  
184 (MQL) for all the compounds that were analysed in bumblebees.

### 185 **2.3. Statistical analysis**

186 Shapiro-Wilk test was used to assess whether data (i.e. concentration of pesticides detected in  
187 the bumblebees per species, habitat, sampling period and bumblebee body mass) met the  
188 assumptions of normality. Since pesticide concentrations did not meet the assumptions of  
189 normality, non-parametric Kruskal-Wallis tests were used to compare this variable among the  
190 5 different bee species and 3 sampling periods. Pairwise comparisons between bee species,  
191 sampling periods and habitats were performed using Mann–Whitney U-tests. Bumblebee body  
192 mass and frequencies of pesticide detection in the 5 species were normally distributed, so one-  
193 way ANOVA analyses were carried out to compare weights of the 5 species, using Bonferroni

194 post-hoc tests for pairwise comparisons, and to compare the frequencies of detection among  
195 them. For the statistical analysis, the MDL value was assigned to concentrations above the  
196 limits of detection ( $\geq$ MDL) but below the limits of quantification ( $>$ MQL), whereas those below  
197 the MDL were considered to be zero. The software used to perform the statistical analysis was  
198 SPSS 21.

### 199 3. Results

#### 200 *Ranges, frequencies and average levels of neonicotinoid and fungicide residues detected in wild* 201 *bumblebee samples*

202 In total, 60.7% of the 150 individuals tested had detectable levels of at least one of the  
203 compounds analysed, with boscalid being the most frequently detected (in 35.3 % of all the  
204 bees analysed), followed by tebuconazole (27.3%), spiroxamine (18.7%), carbendazim (10.7%),  
205 epoxiconazole (8%), imidacloprid (7.3%), metconazole (6.7%) and thiamethoxam (6%) (Table 2)  
206 (Tables S2a-2c). In general, from all the bees where pesticides were detected, the majority  
207 (71.4%) had more than one compound, and 43.4 % of the bees contained two or more, with a  
208 maximum of 7 different pesticides being detected in one specimen (*B. lapidarius* collected  
209 from urban site 2 in June) (Table S2b).

#### 210 *Levels of pesticide exposure for the five bumblebee species studied*

211 Although the combinations of pesticides detected varied notably among individuals of the  
212 same and the other species, some combinations occurred very frequently in farmland  
213 bumblebees. Boscalid with DMI-fungicides was the most common mixture detected in  
214 bumblebees (64%) foraging in farmland in spring (April-May), except for *B. pratorum*  
215 individuals where such combination was not detected (Table S2a). In early summer (June)  
216 spiroxamine with DMI-fungicides was detected in the great majority of farmland bumblebees  
217 (88%), regardless of the species (Table S2b) (Figures 2 and 3). In general, the detection  
218 frequency of the 19 agrochemicals analysed did not significantly differ among the 5 species  
219 (ANOVA,  $F(4, 25) = 0.78$ ,  $P = 0.55$ ). However, when the concentrations of pesticides detected  
220 in the 5 bumblebee species were compared, we found that *B. pratorum*, the species with the  
221 lowest body mass and shortest tongue length range (Table 1), had significantly lower residue  
222 levels (mean  $\pm$  SD =  $1.7 \pm 3.6$  ng/g) than *B. hortorum* ( $4.7 \pm 10.1$  ng/g) (M-W test:  $U(58) = 291$ ;  
223  $P = 0.013$ ), *B. terrestris* ( $6.8 \pm 10.4$  ng/g) ( $U(58) = 275$ ;  $P = 0.006$ ), *B. lapidarius* ( $7.2 \pm 11.8$ )  
224 ( $U(58) = 260$ ;  $P = 0.003$ ) and also tended to have lower concentrations than *B. pascuorum* ( $2.8$   
225  $\pm 4.9$ ) ( $U(58) = 330$ ;  $P = 0.056$ ).

226 In order to evaluate the relationship between bee body size and levels of exposure, we  
227 compared the body mass of the five bumblebee species, and we found that *B. pratorum* was  
228 significantly lighter than the two species with the highest pesticide concentrations, *B. terrestris*  
229 (1-way ANOVA with Bonferroni's multiple comparison test:  $P = 0.03$ ) and *B. lapidarius* ( $P =$   
230  $0.017$ ).

#### 231 *Levels of pesticide exposure in arable and urban habitats*

232 In general, bees foraging in agricultural landscapes had significantly higher levels of  
233 agrochemicals ( $6.8 \pm 9.5$  ng/g) than those foraging in urban sites ( $2.5 \pm 7.8$  ng/g)(M-W test:  
234  $U(148) = 1635.5$ ;  $Z = -4.6$ ;  $P < 0.001$ )(Figure 2). However, the highest levels and frequencies of  
235 detection for neonicotinoids (10 ng/g of imidacloprid on a *B. terrestris* specimen) and the most  
236 frequently detected fungicide boscalid (54.5 ng/g in a *B. lapidarius* specimen) were recorded in  
237 urban bumblebees collected during the early summer (June) (Table S2b). Overall,  
238 neonicotinoids were found in more bees in urban sites than in farmland (9.3% versus 2.7%),  
239 with all five neonicotinoids registered for use in the UK found in at least one urban bee.

#### 240 *Changing levels of pesticide exposure through the season*

241 The levels of exposure to agrochemicals for wild bumblebees were examined for the period of  
242 highest foraging activity in the studied area (East Sussex, England), and we found that the  
243 frequencies of detection decreased both in arable and urban habitats for the 5 species  
244 evaluated in midsummer (July) (Figure 3), when only 28% of the bees collected had at least  
245 one agrochemical, compared to 76% in late spring (April-May) and 78% in early summer (June).  
246 Consequently, the average concentrations detected were lower in midsummer (July:  $0.6 \pm 2.3$   
247 ng/g) than in spring (April-May:  $5.9 \pm 7.6$  ng/g) (M-W test:  $U(98) = 474$ ;  $Z = -5.67$ ;  $P < 0.001$ )  
248 and early summer (June:  $7.5 \pm 12.4$  ng/g) (M-W test:  $U(98) = 462.5$ ;  $Z = -5.74$ ;  $P < 0.001$ ) (Table  
249 2).

## 250 **4. Discussion**

251 Our field study revealed that free-flying wild bumblebees are exposed to multiple pesticide  
252 residues, with different levels and frequencies of detection according to the species, sampling  
253 period and landscape context. Several studies have reported the presence of mixtures of  
254 agrochemicals in honeybee matrices (Lambert et al., 2013; Mullin et al., 2010; Pettis et al.,  
255 2013), where more than 170 compounds have been detected so far (Sánchez-Bayo and Goka,  
256 2014), but little is known about the exposure of wild bees. Recent research detected up to 19

257 different chemicals in native bees collected from wheat fields and grassland in Colorado (USA)  
258 (Hladik et al., 2016), reporting levels as high as 310 ng/g for thiamethoxam, 87 ng/g for  
259 clothianidin and 82 ng/g for imidacloprid. The maximum concentrations detected in our  
260 bumblebee samples were much lower (i.e. 2.35 ng/g for thiamethoxam, 1.4 ng/g for  
261 clothianidin and 10 ng/g for imidacloprid), which could be explained by the differences  
262 between both experimental designs. While we collected free-flying bumblebees with nets and  
263 analysed them individually, Hladik et al. (2016) performed the analysis on composite samples  
264 containing approximately 10 individuals of different wild bee species that were collected using  
265 bee monitoring traps. The collection and analysis of composite samples containing individuals  
266 from different bee genera might conceivably increase the chances of including particular  
267 specimens that could have been exposed to very high concentrations of pesticides due to their  
268 foraging and feeding behaviour, metabolic rates and morphological traits. Although the  
269 bumblebee species analysed in our study can present differences in their foraging distances,  
270 they seem to use and benefit similarly from the resources available in farmland (Wood et al.,  
271 2015), while bees from other genera, specially solitary bees, may present more variation in  
272 their foraging choices (Wood et al., 2016), and thus, a wider range of levels of exposure to  
273 agrochemicals. On the other hand, the dissimilarities in pesticide use patterns between the UK  
274 and the USA could partly explain the differences found, as for instance, the maximum  
275 application rate for thiamethoxam in oil seed crops in the UK is 33.6 g a.i./ha (European Food  
276 Safety Authority, 2013), and 157 g a.i./ha in the USA (USEPA, 2011). Studying the link between  
277 pesticide application rates and the levels of exposure for bees, and how different bee species  
278 can be more or less susceptible to exposure is essential for a full understanding of the risk  
279 posed by pesticides.

280 The comparison of pesticide concentrations among the five bumblebee species that we  
281 studied showed that *B. pratorum*, the species with the smallest body mass and tongue length  
282 range, had lower residue levels than the other four species. Different explanations are  
283 plausible; for example smaller bees may consume lower amounts of food, and hence they  
284 would be less exposed to these active compounds present in pollen and nectar. Smaller body  
285 size may lead to greater mass-specific metabolic rates (Heinrich, 1993), and so pesticides might  
286 be metabolised faster in smaller bees such as *B. pratorum*, whose body mass was significantly  
287 lower than that of the two species with the highest pesticide concentrations, *B. terrestris* and  
288 *B. lapidarius*.

289 Neonicotinoid insecticides can be metabolised relatively fast, with metabolites being the main  
290 residues detected in bees a few minutes after ingestion of the parent compound (Brunet et al.,  
291 2005; Suchail et al., 2004). Also the metabolites of some fungicides have been detected in bees  
292 and bee-collected pollen (Jabot et al., 2016; Mullin et al., 2010; Stoner and Eitzer, 2013), so the  
293 analysis of neonicotinoid and fungicide metabolites could have revealed the presence of other  
294 potentially toxic compounds that bees might have been exposed to. Some neonicotinoid  
295 metabolites have been proven to be highly toxic for bees (Simon-Delso et al., 2015; Suchail et  
296 al., 2004, 2001), while the possible effects of both parent fungicides and their metabolites has  
297 been scarcely studied in bees, with some reports showing detrimental effects (Bernauer et al.,  
298 2015; Pettis et al., 2013; Syromyatnikov et al., 2016; vanEngelsdorp et al., 2009). It is possible  
299 that the action of fungicides on bees may not be directly toxic, as is the case with insecticides,  
300 but may alter the beneficial microbiome present in the pollen and nectar (Bartlewicz et al.,  
301 2016; vanEngelsdorp et al., 2009; Yoder et al., 2013) and as a consequence, in the bee gut,  
302 which could have important implications for bee nutrition and health (Engel et al., 2016;  
303 Mattila et al., 2012; Pettis et al., 2013).

304 The tongue length of the different bumblebee species could be hypothesized as a possible  
305 predictor of residue exposure, since this trait determines whether or not, and how fast, a bee  
306 can manipulate a particular flower to extract nectar, and thus it is crucial for the division of  
307 resources between different species (Brian, 2016; Cariveau et al., 2016; Goulson et al., 2008;  
308 Harder, 2013). Long-tongued bees generally forage from long-corolla flowers, and short-  
309 tongued bees from short corolla flowers (Hobbs, 1962). In the case of *B. pratorum*, as a short-  
310 tongued bee, the most commonly visited flowers should be those with short corolla (*e.g.* many  
311 Rosaceae and Asteraceae flowers), which have both nectaries and stamens more exposed to  
312 environmental conditions and wind-blown aerosols. Oilseed rape flowers are shallow and  
313 more frequently visited by short-tongued bees, even though the long-tongued bumblebee *B.*  
314 *hortorum* often collects pollen from this plant (Stanley et al., 2013), and all our *B. hortorum*  
315 specimens sampled in farmland in late spring were foraging in these crop flowers when  
316 collected (Table S2a). Some of the pesticides analysed in this study have been reported to  
317 degrade after exposure to sun light and/or high temperatures (*i.e.* some neonicotinoids,  
318 carbendazim, carboxin, epoxiconazole and prochloraz) (Bonmatin et al., 2015; Burrows et al.,  
319 2002; Mazellier et al., 2002), so it is possible that the flowers with pollen and nectar more  
320 exposed to the environmental conditions might have lower concentrations of the parent active  
321 compounds. Therefore, the bees feeding on these shallow flowers would be less exposed to  
322 them. However, the tongue length range of *B. pratorum* is not very different to that of the

323 short-tongued bumblebees *B. terrestris* and *B. lapidarius* (Table 1). Moreover, the range of  
324 flowers visited by the three species did not differ remarkably, since more than half (53%) of  
325 the plant species visited by *B. terrestris* and *B. lapidarius* were also visited by *B. pratorum* in  
326 May and June (*i.e.* when concentrations detected were higher) (Tables S2a-2b). Nevertheless,  
327 as mentioned above, *B. terrestris* and *B. lapidarius* showed significantly higher levels of  
328 pesticide concentrations than *B. pratorum*, so the tongue length doesn't seem to be a suitable  
329 predictor of residue exposure for the group of bumblebees species studied here. Otherwise, a  
330 bigger sample size might be needed to test this hypothesis.

331 Regarding the toxicity of the pesticides detected, it is worth noting that the bumblebee  
332 specimens collected in the present study were individually processed as whole samples to  
333 include residues on external as well as internal parts of the bees, so it is not possible to  
334 differentiate if the pesticides detected were on the cuticle (contact toxicity) or inside the  
335 organism (oral). Thus, both routes of exposure should be considered when the levels detected  
336 in the samples are compared to lethal doses reported for the compounds analysed. Moreover,  
337 as the bee gut was not removed before processing the samples, there is a chance that some of  
338 the residues detected were present in the nectar and pollen contained in the digestive tract,  
339 although we consider that the bulk of the bee weights were formed by bee tissues. None of  
340 the residues detected in bumblebees were found to overlap with contact or oral acute LD<sub>50</sub>  
341 values tested on bumblebees or honeybees (Table 2), which is to be expected since the bees  
342 screened for pesticides in this study were performing foraging tasks and appeared to be  
343 healthy at the time of collection; it would be very unlikely to catch bees alive had they been  
344 exposed to lethal doses. Additionally, we cannot determine what doses the bees had been  
345 exposed to since pesticides are metabolized at varying rates (and we do not know the time of  
346 exposure), so that the residues we detected represent an unknown proportion of the dose  
347 received and actual exposures may have probably been higher. It should also be mentioned  
348 that bees are subjected to chronic exposure when foraging on contaminated flowers, and  
349 acute LD<sub>50</sub>s are frequently higher than chronic LD<sub>50</sub>s, particularly for neonicotinoids (Alkassab  
350 and Kirchner, 2016; Rondeau et al., 2014). Thus, the potential risk of chronic exposure to the  
351 levels of pesticides detected in the bumblebees cannot be ruled out. Studies where pesticide  
352 mixtures are analysed in bees showing health problems or mortality in the field would be of  
353 high interest (Kasiotis et al., 2014), since this could provide key information about the hazard  
354 posed by specific mixtures and the threshold levels of risk. Nevertheless, such studies are  
355 highly challenging to perform in the field due to sampling difficulties, and because the  
356 detection of accurate levels at the time of death are problematic since bee samples need to be

357 fresh to avoid degradation of pesticides following exposure to environmental factors and  
358 microbial activity (Katagi, 2004; Liu et al., 2011).

359 Our results revealed that 43.3% of bees contained two or more pesticides, suggesting that  
360 simultaneous exposure is likely to be encountered regularly in a field realistic scenario.  
361 Although contamination with mixtures of pesticides has been detected in flower pollen, bee  
362 collected pollen and bees (David et al., 2016; Hladik et al., 2016; Long and Krupke, 2016; Mullin  
363 et al., 2010), evaluating the impact of the exposure to such combinations of agrochemicals  
364 poses a major challenge and warrants more research. A few laboratory and field studies have  
365 explored the impact of simultaneous exposure to different chemicals on bees, some of them  
366 reporting detrimental effects of certain combinations (Gill et al., 2012; Iwasa et al., 2004;  
367 Johnson et al., 2013; Schmuck et al., 2003; Sgolastra et al., 2016; Thompson and Wilkins,  
368 2003). Given the mixtures detected in bees in the present study and in previous research  
369 (Hladik et al., 2016; Lambert et al., 2013), the number of possible combinations of pesticides is  
370 very high and variable. Therefore, focussing on the most frequent ones when performing  
371 assays of bee exposure might be the most sensible approach. Moreover, these scenarios are  
372 especially important to consider in cases when two or more pesticides that exhibit synergy are  
373 detected simultaneously in bees. For instance, the toxicity of certain insecticides (*e.g.*  
374 neonicotinoids and pyrethroids) can be enhanced in the presence of demethylation-inhibiting  
375 (DMI) fungicides (*e.g.* epoxiconazole, tebuconazole). In our study, 55.6 % of the bumblebees  
376 where neonicotinoid insecticides were detected also contained DMI-fungicides, so exposure to  
377 these combinations seems to be likely in the field although it is not known if these  
378 concentrations were high enough to induce biological effects. These DMI-fungicides can act as  
379 synergists by inhibiting the detoxification system in bees (Iwasa et al., 2004; Johnson et al.,  
380 2013; Pilling et al., 1995), and thus the insecticide residues are metabolised or eliminated more  
381 slowly. It is also important to remark that this is a limited list of pesticides; due to analytical  
382 constrains, insecticides such as pyrethroids, usually detected using gas chromatographic  
383 methods and which are known to interact with neonicotinoids and DMI-fungicides, could also  
384 be present in these bees.

385 As for the differences in levels of exposure for bees living in farmland or urban habitats, we  
386 found that concentrations of pesticides in wild bumblebees foraging in agricultural land were  
387 higher than in urban land, as reported for commercial bumblebee colonies in a previous study  
388 (David et al., 2016). However, the maximum values for neonicotinoids were recorded in  
389 bumblebees collected in urban gardens (*i.e.* 10 ng/g of imidacloprid, 2.35 ng/g of

390 thiamethoxam and 1.4 ng/g of clothianidin), even though these maximum values were high in  
391 comparison to the levels detected for these compounds in the rest of the bee specimens  
392 collected, so it is possible that these particular bees had been visiting freshly sprayed plants  
393 just before they were sampled. The use of imidacloprid, clothianidin and thiamethoxam has  
394 been banned since December 2013 on ornamental plants flowering in the year of treatment  
395 (as well as on flowering crops) (European Commission, 2013), and so the high levels of  
396 imidacloprid in particular are hard to explain. Nonetheless, the result is corroborated by David  
397 et al. (2016) who detected high levels (20 ng/g) of imidacloprid in pollen collected by  
398 commercial bumblebee colonies placed in urban areas. The imidacloprid may be persisting in  
399 urban environments from applications before the ban, or it might still be available in some  
400 stores and presumably may be used in gardens for months or even years after the ban.  
401 Alternatively, this exposure of bees could originate from other uses of imidacloprid; it is widely  
402 used in baits to kill ants and for flea control in dogs and cats. Orally applied imidacloprid in  
403 dogs and other mammals is completely eliminated in the urine (70-80%) and faeces (20-30%)  
404 in 48 h as the main metabolites 6-chloronicotinic acid and its glycine conjugate together with  
405 significant amounts of the parent compound, whereas topical application spreads over the  
406 skin for 24 hours and the compound is stored in the oil glands of the skin and slowly shed with  
407 hair and sebum (European Food Safety Authority, 2006; Hovda and Hooser, 2002). The  
408 environmental impacts of the use of pesticides in ornamental gardens and on pets has been  
409 scarcely studied (Brown et al., 2013; Fevery et al., 2016). There is no policing of homeowner  
410 use of garden pesticides to ensure that they follow label instructions, and amounts used in  
411 gardens are not known. Similarly, we can find no information on the number of dogs and cats  
412 treated with imidacloprid as a prophylactic flea treatment, and the environmental fate of such  
413 chemicals has not been studied. Urban gardens are an important food source and refuge for  
414 bees and other wildlife in cities and towns because they represent the only green space in  
415 these large environments, and they can host a great diversity of pollinators and high density of  
416 bumblebee nests (Baldock et al., 2015; Fetridge et al., 2008; Goulson et al., 2010), resulting in  
417 enhanced pollination services for both urban and nearby agricultural landscapes (Kaluza et al.,  
418 2016; Samnegård et al., 2011; Theodorou et al., 2016). If bees are sometimes exposed to high  
419 doses of harmful pesticides through forage collected in ornamental gardens, this might be a  
420 matter of high ecological concern, meaning that more attention should be paid to this route of  
421 contamination for bees.

422 The frequencies of detection and concentrations of agrochemicals in bee tissues decreased  
423 significantly in midsummer (July), agreeing with findings reported before (Botías et al., 2015;

424 David et al., 2016) where residue levels decreased in the pollen collected by bees as the season  
425 progressed. Midsummer is usually the period for crop harvesting in the studied area, and  
426 fewer pesticides are normally applied during this period to crops, so this may partly explain  
427 this finding. Concurrently, the decrease in residue levels may be due to a reduction in plant  
428 tissue concentrations, and thus on pollen and nectar collected by bees through summer  
429 because of plant/soil metabolism, photolysis and increasing temperatures (Bonmatin et al.,  
430 2015; Lassalle et al., 2014; Mazellier et al., 2002).

#### 431 *Conclusions and perspectives*

432 The extensive incidence of multiple pesticide residues and the scarcity of scientific literature  
433 on the biological consequences of exposure to such combinations on wild bees calls for more  
434 attention on regulatory strategies concerning monitoring procedures and registration of  
435 agrochemicals as they relate to pollinator protection, since the combined toxicity and  
436 synergism of all these chemicals may pose a real threat to the health and survival of managed  
437 and wild bees. Thus, investigations on the toxicological effects of field-realistic levels and  
438 mixtures are crucial to prevent potential exposure of bees to damaging combinations of  
439 agrochemicals. Also, the possible impact of metabolites and of agrochemicals other than  
440 insecticides on bees should not be overlooked, as they could have direct or indirect  
441 detrimental effects.

442 Understanding the factors involved in the degree of exposure, such as the type of flowers that  
443 tend to incorporate more residues or where certain pesticides are more persistent, as well as  
444 the morphological and physiological traits of pollinators that makes them more or less  
445 susceptible to the exposure, is also crucial to mitigate the damaging effects for the most  
446 vulnerable species.

447 Finally, the widespread detection of pesticides in bumblebees foraging in urban areas indicates  
448 a pressing need for further research on the prevalence and doses present in ornamental  
449 gardens, and on the environmental fate of pesticides upon domestic uses in order to better  
450 inform homeowners and garden centers of the potential risk the use of these products poses.  
451 Furthermore, surveillance programs on domestic uses would improve the current lack of safety  
452 and control in the application of agrochemicals in non-commercial agricultural situations.

453 Our results show evidence that wild bumblebees are frequently exposed to mixtures of  
454 agrochemicals when they forage in arable and urban habitats, with peak concentrations  
455 decreasing through the season. The effects of exposure to pesticide mixtures in wild bees

456 remain to be determined, but studying the temporal distribution of such combinations in  
457 habitats favored by bees is crucial to identify timing and routes of pesticide exposure, which  
458 may help us to properly direct our conservation efforts regarding pesticide regulation and bee  
459 health protection.

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720 **Tables and Figures**

721 Table 1. Mean ( $\pm$  standard deviation) and range of body mass values (mg), and tongue length  
722 range (mm) for the five bumblebee species analysed.

BUMBLEBEE SPECIES	BODY MASS (mg)		TONGUE LENGTH RANGE (mm) (Brodie, 1996; Pry-Jones and Corbet, 2011)
	MEAN $\pm$ S.D.	RANGE	
<i>B. hortorum</i>	105 $\pm$ 45	40 - 223	12 - 13.5
<i>B. pascuorum</i>	97 $\pm$ 34	29 - 171	7.6 - 8.6
<i>B. terrestris</i>	142 $\pm$ 46	45 - 236	5.8 - 8.2
<i>B. lapidarius</i>	117 $\pm$ 49	40 - 226	6 - 8.1
<i>B. pratorum</i>	79 $\pm$ 32	21 - 161	6.4 - 7.1

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738 Table 2. Frequencies of detection (%), maximum concentrations (Max), average (Avg) and  
739 median values (Mdn)(ng/g) of neonicotinoid insecticides and fungicides detected in wild  
740 bumblebees. The analytical methods do not allow us to differentiate what fraction of the  
741 pesticide was on the surface (contact toxicity) or inside the bumblebee (oral) since all  
742 specimens were individually processed as whole samples. Therefore, LD<sub>50</sub> values of contact (C)  
743 and/or oral (O) toxicity for honeybees (hb) or bumblebees (bb), according to availability of  
744 data (Sánchez-Bayo and Goka, 2014), are reported for each of the pesticides that were  
745 detected in wild bumblebees.

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	Contact (C) and/or oral (O) LD50s reported for bumblebees (bb) or honeybees (hb)	CONCENTRATIONS (ng/g)																								
		LATE SPRING (27/4/14 - 14/5/14)								EARLY SUMMER (5/6/14 - 23/6/14)								MIDSUMMER (15/7/14 - 2/8/14)								
		ARABLE (N=25)				URBAN (N=25)				ARABLE (N=25)				URBAN (N=25)				ARABLE (N=25)				URBAN (N=25)				
Freq. %	Max	Avg	Mdn	Freq. %	Max	Avg	Mdn	Freq. %	Max	Avg	Mdn	Freq. %	Max	Avg	Mdn	Freq. %	Max	Avg	Mdn	Freq. %	Max	Avg	Mdn			
<b>Neonicotinoid insecticides</b>																										
Thiamethoxam	C = 25 ng/hb; O = 5 ng/hb	8%	<0.90			12%	<0.90	<0.30	<0.30	0%					16%	2.35	<0.30	<0.30	0%				0%			
Clothianidin	C = 16 ng/bb	4%	<1.4			0%				0%					4%	1.4	<0.48	<0.48	0%				0%			
Imidacloprid	C = 20 ng/bb; O = 27 ng/bb	0%				16%	<2.2	<0.72	<0.72	0%					24%	10	<0.72	<0.72	4%	<2.2	<0.72	<0.72	0%			
Thiacloprid	C = 36,000 ng/hb; O = 17,000 ng/bb	0%				0%				0%					12%	1.17	<0.07	<0.02	0%				0%			
Acetamiprid	C = 100,000 ng/hb; O = 22,000 ng/bb	0%				0%				0%					4%	0.17	<0.01	<0.01	0%				0%			
<b>MBC-Fungicide</b>																										
Carbendazim	C = > 50,000 ng/hb	8%	1.2	<0.14	<0.05	24%	28.8	1.49	<0.05	8%	1.2	<0.14	<0.05	12%	4.37	0.23	<0.05		8%	<0.14	<0.05	<0.05	8%	0.92	<0.14	<0.05
<b>SDHI-Fungicides</b>																										
Carboxin		0%				0%				0%					0%					0%				0%		
Boscalid	C = > 200,000 ng/hb; O = 166,000 ng/hb	76%	31.7	7.03	6.2	44%	13.4	1.35	<0.24	44%	5.94	1.39	<0.05	36%	54.4	2.65	<0.05		12%	8.51	0.50	<0.05	0%			
<b>Amine fungicide</b>																										
Spiroxamine	C = 4,200 ng/hb; O = 92,000 ng/hb	4%	3.8	0.15	<0.05	0%				96%	37.7	6.51	2.91	0%					12%	2.19	0.20	<0.05	0%			
<b>DMI fungicides</b>																										
Triticonazole		0%				0%				0%				0%						0%				0%		
Epoxiconazole	C = > 100,000 ng/hb	48%	6.07	1.03	<0.96	0%				4%	<2.9	<0.96	<0.96	0%					0%					0%		
Tebuconazole	C = > 200,000 ng/hb; O = 83,000 ng/hb	12%	1.7	<0.12	<0.12	8%	<0.36	<0.12	<0.12	88%	11.7	2.91	1.59	28%	1.5	<0.36	<0.12		32%	4.95	<0.36	<0.12	0%			
Flusilazole		0%				0%				0%				0%						0%				0%		
Prochloraz	C = > 50,000 ng/hb; O = 60,000 ng/hb	0%				0%				4%	<0.90	<0.30	<0.30	4%	<0.90	<0.30	<0.30		0%				0%			
Metconazole	C = > 100,000 ng/hb; O = 80,000 ng/hb	20%	<0.72	<0.24	<0.24	8%	4	<0.24	<0.24	0%				4%	2.58	<0.24	<0.24		0%				8%	<0.72	<0.24	<0.24
<b>QoI-fungicides</b>																										
Pyraclostrobin	C = > 100,000 ng/hb; O = 73,000 ng/hb	4%	<0.72	<0.24	<0.24	0%				0%				0%					0%					0%		
Trifloxystrobin	C = > 200,000 ng/hb; O = 200,000 ng/hb	4%	2.74	0.11	<0.01	4%	2.74	0.11	<0.01	12%	<0.04	<0.01	<0.01	0%					0%					0%		
Fluoxastrobin	C = > 200,000 ng/hb; O = 843,000 ng/hb	12%	<0.72	<0.24	<0.24	4%	<0.72	<0.24	<0.24	4%	<0.72	<0.24	<0.24	0%					0%					0%		
<b>Synergist</b>																										
Piperonyl butoxide		0%				0%				0%				0%					0%					0%		

Figure 1. Location of sites in East Sussex (UK), where wild bumblebees were collected from farmland sites (circles) and urban sites (squares). The sites selected to collect bees in agricultural land consisted of arable fields within mixed farms, where the predominant crops were oilseed rape, winter wheat and spring barley, and part of the land was pasture. The urban sampling sites consisted of ornamental public gardens and parks surrounded by houses that had private gardens in most cases.



Figure 2. Sum of total concentrations of neonicotinoid insecticides and fungicides detected in 5 different species of wild bumblebees collected in arable (A) and urban (U) sites in three sampling periods (May, June and July). Concentrations reported refer to the amount of active substance (ng) detected per bumblebee body weight (g).

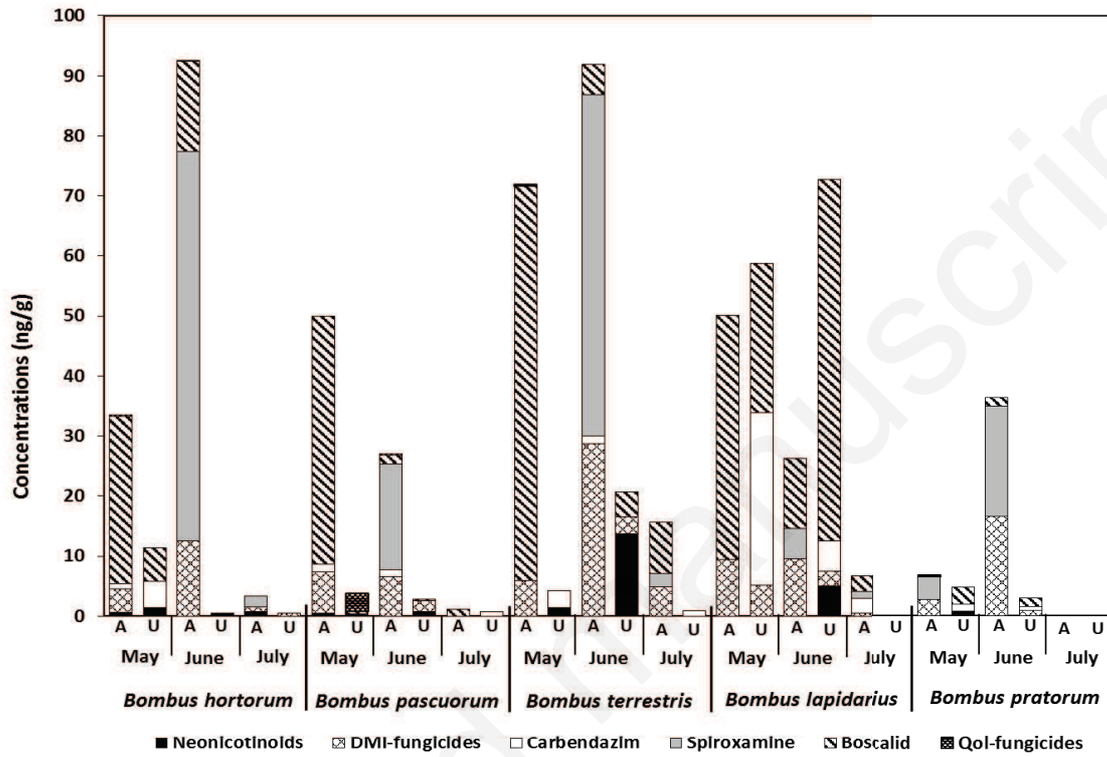


Figure 3. Frequencies of detection (%) of pesticide residues in wild bumblebees collected in arable and urban habitats in May, June and July.

