

# Sussex Research

### Combined stress from parasites, pesticides and lack of flowers drives bee declines

Dave Goulson, Beth Nicholls, Cristina Botias Talamantes, Ellen Rotheray

#### Publication date

27-03-2015

#### Licence

This work is made available under the Copyright not evaluated licence and should only be used in accordance with that licence. For more information on the specific terms, consult the repository record for this item.

#### Citation for this work (American Psychological Association 7th edition)

Goulson, D., Nicholls, B., Botias Talamantes, C., & Rotheray, E. (2015). *Combined stress from parasites, pesticides and lack of flowers drives bee declines* (Version 1). University of Sussex. https://hdl.handle.net/10779/uos.23416976.v1

Published in

Science

Link to external publisher version https://doi.org/10.1126/science.1255957

#### Copyright and reuse:

This work was downloaded from Sussex Research Open (SRO). This document is made available in line with publisher policy and may differ from the published version. Please cite the published version where possible. Copyright and all moral rights to the version of the paper presented here belong to the individual author(s) and/or other copyright owners unless otherwise stated. For more information on this work, SRO or to report an issue, you can contact the repository administrators at sro@sussex.ac.uk. Discover more of the University's research at https://sussex.figshare.com/

## Sussex Research Online

# Combined stress from parasites, pesticides and lack of flowers drives bee declines

Article (Unspecified)

Goulson, Dave, Nicholls, Beth, Botias Talamantes, Cristina and Rotheray, Ellen (2015) Combined stress from parasites, pesticides and lack of flowers drives bee declines. Science, 347 (6229). ISSN 0036-8075

This version is available from Sussex Research Online: http://sro.sussex.ac.uk/id/eprint/54228/

This document is made available in accordance with publisher policies and may differ from the published version or from the version of record. If you wish to cite this item you are advised to consult the publisher's version. Please see the URL above for details on accessing the published version.

#### Copyright and reuse:

Sussex Research Online is a digital repository of the research output of the University.

Copyright and all moral rights to the version of the paper presented here belong to the individual author(s) and/or other copyright owners. To the extent reasonable and practicable, the material made available in SRO has been checked for eligibility before being made available.

Copies of full text items generally can be reproduced, displayed or performed and given to third parties in any format or medium for personal research or study, educational, or not-for-profit purposes without prior permission or charge, provided that the authors, title and full bibliographic details are credited, a hyperlink and/or URL is given for the original metadata page and the content is not changed in any way.

1	Combined stress from parasites, pesticides and lack of flowers drives bee declines
2	
3	
4	Dave Goulson, Elizabeth Nicholls, Cristina Botías and Ellen L. Rotheray
5	
6	
7	School of Life Sciences, University of Sussex, Falmer, Brighton, BN1 9QG
8	Email: D.Goulson@sussex.ac.uk
9	
10	

#### 11 Abstract

12 Bees are subject to numerous pressures in the modern world. The abundance and diversity of 13 flowers has declined, bees are chronically exposed to cocktails of agrochemicals, and they are 14 simultaneously exposed to novel parasites accidentally spread by humans. Climate change is 15 likely to exacerbate these problems in the future. Stressors do not act in isolation; for example pesticide exposure can impair both detoxification mechanisms and immune responses, 16 17 rendering bees more susceptible to parasites. It seems certain that chronic exposure to 18 multiple, interacting stressors is driving honey bee colony losses and declines of wild 19 pollinators, but such interactions are not addressed by current regulatory procedures and 20 studying these interactions experimentally poses a major challenge. In the meantime, taking 21 steps to reduce stress on bees would seem prudent; incorporating flower-rich habitat into 22 farmland, reducing pesticide use through adopting more sustainable farming methods, and 23 enforcing effective quarantine measures on bee movements are all practical measures that 24 should be adopted. Effective monitoring of wild pollinator populations is urgently needed to 25 inform management strategies into the future.

26

#### 27 Is there a 'pollination crisis'?

28 Insect pollination is vitally important to terrestrial ecosystems and to crop production. The 29 oft-quoted statistics are that 75% of our crop species benefit from insect pollinators (1), 30 which provide a global service worth \$215 billion to food production (2). Hence the potential 31 that we may be facing a "pollination crisis" (3, 4) in which crop yields begin to fall because 32 of inadequate pollination has generated understandable debate and concern and stimulated 33 much research in recent decades. Nonetheless, knowledge gaps remain substantial, both with 34 regard to the extent and causes of pollinator declines. Indeed, for most regions of the globe 35 and for most wild pollinator taxa, we have no data as to whether there have actually been 36 declines. Our best estimates are for numbers of domesticated honey bee colonies, which can 37 be obtained for many countries with varying reliability. Overall, these suggest that numbers 38 of managed honey bee colonies have decreased in Europe (25% loss of colonies in central 39 Europe between 1985 and 2005 (5)), and markedly in North America (59% loss of colonies between 1947 and 2005 (6, 7)). However, overall global stocks actually increased by  $\sim 45\%$ 40 41 between 1961 and 2008, due to a major increase in numbers of hives in countries such as 42 China and Argentina (8). Conversely, there are widespread reports of unusually high rates of 43 honey bee colony loss from many parts of the world, sometimes ascribed to a syndrome

known as Colony Collapse Disorder (CCD) (9). It seems that socioeconomic factors (such as
increasing demand for pollination or honey (10), are at present sufficient to incentivise
beekeepers to overcome problems with bee health, when examined at a global scale (but not
locally in North America and Europe).

48 If we turn to wild pollinators, the best data available are for bumblebees (11). In 49 Europe, many species have undergone substantial range contractions and localized extinction, 50 with four species going extinct throughout the continent (11, 12) (Figure 1a). In North 51 America, some formerly abundant and widespread species such as *Bombus terricola* and *B*. 52 occidentalis underwent severe declines from the late 1990s onwards, and now occupy only a 53 small fraction of their former range (Figure 1b) (13, 14). Bombus franklini, a species formerly 54 found in northern California and Oregon, has not been recorded since 2006 and is presumed 55 extinct. In a study of the bumblebee fauna of Illinois over the past 100 years, Grixti et al. (15) describe substantial declines in species diversity, particularly in the period 1940-1960, with 56 the extirpation of four species during the 20<sup>th</sup> century. In South America, the recent invasion 57 58 by the European species *B. terrestris* is causing precipitous declines in the native *B*. 59 dahlbomii (16). There is some evidence of loss of species richness from lowland areas of 60 Sichuan in China (17, 18), and a few reports of declines in Japan from the mid 1990s 61 onwards (19, 20), but elsewhere in the world, few data are available. For the remaining wild 62 bees, data are exceedingly sparse, though they comprise the large majority of the world's 63 approximately 22,000 bee species. Analysis of historic records suggests that diversity of both bumblebees and other wild bees declined in the UK, Netherlands and Belgium during the 64 65 twentieth century, but that these declines have decelerated since 1990 (21, 22). In surveys in Illinois, 50% of wild bee species went extinct over a 120 year period to 2010 (23). Given that 66 67 bee diversity has declined in both Europe and the Americas, it is probably reasonable to 68 assume that declines are also occurring elsewhere across the globe

The biggest knowledge gap is regarding bee abundance; although we have maps of past and present distributions of bees for some well-studied countries such as the UK, we have almost no data on how populations have changed over time. Hence we do not know whether common species such as *Bombus terrestris* in Europe or *Bombus impatiens* in North America are less abundant than formerly, or whether they are currently in decline. Most pollination is delivered by a small number of these abundant species which tend to have large distributions. Declines in their abundance would not be detected in distribution maps until they become extinct in parts of their range, which is rather late to introduce conservationmeasures.

78 Another way to examine the likelihood or proximity of a pollination crisis is to 79 examine delivery of pollination services. Although global honey bee stocks have increased by 80 ~45%, demand has risen more than supply, for the fraction of global crops that require animal 81 pollination has tripled over the same time period (8), making food production more 82 dependent on pollinators than before. It has also emerged that the majority of crop 83 pollination, at a global scale, is delivered by wild pollinators rather than honey bees. Yields 84 correlate better with wild pollinators abundance than with abundance of honey bees (24-26); 85 hence increasing honey bee numbers alone is unlikely to provide a complete solution to the 86 increasing demand for pollination. Reliance on a single species is also a risky strategy (27). 87 Whilst Aizen et al. (28) concluded from a global analysis of changing crop yields over time that there was not yet any clear evidence that a shortage of pollinators was reducing yield, a 88 89 subsequent analysis of the same data set by Garibaldi et al. (29) shows that yields of 90 pollinator-dependent crops are more variable, and have increased less, than crops that do not 91 benefit from pollinators, to the extent that a shortage of pollinators is reducing the stability of agricultural food production. In a meta-analysis of 29 studies on diverse crops and contrasting 92 93 biomes, Garibaldi et al. (30) found that wild pollinator visitation and yields generally drop 94 with increasing distance from natural areas, suggesting that yields on some farms are already 95 impacted by inadequate pollination.

To summarize, in the past 50 years global honey bee stocks have increased, while wild bees appear to have declined substantially, as evidenced by data for bumblebees, and very scant data for other bee species. The demand for insect pollinators in farming has tripled in the same period. There is clearly no major pollination crisis yet, but there is evidence for localized limitation of crop yield as a result of inadequate pollination.

101

#### 102 Drivers of wild bee declines and honey bee colony losses

103

#### 104 <u>Habitat loss</u>

105 Bee declines have been attributed to many factors, some more plausible than others; however,

106 the clear consensus is that loss of habitat has been a long-term contributor to declines (11,

107 *31–33*). Bees require appropriate floral resources during the adult flight season, which may

108 be short for some solitary species or year-long for social species in tropical environments.

109 They also require undisturbed nest sites, with different species occupying diverse locations 110 (e.g.' cavities underground, hollow-stemmed twigs, burrows in the soil, even abandoned snail 111 shells). The conversion of natural and semi-natural flower-rich habitat to farmland has been a 112 major driver of long-term declines in bees. For example, in the UK, approximately 97% of flower-rich grasslands were lost in the  $20^{\text{th}}$  century (34), and this has resulted in major range 113 114 contractions of bee species associated with this habitat, particularly long-tongued bumblebees 115 (11). Declines of similar magnitude have affected the grasslands of North America, although these began in the early  $19^{\text{th}}$  century (35). 116

Urbanization also contributes to loss of natural habitat, but the net effect on bees is 117 118 less clear. Gardens can support high densities of wild bees, particularly bumblebees and some 119 solitary bee species, but highly urbanized environments have few bees (36, 37), and the 120 building of roads and other infrastructure undoubtedly contributes to the ongoing 121 fragmentation and degradation of habitats. Increased traffic can also cause direct mortality 122 through collisions (38), although the numbers of bees killed in this way is not known. The 123 planting of road verges and traffic islands with wildflowers is often promoted as a means of 124 boosting urban pollinator populations, but it might increase mortality by vehicle collisions. 125

#### 126 Parasites and disease

127 Bees naturally suffer from a broad range of parasites, parasitoids and pathogens, the latter 128 including protozoans, fungi, bacteria and viruses. By far the majority of research has focused 129 on those associated with honey bees and to a lesser extent with bumblebees, with very little 130 known about the pathogens of other wild bee species. Some bee diseases, such as Deformed 131 Wing Virus (DWV) and Nosema ceranae, have broad host ranges and are able to infect both 132 honey bees and bumblebees while others, such as Crithidia bombi or Paenibacillus larvae, 133 seem to be more host-specific (39-41). While natural pathogens undoubtedly play an 134 important but poorly-understood role in influencing the population dynamics of their bee 135 hosts, we will focus here on the impacts that non-native parasites and pathogens may have. 136 The spread of most honey bee parasites and pathogens has occurred inadvertently as 137 a result of transporting honey bees long distances. Much of this happened in historic times,

but it continues, despite some improvements in quarantine procedures. The best-known
example is the mite *Varroa destructor*, originally associated with the Asian honey bee *Apis*

- 140 *cerana*. Varroa has since jumped hosts to the European honey bee *Apis mellifera*, which has
- 141 little resistance to this pest. Since the 1960s Varroa has spread from Asia to Europe, the

Americas and most recently to New Zealand. The mite acts as a vector for pathogens such as DWV, and the combined effect of the mite and the diseases it transmits is a major contributor to honey bee colony losses in North America and Europe (42, 43). Fortunately, the mite

145 appears unable to survive on bees outside the genus *Apis*.

146 A strikingly similar series of events has also seen the microsporidian N. ceranae jump from A. cerana to A. mellifera, and in the past 20 years it has spread to Europe and the 147 148 Americas, where it is now prevalent at high frequency (Figure 2) (44, 45). It has also been 149 detected in wild bumblebees in Europe, China and South America (41, 46, 47), and solitary 150 bees in Europe (48). In the lab N.ceranae appears to have higher virulence in bumblebees 151 than it does in honey bees (41), though the impact it has had on wild populations is unknown. 152 Asia is not the only source of non-native diseases; the African honey bee parasite Aethina 153 tumida (small hive beetle) recently invaded North America, Egypt, Australia and Europe, and 154 attacks *B. impatiens* colonies causing considerable damage (49, 50). It seems highly likely 155 that it also attacks other wild bumblebee species that are not so readily cultured and therefore 156 less well studied.

157 Bee diseases are also being redistributed around the globe by the commercial trade in 158 bumblebee colonies, which are mainly used for pollination of greenhouse crops such as 159 tomatoes. This trade began in the 1980s in Europe, and now more than 1 million nests of the 160 European Bombus terrestris are reared each year and exported to various countries. In North 161 America, the eastern American species *Bombus impatiens* is reared for this purpose. 162 Unfortunately, it does not seem possible yet to rear colonies that are free of disease, not least 163 because the bees are reared on honey-bee collected pollen, providing a route for exposure to many bee pathogens. Commercial colonies of *B. terrestris* are commonly infected with one or 164

165 more parasites, including Nosema bombi, N. ceranae, Apicystis bombi and DWV (41).

166 There is evidence that non-native pathogens or pathogen strains associated with these 167 colonies are having devastating impacts on wild bumblebee populations. In North America, 168 the accidental importation of a non-native strain of the parasite Nosema bombi via 169 commercial bumblebees has been implicated in the dramatic decline of several bumblebee 170 species, though convincing causal evidence remains elusive (51, 52). The evidence from 171 South America is clearer; here, B. terrestris were deliberately introduced by the Chilean 172 government despite the presence of native *Bombus* species, with *terrestris* spreading rapidly 173 to occupy a vast area of southern South America. The arrival of *B. terrestris* appears to have

174 led to the rapid local extinction of the native *B. dahlbomii* at a speed only plausibly explained

by pathogen spillover (*16*). Although the parasite responsible has yet to be ascertained with

176 certainty, both A. bombi and C. bombi have been shown to be highly prevalent in the invasive

- 177 species (16, 53). There is a clear parallel with the devastating impact that the arrival of
- 178 European diseases had on the native Americans 500 years ago.

179 Even when commercial bees are free of disease on arrival, or are infected only with 180 indigenous parasites, they can still affect native pollinators. High-density populations of 181 managed bees may provide conditions for the rapid multiplication of parasites that then spill 182 over into wild populations (54–56). A combination of field observations and modelling 183 suggest that waves of *Crithidia bombi* infection travel outwards from greenhouses containing 184 commercial bumblebees. Prediction indicates that waves can spread at ~2 km per week, with 185 up to 100% of wild bees within the spreading radius becoming infected, although this is not yet well supported by direct evidence (55). 186

187 In general we know little about the natural geographic range, host range, prevalence188 or virulence of most bee pathogens, and so it would seem wise to take very careful

189 precautions to prevent further introductions of bee pathogens from outside their native range,

190 in addition to minimizing any spillover from commercial pollination operations (52, 57).

191

#### 192 <u>Pesticides</u>

193 The most controversial and debated cause of bee declines are pesticides. When appropriately 194 used, pesticides can provide an economic benefit, but bring the welfare of bees into direct 195 conflict with industrial agriculture. Herbicides are highly effective at minimizing weed problems in most cropping systems, enabling farmers to grow near-pure monocultures, but 196 197 their use inevitably reduces the availability of flowers for pollinators and can contribute 198 substantially to rendering farmland an inhospitable environment for bees (11, 58, 59). 199 Understandably, most attention has been paid to the direct toxic effects of pesticides on bees, 200 particularly the impacts of insecticides. One hundred and sixty one different pesticides have 201 been detected in honey bee colonies (60, 61) and based on their toxicity, frequency in hives 202 and the concentrations detected, Sanchez-Bayo and Goka (61) predict that three 203 neonicotinoids (thiamethoxam, imidacloprid and clothianidin), and the organophosphates 204 phosmet and chlorpyrifos pose the biggest risk to honey bees at a global scale. It is clear that 205 bees are often chronically exposed to cocktails of pesticides throughout their development 206 and adult life (61-64), but the effects of this are poorly understood and are not examined by

207 current regulatory risk-assessment procedures (65).

208 Neonicotinoids are the newest of the main classes of insecticide, and the group most 209 strongly implicated in bee declines (65, 66). They are neurotoxins that target the insect 210 central nervous system, binding to postsynaptic nicotinic acetylcholine receptors and causing 211 over-stimulation, paralysis and death (67). These insecticides are commonly applied as seed-212 treatments and are systemic within plants, spreading through plant tissues and into the pollen 213 and nectar of flowering crops such as canola. They are also water soluble but highly 214 persistent in soil and soil water, and as a result have been found at significant concentrations 215 in the pollen and nectar of wildflowers near crops (63). Thus there is a clear route for 216 ingestion by bees. Oral toxicity is high, with the short-term LD<sub>50</sub> for the most commonly-217 used neonicotinoids in the region of 4-5ng/honey bee (68) ( $LD_{50}$  = Lethal Dose 50%, the 218 dose that kills 50% of test organisms). Long-term chronic exposure results in mortality in 219 overwintering honey bees when feeding on food contaminated with concentrations as low as 220 0.25ppb (69). Sub-lethal effects of neonicotinoid exposure have also been observed in both 221 honey bees and bumblebees, including a reduction in learning, foraging ability and homing 222 ability, all of which are essential to bee survival (70-74). Yang et al. (75) recently showed that even low exposure during the larval stage (0.04ng/larva equating to less than 1/100<sup>th</sup> of 223 224 the LC<sub>50</sub> for adult bees) can have a lasting impact on learning in adult honey bees (LC<sub>50</sub> = 225 Lethal Concentration 50%, the concentration that kills 50% of test organisms).

226 It seems very likely that bees living in most arable farmland are routinely exposed to 227 sufficient neonicotinoids to suffer both lethal and sublethal effects. However, whether this 228 translates into a detrimental effect at the colony-level remains disputed. In bumblebees, 229 exposure of colonies to field-realistic concentrations of imidacloprid greatly impaired colony 230 growth and reduced queen production by 85% (76). In contrast, field studies with honey bee 231 colonies have proved more challenging to perform convincingly (77, 78), not least because of 232 the huge areas over which honey bees forage, the lack of a clear end-point to colony 233 development, and their long-term storage of food reserves. This lack of clear evidence of 234 harm in the field is often misinterpreted as evidence that toxicological studies on individual 235 bees do not translate into colony losses in the field, rather than as the absence of evidence. 236

237

238 <u>Monotonous diets</u>

239 Intensively farmed areas provide few wildflowers, but do provide spatially and temporally

240 isolated gluts of flowers in the form of mass-flowering crops such as sunflowers and canola.

241 If a human were to consume nothing but sardines one month, chocolate the next, turnips the 242 month after and so on, one could reasonably expect them to fall ill. This may seem a 243 frivolous example, but it is a reasonable parallel to the experience of some honey bee 244 colonies, particularly those in North America that are transported backwards and forwards 245 across the continent each year to provide pollination for major crops such as almonds in California, blueberries in Maine and citrus in Florida. Where the nectar or pollen of crop 246 247 flowers contain toxins, such as the glycoside amygdalin found in almonds (79), bees might potentially consume harmful concentrations of such substances. 248

249 More generally it seems certain that bees inhabiting intensive farmland have a more 250 monotonous diet than they would have experienced in their evolutionary past, but how this 251 impacts upon their fitness remains unclear. The pollen of different plant species varies greatly 252 in protein content, amino acid composition, lipid, starch, vitamin and mineral content. Nectar 253 commonly contains varying and low concentrations of a range of nutrients and other 254 compounds of largely unknown importance (80-84). Thus, we might expect the type and 255 range of flowers available will affect individual bee health and colony fitness in multiple 256 ways; for example in honey bees, both pollen quality and diversity influence longevity, 257 physiology and resistance/tolerance to disease (85–88). However, this topic has been little 258 investigated, particularly for wild bees. The perception that honey bees may be receiving an 259 inadequate diet has led to the development of protein supplements, but once again there has 260 been little research on the long-term effectiveness of such supplements on colony health (89, 261 90).

262 Interpreting the effects of availability of mass-flowering crops on bees and their colonies is further complicated since visiting such crops often exposes bees to pesticide 263 264 residues, so that positive effects of increased food availability may be offset by negative 265 effects of the pesticide. Some studies have found positive effects of proximity to canola on 266 bumblebee colony growth and abundance (91, 92) and on numbers of nesting red mason bees 267 Osmia bicornis (93, 94) while others found no relationship for bumblebees (58, 95) nor for 268 solitary bees (96). Interestingly, none of these studies considered what role pesticides might 269 have played in mediating the effect of the crop, or even report which pesticides were applied 270 to crops in the study area, an omission that now seems naive given the recent focus on 271 impacts of neonicotinoid insecticides on bees.

272

273 Shipping fever

274 It seems reasonable to hypothesize that the long-distance transport of bees, as routinely occurs for honey bees in North America and for commercial bumblebee colonies, places 275 276 stress on the colonies. For several days they may be confined and subject to vibration, high 277 temperatures, high levels of carbon dioxide and irregular disturbance. It has long been known 278 that such stress can activate bacterial and viral infections and generally reduce condition in 279 vertebrate livestock (97), but this has not been investigated in bees, although Bakonyi et al. 280 (98) suggest that shipping stress may have contributed to honey bee colony losses in 281 Hungary. This is clearly an area where further research is needed.

- 282
- 283

#### 284 <u>Competition</u>

285 The role of competition in determining the relative abundance of species is notoriously hard 286 to ascertain in mobile organisms such as bees, but it seems likely that competition for floral 287 resources and perhaps also for nest sites does occur in natural communities, and that it can be 288 exacerbated by the introduction of non-native species, particularly when the latter are present 289 at high densities (57). For example, there is evidence that high concentrations of domestic 290 honey bee hives can displace wild bumblebees from their preferred foodplants and from 291 whole areas if hive densities are sufficiently high (99, 100). This can result in a reduction in 292 the size of bumblebee workers (101), and reduced reproductive success of bumblebee 293 colonies (102). Although in general the interests of honey bee keepers and wild bee 294 conservationists are aligned (all would agree on the benefits of increasing floral resources, 295 reducing exposure to pesticides and preventing invasions of alien pathogens), there may 296 occasionally be conflict where bee keepers wish to place hives in areas with significant 297 populations of rare wild bees (57).

298

#### 299 Climate change

300 Climate change is widely accepted to pose one of the largest threats to biodiversity

- 301 worldwide, but likely impacts on pollinators and pollination are not well understood. One
- 302 danger is that the phenology of pollinators may diverge from that of the plants they pollinate,
- 303 with potentially disastrous consequences for both, but there is little evidence that this is
- happening to a significant extent yet (103). Advances in flowering and bee emergence are
- 305 often broadly similar, and in any case few plants are dependent on a single pollinator so that

any mismatch with one pollinator is likely to be compensated by increased availability ofanother (*103*, *104*).

Another potential effect of climate change is as a driver of range shifts, leading to a spatial mismatch between plants and pollinators. Range shifts in response to climate have been demonstrated in butterflies (105) and are to be expected in bees (13); for example there is already evidence that the lower altitudinal limit of some montane bumblebees has shifted uphill in Spain (106). We would predict declines in bumblebees at the southern edge of their range since they tend to be poorly adapted to high temperatures.

Of course climate change is not solely associated with warming; extreme weather events such as storms, floods and droughts are predicted to increase, and we would expect these to have major impacts on local bee communities. For example, flooding is likely to be harmful to the many bee species that nest or hibernate underground.

318 Overall, although there is little strong evidence that climate change has yet had any 319 great effect on bees, it is likely to provide a growing source of stress in the future that would 320 exacerbate the impact of other factors, such as habitat loss.

321

#### 322 Interactions between stressors

323 Unfortunately the public debate on bee health has often become polarized, with claims that, 324 for example, *Varroa* or neonicotinoid insecticides are the sole or primary cause of bee 325 declines or honey bee colony losses. If a middle-aged man who is overweight, does little exercise, and smokes and drinks heavily were to die of a heart attack, we would not be 326 327 surprised and we might not spend too long arguing over which single risk factor was most 328 important in bringing about his untimely demise. Similarly, wild bee declines and honey bee 329 colony losses are clearly due to multiple, interacting and sometimes synergistic factors, and 330 the combination of factors involved no doubt varies in time and space.

331 In general, the combined effect of multiple stressors are likely to be more harmful than 332 one stressor alone (107–109) (Figure 3). In the worst-case scenario, sublethal stressors that do 333 not incur any significant harmful effects in isolation could, in combination, result in lethal 334 effects. As we have already seen, bees are often exposed chronically to mixtures of pesticides 335 and other chemicals. Some, such as ergosterol biosynthesis inhibitors (EBI) fungicides, have 336 very low toxicity in themselves but may increase the toxicity of some neonicotinoids and pyrethroids up to 1,000-fold (110-112). Piperonyl butoxide is often added to pesticide 337 338 formulations and also acts synergistically with some neonicotinoids, increasing toxicity up to 339 244-fold (111). Intriguingly, whilst imidacloprid alone has been shown to impair olfactory 340 learning (113), combined exposure to imidacloprid and coumaphos has been shown to result 341 in a slight increase in learning in honey bees (114). So while regulatory processes examine 342 the effects on bees of exposure to a single pesticide at a time, in reality bees are 343 simultaneously exposed to many pesticides, some of which have combined effects that cannot 344 be predicted from studies of their effects when used in isolation.

345 Several recent studies indicate that interactive effects between pesticides and pathogens could be especially harmful for bees (115–121). For instance, developmental 346 347 exposure to neonicotinoid insecticides renders honey bees more susceptible to the impact of 348 the invasive pathogen N. ceranae (122). Imidacloprid can act synergistically with Nosema 349 spp. by increasing the prevalence of *Nosema* infections in hives (116) and increasing 350 Nosema-induced mortality (115). Similarly, Aufauvre et al. (118) showed that mortality of 351 honey bees was greater when bees were exposed to the insecticide fipronil and infected by N. 352 ceranae than when only a single stress factor was present. There is evidence that exposure to 353 pesticides may impair the immune function of insects, which would explain these effects (43, 354 123–125). For example, Di Prisco *et al.* (126) recently showed that exposure to 355 neonicotinoids (clothianidin or imidacloprid) leads to immunosuppression in honey bees, 356 which, in turn, promotes the replication of the deformed wing virus in insects with covert 357 infections. This effect was found at very low concentrations, well below those that bees are 358 likely to encounter in the field.

359 Interactions between stressors are not confined to pesticides and pathogens. The 360 ability of bees to survive parasite infections is compromised by nutritional stress. For 361 example, Crithidia bombi causes little mortality in well-fed bumblebees, but becomes 362 virulent in bumblebees with a restricted diet (127). Activating the immune response has a 363 metabolic cost; bumblebees increase their food consumption when immune responses 364 areupregulated (128), and artificially stimulating the immune response by injecting latex 365 beads caused mortality in starving bumblebees but not in those that were well fed (129). Increased food consumption in infected bees could also increase exposure to pesticides. 366 367 Activating immunity has been shown to impair learning in both honey bees (130) and bumblebees (131, 132), and impaired learning will reduce the bees' ability to locate floral 368 369 resources and extract rewards, so exacerbating nutritional stresses. 370 Although to our knowledge this has not yet been examined, it seems highly likely that

nutritional stress may also modulate the ability of bees to cope with pesticides, and this may

explain in part why the observed LD50 of toxins in bees is highly variable across studies(65).

374 In summary, stressors do not act in isolation. Bees of all species are likely to 375 encounter multiple stressors during their lives, and each is likely to reduce the ability of bees 376 to cope with the others. A bee or bee colony that appears to have succumbed to a pathogen 377 may not have died if it had not also been exposed to a sublethal dose of a pesticide and/or 378 been subject to food stress (which might in turn be due to drought or heavy rain induced by 379 climate change, or competition from a high density of honey bee hives placed nearby). 380 Unfortunately, conducting well-replicated studies of the effects of multiple, interacting 381 stressors on bee colonies is exceedingly difficult. The number of stressor combinations 382 rapidly become large, and exposure to stressors is hard or impossible to control with free-383 flying bees. Nonetheless, a strong argument can be made that it is the interaction between 384 parasites, pesticides and diet that lies at the heart of current bee health problems.

385

#### 386 Sustainable pollination into the future

387 There is universal agreement that we must ensure adequate pollinator populations into the 388 future if we wish to continue to grow a diversity of insect-pollinated crops and also ensure the 389 integrity of natural ecosystems. It is also clear that moving towards heavy reliance on a few 390 species of managed pollinators, such as honey bees or one or two species of bumblebee, runs 391 the risk of supply failure; for example, should honey bee stocks in North America fall much 392 further, the viability of almond production in California would be threatened (133). Wild 393 pollinators provide a service that is largely free, and globally already contributes the majority 394 of crop pollination (24, 25, 134, 135). Maintaining a diversity of pollinator species improves 395 crop success via functional complementarity; different species visit different parts of the crop 396 or crop plant, at different times of the day or year, and respond differently to environmental 397 perturbations (1, 136–141). A diversity of pollinators can buffer impacts of climate change 398 which might otherwise result in a mismatch in phenology of pollinators with the flowering of 399 crops (104). It is thus essential that we take steps to conserve a broad community of 400 pollinators in farmland.

Fortunately, although the causes of pollinator ill-health may be complex and multicausal, conserving pollinators need not be difficult or expensive. If we accept that declines are due to interacting stressors, then it follows logically that removing or reducing any of the

13

stressors we have described is likely to benefit bee populations. Measures can be taken that
are likely to simultaneously benefit a broad suit of species, both domesticated and wild:

406

407 1) Increase abundance, diversity and continuity of floral resources. Schemes such as the 408 sowing of flower-rich field margins or hedgerows, or retaining patches of semi-409 natural habitat among or near farmland provide clear benefits to bee diversity and 410 abundance (30, 141–148) (Fig. 4A). This in turn increases pollination to nearby crops and provides an economic incentive to farmers growing insect-pollinated crops (149). 411 412 Many countries also offer financial incentives to farmers for taking measures to boost 413 biodiversity that help to offset implementation and opportunity costs. However, take-414 up of schemes to boost pollinators remains low in most countries, perhaps reflecting a 415 lack of understanding of the economic and environmental benefits, or a lack of 416 familiarity with implementation of such measures. Education and outreach in this area 417 could pay great dividends for pollinator conservation.

418Planting of appropriate flowers in gardens and amenity areas can also419contribute to pollinator conservation (150, 151) (Fig. 4B). There is evidence that420urban areas can support higher populations of some pollinators than farmland e.g.,421(36) and boost bee numbers in adjacent farmland (58). Many lists of bee or wildlife-422friendly flowers are available on the internet but they tend to be based on anecdote423rather than evidence, and there is a need to develop regionally appropriate, evidence-424based advice as to the best plants to grow (152).

425

2) <u>Provide nest sites.</u> Wild bees use a diversity of habitats for nesting, including
burrowing into bare soil, using existing cavities underground, holes in wood, or
hollow plant stems. Semi-natural habitats, hedgerows and permanently uncropped
field margins cater for many of these, meaning that schemes to boost floral diversity
are also likely to boost nesting opportunities (*141*). Additional nest sites can also be
provided by providing bundles of hollow reeds or canes, or patches of bare soil (*153*).

432

3) <u>Reduce exposure to pesticides</u>. Bees are currently chronically exposed to a cocktail of
pesticides some of which act synergistically. Since the late 1990s, the cost of
pesticides has fallen markedly relative to labor and fuel costs and the value of the
crops (*154*). As a result, current levels of pesticide use are generally high, and not

437 always justified by evidence that they are necessary to maintain yield (66, 155). The 438 widespread, prophylactic use of systemic insecticides, such as neonicotinoids as seed 439 dressings, exposes bees and other non-target wildlife, results in accumulation of 440 pesticides in the environment, and places strong selection pressure on pests to evolve 441 resistance. A return to the principles of Integrated Pest Management (156), which 442 utilizes preventative methods, such as crop rotation, and views the use of pesticides as 443 a last resort in the battle against insect pests, could greatly reduce exposure of bees, benefit the environment, and improve farming profitability. Some European countries 444 445 have independently developed national pesticide reduction programs (156), and the 446 European Union Sustainable Use of Pesticides Directive (Anon 2009) required 447 member states to implement national action plans to minimize pesticide use by 448 January 2014. In most EU states this directive appears to have had little or no impact 449 on farming practices.

450 Current risk assessment procedures, which examine the short-term impact of a 451 single pesticide in isolation, are clearly not adequate to encapsulate the true scenario 452 faced by bees living in farmland. Improvements are needed to make them more 453 realistic, whilst keeping the cost of regulatory tests affordable, posing a considerable 454 challenge to the ingenuity of scientists and regulators.

455 The EU moratorium on the use of three neonicotinoids (which started in 456 December 2013) is an attempt to use policy change to reduce exposure of bees to 457 stressors, following a review by the European Food Standards Agency (157–159) 458 which declared that neonicotinoids pose an "unacceptable risk" to bees. However, if 459 this simply leads farmers to replace neonicotinoids with other pesticides this may not 460 be of great benefit to bees or the environment. Funding for research and for the 461 provision of clear, independent advice for farmers with regard to how to reduce 462 pesticide use generally by adopting IPM practices might provide a better and more 463 sustainable long-term solution.

464

4) <u>Prevent further introductions of non-native bees, parasites and pathogens.</u> The
careless disregard with which we ship bees from country to country has resulted in the
irreversible spread of many serious parasites and pathogens. Strict quarantine controls
should be implemented on the movement of all commercial bees, and there is an
urgent need to develop means of rearing commercial bumblebees that are free from

470 471

472

473

474

disease. Deliberate introductions of non-native bee species (such as the recent introduction of the European *Bombus terrestris* to South America) should of course be prevented. The companies that rear commercial bees should bear some responsibility here, and refuse to sell bees to regions where they are not native. There is clear hypocrisy in the policies of countries that prevent importation of non-native species but allow exportation of species to places where they do not naturally occur.

475 476

477 5) <u>Develop monitoring programs.</u> We have good distribution maps for pollinators in 478 some countries, particularly for bumblebees, and citizen science schemes such as 479 "Bumble Bee Watch" in North America and "Beewatch" in the UK can help to track 480 changes in these distributions. However, the lack of long-term data on pollinator 481 abundance is a glaring knowledge gap that urgently needs to be filled. It will probably never be possible to monitor all pollinator species at a global scale, but it would be 482 483 practical to systematically collect data on the abundance of a subset of the more 484 abundant and economically important pollinators. Citizen science surveys can provide 485 a cost effective means for large-scale population monitoring; for example the UK 486 butterfly monitoring scheme employs volunteers to walk regular transects using a 487 standard methodology to count the butterflies seen, and has generated a large and 488 long-term data set which has provided powerful insights into insect population change 489 (160, 161). "Beewalks", a similar scheme to count bumblebees, has been launched by 490 the Bumblebee Conservation Trust to obtain population data for bumblebees in the UK, although it is still in its infancy. In the USA, "The Great Sunflower Project" asks 491 492 volunteers to count pollinators in flower patches in their local area. However, such 493 schemes are limited by the taxonomic skills of volunteers, particularly for the many 494 pollinator taxa that are hard or impossible to identify in the field. LeBuhn et al. (162) 495 argue that a pan-tapping network, which could employ citizen scientists to place out 496 the traps but experts to identify the catch, would be the most cost-effective means for 497 monitoring a large cross-section of pollinator species on a large geographic scale. 498 Indeed, for a relatively modest sum it would be possible to set up an international pan-499 trapping network to monitor pollinators following a standard methodology. Until 500 good population data become available, we cannot identify species or regions under 501 most threat and hence we cannot prioritize management.

502	In the absence of pollinator monitoring, we have no early warning system to tell us how close		
503	we may be to a pollination crisis. With a growing human population and rapid growth in		
504	global demand for pollination services, we cannot afford to see crop yields begin to fall, and		
505	we would be well advised to take pre-emptive action to ensure that we have adequate		
506 507	pollination services into the future.		
508	Ackı	nowledgements	
509	Thanks are due to Rich Hatfield and Sarina Jepsen for their help with Figure 1b.		
510 511	1.	AM. Klein, I. Steffan-Dewenter, T. Tscharntke, Fruit set of highland coffee increases with the diversity of pollinating bees. <i>Proc. Biol. Sci.</i> <b>270</b> , 955–961 (2003).	
512 513 514	2.	N. Gallai, JM. Salles, J. Settele, B. E. Vaissière, Economic valuation of the vulnerability of world agriculture confronted with pollinator decline. <i>Ecol. Econ.</i> <b>68</b> , 810–821 (2009).	
515 516	3.	C. Holden, Report warns of looming pollination crisis in North America. <i>Science</i> . <b>314</b> , 397 (2006).	
517	4.	M. Gross, Bee gloom deepens. Curr. Biol. 18, 1073 (2008).	
518 519	5.	S. G. Potts <i>et al.</i> , Declines of managed honey bees and beekeepers in Europe. <i>J. Apic. Res.</i> , 15–22 (2010).	
520 521	6.	National Resource Council, <i>Status of Pollinators in North America</i> (National Academies Press, 2007).	
522 523	7.	D. van Engelsdorp, J. Hayes, R. M. Underwood, J. Pettis, A survey of honey bee colony losses in the U.S., fall 2007 to spring 2008. <i>PLoS One</i> . <b>3</b> , e4071 (2008).	
524 525	8.	M. A. Aizen, L. D. Harder, The global stock of domesticated honey bees is growing slower than agricultural demand for pollination. <i>Curr. Biol.</i> <b>19</b> , 915–918 (2009).	
526 527	9.	D. vanEngelsdorp <i>et al.</i> , Colony collapse disorder: A descriptive study. <i>PLoS One.</i> <b>4</b> , e6481 (2009).	
528 529	10.	K. M. Smith <i>et al.</i> , Pathogens, pests, and economics: drivers of honey bee colony declines and losses. <i>Ecohealth</i> . <b>10</b> , 434–45 (2013).	
530 531	11.	D. Goulson, G. C. Lye, B. Darvill, Decline and conservation of bumble bees. <i>Annu. Rev. Entomol.</i> <b>53</b> , 191–208 (2008).	
532 533	12.	A. Kosior <i>et al.</i> , The decline of the bumble bees and cuckoo bees (Hymenoptera: Apidae: Bombini) of Western and Central Europe. <i>Oryx.</i> <b>41</b> , 79–88 (2007).	

- 534 13. P. H. Williams, J. L. Osborne, Bumblebee vulnerability and conservation world-wide.
   535 *Apidologie*. 40, 367–387 (2009).
- 536 14. P. H. Williams, R. W. Thorp, L. L. Richardson, S. R. Colla, *Bumble Bees of North*537 *America: An Identification Guide* (Princeton University Press, 2014).
- J. C. Grixti, L. T. Wong, S. A. Cameron, C. Favret, Decline of bumble bees (*Bombus*)
  in the North American Midwest. *Biol. Conserv.* 142, 75–84 (2009).
- R. Schmid-Hempel *et al.*, The invasion of southern South America by imported
  bumblebees and associated parasites. *J. Anim. Ecol.* 83, 823–837 (2014).
- 542 17. Z. Xie, P. H. Williams, Y. Tang, The effect of grazing on bumblebees in the high
  543 rangelands of the eastern Tibetan Plateau of Sichuan. J. Insect Conserv. 12, 695–703
  544 (2008).
- 545 18. P. Williams, Y. Tang, J. Yao, S. Cameron, The bumblebees of Sichuan (Hymenoptera:
  546 Apidae, Bombini). *Syst. Biodivers.* 7, 101–189 (2009).
- 547 19. C. Matsumura, J. Yokoyama, I. Washitani, Invasion status and potential ecological
  548 impacts of an invasive alien bumblebee, *Bombus terrestris* L. (Hymenoptera: Apidae
  549 ) naturalized in Southern Hokkaido, Japan. *Glob. Environ. Res.* 8, 51 66 (2004).
- M. N. Inoue, J. Yokoyama, I. Washitani, Displacement of Japanese native bumblebees
  by the recently introduced *Bombus terrestris* (L.) (Hymenoptera: Apidae). *J. Insect Conserv.* 12, 135–146 (2008).
- L. G. Carvalheiro *et al.*, Species richness declines and biotic homogenisation have
  slowed down for NW-European pollinators and plants. *Ecol. Lett.* 16, 870–878 (2013).
- J. C. Biesmeijer *et al.*, Parallel declines in pollinators and insect pollinated plants in
  Britain and the Netherlands. *Science*. **313**, 351–354 (2006).
- L. A. Burkle, J. C. Marlin, T. M. Knight, Plant-pollinator interactions over 120 years:
  loss of species, co-occurrence, and function. *Science*. 339, 1611–1615 (2013).
- 559 24. T. D. Breeze, a. P. Bailey, K. G. Balcombe, S. G. Potts, Pollination services in the UK:
  560 How important are honeybees? *Agric. Ecosyst. Environ.* 142, 137–143 (2011).
- 561 25. L. A. Garibaldi *et al.*, Wild pollinators enhance fruit set of crops regardless of honey
  562 bee abundance. *Science*. 339, 1608–1611 (2013).
- R. E. Mallinger, C. Gratton, J. Appl. Ecol., in press, doi:10.1111/1365-2664.12377. IS
  THIS NOW PUBLISHED? PLEASE ADD DETAILS CANNOT BE INCLUDED IF
  NOT PUBLISHED It is available online, but there are no page numbers yet:
  http://onlinelibrary.wiley.com/doi/10.1111/1365-2664.12377/abstract. Is this
  acceptable?
- 568 27. C. A. Kearns, D. W. Inouye, N. M. Waser, Endangered mutalisms: The conservation
  569 of plant-pollinator interactions. *Annu. Rev. Ecol. Syst.* 29, 83–112 (1998).

M. A. Aizen, L. A. Garibaldi, S. A. Cunningham, A. M. Klein, Long-term global 570 28. 571 trends in crop yield and production reveal no current pollination shortage but increasing pollinator dependency. Curr. Biol. 18, 1572–1575 (2008). 572 L. A. Garibaldi, M. A. Aizen, A. M. Klein, S. A. Cunningham, L. D. Harder, Global 573 29. growth and stability of agricultural yield decrease with pollinator dependence. Proc. 574 575 Natl. Acad. Sci. U. S. A. 108, 5909-5914 (2011). 576 30. L. A. Garibaldi et al., Stability of pollination services decreases with isolation from 577 natural areas despite honey bee visits. Ecol. Lett. 14, 1062–1072 (2011). M. J. F. Brown, R. J. Paxton, The conservation of bees: a global perspective. 578 31. 579 Apidologie. 40, 410–416 (2009). 580 32. S. G. Potts et al., Global pollinator declines: trends, impacts and drivers. Trends Ecol. 581 *Evol.* **25**, 345–53 (2010). 582 A. J. Vanbergen, Threats to an ecosystem service: pressures on pollinators. Front. 33. 583 Ecol. Environ., 251–259 (2013). 584 34. D. C. Howard, J. W. Watkins, R. T. Clarke, C. L. Barnett, G. J. Stark, Estimating the extent and change in broad habitats in Great Britain. J. Environ. Manage. 67, 219-227 585 586 (2003). 587 F. Samson, F. Knopf, Prairie conservation in North America. Bioscience. 44, 418-421 35. 588 (1994). 589 36. J. L. Osborne et al., Quantifying and comparing bumblebee nest densities in gardens 590 and countryside habitats. J. Appl. Ecol. 45, 784-792 (2008). 591 37. A. J. Bates et al., Changing bee and hoverfly pollinator assemblages along an urbanrural gradient. PLoS One. 6, e23459 (2011). 592 593 38. P. Skórka, M. Lenda, D. Moroń, K. Kalarus, P. Tryjanowski, Factors affecting road 594 mortality and the suitability of road verges for butterflies. Biol. Conserv. 159, 148-157 595 (2013). 596 39. E. Genersch, American Foulbrood in honeybees and its causative agent, Paenibacillus 597 larvae. J. Invertebr. Pathol. 103, S10-S19 (2010). 598 40. E. Genersch, C. Yue, I. Fries, J. R. De Miranda, Detection of *Deformed wing virus*, a 599 honey bee viral pathogen, in bumble bees (Bombus terrestris and Bombus pascuorum) 600 with wing deformities. J. Invertebr. Pathol. 91, 61–63 (2006). P. Graystock, K. Yates, B. Darvill, D. Goulson, W. O. H. Hughes, Emerging dangers: 601 41. 602 deadly effects of an emergent parasite in a new pollinator host. J. Invertebr. Pathol. 603 **114**, 114–119 (2013). P. Rosenkranz, P. Aumeier, B. Ziegelmann, Biology and control of Varroa destructor. 604 42. 605 J. Invertebr. Pathol. 103, S96-S119 (2010).

- F. Nazzi *et al.*, Synergistic parasite-pathogen interactions mediated by host immunity
  can drive the collapse of honeybee colonies. *PLoS Pathog.* 8, e1002735 (2012).
- 44. J. Klee *et al.*, Widespread dispersal of the microsporidian *Nosema ceranae*, an
  emergent pathogen of the western honey bee, *Apis mellifera*. J. Invertebr. Pathol. 96,
  1-10 (2007).
- 45. Y. Chen, J. D. Evans, I. B. Smith, J. S. Pettis, *Nosema ceranae* is a long-present and
  wide-spread microsporidian infection of the European honey bee (*Apis mellifera*) in
  the United States. *J. Invertebr. Pathol.* 97, 186–8 (2008).
- 46. J. Li *et al.*, Diversity of *Nosema* associated with bumblebees (Bombus spp.) from
  China. *Int. J. Parasitol.* 42, 49–61 (2012).
- 616 47. S. Plischuk *et al.*, South American native bumblebees (Hymenoptera: Apidae) infected
  617 by *Nosema ceranae* (Microsporidia), an emerging pathogen of honeybees (*Apis*618 *mellifera*). *Environ. Microbiol. Rep.* 1, 131–5 (2009).
- 48. J. Ravoet *et al.*, Widespread occurrence of honey bee pathogens in solitary bees. *J. Invertebr. Pathol.* 122, 55–8 (2014).
- 49. S. Spiewok, P. Neumann, Infestation of commercial bumblebee (*Bombus impatiens*)
  field colonies by small hive beetles (*Aethina tumida*). *Ecol. Entomol.* 31, 623–628
  (2006).
- 50. D. Hoffmann, J. S. Pettis, P. Neumann, Potential host shift of the small hive beetle
  (*Aethina tumida*) to bumblebee colonies (*Bombus impatiens*). *Insectes Soc.* 55, 153–
  162 (2008).
- 51. S. A. Cameron *et al.*, Patterns of widespread decline in North American bumble bees. *Proc. Natl. Acad. Sci. U. S. A.* 108, 662–667 (2011).
- 52. I. Meeus, M. J. F. Brown, D. C. De Graaf, G. Smagghe, Effects of invasive parasites
  on bumble bee declines. *Conserv. Biol.* 25, 662–671 (2011).
- 631 53. M. P. Arbetman, I. Meeus, C. L. Morales, M. A. Aizen, G. Smagghe, Alien parasite
  632 hitchhikes to Patagonia on invasive bumblebee. *Biol. Invasions.* 15, 489–494 (2013).
- 54. S. R. Colla, M. C. Otterstatter, R. J. Gegear, J. D. Thomson, Plight of the bumble bee:
  Pathogen spillover from commercial to wild populations. *Biol. Conserv.* 129, 461–467
  (2006).
- 636 55. M. C. Otterstatter, J. D. Thomson, Does pathogen spillover from commercially reared
  637 bumble bees threaten wild pollinators? *PLoS One.* 3, DOI:
  638 10.1371/journal.pone.0002771 (2008).
- 639 56. M. Y. Yoneda, H. F. Uruta, K. T. Suchida, K. O. Kabe, K. G. Oka, Commercial
  640 colonies of *Bombus terrestris* (Hymenopter: Apidae) are reservoirs of the tracheal mite
  641 Locustacarus buchneri (Acari: Podapolipidae). *Appl. Entomol. Zool.* 43, 73–76 (2008).

- 57. D. Goulson, Effects of introduced bees on native ecosystems. *Annu. Rev. Ecol. Evol. Syst.* 34, 1–26 (2003).
- 58. D. Goulson *et al.*, Effects of land use at a landscape scale on bumblebee nest density
  and survival. *J. Appl. Ecol.* 47, 1207–1215 (2010).
- 59. L. A. Morandin, M. L. Winston, Wild bee abundance and seed production in
  conventional, organic, and genetically modified canola. *Ecol. Appl.* 15, 871–881
  (2005).
- 649 60. A. M. Chauzat *et al.*, A survey of pesticide residues in pollen loads collected by honey
  650 bees in France. *J. Econ. Entomol.* 99, 253–262 (2006).
- 651 61. F. Sanchez-Bayo, K. Goka, Pesticide residues and bees--a risk assessment. *PLoS One*.
  652 9, e94482–e94482 (2014).
- 653 62. C. A. Mullin *et al.*, High levels of miticides and agrochemicals in North American 654 apiaries: implications for honey bee health. *PLoS One*. **5**, e9754–e9754 (2010).
- 655 63. C. H. Krupke, G. J. Hunt, B. D. Eitzer, G. Andino, K. Given, Multiple routes of
  656 pesticide exposure for honey bees living near agricultural fields. *PLoS One*. 7,
  657 e29268–e29268 (2012).
- 64. D. Paradis, G. Bérail, J. M. Bonmatin, L. P. Belzunces, Sensitive analytical methods
  for 22 relevant insecticides of 3 chemical families in honey by GC-MS/MS and LCMS/MS. Anal. Bioanal. Chem. 406, 621–633 (2014).
- 661 65. L. W. Pisa *et al.*, Effects of neonicotinoids and fipronil on non-target invertebrates.
  662 *Environ. Sci. Pollut. Res. Int.*, 1–35 (2014).
- 663 66. D. Goulson, An overview of the environmental risks posed by neonicotinoid
  664 insecticides. J. Appl. Ecol. 50, 977–987 (2013).
- 665 67. M. Tomizawa, J. E. Casida, Neonicotinoid insecticide toxicology: mechanisms of
  666 selective action. *Annu. Rev. Pharmacol. Toxicol.* 45, 247–268 (2005).
- 667 68. S. Suchail, D. Guez, L. P. Belzunces, Characteristics of imidacloprid toxicity in two
  668 *Apis mellifera* subspecies. *Environ. Toxicol. Chem.* 19, 1901–1905 (2000).
- 669 69. G. Rondeau *et al.*, Delayed and time-cumulative toxicity of imidacloprid in bees, ants
  670 and termites. *Sci. Rep.* 4, 5566 (2014).
- 671 70. E. C. Yang, Y. C. Chuang, Y. L. Chen, L. H. Chang, A. E. C. Yang, Abnormal
  672 foraging behavior induced by sublethal dosage of imidacloprid in the honey bee
  673 (Hymenoptera: Apidae). *J. Econ. Entomol.* 101, 1743–1748 (2008).
- V. Mommaerts *et al.*, Risk assessment for side-effects of neonicotinoids against
  bumblebees with and without impairing foraging behavior. *Ecotoxicology*. 19, 207–
  215 (2010).

- M. Henry *et al.*, A common pesticide decreases foraging success and survival in honey *Science.* 336, 348–350 (2012).
- H. Feltham, K. Park, D. Goulson, Field realistic doses of pesticide imidacloprid reduce
  bumblebee pollen foraging efficiency. *Ecotoxicology*. 23, 317–323 (2014).
- 74. P. Han, C. Y. Niu, C. L. Lei, J. J. Cui, N. Desneux, Quantification of toxins in a
  682 Cry1Ac + CpTI cotton cultivar and its potential effects on the honey bee *Apis mellifera*683 L. *Ecotoxicology*. 19, 1452–1459 (2010).
- E. C. Yang, H. C. Chang, W. Y. Wu, Y. W. Chen, Impaired olfactory associative
  behavior of honeybee workers due to contamination of imidacloprid in the larval stage. *PLoS One.* 7, e49472 (2012).
- 687 76. P. R. Whitehorn, S. O'Connor, F. L. Wackers, D. Goulson, Neonicotinoid pesticide
  688 reduces bumble bee colony growth and queen production. *Science*. 336, 351–352
  689 (2012).
- 690 77. G. C. Cutler, C. D. Scott-dupree, Exposure to clothianidin seed-treated canola has no
  691 long-term impact on honey bees. *J. Econ. Entomol.* 100, 765–772 (2007).
- 692 78. E. Pilling, P. Campbell, M. Coulson, N. Ruddle, I. Tornier, A four-year field program investigating long-term effects of repeated exposure of honey bee colonies to
  694 flowering crops treated with thiamethoxam. *PLoS One.* 8, e77193–e77193 (2013).
- 695 79. I. London-Shafir, S. Shafir, D. Eisikowitch, Amygdalin in almond nectar and pollen696 facts and possible roles. *Plant Syst. Evol.* 238, 87–95 (2003).
- 80. R. G. Stanley, H. F. Linskens, *Pollen: Biology, Biochemistry, Management* (SpringerVerlag, Berlin, 1974).
- T. H. Roulston, S. L. Buchmann, A phylogenetic reconsideration of the pollen starchpollination correlation. *Evol. Ecol. Res.* 2, 627–643 (2000).
- 82. M. E. Hanley, M. Franco, S. Pichon, B. Darvill, D. Goulson, Breeding system,
  pollinator choice and variation in pollen quality in British herbaceous plants. *Funct. Ecol.* 22, 592–598 (2008).
- 704 83. G. A. Wright *et al.*, Caffeine in floral nectar enhances a pollinator's memory of
  705 reward. *Science*. 339, 1202–4 (2013).
- 84. W. Mao, M. a Schuler, M. R. Berenbaum, Honey constituents up-regulate
  detoxification and immunity genes in the western honey bee *Apis mellifera*. *Proc. Natl. Acad. Sci. U. S. A.* **110**, 8842–6 (2013).
- A. Maurizio, The influence of pollen feeding and brood rearing on the length of life
  and physiological condition of the honeybee. *Bee World.* 31, 9–12 (1950).

711 86. L. N. Standifer, A comparison of the protein quality of pollens for growth-stimulation 712 of the hypopharyngeal glands and longevity of honey bees, Apis mellifera L. 713 (Hymenoptera: Apidae). Insectes Soc. 14, 415–425 (1967). 714 87. L. S. Schmidt, J. O. Schmidt, H. Rao, W. Wang, L. Xu, Feeding preference and survival of young worker honeybees (Hymenoptera: Apidae) fed rape, sesame, and 715 716 sunflower pollen. J. Econ. Entomol. 88, 1591–1595 (1995). 717 88. G. Di Pasquale et al., Influence of pollen nutrition on honey bee health: do pollen 718 quality and diversity matter? PLoS One. 8, e72016 (2013). 719 89. M. M. Morais et al., Evaluation of inexpensive pollen substitute diets through 720 quantification of haemolymph proteins. J. Apic. Res. 52, 119 (2013) 721 90. R. Kumar, R. C. Mishra, O. P. Agrawal, Effect of feeding artificial diets to honey bees 722 during dearth period under Panchkula (Haryana) conditions. J. Entomol. Res. 37, 41-723 46 (2013). 724 91. M. E. Knight et al., Bumblebee nest density and the scale of available forage in arable 725 landscapes. Insect Conserv. Divers. 2, 116–124 (2009). 726 92. C. Westphal, I. Steffan-Dewenter, T. Tscharntke, Mass flowering oilseed rape 727 improves early colony growth but not sexual reproduction of bumblebees. J. Appl. 728 Ecol. 46, 187–193 (2009). 729 93. F. Jauker, F. Peter, V. Wolters, T. Diekötter, Early reproductive benefits of mass-730 flowering crops to the solitary bee Osmia rufa outbalance post-flowering 731 disadvantages. Basic Appl. Ecol. 13, 268–276 (2012). 732 94. A. Holzschuh, C. F. Dormann, T. Tscharntke, I. Steffan-Dewenter, Mass-flowering 733 crops enhance wild bee abundance. Oecologia. 172, 477-484 (2013). 734 95. F. Herrmann, C. Westphal, R. F. A. Moritz, I. Steffan-Dewenter, Genetic diversity and 735 mass resources promote colony size and forager densities of a social bee (Bombus pascuorum) in agricultural landscapes. Mol. Ecol. 16, 1167-1178 (2007). 736 737 96. A. Kovács-Hostyánszki et al., Contrasting effects of mass-flowering crops on bee 738 pollination of hedge plants at different spatial and temporal scales. Ecol. Appl. 23, 739 1938-1946 (2013). 740 W. D. Yates, A review of infectious bovine rhinotracheitis, shipping fever pneumonia 97. 741 and viral-bacterial synergism in respiratory disease of cattle. Can. J. Comp. Med. 46, 742 225-263 (1982). 743 98. T. Bakonyi, R. Farkas, A. Szendroi, M. Dobos Kovacs, M. Rusva1, Detection of acute 744 bee paralysis virus by RT-PCR in honey bee and Varroa destructor field samples: 745 rapid screening of representative Hungarian apiaries. Apidologie. 33, 63–74 (2002). M. L. Forup, J. Memmott, The relationship between the abundance of bumblebees and 746 99. 747 honeybees in a native habitat. Ecol. Entomol. 30, 47–57 (2005).

- K. Walther-Hellwig *et al.*, Increased density of honeybee colonies affects foraging
  bumblebees. *Apidologie*. **37**, 517–532 (2006).
- D. Goulson, K. R. Sparrow, Evidence for competition between honeybees and
  bumblebees; effects on bumblebee worker size. *J. Insect Conserv.* 13, 177–181 (2008).
- D. M. Thomson, Detecting the effects of introduced species: A case study of
  competition between *Apis* and *Bombus*. *Oikos*. **114**, 407–418 (2006).
- P. Willmer, Pollinator-plant synchrony tested by climate change. *Curr. Biol.* 22, R131–R132 (2012).
- I. Bartomeus *et al.*, Biodiversity ensures plant-pollinator phenological synchrony against climate change. *Ecol. Lett.* 16, 1331–1338 (2013).
- M. L. Forister *et al.*, Compounded effects of climate change and habitat alteration shift
  patterns of butterfly diversity. *Proc. Natl. Acad. Sci. U. S. A.* **107**, 2088–2092 (2010).
- E. F. Ploquin, J. M. Herrera, J. R. Obeso, Bumblebee community homogenization after
  uphill shifts in montane areas of northern Spain. *Oecologia*. 173, 1649–1660 (2013).
- A. Sih, A. M. Bell, J. L. Kerby, Two stressors are far deadlier than one. *Trends Ecol. Evol.* 19, 274–276 (2004)
- A. Coors, L. De Meester, Synergistic, antagonistic and additive effects of multiple
  stressors: Predation threat, parasitism and pesticide exposure in *Daphnia magna*. J. *Appl. Ecol.* 45, 1820–1828 (2008).
- R. J. Gill, O. Ramos-Rodriguez, N. E. Raine, Combined pesticide exposure severely
   affects individual- and colony-level traits in bees. *Nature*. 491, 105–108 (2012).
- R. Schmuck, T. Stadler, H. W. Schmidt, Field relevance of a synergistic effect
  observed in the laboratory between an EBI fungicide and a chloronicotinyl insecticide
  in the honeybee (*Apis mellifera* L, Hymenoptera). *Pest Manag. Sci.* 59, 279–286
  (2003).
- T. Iwasa, N. Motoyama, J. T. Ambrose, R. M. M. Roe, Mechanism for the differential toxicity of neonicotinoid insecticides in the honey bee, *Apis mellifera*. *Crop Prot.* 23, 371–378 (2004).
- E. D. Pilling, P. C. Jepson, Synergism between EBI fungicides and a pyrethroid insecticide in the honeybee (*Apis mellifera*). *Pestic. Sci.* **39**, 293–297 (1993).
- A. Decourtye *et al.*, Imidacloprid impairs memory and brain metabolism in the honeybee (*Apis mellifera* L.). *Pestic. Biochem. Physiol.* **78**, 83–92 (2004).
- S. M. Williamson, G. A. Wright, Exposure to multiple cholinergic pesticides impairs olfactory learning and memory in honeybees. *J. Exp. Biol.* 216, 1799–1807 (2013).

- 782 115. C. Alaux *et al.*, Interactions between *Nosema* microspores and a neonicotinoid weaken
  783 honeybees (*Apis mellifera*). *Environ. Microbiol.* 12, 774–782 (2010).
- J. S. Pettis, D. VanEngelsdorp, J. Johnson, G. Dively, Pesticide exposure in honey bees
  results in increased levels of the gut pathogen *Nosema*. *Naturwissenschaften*. 99, 153–
  158 (2012).
- 117. C. Vidau *et al.*, Exposure to sublethal doses of fipronil and thiacloprid highly increases
   mortality of honeybees previously infected by *Nosema ceranae*. *PLoS One*. 6, e21550
   (2011).
- J. Aufauvre *et al.*, Parasite-insecticide interactions: a case study of Nosema ceranae
  and fipronil synergy on honeybee. *Sci. Rep.* 2, 326 (2012).
- G. L. Baron, N. E. Raine, M. J. F. Brown, Impact of chronic exposure to a pyrethroid pesticide on bumblebees and interactions with a trypanosome parasite. *J. Appl. Ecol.*51, 460–469 (2014).
- A. Fauser-Misslin, B. M. Sadd, P. Neumann, C. Sandrock, Influence of combined
  pesticide and parasite exposure on bumblebee colony traits in the laboratory. *J. Appl. Ecol.* 51, 450–459 (2014).
- V. Doublet, M. Labarussias, J. R. de Miranda, R. F. A. Moritz, R. J. Paxton, Bees
  under stress: sublethal doses of a neonicotinoid pesticide and pathogens interact to
  elevate honey bee mortality across the life cycle. *Environ. Microbiol.*, in press,
  doi:10.1111/1462-2920.12426. As above, it is now online at
  http://onlinelibrary.wiley.com/doi/10.1111/1462-2920.12426/abstract
- I 22. J. Y. Wu, M. D. Smart, C. M. Anelli, W. S. Sheppard, Honey bees (*Apis mellifera*)
  reared in brood combs containing high levels of pesticide residues exhibit increased
  susceptibility to *Nosema* (Microsporidia) infection. *J. Invertebr. Pathol.* 109, 326–329
  (2012).
- R. R. James, J. Xu, Mechanisms by which pesticides affect insect immunity. J. *Invertebr. Pathol.* 109, 175–182 (2012).
- F. S. Brunner, P. Schmid-Hempel, S. M. Barribeau, Immune gene expression in
   *Bombus terrestris*: signatures of infection despite strong variation among populations,
   colonies, and sister workers. *PLoS One.* 8, e68181 (2013).
- H. Boncristiani *et al.*, Direct effect of acaricides on pathogen loads and gene
  expression levels in honey bees *Apis mellifera*. J. Insect Physiol. 58, 613–620 (2012).
- 814 126. G. Di Prisco *et al.*, Neonicotinoid clothianidin adversely affects insect immunity and
  815 promotes replication of a viral pathogen in honey bees. *Proc. Natl. Acad. Sci. U. S. A.*816 110, 18466–18471 (2013).
- 817 127. M. J. F. Brown, R. Loosli, P. Schmid-Hempel, Condition-dependent expression of
  818 virulence in a trypanosome infecting bumblebees. *Oikos.* 91, 421–427 (2000).

819 128. E. R. Tyler, S. Adams, E. B. Mallon, An immune response in the bumblebee, *Bombus* 820 terrestris leads to increased food consumption. BMC Physiol. 6, 6 (2006). 821 129. Y. Moret, P. Schmid-Hempel, Survival for immunity: the price of immune system 822 activation for bumblebee workers. Science. 290, 1166–1168 (2000). 823 130. E. B. Mallon, A. Brockmann, P. Schmid-Hempel, Immune response inhibits 824 associative learning in insects. Proc. Biol. Sci. 270, 2471-2473 (2003). 825 131. C. E. Riddell, E. B. Mallon, Insect psychoneuroimmunology: Immune response reduces learning in protein starved bumblebees (Bombus terrestris). Brain. Behav. 826 827 *Immun.* **20**, 135–138 (2006). 828 132. A. Alghamdi, L. Dalton, A. Phillis, E. Rosato, E. B. Mallon, Immune response impairs 829 learning in free-flying bumble-bees. Biol. Lett. 4, 479–481 (2008). 830 D. Vanengelsdorp, M. D. Meixner, A historical review of managed honey bee 133. 831 populations in Europe and the United States and the factors that may affect them. J. Invertebr. Pathol. 103, S80–95 (2010). 832 833 134. R. Winfree, C. Kremen, Are ecosystem services stabilized by differences among species? A test using crop pollination. Proc. Biol. Sci. 276, 229-237 (2009). 834 835 R. Rader, B. G. Howlett, S. A. Cunningham, D. A. Westcott, W. Edwards, Spatial and 135. 836 temporal variation in pollinator effectiveness: Do unmanaged insects provide 837 consistent pollination services to mass flowering crops? J. Appl. Ecol. 49, 126-134 838 (2012). 839 136. P. Hoehn, T. Tscharntke, J. M. Tylianakis, I. Steffan-Dewenter, Functional group 840 diversity of bee pollinators increases crop yield. Proc. Biol. Sci. 275, 2283-2291 841 (2008). 842 137. J. K. Tuell, R. Isaacs, Community and species-specific responses of wild bees to insect 843 pest control programs applied to a pollinator-dependent crop. J. Econ. Entomol. 103, 844 668-675 (2010). 845 138. N. Blüthgen, A. M. Klein, Functional complementarity and specialisation: The role of biodiversity in plant-pollinator interactions. Basic Appl. Ecol. 12, 282-291 (2011). 846 847 139. C. Brittain, N. Williams, C. Kremen, A. Klein, Synergistic effects of non-Apis bees 848 and honey bees for pollination services. Proc. Biol. Sci. 280, 20122767 (2013). 849 140. I. Steffan-Dewenter, U. Münzenberg, C. Bürger, C. Thies, T. Tscharntke, Scaledependent effects of landscape context on three pollinator guilds. *Ecology*. 83, 1421– 850 1432 (2002). 851 852 141. L. A. Garibaldi et al., From research to action: enhancing crop yield through wild pollinators. Front. Ecol. Environ. 12, 439-447 (2014). 853

854 142. R. F. Pywell et al., Effectiveness of new agri-environment schemes in providing 855 foraging resources for bumblebees in intensively farmed landscapes. Biol. Conserv. 856 129, 192–206 (2006). R. F. Pywell et al., Providing foraging resources for bumblebees in intensively farmed 857 143. landscapes. Biol. Conserv. 121, 479-494 (2005). 858 859 144. C. Carvell, W. R. Meek, R. F. Pywell, D. Goulson, M. Nowakowski, Comparing the efficacy of agri-environment schemes to enhance bumble bee abundance and diversity 860 861 on arable field margins. J. Appl. Ecol. 44, 29-40 (2006). M. Rundlöf, H. Nilsson, H. G. Smith, Interacting effects of farming practice and 862 145. 863 landscape context on bumble bees. Biol. Conserv. 141, 417–426 (2008). 864 146. C. M. Kennedy et al., A global quantitative synthesis of local and landscape effects on wild bee pollinators in agroecosystems. Ecol. Lett. 16, 584-99 (2013). 865 S. D. Wratten, M. Gillespie, A. Decourtye, E. Mader, N. Desneux, Pollinator habitat 866 147. enhancement: Benefits to other ecosystem services. Agric. Ecosyst. Environ. 159, 112-867 122 (2012). 868 869 L. A. Morandin, M. L. Winston, V. A. Abbott, M. T. Franklin, Can pastureland 148. increase wild bee abundance in agriculturally intense areas? Basic Appl. Ecol. 8, 117-870 871 124 (2007). 872 B. R. Blaauw, R. Isaacs, Flower plantings increase wild bee abundance and the 149. 873 pollination services provided to a pollination-dependent crop. J. Appl. Ecol. 51, 890-874 898 (2014). 875 150. D. Goulson, Bumblebees: Behaviour, Ecology, and Conservation (Oxford University 876 Press, Oxford, 2010). 151. L. M. Blackmore, D. Goulson, Evaluating the effectiveness of wildflower seed mixes 877 878 for boosting floral diversity and bumblebee and hoverfly abundance in urban areas. 879 Insect Conserv. Divers. 7, 480–484 (2014). 880 152. M. Garbuzov, F. L. W. Ratnieks, Quantifying variation among garden plants in attractiveness to bees and other flower-visiting insects. Funct. Ecol. 28, 364-374 881 882 (2014). 883 153. I. Steffan-Dewenter, S. Schiele, Do resources or natural enemies drive bee population dynamics in fragmented habitats? Ecology. 89, 1375–1387 (2008). 884 154. C. D. Osteen, J. Fernandez-Cornejo, Economic and policy issues of U.S. agricultural 885 886 pesticide use trends. Pest Manag. Sci. 69, 1001-1025 (2013). 887 155. S. Stevens, P. Jenkins, Heavy costs; weighing the value of neonicotinoid insecticides in agriculture. Cent. Food Saf. Rep. (2014) This report has no Doi, it is just available 888 889 online.

- 890 156. B. Freier, E. F. Boller, *Integrated Pest Management in Europe History, Policy,*891 *Achievements and Implementation* In: R. Peshin & A.K. Dhawan, Integrated Pest
  892 management: Dissemination and Impact. Springer, p 435-444. (2009).
- Anon, Directive 2009/128/Ec of the European Parliament and of the Council of 21 October
  2009: establishing a framework for Community action to achieve the sustainable use of
  pesticides (2009).
- EFSA, Conclusion on the peer review of the pesticide risk assessment for bees for the active substance thiamethoxam. *EFSA J. 2013.* 11 (2013) (available at http://www.efsa.europa.eu/en/efsajournal/pub/3067.htm).
- 899 158. EFSA, Conclusion on the peer review of the pesticide risk assessment for bees for the active substance clothianidin. *EFSA J. 2013.* 11 (2013) (available at http://www.efsa.europa.eu/en/efsajournal/pub/3066.htm).
- 902 159. EFSA, Conclusion on the peer review of the pesticide risk assessment for bees for the active substance imidacloprid. *EFSA J. 2013.* 11 (2013) (available at http://www.efsa.europa.eu/en/efsajournal/pub/3068.htm).
- M. S. Warren *et al.*, Rapid responses of British butterflies to opposing forces of
  climate and habitat change. *Nature*. 414, 65–69 (2001).
- R. M. Pateman, J. K. Hill, D. B. Roy, R. Fox, C. D. Thomas, Temperature-dependent alterations in host use drive rapid range expansion in a butterfly. *Science*. 336, 1028–30 (2012).
- 910 162. G. Lebuhn *et al.*, Detecting insect pollinator declines on regional and global scales.
  911 *Conserv. Biol.* 27, 113–20 (2013).
- I. Fries, F. Feng, A. da Silva, S. B. Slemenda, N. J. Pieniazek, *Nosema ceranae* n. sp.
  (Microspora, Nosematidae), morphological and molecular characterization of a
  microsporidian parasite of the Asian honey bee *Apis cerana* (Hymenoptera, Apidae). *Eur. J. Protistol.* 32, 356–365 (1996).
- 916 164. M. Higes, R. Martín, A. Meana, *Nosema ceranae*, a new microsporidian parasite in honeybees in Europe. *J. Invertebr. Pathol.* 92, 93–5 (2006).
- 918 165. C. M. Aurori, D. S. Dezmirean, L. A. Mărghitaş, R. F. A. Moritz, *Nosema apis* and *N*.
  919 *ceranae* in Western Honeybee (*Apis mellifera*) Geographical Distribution and
  920 Current Methods of Diagnosis. *Bull. UASVM Anim. Sci. Biotechnol.* 68, 63–70 (2011).
- 921 166. J. Roberts, D. Anderson, Establishing the disease status of the Asian honeybee in the
  922 Cairns region Rural Industries Research and Development Corporation. *RIRDC Publ.*923 No. 13/082, *RIRDC Proj. No. PRJ-008433. Aust. Gov.* (2013).
- 924 167. J. Martínez, G. Leal, P. Conget, *Nosema ceranae* an emergent pathogen of *Apis mellifera* in Chile. *Parasitol. Res.* 111, 601–7 (2012).

- 926 168. Y. Chen *et al.*, Asymmetrical coexistence of *Nosema ceranae* and *Nosema apis* in honey bees. *J. Invertebr. Pathol.* 101, 204–9 (2009).
- 928 169. V. Chaimanee, N. Warrit, P. Chantawannakul, Infections of *Nosema ceranae* in four different honeybee species. *J. Invertebr. Pathol.* 105, 207–10 (2010).
- 930 170. C. Botías *et al.*, Further evidence of an oriental origin for *Nosema ceranae*931 (Microsporidia: Nosematidae). *J. Invertebr. Pathol.* 110, 108–13 (2012).
- 932 Figures:
- 933 Figure 1. Some wild bee species have undergone major range contractions. Shown here
- 934 for A the bumblebee *Bombus distinguendus* in the UK (data from the National Biodiversity
- 935 Network, UK, <u>https://data.nbn.org.uk/</u> and (B) Bombus affinis in North America (Map
- 936 produced by the Xerces Society, list of data providers can be found at:
- 937 <u>http://www.leifrichardson.org/bbna.html</u>). Photo credit: Dave Goulson, Johanna James-
- 938 Heinz.
- 939

#### 940 Figure 2. World distribution of the microsporidian pathogen Nosema ceranae in

- 941 **different bee hosts.** This parasite was first isolated from the Eastern honeybee (*A. cerana*)
- 942 collected in China in 1996 (163), and was subsequently found infecting Western honey bees
- 943 (A. mellifera) in Europe in 2005 (164). Soon after N.ceranae was detected in A. mellifera in
- 944 many regions of the world including Africa, Asia, the Americas and Oceania (44, 45, 165–
- 168), and more recently in other bee species including several Asian Apis species (169, 170)
- and wild bumblebees species from Europe, China and South America (41, 46, 47). N.
- 947 *ceranae* has also now been detected in solitary bees from Europe (48), confirming a very
- 948 wide range of hosts and high dispersal rate. Although the origins and primary host of *N*.
- 949 *ceranae* are yet to be accurately established, the apparent late and gradual invasions of N.
- 950 *ceranae* into different *A. mellifera* populations have led some authors to suggest that *A.*
- 951 *cerana* may be the primary host of *N. ceranae* and that it may have only recently emerged as
- a parasite of Western bees (170). The mechanism by which *N. ceranae* broadened its host
- range from an Asian bee species to other bee species across the world is unknown, but there
- has been human-mediated contact between Asian and Western bees for at least a century. It
- should be noted that regions in the figure where *Nosema* appears to be absent (white) may be
- 956 due to a lack of sampling in these areas.
- 957

#### 958 Figure 3. Both wild and managed bees are subject to a number of significant and

959 interacting stressors. For example, exposure to some fungicides can greatly increase toxicity

- 960 of insecticides (110–112), whereas exposure to insecticides reduces resistance to diseases
- 961 (115–123, 125, 126). Dietary stresses are likely to reduce the ability of bees to cope with both
- 962 toxins and pathogens (127–129). Photo credit: Beth Nicholls; Flickr Commons, AJC1
- 963 (https://creativecommons.org/licenses/by-nc-sa/2.0/legalcode\_)
- 964

#### 965 Figure 4. Increasing floral abundance in the landscape is very likely to benefit

- 966 **pollinator populations**. A Schemes to boost flower abundance in farmland, such as this
- 967 wildflower strip along a field margin, have been demonstrated to provide clear benefits for
- 968 wild bee populations (e.g. *140–145*); B Urban areas can support high populations of
- 969 pollinators, which may spill over into neighboring farmland. Conversion of amenity
- 970 grasslands in urban areas to wildflower patches has been shown to greatly boost numbers of
- 971 wild pollinators (151).
- 972