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14 15	Acoustic communication within ant societies and its mimicry by mutualistic and socially parasitic myrmecophiles
16 17	K Schönrogge ^{1*} F. Barbero ² J. P. Casacci ² J.Settele ³ JA Thomas ⁴
18	
19	
20	
21	
22 23	1. Centre for Ecology & Hydrology, MacLean Building, Benson Lane, Wallingford, OX10 8BB, UK
24 25 26	2. Department of Life Sciences and Systems Biology, University of Turin, Via Academia Albertina, 10123, Turin, Italy
27 28 29	3. UFZ, Helmholtz Centre for Environmental Research, Department of Community Ecology, Theodor-Lieser-Str. 4, 06120 Halle, Germany
30 31 32 33 34 35 36 37 38 39 40 41	4. Department of Zoology, University of Oxford, Oxford, South Parks Rd, OX1 3PS, United Kingdom
41 42	* Corresponding author:
43	K. Schönrogge (<u>ksc@ceh.ac.uk</u>), Tel: +44 (0)1491 838800, Fax: +44 (0)1491 692424
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49 Abstract:

50 This review focusses on the main acoustic adaptations that have evolved to enhance social communication in ants. We also describe how other invertebrates mimic these acoustic 51 52 signals in order to coexist with ants in the case of mutualistic myrmecophiles, or, in the case of social parasites, corrupt them in order to infiltrate ant societies and exploit their resources. 53 New data suggest that the strength of each ant-myrmecophile interaction leads to distinctive 54 sound profiles and may be a better predictor of the similarity of sound between different 55 myrmecophilous species than their phylogenetic distance. Finally, we discuss the 56 evolutionary significance of vibrations emitted by specialised myrmecophiles in the context of 57 ant multimodal communication involving the use of chemical and acoustic signals in 58 59 combination and identify future challenges for research including how new technology might allow a yet better understanding of the study systems. 60

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63 Keywords: Acoustic communication, ants, mutualists, social parasites, social structure

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Efficient communication to coordinate the actions of up to a million specialised nestmates is 66 fundamental to the success of social insects, especially ants. Various modes of signalling 67 68 have been identified, including the release of semio-chemicals, visual behavioural displays involving movement or posture, tactile interactions, and the comparatively poorly studied use 69 of acoustic signals (Hölldobler & Wilson, 1990, 2009). As hotspots of resources in their 70 environment, ants fiercely defend their colonies using a wide range of weapons (e.g. gland 71 secretions, mandibles, sting), which are deployed in the manner of co-ordinated attacks by 72 73 legions of intercommunicating workers. Nevertheless, ant nests are also magnets for other organisms that have evolved means to overcome the hostility of the host ants. Thus, an 74 75 estimated ~10,000 invertebrate species live as obligate social parasites of ants, able to 76 penetrate and exploit the resources within host colonies in order to complete their life-cycle (Thomas, Schönrogge, Elmes, 2005). The large majority of these adaptations evolved in 77 78 many separate lines, especially among Coleoptera, Diptera, Lepidoptera and other 79 Hymenoptera, from a ten-times greater number of commensals or mutualists (Fiedler, 1998; Hölldobler & Wilson, 1990; Nash & Boomsma, 2008; Pierce et al., 2002; Thomas, 80 Schönrogge et al., 2005). All these myrmecophiles show morphological, behavioural, 81 82 chemical or acoustic adaptations to interact with ants (Cottrell, 1984; Donisthorpe, 1927; Hinton, 1951; Lenoir, D'Ettorre, Errard, & Hefetz, 2001; Malicky, 1969; Wasmann, 1913; 83 84 Wheeler, 1910; Witek, Barbero, & Marko, 2014). Armour, stealth and the secretion of attractive food rewards are frequently sufficient for unspecific or facultative myrmecophiles to 85 access the enemy-free spaces of ants. However, the subversion of the ants' chemical and/or 86 acoustic signalling is generally required to enable true social parasites (sensu Nash & 87 Boomsma, 2008) to live for long periods as undetected intruders in close contact with their 88 hosts. 89

90 A key element of successful co-habitation in ant nests is to circumvent the host's ability to differentiate between nestmates and intruders. Nestmate recognition is a dynamic process, 91 primarily based on the detection of distinctive species- or colony-specific cocktails of 92 cuticular hydrocarbons (CHC) covering the surface of all individuals (Hölldobler & Wilson, 93 94 1990; Howard, 1993; vander Meer & Morel, 1998; Winston, 1992). Social interactions such as allogrooming ensure an exchange between the CHC mixtures among nestmates and give 95 rise to a shared CHC gestalt odour (vander Meer & Morel, 1998). The role that chemical 96 97 communication and nestmate recognition have in maintaining the cohesion of ant societies 98 and those of other social insects has been subject to extensive study, with excellent recent 99 reviews, for example by Martin & Drijfhout (2009) and van Wilgenburg, Symonds, & Elgar 100 (2011): The deployment of chemical communication by obligate social parasites to subvert 101 host recognition systems is equally well reviewed (e.g. Lenoir et al., 2001; von Thienen, 102 Metzler, Choe, & Witte, 2014).

103 In contrast, the function, the origin and role of acoustic signals in ants and their corruption by 104 social parasites are much less well studied. In this review, we therefore focus on the state of 105 the art concerning acoustic signaling in ants, and then consider the acoustic signaling of obligate and facultative myrmecophiles. In both cases we emphasize the insights that have 106 resulted from recent technological advances that allow unalarmed ants and their guests to 107 be recorded and to receive broadcasts of their acoustic signals under semi-natural 108 109 conditions (Barbero, Thomas, et al., 2009; Riva, Barbero, Bonelli, Balletto, Casacci, in 110 press).

We first examine ant sound producing organs and convergent adaptations that allow non-ant organisms to mimic and subvert ant–ant communications, focussing on advances in knowledge since the reviews by Hölldobler & Wilson (1990), Fiedler (1998), Pierce and colleagues (2002), Thomas and colleagues (2005) and Nash & Boomsma (2008), or covered cursorily by Witek and colleagues (2014). We then review recent insights concerning the ant acoustic signals themselves and their corruption by social parasites. This includes both the morphological adaptations to produce acoustic signals, the behavioural responses to them, and thus the impact on ant – social parasite/guest interactions. Much of this builds on the pioneering work of Markl (1965, 1967), DeVries (1991a, 1991b), Hölldobler, Braun, Gronenberg, Kirchner, & Peeters (1994) and Kirchner (1997). Finally we present new data relating the intimacy of interactions of lycaenid butterfly larvae to phylogeny and the similarity of acoustic signalling.

123 Acoustic signalling in ants

The use of acoustics, whether through receiving pressure waves through the air (i.e. sounds 124 125 stricto sensu) or substrate vibrations, is a common means of communication in insects, whose functions include defence, displays of aggression, territorial signalling and mate 126 attraction (Bennet-Clark, 1998; Gerhardt & Huber, 2002). Its advantage as a signal over 127 128 chemical volatiles lies in instantaneous reception that pinpoints a distant, but exact, location 129 to the receiver, for example in social insects to attract help (Markl, 1965, 1967; Roces, Tautz, & Hölldobler, 1993). The physics, use and effects of substrate-borne vibrations of 130 ants and other insects are comprehensively reviewed by P.S. Hill (2009). A simple form 131 involves "drumming", where the substrate is tapped by part of the exoskeleton to produce 132 133 vibrations. Drumming is employed by many ant taxa, but at least four of the eleven subfamilies also stridulate by rasping a 'plectrum' across a 'file' (pars stridens), both 134 chitinous organs being located on opposite segments of the anterior abdomen (see Fig. 1 k-135 136 o, u-y) (Barbero, Thomas, Bonelli, Balletto, & Schönrogge, 2009b; Golden & P.S. Hill, 2016; 137 Ruiz, Martinez, Martinez, & Hernandez, 2006). Although these stridulations produce airborne (as well as substrate-borne) pressure waves that are audible to the human ear, it 138 remains uncertain whether ants can perceive sound as pressure waves through the air 139 (Hickling & Brown, 2000, 2001; Roces & Tautz, 2001). In contrast, there is no controversy 140 141 about the ants' ability to perceive substrate vibrations and two types of sensor have been proposed to receive substrate vibrations: campaniform sensilla measuring the tension in the 142

exoskeleton; and the subgenual organ, a spherical arrangement of sensory cells in the tibia,
as described from *Camponotus ligniperda* (Gronenberg, 1996; Menzel & Tautz, 1994).

Most studies that measure insect acoustics have used accelerometers, moving coil- or particle velocity microphones, often with phase inversion focussing on the vibrational part of the signal rather than pressure waves through the air. Hereafter in this review we use the term "sound" *sensu latu* in its broadest sense, as we do the terms: calls, vibrations, vibroacoustics and stridulations.

150 Early studies suggested that acoustic signals were a minor means of communication among 151 ants, largely confined to activities outside the nest and mainly signalling alarm or calls for rescue, for instance when parts of nests collapse (Markl 1965, 1967). Due to a perceived 152 preponderance of stridulatory organs among soil nesting ant species, Markl (1973) 153 hypothesised that stridulation evolved initially as a burial/rescue signal when volatile 154 chemicals would be ineffective, whereas substrate borne vibrations would at least travel 155 short distances. However, this is not supported by Golden and P.S. Hill (2016), who showed 156 157 that stridulation organs have evolved independently multiple times in ants. In addition, whereas MarkI (1973) suggested that they would probably become vestigial over time in 158 arboreal ant species, due to the rarity of burial by soil, there was instead a strong positive 159 association between the presence of functional stridulation organs and the possession of an 160 161 arboreal life-style (Golden & P.S. Hill, 2016).

Nestmate recruitment is the most frequently reported function for ant–ant acoustic signalling. For example, outside the nest, *Atta cephalotes* uses vibratory signals to attract foraging workers towards newly found food sources (Roces & Hölldobler 1995). The same authors also observed that in the presence of parasitic phorid flies, foragers used acoustics to recruit minor workers for defence, thus also employing vibrations as alarm signals (Roces & Hölldobler, 1995, 1996). Finally, although created by a scraper and file organ located on the first gastric tergite and the post-petiole, Tautz and colleagues (1995) observed that vibrations travelled the length of the body to the mandibles, aiding the cutting of soft young
leaf tissue by stiffening it. Behavioural experiments, however, suggest that this is a
secondary effect and that communication is the main function for these vibrations (Roces &
Hölldobler, 1996).

173 It has recently become clear that acoustic signals are also used to transmit more abstract information, including a species' identity or an individual's caste and status (Barbero, 174 Thomas et al., 2009; Casacci et al., 2013; Ferreira, Cros, Fresneau, & Rybak, 2014). For 175 example, modern molecular analyses revealed the neotropical ponerine ant species, 176 177 Pachycondyla apicalis, to be a species complex of five cryptic lineages. The stridulations of three largely sympatric lineages are also distinctive, suggesting that morphological 178 characters on the pars stridens differ in length, width and ridge gap in each lineage (Ferreira, 179 180 Cros, Fresneau, & Rybak, 2014; Wild, 2005). By contrast, two allopatric lineages had very 181 similar acoustics, suggesting disruptive selection on this trait where sympatric overlap is 182 high.

Acoustic patterns also signal caste and hierarchical status in at least two genera of 183 184 Myrmicinae ants: Myrmica (Barbero, Thomas et al., 2009) and Pheidole (Di Giulio et al., 2015). In both taxa, the queens produce distinctive stridulations which, when played back to 185 kin workers, elicit additional 'royal' protective behaviours compared with responses to worker 186 signals (Barbero, Bonelli, Thomas, Balletto, & Schönrogge, 2009; Barbero & Casacci, 2015; 187 188 Barbero, Thomas et al., 2009; Casacci et al., 2013; Ferreira, Poteaux, Delabie, Fresneau, & Rybak, 2010). In addition, in *Pheidole pallidula* the soldier and minor worker castes also 189 190 make distinctive vibroacoustic signals (Di Giulio et al., 2015). Unlike Pachycondyla species, little inter-specific variation was detected in either the queen- or worker-sounds made by 191 192 closely-related sympatric species of Myrmica (Barbero et al., 2012; Barbero, Thomas et al., 2009; Thomas, Schönrogge, Bonelli, Barbero, & Balletto, 2010), which are instead clearly 193 194 demarcated by unique hydrocarbon profiles (Elmes, Akino, Thomas, Clarke, & Knapp,

195 2002). Although the young stages of tested ants are mute (e.g. DeVries, Cocroft, & Thomas, 1993), Casacci and colleagues (2013) found that acoustic signalling appears to act as a 196 substitute for other forms of communication in developing Myrmica pupae. The various 197 stages of ant brood, from egg to pupa, are afforded ascending levels of priority based on 198 199 tactile and chemical cues (Brian, 1975). Most are mute, but the older "brown", sclerotised pupae of Myrmica species produce calls, emitted as single pulses, similar to those of 200 workers (Casacci et al. 2013). This coincides with a presumed reduced ability to secrete 201 202 brood recognition pheromones during this period, and brown pupae that were experimentally silenced fell significantly behind their mute white siblings in social standing. 203

204 Acoustic signals of myrmecophiles

Derived acoustic signals that enhance interactions with ants are increasingly being confirmed in both juvenile and adult stages of myrmecophiles. To date, most studies involve riodinid and, especially, lycaenid butterfly larvae and pupae (e.g. Barbero, Thomas et al., 2009; DeVries, 1990, 1991a; Pierce et al., 2002). However, similar phenomena were recently described from adults of a socially parasitic beetle, *Paussus favieri* (Di Giulio et al., 2015), where males and females emit mimetic stridulations using a row of scrapers on the proximal abdominal segment rasping across a file located on the hind femora (see Fig. 1p-t).

212 Stridulation organs

With a few exceptions, an ability to produce calls occurs after the third larval moult in riodinid and lycaenid larvae, coinciding with the development of chemical 'ant organs', which perhaps suggests they act synergistically (DeVries, 1991a). In most riodinids, acoustic signals are generated by grooved vibratory papillae. These are typically found in pairs on the prothorax, and grate against specialised epicranial granulations when the larva rotates its head (see Fig 1a-e), especially when walking or under attack, generating low amplitude substrate-borne calls (DeVries, 1991a). The tribe Eurybiini lacks vibratory papillae; instead, 220 caterpillars generate calls by scraping teeth on a prothoracic cervical membrane against the epicranial granulations in at least some mutualists or entomophagous predators of ant-221 tended Homoptera (DeVries & Penz, 2002; Travassos, DeVries, & Pierce, 2008). The 222 detection of dedicated organs in lycaenid larvae that produce calls has been elusive, apart 223 from a file-and-scraper described between the 5th and 6th abdominal segments of Arhopala 224 madytus (C. J. Hill, 1993) and a putative organ in Maculinea rebeli larvae (see Fig.1fg). In 225 other species strong substrate-borne vibrations (and apparently weak air-borne sounds) may 226 227 be generated by muscular contractions of the abdomen, which compress air through the 228 tracheae to produce distinctive rhythms and intensities in the manner of a wind instrument, 229 as described by Schurian and Fiedler (1991) for *Polyommatus dezinus*. These vibroacoustic 230 signals range from low background calls punctuated by pulses in mutualists (DeVries, 231 1991a) to the grunts, drumming and hisses of the host-specific Jalmenus evagoras 232 (Travassos & Pierce, 2000), to the mimetic calls of Maculinea larvae (Barbero, Bonelli et al., 2009; DeVries et al., 1993; Sala, Casacci, Balletto, Bonelli, & Barbero, 2014). 233

In contrast, the pupae of all lycaenids studied (Pierce et al., 2002) and a minority of riodinids (DeVries, 1991a; Downey & Allyn, 1973; 1978; Ross, 1966) have a well-developed file-andscraper organ (two pairs in the case of riodinids) situated between opposite segments of the abdomen, that emit substrate- and air-borne calls often audible to humans (see Fig 1h-j). In lycaenids, the plate against which teeth are rubbed may be complex, consisting of tubercles, reticulations or ridges (Alvarez, Munguira, & Martinez-Ibanez, 2014).

240 Acoustic signalling in ant–myrmecophile interactions

Evidence that the acoustics of myrmecophiles are adaptive to their interactions with ants has progressed from correlative studies to two experimental approaches: muting the myrmecophile or recording and playing back their calls to undisturbed ant colonies.

First, DeVries (1991c) showed that fewer ants attended larvae of the mutualistic riodinid *Thisbe irenea* that had been artificially silenced compared with controls that were able to 246 call, establishing that at least one function of riodinid calls is to attract ants. Similarly, Travassos and Pierce (2000) demonstrated that pupae of the lycaenid Jalmenus evagoras 247 stridulated more frequently in the presence of *Iridomyrmex anceps* ants, and attracted and 248 maintained a larger number of guards than muted ones. The calls convey the pupa's value 249 250 as a provider of nutritious secretions to the ants, which does however, represent a significant cost to the pupae. Tended pupae have been shown to lose 25% of weight and take longer to 251 eclose than untended ones (Pierce, Kitching, Buckley, Taylor, & Benbow, 1987). In further 252 253 behavioural experiments Travassos and Pierce (2000) showed that pupae used acoustic 254 signalling to adjust the number of attendant ants. They provided a path from an *I. anceps* 255 nest to signalling pupae and scored the rate of worker movement in relation to signal 256 strength once the pupa was discovered. This appears to be an important fitness component evolved to attract no more than an adequate number of ant guards against enemy attacks. 257 258 The larvae of J. evagoras produce more varied acoustic signals than pupae - grunts, hisses and drumming – and are also heavily attended and guarded by their mutualist ant (Pierce et 259 260 al., 2002). Hisses are emitted briefly after encountering a worker, whereas grunts are produced throughout ant attendance. The ability of J. evagoras juveniles to produce distinct 261 262 vibrations, some probably with different functions, suggests the evolution of a finely-tuned acoustic system of communication with their hosts, which might be elucidated using play-263 back experiments. 264

265 In parasitic interactions with ant colonies, the clearest evidence to date that some acoustic signals are mimetic involves the highly specialized species of the Myrmica ant - Maculinea 266 butterfly and *Pheidole* ant - *Paussus* beetle systems. Initially, DeVries and colleagues (1993) 267 showed that the calls made by larvae of four Maculinea species differed from those of 268 phytophagous lycaenids in showing distinctive pulses that resembled the stridulations of 269 Myrmica worker ants. This was the first suggestion of mimicry of an adult host attribute by 270 the caterpillars, which appeared to be genus- rather than species-specific. The insects in 271 272 early experiments were unavoidably alarmed, being held with forceps during the recording,

273 but a similar genus-specific result was later obtained using modern equipment and unstressed ants and butterflies. Both the pupae and larvae of Maculinea species closely 274 mimicked three attributes of their hosts' acoustic signals: dominant frequency, pulse length, 275 pulse repetition frequency (Barbero, Bonelli et al., 2009, Barbero, Thomas et al., 2009). 276 277 However, the calls of both stages were significantly more similar to queen ant calls than they were to worker calls, despite each being generated in a different way (see Fig.1f-i). 278 Behavioural bioassays, where the calls of butterflies and ants were played back to 279 280 unstressed Myrmica workers, revealed that the calls of juvenile Maculinea, especially those 281 of pupae, caused workers to respond as they do to queen ant calls. Both types of acoustic 282 stimuli caused worker ants to aggregate, antennate the source of sound, and show 283 significantly higher levels of guarding behaviour than was elicited in response to worker ant 284 calls (Barbero, Thomas et al., 2009).

285 Similar, but more sophisticated communication, was recently described between the carabid 286 beetle Paussus favieri, an obligate social parasite in all stages of its life-cycle, and their host ant *Pheidole pallidula* (Di Giulio et al., 2015). Here the adult beetle can generate three types 287 288 of call when it stridulates, which respectively mimic the calls made by the queens, the soldiers and the minor worker caste of its host. These calls elicit a range of responses when 289 290 played back to worker ants, consistent with the intruder's more diverse activities (compared to juvenile Maculinea) in different parts of the host's society and nest. Thus P. favieri's 291 292 various stridulations can elicit recruitment, including digging (rescue) behaviour, as well as the enhanced level of 'royal' (queen ant) protection observed towards Maculinea pupae and 293 larvae. 294

295 [insert Figure 1]

296 Larval acoustic signals and phylogeny in the Lycaenidae

Various authors (e.g. DeVries, 1991a, 1991b; Fiedler, 1998; Pech, Fric, Konvicka, & Zrzavy,
2004; Pellissier, Litsios, Guisan, & Alvarez, 2012; Pierce et al., 2002) have analysed the

evolution of myrmecophily in lycaenids and riodinids, including social parasitism in the Lycaenidae, and most concluded that it also provided a template for diversification and radiation in these species-rich families. Pierce and colleagues (2002) argued convincingly that social parasitism (including entomophagy of the domestic Hemiptera of ants) has evolved independently in at least 20 lineages.

The analysis of acoustics as a parameter in evolutionary studies of these taxa was 304 pioneered by DeVries (1991a, 1991b). In seminal early papers, DeVries (1991a, 1991b) 305 306 found that only lycaenids and riodinids that interacted with ants produced calls, while several 307 non myrmecophilous members of the tribe Eumaeini were silent. Subsequent studies and 308 reviews confirmed this pattern (e.g. Fiedler, Seufert, Maschwitz, & Idris, 1995) and provided 309 evidence of the use of lycaenid calls in enhancing the interaction with ants (Pierce et al., 310 2002; Barbero, Thomas et al., 2009, Sala et al. 2014). However, some lycaenid and riodinid 311 larvae and pupae also emit sounds when disturbed by putative predators or parasitoids, even if ants are absent. In addition, other species classed as having no interaction with ants 312 do emit sound (e.g. Alvarez et al., 2014; Downey & Allyn, 1973; 1978; Fiedler, 1992, 1994; 313 Schurian & Fiedler, 1991). The most recent study, by Riva and colleagues (in press), found 314 315 that lycaenid sounds are highly specific and are emitted by both non- and myrmecophilous species. Calls by species that are least associated with ants consist of shorter and more 316 distant pulses relative to those of species that are highly dependent on them. 317

318 Here we further explore the hypothesis that the strength of ant-myrmecophile interactions (using Fiedler's 1991 definitions) leads to characteristic sound profiles that may be a better 319 predictor of the similarity of sound between species than their phylogenetic distance. We 320 present a new analysis of the acoustic profiles made by 13 species of European lycaenids, 321 ranging from highly integrated 'cuckoo' social parasites (Maculinea alcon, Ma. rebeli) via one 322 host-specific mutualist (Plebejus argus) and a spectrum of generalist myrmecophiles, to 323 species for which little or no interaction is known (Lycaena spp.). The 13 species (see Fig. 2) 324 325 are a subset of the commensal or mutualistic species used by Riva and colleagues (in

Page **13** of **31**

press), with three species of *Maculinea* added to represent the two levels of intimate
 integration found in this socially parasitic genus (Thomas, Schönrogge et al., 2005).

Fourth instar caterpillars were recorded using customized equipment, as described by Riva 328 329 and colleagues (in press). We analyzed recordings of three individuals per species, randomly selecting two trains of five pulses in each trace. Fourteen sound parameters were 330 measured using Praat v. 5.3.53 (Boersma & Weenink, 2013). These included the lower and 331 higher guartiles of the energy spectrum (Hz), power (dB²), intensity (dB), the root-mean-332 square intensity level (dB) and the relation of the frequency peak energy to the call total 333 334 energy (%). Two temporal variables were measured from the oscillogram: the duration of the pulse (s) and the Pulse Rate (calculated as $1/t_{start}(x) - t_{start}(x+1)$; s-1). Six additional variables 335 336 were estimated on each pulse by inspection of power spectra: the frequency of the first, 337 second and third peak amplitudes (Hz), the intensity of the first two peaks (dB) and the center of gravity (Hz). 338

339 Hierarchical Cluster analyses was performed on a matrix of normalized Euclidean distances over sound parameters, averaged by individual using unweighted pair-group average 340 (UPGMA) in Primer v. 6.1.12 (Primer-E Ltd.). A two-sample t - test was used to compare 341 342 differences between group distances. To test whether species differences reflect degrees of 343 myrmecophily, we used Phylogenetic Regression as implemented in the library "phyreg" (Grafen, 1989) using R (R Core Team, 2015). Principal components, derived by PCA on log-344 transformed sound parameters, were correlated with the degree of myrmecophily while 345 controlling for phylogenetic relatedness among species. To assemble a working phylogeny, 346 347 we used cytochrome oxidase subunit 1 (COI) sequences of the 13 lycaenid species from two 348 recent studies on the Romanian and Iberian butterflies (Dinca et al., 2015; Dinca, Zakharov, Hebert, & Vila, 2011). Geneious Pro 4.7.5 (Biomatters, http://www.geneious.com/) was used 349 350 to align COI sequences and to produce a neighbor-joining (NJ) tree. We also included in the phylogeny Hamearis lucina (Riodinidae) and Pieris rapae (Pieridae) as outgroups. 351

352 Two trees for species' phylogenetic distance and for the similarity of acoustic profiles are presented in Figure 2, together with the score for myrmecophily of each species. Similarities 353 in sound profiles neatly match the spectrum of observed strengths and specificities in 354 myrmecophily across the study species, much more closely than does phylogeny. Overall, 355 356 PC1 of the acoustic parameters explained 56% and PC2 a further 27% of variation, and both were significantly correlated with the differences in myrmecophilous relationships (PC1: $F_{1,13}$ 357 = 11.146, P = 0.005; PC2: $F_{1,13} = 6.959$, P = 0.020) after accounting for phylogeny using 358 359 phylogenetic regression.

360 It is apparent that the sound profiles of Ma. rebeli and Ma. alcon (average Euclidean distance (\pm 1SD) between *Ma. rebeli* and *Ma. alcon* = 1.65 \pm 0.14) are far removed from all 361 other species, including from their congeners Ma. arion and Ma. teleius (Barbero, Bonelli et 362 363 al., 2009; Sala et al., 2014). Indeed, the mean Euclidean distances in the acoustic signals of 364 Ma. alcon or Ma. rebeli from other lycaenid species are among the highest measured to date (mean Euclidean acoustic distance of Ma. alcon vs. lycaenids other than M. rebeli: 7.41 ± 365 1.00; Ma. rebeli vs lycaenids other than Ma. alcon: 7.66 ± 1.01; see also Riva et al. in press). 366 367 This is consistent with the intimate level of social integration these species achieve within host ant nests, an association that is so close that in times of shortage the ants kill their own 368 brood to feed to these 'cuckoos' in the nest (Thomas, Elmes, Schönrogge, Simcox, & 369 Settele, 2005). It is also notable that the acoustics of *Plebejus argus*, the only host-specific 370 371 myrmecophile among the mutualistic species, is less similar to its nearest relative *Plebejus* argyrognomon, and appears to converge with the two 'predatory' Maculinea social parasites 372 even though its 'host' ant, Lasius niger, has no known stridulation organs and belongs to a 373 different subfamily to Myrmica (mean Euclidean acoustic distance of P. argus vs. P. 374 argyrognomon: 4.33 ± 0.30; *P. argus vs M. arion*: 2.51 ± 0.55; paired *t* test: *t*₁₆ = -8.723, *P* < 375 0.001; distance of *P. argus vs. Ma. teleius*: 3.79 ± 0.28 ; paired *t* test: $t_{16} = -3.963$, *P* = 0.001). 376 Scolitantides orion perhaps represents selection in the opposite direction to P. argus, being 377 378 less host specific than its ancestry or relatives might suggest, as, less convincingly, may

379 Polyommatus icarus. Yet despite L. coridon and L. bellargus being close congeners, sounds emitted by L. bellargus are much more similar to those produced by P. argyrognomon 380 (belonging to the same myrmecophilous category - 3) rather than to L. coridon (mean 381 Euclidean acoustic distance of L. coridon vs L. bellargus: 3.87 ± 0.15; P. argyrognomon vs L. 382 383 *bellargus*: 1.54 \pm 0.20; paired *t* test: t_{16} = 27.775, *P* < 0.001). A possible, but untested, explanation is that this reflects a similar disruptive selection via acoustics to that described in 384 sympatric lineages of the ant Pachycondyla, since the juveniles of these congeneric 385 butterflies overlap largely in distribution, sharing the same single species of foodplant and 386 often the same individual plant. 387

However, given the small number of species studied, we caution against over-interpreting 388 389 the apparent patterns depicted in Figure 2, and suggest they be tested by comparative 390 behavioural experimentation. We also recognise that vibrations of less- or nonmyrmecophilous lycaenids (and other taxa) may have very different functions, such as 391 392 repelling natural enemies (Bura, Fleming, & Yack, 2009; Bura, Rohwer, Martin, & Yack, 2011). We tentatively suggest that ancestral species in the Lycaenidae were preadapted to 393 394 myrmecophily through an ability to make sounds, and that once behavioural relationships with ants evolved, the selection regime changed resulting in adaptive mimetic sound profiles, 395 396 at least among obligate myrmecophiles.

397 [insert Figure 2]

398 Conclusions & Future Research

Ants are known to sometimes use multiple cues to moderate kin behaviour, for example by combining posturing, tactile and chemical interactions to convey complex or sequential information and to elicit particular responses between members of their society (Hölldobler & Wilson, 1990). To date little is known of how acoustic signalling might interact with other 403 means of communication, and less still of whether myrmecophiles manipulate behaviour404 using multiple cues.

Sound may be used synergistically with other modes of signalling. Hölldobler and colleagues 405 406 (1994) studied the role of audible vibrational signals made by the Ponerine ant Megaponera foetens, a raider of termite colonies, in the context of trail following and column building. 407 They found that stridulations were emitted only during disturbances and for predator 408 avoidance. It is also known that *M. foetens* has a distinctive pheromone to signal alarm 409 (Janssen, Bestmann, Hölldobler, & Kern, 1995). These observations suggest that vibrations 410 411 may be used to qualify a general alarm signal that is chemical, but again this requires formal testing. This is in contrast to the observations by Casacci and colleagues (2013) described 412 above where acoustic signalling appears to replace chemical and tactile signal apparently 413 with the same function of signalling rank, but this is not truly a case of multimodal 414 415 communication.

416 To date, no direct evidence exists for the behavioural consequences of full synergistic 417 multimodal communication involving acoustics. Yet the interactions of Maculinea butterfly larvae and their Myrmica host ant societies illustrate the importance of both chemical and 418 419 acoustic mimicry. Here, the acceptance (or rejection) of larvae as members of their host 420 colony appears to be based entirely on a mimetic mixture of chemical secretions, but on this 421 cue alone intruders are treated simply like the low-ranking kin brood (Akino, Knapp, Thomas, & Elmes, 1999; Thomas et al., 2013; Thomas, Schönrogge et al., 2005). It is the ability 422 simultaneously to emit acoustic calls that mimic adult hosts, and furthermore mimic queen 423 sounds, that is believed to explain the observed priority 'royal' behaviour that workers 424 regularly afford to social parasites, giving them a status that exceeds that of large ant larvae. 425 Not only do these brood parasites gain priority in the distribution of food by nursery workers 426 to the extent that workers feed younger kin ant brood to the Maculinea larvae when food is 427 428 short, but they are also carried ahead of kin ant brood when moving nest or during rescues

429 (Elmes, 1989; Gerrish, 1994; Thomas, Schönrogge, et al., 2005). Anecdotal observations of the manipulation of Paussus favieri by the beetle Pheidole pallidula suggests a similar 430 chemical-acoustic mechanism (Di Giulio et al., 2015), but as with ant-ant communication 431 itself, the putative use of acoustics in multimodal communication requires rigorous testing. 432 433 About 10,000 species of invertebrates from 11 orders are estimated have evolved adaptations to infiltrate ant societies and live as parasites inside nests (Hölldobler & Wilson, 434 1990). Current studies have largely focussed on the family Lycaenidae among the 435 Lepidoptera and a few selected species of Coleoptera. While the study systems used today 436 437 provide some variety in the type of interactions with their host ants, there is clearly a vast 438 variety still to be discovered to understand respective roles of signalling modes and the 439 social interactions in ants and other social insects.

440 The important role that acoustic signalling has in ant- and other social insect societies is well 441 established and it is perhaps unsurprising that other, interacting species show adaptations 442 that relate to the hosts acoustic traits. In only a few cases, however, has the role of vibroacoustics in mediating myrmecophile - host interactions been investigated experimentally. 443 444 The modalities of signal production, transmission and reception remain largely unknown for most species of myrmecophiles or indeed their hosts, but the greatest future challenge is to 445 446 understand how different modes of signalling interact. Social insects are well known to interpret stimuli in a context-dependent manner, where the same stimulus can trigger a 447 448 different behaviour when encountered under different circumstances (Hölldobler & Wilson 1990). Other aspects of insect social behaviour have been subject to sophisticated and 449 successful experimentation, and it should be possible to unravel this essential aspect of 450 communication. Hunt and Richards (2013) suggested that understanding the suites of 451 modalities in signalling enables a clearer view of the adaptive role of multimodal 452 communication, and while that has been true for rare examples such as the honey bee 453 waggle dance, research into understanding the role of ant acoustics is in its infancy. With the 454 455 development of recording equipment that is portable, affordable, which can focus on individuals and record sound and behaviour at the same time, our understanding of social interactions should become more specific. Such instruments, laser-vibrometers and handheld "noses" for acoustic and chemical analyses, are being developed for engineering applications and could be deployed to record acoustic and chemical signals in behavioural science in the near future. Technological developments in both recording equipment and behavioural experimentation will allow designing studies following the same principles to investigate synergistic effects of multiple chemical signals.

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472 Ethical Note

The authors confirm that their work adheres to the ASAB/ABS and ARRIVE Guidelines. The guide to ethical information required for papers published in the journal has been consulted as well. *Maculinea* caterpillars were collected under permit from The Italian Ministry for the Environment (protocol numbers: 446/05. DPN/2D/2005/13993 & 0012494/PNM/2015). The authors declare that there is no conflict of interest.

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711 Figures

Figure 1. The comparative morphology of sound production organs in myrmecophiles and 712 host ants. (a-e) the riodinids Synargis gela and Thisbe irenea (Riodinidae); larva (f, g) and 713 pupa (h-j) of the obligate lycaenid social parasite Maculinea rebeli and its adult host ant 714 715 Myrmica schencki (k-o); the adult beetle Paususs favieri (p-t) and its host Pheidole pallidula (u-y). (a) Frontal view of Synargis gela head showing typical position of the riodinid vibratory 716 papillae; (b) general view of *Thisbe irenea* anterior edge of segment T-1 showing a vibratory 717 718 papilla (arrow) and the surface of the epicranium where the vibratory papilla strikes; (c) detail 719 of the vibratory papilla showing the annulations on its shaft and the epicranial granulations: 720 (d) enlarged view of the epicranial granulation and vibratory papilla; (e) details showing two 721 sizes of epicranial granulations. (f) Position of (g) the presumed sound producing organ of 722 Maculinea rebeli caterpillars and of its pupa (h), formed by a stridulatory plate (pars stridens) 723 placed on the fifth abdominal segment and a file (*plectrum*) in the sixth abdominal segment. (k,p,u) Respective positions of the stridulatory organs of Myrmica schencki, Paussus favieri 724 725 and *Pheidole pallidula*; the organs are composed of suboval pars stridens (I,q,v) with minute ridges (m,r,w) and a plectrum (n, x) consisting of a medial cuticular prominence (t,y) that 726 727 originates from the posterior edge of the postpetiole in the two ant species or of a curved row of small cuticular spines in P. favieri (s,t). (a, modified by De Vries 1991; b-e modified by 728 729 DeVries 1988; p-y modified by Di Giulio et al. 2015).

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Figure 2. A diagram of the phylogeny (left) and the cluster analysis constructed from a matrix of pairwise normalized Euclidean distances of the sound profiles from three caterpillars of 13 species of lycaenid. Symbols and values refer to the intensity of interaction of the lycaenid species with their host ants (0 = none; 4 = social parasite), following Fiedler (1991).

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