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Application of CO₂ carbon stable isotope analysis to ant trophic ecology: preliminary results

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25 Abstract

Stable isotope analysis of animal tissues is commonly used to infer diet and trophic position. 26 27 However, it requires destructive sampling. The analysis of carbon isotopes from exhaled CO₂ is non-invasive and can provide useful ecological information because isotopic CO₂ signatures can 28 reflect the diet and metabolism of an animal. However, this methodology has rarely been used on 29 invertebrates and never on social insects. Here, we first tested whether this method reflects 30 differences in δ^{13} C-CO₂ between workers of the Mediterranean ant *Crematogaster scutellaris* 31 (Olivier, 1792) (Formicidae: Crematogastrini) fed with beet (C3) and cane (C4) sugar (Beta 32 vulgaris L., Amaranthaceae and Saccharum officinarum L., Poaceae, respectively). We found that a 33 significant difference can be obtained after 24 hours. Consequently, we used this technique on wild 34 35 co-occurring ant species with different feeding preferences to assess their reliance on C3 or C4 sources. For this purpose, workers of C. scutellaris, the invasive garden ant Lasius neglectus (Van 36 Loon, Boomsma & Andrasfalvy, 1990) (Formicidae: Lasiini) and the harvester ant Messor capitatus 37 (Latreille, 1798) (Formicidae: Stenammini) were sampled. No significant differences in their carbon 38 39 isotopic signatures were recorded, suggesting that in our study site no niche partitioning based on the carbon pathway occurs, with all species sharing similar resources. However, further analysis 40 revealed that *M. capitatus*, a seed-eating ant, can be regarded as a C3 specialist, whereas *L*. 41 neglectus and C. scutellaris are generalists that rely on both C3 and C4 pathways, though with a 42 preference for the former. Our results show that this methodology can be successfully applied even 43 44 to small animals like ants and can provide useful information on the diets of generalist omnivores.

46 Introduction

Stable isotopes analysis (SIA) of animal tissues is widely used in animal ecology to infer diet, 47 determine trophic positions and define trophic niches (Fry, 2006; Boecklen et al., 2011). The most 48 widely used isotopes are those of C and N (δ^{13} C and δ^{15} N), which provide information on the main 49 carbon source used by a consumer and its trophic level based on predictable isotope fractionation 50 51 occurring from one trophic level to another (Post, 2002). Moreover, SIA is employed for the study of invasive species impacts (Balzani et al., 2016; Stellati et al., 2019) and interactions (Haubrock et 52 53 al., 2019a, 2020) and to predict the effects of species reintroductions (Haubrock et al., 2019b). Applications in studies of insects include the determination of resource origin (Ouyang et al., 2015), 54 movements (Madeira et al., 2013, 2014; Zhang et al., 2020) and feeding periods (Ouyang et al., 55 56 2014), as well as the study of nutrient uptake (Pollier et al., 2016) and allocation (Levin et al., 2017a,b). In ant ecology, SIA is used to study the trophic role of each species (e.g. Ottonetti et al., 57 2008), to infer how ant assemblages vary in different habitat conditions (Gibb & Cunningham, 58 2011), to estimate the importance of trophobiosis (Brewitt et al., 2015) and to study the effects of 59 natural local baseline variations (Cronin et al., 2015) and colony parameters (Barriga et al., 2013) 60 61 on isotopic signatures. This approach requires destructive sampling and is therefore difficult to use 62 when dealing with rare species or when repeated measurements on the same subjects are desired 63 (Hatch et al., 2002b). However, social insects are somewhat unique in that colonies can be 64 considered the ecological unit of study and these can be repeatedly sampled overtime with relative ease, to assess the effect of changes in resource availability on their diet (Roeder & Kaspari, 2017; 65 Shik et al., 2018). 66

67 Stable isotope analysis can also be performed on gas — a common technique in 68 geochemistry in which C stable isotopes of CO_2 and CH_4 are employed to identify emission sources 69 in natural or anthropised environments (e.g. Venturi et al., 2017, 2019, 2020 and references

therein). In the biological sciences, isotopic gas measurements on respiration-derived CO₂ are 70 71 relatively common. This approach is primarily used in medical studies in which changes in the 72 isotopic composition of CO₂ (breath tests) are used to investigate human pathologies, physiology and nutrition, and to test the metabolic rate during stress exercises (McCue & Welch, 2016). 73 Similarly, this approach can be applied to the animal sciences. The carbon isotope signature (δ^{13} C) 74 of exhaled CO₂ correlates with diet and metabolism as it reflects the signature of the catabolised 75 substrate (Perkins & Speakman, 2001). Moreover, the δ^{13} C values in the breath indicate both the 76 present and past diet and are used to calculate the proportion of lipids and carbohydrates 77 metabolised by an animal (Hatch et al., 2002a,b). Comparing the δ^{13} C values of breath and body 78 79 tissues allows for the detection of diet changes (Podlesak et al., 2005; Voigt et al., 2008a). Shifts in 80 substrate oxidation during starvation or torpor/hibernation have been studied using breath stable isotopes in vertebrates (McCue & Pollock, 2013; Lee et al., 2017; Rosner & Voigt, 2018) and 81 82 invertebrates (McCue et al., 2015). The reasoning behind these studies is that breath is highly metabolically active compared to tissues, which show slower turnover rates, thus reflecting the 83 more recent diet (Perkins & Speakman, 2001). If breath and tissue signatures differ, a dietary shift 84 is likely to have occurred over a certain temporal window (Podlesak et al., 2005; Voigt et al., 85 2008a). On the other hand, lipid δ^{13} C values are depleted with respect to carbohydrate δ^{13} C (Post et 86 87 al., 2007), and this difference is recorded in breath isotope signatures (Voigt et al., 2008b,c). Therefore, animals metabolising their fat stores (due to starvation or torpor) will have lower breath 88 δ^{13} C than non-fasted animals (McCue & Welch, 2016). The advantages of this methodology are that 89 90 it (i) uses a non-invasive and non-destructive sampling technique, (ii) can be performed repeatedly on the same individual, and (iii) can be applied to studies of endangered or protected animals 91 92 (Hatch et al., 2002b). However, most studies have focused their attention on birds and mammals, while investigations on invertebrates are less common (but see e.g. DeNiro & Epstein, 1978; Miller 93 et al., 1985; Engel et al., 2009). To the best of our knowledge, no breath study has addressed the 94 95 trophic ecology of ants.

Most ant species are believed to have an omnivorous and opportunist diet (Hölldobler & 96 97 Wilson, 1990; Blüthgen & Feldhaar, 2010). Still, different food preferences can be found in different taxa. For example, Formicinae generally tend to feed on lower trophic levels (mainly 98 homopteran honeydew or plant extrafloral nectaries) compared to Myrmicinae, which are more 99 100 carnivorous, although generalisations are problematic (Fiedler et al., 2007; Blüthgen & Feldhaar, 2010; Brewitt et al., 2015). Food selection is determined by several other factors such as resource 101 102 distribution and availability, food quality and quantity, handling time and colony nutritional requirements (Csata & Dussutour, 2019). Additionally, resource value can vary in different periods 103 of the year, changing their profitability and, consequently, their uptake by ants (Kay, 2002). The 104 105 application of SIA techniques can, therefore, provide cost-effective information on ant metabolism 106 and can help to elucidate details of their trophic ecology.

The acrobat ant Crematogaster scutellaris (Olivier, 1792) is a Mediterranean Myrmicinae 107 ant that nests in tree trunks (Santini et al., 2011). While it is commonly observed tending 108 109 homopterans to obtain their carbohydrate honeydew, it is also known to be an avid predator of arthropods (Schatz et al., 2003; Frizzi et al., 2016). Indeed, SIA has revealed its high trophic level, 110 111 suggesting a strong reliance on animal prey (Ottonetti et al., 2008). The invasive garden ant Lasius neglectus (Van Loon, Boomsma & Andrasfalvy, 1990) is a Formicinae ant considered invasive in 112 Europe (e.g. Nagy et al., 2009). This species tends to monopolise aphid aggregations and to rely 113 114 primarily on honeydew (Paris & Espadaler, 2009; Frizzi et al., 2018). While seed collection (with elaiosome consumption) is common, few genera, such as the Myrmicinae Messor and Pheidole, 115 actually eat them (Blüthgen & Feldhaar, 2010). However, the harvester ant Messor capitatus 116 (Latreille, 1798) often feeds on plant seeds (Cerdá & Retana, 1994), while Pheidole pallidula 117 retrieves mostly insects and thus is more predaceous (Detrain, 1990). 118

In this study, carbon isotopes of CO₂ emitted from different ant species collected in the field
were analysed to assess their diet and their degree of trophic specialisation on C3 or C4 sources.

First, the application of this method on ants was tested by comparing the δ^{13} C values in the CO₂ 121 122 produced by laboratory colonies of C. scutellaris fed with different food sources. In particular, the colonies were fed with either raw beet sugar (C3 plant) or raw cane sugar (C4 plant), which are 123 known to have distinct isotopic signatures (O'Leary, 1981; Boecklen et al., 2011). Then, the 124 isotopic signatures of the gas produced by wild colonies of three ant species (C. scutellaris, L. 125 neglectus and M. capitatus), which have different feeding preferences, were analysed. We 126 127 hypothesised that if these species feed on distinct carbon pathways, a difference in CO₂ isotope should detected. signature be 128

130 Materials and Methods

In July 2019, we collected workers of the ant C. scutellaris from 10 natural colonies on the 131 University Science Campus in Sesto Fiorentino or in the surrounding area (43°49'00"N, 132 11°11'59''E). The area is a managed suburban park made up by University buildings surrounded by 133 134 overgrown fields, public gardens with ornamental trees (Acer campestre, Celtis australis, Quercus robur, Quercus cerris, Quercus ilex, Fraxinus angustifolia) and some sparse trees of Populus sp., 135 Morus sp. and Ailanthus altissima. Samples from each colony were brought to the laboratory and 136 137 subdivided into two groups of approximately 200 workers each. The ants were housed in plastic aquaria (10 x 20 x 30 cm) with Fluon®-coated walls to prevent escape. One group was fed with raw 138 beet sugar (C3) and the other group was supplied with raw cane sugar (C4). Sugar and water were 139 140 provided ad libitum for two weeks to let the ants equilibrate to their new diet.

After two weeks, 100 workers from each group were placed into closed 3-litre respiratory 141 jars, which have a 3-way valve sealed on the cap. The jar walls were coated with Fluon® in the 142 143 upper portion to prevent ants from reaching the valve. Four jars with no ants were used as controls. The jars were kept in a thermostatic chamber with controlled conditions to reproduce the daily (24 144 hours) light-dark cycle (16 h / 8 h). During the daytime, the temperature was set to a constant 27°C, 145 146 and during the night it was 19°C. The air in the jars was sampled at the beginning of the experiment (t₀) and after 24 hours (t₂₄). At each time, aliquots of air (50 ml) were collected (after mixing the air 147 148 in each jar) using a 60 ml plastic syringe connected to the valve and were stored in 1-litre Supelco Tedlar® gas sampling bags equipped with a push/pull lock valve. 149

In the second set of experiments in July–August 2019, we collected specimens of three cooccurring species, *C. scutellaris*, *M. capitatus* and *L. neglectus*, in the same area to assess the differences in their diets. For each species, we selected four natural colonies and collected 70–200 workers (depending on the species' individual size to obtain enough biomass) from each. Workers were immediately placed into sealed respiratory jars and air was collected according to the procedure described above. No food was provided during or before the experiment. At the end of the experiment, the ants were euthanised by freezing at -20°C and dried in an oven at 60°C for 48 h before being weighed using an electronic balance (accuracy: 0. 01 mg).

Carbon isotopic composition of CO₂ (expressed as δ^{13} C-CO₂, % vs. V-PDB standards) was 158 analysed by Cavity Ring-Down Spectroscopy (CRDS) using a Picarro G2201-i Analyzer (CO2 159 160 operational range: 100-4000 ppm). The Supelco Tedlar® gas sampling bags were directly connected to the analyser inlet via silicon connections. Statistical analyses were performed using R 161 software version 3.6. CO₂ production rates were estimated by multiplying the mean difference in 162 gas concentration within the 24 h by the jar volume (3 l) and dividing by 24 to obtain a rate per 163 hour. The mean individual CO₂ production rate was calculated by dividing by the number of 164 workers in the jar. Both estimations were corrected for the mean dry weight of the workers. 165

Linear mixed effects models and simple linear models were built for the first and second 166 experiments, respectively. Separated models were constructed for t₀ and t₂₄ using log-transformed 167 CO₂ concentration and isotopic signature as the response variables and sugar type or species as 168 169 predictors. The CO₂ in the respiration jars was a mixture of the gas produced by ants during the experiment and that of the air already present in the jar. To estimate the signature of the CO₂ 170 171 produced by ants, Keeling plots (Keeling, 1958, 1961; Carleton et al., 2004) were used separately 172 for each colony. Keeling plots are biplots of time-repeated measurements (at t_0 and t_{24}) with the 173 δ^{13} C values on the y-axis and $1/[CO_2]$ on the x-axis. In the biplot, the y-intercept of the regression line between t₀ and t₂₄ represents the case in which the CO₂ concentration is infinitely high and the 174 175 ambient CO₂ is negligible. With the log-transformed data, an additional linear model with sugar or 176 species (in lab or field, respectively) as the predictor was built.

177 According to the two-endpoint mixing model by Engel et al. (2009), the δ^{13} C values in the 178 breath of an ant is:

179
$$\delta^{13}C_{breath} = p(\delta^{13}C_A) + (1-p)(\delta^{13}C_B)$$
 (eq.1)

180 where p is the proportion of C3 or C4 sources in the diet and the endpoints (A and B) are the isotopic signatures of ants fed exclusively with either C3 or C4 sugars, respectively. The latter were 181 obtained by averaging the pure isotopic signatures estimated by the intercepts in the Keeling plots. 182 This formula was used to produce the values expected for each species using different proportions 183 of C3 or C4 sources. This formula was also used to calculate the δ^{13} C pure signatures for the three 184 species of ants used in the second experiment. The average ant pure signatures were then compared 185 to the threshold values obtained from eq. 1 using t-tests to identify species specialists for either C3 186 or C4 plants. We defined specialists as those consuming more than 90% of either C3 or C4 sources 187 188 in their diet (p > 0.9 in eq. 1).

190 **Results**

For the lab experiments, the mixed effects models showed no difference in CO₂ concentrations at t₀ 191 $(F_{2.9.76} = 3.04, P = 0.09)$ as well as no difference in the δ^{13} C-CO₂ values ($F_{2.12.71} = 0.36, P = 0.71$). 192 At t₂₄, a significant difference was found for CO₂ concentrations ($F_{2,20} = 35.32$, P < 0.001), with 193 significant differences between controls and each treatment (P < 0.001 for both) but no differences 194 between cane and beet treatments (P = 0.97). Overall, the δ^{13} C-CO₂ values differed among groups 195 (F_{2, 13,27} = 181.21, P < 0.001; Figure 1). No differences between controls and cane treatment were 196 found (P = 0.84), while the δ^{13} C-CO₂ values measured in beet treatments significantly differed with 197 respect to controls and can treatment (P < 0.001). The average pure δ^{13} C-CO₂ signatures of C3 and 198 C4 sugar were estimated to be -24.56‰ and -11.41‰, respectively. The linear model of the Keeling 199 plot intercepts revealed significant differences between the two treatments ($F_{1,17} = 485.74$, P < 100200 0.001). 201

The linear model for the wild ants showed no difference in CO_2 concentrations at t₀ (F_{3,11} = 202 0.95, P = 0.45) as well as no difference in δ^{13} C-CO₂ (F_{3,11} = 1.84, P = 0.20). At t₂₄, a significant 203 difference was found for CO₂ concentrations ($F_{3,11} = 24.90$, P < 0.001), with all ant species 204 differing from the controls (P < 0.001). The mean CO₂ production rate per g of dry weight was 205 1282.50 µl h⁻¹ g⁻¹ for *C. scutellaris* (mean individual rate: 12.83 µl h⁻¹ g⁻¹), 2888.75 µl h⁻¹ g⁻¹ for *L*. 206 neglectus (mean individual rate: 28.89 µl h⁻¹ g⁻¹) and 598.29 µl h⁻¹ g⁻¹ for M. capitatus (mean 207 individual rate: 8.55 μ l h⁻¹ g⁻¹). Significant differences in δ^{13} C-CO₂ were found at t₂₄ (F_{3,11} = 38.10, 208 P < 0.001; Figure 2). All species differed from controls (P < 0.001 for all), although no difference 209 among species was found. Similar results were obtained from the linear model of the Keeling plot 210 intercepts among species ($F_{2,9} = 3.41$, P = 0.08). 211

212	Using the iso	otope mixing model, we identify	fied the threshold values for	C3 or C4 specialists,	
213	which were δ^{13} C-C	$CO_2 > -12.73\%$ (C4) and δ^1	13 C-CO ₂ < -23.25‰ (C3).	Intermediate values	
214	indicated generalist	feeders. We found no C4 spec	ialists (Figure 3), and all spe	cies had much more	
215	negative δ^{13} C-CO ₂	values. Both C. scutellaris and	d <i>L. neglectus</i> had δ^{13} C-CO ₂	values not different	
216	from the expected t	hreshold value for C3 special	ists (t-test $P > 0.7$ for both),	with some colonies	
217	beyond that threshold. On the other hand, M. capitatus significantly differed from this threshold (
218	= -5.5331, <i>P</i> = 0.01	2) and all colonies were beyon	nd that value (Figure 3), sug	gesting an exclusive	
219	reliance	on	C3	sources.	

221 Discussion

To the best of our knowledge, this is the first application of a breath test on social insects and one of 222 few on insects overall (Miller et al., 1985; Engel et al., 2009). One of the main achievements of this 223 study was that the methodology used for analysing the carbon isotopic composition of breathed CO₂ 224 can easily be applied to small organisms like ants. Indeed, the isotopic signatures of *Crematogaster* 225 ants fed with either C3 or C4 sugars were recorded to be significantly different after 24 h in the 226 respiration chambers, demonstrating that the experimental setup was efficiently able to allow 227 228 discrimination of distinct dietary carbon sources. The technique also provided insights into the feeding choices of the three species of ants collected in the field. 229

For the lab experiments, 24 hours proved to be a sufficient time for the ants to accumulate 230 significant amounts of CO₂ in the respiration chambers relative to the control jars. After 24 hours, a 231 significant difference in the isotopic signatures of ants fed with the two sugars was found and their 232 δ^{13} C-CO₂ values were in line with those expected based on the known isotopic content of the two 233 sources. C3 plants produce sugars with a mean isotopic signature of -28‰, while C4 plants 234 synthesise sugars with a mean δ^{13} C-CO₂ of -13‰ (Ehleringer & Cerling, 2002; Fry, 2006). 235 Significant differences between the two groups at t₂₄ were also found when comparing the pure 236 signatures estimated by the Keeling plots, with these estimates close to the reference values 237 commonly reported in the literature for C3 and C4 (e.g. Fry, 2006). Unfortunately, the isotopic 238 signature of the air in the control chambers, which reflected the composition of the air in the lab, 239 was close to that of C4 sugar, and this is likely the reason why the two groups did not show 240 significant differences. To prevent this issue, we recommend that future experiments are carried out 241 by using pure chromatographic air to amplify the differences among treatments and controls as 242 much as possible. 243

The results from ants collected in the field showed that the CO₂ production rate was 244 consistent with the data reported in the literature and negatively correlated with worker dimensions, 245 as previously found by Mason et al. (2015). Moreover, in our study site, there was no evidence of 246 niche partitioning based on the carbon pathway, with all sampled species sharing similar resources. 247 This result is confirmed by the pure breath signature estimates. As pointed out by Perkins & 248 Speakman (2001), finding a strong difference in carbon signatures in the field can be difficult, 249 250 especially for generalist animals like ants. However, though field-collected ants did not show a clear reliance on different carbon sources (i.e. C3 vs. C4 pathways) as found in other wild animals 251 (Voigt, 2009; Voigt et al., 2013), the two-endpoint mixing model revealed further information on 252 253 ant feeding preferences. Indeed, we could not classify any of the studied species as a selective C4 254 consumer (Engel et al., 2009), but the degree of reliance on C3 plants seems to vary across species. Both C. scutellaris and L. neglectus did not exceed the C3 threshold, suggesting that they use both 255 256 C3 and C4 sources. Interestingly, however, some variability among colonies was present, as some of them seemed to be C3 specialists, while others evidently consumed some sources from the C4 257 pathway. Indeed, both species were collected on C3 trees (Quercus sp.) on which they were feeding 258 on aphid honeydew, but predation on other insects is common, especially for C. scutellaris (Frizzi 259 260 et al., 2016). Conversely, Messor capitatus, a seed-eating ant, showed a complete reliance on C3 261 plants for its diet. It is important to stress that in our study we did not characterised in detail the resources available to each colony, and therefore we cannot say how much the observed differences 262 reflected an active choice or the local availability. 263

Studying omnivorous bats, Voigt et al. (2008a) pointed out that δ^{13} C of breath and tissues could present a discrepancy as a result of isotopic routing, i.e. the different destinations that macronutrients have in the consumer body. While breath signature is indicative of metabolism, tissue signature indicates the sources used for tissue synthesis. This implies that species from lower trophic levels (only one C source, from plants) will have similar δ^{13} C values in the tissues and breath, while omnivorous species (two C sources, from plants and from animal prey) will have

higher (more positive) δ^{13} C values in the tissues than in the breath, the latter underestimating the 270 271 protein contribution to the diet. Therefore, breath signatures are more related to the carbohydrate portion of the diet. This could explain the similarity found between L. neglectus and C. scutellaris 272 breath. Although C. scutellaris preys on other insects, its main carbohydrate source is the plant-273 274 derived aphid honeydew, as well as for L. neglectus (Frizzi et al., 2016, 2018). One important point to carefully consider when dealing with the analysis of breathed CO₂ is that the δ^{13} C values reflects 275 the catabolised substrate. If an animal is fasted and uses fat stores, its breath will reflect the carbon 276 signature of the diet from which reserves were formed, which do not necessarily coincide with its 277 more recent diet (Hatch et al., 2002a; Welch et al., 2006). Unfortunately, the nutritional status of the 278 279 ants collected in the field was not known and, although there were no reasons for them to be fasting, 280 it is possible that differences in their reliance on C3 resources are affected by different uses of fat stores (Hatch et al., 2002a; Welch et al., 2006). At the same time, it is important to stress that the 281 small differences observed could also be due to different availabilities of C3/C4 sources around the 282 nests of the sampled colonies. 283

The applicability of the method to this taxonomic group was facilitated by its social 284 behaviour, which allows the handling of groups of individuals instead of single ants, providing 285 286 information on the colony as an ecological unit (Lach et al., 2010). Indeed, given that a sufficient amount of breathed CO₂ must accumulate in the respirometric chamber and that a non-negligible 287 volume of air must be sampled for the analysis, this approach can more easily process pools of 288 individuals from the same colony rather than the breath of single small animals. In conclusion, we 289 demonstrated that despite the important limitation of providing only one variable, breathing tests 290 could provide useful information on the trophic behaviour of omnivorous generalists like ants. This 291 study represents a starting point for further investigations of insect breath-particularly in 292 herbivores, but also in other ants in tropical ecosystems, where C4 plants are more abundant, to 293 294 better understand their trophic ecology.

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468 **Figure legends**

Figure 1. CO₂ concentrations (A) and carbon isotopic signatures (B) for controls and treatments
(beet and sugar cane) after 24 hours. In each plot, the groups having different letters were
significantly different after Tukey post hoc test.

Figure 2. CO₂ concentrations (A) and carbon isotopic signatures (B) for controls and ant species
(*Crematogaster scutellaris, Lasius neglectus* and *Messor capitatus*) after 24 hours. In each plot, the
groups having different letters were significantly different after Tukey post hoc test.

475 Figure 3. Estimated pure breath signatures of ant species (*Crematogaster scutellaris*, *Lasius*476 *neglectus* and *Messor capitatus*). The horizontal red lines show the threshold values for C3
477 (continuous) and C4 (dashed) specialists.