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(Article begins on next page)

Mimicry as a diversification driver in ants? Biogeography, ecology, ethology, genetics, and morphology define a second West-Palearctic *Colobopsis* species (Hymenoptera: Formicidae)

ENRICO SCHIFANI^{1,†,*}, DANIELE GIANNETTI^{1,†}, SÁNDOR CSŐSZ^{2,†}, FILIPPO CASTELLUCCI³, ANDREA LUCHETTI³, CRISTINA CASTRACANI¹, FIORENZA A. SPOTTI¹, ALESSANDRA MORI^{1,‡}, DONATO A. GRASSO^{1,‡}

¹ Department of Chemistry, Life Sciences & Environmental Sustainability, University of Parma, Parco Area delle Scienze, 11/a, 43124 Parma, Italy

² MTA-ELTE-MTM, Ecology Research Group, Pázmány Péter sétány 1C, H-1117 Budapest, Hungary

Evolutionary Ecology Research Group, Institute of Ecology and Botany, Centre for Ecological Research, Vácrátót, Hungary

³ Department of Biological, Geological and Environmental Sciences – University of Bologna, via Selmi 3, 40126 Bologna, Italy

 † co-first authors

 \ddagger co-last authors

* corresponding author: enrico.schifani@unipr.it

ORCID NUMBERS: ES - 0000-0003-0684-6229; DG - 0000-0003-1622-1329; SC - 0000-0002-5422-5120; FC - 0000-0002-9944-2196; AL - 0000-0002-2986-721X; CC - 0000-0001-8802-9614; FAS - 0000-0002-4270-7229; AM - 0000-0001-6544-3038; DAG: 0000-0001-9334-4280.

Running title: Mimicry-driven diversification in Colobopsis ants?

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1 ABSTRACT

The West-Palearctic Colobopsis ant populations have long been considered a single species (Colobopsis truncata). We studied the diversity of this species by employing a multidisciplinary approach and combining data from our surveys, museum and private collections, and citizen science platforms. As a result, we have revealed the existence of a second species, which we describe as Colobopsis imitans sp. nov., distributed allopatrically from C. truncata and living in the Maghreb, Sicily and Southern Iberia. While the pigmentation of C. truncata is reminiscent of Dolichoderus quadripunctatus, that of C. imitans sp. nov. is very similar to Crematogaster scutellaris, with which C. imitans sp. nov. lives in close spatial association, and whose foraging trails it habitually follows, similarly to Camponotus lateralis and other ant-mimicking ants. The isolation between C. imitans sp. nov. and C. truncata seems to have occured relatively recently because of the significant, yet not extreme morphometric differentiation, and to mtDNA polyphyly. Both C. imitans sp. nov. and C. truncata appear to employ mimicry of an unpalatable or aggressive ant species as an important defensive strategy; this 'choice' of a different model species is motivated by biogeographic reasons and appears to act as a critical evolutionary driver of their diversification.

16	ADDITIONAL KEYWORDS: ADAPTATION – BATESIAN MIMICRY – COI MTDNA –
17	DISCRIMINANT-FUNCTION ANALYSIS – MEDITERRANEAN – MULTIVARIATE
18	STATISTICS – NORTH AFRICA – SIBLING SPECIES – SPECIATION.

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INTRODUCTION

Ants (Insecta: Formicidae) are a hyperdiverse group of organisms which counts about 13,860 species 31 (Bolton, 2021) and is extraordinarily successful in most terrestrial ecosystems (Hölldobler & Wilson, 32 1990; Gibb et al., 2017; Seifert, 2017). Such high diversification stems from several evolutionary 33 strategies and lifestyles, and enables even hundreds of different ant species to coexist in the same 34 habitat (Hölldobler & Wilson, 1990; 2008). However, only one or very few dominant species 35 generally characterize even the most species-rich ant communities: these species form very populous 36 colonies, with large, often permanent foraging trails, and they defend territories that may extend over 37 hectares (Hölldobler & Wilson, 1990; Andersen, 1995; 1997; Grasso et al., 1998; 1999; 2005; Arnan 38 et al., 2018). These ants are exposed to higher predation risk compared the ones whose workers forage 39 solitarily or in small groups, and are accordingly equipped with effective defensive mechanisms 40 41 (Buschinger & Maschwitz, 1984; Hölldobler & Wilson, 1990; Dornhaus & Powell, 2010; Seifert, 2018). Most ant species live in small colonies and forage solitarily or in small groups, only 42 occasionally form trails, and develop evasive anti-predatory strategies (e.g. Hölldobler & Wilson, 43 1990; Tautz et al., 1994; Andersen, 1995; Dornhaus & Powell, 2010; Helms et al., 2014; Larabee & 44 45 Suarez, 2015; Seifert, 2018; Grasso et al., 2020). Of the species that live in small colonies, the only ones armed with dangerous defences are some predatory ants which retain the primitive feature of a 46 47 powerful functional stinger to hunt (Buschinger & Maschwitz, 1984; Hölldobler & Wilson, 1990; Dornhaus & Powell, 2010). 48

Well-armed ant species, in particular the ones that build large colonies, are a very good model for 49 several mimicking organisms, mostly arthropods. Some of these mimics are myrmecophilous 50 organisms: commonly ant predators or parasites, they have adapted to live within or close to ant 51 colonies by relying on chemical or acoustic mimicry (e.g. Geiselhardt et al., 2007; Barbero et al., 52 2009; Cushing, 2012; Parker & Grimaldi, 2014; Parker, 2016; Scarparo et al., 2019). On the other 53 hand, myrmecomorph species resemble their ant model thanks to morphological and/or behavioural 54 adaptations (e.g. Komatsu 1961; Jackson & Drummond, 1974; Oliveira & Sazima, 1984; Oliveira 55 1988; Cobben, 1986; McIver, 1987; McIver & Stonedahl, 1993; Trjapitzin & Trjapitzin, 1995; Cassis 56 & Wall, 2010; Chandler, 2010; Durkee et al., 2011; Huang et al., 2011; Cushing, 2012; Pekár, 2014; 57 58 Corcobado et al., 2016; Pekár et al., 2017; Harvey et al., 2018; de L. Nascimento & Perger, 2018; Gnezdilov, 2019). The main aim of myrmecomorphism is predation avoidance: compared to the 59 mimics, models usually possess superior defensive mechanisms and are also more numerous. As 60 such, they are usually interpreted as Batesian mimics (e.g. Jackson & Drummond, 1974; McIver, 61 62 1987; Durkee et al., 2011; Huang et al., 2011; Cushing, 2012; Harvey et al., 2018), whose evolution 63 can be favoured by model abundance (Kikuchi & Pfenning, 2010).

30

Many ant mimics are ants themselves. Some are parasites (e.g. inquilines) that act similarly to 64 myrmecophilous organisms (Buschinger, 2009), relying on chemical adaptations to interact with the 65 host species. However, there are some free-living ant species that act as mimics of more aggressive 66 or dominant ant species and are therefore interpreted as Batesian mimics. Nonetheless, solid empirical 67 68 evidence to reject the alternative hypothesis of Müllerian mimicry (see Müller, 1879; Pasteur, 1982; Ritland, 1991) is rarely available (Ito et al., 2004; Wagner, 2014). In these species, chromatic mimicry 69 70 is the prevalent mechanism, while behavioural or morphological adaptations are more rarely 71 documented (Emery, 1886; Forel, 1886; Santschi, 1919; Gobin et al., 1998; Merril & Elgar, 2000; Ito et al., 2004; Ward, 2009; Gallego-Ropero & Feitosa, 2014; Powell et al., 2014; Pekár et al., 2017; 72 Rasoamanana et al., 2017; Seifert, 2019a). In addition, recurrent behaviour among mimicking ant 73 species is interspecific trail-following, which consists in the mimics regularly infiltrating into the 74 foraging trails of the model and may lead to parasitic behaviour with regard to food resources (Emery, 75 1886; Santschi, 1919; Gobin et al., 1998; Ito et al., 2004; Menzel et al., 2010; Powell et al., 2014). 76 Unlike specialized parasites (e.g. Visicchio et al., 2001; Buschinger, 2009; de la Mora et al., 2020), 77 no advanced mechanisms of chemical deception exist in most of these cases, so in the eventuality of 78 direct encounters the model species recognizes and attacks the mimic, which however is well-79 equipped to escape (Goetsch, 1942; 1951; Kaudewitz, 1955; Gobin et al., 1998; Ito et al., 2004; 80 Menzel et al., 2010). The vast majority of the hitherto well-documented cases come from the tropics: 81 mimics mainly belong to diverse lineages from the Formicinae tribe Camponotini (mostly 82 Camponotus Mayr, 1861), while their models are phylogenetically scattered, including 83 Ectatomminae, Myrmicinae 84 Myrmeciinae, (Crematogastrini and Stenammini) and 85 Pseudomyrmecinae. Only one case of ant-mimicking ant species is well-documented in the West-Palearctic zone, i.e. Camponotus lateralis (Olivier, 1792). It chromatically mimics the similar 86 87 Crematogaster species Cr. ionia Forel, 1911, Cr. scutellaris (Olivier, 1792) and Cr. schmidti (Mayr, 1853), and follows their trails (Emery, 1886; Baroni Urbani, 1969; Menzel et al., 2010; Wagner, 88 89 2014; Seifert, 2018; 2019a). The three species are closely related with neighbouring geographic 90 ranges and differ slightly chromatically (Blaimer, 2012). Interestingly, there seems to be a geographic 91 trend in the chromatic pattern of Ca. lateralis, allowing it to better resemble these three Crematogaster species in the regions of sympatric occurrence (Wagner, 2014; Seifert, 2019a). 92

The ant genus *Colobopsis* Mayr, 1861 (Formicinae: Camponotini), recently separated from *Camponotus* (Ward *et al.*, 2016), currently counts 95 valid species and 21 subspecies (Bolton, 2021). It is distributed across the Holarctic, Indomalayan and Australasian regions, and is most diversified in the latter two regions (Janicki *et al.*, 2016; Guénard *et al.*, 2017). *Colobopsis* species are usually arboreal ants that nest in dead wood, form small-sized colonies and behave timidly towards other ants

98 (Wheeler, 1904; Ward et al., 2016). In the Western-Palearctic region, Colobopsis truncata (Spinola, 1808) is the only recognized species of its genus. The queen caste was described by Spinola (1808) 99 100 from north-western Italy (Liguria region), while the other castes were described later (Dufour & Perris, 1840; Forel, 1874; Emery, 1916). Another species, Co. fuscipes (Mayr, 1853) was described 101 102 from Austria by Mayr (1853), but was later reclassified as a junior synonym of *Co. truncata* (Emery 103 & Forel, 1879). Colobopsis truncata is therefore considered to have a wide geographic distribution, 104 from the Caucasus to Iberia and from Central Europe to the Maghreb (Seifert, 2018; Janicki et al., 2016; Guénard et al., 2017). It is an arboreal-nesting species, as is typical of the genus, and it 105 preferably nests on broadleaved trees, where it lives in monogynous and often polydomous colonies, 106 rarely exceeding 500 workers (Seifert, 2018). Queens and soldiers are specialized for phragmosis, 107 and soldiers may also function as repletes (living containers of liquid food), seldom leaving the safety 108 109 of the nests they guard (Brun 1924; Goetsch, 1950; 1953; Seifert, 2018). Minor workers are usually active outside the nest during both day and night, forage solitarily, do not recruit nest mates to food 110 sources, and perform very quick evasive movements when encountering other ants (Seifert, 2018). 111

112 During field observations across Italy we encountered marked divergences between Colobopsis colonies: workers of some colonies resembled Cr. scutellaris and followed its trails, while others 113 resembled *Dolichoderus quadripunctatus* (Linnaeus, 1771), two species having remarkably different 114 115 appearance. This is reflected by several contradictory reports which however contain no comment on these incongruences. Forel (1874) first referred to Swiss ants by considering Co. truncata a Batesian 116 117 mimic of D. quadripunctatus. He later suggested that the two species show pre-adaptations to parabiotic nest-sharing (Forel, 1903). Zimmermann (1934) instead studied ants in Croatia and 118 considered the occasional relationship between Co. truncata and Cr. scutellaris similar to the one 119 120 between Ca. lateralis and Cr. scutellaris. Goetsch (1942) stated that in Spain Co. truncata behaves similarly to Ca. lateralis, following the trails of Cr. scutellaris, , yet is also chromatically very 121 122 different and shows no adaptation to mimicry. Baroni Urbani (1971) reported on a case of trailfollowing between a Co. truncata queen and a Cr. scutellaris trail from central Italy. More recently, 123 Tinaut (1991) claimed that in southern Iberia Co. truncata can be easily confused with D. 124 quadripunctatus during field surveys due to their similarity. However, working in the same area, 125 Carpintero et al. (2001; 2005) instead affirmed that Co. truncata is a visual mimic of Cr. scutellaris 126 and follows its trails, and even speculated that after nuptial flights Co. truncata queens specifically 127 choose trees occupied by Cr. scutellaris to found their colony. In reviewing the distribution of 128 Colobopsis in Iberia, García (2020) mentioned possible chromatic similarity of C. truncata with Cr. 129 scutellaris and its mimic Ca. lateralis, but also D. quadripunctatus. More recently, there have been 130 131 reports on trail-following by Co. truncata on Cr. scutellaris ants in Italy (mentioned by Seifert 2018,

no locality specified; Lake Garda according to Wagner HC, personal communication). Wagner (2019)
described a close association between *Co. truncata* and *D. quadripunctatus* in Austria (Vienna),
highlighted morphological and chromatic similarity between the two, and reported trail-following of *D. quadripunctatus* trails by *Co. truncata*.

We aimed to test whether the abovementioned diversity of traits represented intraspecific variation or indicated the existence of multiple *Colobopsis* species in the West-Palearctic. In order to address the different biological dimensions of this problem, we used a multidisciplinary approach which involves the description of the patterns of diversity within the traditional notion of *Co. truncata*. We relied on the principles of integrative taxonomy (Schlick-Steiner *et al.*, 2010), involving morphological, genetic, ecological, ethological and biogeographical data.

142

MATERIALS AND METHODS

143 We combined qualitative morphology through chromatic pattern evaluation, quantitative morphology 144 through the multivariate analysis of morphometric data and genetics in the form of mtDNA (COI) sequencing, ecological surveys to study the association between *Colobopsis* and its putative model 145 146 species and recorded ethological data to document cross-species trail following. Finally, we compared our results with the known biogeographic patterns of other ant species. We chose this quantitative 147 morphological approach as it is widely regarded as the most practical and reliable single source of 148 evidence for cryptic ant species delimitation, and as a cornerstone in integrative approaches on cryptic 149 species complexes of ants (Seifert, 2009; 2018; Seifert et al., 2014; Wagner et al., 2017; Steiner et 150 al., 2011; 2018; Csősz et al., 2020). Moreover, mtDNA sequencing represents a widespread and cost-151 effective method to gain preliminary information on species identification, biogeography and cryptic 152 speciation (Hebert et al., 2003; 2016; Ratnasingham & Hebert, 2007), which has developed into an 153 aid to myrmecological faunistic, biogeographic and taxonomic studies (Steiner et al., 2005; 2018; 154 Csősz et al., 2015; Seifert et al., 2017; Schär et al., 2018; 2020; Blatrix et al., 2020). Ecological data 155 156 on species associations and ethological data are rarely used in ant taxonomy but appear to be highly relevant to the specific case we are investigating, while biogeography is important to understand 157 species diversity. 158

For our morphological and molecular analyses, we gathered type material of *Co. truncata* and *Co. fuscipes* as well as additional non-type material of *Colobopsis* from the Mediterranean region, and relied on our own efforts and the generous contribution of colleagues to achieve a satisfactory geographic coverage. In particular, the type series of *Co. truncata* consists in a single queen with the label "*Polyergus* (?) | *F.ca truncata* | Spin. in Ligur. | Genova || 6571". This queen could be safely identified as the type since it is the sole *Colobopsis* queen in the Spinola collection at the Museo di 165 Scienze Naturali in Turin (Italy), and matched the description given by the author (Spinola, 1808). Concerning Co. fuscipes, at least two syntypes are stored in the Museum für Naturkunde, University 166 167 of Berlin (Germany), and their pictures are available on AntWeb (AntWeb.org, codes FOCOL2496 and FOCOL2497): these are labelled "Oesterreich | Coll. Rhd || Colobopsis fuscipes Mayr || Type || 168 169 29812 || GBIF-D/FoCol | 2496 | specimen + label | data documented" and "Oesterreich | Coll. Rhd || Colobopsis fuscipes Mayr || Type || GBIF-D/FoCol | 2497 | specimen + label | data documented". 170 171 Although the label is unlikely to be an original by Gustav Mayr (B. Seifert, pers. comm.), we deem their status as types credible. We also retrieved a worker labelled Co. fuscipes in Mayr's collection at 172 the Natural History Museum of Vienna, but with no explicit indication ensuring its type status. In 173 order to gather information on chromatic variation of Euro-Mediterranean Colobopsis, we relied on 174 AntWeb pictures, images from scientific papers or monographs (Glaser, 2009; Wagner, 2014; 2019; 175 Lebas et al., 2016; Seifert, 2018; Scupola, 2018; García, 2020; Salata et al., 2020; Tăuşan et al. 2020), 176 and on georeferenced photographs uploaded on citizen science platforms (iNaturalist.org, 177 biodiversidadvirtual.org) and on biodiversity-related Facebook groups. A complete list of the material 178 examined, their depositories and collecting data is available as a Supplementary Material file to this 179 paper. Ecological and behavioural data were obtained through field surveys across Italy. 180

181

PIGMENTATION: CHROMATIC MIMICRY

182 Preliminary observations highlighted that chromatic pattern provides the most evident difference between allopatric *Colobopsis* populations which resemble two different model ant species in Italy. 183 184 We preemptively described the two chromatic forms and investigated whether these were consistently able to represent Colobopsis diversity across the Mediterranean basin, and whether they occurred 185 186 intracolonially or sympatrically, and also checked for the possible existence of third forms. The two Colobopsis model patterns were pre-emptively established by observing ten workers per each form 187 (10 from Sicily and 10 from mainland Italy), and all subsequent investigations were carried out by 188 inspecting specimens of well-preserved pigmentation primarily belonging to the worker caste. 189 190 Queens and soldiers were also examined and identified only if closely resembling one worker model pattern, while males were not considered due to their very different pigmentation. Furthermore, to 191 192 better describe the differences between the two models, a ratio was calculated between head color and mesosoma color by taking dorsal pictures of specimens and calculating the average value of red 193 (RGB colors) between 5 randomly selected pixels of the head and 5 of the mesosoma via software 194 ImageJ (Schneider et al., 2012). Calculating a ratio rather than considering the absolute values greatly 195 reduces the variation produced by different light conditions and camera settings among different 196 197 pictures, thereby allowing comparison of pictures from various sources. The same ratio was also calculated for the two putative model species Cr. scutellaris and D. quadripunctatus. Chromatic ratios 198

were calculated on 20 workers per chromatic pattern or species from across their respective
geographic range using both directly inspected specimens and images from citizen science platforms.
Any differences were statistically tested by using the software R 4.0.3 and R Studio 1.3.1056 (R Core
Team, 2021), and employing an ANOVA test and subsequent Tukey Post-hoc test for pairwise
comparisons.

The visual examination to verify correspondence to either of the two chromatic models was conducted on 79 directly observed *Colobopsis* colonies (76 of which containing workers) plus images of 136 further specimens (including 76 isolated queens), for a total of 310 workers and 79 queens covering a total of 16 countries from across the W-Palearctic *Colobopsis* distribution (see supplementary material).

209 The two model patterns are defined as follows:

Cr. scutellaris-like pattern (CSL pattern): head, or head and anterior part of the mesosoma (rarely
most of it) uniformly red, rest of the body evidently darker and mostly black. White stripes or dots
on the second gastral tergite often absent or weak (present in 10% of the examined workers). See Fig.
1.

D. quadripunctatus-like pattern (DQL pattern): head, mesosoma and appendages from reddish to blackish (therefore chromatically more variable than the *Cr. scutellaris*-like model), head at least slightly darker than the mesosoma or less frequently concolour, gaster black. White stripes or dots on the second gastral tergite often present (80% of examined workers) and more obvious. Phragmotic heads of soldiers or queens are always reddish in their anterior, heavily sculptured part (approximately one half of the head), while the rest follows the same scheme of workers. See Fig. 1.

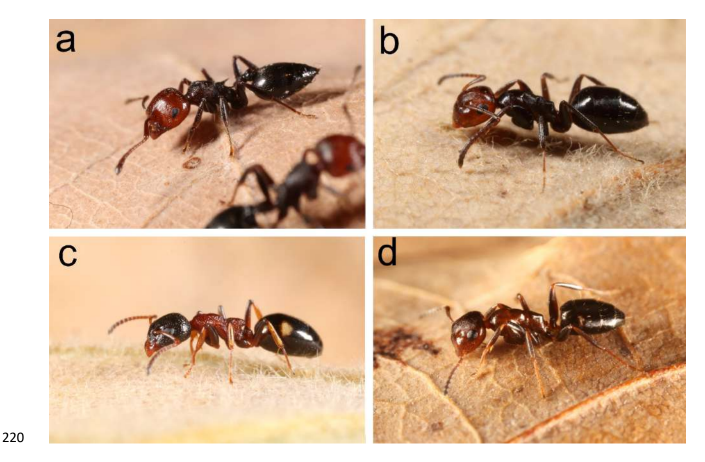


Figure 1. The model species and the two detected chromatic model patterns of *Colobopsis*: a) *Cr. scutellaris*; b) CSL
 Colobopsis from Sicily; c) *D. quadripunctatus* from Tuscany; d) DQL *Colobopsis* from Tuscany.

223 NUMERIC MORPHOLOGY: MULTIVARIATE ANALYSES OF MORPHOMETRIC DATA

A total of 12 continuous morphometric traits were defined following Seifert (2018) (Tab. 1) and 224 225 measured on 115 Colobopsis workers from 44 nest samples (considering minor workers only, and not the soldiers). All measurements were made in µm by using a pin-holding stage, which allowed 226 rotations around the X, Y, and Z axes. An Olympus SZX9 stereomicroscope was used at x150 227 228 magnification for each character; however, with characters larger than the field of view x75 229 magnification was applied. Due to the low number of the much rarer queens, males and soldiers in 230 our possession, we recorded only a reduced set of 7 morphometric traits aimed at providing a brief description of these castes without using them in the following statistical analyses. Morphometric 231 data are provided in µm throughout the whole paper. 232

Repeatability of the recorded size parameters were evaluated via Intraclass Correlation Coefficients (ICC) by using Package ICC (Wolak *et al.*, 2012), see Tab. 1. Variables were tested via matrix scatterplots and Pearson product-moment correlation coefficients for error variance and outliers. Each character resulted highly repeatable, except for NOL, which was considered moderately repeatable.

237 *Exploratory analyses through NC-PART clustering*

238 The prior species hypothesis was generated based on workers through combined application of NC clustering (Seifert et al., 2014) and Partitioning Based on Recursive Thresholding (PART) (Nilsen & 239 240 Lingjaerde, 2013). The script for NC-clustering combined with PART was written in R and can be found in Appendix S1 in Csősz & Fisher (2016). Our exploratory data analysis approach follows the 241 242 protocol described by Csősz & Fisher (2016) with the following specific settings: bootstrap iterations in PART were set to 'b=1000', and the minimum size of clusters was set to 'minSize=5' for both 243 244 'hclust' and 'kmeans'. The optimal number of clusters and the partitioning of samples are accepted as the preliminary species hypothesis in every case in which the two clustering methods, 'hclust' and 245 'kmeans' through PART, have yielded the same conclusion. 246

247 Exploratory analyses via PCA using allometrically corrected data

An alternative prior species hypothesis has been generated via the ordinating Principal Component
Analysis (PCA) that searches for discontinuities in continuous morphometric data and display plots
in a graphic.

Using raw data (without removal of allometric variance) in PCA may lead to weaker performance in 251 ordination because the first vector of the PCA often describes the size component, which is a useless 252 information when cryptic species of similar size have to be separated, hence, in PCA residuals were 253 used. Residuals, in which the head length (CL) was used as covariate, were calculated via a linear 254 regression model according to the following steps: a) scaling properties, intercept and steepness were 255 calculated for each nest sample separately (note: nest samples constituted by a singleton were not 256 257 involved in this phase); b) a grand average for steepness and intercept was calculated from scaling properties of each nest sample; c) residuals are calculated for every nest sample (including singletons) 258 259 based on the grand average. Residuals of every trait calculated against head length (CL) are given (Tab. 1). In contrast to NC-PART clustering, the PCA has no estimation on the number of clusters 260 261 and "classification" of objects has been made based on subjective decision. The coefficients (x any intercept) for removal allometric variance for each trait are given in supplementary material. 262

263 Hypothesis testing by confirmatory analysis

The validity of the prior species hypothesis was tested via Linear Discriminant Analysis (LDA). Classification hypotheses were imposed for all samples that were congruently classified by partitioning methods, while wild-card settings (i.e. no prior hypothesis imposed on its classification) were given to samples that were incongruently classified by the two partitioning methods. Statistical analyses were conducted through the software R 3.6.3 (R Core Team 2021).

CL	Maximum median length of head capsule. The head must be carefully tilted so the maximum length is positioned in the measuring plane.			
CW	0.951			
EL	L Eye length. Maximum diameter of the compound eye.			
dAN	Minimum distance of the inner margins of antennal socket rings.			
ML Diagonal length of the alitrunk in profile. Measured in lateral view from the anteriormost point of anterior pronotal slope to the caudalmost point of the lateral metapleural lobe.				
MW	Maximum width of pronotum.	0.989		
NOL Petiole node length; measured in lateral view, from the center of the petiolar spiracle to the posterior profile.		0.890		
PeW	Petiole width. The maximum width of petiole in dorsal view.	0.994		
PreOC				
SL	Scape length. The maximum straight-line scape length excluding the articular condyle.	0.971		
HTL				
PeSH Petiole scale height measured from the center of petiolar spiracle to 0. top of the crest.				
1 1 4 1 1		(n)		

Table. 1 Abbreviation (Abbr.) of morphometric characters, definition of measurements, and ICC (R) a metric for
 repeatability parameter are given. Definitions of morphometri characters follow Seifert (2018).

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GENETICS: MITOCHONDRIAL COI SEQUENCES

Total genomic DNA was extracted from leg tissues using the NuceloSpin DNA Insect kit (Macherey-272 Nagel, Düren, Germany), following the manufacturer's protocol. A 700 bp region of mitochondrial 273 gene cytochrome c oxidase subunit I (COI) was amplified using the primer couple 274 LCO1490/HCO2198 (Folmer et al., 1994). PCR was carried out in 25 µL reactions using the 275 following profile: initial denaturation step at 95 °C for 5 minutes, 35 amplification cycles 276 (denaturation at 95 °C for 30 seconds, annealing at 52 °C for 30 seconds, elongation at 72 °C for 45 277 seconds), final elongation at 72 °C for 7 minutes. PCR products were sent to Macrogen Europe 278 (Amsterdam, Netherlands) for Sanger sequencing. Chromatograms were checked and edited using 279 SeqTrace (Stucky, 2012). Sequences were aligned using the MUSCLE (Edgar, 2004) algorithm as 280 implemented in AliView (Larsson, 2014). Model selection and Maximum Likelihood phylogenetic 281 analysis were performed on the IQ-TREE web server (Trifinopoulos et al., 2016) using the Eastern-282 Palearctic Co. nipponica (Wheeler, W.M., 1928) and Co. shohki (Terayama, 1999) and the 283 Indomalayan Colobopsis nr. saundersi (Emery, 1889) (GenBank accession numbers AB019417, 284 AB019418 and KU975365, respectively) as outgroups. Ten separate runs were launched, each with 285 1000 replicates of ultrafast bootstrap, and the tree with the best likelihood value out of the ten was 286

chosen. Twenty-three colony samples were sequenced, consisting in 41 workers from 6 countries and
18 localities. Obtained sequences have been submitted to Genbank, under accession numbers
MW462045–MW462085 (see supplementary material).

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ECOLOGY: COEXISTENCE WITH MODEL SPECIES

Field surveys were conducted in the Italian Peninsula (Emilia-Romagna, Tuscany; n sites = 5, DQL 291 pattern) and Sicily (n sites = 8, CSL pattern) to test whether the local *Colobopsis* populations, showing 292 a DQL and a CSL phenotype respectively, actually lived in close proximity with either of the two 293 species indicated as probable mimicry models (see Supplementary material). In each site, we searched 294 for the presence of *Colobopsis* workers on trees until a tree occupied by a *Colobopsis* colony was 295 detected. Then, we performed a 10 minutes-long continuous sampling within a 1.5 m radius of the 296 297 point of the tree trunk where *Colobopsis* was firstly observed, recording the eventual presence of *Cr*. 298 scutellaris or D. quadripunctatus workers.

Occurrences of *Cr. scutellaris* or *D. quadripunctatus* on trees occupied also by *Colobopsis* according to the different DQL and CSL models were statistically tested by using the software IBM SPSS statistics, Italian version 24 and the chi-squared test. All data are presented in the Supplementary Material.

303

ETHOLOGY: INTERSPECIFIC TRAIL-FOLLOWING BEHAVIOUR

Field surveys were conducted in the Italian Peninsula (Emilia Romagna, Tuscany) and Sicily (sites 304 305 as in the section before, also see supplementary material) with the aim of quantifying the occurrence of trail-following behaviour performed by Colobopsis ants in relation to Cr. scutellaris or D. 306 307 quadripunctatus trails. We selected trees where Colobopsis colonies coexisted with either Cr. scutellaris, D. quadripunctatus or both. In accordance with the relevant literature (Gobin et al., 1988; 308 309 Ito et al., 2004; Menzel et al., 2010; Powell et al., 2014), trail-following was defined as the event of 310 Colobopsis workers moving along an established pheromone trail of Cr. scutellaris or D. quadripunctatus within 1 cm from the trail itself. A 10 minutes continuous sampling was used to 311 record the presence or absence of this behaviour on each of the examined tree. 312

To study trail-following on *Cr. scutellaris* trails, we selected a total of 59 trees inhabited by this species: 29 trees hosted *Colobopsis* colonies exhibiting the CSL pattern (Sicily, 4 sites) and 30 hosted *Colobopsis* with the DQL pattern (Emilia-Romagna and Tuscany, 5 sites). Observations on *D. quadripunctatus* trails could be performed only in 23 *Colobopsis* colonies exhibiting the DQL pattern (Emilia-Romagna and Tuscany): since no *D. quadripunctatus* colonies could be found in the studied sites in Sicily (where the species is known to be very rare, see Schifani & Alicata, 2018), no 319 *Colobopsis* colonies exhibiting the CSL pattern could be tested in this regard. All data are summarized
320 in the supplementary material.

321

SPECIES CONCEPT

Integration of the evidence provided by different complementary disciplines into an evolutionarily 322 credible species hypothesis is performed according to the principles emphasized by Schlick-Steiner 323 et al. (2010), i.e. resolving eventual disagreements by invoking solid evolutionary explanations. 324 Biogeography is here treated as an additional source of information, holding an important advisory 325 role to the formation of the final species-hypothesis. We abide by the universal Gene and Gene 326 Expression (GAGE) species concept proposed by Seifert (2020), which, although recently 327 formulated, convincingly summarizes the main theoretical and practical formulae most commonly 328 adopted during the last few decades as a rigorous approach on alpha taxonomy of cryptic ants, 329 330 especially in Europe.

331

RESULTS

332

PIGMENTATION: CHROMATIC MIMICRY

All the examined colonies are safely assignable to one of the two models and no transitional or third 333 forms are detected (see Supplementary Material). No intracolonial coexistence of the two models is 334 detected either. The type series of both Co. truncata and Co. fuscipes show the DQL pattern (Fig. 2). 335 The two models occur strictly allopatrically according to the examined material. Samples from the 336 337 south-western Mediterranean basin, namely Algeria, Sicily (Italy), Morocco, southern Portugal and Andalusia (southern Spain) are assigned to the CSL pattern. All the rest is assigned to the DQL 338 339 pattern, that is samples from Austria, Bulgaria, Croatia, Czech Republic, France, Germany, Greece, Hungary, Israel, Italian peninsula (Apulia, Campania, Emilia-Romagna, Liguria, Tuscany), Romania, 340 341 Serbia, Slovenia, Switzerland, the rest of Spain (Aragon, Catalonia, Balearic Islands, Castilla-La Mancha), and Turkey. In addition, photographs of Co. truncata specimens present in the European 342 ant fauna guides by Lebas et al. (2016) and Seifert (2018), regional faunistic guides by Glaser (2009) 343 (Liechtenstein), Wagner (2014) (Austria's Carinthia), Scupola (2018) (Italy's Veneto) as well as in 344 the Crete's ant fauna monograph by Salata et al. (2020) and in the papers by Wagner (2019) (Austria), 345 García (2020) (Spain) and Tăuşan et al. (2020) (Romania) all show the DQL pattern. 346



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Figure 2. Type material of the so far described West-Palearctic *Colobopsis*, all adhering to the "*D. quadripunctatus*like" pattern. a) holotype queen of *Colobopsis truncata* from Liguria, Italy, preserved at the Turin Natural History
Museum (Italy). b) syntype worker of *Colobopsis fuscipes* from Austria (picture from AntWeb.org, FOCOL2496;
photographer: Christiana Klingenberg), preserved at the Museum für Naturkunde der Humboldt-Universität Berlin
(Berlin, Germany). Note that the queen's red color in the anterior heavily sculptured part of the phragmotic head is not
relevant to evaluate its chromatic pattern. Scale bars: 0.5 mm.

The head red/mesosoma red ratio is statistically different among the two *Colobopsis* patterns and their models ($F_{3,76} = 152.4$, p < 0.001) (also see supplementary material). Pairwise comparisons show no statistically significant difference between CSL *Colobopsis* and *Cr. scutellaris* (p = 0.817; mean \pm sd $= 1.96 \pm 0.36$ for CSL *Colobopsis*; mean \pm sd $= 2.05 \pm 0.38$ for *Cr. scutellaris*) and between DQL *Colobopsis* and *D. quadripunctatus* (p = 0.299; mean \pm sd $= 0.60 \pm 0.27$ for DQL *Colobopsis*; mean \pm sd $= 0.43 \pm 0.20$ for *D. quadripunctatus*), while all other comparisons are significantly different (p< 0.001) (Fig. 3).

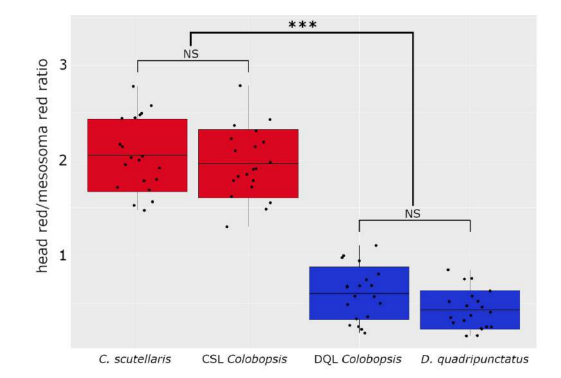


Figure. 3. Chromatic ratios calculated from pictures of the *Colobopsis* CSL and DQL patterns and from pictures of the two putative model species *Cr. scutellaris* and *D. quadripunctatus* (N = 20 for each species or chromatic form). Boxplots show mean and standard deviation, while whiskers represent minimum and maximum values. Dots are measured individuals. Their dispersal on the x-axis is a randomized graphical effect to avoid overlaps.

361

366 NUMERIC MORPHOLOGY: MULTIVARIATE ANALYSES OF MORPHOMETRIC DATA

Two morphological clusters are identified via NC-clustering combined with "kmeans", and "hclust" (Figs. 4). These two clusters correspond to the CSL pattern and DQL pattern specimens, respectively. All but two samples are congruently classified via both partitioning methods. The two incongruently placed samples (ITA:Mondello-VillaMercadante_col-12, ITA:Mondello-VillaMercadante_col-16; both CSL pattern from Sicily) are classified as belonging to the CSL cluster (posterior p = 0.85 and 0.81, geometric means of 3 workers each). Without running samples as wild-cards, the overall classification success is 96.3% using all variables in the analysis.

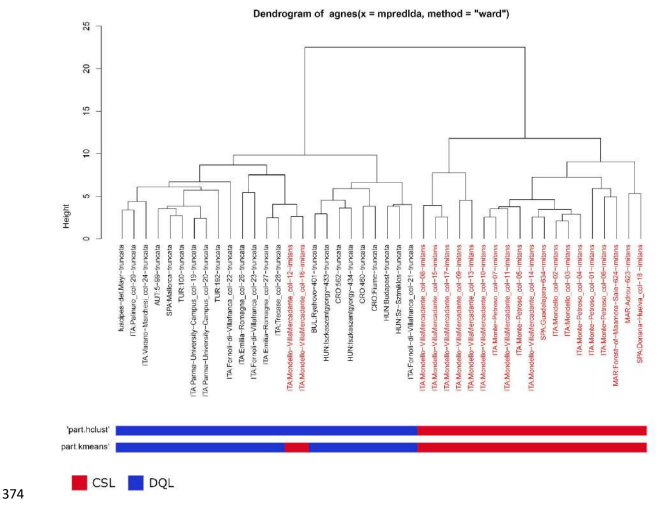


Figure 4. Figure 4. Dendrogram comparing the results of "kmeans", and "hclust" in NC Clustering of Colobopsis
morphometric raw data. Two samples (4.5% of the total) are misplaced by both the dendrogram and one of the
partitioning analyses, NC-part.kmeans; partially different samples being affected in each of the three analyses. The

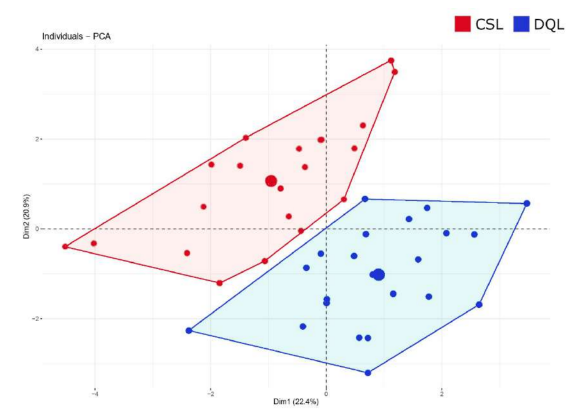


Figure 5. Principal Component Analyses of morphometric data of analyzed *Colobopsis* nest samples according to the
 two clusters evidenced by NC-PART Clustering. Each small dot represents a colony sample. Large dots represent
 centroids.

T-tests were calculated to assess significant differences (p) of body size ratios between specimens of the two different clusters, resulting in significant differences for 7 ratios (Tab. 2). Unfortunately, there is not a single numeric body size ratio available for reliable separation of these clusters on individual level (Tab. 2): the most reduced multivariate function that can reach the goal of attaining an acceptably high rate of classification success (>95%) requires a minimum 6 morphometric characters achieved via backward stepwise method.

- The most simple D(6) function that yields 4.3% of error rate at the individual level is as follows:
 D(6) = 0.03501 *CW 0.03384 * SL 0.03144 * HTL 0.01762 * ML + 0.03653 * PeSH + 0.07458 * EL + 16.61469
- 392 D(6) scores for CSL pattern cluster (n = 55) = mean 1.59 [-4.17, ± 0.66] ± 1.05
- 393 D(6) scores for DQL pattern cluster (n = 60) = mean $1.54 [-0.37, +3.76] \pm 0.96$
- 394

395

379

character	$\mathrm{CSL}\ (\mathrm{n}=55)$	р	DQL (n = 60)
CS	897 ± 51	0.924	898 ± 55
	[725, 1025]		[803, 1042]
PreOc/CL	0.546 ± 0.01	0.000	0.537 ± 0.01
	[0.517, 0.571]		[0.517, 0.570]
CL/CW	1.152 ± 0.02	0.052	1.145 ± 0.02
	[1.113, 1.198]		[1.106, 1.189]
dAN/CS	0.387 ± 0.01	0.128	0.391 ± 0.02
	[0.345, 0.431]		[0.358, 0.426]
SL/CS	0.872 ± 0.03	0.000	0.846 ± 0.02
	[0.798, 0.931]		[0.798, 0.906]
MW/CS	0.682 ± 0.02	0.476	0.680 ± 0.02
	[0.648, 0.717]		[0.648, 0.723]
PeW/CS	0.332 ± 0.02	0.257	0.336 ± 0.02
	[0.268, 0.372]		[0.306, 0.395]
HTL/CS	0.931 ± 0.02	0.000	0.902 ± 0.02
	[0.888, 0.978]		[0.845, 0.957]
ML/CS	1.485 ± 0.03	0.000	1.460 ± 0.02
	[1.395, 1.543]		[1.410, 1.526]
NOL/CS	0.139 ± 0.01	0.008	0.134 ± 0.01
	[0.117, 0.165]		[0.112, 0.157]
PeSH/CS	0.239 ± 0.02	0.000	0.253 ± 0.02
	[0.193, 0.284]		[0.203, 0.293]
EL/CS	0.312 ± 0.01	0.000	0.321 ± 0.01
	[0.290, 0.335]		[0.304, 0.343]

Table 2. Mean of morphometric ratios calculated for CSL pattern and DQL pattern clusters based on individuals (raw data). Morphometric traits are divided by cephalic size (CS), namely the arithmetic mean of CL and CW. The upper row in each data field gives arithmetic mean ± standard deviation, the lower one, in square brackets, lower and upper extremes.
Significant differences are highlighted in bold.

401

GENETICS: MITOCHONDRIAL COI SEQUENCES

402 The Maximum Likelihood phylogenetic analysis on mtDNA COI sequences identifies four main 403 clusters with good nodal support (Fig. 6). The clade A is formed by specimens exhibiting the CSL 404 pattern and collected from Andalusia (Spain) and Morocco. The specimens of the clade B exhibited 405 the DQL pattern and were sampled from Bulgaria, Castilla La Mancha (Spain), Hungary, the Italian Peninsula, and the Occitanic region of France. The clade C groups specimens with the DQL pattern,
sampled in the Spanish regions of Aragona and Catalonia, and from the Balearic Islands. Finally, the
clade D is formed by all specimens from Sicily, showing the CSL pattern, and one of the Spanish
specimens from Catalonia actually exhibiting the DQL pattern.

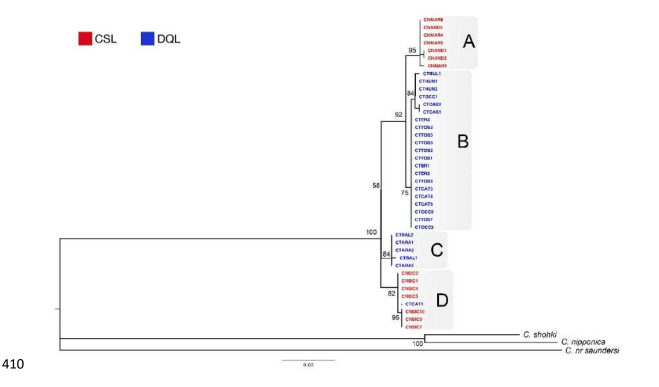
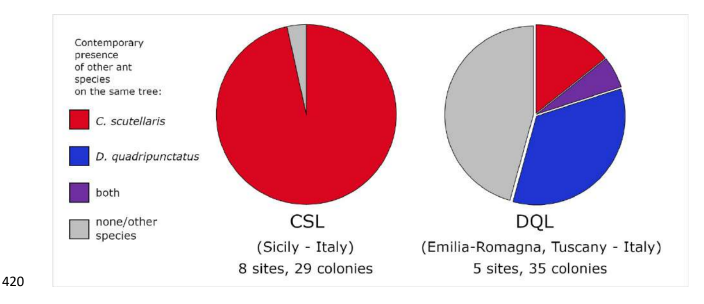


Figure 6. Maximum likelihood phylogenetic tree based on the barcode fragment of the mtCOI gene from the sequenced
 Colobopsis specimens.

413

ECOLOGY: COEXISTENCE WITH MODEL SPECIES

414 *Crematogaster scutellaris* is present in 97% of the investigated trees occupied by CSL *Colobopsis* 415 and in 20% of those occupied by DQL *Colobopsis*, the difference is statistically significant (χ^{2}_{1} = 416 26.23, p < 0.001). *Dolichoderus quadripunctatus* was never detected in trees occupied by CSL 417 *Colobopsis*. On the other hand, *D. quadripunctatus* occurrs on 40% of the investigated trees occupied 418 by DQL *Colobopsis* (6% of which also hosted *Cr. scutellaris*). Results are illustrated in Fig. 7, and 419 detailed data is provided in the supplementary material.



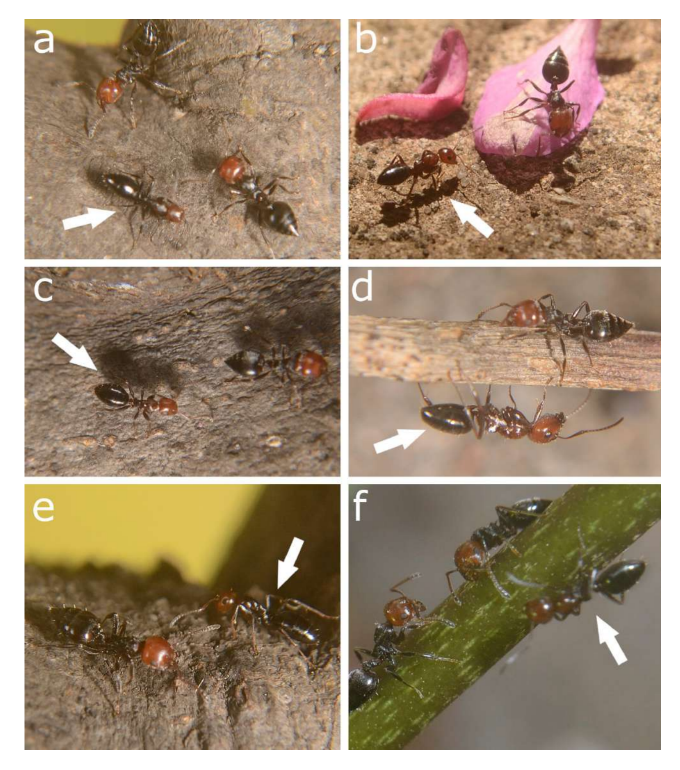
421 Figure 7. Coexistence between *Colobopsis* and their putative model species on the same tree.

422

ETHOLOGY: INTERSPECIFIC TRAIL-FOLLOWING BEHAVIOUR

During field observations, trail-following behaviour was never performed by *Colobopsis* colonies
exhibiting the DQL pattern (neither to *Cr. scutellaris* nor *D. quadripunctatus* trails). Conversely, 77%
of the observed *Colobopsis* colonies exhibiting a CSL pattern had workers following the *Cr. scutellaris* trails (Fig. 8). Detailed data are shown in the supplementary material.

427 CSL *Colobopsis* followed trails of *Cr. scutellaris* by either walking directly on them (more scarcely 428 populated worker trails with considerable gaps) or slightly sideways (crowded trails without or with 429 very small gaps only). If coming into contact with a *Cr. scutellaris* worker, they immediately 430 performed sudden accelerations and evasive movements. Trail-following often began a few moments 431 after the *Colobopsis* worker left its nest and encounter a *Cr. scutellaris* trail on the tree trunk and 432 ended with the *Colobopsis* worker leaving the trail and directing towards some specific twigs, no 433 longer following *Cr. scutellaris* workers.



434

Figure 8. Trail-following behaviour on *Crematogaster scutellaris* trails by other ants (indicated with arrows). On the
left (a,c,e) CSL *Colobopsis*, on the right (b,d,f) *Camponotus lateralis* observed in the same locality performing the same
behaviour (photos taken in Palermo (Sicily) during field surveys).

438 BIOGEOGRAPHY, EVIDENCE DISCUSSION AND FINAL SPECIES HYPOTHESIS

CSL and DQL chromatic patterns are found to effectively split into two the Mediterranean *Colobopsis*into two populations, occurring allopatrically and each covering a vast geographic region (Fig. 9).
The DQL pattern characterizes almost the entire European distribution of *Colobopsis* in addition to

442 Western Asia, while the CSL pattern occurs mainly in the Maghreb region (North-Western Africa), a well-recognized ant biodiversity hotspot (Borowiec, 2014), and in the European regions of greater 443 444 biogeographic proximity to it (Sicily and Southern Iberia) (e.g. Alicata & Schifani, 2019; Tinaut & Ruano, 2021). In particular, this distribution mirrors strikingly well those of some camponotine ant 445 446 species such as Camponotus barbaricus Emery, 1905, Ca. micans (Nylander, 1856) and Ca. ruber Emery, 1925 (Fig. 9; for their distribution see de Haro et al., 1996; Forel, 1890; 1905; Santschi, 1925; 447 448 Finzi, 1940; Menozzi, 1940; Cagniant, 1968; 1996; Collingwood & Yarrow, 1969; Baroni Urbani, 1971; Cagniant & Espadaler, 1993; Schembri & Collingwood, 1995; Janicki et al., 2016; Guénard et 449 al., 2017; Schär et al., 2020) and to a slightly lesser extent the distribution of myrmicine ants as the 450 Aphaenogaster crocea species group, A. sardoa Mayr, 1855 or the Temnothorax algiricus-451 mediterraneus complex (see Mayr, 1853; Emery, 1880; Forel, 1909; Santschi, 1929; Galkowski & 452 453 Cagniant, 2017; Alicata & Schifani, 2019). Ecological and behavioural field surveys across different Italian regions suggest that the two chromatic patterns are related to quite different lifestyles: the CSL 454 pattern often coexists with Cr. scutellaris and very often follows its trails, while the DQL pattern is 455 associated with D. quadripunctatus without the involvement of frequent trail-following. The CSL 456 457 pattern characterized specimens from the Spanish locality where mimicry, close nesting association and extensive trail-following of Cr. scutellaris were described by Carpintero et al. (2001; 2005). On 458 the other hand, the DQL pattern characterized samples from the Austrian region where Wagner (2019) 459 based his suggestions of close association and mimicry between Colobopsis and D. quadripunctatus. 460 461 A survey conducted in the region of Vienna (Austria) employing similar methodologies to ours estimated that 36% of the investigated *Colobopsis* colonies (n = 110) nested on trees occupied by D. 462 463 quadripunctatus, confirming the trend observed in our study (Wagner, pers. comm.). However, interspecific trail-following between DQL pattern Colobopsis and either D. quadripunctatus or Cr. 464 465 scutellaris as reported in Wagner (2019; personal communication) was never observed during our surveys and seems to represent a considerably less frequent phenomenon. Such differences between 466 467 the two groups in chromatic pattern, biogeography and life history traits could arguably be sufficient 468 to suggest a separation of the West-Palearctic Colobopsis into two species even according to a 469 conservative classical taxonomical approach. Moreover, examined specimens from the two chromatic 470 patterns are also classified into two morphometric clusters, whose separation reaches a significant threshold indicating heterospecificity according to the current procedures of cryptic ant species 471 separation (Seifert, 2020). At the same time, the morphometric separation between the two clusters 472 is relatively narrow, possibly indicating that the two species may have separated quite recently. 473 Concerning the mtDNA phylogenetic analysis, each clade is unambiguously monophyletic with 474 475 respect to morphometric and chromatic evidences (A and D = CSL pattern; B and C = DQL pattern),

with only one misplaced DQL specimen (a 2.4% error rate). On the other hand, with respect to 476 mtDNA, CSL and DQL patterns resulted in polyphy. This can be explained with possible retention 477 of ancestral polymorphisms and/or introgression of mtDNA (see Chan & Levin, 2005; Willis et al., 478 2013). These phenomena appear, in fact, largely responsible for the actual estimate of paraphyly 479 emerging from mtDNA phylogenies analyses in about 20% of animal species (Funk & Omland, 2003; 480 Ross, 2014). Mitochondrial DNA introgression is, like in other eukaryotic groups, quite frequent in 481 ants (e.g. Darras & Aron 2015; Beresford et al. 2017; Seifert, 2018), and coalescence during 482 483 speciation commonly results in species undergoing through phases of polyphyly and paraphyly – averagely longer in arthropods than in other groups – before normally reaching monophyly due to the 484 stochastic process of complete lineage sorting (Avise, 2004; Ross 2014). In the presently analyzed 485 taxa, this would support the hypothesis of the recent divergence. Due to their geographic origins, the 486 ambiguous placement of a few specimens during morphometric or genetic analyses also seems better 487 supported by this hypothesis than by hybridization (despite the latter being relatively frequent in 488 European ants, e.g. Steiner et al., 2011; Seifert, 2018; 2019b). 489

490 In conclusion, the CSL and DQL Colobopsis clusters are considerable separate species in accordance with the good practices of ants' alpha-taxonomy: all available sources of evidence suggest monophyly 491 with the exception of mtDNA, whose advisory role to infer species boundaries may be relatively 492 weak in comparison with nuclear genes or nuclear genes' expression products for the arguments given 493 by Seifert (2020). As a result, the formal naming of CSL and DQL Colobopsis species holds a key 494 495 informative value over their biology and life history traits. The type material of Co. truncata, consisting of a single queen, could not be part of the morphometric or genetic analyses, but shows 496 497 very clearly the DQL pattern and its geographic origin is unambiguous (with the type locality at mountains of Orero, near Genoa, in Italy's Liguria, placed in the middle of a highly investigated area 498 499 within the DQL Colobopsis geographic range and about 780 km away from the closest area inhabited by CSL Colobopsis). The same arguments of safe chromatic identification apply for Co. fuscipes, and 500 501 in this case they are supplemented by an even stronger biogeographic argument. As a result, the Colobopsis characterized by the CSL pattern is an undescribed species. Accordingly, a formal 502 503 description is provided below.

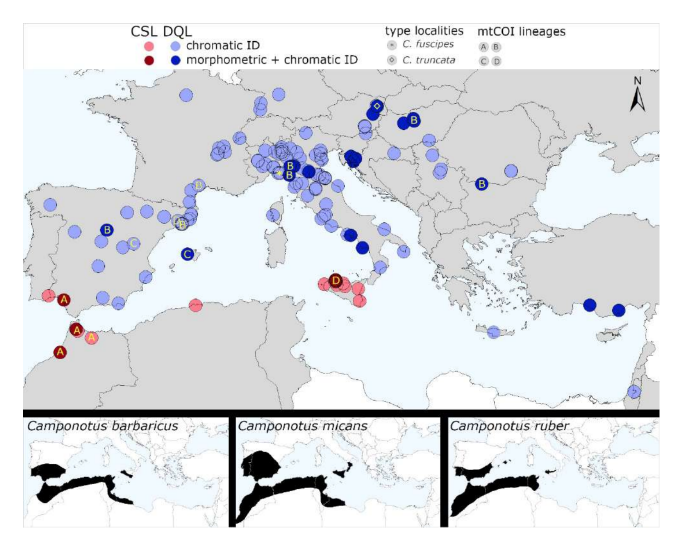


Figure 9. Above: distribution map of examined *Colobopsis* samples – countries where *Colobopsis* presence is known
 from literature are highlighted in grey. Below: approximate distributions of other Camponotini (*Camponotus barbaricus*, of *C. micans* and of *C. ruber*) which resemble that of CSL *Colobopsis*.

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DESCRIPTION OF COLOBOPSIS IMITANS SP. NOV.

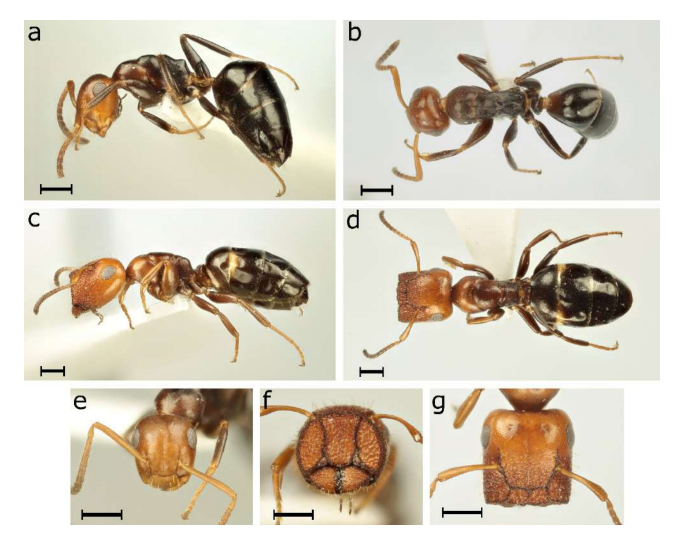
Etymology: imitans is the present participle of the latin verb imitor, meaning "imitating", and is here
used in apposition. It refers to the interpretation that this species resembles *Cr. scutellaris*.

Type series: 1 holytpe worker (Figure 10) and 14 paratype workers from Mondello, Sicily (Italy),
38.1953, 13.3354, 5 m, 14.X.2018, E. Schifani leg. The holotype is stored in the Hungarian Natural
History Museum collection.

514 Worker description: Morphometric indexes are shown in Tab. 2. Head subrectangular, on all sides 515 rounded. A straight, central furrow runs from the frontal triangle to the level at which the frontal 516 carinae end. Eyes large, ocelli extremely reduced. Antennae of 12 segments, without a distinct 517 antennal club. Pronotum significantly wider than the rest of the mesosoma. In lateral profile, 518 pronotum and mesonotum gently convex, propodeum profile often showing a central concavity thus

having a saddle-like appearance. Petiolar scale profile anteriorly roundly concave and posteriorly 519 straight, its dorsal crest excavated in frontal view. Promesonotal and mesoepinotal sutures as well as 520 metathoracic and propodeal spiracles well-visible. All legs with well-developed tibial spurs, but more 521 so in the anterior legs which are characterized by strikingly large femurs (identical to Co. truncata, 522 function unknown). Pigmentation as described in the CSL model. Very fine alveolate to areolate 523 sculpture covering the whole body and appendages. Few erect hairs near the posterior margin of the 524 vertex, between the frons and on the clypeus, and few others on the gaster tergites. See Figs 1, 8, 10, 525 13. 526

Soldier (= phragmotic major worker) description: Measurements (2 specimens from Sicily): CL 527 = 1327, 1419; CW = 1262, 1470; SL = 865, 942; ML = 1752, 1774; MW = 921, 1103; EW = 264, 528 331; EL = 409, 459; CS = 1294, 1444; CL/CW = 0.96, 1.05; SL/CS = 0.65, 0.67; ML/CS = 1.23, 529 1.35. Large phragmotic head with a cylindrical shape, and a flattened anterior part formed by the 530 mandibles, part of the clypeus and of the genae. In the distalmost half, it is characterized by a strong 531 areolate-rugose sculpture and a dense coverage of thick and short erect hairs. Rest of the shape, 532 533 sculpture and pigmentation generally similar to the worker but white dots or stripe on the first gastral tergite sometimes very evident. See Fig. 10. 534



535

Figure 10. *Colobopsis imitans* sp. nov.: a,b,e) worker (holotypus), c,d,f,g) soldier (specimen from the type
locality). Scale bars: 0.5 mm. Pictures also available on AntWeb.org database, specimen codes:
ANTWEB1041481 and ANTWEB1041482.

Queen description: Measurements (3 specimens from Sicily): CL = 1437-1531; CW = 1281-1406; 539 SL = 1156-1218; ML = 2687-3031; MW = 1281-1312; EW = 325-362; EL = 525-537; CS = 1359-540 1468; CL/CW = 1.08-1.11; SL/CS = 0.83-0.87; ML/CS = 1.93-2.22. Large phragmotic head very 541 similar to the soldier not only in shape but also in size (despite larger body size), but well-developed 542 ocelli, eyes much larger and much longer scapi. Immediately distinguishable by the larger, dorsally 543 flatter mesosoma, which is largely unsculptured and shiny. Propodeum profile similar to the end of 544 soldiers' propodeum. Head red as in the worker, but the mesosoma is brownish and the white dots or 545 stripe on the first gaster tergite are/is evident. See Fig. 11. 546

547 **Male description:** Measurements (3 specimens from Sicily): CL = 875-1093; CW = 781-1000; SL = 937-1001; ML = 2281-2437; MW = 1062-1218; EW = 300-387; EL = 462-525; CS = 828-1046; 549 CL/CW = 1.09-1.17; SL/CS = 0.92-1.20; ML/CS = 2.32-2.75. Small, subrectangular head with large 550 ocelli and very large eyes protruding laterally. Toothless mandibles. Relatively large mesosoma, propodeum more gently rounded than in queens or workers. Petiolar node very low and round.
Sculpture very weak, mesosoma shiny. Mandibles very hairy, other hairs on clypeus and gaster. Entire
body ferruginous or brownish, gaster blackish. See Fig. 11. Genitalia as in Fig. 12.

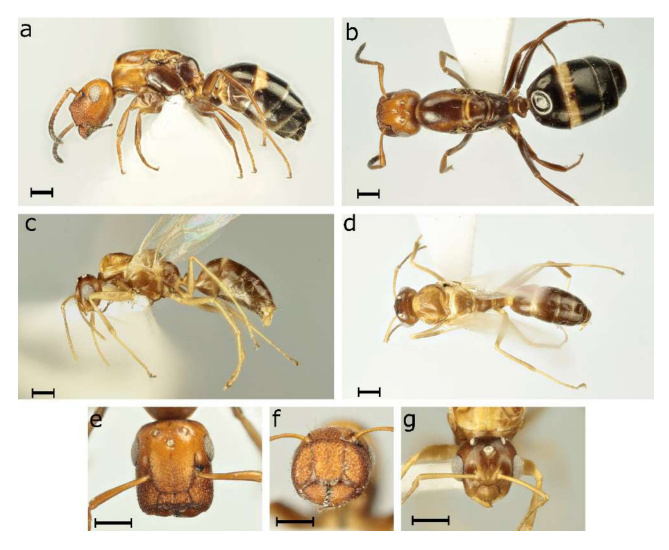


Figure 11. *Colobopsis imitans* sp. nov.: a,b,e,f) queen, c,d,g) male. Specimens from the type locality. Scale
bars: 0.5 mm. Pictures also available on AntWeb.org database, specimen codes: ANTWEB1041483 and
ANTWEB1041484.

554



Figure 12. Male genitalia of *Colobopsis imitans* sp. nov. in ventral and dorsal view, specimen from the type
locality. Scale bars = 0.25 mm.

561 Diagnosis: Generally easy to determine on the basis of worker material due to strong chromatic differentiation from Co. truncata and allopatric distribution (although further investigation is required 562 for possible contact regions in southern Iberia and southern Italy). Some very small worker specimens 563 may appear almost completely black, therefore lacking the typical chromatic pattern, and workers 564 with entirely red mesosoma can seldom be observed. Tentative identifications of isolated soldier or 565 queen specimens should be much more cautious, although their chromatic appearance may sometimes 566 567 appear to be very explicit. The low number of males and the lack of particularly evident distinctive characters from Co. truncata do not allow a safe species-level identification of this caste based on 568 morphology. In respect to workers, the morphometric linear discriminant function provided in the 569 results section should be helpful to determine dubious cases including decolored specimens. Finally, 570 571 DNA barcoding, which can be used for the same purpose and also be employed on the other castes, shows a relatively low error rate but may present some risks due to the polyphyletic pattern that we 572 observed. 573

Biological, ecological and phenological notes: Relatively thermophilous, in Sicily occurring from 574 few meters above the sea level to at least 1015 m a.s.l., in Morocco ascending up to 1290 m a.s.l. and 575 in Spain so far known from coastal lowland areas. Probably very common but also heavily under-576 recorded due to cryptic arboreal lifestyle, low colony population (most-likely monogynous), effective 577 578 mimicry and long periods of inactivity during the most arid and coldest times of the year. Workers and soldiers are unlikely to descend to the ground but were observed to do so at least once, following 579 580 a sparsely populated Cr. scutellaris trail. Soldiers in general are very difficult to be found outside the nest and usually seen acting as gatekeepers of the nest entrance. Founding queens were observed to 581 do the same. Nests are hidden in minute holes on the dead parts of arboreal trunks, where *Co. imitans* 582

583 sp. nov. often seems to act as a secondary user of cavities excavated by xylophagous insects. It exploits Andricus quercustozae (Bosc, 1792) oak galls as nests (occupying about 15% of galls 584 585 collected in Sicily's Bosco della Ficuzza in a recent survey, authors' unpublished data), in a similar way to Co. truncata (see Giannetti et al., 2019; 2021; Fürjes-Mikó et al., 2020). Polydomy appears 586 587 probable due to repeated findings of groups of workers without queens within oak-galls. Observed nesting on several and diverse plant species, including at least: Citrus reticulata Blanco, 1837, Ci. 588 589 sinensis (L.) Osbeck, 1765, Laurus nobilis L., Olea europaea L., Quercus ilex L., Q. pubescens-590 group, Q. suber L., Pyrus communis L., Pittosporum tobira (Thunb.) W.T.Aiton. Apparently dense populations were found in old Citrus orchards and relatively sparse Q. suber woods, but also in 591 deciduous oak forests. However, it occurs in a broad range of habitats from cities to agricultural lands 592 to natural forest habitats, but information available so far is insufficient to depict a satisfactory picture 593 594 of habitat preferences. Despite the earlier claim by Carpintero et al. (2005), there is currently no 595 evidence backing the fascinating hypothesis that C. *imitans* sp. nov. foundress queens prefer trees hosting Cr. scutellaris to found their colonies. A focused investigation on this topic would be 596 interesting. Nuptial flights for Co. imitans sp. nov. occur approximately in the same period of Co. 597 *truncata* (alates in Sicily observed from June 30 to July 13, n = 5, see supplementary material). 598 Winged queens and males were repeatedly seen attracted by artificial lights at night. 599

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FINAL REMARKS

Body pigmentation pattern is the only qualitative character that makes Co. imitans sp. nov. 601 602 identifiable without recurring to quantitative data, as it is otherwise morphologically extremely similar to Co. truncata up to a significant level of crypsis (see Wagner et al., 2018). These 603 604 pigmentation differences among West-Palearctic Colobopsis so far went completely unnoticed, the 605 sole exception being a brief statement by Santschi (1929) noting that the chromatic aspect of the 606 Moroccan Co. truncata is different than the typus one by its lighter head color. The case we 607 documented can be considered one of the few where such element is important for species discrimination in European ants. While body pigmentation has been used without scientific rigor by 608 some past ant taxonomists (see the example described by Boer, 2008), it can be important for the 609 morphological identification of species such as Formica clara Forel, 1886 and F. cunicularia 610 Latreille, 1798 or even fundamental for many Temnothorax spp. (Seifert & Schulz, 2009; Seifert, 611 612 2018) and should not be overlooked in multi-character approaches for taxonomic purposes. Under these conditions, checking pictures uploaded on citizen science platforms and social media proved to 613 be significantly helpful to obtain data on these species distribution, evidencing once more the 614 uncovered potentials of citizen science in the study of ant distribution (e.g. Lucky et al., 2014; Zhang 615

et al., 2019; Castracani *et al.*, 2020; Sheard *et al.*, 2020) and more in general of platforms hosting these kind of data in the study of insect distribution (e.g. Schifani & Paolinelli 2018; Hochmair *et al.*, 2020; Ruzzier *et al.*, 2020; Winterton, 2020). Moreover, behavioural data are seldom considered in integrative taxonomic approaches dealing with ants, but they may prove valuable in some cases (see also Ronque *et al.*, 2016). Finally, while mtDNA has a decent identification performance, our data clearly support the idea that it should not be used as the primary source of information to take taxonomic decision on species delimitation (see Seifert 2020).

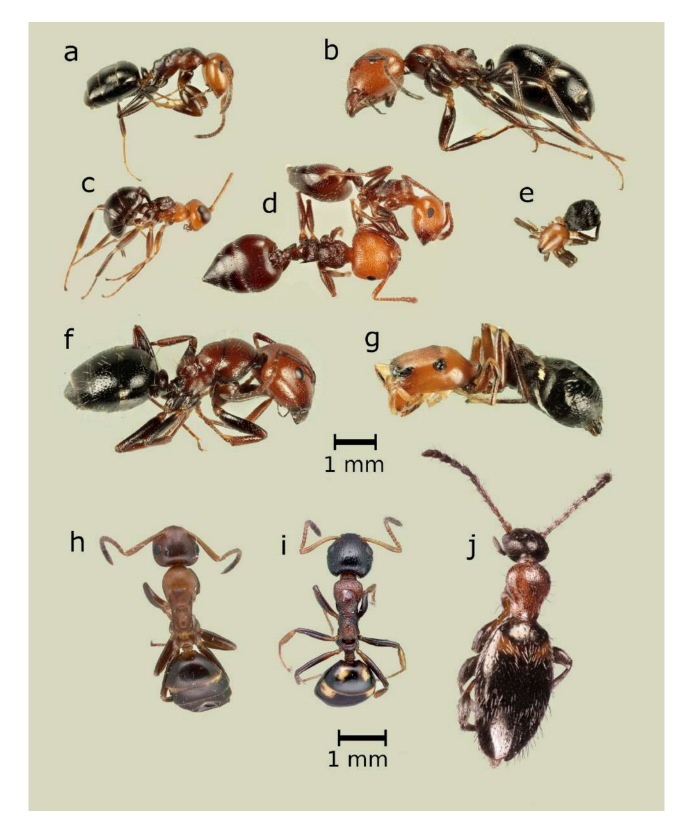
- 623 The taxonomic status of the West-Palearctic Colobopsis populations appears now well-resolved. Still, our analyses missed data from what the existing literature describes as the easternmost distribution 624 of Co. truncata east to the Mediterranean region, which reaches to the Kopet Dag in Turkmenistan 625 (Dlussky et al., 1990; Gratiashvili & Barjadze, 2008; Dubovikoff & Yusupov, 2018; Bračko, 2019; 626 Samin et al., 2020). In biogeographic terms, they are extremely unlikely to represent a disjunct Co. 627 imitans sp. nov. population, while conspecificity with Co. truncata appears likely due to the existence 628 of several ant species with similar distributions (e.g. Wagner et al., 2017; Seifert, 2018). Within the 629 630 Mediterranean, the range limits of Co. imitans sp. nov. and Co. truncata or their eventual sympatry 631 in contact zones should be appropriately investigated in areas of biogeographic transition (southern Iberia, Sicily, Calabria and perhaps Sardinia, see Alicata & Schifani, 2019; García, 2020; Schifani et 632 al., 2020; 2021; Tinaut & Ruano, 2021). 633
- The fact that Co. imitans sp. nov. and Co. truncata greatly differ chromatically is interesting if one 634 635 considers that phylogenetics and morphometry suggest a recent differentiation. In evolutionary terms, the most likely interpretation is to link such differentiation to a shared strategy based on ant-mimicry 636 637 modulated according to the presence or absence of certain good models across different Mediterranean regions. Both D. quadripunctatus and Cr. scutellaris have much more populous 638 colonies than Co. imitans sp. nov. and Co. truncata, while both are likely less palatable for predators 639 and armed with effective toxic substances (Cavill & Hinterberger, 1960; Wagner, 2019). Therefore, 640 641 even though only Cr. scutellaris is truly recognized as an aggressive and dominant species (Santini et al., 2007; Frizzi et al., 2015; Castracani et al., 2017; Seifert, 2018), both appear to possess the 642 required traits to be considered good Batesian models to the non-aggressive and relatively unarmed 643 Colobopsis (which still possess some formic acid). However, across the distribution range of Co. 644 *imitans* sp. nov., *D. quadripunctatus* is almost completely absent: it does not inhabit the Maghreb, its 645 Iberian distribution is concentrated to the North and in Sicily it is considered to bevery rare (Schifani 646 & Alicata, 2018; Cabanillas et al., 2019). Yet it is interesting to note that the opposite is not true for 647 648 *Co. truncata*: the latter is not only sympatric with *D. quadripunctatus* along its entire range (including in the hypothesis that easternmost Colobopsis are Co. truncata: see Reznikova, 2003; Ghahari et al., 649

650 2015), but also sympatric with Cr. scutellaris in south-western Europe and with Cr. schmidti in the east. Finally, it is worth noting that the white dots or stripe that have been linked to mimicry of D. 651 652 quadripunctatus in Co. truncata (Forel, 1886; Wagner, 2019) are/is absent or hardly visible in Co. *imitans* sp. nov. workers but at the same time well-visible in at least a few soldiers and especially 653 654 queens that we inspected. Following the mimicry interpretation of the chromatic patterns, it is 655 imaginable that this character is an ancestral remnant but that selective (predatory) pressures leading 656 to perfect mimicry are stronger on workers than on queens or soldiers which rarely leave the safety 657 of their nest.

Mimicry may be considered as a third defensive strategy of *Colobopsis* unique or very rare among 658 ants after suicidal authothysis and phragmosis (Emery, 1925; Maschwitz & Maschwitz, 1974; 659 Davidson et al., 2012; Shorter & Rueppel, 2012; Ward et al., 2016; Laciny et al., 2018). Apart from 660 the two species we treated, the only existing claims of mimicry in the genus come from 661 morphologically very different and likely unrelated species from Fiji Islands (Santschi 1928; Wheeler 662 1934). However, since several other Palearctic Colobopsis species share a general morphological 663 similarity with Co. imitans sp. nov. and Co. truncata, likely belonging to the same evolutionary 664 665 lineage, it is possible that some of them represent yet undiscovered mimics. It also appears that mimicry may have played a powerful role driving phenotypic diversification of West-Palearctic 666 667 Colobopsis: Co. imitans sp. nov. and Co. truncata can be considered as the only well-documented example among ants that suggests mimicry-driven phenotypic divergence of sister species. In 668 669 comparison, the intraspecific case of mimicry pattern divergence in Ca. lateralis is one of much more modest differentiation (Wagner, 2014; Seifert, 2019a). Similar accounts are not particularly common 670 671 in other organisms either, but recently an interesting scenario of strong diverging aposematic patterns 672 coupled with minimal genetic differentiation was described by for a group of frogs (Tarvin et al., 673 2017).

674 It is still unknown which visual predators may have been responsible for determining the selective pressures that lead to the emergence of ant-mimicry across different ant lineages. Birds and lizards 675 appear to be good candidates to start with (Ito et al., 2004; Wagner, 2014). Our surveys around the 676 colonies and trails of Cr. scutellaris and D. quadripunctatus led us to find several possible mimics of 677 either species that belong to different insect and spider groups already known for ant mimicry (Fig. 678 13). In particular, Santschi (1919) suggested in the the Canarian relative of Ca. ruber (Ca. guanchus 679 Santschi, 1908) the existence of an association similar to that between Ca. lateralis and Cr. 680 scutellaris, while Harvey et al. (2018) described the anti-predatory function of ant-mimicry in Gelis 681 682 spp., Komatsu (1961) reported on *Phrurolithus*-ant associations, Corcobado et al. (2016) reported on Leptorchestes-ant associations, and finally Chandler (2010) mentions myrmecomprhism among 683

Anthicidae. Although some of these findings may have been coincidental and deserve further 684 investigation, it is imaginable that Co. imitans sp. nov. and Co. truncata are each part of a larger 685 cohort of different arthropods that evolved mimicry to resemble Cr. scutellaris or D. quadripunctatus 686 in response to visually hunting generalist insectivores, similarly to the "golden mimicry complex" 687 described by Pekár et al. (2017). Further investigation is also required to understand whether the 688 advantages of mimicry for Co. imitans sp. nov. and Co. truncata may lay in a dilution effect, if 689 Batesian mimicry is truly implied and if Müllerian mimicry also plays a role (see Speed, 1999; Pekár 690 et al., 2017) –, keeping in mind that different evolutionary relations may exist between the same prey 691 692 and different predators.



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Figure 13. Above *Cr. scutellaris* and species showing a very similar chromatic pattern that were collected near *Cr. scutellaris* trails in Sicily (Italy): a) *Co. imitans* sp. nov. (worker from Mondello), b) *Ca. lateralis* (worker from Monte
Pellegrino), c) *Gelis* sp. (Hymenoptera: Braconidae) from Monte Petroso, d) *Cr. scutellaris* from Levanzo island, e) *Phrurolithus* sp. (Araneae: Phrurolitidae) from Mondello, f) *Ca. ruber* (worker from Monte Pellegrino) and g) *Leptorchestes* sp. (Araneae: Salticidae) from Monte Petroso. Below, *D. quadripunctatus* and species with a very similar
chromatic pattern that were collected near its trails or in the same trees in the Italian Peninsula: h) *Co. truncata* (specimen

from Bulgaria, AntWeb code CASENT0280000, photographer Michele Esposito), i) *D. quadripunctatus* (specimen from
 Czech Republic, AntWeb code CASENT0179916, photographer Michele Esposito), j) *Formicomus pedestris* (Rossi,
 1790) (Coleoptera: Anthicidae) from Parma (Italy).

Interspecific but intrageneric trail-following described for some ants is likely relatable to 703 phylogenetic proximity and morphofunctional and behavioural similarities or similar foraging 704 705 strategies among species (e.g. Grasso et al., 2002 and references therein). On the other hand, the significance of the recorded Colobopsis-Crematogaster trail-following is not yet fully clear to us. 706 707 Similar cases (such as that of *Ca. lateralis*) have often been referred to as parabiosis, but we avoided this term since quite different interpretations of its meaning coexist creating ambiguity: it is 708 709 sometimes used to simply indicate trail-following but in other occasions it implies also nest-sharing 710 (see Forel, 1898; Swain, 1980; Vantaux et al., 2007; Menzel et al., 2008; 2010; 2014a; 2104b; Seifert, 2018). Outside of Co. imitans sp. nov. and Co. truncata, many other camponotine ants follow 711 Crematogaster trails without always acting as mimics (Ito et al., 2004; Vantaux et al., 2007; Menzel 712 713 et al., 2008; 2014). Baroni Urbani (1969) and Menzel et al. (2014) speculated that the compounds 714 used as trails pheromones by Crematogaster are generally easily perceived by Camponotus, this capacity representing an important pre-adaptation to trail-following. During our field surveys, we 715 716 unexpectedly observed several workers of Camponotus piceus (Leach, 1825) (a relative of Ca. lateralis with no resemblance of Cr. scutellaris, see Seifert, 2019a) easily following part of a Cr. 717 scutellaris trail to the canopy of a tangerine tree while avoiding Cr. scutellaris attacks. It may be 718 possible that many other similar camponotine ants rarely perform the same without possessing a 719 specific mimicry adaptation, which can partly explain the occasional observations of trail-following 720 between Co. truncata and Cr. scutellaris (Zimmermann 1934; Goetsch 1942; Baroni Urbani 1969; 721 Wagner 2014). Still, in the overwhelming majority of the documented cases inter-specific trail-722 following is associated either with mimicry (as for Colobopsis imitans sp. nov., see Gobin et al., 723 1998; Ito et al., 2004; Menzel et al., 2010; Powell et al., 2014) or with nest-sharing (Vantaux et al., 724 2007; Menzel et al., 2008; 2014). For mimics, it appears to be primarily a way to better hide within 725 726 the ranks of the model species, an example of dilution effect (Lehtonen & Jaatinen, 2016), which may also apply to a certain degree non-mimic ant species as well. However, the trail-followers may be 727 728 able to obtain additional benefits in their success of locating trophic resources, sometimes even establishing somewhat parasitic relationships (see Vantaux et al. 2007; Menzel et al. 2010; 2014a; 729 2014b). It is unclear whether Colobopsis ants may also benefit from a similar mechanism although a 730 parasitic aspect of its trail-following behaviour has been suggested by Baroni Urbani (1969). 731

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