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## Army imposters

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- 10 Short running title: Diversification of myrmecophiles with hosts
- 11 Abstract

12 Colonies of neotropical army ants of the genus *Eciton* Latreille offer some of the most captivating examples of 13 intricate interactions between species, with hundreds of associated species already described in colonies of *Eciton* 14 burchellii Westwood. Among this plethora of species found with Eciton colonies, two genera of staphylinid 15 beetles, Ecitomorpha Wasmann, and Ecitophya Wasmann, have evolved to mimic the appearance and parallel the 16 colouration of the most abundant ant worker cast. Here, we study for the first time the association of these ant-17 mimicking beetles with their ant host in an evolutionary and population genetics framework. The central emphasis 18 is on colonies of *E. burchellii*, the only *Eciton* species that harbours both genera of ant-mimicking beetles. 19 Phylogenetic and population structure analyses using the same mtDNA COI region (802bp) for ants and beetles 20 indicated that specialization patterns of the myrmecophiles were congruent with specialization to a particular Eciton 21 (sub)species. Therefore, current taxonomic treatments of Eciton and its Ecitomorpha and Ecitophya associates 22 need revision. Molecular clock analyses suggested that diversification of the Eciton hosts pre-date that of their 23 guests, with a possible earlier association of *Ecitophya* (found with a large number of *Eciton* species) than with 24 Ecitomorpha (found only with E. burchellii colonies). Population-level analyses revealed that patterns of 25 diversification for the myrmecophiles are also consistent with specialisation to a particular host across broad 26 geographical areas but not at small geographical scales, with gene flow within each species found between host 27 colonies even across landscape features that are strong barriers for *Eciton* female-mediated gene flow.

28 Keywords

29 Gene flow; mimicry; mitochondrial DNA; myrmecophily; myrmecophory; population structure; speciation;

30 taxonomy

#### 31 Introduction

32 The study of associations between species is vital if we are to understand the evolution of biological diversity 33 (Thompson, 2013). This was recognised by Darwin in the closing paragraph of On the Origin of Species by Means 34 of Natural Selection (1859) when he used the term 'entangled bank' to refer to the interaction between species that 35 form biological communities, as highlighted by Thompson (1994). Nature is full of examples of intricate and 36 intimate associations between different species but it is in the world of ants where some of the most numerous and 37 astonishing associations can be found (Hölldobler and Wilson, 1990). Among the ants, army ants harbour the most 38 extensive array of species associations, with multiple vertebrate and invertebrate associate species exploiting the 39 ants and the different environments and homeostatic conditions that their colony life creates (Gotwald Jr., 1995; 40 Hughes et al., 2008). Of all army ants, those of the genus *Eciton*, inhabiting the tropics of the New World, are the 41 ones exhibiting the most captivating display of associates (Gotwald Jr., 1995), with 557 species already recorded 42 with Eciton burchellii and many more still to be described (Ivens et al., 2016; Rettenmeyer et al., 2011). Associates 43 found with *Eciton* army ants include, among others, mites that feed on secretions and hemolymph of the ants; flies 44 that feed on the middens' refuse; beetles that steal prey from the ants or predate on the ants or their brood; the 45 iconic army-ant-following birds that feed on arthropod prey flushed out by the ants during their raids; and 46 butterflies that feed on droppings from ant-following birds (Gotwald Jr., 1995; Kistner, 1979; Rettenmeyer et al., 47 2011; Schneirla, 1971)

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49 As part of this plethora of associates, some have evolved to conceal their presence among the army ants through 50 chemical, tactile and morphological mimicry (Gotwald Jr., 1995; Kistner, 1979). These 'imposters' have evolved 51 different strategies to associate with army ants, with staphylinid beetles having mastered the art of blending in 52 with the ants by evolving to resemble the appearance of their hosts (Hölldobler and Wilson, 1990; Kistner and 53 Jacobson, 1990; Maruyama and Parker, 2017). Two genera of myrmecomorph (ant-like) staphylinid beetles, 54 Ecitomorpha and Ecitophya, both in the subfamily Aleocharinae (tribe Athetini; Elven, Bachmann, & Gusarov, 55 2012), are found with neotropical army ants of the genus *Eciton*. These two genera of beetles are highly specialized 56 to the epigaeic patterns and nomadic life of *Eciton* army ants and both mimic the most abundant worker cast in 57 Eciton colonies (Seevers, 1965); the media workers (Franks, 1985). Both genera present similar morphological 58 modifications that confer resemblance to their host: similarity in surface sculpturing, subpetiolate and ellipsoidal 59 abdomens, slender heads and pronota, and long appendages (Seevers, 1965). Their appearance is considered 60 moderately ant-like as they are not a perfect mimic of the ants, but their colour parallels that of the species or 61 subspecies of *Eciton* with which they are associated (Akre and Rettenmeyer, 1966; Kistner and Jacobson, 1990; 62 Seevers, 1965). The two genera are mainly distinguished by *Ecitophya* presenting a much slender body and longer 63 appendages than *Ecitomorpha*, with a head more than twice as long as wide, a bilobed mentum, slender gula with 64 sutures not converging in the front, longer antennae (more than six times as long as the head width) but with the 65 terminal segments of the antennae not much wider than the preceding segments (terminal segments in *Ecitomorpha* 66 are more club-shaped) (Seevers, 1965).

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68 The mimicry of *Ecitomorpha* and *Ecitophya* to *Eciton* media workers is likely to be both an adaptive response to 69 avoid predators such as ant-following birds (Batesian mimicry), as well as an adaptation for integration into the 70 ant colony and avoidance of host aggression - Wasmannian mimicry (Parker, 2016). The presence of Ecitomorpha 71 and *Ecitophya* in *Eciton* colonies is rare (and sometimes absent), with many colonies presenting less than one of 72 these beetles per 1,000 worker ants. These two myrmecophile genera are considered hunting guests of *Eciton* army 73 ants, as they are found running among ants in raiding columns where they feed on dropped prey or at booty caches 74 (Kistner and Jacobson, 1990). These beetles are also found in emigration columns (Akre and Rettenmeyer, 1966; 75 Kistner and Jacobson, 1990) when the conspicuously nomadic Eciton colonies move to another location to set up 76 their new bivouac (temporary nest). During *Eciton* colonies emigration, *Ecitomorpha* and *Ecitophya* individuals 77 run in the centre of the columns or ride on prey captured by the ants or on ant pupae (Kistner and Jacobson, 1990). 78 Therefore, they are adapted to the movement and life cycle of their host (Akre and Rettenmeyer, 1966), as has 79 been shown for other *Eciton* myrmecophiles (Berghoff et al., 2009; Von Beeren et al., 2016a, 2016b).

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81 Taxonomically, the genus *Ecitophya* was initially divided into five species acknowledging the colour parallel 82 between this myrmecophile and its Eciton hosts (Reichensperger, 1933). Ecitophya rapaxae Mann found 83 associated with the ant Eciton rapax Smith, Ecitophya consecta Mann associated with Eciton vagans Olivier, 84 Ecitophya gracillima Mann that is associated with Eciton hamatum Fabricius, and Ecitophya simulans Wasmann 85 and Ecitophya bicolor Reichensperger associated with E. burchellii. The latter two species were later grouped into 86 a single species, E. simulans, as it was considered that specimens of Ecitophya collected with E. burchellii colonies 87 did not differ sufficiently to be considered as separate species (Kistner and Jacobson, 1990; Seevers, 1965). 88 Another species found associated with Eciton lucanoides Emery was later described as Ecitophya rettenmeveri 89 (Kistner and Jacobson, 1990).

90 Ecitomorpha beetles have only been found with the army ant Eciton burchellii, with the first specimens of 91 Ecitomorpha described by Wasmann in 1889 as Ecitomorpha arachnoides (Akre and Rettenmeyer, 1966). 92 Reichensperger divided this genus into four species taking into account the colour polymorphism within Eciton 93 burchellii: Ecitomorpha arachnoides Wasmann, Ecitomorpha nevermanni Reichensperger, Ecitomorpha 94 breviceps Reichensperger and Ecitomorpha melanotica Mann (Reichensperger, 1935, 1933). However, due to the 95 difficulty in finding consistent morphological characters (besides colouration) supporting the separation of these 96 species, they were subsequently lumped back into a single species, *Em. arachnoides* (Kistner and Jacobson, 1990; 97 Seevers, 1965).

98

99 In this study, we investigated for the first time in an evolutionary and population genetics framework the interaction 100 of Ecitomorpha and Ecitophya ant-mimicking beetles with their Eciton hosts; in particular, E. burchellii, as it is 101 the only Eciton species known to host both genera of beetles. Genetic analyses of ants and beetles collected in 102 Panama, a geographical area where many different *Eciton* species have overlapping ranges (Watkins, 1976), were 103 conducted to test the following hypotheses: (i) considering the strong level of association of Ecitophya and 104 Ecitomorpha with Eciton, phylogenetic patterns of the myrmecophiles will mirror that of their host, (ii) due to 105 the dependence of these two genera on *Eciton's* hunted prey and the pedestrian dispersal capability of the queen 106 and workers, the level of specificity between host and myrmecophile will be observable at broad geographical 107 scales, (iii) if myrmecophiles are truly host-specific and have evolved and diverged via increased specification on 108 a particular *Eciton* host, molecular patterns should support earlier taxonomic classifications of *Ecitophya* and 109 Ecitomorpha by Reichensperger (i.e. each Eciton species will host a particular Ecitophya and Ecitomorpha 110 species).

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112 Methods

#### 113 Study area and sampling

Sampling for this study was targeted on colonies of *E. burchellii* ssp. *foreli* Mayr and *E. b.* ssp. *parvispinum* Forel, the two most-studied *E. burchellii* subspecies. These two subspecies are highly epigaeic and their distribution ranges overlap in Panama, Costa Rica, and Honduras (Watkins, 1976). Descriptions of these species highlight their morphological similarity (Borgmeier, 1955; Santschi, 1925), with main differences reported being the colouration of media workers, *E. b. foreli* having black head and mesosoma but reddish metasoma, and *E. b. parvispinum*'s media workers having complete black bodies. However, studies assessing the genetic differences between these 120 two subspecies in view of their current taxonomic treatment have yet to be conducted. This study focused on 121 sampling of colonies in Panama, an area where the geographical range of both subspecies partly overlaps. Three 122 main areas of Panama were sampled (Fig. 1); in West Panama the Bosque Protector Palo Seco (BPPS) and the 123 adjacent Reserva Forestal Fortuna (RFF), and in Central Panama the Área Protegida San Lorenzo and its buffer 124 zone (APSL). As the Chagres River was found to be a barrier for E. b. foreli when gene flow was estimated with 125 mtDNA markers (Pérez-Espona et al., 2012), this area was divided into two (APSLA and APSLB) to group 126 colonies from each side of the Chagres River. In total, 13 colonies of E. b. foreli (4 in BPPS, 6 in APSLA and 3 127 in APSLB) and 12 colonies of E. b. parvispinum (all in RFF) for which we found associated Ecitophya and/or 128 Ecitomorpha beetles were sampled (Fig. 1; Table 1). In addition, four colonies of E. hamatum and the associated 129 Ep. gracillima: 1 colony in RFF, 2 colonies in BPPS, and 1 in Soberanía National Park (SOB), and one colony of 130 E. lucanoides and its associated Ep. rettenmeveri collected in RFF were opportunistically sampled. The number 131 of Ecitophya and Ecitomorpha beetles sampled and sequenced from E. b. foreli and E. b. parvispinum colonies 132 are summarized in Table 1. Although the number of beetles collected is not directly comparable between colonies, 133 as effort spent searching and collecting the beetles was constrained by the time of the day a colony was 134 encountered, the maximum number of beetles collected in a colony was 78 Em. arachnoides (colony E114 135 collected in RFF with E. b. parvispinum) and 40 Ep. simulans (colony E89 collected in BPPS with E. b. foreli).

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137 The sampling protocol for our study consisted of walking all available trails and adjacent less accessible areas 138 (off trails), through daily extensive walks (9 a.m. until dusk), to collect individuals from as many colonies of E. b. 139 foreli and E. b. parvispinum as possible. Once an E. burchellii colony was encountered, workers from all castes 140 were sampled from raid or emigration columns by removing them with the help of long forceps. Ant columns were 141 then carefully observed at several points of the raid or emigration columns to sample Ecitophya and Ecitomorpha 142 beetles with straight tube aspirators. Ant column observations per colony lasted several hours or until dusk, 143 depending on the time of the day when a colony was encountered. Collections were simultaneously conducted by 144 two people in order to maximise sampling of the beetles. All samples were preserved in 99% ethanol for further 145 examination and subsequent genetic studies.

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147 Laboratory procedures

148 DNA extraction

Abdomens of ants and beetles were carefully dissected before DNA extractions to avoid any contamination from consumed prey. To facilitate DNA extraction, the tissue samples were deposited in a 1.5mL Eppendorf tube and briefly immersed in liquid nitrogen prior to extraction procedures. Genomic DNA was extracted using the DNeasy tissue kit (QIAGEN) following the manufacturer's instructions.

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154 Sequencing of the mitochondrial marker Cytochrome Oxidase subunit I

155 Ants and ant-mimicking beetles from both genera, Ecitomorpha and Ecitophya, were sequenced for the same 156 region of the mitochondrial cytochrome oxidase subunit I gene (COI, cox1). Mitochondrial DNA markers are the 157 most widely used genetic markers in species-level phylogenies and DNA barcoding studies. Due to their haploid 158 nature, maternal inheritance, and smaller population size, sequences derived from mtDNA coalesce over a shorter 159 time scale than those derived from nuclear DNA (Simon et al., 2006), with reciprocal monophyly at the species 160 level reached faster after speciation in mtDNA phylogenies than in nuclear DNA phylogenies (Sunnucks, 2000). 161 The faster mutation rate of mtDNA markers has been shown to offer more powerful resolutions of relationships 162 between closely related taxa in phylogeographic and population-level studies (Avise, 2000; Zhang and Hewitt, 163 2003). In insects, mutation rates of mtDNA markers have been estimated to be 2 to 9 times faster than nuclear 164 protein-coding genes (Moriyama and Powell, 1997) making them more suitable for the study of closely related 165 species that have diverged recently (Lin and Danforth, 2004). COI fragments were amplified from the Eciton 166 samples using a modified version of the primer pairs CI13/CI14 (Hasegawa et al., 2002) and Ben/Jerry (Simon et 167 al., 1994). Details of the modified primers and Polymerase Chain Reactions (PCR) conditions can be found in 168 Pérez-Espona et al. (2012). Fragments of COI were amplified from Ecitomorpha and Ecitophya using the primer 169 pairs C1-J-1634/C1-N-2317 and C1-J-2216/C2-N-3431 (Maus et al., 2001). These beetle PCR amplifications were 170 conducted in a total volume of 25µL, using 10-15ng of template DNA, 1X NH4 buffer, 2.5mM MgCl<sub>2</sub>, 0.6µM of 171 each primer, 1 unit of BIOTAQ polymerase (Bioline, London) and double processed tissue culture distilled water 172 (Sigma-Aldrich, Buchs, Switzerland) to bring the volume up to 25µL. The PCR cycling protocol included an initial 173 denaturation step of 94°C for 3 min, a three-step cycling consisting of a denaturing step of 94°C for 30 s, annealing 174 at 51°C for 30 s and ramping at 0.3°C/s to an extension step of 72°C for 1 min. The cycle was repeated 29 times 175 and was followed by a final extension of 72°C for 10 min. PCR products were run on a 1.5% agarose gel and 176 visualised using ethidium bromide staining. Successful amplifications were purified using EXOSAP (GE 177 Healthcare), and forward and reverse strands for each of the fragments sequenced in two reactions using 6µL of 178 purified PCR product, 4µL of the reaction mix DYEnamic ET Terminator Cycle Sequence Kit (Amersham, GE

179 Healthcare) and 0.5µL of primer. Cycle sequencing consisted of 25 cycles including a denaturation step of 95°C 180 for 20 s, an annealing step of 50°C for 15 s and an extension step of 60°C for 1 min. Sequences were run on a 181 MegaBACE<sup>TM</sup> 1000 capillary sequencer (Amersham GE Healthcare) at The University of Bristol. 182 Electropherograms from the forward and reverse sequencing reads were edited and assembled into contigs using 183 the software Geneious version 10 (Biomatters: http://www.geneious.com). The resulting consensus reads from 184 each individual were sorted into unique haplotypes and subsequently manually aligned in Geneious, together with 185 some additional sequences obtained from four individuals of Eciton dulcium Forel (collected in RFF and APSL) 186 and sequences obtained from GenBank (Accession numbers: AY233691-4, AY233696, GQ980948). These 187 additional sequences were selected as ingroup placeholders and outgroups based on the army ant phylogeny of 188 Brady (2003) and the army ant-mimicking beetle phylogeny in Maruyama and Parker (2017). The resulting 189 alignments were all trimmed to include the same region COI fragment (802bp). All unique sequences were 190 submitted to DDBJ under the accession numbers LC258007-LC258019 for the Eciton sequences, and LC258020-191 LC258064 for the *Ecitophya* and *Ecitomorpha* sequences.

192

### 193 Phylogenetic and molecular clock analyses

194 Unique haplotype alignments for *Eciton* and *Ecitomorpha* with *Ecitophya* were initially evaluated for nucleotide 195 compositional heterogeneity using the Chi-square test in PAUP version 4.0b10 (Swofford, 2002), and using 196 tetrahedral plots and matched-pairs tests for symmetry implemented in SeqVis version 1.5 (Ho et al., 2006). The 197 more conservative Chi-square test provided no significant evidence for compositional heterogeneity in either of 198 the alignments. The more sensitive tetrahedral plots and matched-pairs tests for symmetry also provided no strong 199 evidence for compositional heterogeneity in the Ecitophya-Ecitomorpha alignment; however, there was evidence 200 for some heterogeneity in the *Eciton* alignment: tetrahedral plots contained co-dispersed clusters of data points 201 and the number of the matched-pairs tests for symmetry was >5% at P = 0.05 and >1% at P = 0.01. To explore 202 whether data re-coding could reduce the level of compositional heterogeneity in the Eciton alignment, C and T 203 nucleotides were re-coded as Ys. SeqVis analysis of this re-coded DNA alignment revealed no significant evidence 204 for heterogeneity. Therefore, in addition to analyzing the full Eciton DNA alignment, we also analyzed a data 205 reduced AGY form of the Eciton alignment to account for artefacts that may arise from compositional 206 heterogeneity.

208 Phylogenies were generated from the Eciton and Ecitophya-Ecitomorpha DNA alignments using Maximum 209 Parsimony and Bayesian methods. For the Maximum Parsimony analysis, heuristic searches were conducted using 210 PAUP, with the full Eciton and Ecitophya-Ecitomorpha alignments and the AGY re-coded Eciton alignment. Each 211 heuristic search started from a random tree with 50 random addition replicates, one tree was held per step, saved 212 trees set to a maximum of 10,000 and other settings left at default values. Confidence values were generated using 213 non-parametric bootstrapping (10,000 replicates). For the Bayesian inference, the DNA alignments were analysed 214 as a single partition using a mixed nucleotide substitution model and gamma corrected rate heterogeneity across 215 sites with the software MrBayes version 3.2.6 (Ronquist et al., 2012). Parameter distributions were approximated 216 using reversible jump Metropolis-coupled Markov chain Monte Carlo methods, with three chains of 10,000,000 217 generations, chain heating at 0.05, sampling frequency at 1,000 and other settings at default values. Posterior 218 samples of parameter estimates were assessed using generation plots, distribution plots, the potential scale 219 reduction convergence diagnostic and estimated sample sizes as recommended in the MrBayes manual. These 220 revealed that the total number of generations and default burnin of 25% appeared to be sufficient to acquire final 221 parameter estimates from a stationary distribution.

222

223 To analyse the *Eciton* alignment under AGY coding using Bayesian inference, we used the three-state 'AGY' 224 model implemented in the software mcmcphase within the PHASE version 3.0 software package 225 (https://github.com/james-monkeyshines/rna-phase-3). Heterogeneity across sites was modelled using a gamma 226 correction, and a chain length of 10,000,000 iterations, sampling period at 1,000 and burnin of 25% were used, as 227 in the MrBayes analyses. Perturbation proposal priorities were 10 for the tree and 1 for the substitution model. 228 Within these tree and model components, the proposal priorities were 1 for topology changes, 10 for branch lengths 229 (with an exponential (10) prior), 1 for frequencies, 1 for rate ratios, and 1 for the gamma parameter. Generation 230 versus log probability plots, parameter distribution plots, and repeated analyses starting from different random 231 seeds indicated that these settings generated final posterior estimates from a stationary distribution. Tree files 232 generated using mcmcphase were analysed using the associated program mcmcsummarize and also Geneious.

233

Chronological estimates for key diversification events were obtained with additional Bayesian analyses conducted using MrBayes. For these analyses, we reduced the level of haplotype sampling in order to include only the most abundant haplotypes and key biogeographic placeholders. This was to ameliorate the impact of intra-specific differences in haplotype sampling between the *Eciton* and *Ecitomorpha-Ecitophya* datasets that could have a 238 negative impact on the date estimation procedure. The molecular clock analyses used the same underlying model 239 and chain settings as described for MrBayes above with an outgroup-ingroup division enforced as a strong prior 240 topological constraint. Two different methods of clock calibration were explored: a standard 1% per million years' 241 rate applied to both the *Eciton* and *Ecitomorpha-Ecitophya* datasets, and fixed date calibrations of 26 million years 242 on the most recent common ancestor (MRCA) of Eciton following the results obtained by Brady (2003), and a 243 date of 25 million years on the Ecitomorpha-Ecitophya divergence following the results in Maruyama and Parker 244 (2017). Based on the data in Brady (2003) and Maruyama and Parker (2017), a uniform tree age prior of 10-100 245 million years was used. Clock model options were explored using stepping-stone sampling estimates of the 246 marginal likelihood. The Thorne-Kishino 2002 'TK02' relaxed clock had the smallest log likelihood but this was 247 less than 5 units better than the independent gamma rates 'IGR' relaxed clock and strict clock in both the Eciton 248 and Ecitophya-Ecitomorpha datasets. This preliminary analysis thus provided no strong evidence in favor of either 249 of these alternative clock models and, consequently, all three were used to generate date estimates. Posterior 250 samples of parameter estimates were assessed for stationarity as described above.

251

#### 252 Population-level analyses of E. burchellii and associates

253 Genetic diversity analyses for E. b. foreli and E. b. parvispinum colonies and their associated Ecitophya and 254 Ecitomorpha beetles were conducted using the softwares Arlequin version 3.1 (Excoffier et al., 2005) and DnaSP 255 version 5 (Librado & Rozas, 2009). Genetic diversity was estimated in terms of number of haplotypes, segregating 256 sites (S) and average number of nucleotide differences (k). In order to compare divergence within and between 257 species, S and k were also calculated for the different *Eciton*, *Ecitophya* and *Ecitomorpha* species included in our 258 study. Population structure for the host and each of the associates was estimated using a hierarchical analysis of 259 molecular variance (AMOVA) with the software popART (http://popart.otago.ac.nz). The partitioning of genetic 260 variation was assessed within and among two main geographical areas, West Panama (BPPS) and Central Panama 261 (APSLA, APSLB), and significance values obtained after 1,000 permutations. The software popART was also 262 used to build Median Joining haplotype networks (epsilon = 0) to assess haplotype relationships and identify 263 patterns of haplotype structure for *E. burchellii* and the myrmecophile beetles at different geographical scales.

264

265 Results

#### 266 Diversification and species relationships

267 Phylogenetic and molecular clock analyses provided further insights into the evolution and diversification of the 268 Eciton species and their Ecitophya and Ecitomorpha beetle associates (Figs. 2 & 3, Table 2 & 3). Analyses using 269 the full and reduced datasets using different methods (i.e. standard and AGY coded, Parsimony and Bayesian 270 analyses) resolved the same suites of well-supported haplotype relationships; this indicated that these were robust 271 to the nucleotide compositional heterogeneity found in the Eciton DNA alignment. With the Eciton dataset, 272 haplotypes of E. dulcium and E. hamatum, and the subspecies E. b. foreli, were grouped according to these 273 taxonomic categories with strong statistical support (≥ 99% Parsimony bootstrap support, Bayesian posterior 274 probabilities of 1.0). The subspecies E. b. foreli and E. b. parvispinum were also resolved as a single group but 275 with lower support ( $\geq 60\%$  bootstrap,  $\geq 0.91$  posterior probability). Other basal relationships between *Eciton* taxa 276 included in this study were poorly supported in all phylogenetic analyses. In the *Ecitophya-Ecitomorpha* analyses, 277 haplotypes assigned to *Ecitomorpha* and *Ecitophya* were clearly separated into these genera level categories ( $\geq$ 278 99% bootstrap, posterior probability of 1.0). The *Em. arachnoides* haplotypes formed two well-supported groups 279 that corresponded to their *E. burchellii* host ( $\geq$  98% bootstrap, posterior probability of 1.0); *E. b. foreli* and *E. b.* 280 parvispinum. The Ep. simulans haplotypes also consisted of two well-supported groups following their E. 281 burchellii host (100% bootstrap, posterior probabilities of 1.0). However, these two Ep. simulans groups formed 282 a complex with two groups of Ep. gracillima (associated with E. hamatum). Within this complex, Ep. simulans 283 (associated with E. b. foreli) was clearly grouped with Ep. gracillima (87% bootstrap, posterior probability of 284 0.95). Ep. rettenmeyeri (associated with E. lucanoides) was separated from the other Ecitophya species with 92% 285 parsimony bootstrap and 0.78 posterior probability support. Lineage relationships within Ecitophya, therefore, did 286 not mirror that of their *Eciton* host. The number of segregating sites (S) and the average pairwise nucleotide 287 differences (k) between species revealed further insights into the taxonomy and species relationships of the 288 myrmecophiles and their hosts (Table 3). Levels of divergence estimated as S and k between E. b. foreli and E. b. 289 *parvispinum* were similar (or slightly higher) to those between taxa currently recognised as separate *Eciton* species. 290 Although speciation patterns of *Ecitophya* did not mirror those of the hosts, a strong divergence was observed 291 between Ecitomorpha or Ecitophya associated with each subspecies of E. burchellii. Divergence of Ecitomorpha 292 was only assessed for two E. burchellii subspecies so further subspecies would need to be studied to elucidate 293 further speciation patterns.

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The molecular clock analyses recovered identical sets of *Eciton* and *Ecitophya-Ecitomorpha* haplotype relationships but with different associated date estimates depending on the clock model and calibration method

297 used (Table 2). Analyses calibrated using fixed dates taken from the literature were consistently older than date 298 estimates calibrated using a 1% rate, while differences arising from the use of different clock models were subtler. 299 Regardless of the clock model or calibration method, the relative age differences between Eciton and Ecitophya-300 Ecitomorpha lineage divergences were similar in all molecular clock analyses (Table 2, Figure 3). Although the 301 confidence intervals for the divergence estimates were wide and therefore we should be cautious with 302 interpretations, median estimates of divergences indicated that the diversification of the genus *Eciton* is likely to 303 be older than that of the associated myrmecophile genera Ecitophya and Ecitomorpha. This was the case when 304 considering date estimates for the MRCA of Eciton species taken from literature (used as a fixed 26 MYA 305 calibration date) and when considering independently obtained date estimates derived from the 1% calibration rate 306 (Table 2, Fig 3). Median estimates for diversification of the main *Eciton* species (or subspecies) also appears to 307 pre-date the diversification of the main myrmecophile lineages (Table 2, Fig 3). Poor branch support for most of 308 the basal *Eciton* relationships (Fig 2, Fig 3) may, in part, be due to the rapid diversification of these primary *Eciton* 309 lineages following the MRCA, as other studies using nuclear markers and phylogenomics have also failed to 310 unambiguously resolve these species relationships (see Discussion). Key among the main Eciton diversification 311 events was the MRCA of E. b. parvispinum and E. b. foreli, which pre-dated the diversification of the associated 312 myrmecophiles *Em. arachnoides* and the *Ep. simulans – Ep. gracillima* complex (Table 2, Fig 2, Fig 3). Among 313 the two genera of myrmecophiles, median estimates for the diversification of *Ecitophya*, which is found with more 314 species of *Eciton*, appear to be older than those for *Ecitomorpha*, although confidence intervals did overlap for 315 these estimates (Table 2, Fig 3). The genus *Ecitophya* has been reported in colonies of *E. rapax, E. vagans, E.* 316 burchellii, E. hamatum and E. lucanoides while Ecitomorpha has only been reported for E. burchellii.

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318 Population-level analyses of E. burchellii and its Ecitophya and Ecitomorpha associates

The diversification of the ant-mimicking beetles *Ecitophya* and *Ecitomorpha* with *E. burchellii* was further confirmed by the haplotype networks (Fig. 4). The networks clearly indicated a strong divergence between *E. b. foreli* and *E. b. parvispinum* haplotypes, with 90 segregating sites among haplotypes of both subspecies (Fig. 4). This divergence was mirrored in the associated beetles; however, the divergence between haplotypes in the beetles was smaller than that found for the host (55 segregating sites in *Ep. simulans* and 50 segregating sites in *Em. arachnoides*).

326 The haplotype networks confirmed that the diversification of Ecitophya and Ecitomorpha with E. b. foreli was 327 also prevalent over broad geographical areas. Haplotypes of ants and beetles collected in West Panama (BPPS) 328 markedly differed from those collected in Central Panama (APSLA and APSLB). Divergence time estimates for 329 the separation of haplotype lineages between West and Central Panama were older for E. b. foreli than for the 330 associate Ep. simulans (Fig. 3, Table 2). Although the haplotypes of Em. arachnoides found in the two main 331 geographical study areas were clearly distinct (Fig. 4), with haplotypes derived from two distinct lineages found 332 in West Panama (BPPS), the phylogenetic analyses did not support a strong grouping of haplotypes according to 333 these geographical areas (Fig. 2). In the Central Panama study area (APSL) the Chagres River was a strong gene 334 flow barrier for E. b. foreli females, with median estimates of divergence of E. b. foreli haplotypes either side of 335 the river estimated at 0.4-1.6 MYA (Fig. 3, Table 2). The Chagres River, however, was not a gene flow barrier for 336 either Ecitomorpha or Ecitophya, as haplotypes characteristic of a particular species of these myrmecophiles were 337 shared between their *Eciton* host colonies either side of the river (Fig. 4).

338

339 Estimates of population structure were higher for E. b. foreli than for the associated Ecitophya and Ecitomorpha 340 beetles (Table 4). In the host, genetic differentiation was found at all levels of analyses, with the majority of the 341 genetic variation explained by differences among groups (76.31%), and hardly any variation explained due to 342 differences within populations (0.20%). In Ecitophya and Ecitomorpha, most of the genetic variation was also 343 explained by differences among groups (84.62% and 79.57%, respectively) but genetic variation due to differences 344 within populations (15.51% and 22.03%, respectively) was much larger than that found for the host. Negative 345 values in variation among populations within groups found in *Ecitophya* and *Ecitomorpha* indicate the large 346 haplotype variation within populations and the sharing of haplotypes between these. No genetic variation was 347 attributed to differences among populations collected on either side of the river within each of the beetle species, 348 further corroborating that the Chagres River was not a gene flow barrier for these myrmecophiles.

349

#### 350 Discussion

351 Taxonomy of Eciton and associated Ecitophya and Ecitomorpha

The phylogenetic analyses and haplotype networks generated for this study revealed that current taxonomic treatments of *E. burchellii* and of its *Ecitophya simulans* and *Ecitomorpha arachnoides* associates need further revision to take into account further speciation within these taxa. Morphological descriptions of *E. b. parvispinum* 

355 have highlighted its black colouration, reduced spines in the metanotum and barely protruding epinotum teeth

356 (Borgmeier, 1955; Santschi, 1925), but these characteristics were considered not to warrant species status for E. 357 b. parvispinum. However, our results clearly indicate that genetic divergence between E. b. foreli and E. b. 358 parvispinum is as high as (or higher than) between taxa currently considered separate Eciton species. Divergence 359 estimates calculated in terms of S and k further suggest this speciation, as values between E. b. foreli and E. b. 360 parvispinum were higher than the those obtained between E. hamatum - E. dulcium and E. hamatum - E. 361 lucanoides; therefore, indicating that taxonomic treatments of E. burchellii need to be revised. Our results coupled 362 with those from a recent study phylogeographic study of this species in Central America (Winston et al., 2017), 363 provide strong evidence that speciation within this taxon has long been underestimated.

364

365 Although divergence between the *Eciton* species, and within *E. burchellii*, is clear from our genetic analyses, 366 phylogenetic relationships between *Eciton* species were not fully resolved in our study, despite the suitability and 367 widespread use of COI for resolving relationships between closely related insect taxa (Lin and Danforth, 2004; 368 Sunnucks, 2000). Weak statistical support for relationships between *Eciton* species does not appear solely reflect 369 our choice of COI as a molecular marker, because combined analyses of COI and nuclear markers (Brady, 2003), 370 and even a recent phylogenomic approach (Winston et al., 2017), also failed to unambiguously resolve these Eciton 371 species relationships. Rather, the lack of resolution in phylogenetic trees reported here is more likely indicative of 372 a rapid diversification of lineages within Eciton (Whitfield and Lockhart, 2007).

373

374 Important insights into the taxonomy of *Ecitophya* and *Ecitomorpha* were also revealed by our genetic analyses. 375 The taxonomy of these myrmecophiles has been challenging due to the subtle differences observed in 376 morphological characters, besides colouration, within each of these genera (Kistner and Jacobson, 1990; Seevers, 377 1965). The Ecitomorpha species key by Reichensperger was based on head dimensions, thoracic plates 378 sculpturing, length of the hind tibias and depressions behind the eves (Reichensperger, 1933). Further differences 379 between the species were highlighted in each of the species descriptions; these included different antennae 380 formation, elytra sculpturing, and colouration in different parts of the body of the specimens. Species 381 characterisation according to colouration was also indicated in previous descriptions such as that of Em. 382 melanotica found with E. b. parvispinum (Mann, 1926). Kistner and Jacobson (1990), after analysing a large series 383 of specimens, concluded that variation in colour was not consistent; however, they reported that black specimens 384 of Ecitomorpha were more frequently found in geographical areas where E. b. parvispinum was present, and rarer 385 in areas populated by E. b. foreli. In terms of other morphological features, Kistner and Jacobson (1990) did not 386 find consistency either in the shape of the groove of the pronotum or in spermathecas in the male genitalia that 387 would allow species differentiation. They, therefore, considered that early classifications of the Ecitomorpha 388 species by Reichensperger could not be supported. For Ecitophya, early species classification by Reichensperger 389 (1935, 1933) were based on the general appearance of specimens, measurements and characteristics of the head 390 and legs, differences in antennal segments, elytra sculpturing, presence and characteristics of abdominal bristles, 391 and the colouration of different body parts. In their analysis of *Ecitophya* specimens, that included dissections of 392 male genitalia, Kistner and Jacobson (1990) concluded that only some characteristics of the abdominal bristles, 393 the relative measurements of size of body parts and the relative length of antennal segments were reliable 394 characters for species identification. This reduction in the number of reliable characters to distinguish species led 395 Kistner and Jacobson (1990) to lump some of the earlier species described by Reichensperger into a single species, 396 E. simulans.

397

398 Our study has shown the importance of the use of genetic studies for resolving taxonomic challenges and has 399 provided strong evidence of speciation of *Ecitomorpha* and *Ecitophya* as a result of their specialisation to their 400 Eciton hosts. The speciation patterns found in our study for both myrmecophile genera (i.e. specialisation to an 401 Eciton host) would further support that E. b. foreli and E. b. parvispinum may be considered separate species. The 402 diversification patterns obtained for Ecitophya and Ecitomorpha were more concordant with initial species 403 classifications by Reichensperger (1935, 1933). Divergence estimated as S and k further supported the speciation 404 of Ecitomorpha and Ecitophya according to their host, with values between Ep. simulans found with E. b. foreli 405 or E. b. parvispinum differing by a similar number of segregating sites than between Ep. simulans found with E. 406 b. foreli and those found with Ep. gracillima (found with E. hamatum). Therefore, the current taxonomic treatment 407 of these Eciton burchellii associates as single species, i.e. Ep. simulans and Em. arachnoides, merits revision to 408 take into account the speciation patterns revealed by our genetic data. For Ecitomorpha beetles found associated 409 with E. b. parvispinum we suggest the adoption of the previous name of Em. melanotica Mann. Given our results 410 and those from Winston et al. (2017) supporting speciation of E. burchellii in Central America, we expect that 411 further species of *E. burchellii* and its *Ecitomorpha* and *Ecitophya* associates will be reported, as morphological 412 diversification of E. burchellii within its broad distributional range has long been acknowledged (Kistner and 413 Jacobson, 1990; Reichensperger, 1935, 1933; Seevers, 1965).

415 The lack of consistent and reliable morphological characters to differentiate species within Ecitomorpha and 416 Ecitophya reported by Kistner and Jacobson (1990) cannot be explained simply in terms of the recent and rapid 417 diversification within these genera, as indicated by our genetic analyses. In myrmecophiles, it is expected that 418 morphological variation between species may be reduced or absent due to strong selection on morphological, 419 behavioural and physiological characters as adaptations to exploit their hosts (Schonrogge et al., 2002). Often, 420 species are determined by visual inspection of morphological features; however, chemical and acoustic characters 421 can be more important for species recognition in arthropods. Future studies of Ecitophya and Ecitomorpha 422 diversification, therefore, will benefit from thorough analyses of chemical and acoustic characters, as these cues 423 are likely to be very important for their adaptation to their *Eciton* hosts (Lenoir et al., 2001).

424

### 425 Diversification of Ecitophya and Ecitomorpha with Eciton species

426 Phylogenetic patterns of Ecitophya and Ecitomorpha indicated specialization of these myrmecophiles with their 427 Eciton hosts, confirming previous observations of ant-resemblance and colouration parallels (Akre and 428 Rettenmeyer, 1966; Kistner and Jacobson, 1990; Reichensperger, 1933), and behavioural observations such as the 429 preference of following trails of the host species (Akre and Rettenmeyer, 1968). However, the patterns of 430 speciation of the myrmecophiles did not mirror those of the host revealing that the beetles' phylogenies were not 431 an 'evolutionary print' of the host (Thomas et al., 1996). Furthermore, phylogenetic analyses and estimates of S 432 and k disagreed with previously reported evolutionary relationships between  $E_{citophya}$  species based on 433 morphological characters (Kistner and Jacobson, 1990). Ep. simulans found with colonies of E. b. foreli were more 434 closely related to Ep. gracillima (associated with E. hamatum) than to the Ep. simulans found with E. b. 435 parvispinum. Additionally, Ep. rettenmeyeri (associated with E. lucanoides) was not closely related to Ep. 436 gracillima as previously inferred by the similar colouration between the ant hosts (Kistner and Jacobson, 1990); 437 in fact, Ep. rettenmeyeri was sister to the other Ecitophya.

438

Confidence intervals for time divergence estimates overlapped slightly between the ants and the myrmecophiles and, therefore, we cannot discard with certainty potential coevolutionary processes between hosts and guests. However, median time divergence estimates indicated that diversification patterns of the ants probably pre-dates that of the associated beetles. We acknowledge that our divergence estimates are derived from a single genetic marker and therefore we focus our discussion on *Eciton* and myrmecophile divergences on relative (rather than absolute) date estimates. Comparisons of median time divergence estimates from the molecular clock analyses for both beetle genera suggest that diversification of *Ecitophya* with *Eciton* ants is likely to have occurred earlier than for *Ecitomorpha*. A possible earlier diversification of *Ecitophya*, and subsequently a longer time frame to finetune their interaction with their host, could explain the association of *Ecitophya* with a larger number of *Eciton* species.

449

450 Myrmecophily in Staphylinidae beetles is an ancient phenomenon, with a fossil of *Protoclaviger trichodens* gen. 451 et sp. nov. (Clavigeritae) in amber being dated to the Early Eocene (c. 52 MYA; Parker and Grimaldi, 2014). This 452 early association of Clavigeritae beetles with ants can explain the remarkable diversity of myrmecophilous species 453 within this supertribe (Parker and Grimaldi, 2014). High species diversification due to myrmecophily has also 454 been reported for ant-nest beetles of the genus Paussus L. (subfamily Paussinae; Moore and Robertson, 2014). 455 Although new species of *Ecitophya* and *Ecitomorpha* are likely to be described, due to underestimated speciation 456 within E. burchellii (and maybe in other Eciton species), the diversification of these two Aleocharinae genera is 457 not as exceptional as that found for other myrmecophile Staphilinidae such as *Paussus*. This difference probably 458 reflects the high specialisation of Ecitophya and Ecitomorpha to the genus Eciton, in contrast to Paussus beetles 459 that are found associated with different, and sometimes distantly unrelated, ant genera (Moore and Robertson, 460 2014). Furthermore, our results suggest that contrary to other myrmecophile beetles found associated with Eciton 461 colonies (e.g. Vatesus), Ecitophya and Ecitomorpha species have evolved more host-specific adaptations, probably 462 as a result of stronger selection pressures because they are hunting guests of day-time raiding epigaegic army ants.

463

#### 464 Geographical patterns of diversification of Ecitophya and Ecitomorpha with E. b. foreli

465 Broad phylogeographic patterns of Ecitophya and Ecitomorpha were concordant with their E. b. foreli host, with 466 a clear separation of haplotypes between the West and Central Panama study areas. However, patterns of 467 diversification of the host and these two myrmecophiles differed at smaller geographical scales, indicating that 468 local differences might be the result of the spatial distribution of the host and the capability of dispersal of the 469 myrmecophiles (Tack and Roslin, 2010; Thompson, 2005). The haplotype networks revealed that Ecitophya and 470 Ecitomorpha beetles are not colony- or ant-mtDNA lineage-specific even though vertical transmission of 471 myrmecophiles is likely to occur during colony fission (Schneirla, 1971). Ant colonies with different mtDNA 472 haplotypes shared myrmecophile haplotypes, indicating that horizontal transmission of Ecitophya and 473 Ecitomorpha mtDNA lineages occurs between colonies. Horizontal transmission of myrmecophiles between 474 Eciton colonies has been previously reported for Vatesus (Akre and Torgerson, 1969), a beetle that does not mimic

475 the physical appearance of the ants but most of its life cycle is tightly linked to that of its host (Von Beeren et al., 476 2016a). Horizontal transmission of Ecitophya and Ecitomorpha individuals, as a potential strategy to avoid 477 inbreeding within a colony, might occur by dispersal through flight between colonies such as shown for Vatesus 478 (Chatzimanolis et al., 2004; Von Beeren et al., 2016a), or whenever colony fusion - the aggregation of workers to 479 another colony after losing their queen (Schneirla, 1940; Schneirla & Brown, 1950) - takes place (Kronauer et al., 480 2010). In our study, gene flow between the myrmecophile populations of each species was also found across the 481 Chagres River despite this being a major gene flow barrier for E. b. foreli females (Pérez-Espona et al., 2012). 482 Gene flow across the Chagres River, therefore, indicates a higher dispersal capability of the beetles in contrast to 483 the obligate pedestrian dispersal of *Eciton* queen and worker ants. Reports of flight in these beetles have only been 484 anecdotal and limited to observations of hovering of Ep. consecta over a colony of E. vagans when this colony 485 was spreading to attack the observer (Mann, 1921), and an individual of *Ep. gracillima* found with a colony of *E*. 486 hamatum (Pérez-Espona pers. obs.). In the latter case, hovering was observed when trying to aspirate one 487 individual from a raiding column. This specimen hovered to seek refuge under some fallen leaves but after a few 488 minutes tried to follow the ant trail (when it was successfully collected and included in this study). Fully developed 489 wings with venation characteristic of staphylinids have been described for both Ecitophya and Ecitomorpha 490 (Kistner and Jacobson, 1990). However, due to the close dependence of these beetles with their specific host and 491 the relatively low density of colonies, at least for E. burchellii (Franks, 1982), dispersal of the myrmecophiles 492 between colonies, in particular those located at further distance or separated by landscape features that act as 493 barriers for *Eciton* female dispersal, might be challenging. In such scenarios, it is likely that dispersal between 494 colonies is mediated by large and alate *Eciton* males when they leave their natal colony in search of conspecific 495 colonies to find a queen for mating (Gotwald Jr., 1995; Schneirla, 1971). Males of E. b. foreli are produced in 496 large numbers (c. 3,000) as part of sexual broods (including a small number of queens) when colony fission is 497 imminent (Franks and Hölldobler, 1987; Gotwald Jr., 1995; Schneirla, 1971). Males have been shown to be strong 498 fliers and able to disperse over 1km distances (Jaffé et al., 2009). They are therefore responsible for the majority 499 of gene flow between colonies (Berghoff et al., 2008; Jaffé et al., 2009; Pérez-Espona et al., 2012; Soare et al., 500 2014). Dispersal of myrmecophiles by ant alate reproductives have been shown in Atta leaf-cutting ants, with 501 Attafila cockroaches observed on ant queens departing for nuptial flights (Moser, 1967), and similar strategies 502 have been suggested for the movement of myrmecophiles between Eciton colonies (Kronauer et al., 2010).

504 Ecological and evolutionary studies of myrmecophiles of army ants at the population-level are still in their infancy 505 due to the difficulty of keeping the host and myrmecophiles in laboratory conditions (Kistner and Jacobson, 1990). 506 However, as shown in this and recent studies in Vatesus and Tetradonia beetles (Von Beeren et al., 2016a, 2016b), 507 genetic approaches can shed light on the evolution and levels of specificity of these army ant imposters. Our study 508 was based on variation in mtDNA therefore future studies aiming to further elucidate the level of association of 509 Ecitomorpha and Ecitophya, in particular at small geographical scales, would benefit through the use of highly 510 variable markers such as microsatellites. Using a combination of mitochondrial and microsatellite data in previous 511 studies has been demonstrated that deforestation has a major impact on the connectivity of E. burchellii 512 populations (Pérez-Espona et al., 2012; Soare et al., 2014), threatening the long-term persistence not only of these 513 top neotropical predators but also the multitudes of species associated with them. Further research on *Eciton* and 514 their associates is crucial if we are to provide conservation solutions that would guarantee the maintenance of this 515 manifestation of Darwin's 'entangled bank'.

516

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- 667 Figure legends
- Figure 1. Map of Panama indicating the main study areas. RFF = Reserva Forestal de Fortuna, BPPS = Bosque
  Protector de Palo Seco, APSL = Area Protegida de San Lorenzo and its buffer zone; SOB = Soberanía National
  Park. Colonies collected in APSL and its buffer zone were grouped in the analyses as APSLA and APSLB to
  reflect colonies collected on either side of the Chagres River.

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**Figure 2.** Majority-rule consensus phylograms for unique haplotypes derived from *Eciton* species (a) and the associated myrmecophiles *Ecitophya* and *Ecitomorpha* (b), generated with MrBayes using a mixed nucleotide model and gamma corrected rate heterogeneity. Support values are shown for nodes (marked by filled circles) found in  $\geq$  50% of parsimony bootstrap and posterior probability samples. Coloured bars indicate differences in abdomen (or whole body) colour of different taxa.

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680 Figure 3. All compatible groups consensus chronograms for the selected *Eciton* (a) and associated myrmecophiles 681 Ecitophya and Ecitomorpha (b) haplotypes, generated with MrBayes using a mixed nucleotide model, gamma 682 corrected rate heterogeneity, tk02 relaxed clock and fixed node calibration of 26 (Eciton) or 25 (associated 683 myrmecophiles) MYA (indicated by a star). Nodes are scaled to median date estimates with the 95% Highest 684 Posterior Density indicated by a translucent blue bar. Nodes with date estimates are labelled with roman numerals 685 and correspond to values shown in Table 2. Node support values are given for the presented chronograms, filled 686 circles indicate nodes found in  $\geq$  50% of parsimony bootstrap and posterior probability samples in the main 687 phylogenetic analyses. Coloured bars indicate the abdomen (or whole body)'s colour of different taxa.

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- **Figure 4.** Haplotype networks of *Eciton burchellii* (a) and associated *Ecitophya* (b) and *Ecitomorpha* (c) in the main study areas in Panama (RFF, BPPS, APSLA, APSLB) constructed using a median-joining approach. Study areas are indicated with different colours. The size of the circles is proportional to the number of individuals representing a particular haplotype. Missing intermediated haplotypes are indicated with black dots, nucleotide substitutions between haplotypes are indicated by small lines over the haplotype connecting branches.

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**Table 1.** Details of collection and sequencing of *Ecitophya* and *Ecitomorpha* myrmecophiles found with different *Eciton* army ant species in the study areas in Panama

			Individuals collected		Individuals sequenced		
Ant species	Location	Ant colony	Ecitomorpha	Ecitophya	Ecitomorpha	Ecitophya	
E. b. parvispinum	RFF	E85	6	2	3	2	
E. b. parvispinum	RFF	E86	4	12	2	0	
E. b. parvispinum	RFF	E87	1	0	1	0	
E. b. parvispinum	RFF	E94	3	8	3	4	
E. b. parvispinum	RFF	E95	23	8	4	2	
E. b. parvispinum	RFF	E96	2	0	2	0	
E. b. parvispinum	RFF	E97	12	5	3	2	
E. b. parvispinum	RFF	E99	8	20	2	6	
E. b. parvispinum	RFF	E100	13	10	4	6	
E. b. parvispinum	RFF	E108	15	4	2	1	
E. b. parvispinum	RFF	E109	58	28	1	5	
E. b. parvispinum	RFF	E114	78	32	8	8	
E. b. foreli	BPPS	E89	15	40	4	2	
E. b. foreli	BPPS	E101	21	2	2	2	
E. b. foreli	BPPS	E103	5	13	2	2	
E. b. foreli	BPPS	E104	7	12	2	2	
E. b. foreli	APSLA	E126	8	6	6	2	
E. b. foreli	APSLA	E127	9	3	2	2	
E. b. foreli	APSLA	E132	28	17	14	8	
E. b. foreli	APSLA	E143	14	6	13	5	
E. b. foreli	APSLA	E154	4	0	2	2	
E. b. foreli	APSLA	E156	8	4	2	1	
E. b. foreli	APSLB	E162	3	5	2	3	
E. b. foreli	APSLB	E165	2	3	2	2	
E. b. foreli	APSLB	E166	8	4	2	2	
E. hamatum	BPPS	E88	0	69	0	6	
E. hamatum	RFF	E105	0	13	0	3	
E. hamatum	BPPS	E110	0	8	0	2	
E. hamatum	SOB	E169	0	1	0	1	
E. lucanoides	RFF	E107	0	6	0	1	

700 701 702 703 Table 2. Date estimates for selected nodes in the Ecitophya and myrmecophiles' chronograms generated using MrBayes with TK02, IGR and strict clock models and either a fixed date calibration according to Brady (2003) and Maruyama and Parker (2017) or 1% rate calibration.

I         Eciton         TK02 clock, 26 MYA date calibration         26.0         n.a.         n.a.           I         Eciton         IGR clock, 26 MYA date calibration         12.1         8.8         21.9           I         Eciton         IGR clock, 1% rate calibration         12.1         8.9         16.7           I         Eciton         Strict clock, 1% rate calibration         12.1         8.9         16.7           I         Eciton         Strict clock, 1% rate calibration         15.9         2.2         14.5           II         Eciton         TK02 clock, 1% rate calibration         15.8         12.3         24.0           II         Eciton         IGR clock, 1% rate calibration         7.8         4.3         11.8           II         Eciton         Strict clock, 26 MYA date calibration         7.8         4.3         11.8           II         Eciton         TK02 clock, 1% rate calibration         1.8         0.6         5.3           III         Eciton         TK02 clock, 1% rate calibration         1.8         0.6         5.3           III         Eciton         TK02 clock, 26 MYA date calibration         1.3         0.3         3.5           IV         Eciton         TK12 clock, 1% rate ca	Code	Dataset	Method	Median	95% Lower	95% Upper
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I         Eciton         ICR clock, 26 MYA date calibration         26.0         n.a.         n.a.           I         Eciton         Strict clock, 26 MYA date calibration         26.0         n.a.         n.a.           I         Eciton         Strict clock, 1% rate calibration         11.5         9.2         14.45           II         Eciton         TK02 clock, 26 MYA date calibration         16.8         10.2         23.6           II         Eciton         TK02 clock, 26 MYA date calibration         7.8         5.0         16.4           II         Eciton         GR clock, 1% rate calibration         7.8         5.7         10.3           III         Eciton         Strict clock, 1% rate calibration         7.8         5.7         10.3           III         Eciton         TK02 clock, 26 MYA date calibration         3.5         1.3         7.5           III         Eciton         TGR clock, 1% rate calibration         1.8         0.6         5.3           III         Eciton         Strict clock, 26 MYA date calibration         1.3         3.8         11           Eciton         TGR clock, 1% rate calibration         1.3         0.3         3.5           IV         Eciton         TGR clock, 1% rate calibration </td <td>Ι</td> <td>Eciton</td> <td>TK02 clock, 1% rate calibration</td> <td>13.7</td> <td>8.8</td> <td>21.9</td>	Ι	Eciton	TK02 clock, 1% rate calibration	13.7	8.8	21.9
I         Eciton         IGR clock, 1% rate calibration         12.1         8.9         16.7           I         Eciton         Strict clock, 26 MYA date calibration         11.5         9.2         14.5           II         Eciton         TK02 clock, 1% rate calibration         18.0         12.3         24.0           II         Eciton         TK02 clock, 26 MYA date calibration         18.0         12.3         24.0           II         Eciton         IGR clock, 26 MYA date calibration         16.8         10.2         23.6           III         Eciton         Strict clock, 26 MYA date calibration         7.8         4.3         11.8           III         Eciton         Strict clock, 1% rate calibration         7.8         4.3         11.8           III         Eciton         TK02 clock, 26 MYA date calibration         1.8         0.6         5.3           III         Eciton         Strict clock, 1% rate calibration         1.8         0.6         1.6           III         Eciton         Strict clock, 26 MYA date calibration         1.1         0.6         1.6           IV         Eciton         TK02 clock, 26 MYA date calibration         1.1         0.1         2.3           IV         Eciton	Ι	Eciton	IGR clock, 26 MYA date calibration	26.0	n.a.	n.a.
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IIIEctionTK02 clock, 1% rate calibration1.80.65.3IIIEcitonIGR clock, 26 MYA date calibration3.91.28.3IIIEcitonStrict clock, 26 MYA date calibration1.80.54.0IIIEcitonStrict clock, 1% rate calibration1.10.61.6IVEcitonStrict clock, 26 MYA date calibration1.30.33.5IVEcitonTK02 clock, 26 MYA date calibration1.60.24.1IVEcitonIGR clock, 1% rate calibration0.70.11.9IVEcitonStrict clock, 26 MYA date calibration0.90.41.7IVEcitonStrict clock, 25 MYA date calibration0.80.22.0IXMyrmecophilesTK02 clock, 25 MYA date calibration0.40.20.7IXMyrmecophilesIGR clock, 25 MYA date calibration0.50.11.4IXMyrmecophilesIGR clock, 25 MYA date calibration0.70.21.4IXMyrmecophilesStrict clock, 1% rate calibration0.70.21.4IXMyrmecophilesStrict clock, 25 MYA date calibration0.70.11.2IXMyrmecophilesStrict clock, 25 MYA date calibration0.70.21.4IXMyrmecophilesStrict clock, 25 MYA date calibration25.0n.a.n.a.IXMyrmecophilesStrict clock, 25 MYA date calibration25.0n.a.n.a.IX<	III	Eciton	TK02 clock, 26 MYA date calibration	3.5	1.3	7.5
IIIEcitonIGR clock, 26 MYA date calibration $3.9$ $1.2$ $8.3$ IIIEcitonIGR clock, 1% rate calibration $1.8$ $0.5$ $4.0$ IIIEcitonStrict clock, 26 MYA date calibration $2.4$ $1.3$ $3.8$ IIIEcitonStrict clock, 1% rate calibration $1.1$ $0.6$ $1.6$ IVEcitonTK02 clock, 26 MYA date calibration $0.7$ $0.1$ $2.3$ IVEcitonIGR clock, 1% rate calibration $0.7$ $0.1$ $2.3$ IVEcitonIGR clock, 1% rate calibration $0.7$ $0.1$ $1.9$ VEcitonStrict clock, 26 MYA date calibration $0.9$ $0.4$ $1.7$ IVEcitonStrict clock, 26 MYA date calibration $0.8$ $0.2$ $2.0$ IXMyrmecophilesTK02 clock, 25 MYA date calibration $0.5$ $0.1$ $1.4$ IXMyrmecophilesIGR clock, 1% rate calibration $0.5$ $0.1$ $1.4$ IXMyrmecophilesStrict clock, 25 MYA date calibration $0.7$ $0.2$ $1.4$ IXMyrmecophilesStrict clock, 25 MYA date calibration $0.7$ $0.1$ $0.7$ VMyrmecophilesStrict clock, 25 MYA date calibration $0.7$ $0.1$ $0.7$ VMyrmecophilesStrict clock, 25 MYA date calibration $0.7$ $0.1$ $0.7$ VMyrmecophilesStrict clock, 25 MYA date calibration $1.3$ $9.3$ $16.1$ VMyrmecophilesStrict clock, 25 M	III	Eciton	TK02 clock, 1% rate calibration	1.8	0.6	5.3
IIIEcitonIGR clock, 1% rate calibration1.80.54.0IIIEcitonStrict clock, 26 MYA date calibration2.41.33.8IIIEcitonStrict clock, 1% rate calibration1.10.61.6IVEcitonTK02 clock, 1% rate calibration1.30.33.5IVEcitonIGR clock, 26 MYA date calibration0.70.12.3IVEcitonIGR clock, 26 MYA date calibration0.70.11.9IVEcitonStrict clock, 16% rate calibration0.70.11.9IVEcitonStrict clock, 16% rate calibration0.90.41.7IVEcitonStrict clock, 16% rate calibration0.80.22.0IXMyrmecophilesTK02 clock, 25 MYA date calibration0.50.11.4IXMyrmecophilesIGR clock, 25 MYA date calibration0.70.21.4IXMyrmecophilesStrict clock, 25 MYA date calibration0.70.21.4IXMyrmecophilesStrict clock, 16% rate calibration0.70.21.4IXMyrmecophilesStrict clock, 16% rate calibration0.70.10.7VMyrmecophilesStrict clock, 25 MYA date calibration0.40.10.7VMyrmecophilesStrict clock, 25 MYA date calibration25.0n.a.n.a.VMyrmecophilesIGR clock, 16% rate calibration25.0n.a.n.a.VMyrmecophiles <td>III</td> <td>Eciton</td> <td>IGR clock, 26 MYA date calibration</td> <td>3.9</td> <td>1.2</td> <td>8.3</td>	III	Eciton	IGR clock, 26 MYA date calibration	3.9	1.2	8.3
IIIEcitonStrict clock, 26 MYA date calibration2.41.33.8IIIEcitonStrict clock, 1% rate calibration1.10.61.6IVEcitonTK02 clock, 26 MYA date calibration1.30.33.5IVEcitonIGR clock, 26 MYA date calibration0.70.12.3IVEcitonIGR clock, 1% rate calibration0.70.11.9IVEcitonStrict clock, 26 MYA date calibration0.70.11.9IVEcitonStrict clock, 26 MYA date calibration0.90.41.7IVEcitonStrict clock, 1% rate calibration0.80.22.0IXMyrnecophilesTK02 clock, 1% rate calibration0.50.11.4IXMyrnecophilesIGR clock, 1% rate calibration0.50.11.2IXMyrnecophilesStrict clock, 25 MYA date calibration0.70.21.4IXMyrnecophilesStrict clock, 25 MYA date calibration0.40.10.7VMyrnecophilesStrict clock, 25 MYA date calibration25.0n.a.n.a.VMyrnecophilesIGR clock, 1% rate calibration12.39.316.1VMyrnecophilesStrict clock, 25 MYA date calibration12.39.815.4VMyrnecophilesStrict clock, 25 MYA date calibration12.39.815.4VMyrnecophilesStrict clock, 1% rate calibration12.39.815.4VMyr	III	Eciton	IGR clock, 1% rate calibration	1.8	0.5	4.0
IIIEcitonStrict clock, 1% rate calibration1.10.61.6IVEcitonTK02 clock, 26 MYA date calibration0.70.12.3IVEcitonIGR clock, 1% rate calibration0.70.12.3IVEcitonIGR clock, 1% rate calibration0.70.11.9IVEcitonStrict clock, 26 MYA date calibration0.90.41.7IVEcitonStrict clock, 1% rate calibration0.40.20.7IXMyrnecophilesTK02 clock, 25 MYA date calibration0.80.22.0IXMyrnecophilesIGR clock, 25 MYA date calibration1.10.12.9IXMyrnecophilesIGR clock, 25 MYA date calibration0.50.11.2IXMyrnecophilesStrict clock, 1% rate calibration0.70.21.4IXMyrnecophilesStrict clock, 25 MYA date calibration0.70.21.4IXMyrnecophilesStrict clock, 25 MYA date calibration0.70.21.4IXMyrnecophilesStrict clock, 1% rate calibration14.39.523.1VMyrnecophilesStrict clock, 1% rate calibration1.439.523.1VMyrnecophilesIGR clock, 25 MYA date calibration12.39.316.1VMyrnecophilesStrict clock, 1% rate calibration12.39.316.1VMyrnecophilesStrict clock, 25 MYA date calibration12.39.815.4 <t< td=""><td>III</td><td>Eciton</td><td>Strict clock, 26 MYA date calibration</td><td>2.4</td><td>1.3</td><td>3.8</td></t<>	III	Eciton	Strict clock, 26 MYA date calibration	2.4	1.3	3.8
IVEcitonTK02 clock, 26 MYA date calibration1.30.33.5IVEcitonTK02 clock, 1% rate calibration0.70.12.3IVEcitonIGR clock, 26 MYA date calibration0.60.24.1IVEcitonStrict clock, 26 MYA date calibration0.70.11.9IVEcitonStrict clock, 1% rate calibration0.40.20.7IXMyrmecophilesTK02 clock, 25 MYA date calibration0.40.20.7IXMyrmecophilesTK02 clock, 1% rate calibration0.50.11.4IXMyrmecophilesIGR clock, 25 MYA date calibration0.50.11.2IXMyrmecophilesStrict clock, 1% rate calibration0.70.21.4IXMyrmecophilesStrict clock, 1% rate calibration0.70.21.4IXMyrmecophilesStrict clock, 25 MYA date calibration0.40.10.7VMyrmecophilesStrict clock, 25 MYA date calibration25.0n.a.n.a.VMyrmecophilesStrict clock, 25 MYA date calibration12.39.316.1VMyrmecophilesStrict clock, 25 MYA date calibration12.39.316.1VMyrmecophilesStrict clock, 25 MYA date calibration12.39.316.1VMyrmecophilesStrict clock, 25 MYA date calibration12.39.815.4VMyrmecophilesStrict clock, 25 MYA date calibration12.39.815.4<	III	Eciton	Strict clock, 1% rate calibration	1.1	0.6	1.6
IVEcitonTK02 clock, 1% rate calibration $0.7$ $0.1$ $2.3$ IVEcitonIGR clock, 26 MYA date calibration $1.6$ $0.2$ $4.1$ IVEcitonStrict clock, 26 MYA date calibration $0.7$ $0.1$ $1.9$ IVEcitonStrict clock, 25 MYA date calibration $0.4$ $0.2$ $0.7$ IXMyrmecophilesTK02 clock, 25 MYA date calibration $0.8$ $0.2$ $2.0$ IXMyrmecophilesIGR clock, 1% rate calibration $0.5$ $0.1$ $1.4$ IXMyrmecophilesIGR clock, 25 MYA date calibration $0.5$ $0.1$ $1.4$ IXMyrmecophilesStrict clock, 1% rate calibration $0.5$ $0.1$ $1.2$ IXMyrmecophilesStrict clock, 25 MYA date calibration $0.7$ $0.2$ $1.4$ IXMyrmecophilesStrict clock, 25 MYA date calibration $0.4$ $0.1$ $0.7$ VMyrmecophilesTK02 clock, 1% rate calibration $25.0$ $n.a.$ $n.a.$ VMyrmecophilesIGR clock, 1% rate calibration $25.0$ $n.a.$ $n.a.$ VMyrmecophilesStrict clock, 25 MYA date calibration $25.0$ $n.a.$ $n.a.$ VMyrmecophilesStrict clock, 25 MYA date calibration $25.0$ $n.a.$ $n.a.$ VMyrmecophilesStrict clock, 25 MYA date calibration $25.0$ $n.a.$ $n.a.$ VMyrmecophilesStrict clock, 25 MYA date calibration $2.3$ $9.3$ $16.1$ V <td>IV</td> <td>Eciton</td> <td>TK02 clock, 26 MYA date calibration</td> <td>1.3</td> <td>0.3</td> <td>3.5</td>	IV	Eciton	TK02 clock, 26 MYA date calibration	1.3	0.3	3.5
IVEcitonIGR clock, 26 MYA date calibration1.60.24.1IVEcitonIGR clock, 1% rate calibration0.70.11.9IVEcitonStrict clock, 1% rate calibration0.90.41.7IVEcitonStrict clock, 1% rate calibration0.40.20.7IXMyrmecophilesTK02 clock, 25 MYA date calibration0.80.22.0IXMyrmecophilesIGR clock, 1% rate calibration0.50.11.4IXMyrmecophilesIGR clock, 1% rate calibration0.50.11.2IXMyrmecophilesStrict clock, 1% rate calibration0.70.21.4IXMyrmecophilesStrict clock, 25 MYA date calibration0.40.10.7VMyrmecophilesTK02 clock, 25 MYA date calibration25.0n.a.n.a.VMyrmecophilesIGR clock, 1% rate calibration25.0n.a.n.a.VMyrmecophilesStrict clock, 25 MYA date calibration25.0n.a.n.a.VMyrmecophilesStrict clock, 1% rate calibration12.39.316.1VMyrmecophilesStrict clock, 25 MYA date calibration25.0n.a.n.a.VMyrmecophilesStrict clock, 25 MYA date calibration2.19.815.4VIMyrmecophilesStrict clock, 25 MYA date calibration3.04.214.1VIMyrmecophilesIGR clock, 25 MYA date calibration6.73.411.7	IV	Eciton	TK02 clock, 1% rate calibration	0.7	0.1	2.3
IVEcitonIGR clock, 1% rate calibration $0.7$ $0.1$ $1.9$ IVEcitonStrict clock, 26 MYA date calibration $0.9$ $0.4$ $1.7$ IVEcitonStrict clock, 1% rate calibration $0.4$ $0.2$ $0.7$ IXMyrmecophilesTK02 clock, 25 MYA date calibration $0.8$ $0.2$ $2.0$ IXMyrmecophilesICR clock, 25 MYA date calibration $0.5$ $0.1$ $1.4$ IXMyrmecophilesICR clock, 1% rate calibration $0.5$ $0.1$ $1.2$ IXMyrmecophilesStrict clock, 1% rate calibration $0.7$ $0.2$ $1.4$ IXMyrmecophilesStrict clock, 1% rate calibration $0.4$ $0.1$ $0.7$ VMyrmecophilesTK02 clock, 25 MYA date calibration $14.3$ $9.5$ $23.1$ VMyrmecophilesTK02 clock, 1% rate calibration $12.3$ $9.3$ $16.1$ VMyrmecophilesIGR clock, 1% rate calibration $12.3$ $9.8$ $15.4$ VMyrmecophilesStrict clock, 1% rate calibration $12.3$ $9.8$ $15.4$ VIMyrmecophilesIGR clock, 1% rate calibration $4.5$ $1.9$ $11.7$ VIMyrmecophilesIGR clock, 1% rate calibration $4.5$ $1.9$ $11.2$ VMyrmecophilesStrict clock, 25 MYA date calibration $12.3$ $9.8$ $15.4$ VIMyrmecophilesIGR clock, 1% rate calibration $3.2$ $1.7$ $5.2$ VIMyrmecophilesI	IV	Eciton	IGR clock, 26 MYA date calibration	1.6	0.2	4.1
IVEcitonStrict clock, 26 MYA date calibration $0.9$ $0.4$ $1.7$ IVEcitonStrict clock, 1% rate calibration $0.4$ $0.2$ $0.7$ IXMyrmecophilesTK02 clock, 25 MYA date calibration $0.8$ $0.2$ $2.0$ IXMyrmecophilesTK02 clock, 1% rate calibration $0.5$ $0.1$ $1.4$ IXMyrmecophilesIGR clock, 25 MYA date calibration $0.5$ $0.1$ $1.4$ IXMyrmecophilesStrict clock, 25 MYA date calibration $0.7$ $0.2$ $1.4$ IXMyrmecophilesStrict clock, 1% rate calibration $0.4$ $0.1$ $0.7$ VMyrmecophilesStrict clock, 1% rate calibration $0.4$ $0.1$ $0.7$ VMyrmecophilesStrict clock, 1% rate calibration $14.3$ $9.5$ $23.1$ VMyrmecophilesIGR clock, 1% rate calibration $12.3$ $9.3$ $16.1$ VMyrmecophilesIGR clock, 1% rate calibration $12.3$ $9.3$ $16.1$ VMyrmecophilesStrict clock, 1% rate calibration $12.3$ $9.8$ $15.4$ VIMyrmecophilesStrict clock, 25 MYA date calibration $12.3$ $9.8$ $15.4$ VIMyrmecophilesTK02 clock, 1% rate calibration $4.5$ $1.9$ $11.2$ VIMyrmecophilesTK02 clock, 1% rate calibration $3.2$ $1.7$ $5.2$ VIMyrmecophilesStrict clock, 1% rate calibration $2.9$ $2.0$ $4.0$ VIMyrmeco	IV	Eciton	IGR clock, 1% rate calibration	0.7	0.1	1.9
IVEcitonStrict clock, 1% rate calibration $0.4$ $0.2$ $0.7$ IXMyrmecophilesTK02 clock, 25 MYA date calibration $0.8$ $0.2$ $2.0$ IXMyrmecophilesIGR clock, 25 MYA date calibration $0.5$ $0.1$ $1.4$ IXMyrmecophilesIGR clock, 1% rate calibration $0.5$ $0.1$ $1.2$ IXMyrmecophilesStrict clock, 1% rate calibration $0.5$ $0.1$ $1.2$ IXMyrmecophilesStrict clock, 1% rate calibration $0.7$ $0.2$ $1.4$ IXMyrmecophilesStrict clock, 1% rate calibration $0.4$ $0.1$ $0.7$ VMyrmecophilesTK02 clock, 1% rate calibration $14.3$ $9.5$ $23.1$ VMyrmecophilesTK02 clock, 1% rate calibration $12.3$ $9.3$ $16.1$ VMyrmecophilesStrict clock, 25 MYA date calibration $12.3$ $9.8$ $15.4$ VMyrmecophilesStrict clock, 25 MYA date calibration $12.3$ $9.8$ $15.4$ VIMyrmecophilesStrict clock, 25 MYA date calibration $12.3$ $9.8$ $15.4$ VIMyrmecophilesIGR clock, 1% rate calibration $3.2$ $1.7$ $5.2$ VIMyrmecophilesIGR clock, 25 MYA date calibration $5.9$ $3.8$ $8.1$ VIMyrmecophilesIGR clock, 25 MYA date calibration $5.9$ $3.8$ $8.1$ VIMyrmecophilesIGR clock, 1% rate calibration $7.8$ $4.4$ $11.7$ VI <t< td=""><td>IV</td><td>Eciton</td><td>Strict clock, 26 MYA date calibration</td><td>0.9</td><td>0.4</td><td>1.7</td></t<>	IV	Eciton	Strict clock, 26 MYA date calibration	0.9	0.4	1.7
IXMyrmecophilesTK02 clock, 25 MYA date calibration $0.8$ $0.2$ $2.0$ IXMyrmecophilesIGR clock, 25 MYA date calibration $0.5$ $0.1$ $1.4$ IXMyrmecophilesIGR clock, 1% rate calibration $1.1$ $0.1$ $2.9$ IXMyrmecophilesIGR clock, 1% rate calibration $0.5$ $0.1$ $1.2$ IXMyrmecophilesStrict clock, 25 MYA date calibration $0.7$ $0.2$ $1.4$ IXMyrmecophilesStrict clock, 25 MYA date calibration $0.4$ $0.1$ $0.7$ VMyrmecophilesTK02 clock, 25 MYA date calibration $25.0$ $n.a.$ $n.a.$ VMyrmecophilesIGR clock, 1% rate calibration $25.0$ $n.a.$ $n.a.$ VMyrmecophilesIGR clock, 1% rate calibration $25.0$ $n.a.$ $n.a.$ VMyrmecophilesStrict clock, 25 MYA date calibration $25.0$ $n.a.$ $n.a.$ VMyrmecophilesStrict clock, 1% rate calibration $25.0$ $n.a.$ $n.a.$ VMyrmecophilesStrict clock, 25 MYA date calibration $25.0$ $n.a.$ $n.a.$ VMyrmecophilesStrict clock, 1% rate calibration $25.0$ $n.a.$ $n.a.$ VMyrmecophilesStrict clock, 1% rate calibration $25.0$ $n.a.$ $n.a.$ VMyrmecophilesStrict clock, 1% rate calibration $25.0$ $n.a.$ $n.a.$ VIMyrmecophilesStrict clock, 1% rate calibration $25.0$ $n.a.$ $n.a.$ <	IV	Eciton	Strict clock, 1% rate calibration	0.4	0.2	0.7
IXMyrmecophilesTK02 clock, 1% rate calibration $0.5$ $0.1$ $1.4$ IXMyrmecophilesIGR clock, 25 MYA date calibration $1.1$ $0.1$ $2.9$ IXMyrmecophilesIGR clock, 1% rate calibration $0.5$ $0.1$ $1.2$ IXMyrmecophilesStrict clock, 25 MYA date calibration $0.7$ $0.2$ $1.4$ IXMyrmecophilesStrict clock, 25 MYA date calibration $0.4$ $0.1$ $0.7$ VMyrmecophilesTK02 clock, 25 MYA date calibration $25.0$ $n.a.$ $n.a.$ VMyrmecophilesIGR clock, 25 MYA date calibration $25.0$ $n.a.$ $n.a.$ VMyrmecophilesIGR clock, 25 MYA date calibration $25.0$ $n.a.$ $n.a.$ VMyrmecophilesIGR clock, 1% rate calibration $25.0$ $n.a.$ $n.a.$ VMyrmecophilesStrict clock, 25 MYA date calibration $25.0$ $n.a.$ $n.a.$ VMyrmecophilesStrict clock, 1% rate calibration $25.0$ $n.a.$ $n.a.$ VMyrmecophilesStrict clock, 1% rate calibration $25.0$ $n.a.$ $n.a.$ VIMyrmecophilesTK02 clock, 1% rate calibration $4.5$ $1.9$ $11.2$ VIMyrmecophilesIGR clock, 25 MYA date calibration $3.2$ $1.7$ $5.2$ VIMyrmecophilesIGR clock, 25 MYA date calibration $5.9$ $3.8$ $8.1$ VIMyrmecophilesStrict clock, 25 MYA date calibration $1.4$ $1.4$ <td< td=""><td>IX</td><td>Myrmecophiles</td><td>TK02 clock, 25 MYA date calibration</td><td>0.8</td><td>0.2</td><td>2.0</td></td<>	IX	Myrmecophiles	TK02 clock, 25 MYA date calibration	0.8	0.2	2.0
IXMyrmecophilesIGR clock, 25 MYA date calibration1.10.12.9IXMyrmecophilesIGR clock, 1% rate calibration0.50.11.2IXMyrmecophilesStrict clock, 25 MYA date calibration0.70.21.4IXMyrmecophilesStrict clock, 1% rate calibration0.40.10.7VMyrmecophilesTK02 clock, 25 MYA date calibration25.0n.a.n.a.VMyrmecophilesIGR clock, 25 MYA date calibration25.0n.a.n.a.VMyrmecophilesIGR clock, 25 MYA date calibration25.0n.a.n.a.VMyrmecophilesStrict clock, 1% rate calibration25.0n.a.n.a.VMyrmecophilesStrict clock, 25 MYA date calibration12.39.815.4VIMyrmecophilesStrict clock, 25 MYA date calibration3.04.214.1VIMyrmecophilesIGR clock, 1% rate calibration3.21.75.2VIMyrmecophilesStrict clock, 25 MYA date calibration3.21.75.2VIMyrmecophilesStrict clock, 1% rate calibration2.92.04.0VIIMyrmecophilesStrict clock, 25 MYA date calibra	IX	Myrmecophiles	TK02 clock, 1% rate calibration	0.5	0.1	1.4
IXMyrmecophilesIGR clock, 1% rate calibration0.50.11.2IXMyrmecophilesStrict clock, 25 MYA date calibration0.70.21.4IXMyrmecophilesStrict clock, 25 MYA date calibration0.40.10.7VMyrmecophilesTK02 clock, 25 MYA date calibration14.39.523.1VMyrmecophilesIGR clock, 25 MYA date calibration12.39.316.1VMyrmecophilesStrict clock, 25 MYA date calibration12.39.316.1VMyrmecophilesStrict clock, 25 MYA date calibration12.39.815.4VIMyrmecophilesStrict clock, 1% rate calibration12.39.815.4VIMyrmecophilesTK02 clock, 1% rate calibration4.51.911.2VIMyrmecophilesTK02 clock, 1% rate calibration4.51.911.2VIMyrmecophilesIGR clock, 1% rate calibration3.21.75.2VIMyrmecophilesIGR clock, 1% rate calibration2.92.04.0VIMyrmecophilesStrict clock, 25 MYA date calibration7.84.414.2VIMyrmecophilesTK02 clock, 1% rate calibration7.84.414.2VIMyrmecophilesIGR clock, 1% rate calibration7.84.414.2VIMyrmecophilesIGR clock, 25 MYA date calibration14.710.419.9VIIMyrmecophilesIGR clock, 25 MYA date calibration7.14	IX	Myrmecophiles	IGR clock, 25 MYA date calibration	1.1	0.1	2.9
IXMyrmecophilesStrict clock, 25 MYA date calibration0.70.21.4IXMyrmecophilesStrict clock, 1% rate calibration0.40.10.7VMyrmecophilesTK02 clock, 25 MYA date calibration25.0n.a.n.a.VMyrmecophilesIGR clock, 1% rate calibration14.39.523.1VMyrmecophilesIGR clock, 1% rate calibration12.39.316.1VMyrmecophilesStrict clock, 25 MYA date calibration12.39.316.1VMyrmecophilesStrict clock, 25 MYA date calibration12.39.815.4VIMyrmecophilesTK02 clock, 1% rate calibration12.39.815.4VIMyrmecophilesTK02 clock, 1% rate calibration8.04.214.1VIMyrmecophilesTK02 clock, 1% rate calibration6.73.411.7VIMyrmecophilesIGR clock, 1% rate calibration3.21.75.2VIMyrmecophilesStrict clock, 25 MYA date calibration2.92.04.0VIMyrmecophilesStrict clock, 25 MYA date calibration13.89.918.0VIMyrmecophilesTK02 clock, 1% rate calibration7.84.414.2VIMyrmecophilesTK02 clock, 25 MYA date calibration14.710.419.9VIIMyrmecophilesStrict clock, 25 MYA date calibration7.14.89.7VIIMyrmecophilesIGR clock, 1% rate calibration7.1<	IX	Myrmecophiles	IGR clock, 1% rate calibration	0.5	0.1	1.2
IXMyrmecophilesStrict clock, 1% rate calibration0.40.10.7VMyrmecophilesTK02 clock, 25 MYA date calibration25.0n.a.n.a.n.a.VMyrmecophilesIGR clock, 25 MYA date calibration25.0n.a.n.a.n.a.VMyrmecophilesIGR clock, 1% rate calibration25.0n.a.n.a.n.a.VMyrmecophilesIGR clock, 1% rate calibration25.0n.a.n.a.n.a.VMyrmecophilesStrict clock, 25 MYA date calibration25.0n.a.n.a.n.a.VMyrmecophilesStrict clock, 25 MYA date calibration12.39.815.4VIMyrmecophilesStrict clock, 1% rate calibration8.04.214.1VIMyrmecophilesTK02 clock, 25 MYA date calibration6.73.411.7VIMyrmecophilesIGR clock, 1% rate calibration3.21.75.2VIMyrmecophilesStrict clock, 25 MYA date calibration3.21.75.2VIMyrmecophilesStrict clock, 25 MYA date calibration2.92.04.0VIIMyrmecophilesStrict clock, 25 MYA date calibration13.89.918.0VIIMyrmecophilesTK02 clock, 25 MYA date calibration14.710.419.9VIIMyrmecophilesTK02 clock, 25 MYA date calibration14.211.217.4VIIMyrmecophilesTK02 clock, 1% rate calibration14.710.419.9 <td< td=""><td>IX</td><td>Myrmecophiles</td><td>Strict clock, 25 MYA date calibration</td><td>0.7</td><td>0.2</td><td>1.4</td></td<>	IX	Myrmecophiles	Strict clock, 25 MYA date calibration	0.7	0.2	1.4
VMyrmecophilesTK02 clock, 25 MYA date calibration25.0n.a.n.a.VMyrmecophilesTK02 clock, 1% rate calibration14.39.523.1VMyrmecophilesIGR clock, 25 MYA date calibration25.0n.a.n.a.VMyrmecophilesIGR clock, 25 MYA date calibration25.0n.a.n.a.VMyrmecophilesStrict clock, 25 MYA date calibration25.0n.a.n.a.VMyrmecophilesStrict clock, 25 MYA date calibration25.0n.a.n.a.VMyrmecophilesStrict clock, 25 MYA date calibration25.0n.a.n.a.VIMyrmecophilesStrict clock, 1% rate calibration25.0n.a.n.a.VIMyrmecophilesStrict clock, 1% rate calibration4.214.1VIMyrmecophilesIGR clock, 1% rate calibration4.51.911.2VIMyrmecophilesIGR clock, 1% rate calibration3.21.75.2VIMyrmecophilesStrict clock, 25 MYA date calibration5.93.88.1VIMyrmecophilesStrict clock, 25 MYA date calibration13.89.918.0VIIMyrmecophilesTK02 clock, 1% rate calibration14.710.419.9VIIMyrmecophilesIGR clock, 1% rate calibration7.84.414.2VIIMyrmecophilesIGR clock, 25 MYA date calibration14.710.419.9VIIMyrmecophilesStrict clock, 25 MYA date calibration	IX	Myrmecophiles	Strict clock, 1% rate calibration	0.4	0.1	0.7
VMyrmecophilesTK02 clock, 1% rate calibration14.39.523.1VMyrmecophilesIGR clock, 25 MYA date calibration25.0n.a.n.a.n.a.VMyrmecophilesIGR clock, 25 MYA date calibration12.39.316.1VMyrmecophilesStrict clock, 25 MYA date calibration25.0n.a.n.a.VMyrmecophilesStrict clock, 25 MYA date calibration25.0n.a.n.a.VMyrmecophilesStrict clock, 1% rate calibration12.39.815.4VIMyrmecophilesTK02 clock, 1% rate calibration8.04.214.1VIMyrmecophilesIGR clock, 25 MYA date calibration6.73.411.7VIMyrmecophilesIGR clock, 25 MYA date calibration5.93.88.1VIMyrmecophilesStrict clock, 1% rate calibration2.92.04.0VIMyrmecophilesStrict clock, 1% rate calibration13.89.918.0VIIMyrmecophilesTK02 clock, 25 MYA date calibration14.710.419.9VIIMyrmecophilesTK02 clock, 25 MYA date calibration14.710.419.9VIIMyrmecophilesIGR clock, 1% rate calibration7.14.89.7VIIMyrmecophilesStrict clock, 1% rate calibration7.14.89.7VIIMyrmecophilesStrict clock, 25 MYA date calibration7.84.612.0VIIMyrmecophilesStrict clock, 1% rate ca	V	Myrmecophiles	TK02 clock, 25 MYA date calibration	25.0	n.a.	n.a.
VMyrmecophilesIGR clock, 25 MYA date calibration25.0n.a.n.a.VMyrmecophilesIGR clock, 1% rate calibration12.39.316.1VMyrmecophilesStrict clock, 25 MYA date calibration25.0n.a.n.a.VMyrmecophilesStrict clock, 25 MYA date calibration12.39.815.4VIMyrmecophilesTK02 clock, 25 MYA date calibration8.04.214.1VIMyrmecophilesTK02 clock, 25 MYA date calibration4.51.911.2VIMyrmecophilesIGR clock, 25 MYA date calibration6.73.411.7VIMyrmecophilesIGR clock, 1% rate calibration3.21.75.2VIMyrmecophilesStrict clock, 25 MYA date calibration2.92.04.0VIMyrmecophilesStrict clock, 1% rate calibration2.92.04.0VIIMyrmecophilesTK02 clock, 1% rate calibration13.89.918.0VIIMyrmecophilesIGR clock, 1% rate calibration7.84.414.2VIIMyrmecophilesIGR clock, 25 MYA date calibration14.710.419.9VIIMyrmecophilesStrict clock, 25 MYA date calibration14.211.217.4VIIMyrmecophilesIGR clock, 25 MYA date calibration14.211.217.4VIIMyrmecophilesStrict clock, 25 MYA date calibration7.14.89.7VIIMyrmecophilesStrict clock, 1% rate calibratio	V	Myrmecophiles	TK02 clock, 1% rate calibration	14.3	9.5	23.1
VMyrmecophilesIGR clock, 1% rate calibration12.39.316.1VMyrmecophilesStrict clock, 25 MYA date calibration25.0n.a.n.a.VMyrmecophilesStrict clock, 1% rate calibration12.39.815.4VIMyrmecophilesTK02 clock, 25 MYA date calibration8.04.214.1VIMyrmecophilesTK02 clock, 1% rate calibration4.51.911.2VIMyrmecophilesIGR clock, 25 MYA date calibration6.73.411.7VIMyrmecophilesIGR clock, 1% rate calibration3.21.75.2VIMyrmecophilesStrict clock, 1% rate calibration2.92.04.0VIMyrmecophilesTK02 clock, 25 MYA date calibration13.89.918.0VIMyrmecophilesTK02 clock, 25 MYA date calibration14.710.419.9VIIMyrmecophilesTK02 clock, 1% rate calibration14.710.419.9VIIMyrmecophilesIGR clock, 25 MYA date calibration14.211.217.4VIIMyrmecophilesIGR clock, 25 MYA date calibration14.211.217.4VIIMyrmecophilesStrict clock, 25 MYA date calibration14.211.217.4VIIMyrmecophilesIGR clock, 25 MYA date calibration14.211.217.4VIIMyrmecophilesStrict clock, 25 MYA date calibration7.84.612.0VIIIMyrmecophilesTK02 clock, 25 MYA date calib	V	Myrmecophiles	IGR clock, 25 MYA date calibration	25.0	n.a.	n.a.
VMyrmecophilesStrict clock, 25 MYA date calibration25.0n.a.n.a.VMyrmecophilesStrict clock, 1% rate calibration12.39.815.4VIMyrmecophilesTK02 clock, 25 MYA date calibration8.04.214.1VIMyrmecophilesTK02 clock, 1% rate calibration4.51.911.2VIMyrmecophilesIGR clock, 25 MYA date calibration6.73.411.7VIMyrmecophilesIGR clock, 1% rate calibration3.21.75.2VIMyrmecophilesStrict clock, 25 MYA date calibration5.93.88.1VIMyrmecophilesStrict clock, 1% rate calibration2.92.04.0VIIMyrmecophilesTK02 clock, 1% rate calibration13.89.918.0VIIMyrmecophilesTK02 clock, 1% rate calibration14.710.419.9VIIMyrmecophilesIGR clock, 25 MYA date calibration14.710.419.9VIIMyrmecophilesIGR clock, 1% rate calibration14.211.217.4VIIMyrmecophilesStrict clock, 25 MYA date calibration14.211.217.4VIIMyrmecophilesStrict clock, 25 MYA date calibration6.95.38.8VIIIMyrmecophilesStrict clock, 25 MYA date calibration7.84.612.0VIIMyrmecophilesStrict clock, 1% rate calibration7.84.612.0VIIMyrmecophilesStrict clock, 1% rate calibration	V	Myrmecophiles	IGR clock, 1% rate calibration	12.3	9.3	16.1
VMyrmecophilesStrict clock, 1% rate calibration12.39.815.4VIMyrmecophilesTK02 clock, 25 MYA date calibration8.04.214.1VIMyrmecophilesTK02 clock, 25 MYA date calibration4.51.911.2VIMyrmecophilesIGR clock, 25 MYA date calibration6.73.411.7VIMyrmecophilesStrict clock, 25 MYA date calibration5.93.88.1VIMyrmecophilesStrict clock, 25 MYA date calibration2.92.04.0VIIMyrmecophilesTK02 clock, 1% rate calibration13.89.918.0VIIMyrmecophilesTK02 clock, 25 MYA date calibration14.710.419.9VIIMyrmecophilesIGR clock, 25 MYA date calibration14.710.419.9VIIMyrmecophilesIGR clock, 25 MYA date calibration14.211.217.4VIMyrmecophilesIGR clock, 25 MYA date calibration14.710.419.9VIIMyrmecophilesIGR clock, 1% rate calibration14.211.217.4VIIMyrmecophilesStrict clock, 1% rate calibration6.95.38.8VIIIMyrmecophilesTK02 clock, 1% rate calibration7.84.612.0VIIMyrmecophilesTK02 clock, 1% rate calibration7.84.612.0VIIIMyrmecophilesTK02 clock, 1% rate calibration7.84.612.0VIIIMyrmecophilesTK02 clock, 1% rate calibration </td <td>V</td> <td>Myrmecophiles</td> <td>Strict clock, 25 MYA date calibration</td> <td>25.0</td> <td>n.a.</td> <td>n.a.</td>	V	Myrmecophiles	Strict clock, 25 MYA date calibration	25.0	n.a.	n.a.
VIMyrmecophilesTK02 clock, 25 MYA date calibration8.04.214.1VIMyrmecophilesTK02 clock, 1% rate calibration4.51.911.2VIMyrmecophilesIGR clock, 25 MYA date calibration6.73.411.7VIMyrmecophilesIGR clock, 1% rate calibration3.21.75.2VIMyrmecophilesStrict clock, 25 MYA date calibration5.93.88.1VIMyrmecophilesStrict clock, 1% rate calibration2.92.04.0VIIMyrmecophilesTK02 clock, 25 MYA date calibration13.89.918.0VIIMyrmecophilesTK02 clock, 1% rate calibration7.84.414.2VIIMyrmecophilesIGR clock, 25 MYA date calibration14.710.419.9VIIMyrmecophilesIGR clock, 25 MYA date calibration14.710.419.9VIIMyrmecophilesIGR clock, 1% rate calibration14.211.217.4VIIMyrmecophilesStrict clock, 25 MYA date calibration6.95.38.8VIIIMyrmecophilesTK02 clock, 1% rate calibration7.84.612.0VIIIMyrmecophilesTK02 clock, 1% rate calibration7.84.612.0VIIIMyrmecophilesTK02 clock, 1% rate calibration4.42.19.5VIIIMyrmecophilesIGR clock, 25 MYA date calibration8.45.113.3VIIIMyrmecophilesIGR clock, 1% rate calibration4.0<	V	Myrmecophiles	Strict clock, 1% rate calibration	12.3	9.8	15.4
VIMyrmecophilesTK02 clock, 1% rate calibration4.51.911.2VIMyrmecophilesIGR clock, 25 MYA date calibration6.73.411.7VIMyrmecophilesIGR clock, 1% rate calibration3.21.75.2VIMyrmecophilesStrict clock, 25 MYA date calibration5.93.88.1VIMyrmecophilesStrict clock, 1% rate calibration2.92.04.0VIIMyrmecophilesTK02 clock, 25 MYA date calibration13.89.918.0VIIMyrmecophilesTK02 clock, 1% rate calibration7.84.414.2VIIMyrmecophilesIGR clock, 25 MYA date calibration14.710.419.9VIIMyrmecophilesIGR clock, 25 MYA date calibration14.211.217.4VIIMyrmecophilesStrict clock, 25 MYA date calibration14.211.217.4VIIMyrmecophilesStrict clock, 25 MYA date calibration14.211.217.4VIIMyrmecophilesStrict clock, 25 MYA date calibration6.95.38.8VIIIMyrmecophilesStrict clock, 25 MYA date calibration7.84.612.0VIIIMyrmecophilesTK02 clock, 1% rate calibration7.84.612.0VIIIMyrmecophilesIGR clock, 25 MYA date calibration7.84.612.0VIIIMyrmecophilesIGR clock, 25 MYA date calibration8.45.113.3VIIIMyrmecophilesIGR clock, 25 MYA dat	VI	Myrmecophiles	TK02 clock, 25 MYA date calibration	8.0	4.2	14.1
VIMyrmecophilesIGR clock, 25 MYA date calibration6.73.411.7VIMyrmecophilesIGR clock, 1% rate calibration3.21.75.2VIMyrmecophilesStrict clock, 25 MYA date calibration5.93.88.1VIMyrmecophilesStrict clock, 1% rate calibration2.92.04.0VIIMyrmecophilesTK02 clock, 25 MYA date calibration13.89.918.0VIIMyrmecophilesTK02 clock, 1% rate calibration7.84.414.2VIIMyrmecophilesIGR clock, 25 MYA date calibration14.710.419.9VIIMyrmecophilesIGR clock, 1% rate calibration7.14.89.7VIIMyrmecophilesStrict clock, 25 MYA date calibration14.211.217.4VIIMyrmecophilesStrict clock, 25 MYA date calibration6.95.38.8VIIIMyrmecophilesStrict clock, 25 MYA date calibration7.84.612.0VIIMyrmecophilesStrict clock, 1% rate calibration7.84.612.0VIIIMyrmecophilesTK02 clock, 1% rate calibration7.84.612.0VIIIMyrmecophilesIGR clock, 25 MYA date calibration7.84.612.0VIIIMyrmecophilesIGR clock, 25 MYA date calibration7.84.612.0VIIIMyrmecophilesIGR clock, 25 MYA date calibration8.45.113.3VIIIMyrmecophilesIGR clock, 25 MYA date calibr	VI	Myrmecophiles	TK02 clock, 1% rate calibration	4.5	1.9	11.2
VIMyrmecophilesIGR clock, 1% rate calibration3.21.75.2VIMyrmecophilesStrict clock, 25 MYA date calibration5.93.88.1VIMyrmecophilesStrict clock, 1% rate calibration2.92.04.0VIIMyrmecophilesTK02 clock, 25 MYA date calibration13.89.918.0VIIMyrmecophilesTK02 clock, 1% rate calibration7.84.414.2VIIMyrmecophilesIGR clock, 25 MYA date calibration14.710.419.9VIIMyrmecophilesIGR clock, 25 MYA date calibration7.14.89.7VIIMyrmecophilesStrict clock, 25 MYA date calibration14.211.217.4VIIMyrmecophilesStrict clock, 25 MYA date calibration6.95.38.8VIIIMyrmecophilesStrict clock, 25 MYA date calibration7.84.612.0VIIMyrmecophilesTK02 clock, 1% rate calibration7.84.612.0VIIIMyrmecophilesTK02 clock, 1% rate calibration7.84.612.0VIIIMyrmecophilesTK02 clock, 1% rate calibration4.42.19.5VIIIMyrmecophilesIGR clock, 25 MYA date calibration8.45.113.3VIIIMyrmecophilesIGR clock, 25 MYA date calibration8.45.113.3VIIIMyrmecophilesIGR clock, 1% rate calibration4.02.66.0VIIIMyrmecophilesStrict clock, 25 MYA date calibrat	VI	Myrmecophiles	IGR clock, 25 MYA date calibration	6.7	3.4	11.7
VIMyrmecophilesStrict clock, 25 MYA date calibration5.93.88.1VIMyrmecophilesStrict clock, 1% rate calibration2.92.04.0VIIMyrmecophilesTK02 clock, 25 MYA date calibration13.89.918.0VIIMyrmecophilesTK02 clock, 1% rate calibration7.84.414.2VIIMyrmecophilesIGR clock, 25 MYA date calibration14.710.419.9VIIMyrmecophilesIGR clock, 1% rate calibration7.14.89.7VIIMyrmecophilesStrict clock, 25 MYA date calibration14.211.217.4VIIMyrmecophilesStrict clock, 1% rate calibration6.95.38.8VIIIMyrmecophilesTK02 clock, 1% rate calibration7.84.612.0VIIIMyrmecophilesTK02 clock, 1% rate calibration4.42.19.5VIIIMyrmecophilesTK02 clock, 25 MYA date calibration8.45.113.3VIIIMyrmecophilesTK02 clock, 1% rate calibration4.42.19.5VIIIMyrmecophilesIGR clock, 25 MYA date calibration8.45.113.3VIIIMyrmecophilesIGR clock, 25 MYA date calibration4.02.66.0VIIIMyrmecophilesIGR clock, 1% rate calibration7.45.49.6VIIIMyrmecophilesStrict clock, 25 MYA date calibration7.45.49.6VIIIMyrmecophilesStrict clock, 1% rate calibration <td>VI</td> <td>Myrmecophiles</td> <td>IGR clock, 1% rate calibration</td> <td>3.2</td> <td>1.7</td> <td>5.2</td>	VI	Myrmecophiles	IGR clock, 1% rate calibration	3.2	1.7	5.2
VIMyrmecophilesStrict clock, 1% rate calibration2.92.04.0VIIMyrmecophilesTK02 clock, 25 MYA date calibration13.89.918.0VIIMyrmecophilesTK02 clock, 1% rate calibration7.84.414.2VIIMyrmecophilesIGR clock, 25 MYA date calibration14.710.419.9VIIMyrmecophilesIGR clock, 1% rate calibration7.14.89.7VIIMyrmecophilesStrict clock, 25 MYA date calibration14.211.217.4VIIMyrmecophilesStrict clock, 1% rate calibration6.95.38.8VIIIMyrmecophilesTK02 clock, 25 MYA date calibration7.84.612.0VIIIMyrmecophilesTK02 clock, 1% rate calibration4.42.19.5VIIIMyrmecophilesIGR clock, 25 MYA date calibration8.45.113.3VIIIMyrmecophilesIGR clock, 25 MYA date calibration8.45.113.3VIIIMyrmecophilesIGR clock, 25 MYA date calibration4.02.66.0VIIIMyrmecophilesIGR clock, 25 MYA date calibration7.45.49.6VIIIMyrmecophilesStrict clock, 25 MYA date calibration7.45.49.6VIIIMyrmecophilesStrict clock, 1% rate calibration7.45.49.6VIIIMyrmecophilesStrict clock, 1% rate calibration7.45.49.6	VI	Myrmecophiles	Strict clock, 25 MYA date calibration	5.9	3.8	8.1
VIIMyrmecophilesTK02 clock, 25 MYA date calibration13.89.918.0VIIMyrmecophilesTK02 clock, 1% rate calibration7.84.414.2VIIMyrmecophilesIGR clock, 25 MYA date calibration14.710.419.9VIIMyrmecophilesIGR clock, 1% rate calibration7.14.89.7VIIMyrmecophilesStrict clock, 25 MYA date calibration14.211.217.4VIIMyrmecophilesStrict clock, 1% rate calibration6.95.38.8VIIIMyrmecophilesTK02 clock, 25 MYA date calibration7.84.612.0VIIIMyrmecophilesTK02 clock, 1% rate calibration4.42.19.5VIIIMyrmecophilesTK02 clock, 1% rate calibration8.45.113.3VIIIMyrmecophilesIGR clock, 25 MYA date calibration8.45.113.3VIIIMyrmecophilesIGR clock, 25 MYA date calibration4.02.66.0VIIIMyrmecophilesStrict clock, 25 MYA date calibration7.45.49.6VIIIMyrmecophilesStrict clock, 25 MYA date calibration7.45.49.6VIIIMyrmecophilesStrict clock, 1% rate calibration7.45.49.6	VI	Myrmecophiles	Strict clock, 1% rate calibration	2.9	2.0	4.0
VIIMyrmecophilesTK02 clock, 1% rate calibration7.84.414.2VIIMyrmecophilesIGR clock, 25 MYA date calibration14.710.419.9VIIMyrmecophilesIGR clock, 1% rate calibration7.14.89.7VIIMyrmecophilesStrict clock, 25 MYA date calibration14.211.217.4VIIMyrmecophilesStrict clock, 1% rate calibration6.95.38.8VIIIMyrmecophilesTK02 clock, 25 MYA date calibration7.84.612.0VIIIMyrmecophilesTK02 clock, 1% rate calibration4.42.19.5VIIIMyrmecophilesIGR clock, 25 MYA date calibration8.45.113.3VIIIMyrmecophilesIGR clock, 1% rate calibration4.02.66.0VIIIMyrmecophilesStrict clock, 25 MYA date calibration7.45.49.6VIIIMyrmecophilesStrict clock, 1% rate calibration7.45.49.6VIIIMyrmecophilesStrict clock, 1% rate calibration3.72.74.8	VII	Myrmecophiles	TK02 clock, 25 MYA date calibration	13.8	9.9	18.0
VIIMyrmecophilesIGR clock, 25 MYA date calibration14.710.419.9VIIMyrmecophilesIGR clock, 1% rate calibration7.14.89.7VIIMyrmecophilesStrict clock, 25 MYA date calibration14.211.217.4VIIMyrmecophilesStrict clock, 1% rate calibration6.95.38.8VIIIMyrmecophilesTK02 clock, 25 MYA date calibration7.84.612.0VIIIMyrmecophilesTK02 clock, 1% rate calibration4.42.19.5VIIIMyrmecophilesIGR clock, 25 MYA date calibration8.45.113.3VIIIMyrmecophilesIGR clock, 1% rate calibration4.02.66.0VIIIMyrmecophilesStrict clock, 25 MYA date calibration7.45.49.6VIIIMyrmecophilesStrict clock, 1% rate calibration7.45.49.6VIIIMyrmecophilesStrict clock, 1% rate calibration3.72.74.8	VII	Myrmecophiles	TK02 clock, 1% rate calibration	7.8	4.4	14.2
VIIMyrmecophilesIGR clock, 1% rate calibration7.14.89.7VIIMyrmecophilesStrict clock, 25 MYA date calibration14.211.217.4VIIMyrmecophilesStrict clock, 1% rate calibration6.95.38.8VIIIMyrmecophilesTK02 clock, 25 MYA date calibration7.84.612.0VIIIMyrmecophilesTK02 clock, 1% rate calibration4.42.19.5VIIIMyrmecophilesIGR clock, 25 MYA date calibration8.45.113.3VIIIMyrmecophilesIGR clock, 1% rate calibration4.02.66.0VIIIMyrmecophilesStrict clock, 25 MYA date calibration7.45.49.6VIIIMyrmecophilesStrict clock, 1% rate calibration3.72.74.8	VII	Myrmecophiles	IGR clock. 25 MYA date calibration	14.7	10.4	19.9
VIIMyrmecophilesStrict clock, 25 MYA date calibration14.211.217.4VIIMyrmecophilesStrict clock, 1% rate calibration6.95.38.8VIIIMyrmecophilesTK02 clock, 25 MYA date calibration7.84.612.0VIIIMyrmecophilesTK02 clock, 1% rate calibration4.42.19.5VIIIMyrmecophilesIGR clock, 25 MYA date calibration8.45.113.3VIIIMyrmecophilesIGR clock, 1% rate calibration4.02.66.0VIIIMyrmecophilesStrict clock, 25 MYA date calibration7.45.49.6VIIIMyrmecophilesStrict clock, 1% rate calibration3.72.74.8	VII	Myrmecophiles	IGR clock, 1% rate calibration	7.1	4.8	9.7
VIIMyrmecophilesStrict clock, 1% rate calibration6.95.38.8VIIIMyrmecophilesTK02 clock, 25 MYA date calibration7.84.612.0VIIIMyrmecophilesTK02 clock, 1% rate calibration4.42.19.5VIIIMyrmecophilesIGR clock, 25 MYA date calibration8.45.113.3VIIIMyrmecophilesIGR clock, 1% rate calibration4.02.66.0VIIIMyrmecophilesStrict clock, 25 MYA date calibration7.45.49.6VIIIMyrmecophilesStrict clock, 1% rate calibration3.72.74.8	VII	Myrmecophiles	Strict clock, 25 MYA date calibration	14.2	11.2	17.4
VIIIMyrmecophilesTK02 clock, 25 MYA date calibration7.84.612.0VIIIMyrmecophilesTK02 clock, 1% rate calibration4.42.19.5VIIIMyrmecophilesIGR clock, 25 MYA date calibration8.45.113.3VIIIMyrmecophilesIGR clock, 1% rate calibration4.02.66.0VIIIMyrmecophilesStrict clock, 25 MYA date calibration7.45.49.6VIIIMyrmecophilesStrict clock, 1% rate calibration3.72.74.8	VII	Myrmecophiles	Strict clock, 1% rate calibration	6.9	5.3	8.8
VIIIMyrmecophilesTK02 clock, 1% rate calibration4.42.19.5VIIIMyrmecophilesIGR clock, 25 MYA date calibration8.45.113.3VIIIMyrmecophilesIGR clock, 1% rate calibration4.02.66.0VIIIMyrmecophilesStrict clock, 25 MYA date calibration7.45.49.6VIIIMyrmecophilesStrict clock, 1% rate calibration3.72.74.8	VIII	Myrmecophiles	TK02 clock, 25 MYA date calibration	7.8	4.6	12.0
VIIIMyrmecophilesIGR clock, 25 MYA date calibration8.45.113.3VIIIMyrmecophilesIGR clock, 1% rate calibration4.02.66.0VIIIMyrmecophilesStrict clock, 25 MYA date calibration7.45.49.6VIIIMyrmecophilesStrict clock, 1% rate calibration3.72.74.8	VIII	Myrmecophiles	TK02 clock, 1% rate calibration	4.4	2.1	9.5
VIIIMyrmecophilesIGR clock, 1% rate calibration4.02.66.0VIIIMyrmecophilesStrict clock, 25 MYA date calibration7.45.49.6VIIIMyrmecophilesStrict clock, 1% rate calibration3.72.74.8	VIII	Myrmecophiles	IGR clock, 25 MYA date calibration	8.4	5.1	13.3
VIIIMyrmecophilesStrict clock, 25 MYA date calibration7.45.49.6VIIIMyrmecophilesStrict clock, 1% rate calibration3.72.74.8	VIII	Myrmecophiles	IGR clock, 1% rate calibration	4.0	2.6	6.0
VIII Myrmecophiles Strict clock, 1% rate calibration 37 2.7 4.8	VIII	Myrmecophiles	Strict clock, 25 MYA date calibration	7.4	5.4	9.6
	VIII	Myrmecophiles	Strict clock, 1% rate calibration	3.7	2.7	4.8

Table 3. Number of segregating sites (S) and average number of nucleotide differences (k) between Eciton species, 707 between the Ecitophya and Ecitomorpha, and between species within each of these genera; based on unique

haplotypes (802bp mtDNA COI).

	Comparison	S	k
Ants			
Eciton	b. parvispinum – dulcium	104	103.5
	b. parvispinum – hamatum	100	94.33
	b. foreli – dulcium	100	91.67
	b. parvispinum – lucanoides	88	88.00
	lucanoides – dulcium	86	85.50
	b. foreli – lucanoides	90	81.83
	b. foreli – b. parvispinum	90	81.00
	b. foreli – hamatum	92	79.28
	hamatum - dulcium	81	74.33
	hamatum – lucanoides	80	73.33
Beetles	Ecitomorpha – Ecitophya	185	103.98
Ecitomorpha	arachnoides (foreli) – arachnoides (parvispinum)	19	33.90
Ecitophya	simulans (foreli) – rettenmeyeri	82	77.30
	gracillima – rettenmeyeri	97	74.67
	simulans (parvispinum) – rettenmeyeri	79	73.71
	simulans (parvispinum) – gracillima	72	42.43
	simulans (foreli) – simulans (parvispinum)	54	41.81
	simulans (foreli) – gracillima	55	24.83

711 712 713 714 715  $\begin{array}{c} 716\\ 717\\ 718\\ 719\\ 720\\ 721\\ 722\\ 723\\ 724\\ 725\\ 726\\ 727\\ 728\\ 729\\ 730\\ 731\\ 732\\ 733\\ 734\\ 735\\ 736\end{array}$ 

740 741 742 **Table 4.** Population structure estimates derived from haplotypic data from *E. burchelli foreli* and their associated populations of *Ecitophya* and *Ecitomorpha* myrmecophile beetles between the areas studied in West (BPPS) and Central Panama (APSLA, APSLB). \* P < 0.001, NS non-significant

Source of variation	df	SSQ	Variance	% variation	Fixation index			
Eciton burchellii foreli								
A	1	1120 650	76.000	76.01	÷ 0.0070#			
Among groups	1	1139.658	/6.888	/6.31	$\Phi_{\rm ST} = 0.9979^*$			
Among populations within groups	1	210.061	23.660	23.48	$\Phi_{SC} = 0.9914*$			
Within populations	28	5.765	0.206	0.20	$\Phi_{\rm CT} = 0.7631$			
Total	30	1355.484	100.574					
Ecitophya simulans								
Among groups	1	107.796	8.859	84.62	Φ <sub>ST</sub> =0.8449*			
Among populations within groups	1	1.522	-0.014	-0.13	$\Phi_{SC} = -0.0088NS$			
Within populations	24	38.978	1.624	15.51	$\Phi_{\rm CT} = 0.8462^{*}$			
Total	26	148.296	10.469					
Ecitomorpha arachnoides								
Among groups	1	218.331	14.713	79.57	$\Phi_{ST} = 0.7797*$			
Among populations within groups	1	1.032	-0.296	-1.59	$\Phi_{SC} = -0.0783 NS$			
Within populations	48	195.5	4.073	22.03	$\Phi_{\rm CT} = 0.7957*$			
Total	50	414.863	18.491		01			