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Ecological specialization is associated with genetic structure in the ant-associated butterfly family Lycaenidae

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The role of specialization in diversification can be explored along two geological axes in the butterfly family Lycaenidae. In addition to variation in hostplant specialization normally exhibited by butterflies, the caterpillars of most Lycaenidae have symbioses with ants ranging from no interactions through to obligate and specific associations, increasing niche dimensionality in ant-associated taxa. Based on mitochondrial sequences from 8282 specimens from 967 species and 249 genera, we show that the degree of ecological specialization of lycaenid species is positively correlated with genetic divergence, haplotype diversity and an increase in isolation by distance. Nucleotide substitution rate is higher in carnivorous than phytophagous lycaenids. The effects documented here for both micro- and macroevolutionary processes could result from increased spatial segregation as a consequence of reduced connectivity in specialists, niche-based divergence or a combination of both. They could also provide an explanation for the extraordinary diversity of the Lycaenidae and, more generally, for diversity in groups of organisms with similar multi-dimensional ecological specialization.

1. Background

'What drives the diversification of life?' is a fundamental question in biology. Although the role of long-term geographical isolation in giving rise to the divergence between allopatric populations is well known [1], the idea that speciation may also result from ecologically based divergent selection driving the evolution of reproductive isolation has seen a recent revival (e.g. 'speciation-with-gene-flow') [2]. For example, ecological specialization of conspecific herbivorous insects feeding on different host-plants (host races) [3] may enhance diversity within species [4] and promote the evolution of barriers to gene flow via host-associated differentiation [5]. The formation of host races appears to be associated with subsequent speciation and radiation in some cases [6] (but see [7,8]).

Rates of molecular evolution have been shown to accelerate in specialized parasitic [9,10] as well as mutualistic symbiont lineages [11,12] compared with generalists. However, theoretical predictions disagree as to how coevolution should affect rates of molecular evolution in those two cases. According to the 'red queen hypothesis', coevolving organisms must continuously adapt and



Figure 1. Distribution map of the 8282 samples of lycaenid butterflies used in this study grouped by their life-history strategy in regard to specialization. (Online version in colour.)

evolve in order to survive in an environment of ever-evolving antagonistic organisms if their interaction is locked [13,14]. Thus, accelerated rates of molecular evolution can be expected in antagonistic (parasitic) relationships. On the other hand, this is not always true in 'red king' scenarios involving mutualists where the symbiont evolving mutualististic traits more quickly could end up providing more help than it receives in return [15].

The butterfly family Lycaenidae is globally distributed and unusually diverse, particularly considering that it is among the youngest families of Lepidoptera (approx. 90 Myr old) [16]. The juveniles of many lycaenid species engage in symbiotic associations with ants (myrmecophily). Their caterpillars typically possess multiple, specialized ant-associated organs that form the basis for this association by appeasing ants that might otherwise be predators [17]. In return, the ants protect the caterpillars from natural enemies [18]. Degree of myrmecophily in the Lycaenidae ranges from no interactions, to obligate ones [19]. Most described lycaenid– ant interactions are facultative and appear to be mutualistic. These loose, facultative symbioses are not specific, and the caterpillars of a single lycaenid species can be found associating with up to 11 different species of ants [20].

About 30% of the Lycaenidae are obligately ant-associated in the sense that juveniles are never found without ants and often only associate with a single or closely related group of ant species [19]. Obligate lycaenid–ant symbioses can be mutualistic or parasitic. Parasitic caterpillars typically enter the ant nest where they either eat the brood directly or solicit the workers for food regurgitations (kleptoparasitism). Myrmecophilous and non-myrmecophilous species occur in most lycaenid subfamilies, although a few, such as the Lycaeninae and Lipteninae, are rarely ant associated [19]. The often specific interactions between lycaenids and ants add another dimension to the lycaenids' ecological specialization beyond the host-plant associations of most Lepidoptera, providing a potential explanation for their disproportionally high diversity [21,22].

The hypothesis of 'ecological islands' [23,24] (suitable areas where host plants and ant associates intersect)

affecting genetic structure has been tested with data from 34 species of Lycaenidae collected in Romania [25]. Effective population sizes were found to be smaller in mutualistic species with a 'high degree of myrmecophily' compared with species with a 'low degree of myrmecophily' [25]: ant specialization may drive diversification in the Lycaenidae by decreasing effective population sizes. In addition, several studies have investigated the possibility of genetic differentiation between conspecific lycaenid populations associated with different ants, without finding any evidence [26–28].

Here we re-evaluate the possible association between genetic structure and myrmecophily, and more generally ecological specialization in the Lycaenidae, using a substantially larger and more representative dataset than has been assembled before. We explore the hypothesis that host-ant and host-plant specialization, as well as dual specialization on ants and plants simultaneously and carnivory, are associated with both micro- and macroevolutionary patterns.

2. Methods

A total of 8282 samples representing eight subfamilies, 249 genera and 967 species of Lycaenidae from all over the world were analysed (figure 1; electronic supplementary material, table S1). The degree of specialization on host ants and host plants for each species is shown in electronic supplementary material, table S2. Phylogenetic relationships between species based on the withinspecies consensus of the mitochondrial gene cytochrome oxidase subunit I (COI) are shown in electronic supplementary material, figure S1. Publicly available COI sequences for the family Lycaenidae were downloaded from the Barcoding Of Life Data Systems (BOLD) or GenBank. Additionally, 707 samples were sequenced to complete the dataset for underrepresented species. DNA extraction, PCR reactions and sequencing were similar to the conditions described in [29] using the primer pair 'LCO1490' (forward) [30] and 'Nancy' (reverse) [31]. GENEIOUS v. 6.1.2 (created by Biomatters; http://www.geneious.com/) was used for assembly of chromatograms and to edit alignments. A maximum-likelihood phylogenetic tree for the investigated species (electronic supplementary material, figure S1) was obtained based on the majority-rule consensus DNA barcodes for each species using



Figure 2. Coefficient plots from the three generalized linear mixed models investigating microevolution using the full dataset. The response variables are maxima of intraspecific genetic divergence (a), haplotype diversity, h (b) and IBD regression coefficients, r (c). Coefficients adjusted for comparison by division by two s.d. are shown for explanatory variables included in the models (mean: dots, standard deviations, s.d.: bold lines, 2s.d.: thin lines).

the program RAxML [32] through the raxmlGUI v. 1.5b1 interface [33] using the GTR + G model of sequence evolution with 100 bootstraps. Monophyly of subfamilies and their relationships were constrained according to [34]. Species for which at least five sequenced specimens with published sampling location existed were selected for microevolutionary analysis of intraspecific lineage accumulation. Location descriptions lacking coordinates were geo-referenced (Google Maps). Degree of ant specialization was quantified using a binary categorical predictor with the levels 'ant specialized' (obligate association with ants) and 'not ant specialized' (not obligatorily associated with ants), based on literature records and expert opinions (electronic supplementary material, table S1). Plant specialization was similarly used as a binary categorical variable with the levels 'plant specialized' (feeding on only one plant genus) and 'not plant specialized' (feeding on more than one plant genus). To investigate a third type of specialization, 'carnivorous' (defined here as strictly carnivorous during all larval stages) versus 'not carnivorous', diet was included as a binary categorical predictor, based on literature records (electronic supplementary material, table S1), the Lepidoptera host plant database of the Natural History Museum (http://www.nhm.ac.uk/our-science/data/hostplants/) and expert opinions (electronic supplementary material, table S2; Acknowledgements). As a possible confounder, body size (measured as wingspan, based on data published in field guides) was included (electronic supplementary material, table S2). Associations between levels of specialization and multiple genetic parameters, related to both micro- and macroevolution processes, were analysed.

(a) Microevolution

Matrices of divergence (K80 model of DNA evolution [35]) between all combinations of individual sequences and haplotype diversity [36] were calculated for species using the R packages 'ape' [37], 'adephylo' [38] and 'Biostrings' [39] in R v. 3.2.2 [40]. Average geographical distances between samples were calculated per species using the 'sp' package [41]. Associations between measures of intraspecific lineage accumulation (maximum divergence and haplotype diversity, h), and specialization categories were analysed using generalized linear mixed models in the R package 'MCMCglmm' [42]. Maximum divergence (square root transformed) and h were included as independent, Gaussian variables. Ant and plant specialization categories, double specialization (on ants and plants), carnivory, sample size and average geographical distance between samples of the same species were

included as fixed effects. Phylogeny was included as a random effect (as a phylogenetic covariance matrix [43]). Possible correlations between significant predictors were assessed using Pearson's product-moment correlation. There was a significantly positive correlation between body size and specialization (Pearson's product–moment correlation: $t_{129} = 2.9$, p = 0.004), but not between body size and average geographical distance (Pearson's product–moment correlation: $t_{129} = 1.24$, p = 0.217). Isolation by distance (IBD) was investigated for each species using Mantel tests comparing matrices of pairwise genetic distances (K80 model of DNA evolution [35]) between individuals (R package 'Biostrings' and 'ape') with their geographical distances (R package 'sp') using the R package 'adegenet' [38]. The association between the resulting correlation coefficients (r) and specialization was investigated in 'MCMCglmm', with Gaussian residual distribution. The dependent variable r was predicted by the fixed effects wingspan, average geographical distance, sample size, carnivory, ant and plant specialization, and their interaction. Phylogeny was included as a random effect.

(b) Macroevolution

Molecular substitution rate was analysed as the branch length of each species (tip) to the root, extracted from the ML phylogeny described above. Branch lengths to root were log-transformed for the analysis in order to approach a normal distribution. Ant- and plant specialization, their interaction, carnivory and average geographical distance between samples were included as fixed predictors, and a phylogeny without branch lengths was included as a random effect.

All MCMCglmm models were run for 1.3×10^6 iterations with a burn-in period of 3×10^5 iterations (thinning interval: 500 iterations). In order to investigate a possible bias introduced by unequal sample sizes, models for analysis of microevolutionary processes were repeated with average values of each parameter obtained from 10^3 iterations of a random selection of five sequences per species. In addition, all models were repeated comparing the subset of mutualistic obligate ant symbionts to parasitic ones. The R scripts used are available in electronic supplementary material, appendix S1.

3. Results

Species whose larvae specialize on plants had higher maxima of intraspecific genetic divergence than their **Table 1.** Results from generalized linear mixed models using the 'MCMCgImm' R package [42] with 1.3×10^6 iterations, and a burn-in period of 3×10^5 iterations. Phylogeny (electronic supplementary material, figure S1) was included as a random effect in all models. Significant *p*-values are printed in italics.

		nostorior	lower 95%	upper 95% confidence	effective	мсмс
response variable	fixed effects	mean	limit	limit	size	<i>p</i> -value
microsulution						
microevolution	ant enocialization	0 02002	0.00110	0.06427	2000	0.062
samples		0.02902	0.00110	0.00437	2000	0.005
	plant specialization	0.01957	0.00142	0.03466	2000	0.043
		0.02009	- 0.07755	0.02400	2121	0.510
		0.02110	0.00001	0.0002	2000	0.424
	samples	0.00002	0.00001	0.00005	2310	< 0.001
	sample size	0.00009	-0.00002	0.00020	1836	0.103
	body size (wingspan mm)	0.00164	0.00059	0.00273	2000	0.004
	obligate ant parasites versus obligate ant mutualists	0.00179	-0.08052	0.07748	2000	0.955
	average geographical distance between samples	0.00003	-0.00001	0.00007	2000	0.064
	sample size	0.00006	-0.00041	0.00056	2000	0.827
	body size (wingspan mm)	0.00229	-0.00095	0.00572	2000	0.164
haplotype diversity (h), all	ant specialization	0.07692	-0.08681	0.22820	1741	0.326
samples	plant specialization	0.13150	0.03575	0.22000	2000	0.008
	ant specialization and plant specialization	0.01256	-0.23530	0.27780	1904	0.920
	carnivory	0.09692	-0.12020	0.32950	2000	0.394
	average geographical distance between	0.00006	0.00002	0.00010	2000	0.003
	samples					
	sample size	-0.00010	-0.00063	0.00045	2000	0.732
	body size (wingspan mm)	0.00617	0.00079	0.01159	2215	0.025
	obligate ant parasites versus obligate ant mutualists	-0.04421	-0.32710	0.24460	2548	0.724
	average geographical distance between samples	0.00005	-0.00007	0.00019	2000	0.448
	sample size	-0.00066	-0.00241	0.00119	2085	0.456
	body size (wingspan mm)	0.00249	-0.00893	0.01324	2000	0.624
IBD coefficient (r)	ant specialization	0.19730	-0.01266	0.39530	2000	0.063
	plant specialization	0.08445	-0.02880	0.22410	2000	0.191
	ant specialization and plant specialization	-0.13190	-0.47370	0.26230	2333	0.476
	carnivory	0.25710	-0.02395	0.52020	1763	0.072
	average geographical distance between samples	0.00002	-0.00004	0.00007	1827	0.595
	sample size	-0.00024	-0.00101	0.00054	2000	0.540
	body size (wingspan mm)	0.00347	-0.00400	0.01189	2367	0.381
	obligate ant parasites versus obligate ant mutualists	-0.10310	-0.42890	0.24340	2191	0.521
	average geographical distance between samples	0.00005	-0.00010	0.00021	2000	0.504
	sample size	-0.00103	-0.00302	0.00100	1768	0.312
	body size (wingspan mm)	0.00403	- 0.00899	0.01730	2295	0.518
macroevolution						
molecular substitution rate	ant specialization	-0.01577	-0.05338	0.02656	2000	0.460
(branch length to root)	plant specialization	0.01510	-0.00853	0.03656	2000	0.210
	ant specialization and plant specialization	0.05319	-0.00406	0.10790	1521	0.067
	carnivory	0.09501	0.02730	0.16563	2000	0.012



Figure 3. Coefficient plot from the generalized linear mixed model investigating macroevolution using the full dataset. The response variable is molecular substitution rate (branch length). Coefficients are shown for explanatory variables included in the model (mean: dots; standard deviations, s.d.: bold lines, 2 s.d.: thin lines). (Online version in colour.)

generalist counterparts, and a similar trend was found for ant specialized and carnivorous species (figure 2a and table 1; electronic supplementary material, appendix S2, table S3). Maximum divergence also increased with average geographical distance and body size (table 1 and figure 2*a*). The species mean for maximum divergence was $0.015 \pm$ 0.014 (s.d.), ranging between 0 and 0.086. With the mean of 1000 iterations of five randomly chosen specimens per species, all results were consistent with results obtained from the full dataset (electronic supplementary material, appendix S2, table S3). Haplotype diversity was higher for species with a high degree of plant specialization than for generalist plant feeders (table 1). Haplotype diversity was also correlated with average geographical distance and body size (table 1). Average h was 0.592 ± 0.274 , ranging between 0 and 0.969. Results for h were equivalent to those obtained for the full dataset when five randomly selected sequences per species were used (mean of 1000 iterations; electronic supplementary material, appendix S2, table S3). There was a trend of regression coefficients obtained in Mantel tests for IBD being higher in carnivorous than in non-carnivorous species, and in ant-specialized compared with not ant-specialized species (table 1 and figure 2*c*). Average geographical distance between conspecific individuals was also not correlated with the strength of IBD of that species (table 1). The average for r was 0.398 ± 0.351 ranging between -0.328 and 1. When five specimens were randomly selected per species (1000 iterations), the results were similar to those obtained in the analysis of all specimens (electronic supplementary material, appendix S2, table S3). None of the investigated microevolutionary response variables differed significantly between species whose larvae are either obligate mutualists or parasites of ants (table 1). Molecular substitution rate (branch length) was significantly higher in carnivorous compared with non-carnivorous species (table 1 and figure 3).

4. Discussion

We found a positive association between genetic structure and ecological specialization. Species' maximum genetic divergence and h are higher in lycaenids that are specialists on host-plants compared to generalists (figure 2a). Alternatively, specialization may be promoted in species with increased genetic structure (due to dispersal limitation, fragmentation, etc.). These results could reflect reduced gene flow among populations due to decreased dispersal, stronger fragmentation and/or disruptive selection in specialists. Decreased dispersal in specialists may result from selection due to coevolution with local ant/plant associates. That is, selection may favour symbionts specializing on local partners because of increased fitness benefits of co-adaptation. Disruptive effects of distance and/or biogeographical barriers may be magnified in specialists because of fewer 'stepping stones' (suitable habitat patches), limiting dispersal. Disruptive selection in species specializing on different host plants in sympatry is known for phytophagous insects [44]. Such host races could exist within butterfly species living in close association with different ant species. Here we found that h was indeed higher in plant specialists than generalists (figure 2b and table 1). However, patterns consistent with IBD increased in carnivores, suggesting stronger spatial segregation in these species, and there was a general trend of stronger IBD patterns with increased specialization (figure 3). This suggests that diversification is not driven purely by ant specialization in sympatry, but that distance or biogeographic barriers play a role. Certain highly specialized lycaenid species have been shown to use ants themselves as cues in egg-laying, a mechanism that could potentially produce an amplification of their host-plant range [45,46]. This could in turn lead to greater opportunities for diversification if different host-plants occupy different ecological niches.

Molecular substitution rate (branch length from tip to root in the species tree) increased significantly in strictly carnivorous species, but not in moderately specialized species (on only ants or plants; figure 3). Thus, the effects at the macroevolutionary scale (rate of molecular evolution) were only visible in extreme specialists (carnivorous). Nevertheless, at least in the most extreme cases, the microevolutionary mechanisms documented here seem to be translated to macroevolutionary scales. Accelerated rates of molecular evolution have previously been documented in parasites, and this phenomenon has been explained by their typically small population sizes [9,10].

Specialists could be more susceptible to fragmentation and reduced population sizes than generalists during broad-scale environmental changes such as climatic oscillations. It is frequently argued that specialist species are more likely to go extinct due to narrower and more complex habitat requirements (specialization as an evolutionary dead end; but see [47,48]). However, interactions involving that are simultaneously specialized lycaenids with respect to both attendant ants and host plants simultaneously were not significantly correlated with divergence, h and r (table 1). Perhaps these dual specialists have reached a point of ecological refinement where they are simply constrained from further population subdivision.

In summary, increased genetic structuring of specialists could be a result of smaller population sizes, reduced dispersal, ant-induced host-plant shifts, divergent selection on

different ecological niches, relaxed selection or any combination of these factors. The assessment of ecological niche width and its translation to total range and connectivity of habitats in geographical space and time (the factors potentially linked to genetic population structure) are difficult. Moreover, the dimensions analysed here for life-history specialization do not cover those species that specialize in additional dimensions. Future comparative studies, including detailed host use data (for both ants and plants) at a population level for a large number of species with different degrees of specialization, may help to identify mechanisms leading to the observed increase in genetic divergence in specialists. Comparative analyses at an intraspecific level shed light on microevolutionary processes that may give rise to macroevolutionary patterns of diversification. Our results support the importance of ecological specialization in general, and of species interactions in particular, in generating biodiversity.

Ethics. This study has been conducted using published DNA sequences or collection material that belongs to the Museum of Comparative Zoology, and procedures have been subject to relevant institutional oversight.

Data accessibility. All primary data are accessible via the electronic supplementary material, GenBank or BOLD (see electronic

supplementary material, table S1 for individual GenBank Accession numbers and BOLD access codes).

Authors' contributions. N.E.P., R.V., R.E., D.R.N. and S.S. conceived of the study; N.E.P., R.V., R.E., S.S., A.K. and M.E. contributed with DNA material; S.S., K.G.A., A.K., J.H.B. and M.E. performed laboratory work; S.S., G.T. and D.R.N. analysed data; all authors contributed to writing the manuscript.

Competing interests. We declare we have no competing interests.

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References

- Dobzhansky T, Dobzhansky T. 1937 Genetics and the origin of species. New York, NY: Columbia University Press.
- 2. Nosil P. 2012 *Ecological speciation*. Oxford, UK: Oxford University Press.
- Bush G. 1969 Sympatric host race formation and speciation in frugivorous flies of the genus *Rhagoletis* (Diptera, Tephritidae). *Evolution* 23, 237–251. (doi:10.2307/2406788)
- Jaenike J. 1990 Host specialization in phytophagous insects. *Annu. Rev. Ecol. Evol. Syst.* 21, 243–273. (doi:10.1146/annurev.es.21.110190.001331)
- Pashley D. 1986 Host-associated genetic differentiation in fall armyworm (Lepidoptera: Noctuidae): a sibling species complex. *Ann. Entomol. Soc. Am.* 79, 898–904. (doi:10.1093/aesa/79.6.898)
- Berlocher S, Feder J. 2002 Sympatric speciation in phytophagous insects: moving beyond controversy? *Annu. Rev. Entomol.* 47, 773–815. (doi:10.1146/ annurev.ento.47.091201.145312)
- 7. Coyne J, Orr H. 2004 *Speciation*. Sunderland, MA: Sinauer Associates.
- Linnen CR, Farrell BD. 2010 A test of the sympatric host race formation hypothesis in Neodiprion (Hymenoptera: Diprionidae). *Proc. Soc. B* 277, 3131–3138. (doi:10.1098/rspb.2010.0577)
- Paterson S *et al.* 2010 Antagonistic coevolution accelerates molecular evolution. *Nature* 464, 275–278. (doi:10.1038/nature08798)
- Bromham L, Cowman PF, Lanfear R. 2013 Parasitic plants have increased rates of molecular evolution across all three genomes. *BMC Evol. Biol.* 13, 126. (doi:10.1186/1471-2148-13-126)
- 11. Lutzoni F, Pagel M. 1997 Accelerated evolution as a consequence of transitions to mutualism. *Proc. Natl*

Acad. Sci. USA **94**, 11422–11427. (doi:10.1073/ pnas.94.21.11422)

- Rubin BER, Moreau CS. 2016 Comparative genomics reveals convergent rates of evolution in ant – plant mutualisms. *Nat. Commun.* 7, 12679. (doi:10.1038/ ncomms12679)
- Van Valen L. 1973 A new evolutionary law. *Evol. Theory* 1, 1–30.
- Suchan T, Alvarez N. 2015 Fifty years after Ehrlich and Raven, is there support for plant – insect coevolution as a major driver of species diversification? *Entomol. Exp. Appl.* **157**, 98–112. (doi:10.1111/eea.12348)
- Bergstrom CT, Lachmann M. 2003 The Red King effect: when the slowest runner wins the coevolutionary race. *Proc. Natl Acad. Sci. USA* **100**, 593–598. (doi:10.1073/pnas.0134966100)
- Kawahara AY, Breinholt JW. 2014 Phylogenomics provides strong evidence for relationships of butterflies and moths. *Proc. R. Soc. B* 281, 20140970. (doi:10.1098/rspb.2014.0970)
- Malicky H. 1970 New aspects of the association between lycaenid larvae (Lycaenidae) and ants (Formicidae, Hymenoptera). *J. Lepid Soc.* 24, 190–202.
- Pierce N, Mead P. 1981 The evolution and biogeography of association between Lycaenid butterflies and ants. *Science* **211**, 1185–1187. (doi:10.1126/science.211.4487.1185)
- Pierce N, Braby M, Heath A, Lohman D, Mathew J, Rand D, Travassos M. 2002 The ecology and evolution of ant association in the Lycaenidae (Lepidoptera). *Annu. Rev. Entomol.* 47, 733–771. (doi:10.1146/annurev.ento.47.091201. 145257)

- Fiedler K. 1991 Systematic, evolutionary, and ecological implications of myrmecophily within the Lycaenidae (Insecta: Lepidoptera: Papilionoidea). Bonn, Germany: Zoologisches Forschungsinstitut und Museum Alexander Koenig.
- Pierce N. 1987 The evolution and biogeography of association between Lycaenid butterflies and ants. In Oxford surveys in evolutionary biology (eds. P Harvey, L Partridge), pp. 89–116. New York, NY: Oxford University Press.
- Pellissier L, Kostikova A, Litsios G, Salamin N, Alvarez N. 2017 High rate of protein coding sequence evolution and species diversification in the Lycaenids. *Front. Ecol. Evol.* 5, 90. (doi:10.3389/fevo. 2017.00090)
- Janzen DH. 1968 Host plants as islands in evolutionary and contemporary time. *Am. Nat.* 102, 592–595. (doi:10.1086/282574)
- Opler PA. 1974 Oaks as evolutionary islands for leafmining insects: the evolution and extinction of phytophagous insects is determined by an ecological balance between species diversity and area of host occupation. *Am. Sci.* 62, 67–73.
- Pellissier L, Litsios G, Guisan A, Alvarez N. 2012 Molecular substitution rate increases in myrmecophilous lycaenid butterflies (Lepidoptera). *Zool. Scr.* 41, 651–658. (doi:10.1111/j.1463-6409. 2012.00556.x)
- Costa JT, Mcdonald JH, Pierce NE. 1996 The effect of ant association on the population genetics of the Australian butterfly *Jalmenus evagoras* (Lepidoptera: Lycaenidae). *Biol. J. Linn. Soc.* 58, 287–306. (doi:10.1111/j.1095-8312.1996.tb01436.x)
- 27. Eastwood R, Pierce NE, Kitching RL, Hughes JM. 2006 Do ants enhance diversification in lycaenid

butterflies? Phylogeographic evidence from a model myrmecophile, *Jalmenus evagoras. Evolution* **60**, 315–327. (doi:10.1111/j.0014-3820.2006. tb01109.x)

- Sielezniew M, Rutkowski R. 2012 Population isolation rather than ecological variation explains the genetic structure of endangered myrmecophilous butterfly Phengaris (=Maculinea) arion. J. Insect Conserv. 16, 39–50. (doi:10.1007/ s10841-011-9392-9)
- Kaliszewska ZA, Lohman DJ, Sommer K, Adelson G, Rand DB, Mathew J, Talavera G, Pierce NE. 2015 When caterpillars attack: Biogeography and life history evolution of the Miletinae (Lepidoptera: Lycaenidae). *Evolution* 69, 571–588. (doi:10.1111/ evo.12599)
- Folmer O, Black M, Hoeh W, Lutz R, Vrijenhoek R. 1994 DNA primers for amplification of mitochondrial cytochrome c oxidase subunit 1 from diverse metazoan invertebrates. *Mol. Mar. Biol. Biotechnol.* 3, 294–299.
- Simon C, Frati F, Beckenbach A, Crespi B, Liu H, Flook P. 1994 Evolution, weighting, and phylogenetic utility of mitochondrial gene sequences and a compilation of conserved polymerase chain reaction primers. *Ann. Entomol. Soc. Am.* 87, 651–701. (doi:10.1093/aesa/87.6.651)
- Stamatakis A. 2014 RAxML version 8: a tool for phylogenetic analysis and post-analysis of large phylogenies. *Bioinformatics* **30**, 1312–1313. (doi:10.1093/bioinformatics/btu033)

- Silvestro D, Michalak I. 2012 raxmlGUI: a graphical front-end for RAxML. *Organ. Divers. Evol.* **12**, 335–337. (doi:10.1007/s13127-011-0056-0)
- Wahlberg N *et al.* 2005 Synergistic effects of combining morphological and molecular data in resolving the phylogeny of butterflies and skippers. *Proc. Soc. B* 272, 1577–1586. (doi:10.1098/rspb.2005.3124)
- Kimura M. 1980 A simple method for estimating evolutionary rates of base substitutions through comparative studies of nucleotide sequences. *J. Mol. Evol.* 16, 111–120. (doi:10.1007/BF01731581)
- Nei M, Tajima F. 1981 DNA polymorphism detectable by restriction endonucleases. *Genetics* 97, 145–163.
- Paradis E, Claude J, Strimmer K. 2004 APE: analyses of phylogenetics and evolution in R language. *Bioinformatics* 20, 289–290. (doi:10.1093/ bioinformatics/btq412)
- Jombart T. 2008 adegenet: a R package for the multivariate analysis of genetic markers. *Bioinformatics* 24, 1403–1405. (doi:10.1093/ bioinformatics/btn129)
- Pages H, Aboyoun P, Gentleman R, DebRoy S. 2014 Biostrings: string objects representing biological sequences, and matching algorithms. R package version 2.38.4 See https://bioconductor.org/ packages/release/bioc/html/Biostrings.html.
- R Development Core Team. 2014 R: a language and environment for statistical computing. Vienna, Austria: R Foundation for Statistical Computing. See http://www.R-project.org/.

- Pebesma E, Bivand R. 2005 Classes and methods for spatial data in R. *R News* 5, 9–13.
- Hadfield J. 2010 MCMC methods for multi-response generalized linear mixed models: the MCMCglmm R package. J. Stat. Softw. 33, 1–22. (doi:10.18637/ jss.v033.i02)
- Hadfield JD, Nakagawa S. 2010 General quantitative genetic methods for comparative biology: phylogenies, taxonomies and multi-trait models for continuous and categorical characters. *J. Evol. Biol.* 23, 494-508. (doi:10.1111/j.1420-9101.2009. 01915.x)
- Nosil P. 2007 Divergent host plant adaptation and reproductive isolation between ecotypes of *Timema cristinae* walking sticks. *Am. Nat.* **169**, 151–162. (doi:10.1086/510634)
- Pierce NE, Elgar MA. 1985 The influence of ants on host plant selection by *Jalmenus evagoras*, a myrmecophilous lycaenid butterfly. *Behav. Ecol. Sociobiol.* 16, 209–222. (doi:10.1007/ BF00310983)
- Fiedler K. 1994 Lycaenid butterflies and plants: is myrmecophily associated with amplified hostplant diversity? *Ecol. Entomol.* **19**, 79–82. (doi:10.1111/j. 1365-2311.1994.tb00393.x)
- 47. Futuyma DJ, Moreno G. 1988 The evolution of ecological specialization. *Annu. Rev. Ecol. Evol. Syst.* 19, 207–233. (doi:10.1146/annurev.es.19.110188. 001231)
- 48. Thompson J. 1994 *The coevolutionary process*. Chicago, IL: University of Chicago Press.