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Abstract

Comparative studies of insect behaviour based on evolutionary trees are currently blossoming, because of the increasing ease of phylogeny estimation, the availability of new trait data to analyse, and a vast and growing array of statistical techniques for exploring data and testing hypotheses. These studies address not only the selective forces and constraints on insect behaviour, which are the realm of traditional behavioral ecology, but also their ecological and evolutionary consequences. Recent studies have significantly increased our understanding of foraging behaviour, interspecific interactions, locomotion and dispersal, communication and signalling, mate choice and sexual selection, parental care and the evolution of sociality. The curating of trait data remains a significant challenge to maximize the future potential of insect comparative studies.

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Highlights

- Insect comparative studies help us to learn about the evolution of phenotypic diversity
- They use cross-taxonomic data, and require a phylogeny and analytical methods
- Trait data is still difficult to acquire and collate
- Phylogenetic data on insects have grown, as have analytical methods
- Recent studies have improved our understanding of the causes and consequences of all the major traits studied by behavioural ecologists

Comparative analysis of behavioural traits in insects

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Abstract

- 2 Comparative studies of insect behaviour based on evolutionary trees are currently
- 3 blossoming, because of the increasing ease of phylogeny estimation, the availability
- 4 of new trait data to analyse, and a vast and growing array of statistical techniques for
- 5 exploring data and testing hypotheses. These studies address not only the selective
- 6 forces and constraints on insect behaviour, which are the realm of traditional
- behavioural ecology, but also their ecological and evolutionary consequences. Recent
- 8 studies have significantly increased our understanding of foraging behaviour,
- 9 interspecific interactions, locomotion and dispersal, communication and signalling,
- mate choice and sexual selection, parental care and the evolution of sociality. The
- curating of trait data remains a significant challenge to maximize the future potential
- of insect comparative studies.

Introduction

13

14 Comparative analyses take variation across taxa as a source of data with which to test 15 hypotheses [1,2] (Figure 1). Such tests often draw together large numbers of 16 observations to provide a more holistic picture than studies on individual species can, 17 and they relate to real-world situations (the data are often traits observable in the 18 field), which is not necessarily true of experimental studies. Cross taxonomic data are 19 also often readily available, and can show much wider variation than is obtained from 20 single species or experimental studies. For these reasons they have wide appeal. The 21 main limitations of comparative studies are that they are observational and often 22 correlative in nature, hence cannot so robustly inform causation, and they are mostly 23 limited to existing variation, which experimental studies are not. However, 24 comparative analyses and experimental studies do overlap in the form of meta-25 analyses, where the results of many experimental studies, often involving several 26 species, can be brought together to give holistic experimental tests of hypotheses 27 [3,4].28 Because cross-taxonomic variation potentially has an evolutionary basis, and 29 the hypotheses that are tested are frequently explicitly evolutionary in nature, this 30 brings analytical challenges that were first widely formalized in the 1980s, coincident 31 with the development of computational methods for reconstructing phylogenies. It 32 was then recognized that phylogenies themselves can help overcome those challenges 33 [1]. In those days, phylogeny-based comparative approaches were often presented as 34 ways to avoid falling into naïve statistical traps (such as treating species as 35 independent data points), but more recently, since the development of likelihood-36 based and Bayesian computational techniques, along with information theoretic 37 approaches, the emphasis has been on finding appropriate evolutionary models that 38 explain the data well [2]. Techniques have diversified to incorporate an increasingly 39 sophisticated range of data types and approaches (Table 1), although these can mostly 40 be reduced to a small number of basic tasks, such as reconstructing of ancestral states 41 and detecting evolutionary associations between traits (Figure 1, Table 1). To apply 42 these techniques, a well resolved, and preferably dated, phylogeny is often essential 43 [2]. 44 In the past, and still to some extent today [5], phylogenetic requirements could 45 present an obstacle for comparative studies, especially of insects. However, good

quality phylogenetic information for insects is becoming more and more routinely produced thanks to widely available molecular markers [6,7], the development of whole genome and transcriptome approaches [8,9], and an increasingly better-known fossil record that provides the calibration points for dating analyses [10,11]. In addition to studies collecting primary morphological or molecular data, pipelines are now available that harvest existing molecular data from publically available databanks to produce trees [6,7,12], as well as compile existing phylogenies into larger metatrees [13,14]. Large numbers of insect comparative studies now incorporate the development of bespoke phylogenies as an explicit step, and it is common for studies whose main output is a phylogeny to piggyback a comparative study as a selling point [15-17].

In addition to more trees on which to base studies, there are also more traits to analyze. Whole genomes and transcriptomes now allow us to investigate the evolution of the genes that control phenotypic traits of interest [18,19]. There has been a flowering of studies of macroevolutionary (speciation and extinction) rates, which can be inferred from the branching pattern on phylogenies [20-23], or, in the case of extinction, from conservation designations [24]. From the perspective of behavioural traits, this means that we are better able to explore not only the causes of variation in behaviour across taxa (e.g. such as the selective pressures and constraints controlling them), but also their consequences (both ecological and evolutionary). Entomologists also study esoteric but fascinating questions, such as the function of halters in locomotion [25] and the choreography of silk spinning [26], which result from the unique variation in phenotypes produced by one of the world's most impressive adaptive radiations.

Here I collate recent comparative studies addressing the causes and consequences of variation in insect behavioural traits to illustrate the range of potential applications of comparative methods to such studies, and what they can tell us. I choose studies to illustrate a wide range of focal behaviours, although many studies illustrate well how these different categories of behaviour overlap and interact or influence each other [27-33].

Recent comparative studies of behaviour

Foraging behaviour

Finding food, and a habitat that provides it, is necessary for all animals, and several recent studies have addressed how insects do this [3,4,34]. Patterns of host use in phytophagous insects are basis of terrestrial food webs, and may be shaped by experience, such that species are more likely to accept hosts they have previously encountered. Such conditioning may be adaptive if it facilitates decision-making in a complex community of potential hosts. Across 196 studies that had tested for this conditioning, such responses are indeed the norm, are just as common in monophagous as polyphagous species, and are just as likely to be produced by larval and adult experiences, but pupal experiences less so [3]. Closely related species also show similar responses. Thus, previous conditioning likely exerts a powerful effect on realized patterns of host use in nature.

Other studies have addressed the consequences of foraging choices and habitat selection [20,22,23,31,34,35]. For example, the phylogeny of skipper butterflies suggests that they fed ancestrally on dicot (broadleaved) plants, but some groups transitioned to monocot plants (grasses and allies) on which net diversification has been faster [22]. This is mainly attributable to two increases in net diversification rate within the monocot feeding clades which may have been triggered by climatic events which favoured the expansion of grasses first in forested, and then in more open habitats. This scenario suggests that behavioural (host choice) and abiotic forces (climate) have interacted to produce macroevolutionary effects mediated through the hosts, and intuitively this seems likely to be common in phytophagous insects.

Interspecific interactions

Recent comparative studies of predator-prey interactions have uncovered interesting associated trait variation [27,32,36]. In tiger moths and their relatives for example, hidden contrast colours (e.g. brightly coloured hindwings used to startle predators if crypsis fails) are more common in larger species [27]. A theoretical model shows that contrast colours can evolve in larger species if larger species are easier for predators to detect when cryptic, and if larger signals can more effectively startle predators [27]. Experiments with robotic moth models show that this is indeed the case.

The origin of some specialized trophic interactions, common in insects, is the focus of enduring interest [37,38]. A large data-base of global host records of

phytophagous insects indicates that Lepidoptera which attack particular host orders are less likely use others (the main trade-off in host-use is between woody and non-woody plants, with insects being largely restricted to one of these groups but not both). These negative associations between host-use were mainly seen when comparing higher insect taxa, not closely related species, so the failure to detect host-use trade-offs in laboratory selection experiments does not necessarily mean they do not emerge over longer timescales. In Hemiptera, trade-offs in host use were not generally detected, so cannot explain host specialization in that group [39]. The specificity of interspecific interactions such as these can also have wider ecological and evolutionary consequences [20,40]: a phylogeny of ambrosia beetles for example suggests that genera with broader host ranges tend to have diversified faster [20]. This might be because host switching facilitates reproductive isolation between incipient species.

Dispersal and locomotion

Dispersal and locomotion ability are traits of ecological importance addressed by several recent studies [23,24,41]. In the semi-aquatic bug group Gerromorpha, there is a variety of locomotion styles, from tripod-walking to rowing across the water surface (water striders). Phylogenetic reconstructions show that the ancestral habitat of the group was probably terrestrial or waterside vegetation, and a transition to living on the water surface was associated with an increase in locomotion speed across species, necessitated perhaps by increased predation risk and the need to move faster than the water when water is flowing, in order to maintain position. The increase in speed is correlated with the lengthening of legs and increasing body size, and adoption of a rowing action which decreases stroke rate, increasing efficiency [41]. The macroevolutionary consequences of locomotory and dispersal behaviour are also known to be far-reaching. In European butterflies, it is one of the life history traits that predicts a lower extinction risk, alongside high voltinism, and overwintering in later life history stages, presumably because it facilitates metapopulation persistence [24].

Communication and signalling

Explaining the diversity of animal signals is another enduring challenge to which recent comparative studies have contributed [29,33]. In ladybird beetles, for example,

the wing cases (elytra) show a variety of colour patterns (often red or yellow against black), which are thought to warn predators of their toxicity. Comparing toxicity (determined experimentally) and colour patterns across several species, it can be shown that brighter colours are correlated with increased toxicity (Figure 2) suggesting that aposematic signalling is honest, implying evolutionary mechanisms which maintain this honesty. A further experiment with model ladybirds showed that brighter colours are also more effective at deterring predators [33]. Fireflies are another group of beetles that show colour variety in their signals, this time produced by bioluminescence. Again, the signal is correlated with other species traits. Male fireflies that are active in early evening in vegetated habitats (when the background vegetation still reflects green) are predicted to produce light that contrasts better with the green background. This indeed is the case, as they tend to produce yellow light instead of green. Sedentary females and later active males however can use green light to their advantage in the dark to maximize reflectance from vegetation so their signals are more obvious, and their light is indeed greener [29].

Mate choice and sexual selection

Mate choice and sexual selection have produced impressive phenotypic variation addressed by comparative studies [18,21,30,42]. Male orchid bees attract females by perfumes which they construct by collecting volatile substances from orchid flowers, and which they store in specialized leg pouches. Traits like these that evolve by persistent strong sexual selection are predicted to evolve more rapidly than other traits. Mapping perfume traits across a phylogeny of 65 species, perfume chemistry was shown to display faster rates of evolution and a higher disparity compared to non-signalling traits, matching theoretical expectations. The complexity of the perfume increased with the number of sympatric congeners, suggesting that other species are one selective agent increasing signal diversity [30]. Such sexually selected signals have also long been suspected to increase diversification rates. Recent evidence from insect and other animal species showing bioluminescent displays is consistent with this: those using lights in their courtship displays are more species rich than their relatives without these displays, but this is not true for non-courtship displays, suggesting that sexual selection indeed promotes species richness [21].

Parental care

Compared to birds and mammals, insects are not widely known for their parental care, but it is found in hundreds of species in many different taxonomic groups. The selective pressures that lead to care in one or either sex may depend on the ancestral states from which different types of care evolved. Across a dataset of over 2000 insect species, the ancestral care state was found to be no-care, and female-only-care was the most common type of care, evolving directly from no care, and sometimes transitioning to biparental care. Male-only-care also evolved from no-care in Hemimetabola, although quite rarely, whilst in a few cases biparental care evolved directly from no-care [5] (Figure 3). These findings are very different to those in vertebrates, where biparental care and male-only-care are more common, and femaleonly-care often evolves from biparental care. The lack of male care in insects may be the result of widespread sperm competition, which encourages males to desert and remate. The selective pressures favouring care have also been addressed in some studies: in acanthosomatine bugs maternal care is found in species which lack a protective covering for their eggs, suggesting that predation on offspring drives the evolution of care, and that there is a trade-off between different mechanisms of providing offspring protection [32].

Social evolution

Insect comparative studies have provided important evidence about the causes [43] and consequences [19,28,44] of sociality. *Polistes* paper wasps sometimes nest solitarily and sometimes cooperatively. Sheehan et al. [43] georeferenced data on nest foundress number worldwide, and then correlated it against climate data for those locations. They showed that cooperative nesting was more common in locations with short term instability in temperature, whilst the number of foundresses was lower in harsh environments. This suggests that cooperation is driven by selective responses to environmental conditions, but that the forces that regulate cooperation and foundress number may be different.

Kapheim et al. [19] compared the genomes of ten bee species with a variety of social structures to investigate the genetic mechanisms and consequences behind social evolution. They found that increasing social complexity (i.e. from solitary at one extreme to obligate complex eusociality at the other) was associated with

211	increased capacity for gene regulation (more transcription factor binding sites in
212	promotor regions, as well as the number of genes predicted to be methylated, which
213	affects gene expression, and more rapid evolution of regulatory genes). Hence social
214	evolution appears to have produced more complex gene networks.
215	
216	Other behaviours
217	Entomologists often study more unique behaviours that do not fit easily into the
218	standard pantheon of animal behavioural repertoires. Silk spinning, unknown in
219	vertebrates, is widely used by spiders and insects to form structures (extended
220	phenotypes) with obvious adaptive purposes, and likely macroevolutionary
221	consequences [45]. One of the lesser known insect groups that does this is the
222	Embioptera (webspinners), relatives of stick insects [8] which live in silk-lined
223	burrows mainly in the tropics. The group is morphologically very uniform, but varies
224	in silk-spinning behaviours, produced from modified forelimbs. By coding the
225	movements of the legs and transitions between spin-steps and correlating them with
226	other traits across a phylogeny, body size was shown to explain much of the diversity
227	in spinning choreography across species [26], and there were also some differences
228	between species inhabiting different microhabitats (such as tree trunks versus soil).
229	Web spinning behaviours therefore seem to have evolved in concert with both
230	transitions in microhabitat and morphology.
231	
232	Conclusion
233	The insects contain the greatest adaptive radiations that can be seen with the naked
234	eye [6,8,35]. There is arguably no greater resource to learn about the evolution of
235	phenotypic diversity. To exploit it to the full we need to have access to phylogenies,
236	trait data and analytical methods. Phylogenies and analytical techniques are much
237	more accessible and useful to comparative biologists now than only a few years ago,
238	but insect trait data still lie scattered across a vast heterogeneous landscape of natural
239	history books, encyclopedias, museum collections, scientific journals and other
240	sources. To exploit the promise of insect comparative studies to the full, we need
241	global digital data depositories that will collate, store and curate this information.
242	Such trait-data hubs will finally bring comparative entomology fully into the

information age.

Table 1. Analytical methods applied by recent comparative studies of insect behaviour.

behaviour.			
Method	Type of output	Computer applications	Recent studies
Ancestral state	A set of most	'Mesquite'	[35,43]
reconstruction with	parsimonious		
parsimony	ancestral states		
Bayesian ancestral	A set of most likely	'RASP' [46]	[42]
state reconstruction	ancestral states and		
	models of change		
Maximum likelihood	A set of most likely	'ape' [47] and	[35,40,41]
ancestral state	ancestral states for	'corHMM' [48] in	[, -,]
reconstruction	a given model of	R, 'Mesquite'	
	change	ii, iiiosquite	
Reconstruction of	A set of most likely	'BioGeoBEARS'	[43]
ancestral	ancestral ranges	in R [49],	[15]
biogeographic ranges	and models of		
ologeograpine ranges	change	'Lagrange' [50]	
Blomberg's K	Metric of	'geiger' in <i>R</i> [51]	[26,33,52]
estimate of		geiger in K [31]	[20,33,32]
	phylogenetic		
phylogenetic	similarity across		
similarity	species for a trait	37/4	F01 053
Sister-clade	Evolutionary	N/A	[21,35]
comparisons	associations		
	between traits		
Phylogenetically	Evolutionary	'caper' and 'ape' in	[19,26,41]
independent contrasts	associations	<i>R</i> [47,53],	
(PICs)	between traits	'Mesquite'	
Lynch's phylogenetic	Evolutionary	'ape' in <i>R</i> [47]	[31]
mixed model [54]	associations		
	between traits		
Phylogenetic	Evolutionary	'adephylo' in R	[4]
autoregression	associations	[55]	
	between traits		
Pagel's discrete	Evolutionary	'Mesquite'	[32]
character association	associations	-	
test [56]	between traits		
Concentrated changes	Evolutionary	'MacClade'	[32]
test for binary	associations		. ,
characters	between traits		
Bayesian modelling	Evolutionary	'BayesTraits' [57]	[5]
of trait evolution	associations) [2.]	c J
	between traits and		
	transition rates		
	between states		
Generalized	Evolutionary	'ape' in <i>R</i> [47]	[23]
estimating equations	associations	ape mr[+/]	[23]
[58]	between traits		
Phylogenetic	Evolutionary	'phytools' in <i>R</i> [60]	[36]
ANOVA [59]	associations	phytoois in A [00]	[20]
ANOVA [37]	associations		

	1 4 4		
Phylogenetic	between traits Evolutionary	'caper' in R	[24,27,43,44,61]
Generalized Least	associations	caper in K	[24,27,43,44,01]
Squares (PGLS) and	between traits		
Pagel's λ metric of			
phylogenetic			
constraint			
Phylogenetic	Evolutionary	Bespoke <i>R</i> script	[34]
Generalized Least	associations	[34] based on [62]	r- 1
Squares extended to	between traits,		
incorporate	metric of		
measurement error	phylogenetic		
	constraint		
Phylogenetic	Evolutionary	'MCMCglmm' in	[3,28,39]
multivariate mixed	associations	<i>R</i> [43,63]	
models	between traits		
Phylogenetic logistic	Evolutionary	'Phylolm' in R	[24]
regression [64]	associations		
	between traits		
Stochastic linear	Evolutionary	'SLOUCH' in R	[20]
Ornstein-Uhlenbeck	associations	[65]	
modelling	between traits		
Ornstein-Uhlenbeck	Model that best	'OUwie' in <i>R</i> [66]	[40]
modelling of	describes how		
predator-regime	predators affect		
specific dynamics	evolution of a		
	continuous trait		
Phylogenetic	Reduction of	'phytools' in R [60]	[26,36]
principle component	multivariate cross-		
analysis	taxonomic data to		
	principle		
36.1	components	0.07	F2.03
Multivariate trait	Tempo and mode	'MVmorph' in R	[30]
evolution modelling	of evolution of	[67]	
D : 1 : C	multivariate traits	(1,). D	[22]
Bayesian analysis of	Effect of a binary	'diversitree' in <i>R</i>	[22]
speciation and	trait on speciation	[69]	
extinction (BiSSE)	and extinction rates		
[68]	True of assents that	(Iona) [70]	[20]
Event-based analysis	Type of events that	'Jane' [70]	[38]
of co-phylogenetic	best explain the co-		
structure	phylogenetic structure		
Distance-based	Assessment of the	'Parafit' [71],	[38]
analysis of co-	congruence of two	'PACo' [72]	[38]
phylogenetic structure	phylogenies	11100 [12]	
Network analysis with	Rates of co-	'abctools' in R [73]	[37]
Approximate	speciation and host	4000015 III K [75]	[-,]
Bayesian	shifting across		
Computation	phylogenies of		
	F-17 10 Demies 01		

interacting species

Figure legends 247 248 249 Figure 1. A schematic diagram of the stages of a comparative analysis. Trait data are 250 gathered across taxa (X and Y represent different traits, and the numeric subscripts 251 indicate that each species is assigned a value for each trait from observation), and 252 phylogenetic information assembled. This information is then integrated through one 253 or more of a battery of analytical methods (Table 1) to produce a variety of outcomes 254 (Table 1), the most common of which are ancestral state reconstruction (i.e. 255 estimating the values of X and Y for ancestors of the living species for which we have 256 data) and detecting evolutionary correlations between traits (i.e. whether evolutionary 257 change in Y tends to be associated with evolutionary change in X). 258 259 Figure 2. Association between the colour intensity (i.e. saturation; how much colour 260 there is compared to white light) of ladybird individuals belonging to different 261 species, and the toxicity of those individuals as measured by their killing effect on 262 Daphnia (linear mixed-effects model: $F_{1.54} = 5.57$, lower p < 0.05). Saturation is measured as the Euclidean distance between each colour and the achromatic centre of 263 264 a cone-sensitivity weighted tetrahedral colour space. Briefly, the colour of each 265 ladybird is plotted in four dimensions (tetrahedral space) where the four axes 266 represent how much the four different cone cells of a bird are stimulated by the colour 267 (standardized so 1 = full stimulation, 0 = no stimulation). Saturation measures the 268 distance between the colour and the centre of the space representing white light, and 269 the units are therefore standardized probabilities of absorption (for full details of the 270 method see Figure 3 of [74]). The black line is a linear regression through the points 271 (for indicative purposes only), and the grey area encompasses the regression standard 272 errors. Reproduced from Figure 3 of [33] under the Creative Commons Attribution 273 License (CC BY 4.0). 274 275 Figure 3. Transition rates (events per unit branch length on the phylogeny) between 276 parental care states, estimated from Bayesian analysis of phylogenies of 277 hemimetabolous insects. NC = no care MC = male-only-care, FC = female-only-care, 278 and BP = biparental care. Histograms show the frequency distributions of rate

estimates over the modelled posterior distribution. f(Z) = frequency at which the

transition rate was zero. Arrow thickness is proportional to transition rates, and dotted lines are where the median rate $(\bar{x}) \pm \text{standard deviations (given in the histograms)}$ overlap zero. The highest transition rates are between no care and female-only-care (and vice-versa), and also male-only-care to no care. Reproduced from Figure 4a of [5] under the Creative Commons Attribution License (CC BY).

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