Phylogenetic analysis of non-stereotyped behavioural sequences with a successive event-pairing method

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A new method is proposed which uses transitions among acts in non-stereotyped behavioural sequences as phylogenetic characters. This method is derived from the event-pairing method designed for the phylogenetic study of developmental sequences and from ethological analyses of transition matrices. It is applied to study the phylogenetic relationships within a well-known group, the presocial Zetoborinae cockroaches. The analysis is carried out with three data sets: a behavioural data set with transitions among acts in behavioural dyadic sequences, together with morphological and molecular data sets. Non-stereotyped behaviour proved to be phylogenetically informative and to display low homoplasy. This new method opens an avenue for studying the evolution of behaviour in the framework of phylogenetic analysis, which was restricted until now to the study of stereotyped sequences and/or isolated features involved in courting or building activities. © 2008 The Linnean Society of London, Biological Journal of the Linnean Society, 2008, 94, 853–867.


INTRODUCTION

Studies on behaviour and phylogeny have a long common history. From the early times, comparative studies have shown that behaviour can be remarkably informative regarding the relationships of taxa, as reviewed by Hinde & Tinbergen (1958). These comparative studies focused on highly stereotyped and ritualized behaviours, such as courting or nest building (Wenzel, 1992). Being stereotyped, these behaviours were easily compared among species to assess their patterns of evolution (Lorenz, 1941; Hinde, 1955; Tinbergen, 1959). In this context, the homology of stereotyped behaviour has been repeatedly discussed by several seminal papers (Baerends, 1958; Atz, 1970; Hodos, 1976; Lauder, 1986; Wenzel, 1992; Greene, 1994) with reference to the classical criteria of homology proposed by Remane (1952) and especially the criteria of position and special quality. The rationale behind using behaviour for phylogenetic inference was that behavioural acts that are repeatedly observed in various individuals and populations without strong variation or evidence for learning can be presumptively considered as inherited, in the same way as other phenotypic traits, such as morphology or development (Wenzel, 1992). More recently, many studies confirmed the old perception (de Queiroz & Wimberger, 1993; Proctor, 1996; Brooks & McLennan, 2002) that behaviour is phylogenetically informative and that it can be studied within such an evolutionary perspective (e.g. Coddington, 1986; McLennan, Brooks & McPhail, 1988; Wenzel, 1993; Kennedy, Spencer & Gray, 1996; Johnson et al., 2000; Cap, Aulagnier & Deleporte, 2002; Noll, 2002; Price & Lanyon, 2002; Desutter-Grandcolas & Robillard,
In phylogenetic analyses, coded ‘present’ vs. ‘absent’ lar behaviours are used as characters or attributes among the different studies. In the best case, particular treatments of behavioural characters greatly vary (as reported by Proctor, 1996; Desutter-Grandcolas & Robillard, 2003; Grandcolas & D’Haese, 2004). However, when behavioural sequences are recoded in all possible pairwise combinations of events, thereby encoding the relative position of each item in the sequence. Each developmental event is coded as possibly occurring before, simultaneously or after any other event (Jeffery et al., 2005). Event-pairing coding has been recently challenged by Schulmeister & Wheeler (2004) who suggested that treating features of sequences as if they were independent can produce inconsistent reconstructions. This is mainly because developmental sequences are more constrained in terms of temporal linearity than are ethological sequences with respect to the biological process involved. For example, many developmental events cannot occur earlier or later within a sequence because the structures where they should take place are not yet developed or cannot develop twice (Schulmeister & Wheeler, 2004).

The limitations of event-pairing coding as applied to the study of development are not a problem for non-stereotyped behavioural sequences, where the same event can be expressed several times within the sequence. There is not a one-to-one relationship between a linear sequence and a species, but many different sequences for the same species. Our goal is to code in a phylogenetic context the occurrence and frequency of transitions between two acts among many differently ordered sequences. Only transitions between two successive events are considered here and not the relative position between all pairs of events. This methodology is named the successive event-pairing method to avoid confusion with the event-pairing method.

Establishing a matrix of characters coding the occurrence of two successive events in a behavioural sequence is already part of the current statistical analyses of behavioural sequences (Fig. 1). Ethologists build matrices of transition where each cell is filled with the frequency of a transition between two particular acts (Martin & Bateson, 1986; Gottman & Roy, 1990; Bakeman & Quera, 1995). These frequencies are then organized in flow charts generated by hand or according to correspondence analyses (van der Heijden, 1987) to investigate how different acts are organized in different kinds of sequences and to compare them between different species. To adapt this procedure to the successive event-pairing
method, we only need to consider that the cells of matrices of behavioural transitions can be used as phylogenetic characters (Fig. 1). This is justified on the basis of the classical criteria of homology applied to comparative ethology (Wenzel, 1992). Homology based on the cells of such matrices fits the criterion of position as it defines a particular succession of two acts, thus specifying the position of one act relative to another. In this context, events A occurring after an event B or after an event C are not considered homologous, strictly speaking: answering by A after B or after C is not the same behaviour and will not necessarily be shown by all species even if the fixed motor pattern involved in displaying A is the same in each case. This is easy to understand if one considers a real example where a species would tend to answer to conspecific aggression by escape, while another species would answer by reciprocal aggression. Ethologists have long known that this kind of difference can be species-specific or common to related species. The occurrence of transitions between two particular acts can be treated as presence–absence characters. The frequencies of transitions can also be used as it is very different to observe that a given transition is very rare or very common. Either a low or a high frequency can be considered characteristic of species and therefore used in phylogenetic analysis as characters. Frequencies can be discretized and coded in different character states using gap coding (Archie, 1985; Stevens, 1991). Recently, Goloboff, Mattoni & Quinteros (2006) argued that continuous characters need not to be discretized. However, their methodology treats continuous characters as additive characters, which requires an assumption of progressive evolution that we do not want to follow here. The study of quantitative characters has proved to be a
difficult problem in phylogenetics and the present work is not aimed at solving it. We will focus this work on the most commonly used approaches: discretization and gap coding.

Our method requires that all these behavioural patterns, both the acts and the trends of succession among acts, are largely heritable and that their plasticity and variability are low. This is the most general and necessary assumption that should be substantiated to legitimate phylogenetic studies of behaviour. It can be partly carried out in evaluating the congruence of the phylogenetic tree based on behavioural data with molecular and morphological data. Obviously, assessing the minimal sampling effort is needed to document correctly the behavioural acts, their transitions and frequencies and their independence. These assumptions are no different than for other phenotypic characters (morphology, cytology, etc.), as already argued by Wenzel (1992).

MATERIAL AND METHODS

A N ILLUSTRATIVE CASE STUDY: GREGARIOUS BEHAVIOUR IN ZETOBORINAE COCKROACHES

Non-stereotyped behavioural sequences are most often observed in the context of social relationships. The observed behavioural sequences are not a series of acts successively emitted by the same individual, a situation which could occur with other behaviours such as territorial displays, grooming activities, etc., but a series of acts emitted by two individuals in alternation. These behavioural relationships are rarely stereotyped and there can be different answers to a particular act from a conspecific, depending on the context, and several different acts can initiate a sequence of interaction between two conspecifics.

Gregarious behaviour in cockroaches is a famous example of presocial behaviour (Schal, Gautier & Bell, 1984; Gautier, Deleporte & Rivault, 1988; Nalepa & Bell, 1997; Grandcolas, 1999; van Baaren et al., 2002, 2003b). It has been recently analysed in a molecular and morphological comparative framework in the subfamilies Zetoborinae and Blaberinae (Grandcolas, 1991, 1993a, b, 1998; Pellens, Legendre & Grandcolas, 2007a; Pellens et al., 2007b), which provided both a phylogenetic reference and a natural history context for the interpretation of social behaviour observed in the laboratory (Grandcolas, 1991; van Baaren & Deleporte, 2001; van Baaren et al., 2002, 2003a). In a first attempt to understand the evolution of social behaviour, relevant categories such as ‘gregarious’, ‘solitary’ and ‘subsocial’, have been mapped onto a phylogenetic tree based on morphology (Grandcolas, 1993a, 1998). Ethological studies have shown that the gregarious behaviours should be analysed in detail and contrasted between species, not only considering broad behavioural categories (Grandcolas, 1991; van Baaren & Deleporte, 2001; van Baaren et al., 2002, 2003a). A small group of closely related species has been already well studied and will be used as an example of how the successive event-pairing method can be applied to behavioural sequences to infer a phylogenetic tree congruent with other data and to propose hypotheses of behavioural evolution. We will not discuss the issues of behavioural plasticity and repertoire sampling which have been explored and controlled for in the specific papers cited earlier.

Dyadic interactions in four species – namely Thanatophyllum akeinetum Grandcolas, 1991, Schultesia lampyridiformis Roth, 1973, Lanxobatta emarginata Burmeister, 1838 and Phortioeca nimbata Burmeister, 1838 – have been observed among 11 to 17 groups of six nymphs placed in standard conditions for each species, according to the protocol described in Grandcolas (1991) and van Baaren et al. (2002). Each group has been placed in an open-field arena. The observations began 1 h later, lasted 15 min and have been recorded on a Samsung Digital Camcorder VP-D11. The observations have been carried out on nymphs in the middle of their development as this is the most characteristic and intense period of gregarious behaviour (Grandcolas, 1993b; van Baaren & Deleporte, 2001). One additional outgroup species Eublaberus distanti Kirby, 1903 from the closely related subfamily Blaberinae has also been observed. Transition matrices have been constructed using the behavioural sequences reported as described in Figure 1. Phylogenetic analyses have been carried out with these behavioural matrices and also by comparison with a morphological and molecular data set. The morphological data are taken from Grandcolas (1993a, 1998). Molecular data for all the species and behavioural data for Eublaberus distanti have been acquired for the present study.

The sampling effort for behaviour has been critically evaluated with respect to the previous behavioural studies that were conducted and published on the same insect species (e.g. van Baaren et al., 2002). Accumulation curves for the occurrence of transitions according to the number of observations have been computed to show that the sampling effort is large enough to observe the transitions, either uncommon or frequent, occurring in each species.

Character independence has also been evaluated by checking whether frequencies of transitions involving a similar behavioural act are not misleadingly correlated. This can be easily tested with a $\chi^2$ goodness-of-fit test (Chatfield & Lemon, 1970; Zar, 1999), which verifies whether the frequency of a transition between two acts can be determined by the total frequency
of each act involved. Basically, this test compares expected frequencies with observed frequencies. Following Zar (1999), some data have been pooled together in order to have an average expected frequency of at least six, which avoids bias in $\chi^2$ computation.

**PRIMERS, PCR AND SEQUENCING**

Leg muscle tissue was excised from roach specimens preserved in 100% ethanol. DNA was extracted using the Qiagen DNeasy protocol for animal tissue. Mitochondrial ribosomal DNA large subunit (16S, ~385 bp), nuclear ribosomal DNA small subunit (18S, ~1875 bp) and nuclear ribosomal DNA large subunit (28S) domains A (~360 bp) and C (~330 bp) were amplified. 18S was amplified and sequenced in four overlapping fragments corresponding to GA, AD1D2, BCE and EF domains. PCR reactions were lead on a DNA Engine DYAD™, Peltier Thermal Cycler with the following conditions: an initial heating step of 94 °C for 2 min followed by 40 cycles of 94 °C for 60 s, 55 °C for 60 s and 72 °C for 75 s. Then a final elongation at 72 °C was carried out over 7 min. The different primers used which have already been published are listed in Table 1. Electrophoresis gel was used to visualize PCR products and to check that there was no contamination as a result of a negative control. PCR products were purified via the Montage PCR96 Cleanup Kit (Millipore) and sequenced using ABI Big Dye 3.1 with the following sequence profile: 27 cycles of 96 °C for 10 s, 50 °C for 5 s and 60 °C for 4 min. Sequencing reactions products were purified with Sephadex™ columns and fractionated on an ABI 3730 XL DNA sequencer. Each sequence was edited using Sequencher 4.0 (Genecodes, 1999) and blasted on GenBank (http://www.ncbi.nlm.nih.gov/blast/) to check for contamination. All the sequences (16S/18S/28SA/28SC, respectively) were deposited on GenBank under the following accession numbers: *Eublaberus distanti* (EU367504/EU367508/EU367511/EU367516), *Lanxoblatta emarginata* (EU367505/EU367509/EU367512/EU367517), *Phortioeca nimbata* (EU367506/EU367510/EU367513/EU367520), *Schultesia lampyridiformis* (EF363280/EF363251/EU367515/EU367520) and *Thanatophyllum akinetum* (EU367507/EU367503/EU367514/EU367519).

**PHYLOGENETIC ANALYSES**

Even if repertoires are basically identical among species and therefore cannot convey a great deal of phylogenetic information, their analysis has been carried out for a matter of comparison (analysis A). Using the successive event-pairing method, a second phylogenetic analysis has been performed on the presence–absence of transitions taken as characters (analysis B). In a third analysis (analysis C), characters based on the discretized frequencies of the transitions were added to the B data set. The marginal frequencies of each initiating act have been calculated. For instance, in species 1, one event A and two events C have been observed in answer to an event B (Fig. 1). Then, the frequencies for the transitions B/A, B/B, B/C and B/D are 0.33, 0.00, 0.67 and 0.00, respectively. All the frequencies for the same initiating act have been pooled throughout all the species to form a distribution. Each distribution of frequencies was discretized and coded using gap coding (Archie, 1985; Stevens, 1991).

The morphological data set has been taken from the matrix of 78 characters used by Grandcolas (1993a), considering only the five species of the present study.

<table>
<thead>
<tr>
<th>Genes</th>
<th>Primers</th>
<th>Sequences (5′-3′)</th>
<th>Sources</th>
</tr>
</thead>
<tbody>
<tr>
<td>16S</td>
<td>16SAr</td>
<td>CGCCTGTTTATCAAAAACAT</td>
<td>Xiong &amp; Kocher (1991)</td>
</tr>
<tr>
<td></td>
<td>16SF</td>
<td>TTA CGCTGTATCCCTAA</td>
<td>Kambhampati (1995)</td>
</tr>
<tr>
<td>18S GA</td>
<td>1F</td>
<td>TACCTGGTTGATCCCTGCCAGTAG</td>
<td>Giribet et al. (1996)</td>
</tr>
<tr>
<td></td>
<td>b5.0</td>
<td>TAACCGCAACAACTTTAAT</td>
<td>Whiting et al. (1997)</td>
</tr>
<tr>
<td>18S AD1D2</td>
<td>2F</td>
<td>AGGGTTGATTTCGGAGGGGACG</td>
<td>Hillis &amp; Dixon (1991)</td>
</tr>
<tr>
<td></td>
<td>b2.9</td>
<td>TATCTGATCGCCTTGGAGACCTCT</td>
<td>Jarvis, Haas &amp; Whiting (2004)</td>
</tr>
<tr>
<td>18S BCE</td>
<td>a1.0</td>
<td>GGTTGAAATTTTGGACCCTGC</td>
<td>Whiting et al. (1997)</td>
</tr>
<tr>
<td></td>
<td>7R</td>
<td>GAC ATACAGACCTGTATTGC</td>
<td>Whiting (2002)</td>
</tr>
<tr>
<td>18S EF</td>
<td>a3.5</td>
<td>TTG TGCA TGGCCGGTCTTGTAG</td>
<td>Whiting (2002)</td>
</tr>
<tr>
<td></td>
<td>9R</td>
<td>GAACTGCCGAGTTCTGGACCTAC</td>
<td>Giribet et al. (1996)</td>
</tr>
<tr>
<td>28S A</td>
<td>Rd1.2a</td>
<td>CCSSGGTATTTAAGCATAATTA</td>
<td>Whiting (2002)</td>
</tr>
<tr>
<td></td>
<td>Rd3b</td>
<td>CCY TAAACGCTTTACCTAC</td>
<td>Jarvis et al. (2004)</td>
</tr>
<tr>
<td>28S C</td>
<td>28SA</td>
<td>GACCGGTCTTGGAGCAGCAG</td>
<td>Whiting et al. (1997)</td>
</tr>
<tr>
<td></td>
<td>28SB</td>
<td>TCGGA GAAACCCAGCTAC</td>
<td>Whiting et al. (1997)</td>
</tr>
</tbody>
</table>
As it represents few parsimony-informative characters (9), it has been analysed together with the molecular data (analysis D). The molecular data set represents portions of genes 16S, 18S and 28S. Molecular sequences, which present a very low variability in length, were aligned using Muscle 3.6 (Edgar, 2004). Finally, behavioural data (including frequencies) have been combined together with the morphological + molecular data set (analysis E).

All characters were equally weighted and coded as non-additive. All parsimony analyses have been performed using PAUP4.0b10 (Swofford, 1998) with an exhaustive search to ensure finding the most parsimonious tree. Consistency index (CI; Kluge & Farris, 1969), retention index (RI; Farris, 1989) and number of parsimony-informative characters were recorded. Bremer support values were computed with the help of TreeRot.v2b (Sorenson, 1999). Bootstrap values were computed for 1000 replicates using PAUP4.0b10. Character optimizations on the phylogenetic trees were performed with fast procedure (i.e. accelerated transformation ACCTRAN) using Winclada version 1.00.08 (Nixon, 2002).

Because the monophyly of the ingroup has been established previously (Grandcolas, 1993a, 1998), the ingroup was designated as monophyletic in tree visualization.

Analyses C and E include characters based on the frequencies of transitions and consequently include inapplicable characters. If a particular behavioural transition is not observed in one or several species, the character based on its frequencies is inapplicable for that or those species. Those characters were coded with a dash (‘-’) in the matrices but were interpreted as missing data during the tree search. This ‘reductive coding’ better reflects the information content of the data (Strong & Lipscomb, 1999).

The behavioural data were tested for significant structure using the permutation tail probability test (PTP; Faith & Cranston, 1991) and the g1 statistics (Hillis, 1991; Hillis & Huelsenbeck, 1992) in PAUP4.0b10 (Swofford, 1998). However, even if there is a structure in a phylogenetic tree based on behavioural data, this structure can be a reflection of common evolutionary ecological pressures rather than of phylogenetic relationships (Kennedy et al., 1996). If the behavioural tree is congruent with a tree based on other data sets (morphological + molecular here), it is most likely to be as a result of a common phylogenetic signal in the different data sets. Behavioural and morphological + molecular trees were tested for congruence using the triplets tree comparison metric (Symmetric Difference of triplets, SDt hereafter) as implemented in the software Component v2.0 (Page, 1992); the lower the value, the more congruent the trees. This value can be compared with a null distribution calculated after generating all topologies with five leaves unrooted (options ‘generate all’ and ‘tree-to-tree distances/triplets/SD’).

RESULTS

Twenty-four different acts were identified from the behavioural sequences of the different species (Tables 2 and 3). Eublaberus distanti and Schultesia lampyridiformis have the largest behavioural repertoire (19 acts), whereas Lanzoblatta emarginata and Phortioeca nimba have the smallest and the same one (15 acts). The outgroup E. distanti displays some autapomorphic acts and notably the sudden jump (SJ) and the act PS (when an individual puts its pronotum under the other individual and stands up suddenly), both of which are known in Blaberus, the sister genus of Eublaberus (Gautier, 1974).

Transition matrices (Appendix S1) have been constructed using the behavioural sequences. Accumulation curves showed that the total number of observations is large enough to observe all frequent or uncommon transitions, the number of which has reached a plateau (e.g. for Eublaberus distanti, Fig 2). According to the matrices, S. lampyridiformis was the most active cockroach with more than 1300 behavioural transitions (Appendix S1) representing a mean of 65 interactions/h. Conversely, P. nimba and E. distanti were the less active (c. 30 interactions/h). We found no correlation between the amount of activity and the number of kinds of transitions among the five species, which shows again that there is no bias related to a sampling effect depending on different species activities ($r = 0.70$, $P > 0.15$). According to the behavioural matrices (Appendix S1), some transitions were never observed.
Table 2. List of the different behaviours displayed by the cockroaches (also established according to Gautier, 1974; Grandcolas, 1991; van Baaren et al., 2002, 2003a, b)

<table>
<thead>
<tr>
<th>Behaviours promoting interactions</th>
<th>MT</th>
<th>AC</th>
<th>MA</th>
<th>CB</th>
<th>PS</th>
<th>PP</th>
<th>KL</th>
<th>SP</th>
<th>SA</th>
<th>SD</th>
<th>SJ</th>
<th>BI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Moving towards the other individual</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Antennal contact with the body of the other individual</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Mutual antennation</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Climbing onto the body of the other individual with one to six legs</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
</tbody>
</table>

Table 3. Behavioural repertoires of the five cockroaches used in this study (also established according to Gautier, 1974; Grandcolas, 1991; van Baaren et al., 2002, 2003a, b)

<table>
<thead>
<tr>
<th>MT</th>
<th>AC</th>
<th>MA</th>
<th>CB</th>
<th>PS</th>
<th>PP</th>
<th>KL</th>
<th>SP</th>
<th>SA</th>
<th>SD</th>
<th>SJ</th>
<th>BI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Eublaberus distanti</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Lanxobatta emarginata</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
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<tr>
<td>Phortioeca nimbata</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
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<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
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<tr>
<td>Thanatophyllum akinetum</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
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<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
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<tr>
<td>Schultesia lampyridiformis</td>
<td>X</td>
<td>X</td>
<td>X</td>
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<td>X</td>
<td>X</td>
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<table>
<thead>
<tr>
<th>GD</th>
<th>GA</th>
<th>TP</th>
<th>FP</th>
<th>RO</th>
<th>WD</th>
<th>ES</th>
<th>WP</th>
<th>GA</th>
<th>GL</th>
<th>WA</th>
<th>NO</th>
</tr>
</thead>
<tbody>
<tr>
<td>Eublaberus distanti</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
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<td>Lanxobatta emarginata</td>
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<td>X</td>
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<tr>
<td>Phortioeca nimbata</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
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<td>Thanatophyllum akinetum</td>
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<td>X</td>
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<tr>
<td>Schultesia lampyridiformis</td>
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<td>X</td>
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</table>

(empty cells of the matrices), while others were rarely or commonly observed (cells filled with a small or a large integer, respectively). As a mean, the behavioural sequences include from 6 to 7 acts depending on the species, ranging from very short (two acts) to quite long (25 acts). Transition matrices show that many transitions are not observed in some species and that others are rare or frequent. Statistical independence among different transitions for a same species was assessed according to \( \chi^2 \) goodness-of-fit tests which were highly significant (\textit{Eublaberus}: \( \chi^2 = 380.3, \text{ddl} = 56, \chi^2_{0.05} = 74.5, P << 0.05; \textit{Lanxobatta}: \chi^2 = 297.3, \text{ddl} = 81, \chi^2_{0.05} = 103.0, P << 0.05; \textit{Phortioeca}: \chi^2 = 222.1, \text{ddl} = \)
The phylogenetic analysis based on behavioural repertoires of the different species (Table 3) resulted in four most parsimonious trees (six informative characters, \( L = 15 \text{ steps; } CI = 0.80, RI = 0.50 \)), the strict consensus of which is totally unresolved (Fig. 3A). Even if the repertoires are rich, their phylogenetic analysis is not decisive enough for establishing a resolved tree.

Phylogenetic matrices were built up from the transitions matrices as explained in Figure 1. The data set (B) based on the presence-absence of transitions taken as characters comprised 576 characters, 74 of which were parsimony informative (i.e. 12.8 %). Its analysis resulted in a single most parsimonious tree with a slightly different topology: the positions of

Figure 3. Analysis A, strict consensus of the four most parsimonious trees obtained (\( L = 15 \), CI = 0.80, RI = 0.50) with the data set based only on the behavioural repertoires. Analyses B–E, most parsimonious trees obtained with the four different data sets (B, presence–absence of behavioural transitions, \( L = 217 \), CI = 0.86, RI = 0.58; C, presence–absence and frequencies of behavioural transitions, \( L = 300 \), CI = 0.87, RI = 0.55; D, morphological and molecular data set, \( L = 351 \), CI = 0.88, RI = 0.50; E, molecular, morphological and behavioural data – including frequencies, \( L = 659 \), CI = 0.86, RI = 0.48). Numbers above and below branches are branch lengths (under fast optimization) and bootstrap/Bremer values, respectively.
S. lampyridiformis and T. akinetum are inverted with regard to the analyses B and C (L = 351 steps, CI = 0.88, RI = 0.50; Fig. 3D). The matrix comprised 3034 characters among which 88 were parsimony informative (2.9%). Finally, the combined analysis resulted in one most parsimonious tree similar to those based on behavioural characters (L = 659, CI = 0.86, RI = 0.48; Fig. 3E). Among the 4186 characters, 177 were parsimony informative (4.3%). This latter topology showed maximal bootstrap values and very high Bremer values (18 and 24). Statistics of all those analyses have been summarized in Table 4. They demonstrated that behavioural data were not more homoplastic than molecules and morphology and that behaviour brought proportionally more informative characters (up to 12.8 % of informative characters vs. 2.9 %). Table 5 included the partitioned Bremer support values and the number of informative characters for each partition. It revealed that behaviour (including frequencies) brought more than 80% of the signal [(32 + 2)*100/42 = 80.95]. Even when the support of each character was normalized by the number of informative characters, behavioural data appeared the most informative together with morphological data.

The g1 statistics and the PTP test indicated that behavioural data were highly structured (data set B: g1 = -1.406, P < 0.01; PTP test for 1000 replicates, P = 0.001) and the comparison between trees of the analyses B and D resulted in a SDt value of 0.2. Compared with the null distribution, this result suggested that those two trees (behavioural and morphological + molecular) were rather congruent. Only 12 pairwise comparisons between topologies out of 105 have a SDt value lower than 0.2.

The comparison of the trees and their statistics produced by the analyses B and C revealed to what extent presence–absence of behavioural transitions and their frequencies were informative. First, the B data set included 74 informative characters, whereas the C data set included 89 informative characters. Therefore, frequencies brought few supplementary informative characters (89 – 74 = 15) when compared with presence–absence of behavioural transitions.

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**Table 4.** Phylogenetic analyses with their data sets, results and statistics

<table>
<thead>
<tr>
<th>Analyses</th>
<th>data sets</th>
<th>Number of MPT</th>
<th>L (steps)</th>
<th>CI</th>
<th>RI</th>
<th>% of informative characters</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>Repertoire</td>
<td>4</td>
<td>15</td>
<td>0.80</td>
<td>0.50</td>
<td>6</td>
</tr>
<tr>
<td>B</td>
<td>Behaviour</td>
<td>1</td>
<td>217</td>
<td>0.86</td>
<td>0.58</td>
<td>74</td>
</tr>
<tr>
<td>C</td>
<td>Behaviour including frequencies</td>
<td>1</td>
<td>300</td>
<td>0.87</td>
<td>0.55</td>
<td>89</td>
</tr>
<tr>
<td>D</td>
<td>Molecular and morphology</td>
<td>1</td>
<td>351</td>
<td>0.88</td>
<td>0.50</td>
<td>88</td>
</tr>
<tr>
<td>E</td>
<td>Behaviour, molecular and morphology</td>
<td>1</td>
<td>659</td>
<td>0.86</td>
<td>0.48</td>
<td>177</td>
</tr>
</tbody>
</table>

CI, consistency index; L, length of the MPT; MPT, most parsimonious trees; RI, retention index.

**Table 5.** Partitioned Bremer values

<table>
<thead>
<tr>
<th>Partitions</th>
<th>Node (L, P)</th>
<th>Node [T, (L, P)]</th>
<th>Σ Bremer</th>
<th>Number of informative characters</th>
<th>%</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Behaviour</td>
<td>1</td>
<td>31</td>
<td>32</td>
<td>74</td>
<td>1.03</td>
<td>0.91</td>
</tr>
<tr>
<td>Behaviour (frequencies only)</td>
<td>1</td>
<td>1</td>
<td>2</td>
<td>15</td>
<td>0.32</td>
<td></td>
</tr>
<tr>
<td>Morphology</td>
<td>6</td>
<td>1</td>
<td>7</td>
<td>9</td>
<td>1.85</td>
<td>0.22</td>
</tr>
<tr>
<td>16S</td>
<td>1</td>
<td>–5</td>
<td>–4</td>
<td>46</td>
<td>–0.21</td>
<td></td>
</tr>
<tr>
<td>18S</td>
<td>4</td>
<td>–2</td>
<td>2</td>
<td>15</td>
<td>0.32</td>
<td></td>
</tr>
<tr>
<td>28S</td>
<td>5</td>
<td>–2</td>
<td>3</td>
<td>18</td>
<td>0.40</td>
<td></td>
</tr>
<tr>
<td>TOT</td>
<td>18</td>
<td>24</td>
<td>42</td>
<td>177</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

‘Σ Bremer’ is the total Bremer support for each partition and ‘%’ is the percentage of Bremer values that each partition supports normalized by the number of parsimony informative characters. Then, for each partition i, \( \%_i = \left( \frac{\Sigma \text{Bremer}_i}{\Sigma \text{Bremer}_\text{TOT} \times 100} \right) / N_i \); with \( N_i \) the number of informative characters for the partition i. For the last column, partitions are: behaviour including frequencies and morphology + molecular.

Second, Bremer values increased (increment of one for both clades) which means that the two data sets are not contradictory. However, frequencies gave less support to the clade \( T.\ akinetum, \ (L.\ emarginata - P.\ nimbata) \) than did the presence–absence of behavioural transitions. The comparison of branch lengths confirmed this point. Indeed, the increase of branch length between the two topologies was, in proportion, the smallest for this clade. On the contrary, frequencies brought relatively more information than presence-absence of behavioural transitions for the clade \( L.\ emarginata - P.\ nimbata \) with a Bremer support value of one for 15 informative characters vs. a Bremer support value of one for 74 informative characters, respectively (see Table 5).

Using the topology retrieved in the combined analysis (Fig. 3), we have looked at the behavioural transitions supporting different nodes of the tree. First, every branch was supported by a reasonable amount of changes. By comparison with repertoires which were basically identical for all species, this means that transitions offered a large amount of information with common states and differences among every species and group of species.

Second, acts supporting Zetoborinae with respect to the Blaberinae outgroup corresponded mainly to absences of agonistic or avoiding acts: slip ones pronotum under the other and stand up suddenly (PS), push the other with ones pronotum (PP), sudden jump (SJ) and sudden withdrawal of the antennae (WA), move away but stop in proximity (WP), rotation (RO) or freezing posture (FP), respectively. In the other way, Zetoborinae displayed other ‘negative’ acts in answer to ‘positive’ sollicitation (transitions which are not expressed by *Eublaberus*). Those ‘negative’ acts were: stilt posture with antennation (SA), goes down (GD), goes down and hides its antennae (GA) and withdrawal (WD).

Third, species with contrasted social behaviour (e.g. the solitary *T. akinetum* vs. other gregarious species) did not show a special amount of difference but particular changes. The behavioural repertoire of *T. akinetum* revealed only one autapomorphic behaviour: the absence of stilt posture (SP). Therefore, despite its solitary behaviour, *Thanatophyllum* did not display an especially idiosyncratic repertoire with regard to gregarious species. However, according to the successive event-pairing analysis, we were able to determine that *T. akinetum* was the only species displaying the following behavioural transitions, all of which limiting inter-individual interactions: move towards/bite (MT/BI), antennal contact/leg kick (AC/KL), stilt posture with antennation/move away but stop in proximity (SA/WP), bite/escape (BI/ES), rotation/leg kick (RO/KL), move away but stop in proximity/goes down and hides its antennae (WP/GA), nothing/goes down and hides its antennae (NO/GA).

We also looked at its gregarious sister group, composed of two closely related species which were supposed to have very similar social gregarious behaviour, *L. emarginata* and *P. nimbata*. These species were not only very close morphologically, they also lived in similar habitats (under loose bark of trees, Grandcolas, 1993b) and exhibit the same behavioural repertoire. Thirteen transitions supported this clade and notably five transitions involving acts of antennation (antennal contact and mutual antennation, AC and MA, respectively). Moreover, on the five characters based on frequencies and supporting this clade, four involved antennation as an answer to an act: move towards (MT) twice, stilt posture with antennation (SA) and nothing (NO).

**DISCUSSION**

**BEHAVIOURAL SEQUENCES IN PHYLOGENETIC ANALYSES**

Since Whitman (1898), Heinroth (1909), Lorenz (1941) and others pioneered the comparative study of behaviour, many phylogenetic analyses of behaviour have been carried out. The concept of behavioural homology has been discussed and finally found to be similar to the homology of other phenotypic characters (see Baerends, 1958; Atz, 1970; Hodos, 1976; Lauder, 1986; Wenzel, 1992). As morphological and molecular characters, behavioural ones proved to be phylogenetically informative when they were accurately defined and adequately sampled. As emphasized by Wenzel (1992), all these analyses took benefit and became more rigorous with the rise of phylogenetic methods, such as cladistics.

However, in spite of these very significant advances, the great majority of phylogenetic analyses of behaviour still considered behavioural acts in isolation (see Wenzel, 1992 for a review). Behavioural acts are regarded most often as equivalent if they are similar, even if they occur in different sequences or different places within the sequences. This is a much lower standard than for other kinds of characters (molecules, morphology) where the position of the feature is always carefully established (i.e. DNA sequence alignment, position criterion for morphological structures).

We recently emphasized that the most up-to-date phylogenetic algorithms designed to compare DNA sequences – such as direct optimization (Wheeler, 1996, Wheeler et al., 2006) – can be successfully applied to the stereotyped behavioural sequences (Robillard et al., 2006b). In the present paper, we make one more step and propose to compare non-
stereotyped and, therefore strictly speaking, not species-specific sequences. Thus, instead of focusing directly on the relationships among different behavioural sequences to infer relationships among species, we aimed at inferring relationships among characteristic parts of these sequences by applying a method of successive event-pairing which codes successions of acts.

The method is conceptually very straightforward by coding the occurrence of transitions between the acts in the sequences. It applies a better and more accurate concept of behavioural homology, each act being considered according not only to its special quality but also to its position within sequences.

Three main assumptions are implied and must be discussed with regard to behavioural characters: heritability, sampling and independence. All kind of characters used in phylogenetic analyses, either morphological, molecular or behavioural, should be checked from this point of view. In practice, this is usually made only for non-traditional characters such as behavioural or physiological ones, the variability of which is intuitively more questioned by scientists (Wenzel, 1992). Heritability is the first and most basic concern as non-heritable traits would be nonsensical if used in a phylogenetic context of descent with modification (see Grandcolas & D’Haese, 2003 for a review). With the exception of breeding and genetic studies, the only a priori way to assess the heritability of characters is to control for epigenetic effects by observing every species in the same conditions and by varying and repeating the conditions of observation. A posteriori, phylogenetic congruence between behaviour and other markers, including those molecular ones reputed to be neutral, is also a mean to assess heritability. In the present case, both criteria have been employed: repertoires and kinds of interactions have been found stable in repeated studies (Gautier, 1974; Grandcolas, 1991; van Baaren et al., 2002, 2003a) and the present study and the different data sets are reasonably congruent.

As for the sampling effort, behaviour is not more difficult to sample than morphology or molecules and it only requires to have living specimens placed in controlled and relevant conditions and to follow classical protocols (e.g. Martin & Bateson, 1986; Wenzel, 1992). In our case, accumulation curves were computed showing that the sampling of behavioural transitions has reached a plateau in every species, which allowed a sound interspecific comparison. Given that samples are large enough, the successive event-pairing method also has the advantage to bring more characters – potentially up to a square power more – than the acts considered in isolation.

Character independence has been mentioned for a long time as a potential problem in phylogenetics, but also as one without solution. The most obvious cases of dependence between characters must be checked for and discarded, but some dependence will necessarily exist between characters observed in a same organism which cannot be extirpated (Wiley, 1981; Simmons & Freudenstein, 2002). For instance, molecular phylogenetic studies usually do not consider that base pairs in the stem regions of ribosomal RNA are not truly independent. This problem does not occur with our case for the successive event-pairing method as shown by the $\chi^2$ goodness-of-fit tests.

Finally, this method successfully proved to be efficient and informative according to the present example of sequences of dyadic interactions in cockroach groups taken from a well-studied case in the literature (van Baaren & Deleporte, 2001; van Baaren et al., 2002, 2003a). In contrast with the undecided analysis of the repertoires which were very similar, the analyses of the event-pairing data provided a high number of independent characters, which resulted in a phylogenetic tree fully resolved and highly consistent, not much different and not less consistent than the one retrieved with morphology and molecules. For example, 60 characters (or 89 with the data set including frequencies) potentially support the monophyly of the Zetoborinae, a subfamily which is well supported based on other data (Roth, 1970; Grandcolas, 1993a, 1998; Pellens et al., 2007a, b). Our study also refuted the old and recurrent belief that behaviour is more homoplastic than other phenotypic traits, as already argued by McLennan et al. (1988), Wenzel (1992), De Queiroz & Wimberger (1993) and Proctor (1996). Consistency and retention indices and numbers of informative characters were similar between analyses C (behaviour) and D (morphology and molecules). In brief, our analysis suggested that even non-stereotyped sequences, which are usually considered more variable and less species-specific, also contain phylogenetic information. It is worth noting that phylogenetic analysis of habitats in the subfamily Zetoborinae revealed much less consistency than the present behavioural analysis (Grandcolas, 1998; Pellens et al., 2007a), in that there was more homoplasy in habitat changes than in social behaviour for this clade.

**Gregarious behaviour in Zetoborinae cockroaches**

Behavioural transitions analysed via our method included many informative characters, few of which were homoplastic. These characters also supported relationships among species in a way which did not appear biased, as also shown by the comparison with morphological and molecular analyses. For example,
species did not cluster in broad behavioural categories or potentially non-natural classes, such as gregarious vs. solitary species. The solitary species *Thanatophyllum akitetum* was not the sister group of gregarious species, but it was nested within ingroup taxa, suggesting that some synapomorphic behavioural transitions were shared by two gregarious species and the solitary one, but not with the other gregarious species, including the outgroup. Additionally, the analysis identified some autapomorphic characters which made sense in the context of solitary life habits. With the same perspective, the acts supporting Zetoborinae – both the solitary and gregarious – with respect to the gregarious Blaberinae outgroup corresponded mainly to absences of agonistic or avoiding acts in response to varied acts such as antennations or moving away (PS, PP, SJ and WA, WP, RO or FP respectively). Zetoborinae displayed other negative acts in answer to positive solicitation (transitions which are not expressed by *Eublaberus*). Those negative acts were SA (agonistic), GD, GA, RO, WD (avoiding). Thus, *Eublaberus* and the Zetoborinae studied here displayed different agonistic and avoiding acts in several particular situations. This suggests that Zetoborinae were more disposed than *Eublaberus* to display promoting acts (mostly antennation) in front of negative acts and that they gained the behaviour of displaying ‘avoiding’ acts (GD, GA, RO, WD) to stop interactions rather than agonistic behaviours (or that *Eublaberus* lost it and acquired aggressive behaviour). Zetoborinae did not appear to be prone to more or less social behaviour than *Eublaberus*; they displayed a different kind of social behaviour. Finally, *Lanxobliatta* and *Phortioeca*, two genera known to be closely related (Roth, 1970; Grandcolas, 1993a, 1998), clustered together on the basis of similar behavioural transitions involving antennation.

Using non-stereotyped behaviours in phylogenetic analyses opens new avenues for studies in behaviour evolution. In contrast to restricted stereotyped behaviours such as courtships or nest-building, non-stereotyped behaviours represent most of the behavioural activity of many species, such as feeding, foraging, playing, interacting, etc. (McFarland, 1993). Since long ago, these non-stereotyped behaviours were generally considered as less species-specific in their characteristics and composed of acts often widespread in related species, hence not chiefly adequate for comparative studies (e.g. Hinde & Tinbergen, 1958). Conversely, they were commonly analysed in psychological or sociological studies where the analysis of non-stereotyped sequences is a frequent and important matter (Abbott, 1995; Abbott & Tsay, 2000; Schlich, 2001; Elzinga, 2003; Van der Aalst et al., 2003; Hay et al., 2004).

The successive event-pairing methodology and the present case study show that non-stereotyped behaviours are informative from a phylogenetic and evolutionary point of view. This brings many characters potentially useful to study phylogenetic relationships, the adequacy of which can be established with a simple but careful preliminary statistical treatment. Once the phylogenetic tree is reconstructed, the occurrence of each kind of transition among two acts can be mapped on the tree to understand where and how it evolved, allowing for the more detailed elucidation of behavioural evolution.

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REFERENCES


Goloboff PA, Mattoni CI, Quinteros SA. 2006. Continuous characters analysed as such. Cladistics 22: 589–601.


Tinbergen N. 1959. Comparative studies of the behaviour of gulls (Laridae); a progress report. Behaviour 15: 1–70.


SUPPLEMENTARY MATERIAL

The following material is available for this article online:

Appendix S1. Transitions matrices of the five cockroaches.

This material is available as part of the online article from:
(This link will take you to the article abstract).

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