CLADISTIC INFERENCE AND EVOLUTIONARY SCENARIOS: LOCROMOTORY STRUCTURE, FUNCTION, AND PERFORMANCE IN WATER STRIDERS

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Abstract — A research methodology that aims to reveal how historical changes in environmental conditions (or selective regimes) have shaped the adaptive evolution of clades is applied to the adaptive evolution of water striders and their allies (Hemiptera—Heteroptera, Gerromorpha), a group of semiaquatic insects which includes species that are conspicuously adapted to life on the surface film of water. Based upon reconstructed phylogenies for the higher gerromorphan taxa, the hypothesis that the hygropetric zone is the ancestral one is confirmed for the Mesovelidae, Hebridae and the clade comprising the Paraphrynovelidae, Macrovelidae and Hydrometridae, but not for the Hermatobadidae and Veliidae. There is no support for the hypothesis that the intersection zone was a sort of transitional zone during the ecological evolution of pleustonic bugs. It is shown that the unique morphological and behavioural traits of the most derived members of this group evolved after inferred historical changes in environmental conditions and therefore qualify as adaptations in the sense of Gould and Vrba (1982), Coddington (1988) and Baum and Larson (1991). Other predictions about the adaptive evolution of gerromorphan bugs do not pass the cladistic test. The study illustrates that cladistic inference is a valuable tool in clarifying and sharpening retrospective explanations of complex evolutionary scenarios.

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Introduction

One of the central themes in evolutionary biology is the search for a functional (adaptive) fit between the traits (morphological, physiological, behavioural or ecological) of an organism and its environment (Gould and Lewontin, 1979; Futuyma, 1986). Most studies of adaptation, however, concentrate on the maintenance of traits under actual environmental conditions where the interaction between the organism and its environment can be studied and measured directly. In the most favourable circumstances, the outcome of experimental manipulations of adaptive traits can be predicted and tested (e.g. Arnqvist and Rowe, 1995). Along with the increasing acceptance of cladistic approaches to infer phylogenetic relationships among taxa has come an emphasis on the concept of adaptation having an important historical component (e.g. Coddington, 1988; Wanntorp et al., 1990; Brooks and McLennan, 1991; Harvey and Pagel, 1991; and several chapters in Eggleton and Vane-Wright, 1994).

The definition of adaptation has been a much debated issue in evolutionary biology (e.g. Baum and Larson, 1991; Lauder et al., 1993; Miles and Dunham, 1993; Losos and Miles, 1994). Coddington (1988: 5) was among the first investigators to define adaptation in cladistic terms: “adaptation is apomorphic function due to natural selection”, thereby agreeing with Gould and Vrba (1982) that an
evolutionary definition of adaptation must have a historical component specifying selection as the evolutionary agent responsible for its initial appearance and subsequent maintenance. An adaptive trait can, in principle, be treated like any other character. Adopting a phylogenetic (cladistic) approach, however, provides a method for distinguishing the primitive (plesiomorphic) state from the derived (apomorphic) state(s) of a particular adaptive trait.

Following Gould and Vrba (1982), an adaptation can be any trait that enhances fitness in the organisms carrying this trait and whose modification in time was a result of selection for its current biological role. A trait that has current utility, but which did not result from natural selection for its present biological role, was termed an *exaptation*. Since historical genesis by natural selection is central to this definition of adaptation, a methodology testing hypotheses of adaptive scenarios must incorporate some reference to the environmental conditions1 in which a particular trait evolved.

A research methodology that aims to reveal how historical changes in environmental conditions have shaped the adaptive evolution of clades has been described in detail by Baum and Larson (1991) and outlined by Lauder et. al. (1993). It is here assumed that a corroborated phylogenetic hypothesis is available for the focal group of organisms. The methodology is composed of four steps (here restated in cladistic terminology):

1. Identify environmental conditions that might have influenced the evolution of clades. Infer historical changes in environmental conditions with reference to the reconstructed phylogeny of the focal group.
2. Estimate functional attributes2 that might have conferred increased fitness to the organisms in a given environment.
3. Describe the characters or traits associated with such functional attributes and infer plesiomorphic–apomorphic sequences of change by character optimization on the reconstructed phylogeny of the focal group.
4. Measure the relative performance in plesiomorphic (ancestral) and apomorphic (derived) states of functional attributes.

If the apomorphic state of an attribute can be shown to have a superior performance relative to its plesiomorphic state, and can be inferred to have evolved *after* historical changes in environmental conditions, then, and only then, is the trait or attribute an adaptation (Fig. 1a). If the trait with superior performance evolved simultaneously with the environmental change (Fig. 1b), then the adaptation is ambiguous (unless there are other ways to decide which came first). Finally, the trait with superior performance is an exaptation (Gould and Vrba, 1982) if it evolved *before* historical changes in environmental conditions (Fig. 1c).

In the present study, this research methodology is applied to water striders and their allies (Hemiptera–Heteroptera, Gerromorpha), a group of semiaquatic

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1"Environmental conditions" in this context is equivalent to the concept of "selective regime" introduced by Baum and Larson (1991: 4) and defined as: "the aggregate of all such environmental and organismic factors that combine to determine how natural selection will act upon character variation".

2I follow Deleporte (1993) in making a distinction between *traits* (=characters) and *attributes*, the latter being organismic traits (e.g. behaviour) for which primary homology (de Pinna, 1991) can be roughly assessed, but which are not introduced into the data matrix supporting phylogeny construction.
insects which includes species that are conspicuously adapted to life on the surface film of water, especially with respect to locomotion, feeding, reproductive behaviour and life history (Andersen, 1982a; Spence and Andersen, 1994). It is shown that the unique morphological and behavioural traits of the most derived members of this group evolved after inferred historical changes in environmental conditions and therefore qualify as adaptations as earlier suggested (Andersen, 1979, 1982a). Other predictions about the adaptive evolution of gerromorphan bugs do not pass the cladistic test. It is argued that cladistic inference is a valuable tool in clarifying and sharpening retrospective explanations of adaptive evolution and for suggesting strategies for further research.


Phylogenetic Background

The group of hemipterans known as water striders or pond skaters are familiar insects, readily observed as they skate along the surface of ponds and streams. The water striders and their allies, collectively called semiaquatic bugs (infraorder Gerromorpha), are distributed world-wide with approximately 1500 species. The current classification of semiaquatic bugs comprises eight families. Andersen (1982a) reconstructed the phylogeny of the group to the family and subfamily level, and in some families down to the level of genera. The cladogram (Fig. 2) summarizes the phylogenetic relationships between higher gerromorphan taxa (families, subfamilies, tribes).

The evolution of structural and functional traits was inferred by reference to a representative sample of gerromorphan bugs for which comparative data on locomotory structure and function are available (Andersen, 1976, 1982a, 1989b; Foster and Treherne, 1980; Foster, 1989). The genera studied are: Mesovelia (Mesoveliidae), Hebrus (Hebridae), Macrovelia (Macroveliidae), Hydrometra (Hydrometridae), Hermatobates (Hermatobatidae), Microvelia (Veliidae, Microveliinae), Halovelia (Veliidae, Haloveliinae), Velia (Veliidae, Velinae), Rhagovelia (Veliidae, Rhagoveliinae), Gerris (Gerridae, Gerrinae), Eotrechus (Gerridae, Eotrechinae), Ptilomera (Gerridae, Ptilomerinae) and Halobates (Gerridae, Halobatinae). The cladogram (Fig. 3) depicts the phylogenetic relationships among these genera as inferred from available reconstructions using cladistic methods (Andersen, 1982a).

Comparative Data

Environmental Zones

Semiaquatic bugs live in a wide range of humid terrestrial and aquatic habitats, including both freshwater and marine habitats (Andersen, 1979, 1982a). Most species live on the water surface and their body and legs are covered by hydrofuge hair layers which prevent wetting (Andersen, 1977). In their natural environment, these bugs chiefly use their legs for local movements and dispersal. Secondary loss of wings and flight musculature is very common and natural populations of most species are predominantly flightless (Andersen, 1982a, 1993). For feeding, various activities associated with reproduction and escape from predators, these bugs rely on their ability as pedestrians. It is therefore reasonable to assume that the relative proportion of solid matter and free water surface is an important element of the environmental conditions or “selective regime” (sensu Baum and Larson, 1991) of semiaquatic bugs and their ancestors. Following previously used terminology (Andersen, 1982a), the following environmental zones are delimited: (1) the hygropetric zone where the substrate for locomotion is mainly solid; (2) the intersection zone with a varying mixture of solid substrate and water surface; and (3) the pleustonic zone which is essentially the air–water interface.

Locomotory Function and Performance

A number of semiaquatic bugs have been subject to studies of locomotory
behaviour and function (Darnhofer-Demar, 1969b; Andersen, 1976, 1982a, 1989b; Cheng, 1977; Foster and Treherne, 1980; Foster, 1989). Walking on alternating tri-pods of support (legs belonging to the same segment are moved asynchronously) is the ordinary way of locomotion in terrestrial bugs and undoubtedly the ancestral one in gerromorphan bugs. On a solid substrate, these bugs prefer to walk (except the gerrids which usually leap). On the water surface, locomotion is achieved in three different ways: (1) by walking as described above; (2) by rowing with simultaneous strokes of the middle legs while the two other leg pairs slide on the surface film; or (3) by skating (or "jump-and-slide" movements), where the simultaneous power strokes of the middle legs make the insect take off from the water surface; the movement is extended after touch down by passive sliding on the almost frictionless surface film (in contrast to leaping in land insects).

Locomotory function and performance on the water surface is listed in Table 1 for selected species of semiaquatic bugs. This tabulation provides crude estimates of locomotory performance for the terminal taxa of the cladogram (Fig. 3). In general, all semiaquatic bugs except gerrids perform well when moving on land. Walking is the preferred mode in most species, but gerrids tend to leap when pursued. Individuals of Hebrus, Macrovelia and Hydrometra move relatively slowly along the surface film, while those of Mesovelia move fast (relative to their body size). Water striders belonging to the family Gerridae have a superior locomotory performance on the water surface, followed by those belonging to the Veliidae, in particular Halovelia (Andersen, 1989b) and Rhagovelia. Individuals of Hermatobates move fast both on land and water (Cheng, 1977; Foster, 1989).
Structural Traits

Semiaquatic bugs are structurally quite diverse (representative species shown in Fig. 4). Most species are relatively small insects (total length 2-5 mm), but there is quite extensive variation in size within the large families Veliidae (1.2-9.8 mm) and, in particular, the Gerridae (1.7-35 mm), both between species and between sexes (Andersen, 1994). Morphological structures associated with locomotion are summarized in Table 2. In *Mesovelia* spp., the thoracic segments are almost equal in size and, at least in wingless individuals (Fig. 4) of the same general structure. The legs originate from the thoracic venter at almost equal distances from each other. The inclination of the meso- and metacoxal axes (the line drawn between the points of coxal and trochanteral articulation) is moderate (40° and 25°, respectively). The hind legs are much longer than the middle legs with slightly thickened femora. The most powerful muscles involved in leg movements are the coxal rotators and trochanteral depressors (Andersen, 1976). The locomotory structures of *Mesovelia* are quite similar to those of most terrestrial bugs and are therefore inferred to be plesiomorphic.

| Mes | Heb | Mac | Hyd | Hem | Mic | Hal | Vel | Rha | Ger | Eot | Pti | Hab |
|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|
| Thorax structure | | | | | | | | | | | | |
| 1. Mesothorax prolonged | - | - | - | + | + | + | + | + | + | + | + | + |
| 2. Metathorax prolonged | - | - | + | - | - | - | - | - | - | - | - | - |
| 3. Coxae laterally displaced on thorax | - | + | + | + | + | + | + | + | + | + | + | + |
| 4. Mesocoxal axes vertical | - | - | + | - | - | - | - | - | - | - | - | - |
| 5. Meso- and metacoxal axes horizontal | - | - | + | - | - | - | + | + | + | + | + | + |
| 6. Mesotrochanteral depressors more powerful than coxal rotators | - | - | + | - | - | - | - | + | + | + | + | + |
| Leg structure | | | | | | | | | | | | |
| 7. Middle and hind legs prolonged, femora slender | - | - | - | - | - | - | - | - | + | + | + | + |
| 8. Middle legs slightly longer than hind legs | - | - | - | - | + | + | + | + | + | + | + | + |
| 9. Middle legs much longer than hind legs | - | - | - | - | - | - | + | - | + | + | + | + |
| 10. Number of tarsal segments reduced (from three to two segments) | - | + | + | - | + | + | + | - | + | + | + | + |
| 11. Claws inserted preapically on tarsi | - | - | + | + | + | + | + | + | + | + | + | + |
| 12. Pretarsal structures modified as swimming fans | - | - | - | - | - | + | - | - | - | - | - | - |
| 13. Pretarsal structures reduced | - | - | - | - | - | - | - | - | + | + | + | + |
| Other structural traits | | | | | | | | | | | | |
| 14. Adults obligatory flightless | - | - | - | - | + | - | - | - | - | - | - | - |
| 15. Abdomen reduced in length | - | - | - | + | - | - | - | - | - | - | - | - |
The thoracic structure of Hebrus (Fig. 4), Macrovelia, Microvelia, Halovelia, Velia (Fig. 4) and Rhagovelia is quite similar to the ancestral type mentioned above, except that the points of insertion of the meso- and metacoxae are displaced laterally on the body. The leg structure (apart from the tarsus) is also quite unspecialized, except that the middle legs of Velia and, in particular Halovelia and Rhagovelia, are slightly longer than the hind legs.

Compared to the ancestral type, the thorax is more or less modified in several groups of semiaquatic bugs. Hydrometra species have an extremely long and slender head and body, including the thorax (Fig. 4). The middle legs originate laterally on the thorax, and the mesocoxal axis is vertical (=90°). All femora are extremely slender. The lateral position of the coxae increases the action angle of the vertical leg movements whereas the direction of pull of the trochanteral muscles falls almost exactly in the frontal plane (Andersen, 1976).

The thorax of Hermatobates spp. and members of the family Gerridae (Gerris, Eotrechus, Ptthomera and Halobates) is much more radically modified (Fig. 4). The mesothorax is greatly prolonged and the coxal cavities of the middle and hind legs are far removed from those of the front coxae. The meso- and metacoxae are inserted clearly on the sides of thorax and the coxal axes have an almost horizontal inclination (=0°). The middle and hind legs are long and slender with the middle pair being much longer than the hind pair. The joints of the two posterior pairs of legs allow extremely wide movements of the femora. Associated with the restructuring of the gerrid thorax are radical changes of the leg musculature. The most powerful muscles are the mesotrochanteral depressors, which produce the horizontal thrust of the middle legs, while the vertical component of the leg movements is produced by the two coxal rotators (Darnhofer-Demar, 1969b; Andersen, 1976). In the coral treaders, Hermatobates, and the sea skaters, Halobates, the middle and hind legs appear to originate from near the posterior end of the extremely short abdomen (Andersen and Polhemus, 1976; Andersen, 1982a) and the action angles of the middle and hind legs are even more extreme. The prolongation of the gerrid mesothorax has moved the gravity center of the body forward to a position in front of the fulcrum of the middle legs.

Mesovelia, Macrovelia and Hydrometra have three tarsal segments (the ancestral number in bugs) and the well developed, hook-shaped claws are inserted apically on the last segment. Microvelia, Halovelia, Gerris and other species of the families Veliidae and Gerridae often have less than three tarsal segments and the claws are inserted preapically, that is in a cleft before the apex of the last tarsal segment. When retracted, the claws are lifted off from the water surface film (Andersen, 1976). The middle and hind leg claws are relatively weak in Gerris and more or less reduced in Ptthomera and Halobates. In contrast, eotrechine gerrids have well developed claws which in Eotrechus spp. seem to be inserted apically on the tarsi (Andersen, 1982b; Polhemus and Andersen, 1984). Other pretarsal structures include the arolium which, in several veliids, form swimming fans, alone or together with the modified claws (Andersen, 1982a). The most elaborate swimming fan is found in species belonging to the genus Rhagovelia and its allies and consists of a fan of feathered hairs which is unfolded during the thrust stroke of the middle legs (Andersen, 1976).

Many species of semiaquatic bugs that are capable of flight are polymorphic for wing and flight muscle development (Andersen, 1982a, 1993). However, natural
populations of most water striders are predominantly flightless and adults of no marine species have wings (Andersen and Polhemus, 1976). Since the lack of flight muscles and other thoracic modifications associated with the loss of wings may influence locomotion, this character is included in Table 2.

**Cladistic Analyses**

**Environmental Changes**

Figure 2 shows the three environmental zones, the hygropetric, intersection and the pleustonic zone, superimposed on a cladogram of relationships between higher taxa (for details about the reconstruction of this phylogeny, see Andersen, 1982a). The environmental zone of possible outgroups of the Gerromorpha (Enicocephalomorpha, Dispsocoromorpha: Schuh, 1986; Wheeler et al., 1993; Schuh and Slater, 1995) is definitely more terrestrial than aquatic and the hygropetric zone is therefore inferred to be the ancestral one in semiaquatic bugs. Optimizing the three environmental zones on the cladogram using MacClade 3.05 (Maddison and Maddison, 1992) yields the following results indicated by different shading of the branches of the cladogram (Fig. 2).

![Cladogram with environmental zones](image-url)

Fig. 2. Environmental zones superimposed upon the reconstructed phylogeny of higher taxa (families, subfamilies, tribes) of semiaquatic bugs. Further explanations in text.
In the most parsimonious optimization, transitions between the hygropetric zone and the pleustonic zone are inferred to have occurred once in the Hydrometridae (the genus Bacillometra) and once on the internode below the clade comprising the Hermatobatidae, Veliidae and Gerridae (the superfamily Gerroidea). However, delaying this transition to having occurred independently in the three families (Fig. 2, light arrows) only requires one extra step. Besides, the assignment of the Hermatobatidae to the pleustonic zone is somewhat ambiguous. The coral treadsers, Hermatobates spp., inhabit the intertidal zone of coral coasts (like the coral bugs, Halovelia: Andersen, 1989b). They retreat to crevices and holes in coral blocks during high tides (Foster, 1989) and move around with the same agility on the surface of corals and tidal pools.

Transitions between environmental zones have been complex but the optimization (Fig. 2) generally favours direct changes between zones, e.g. hygropetric—pleustonic, and disfavours the idea of the intersection zone as intermediate between the hygropetric and the pleustonic zone. There are, however, three cases where the intersection zone may fall between the two others zones (Fig. 2): a forward change in the Hydrometridae and Veliidae-Microveliinae, and a reversal in the Veliidae-Veliinae.

Transitions between the hygropetric zone and the intersection zone have occurred in Mesovelia, the Hebridae (Merragata) and Hydrometra. The intersection zone is also occupied by species belonging to the veliid subfamilies Microveliinae and Veliinae but use of the intersection zone may have evolved after transition from the pleustonic zone. Reversed transitions from the pleustonic zone to the hygropetric zone have seemingly occurred in some microveliine genera, e.g. Hebrovelia (Andersen, 1982a), Aegilipsovelia (Polhemus, 1970), Baptista (Andersen, 1989a), Aegilipsicola (Polhemus and Polhemus, 1994), etc. The most remarkable examples of this type of reversal are found in the gerrid subfamily Eotrechinae where species belonging to the genera Eotrechus and Onychotrechus live on humid soil and seeping rock faces (Andersen, 1980, 1982b; Polhemus and Andersen, 1984).

Marine habitats have been invaded by semiaquatic bugs about a dozen times independently of each other (Fig. 2). The family Hermatobatidae contains only marine species (Foster, 1989; Andersen, 1982a, 1991a). The most speciose genera of marine bugs are Halovelia, Xenobates (Veliidae-Haloveliniinae: Andersen, 1989b, 1991a), and Halobates (Gerridae-Halobatinae: Andersen, 1991b). The transition between limnic and marine habitats probably took place through estuaries and mangrove swamps (Andersen and Weir, 1994). Five species of ocean skaters, Halobates, are the only insects that inhabit the surface of the open ocean. This unique life-style may have evolved from mangrove-inhabiting species becoming gradually better adapted to more exposed sea conditions (Andersen, 1991b).

Changes in locomotory structure, function and performance

The evolution of structural traits associated with locomotion (Table 2) was inferred by reference to the phylogenetic relationships between selected genera.
Fig. 3. Cladogram of relationships between gerromorphan taxa (genera) selected for optimization of structural and functional traits associated with locomotion (data of Tables 1, 2). Environmental zones superimposed upon the cladogram. Inferred changes from plesiomorphic to apomorphic states marked on internodes.

The character optimization was carried out on the distribution of structural traits (Table 2) using MacClade 3.05. Each trait was polarized by rooting the cladogram between *Mesovelia* (the overall most plesiomorphic taxon; see presentation of structural and functional characters above) and the rest of the terminal taxa. When the resolution at some nodes was ambiguous (e.g. for characters 1, 5, 6, 8, 9, and 10), parallelisms were favoured over reversals. Different modes of locomotion on land and on the water surface (Table 1) were also optimized on the cladogram (Fig. 3).

The final step was to compare the inferred changes in structural and functional traits associated with locomotion (Fig. 3) with the inferred transitions between environmental zones (Fig. 2; see above) as well as the performance estimates for

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3To avoid circular arguments in cladistic inference applied to evolutionary scenarios, I follow the procedure suggested by Deleporte (1993) rather than the strict rule of independence between the traits and attributes of focal interest and the data used for phylogeny construction (e.g. Coddington, 1988; Brooks and McLennan, 1991). In the actual case, habitat zones (i.e. hygropric, intersection, pleustonic, limnic, marine) were not entered as data in phylogeny reconstruction. Several morphological features associated with locomotion were used (Andersen, 1982a), but this only amounted to about 10% of the total number of characters used in the cladistic analyses of relationships between the families of Gerromorpha, and the subfamilies of Veliidae and Gerridae.

4This procedure does not affect the total number of evolutionary steps of the optimized character, only the number of parallelisms versus reversals. It is, for example, more likely that the similar structure of the meso- and metathorax of the Hermatobatidae and Gerridae (Andersen, 1982a) evolved in a parallel way in the two families, instead of just once with a subsequent reversal to the more plesiomorphic thorax of the Veliidae.
selected species (Table 1). The results of these comparisons are summarized below and in the evolutionary diagram (Fig. 4) where the inferred major transition between the hygropetric zone and the pleustonic zone is marked. The main results of the cladistic analyses are as follows:

(1) In the basal gerromorphan lineages, locomotory structures are plesiomorphic and locomotion both on land and water is close to the supposed ancestral state of hexapods (“walking”). Nevertheless, *Mesovelia* performs very well on the surface film although “walking” on this substrate is less efficient than “rowing” (Andersen, 1976). Thus, evolution of effective locomotion on the water surface may have occurred without a major transition between zones.

(2) The apomorphic thorax structure of *Hydrometra* spp. is not associated with increased locomotory performance on the surface film and may therefore have other functional implications, e.g. in the context of feeding or escaping predators (Andersen, 1982a).

(3) The movement of the claws from an apical to a preapical position on the tarsi (Andersen, 1976) has occurred after the inferred transition between the hygropetric and pleustonic zone and may therefore be an adaptation towards life on the surface film of water.

(4) Extensive changes in thorax and leg structures have occurred in several gerromorphan lineages (*Hydrometridae, Hermatobatidae, Gerridae*). An important
functional change was the neuromuscular reprogramming involved in changing the coordination of leg movements from walking to synchronized movements of leg of the same thoracic segment ("rowing"). The completely remodelled thorax and leg structure of gerrids is associated with the highly efficient jump-and-slide locomotion ("skating") characteristic of water striders. Since most changes in locomotory structure and function have occurred after the inferred transition between the hygropetric and pleustonic zones, these apomorphic traits qualify as adaptations (Fig. 1a).

(5) The apomorphic swimming fan of *Ragovelia* is associated with increased locomotory performance. Similar swimming fans, although of different pretarsal design, have evolved several times in the Veliidae (Andersen, 1979, Fig. 12). Multiple independent evolution of functionally equivalent structures is usually taken as evidence of adaptation (Futuyma, 1986; Ridley, 1992).

(6) The family Gerridae displays a much larger size variation (1.7–35 mm) than its sister group, the Veliidae (1.2–9.8 mm). A plausible explanation is that the evolution of the highly efficient locomotion ("skating") found in gerrids has overcome functional obstacles for an increase in body size for pleustonic insects.

(7) Marine habitats have been invaded many times during the evolutionary history of gerromorphan bugs. Apart from the physiological adaptations required for insects to cope with a saline environment, the permanent loss of wings and flight capacity were apparently the only structural changes associated with the transition from limnic to marine habitats. However, behavioural changes were doubtless required for adapting to life in the intertidal zone of mangroves and coral reefs (Andersen and Polhemus, 1976).

(8) The occurrence of some gerrids (*Eotrechus, Onychotrechus*) in hygropetric habitats is inferred to represent reversals from pleustonic habitats. Yet on modest structural changes (reversals) in leg structures were associated with this major shift between environmental zones (paralleled in several veliid genera; see above).

In conclusion, most structural traits and functional attributes (items 3–8 above) are inferred to have evolved after the major lineages of gerromorphan bugs entered the pleustonic zone. Associated with these apomorphic traits and attributes is an increase in locomotory performance on the free water surface. The most probable explanation for this association is adaptation through natural selection.

**Discussion**

This study is essentially a reanalysis of comparative data presented in earlier papers (Andersen, 1976, 1979) and a monographic treatment of the semiaquatic bugs (Andersen, 1982a). These works were published before the influential papers by Gould and Vrba (1982), Coddington (1988) and Baum and Larson (1991) laid the foundation for a more precise definition of organismic adaptation and, in particular, how phylogenetic or cladistic methods can be used profitably in studies of adaptation and organismic evolution. I am therefore now able to examine some of my own previous hypotheses using more refined analytical tools.

The evolutionary history of water striders and their allies probably began in the early Mesozoic. The oldest fossil gerromorphans are of lower Cretaceous age.
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(Jell and Duncan, 1986) and indisputably belong to the families Mesoveliidae and Veliidae, some of the structurally least modified forms. Species belonging to the most apomorphic families, Hydrometridae and Gerridae, occur in Upper Paleocene deposits of Denmark (Andersen, 1982a), setting a minimum age of these families of about 55 my. A Middle Eocene fossil from western Canada was designated as belonging to a modern species group of the genus Limnopus (Andersen et al., 1993), suggesting that gerrid specializations for life on freshwater surfaces were not associated with high rates of speciation or extinction. Finally, a fossil sea skater (Halobates) from the Eocene of northern Italy (Andersen et al., 1994) suggests an early invasion of marine habitats in water striders. With this background, it is a reasonable assumption that semiaquatic bugs displayed most of their present structural and functional diversity at the beginning of the Tertiary. However, in the absence of direct historical evidence, our knowledge of the environmental conditions that shaped this evolution can only be inferred from comparative studies of extant forms.

Previous ideas about the ecological evolution of semiaquatic bugs (China, 1955) suggested an early colonization of the water surface by "littoral" bugs (the Saldidae or shore bugs were thought to be the sister group of the Gerromorpha). These bugs included the ancestor of all families except the Mesoveliidae and Hebridae. Terrestrial or hygropetric forms of the families Macroveliidae and Hydrometridae were seen as examples of a secondary return to land. Based upon reconstructed phylogenies of higher taxa, several elements of China's (1955) scenario could be falsified (Andersen, 1979, 1982a). Hygropetric habitats were inferred to be ancestral to all gerromorphan lineages except the Gerridae, and the adaptive transitions between the hygropetric and pleustonic zones (following the concept of "adaptive zone" introduced by Simpson, 1953) were visualized (Andersen, 1982a, Figs. 606–607) as being a gradual process where different lineages, independently of each other, passed through a "transitional zone" (the intersection zone) and reached the pleustonic zone where extensive adaptive radiation took place.

The present study supports only some parts of this scenario. The hypothesis that the hygropetric zone is the ancestral one is confirmed for the Mesoveliidae, Hebridae and the clade comprising the Paraphrynoveliiidae, Macroveliidae and Hydrometridae, but not for the Hermatobatidae and Veliidae. There is only weak support for the hypothesis that the intersection zone was a sort of transitional zone during the ecological evolution of pleustonic bugs. On the contrary, most of the unique structural and behavioural adaptations of water striders seem to have evolved after the inferred direct transition between the hygropetric and pleustonic zones. Additional and more exact studies of locomotory behaviour and performance in a variety of semiaquatic bugs are required to add more precision to alternative hypotheses. The cladistic analysis could also be extended profitably to encompass other aspects of water strider biology (reviewed by Andersen, 1979, 1982a; Spence and Andersen, 1994), e.g. feeding, reproductive behaviour, escape from predators and parasites and life history tactics, as exemplified by recent studies of wing polymorphism, sexual size dimorphism and mating strategies in water striders (Andersen, 1993, 1994).

Among the gerromorphan lineages, the velliids and gerrids are by far the most speciose (Fig. 4). The question of evolutionary "success" (e.g. through adaptive radiation) has in recent literature been associated with comparisons between
species numbers in sister groups (Brooks and McLennan, 1991, 1993). If species richness is a measure of success, then both the Veliidae and Gerridae are the most successful clades of semiaquatic bugs (with about 640 and 520 species, respectively). As could be predicted, the diversification of both clades occurred after they entered the pleustonic zone (Fig. 2). Regarding adaptations towards life on the water surface, the two families nevertheless differ significantly. The evolutionary success of the Gerridae was followed by a complete remodelling of thorax and leg structures and neuromuscular reprogramming associated with the highly efficient “skating” locomotion. In contrast to this, the Veliidae did not experience extensive changes of locomotory structure and function when they emerged as a monophyletic group (Fig. 3). Subsequently, locomotory diversification and extensive speciation took place in a number of lineages (e.g. the speciose genera Halovelia and Rhagovelia).

In a recent study, Brooks and McLennan (1993: 773) listed four criteria that are necessary to identify adaptive radiation: (1) the group in question contains more species that its sister group; (2) species richness is a derived characteristic within the larger clade; (3) an apomorphy present in the more species-rich group enhances the potential that adaptively driven speciation (i.e. sympatric speciation or speciation by peripheral isolation) will occur; and (4) adaptively driven speciation modes played the dominant role in the speciation of the more species-rich group. None of these apply readily to the evolution of gerromorphan bugs at the level of the present study (families, subfamilies). Admittedly, the clade Veliidae+Gerridae (ca. 1160 species) is far more speciose than its sister group, the Hermatobatidae (8 species), but the frequency of different speciation modes (especially sympatric versus allopatric speciation) is difficult to assess for the large and widespread families. The diversity of different life forms found within these families, however, suggests a significant adaptive radiation most likely founded on locomotory adaptations.

As exemplified by this case study, the application of the research methodology suggested by Coddington (1988) and Baum and Larson (1991) is able to disclose weaknesses in currently accepted scenarios for adaptive evolution and suggest alternative explanations as well as the most profitable strategy for further studies. Because data on the function or “performance” (absolute or relative) of organismic traits are hard to obtain for a diversity of taxa, this research methodology has rarely been employed in studies exceeding a small number of species. Not surprisingly, the majority of case studies found in the literature use vertebrate species as their focal group (reviewed by Miles and Dunham, 1993; see also Eggleton and Vane-Wright, 1994). However, studies using invertebrates, in particular insects, are accumulating at an increasing rate (e.g. Coddington, 1986; Mitter et al., 1988; Carpenter, 1989; Ambruster, 1992; Carpenter et al., 1993; Wenzel, 1993; see review by Miller and Wenzel, 1995).

The use of phylogenies to test hypotheses of organismic adaptation has recently been criticized (Lauder et al., 1993; Frumhoff and Reeve, 1994; Leroi et al., 1994) on the grounds that the phylogenetic or cladistic approach can only test hypotheses about patterns but not about the evolutionary processes that shaped these patterns. Already, Coddington (1988) has admitted that his cladistic test of adaptational hypotheses may fail if the trait examined had evolved by causes other than natural selection (e.g. developmental canalization or genetic drift). Leroi et al.
(1994) add further examples and argue that many complex mechanisms of evolution can only be disclosed if selection and genetic variation are measured directly in many closely related species. This is probably true, and most contemporary workers realize that the new methods in comparative biology (Brooks and McLennan, 1991; Harvey and Pagel, 1991; Coddington, 1994) do not hold the promise of being able to completely resolve the processes of evolution. The conclusion by Leroi et al. (1994: 398) “Indeed, process is not all that is interesting in evolution; we can still learn a great deal about the nature of organismic diversity by describing and analyzing patterns of character evolution only” is nevertheless too discouraging. I am a little more confident about the power of resolution of cladistic inference when applied to cases like that presented in this paper. After all, viewing organismic adaptation as an apomorphic function due to natural selection is deeply rooted in the Darwinian theory of evolution.

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REFERENCES

AMBRUSTER, W. S. 1992. Phylogeny and the evolution of plant–animal interactions. BioScience 42: 12–20.

ANDERSEN, N. M. 1976. A comparative study of locomotion on the water surface in semiaquatic bugs (Insecta, Hemiptera, Gerromorpha). Vidensk. Medd. Dan. Naturhist. Foren. 139: 337–396.

ANDERSEN, N. M. 1977. Fine structure of the body hair layers and morphology of the spiracles of semiaquatic bugs (Insecta, Hemiptera, Gerromorpha) in relation to life on the water surface. Vidensk. Medd. Dan. Naturhist. Foren. 140: 7–37.

ANDERSEN, N. M. 1979. Phylogenetic inference as applied to the study of evolutionary diversification of semiaquatic bugs (Hemiptera: Gerromorpha). Syst. Zool. 28: 554–578.

ANDERSEN, N. M. 1980. Hygropteric water striders of the genus Onychotrechus Kirkaldy with description of a related genus (Insecta, Hemiptera, Gerridae). Steenstrupia 6: 113–146.

ANDERSEN, N. M. 1982a. The Semiaquatic Bugs (Hemiptera, Gerromorpha). Phylogeny, adaptations, biogeography, and classification. Entomonomograph 3: 1–455.

ANDERSEN, N. M. 1982b. Semiterrestrial after striders of the genera Eotrechus Kirkaldy and Chimarrhometra Bianchi (Insecta, Hemiptera, Gerridae). Steenstrupia 9: 1–25.

ANDERSEN, N. M. 1989a. The Old World Microveliinae (Hemiptera: Veliidae). II. Three new species of Baptista Distant and a new genus from the Oriental region. Entomol. Scand. 19: 363–380.

ANDERSEN, N. M. 1989b. The coral bugs, genus Halovelia Bergroth (Hemiptera, Veliidae). II. Taxonomy of the H. malaya-group, cladistics, ecology, biology, and biogeography. Entomol. Scand. 20: 179–227.

ANDERSEN, N. M. 1991a. Cladistic biogeography of marine water striders (Hemiptera, Gerromorpha) in the Indo-Pacific. Aust. Syst. Bot. 4: 151–163.

ANDERSEN, N. M. 1991b. Marine insects: genital morphology, phylogeny and evolution of sea skaters, genus Halobates (Hemiptera, Gerridae). Zool. J. Linn. Soc. 103: 21–60.
ANDERSEN, N. M. 1993. The evolution of wing polymorphism in water striders (Gerridae): a phylogenetic approach. Oikos 67: 433-443.

ANDERSEN, N. M. 1994. The evolution of sexual size dimorphism and mating systems in water striders (Hemiptera: Gerridae): a phylogenetic approach. Ecocene 1: 208-214.

ANDERSEN, N. M. AND J. T. POLHEMUS. 1976. Water-striders (Hemiptera: Gerridae, Velidae, etc.). In: L. Cheng (ed.). Marine Insects. North-Holland Publishing Company, Amsterdam, pp. 187-224.

ANDERSEN, N. M. AND T. A. WEIR. 1994. Austroboles rivularis gen. et sp. nov., a freshwater relative of Halobates (Hemiptera, Gerridae) with a new perspective on the evolution of sea skaters. Invertebr. Taxon. 8: 1-15.

ANDERSEN, N. M., A. Farma, A. Minelli and G. Piccoli. 1994. A fossil Halobates from the Mediterranean and the origin of sea skaters (Hemiptera, Gerridae). Zool. J. Linn. Soc. 112: 479-489.

ANDERSEN, N. M., J. R. SPENCE and M. V. H. WILSON. 1993. 50 million years of structural stasis in water striders (Hemiptera, Gerridae). Am. Entomol. 39: 174-176.

ARNQUIUST, G. AND L. Rowe. 1995. Sexual conflict and arms races between the sexes: a morphological adaptation for control of mating in a female insect. Proc. R. Soc. Lond. B. Biol. Sci. 261: 123-127.

BAUM, D. A. AND A. LARSON. 1991. Adaptation reviewed: a phylogenetic methodology for studying character macroevolution. Syst. Zool. 40: 1-18.

BROOKS, D. R. AND D. A. MCLENNAN. 1991. Phylogeny, Ecology, and Behaviour. A Research Program in Comparative Biology. The University of Chicago Press, Chicago, London.

BROOKS, D. R. AND D. A. MCLENNAN. 1993. Comparative study of adaptive radiations with an example using parasitic flatworms (Platyhelminthes: Cercocmeria). Am. Nat. 142: 755-778.

CARPENTER, J. M. 1989. Testing scenarios: wasp social behaviour. Cladistics 5: 131-144.

CARPENTER J. M., J. E. STRASSMANN, S. Turillazzi, C. R. HUGHES, C. R. Solfs and R. SERVO. 1993. Phylogenetic relationships among paper wasps social parasites and their hosts (Hymenoptera: Vespidae; Polistinae). Cladistics 9: 129-146.

CHENG, L. 1977. The elusive sea bug Hermatobates (Heteroptera). Pan-Pac. Entomol. 53: 87-97.

CHINA, W. E. 1955b. The evolution of the water bugs. Symposium on Organic Evolution. Bull. Nat. Inst. Sci., India 7: 91-103.

CODDINGTON, J. A. 1986. The monophyletic origin of the orb web. In: W. A. Shear (ed.). Spider Webs and Spider Behaviour. Stanford University Press, Palo Alto, California, pp. 319-363.

CODDINGTON, J. A. 1988. Cladistic tests of adaptational hypotheses. Cladistics 4: 3-22.

CODDINGTON, J. A. 1994. The roles of homology and convergence in studies of adaptation. In: P. Eggleton and R. I. Vane-Wright (eds). Phylogenetics and Ecology. Academic Press, London, etc., pp. 53–78.

DARNHOFER-DEMAR, B. 1969a. Zur Fortbewegung des Wasserlaufers Genis kozmtris L. auf der Wasseroberfläche. Zool. Anz. Suppl. 32: 430-439.

DARNHOFER-DEM MAR, B. 1969b. Zur Funktionsmorphologie der Wasserläufer. I. Die Morphologie des Lokomotionsapparates von Gerris lacustris L. (Heteroptera: Gerridae). Zool. Jb. Anat. 86: 28-66.

DELEPORTE, P. 1993. Characters, attributes and tests of evolutionary scenarios. Cladistics 9: 427-432.

DE PINNA, M. C. C. 1991. Concepts and tests of homology in the cladistic paradigm. Cladistics 7: 367-394.

EGGLETON, P. AND R. I. VANE-WRIGHT (eds). 1994. Phylogenetics and Ecology. Linnean Society Symposium Series, No. 17, Academic Press, London, San Diego, New York, Boston, Sydney, Tokyo, Toronto.

FOSTER, W. A. 1989. Zonation, behaviour and morphology of the intertidal coral-treader Hermatobates (Hemiptera: Hermatobatidae) in the south-west Pacific. Zool. J. Linn. Soc. 96: 87-105.

Foster, W. A. AND J. E. TREHERNE. 1980. Feeding, predation and aggregation behaviour in a marine insect, Halobates robustus Barber (Hemiptera: Gerridae), in the Galapagos Islands. Proc. R. Soc. Lond. B Biol. Sci. 209: 539-553.
FRUMHOF, P. C. AND H. K. REEVE. 1994. Using phylogenies to test hypotheses of adaptation: a critique of some current proposals. Evolution 48: 172–180.

FUTUYMA, D. J. 1986. Evolutionary Biology (2nd edn). Sinauer Associates, Sunderland, Mass.

GOULD, S. J. AND R. LEWONTIN. 1979. The spandrels of San Marco and the Panglossian paradigm: A critique of the adaptationist programme. Proc. R. Soc. Lond. B Biol. Sci. 205: 581–598.

GOULD, S. J. AND E. S. VRBA. 1982. Exaptation—A missing term in the science of form. Paleobiology 8: 4–15.

HARVEY, P. H. AND M. D. PAGEL. 1991. The Comparative Method in Evolutionary Biology. Oxford Series in Ecology and Evolution. Oxford University Press, Oxford, New York, Tokyo.

JELL, P. A. AND P. M. DUNCAN. 1986. Invertebrates, mainly insects, from the freshwater, Lower Cretaceous, Koonwarra Fossil Bed (Korumburra Group), South Gippsland, Victoria. Mem. Ass. Australas. Palaeontols 3: 111–205.

LAUDER, G. V. 1990. Functional morphology and systematics: Studying functional patterns in an historical context. Annu. Rev. Ecol. Syst. 21: 317–340.

LAUDER, G. V., A. M. LEROI AND M. R. ROSE. 1993. Adaptations and history. Trends Ecol. Evol. 8: 294–297.

LEROI, A. M., M. R. ROSE AND G. V. LAUDER. 1994. What does the comparative method reveal about adaptation? Am. Nat. 143: 381–402.

LOSOS, J. B. AND D. B. MILES. 1994. Adaptation, constraint, and the comparative method: Phylogenetic issues and methods. In: P. C. Wainwright and S. M. Reilly (eds). Ecological Morphology. The University of Chicago Press, Chicago, London, pp. 60–98.

MADDISON, W. AND D. MADISON. 1992. MacClade. VER. 3.0 Analysis of Phylogeny and Character Evolution. Sinauer Associates, Sunderland, Massachusetts.

MILES, D. B. AND A. E. DUNHAM. 1993. Historical perspectives in ecology and evolutionary biology: the use of phylogenetic comparative analysis. Annu. Rev. Ecol. Syst. 24: 587–619.

MILLER, J. S. AND J. W. WENZEL. 1995. Ecological characters and phylogeny. Annu. Rev. Entomol. 40: 389–415.

MITTER, C., B. FARRELL AND B. WIEGMANN. 1988. The phylogenetic study of adaptive zones: has phytophagy promoted insect diversification? Am. Nat. 132: 107–128.

POLHEMUS, J. T. 1970. A new genus of Veliidae from Mexico (Hemiptera). Proc. Entomol. Soc. Wash. 72: 443–448.

POLHEMUS, J. T. AND N. M. ANDERSEN. 1984. A revision of Amemboa Esaki with notes on the phylogeny and ecological evolution of eotrechine water striders (Insecta, Hemiptera, Gerridae). Steenstrupia 10: 65–111.

POLHEMUS, J. T. AND H. C. CHAPMAN. 1979. Family Macroversilliidae. In: A. S. Menke (ed.). The semiaquatic and aquatic Hemiptera of California (Heteroptera: Hemiptera). Bull. Calif. Insect. Surv. 21: 46–48.

POLHEMUS, J. T. AND D. A. POLHEMUS. 1994. Four new genera of Microveliinae (Heteroptera) from New Guinea. Tijdschr. Entomol. 137: 57–74.

SCHUH, R. T. 1986. The influence of cladistics on heteropteran classification. Annu. Rev. Entomol. 31: 67–93.

SCHUH, R. T. AND J. A. SLATER. 1995. True Bugs of the World (Hemiptera: Heteroptera). Comstock Publishing Associates, Cornell University Press, Ithaca, London.

SIMPSON, G. G. 1953. The Major Features of Evolution. Columbia University Press, New York, London.

SPENCE, J. R. AND N. M. ANDERSEN. 1994. Biology of water striders: interactions between systematics and ecology. Annu. Rev. Entomol. 39: 101–128.

WANNTORP, H., D. R. BROOKS, T. NILSSON, S. NYLIN, F. RONQUIST, S. G. STEARNS AND N. WEDELL. 1990. Phylogenetic approaches in ecology. Oikos 57: 119–132.

WENZEL, J. W. 1993. Application of the biogenetic law to behavioral ontogeny: a test using nest architecture in paper wasps. J. Evol. Biol. 6: 229–247.

WHEELER, W. C., R. T. SCHUH AND R. BANG. 1993. Cladistic relationships among higher groups of Heteroptera: congruence between morphological and molecular data sets. Entomol. Scand. 24: 124–138.