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Phylogeny and biogeography of southern African spoon-winged lacewings (Neuroptera: Nemopteridae: Nemopterinae)

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ABSTRACT

Nemopteridae are a charismatic family of lacewings characterised by uniquely extended hind wings. They are an ancient widespread group in the drier regions of the world. The family comprises two sub-families, Crocinae (thread-wings) and Nemopterinae (spoon- and ribbon-wings). The present distribution of the family has been largely influenced by the vicariant events of plate tectonics, resulting in relict populations in some parts of the world and extensive evolutionary radiations in others, particularly southern Africa where the vast majority of the species are endemic to the Western and Northern Cape Provinces of South Africa. This study aimed to establish the validity of the 11 currently recognised genera and infer their biogeographic history using molecular sequence data from four gene regions. The hypothesis that the Cape nemopterines co-evolved with certain taxa in the Cape Floristic Region was also tested. Phylogenetic analysis supports seven of the 11 currently recognised genera. The crown age of the Nemopterinae is estimated to be at ca. 145.6 Mya, indicating that the group has been present since the late Jurassic. Most of the genera appear to have diversified during the middle Eocene and into the middle Miocene (ca. 44–11 Mya) with recent rapid radiation of several of the genera occurring during the late Miocene (ca. 6–4.5 Mya). While these data support an initial radiation with the Rushioideae (Aizoaceae) it is recommended that further study including observations and gut content be carried out.

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1. Introduction

Nemopteridae are a charismatic family of lacewings characterised by uniquely extended hind wings. They are the only entire insect family with this innovation that has further evolved into a range of striking forms with specialised functions that include aerodynamics, camouflage, mate recognition and tactile responses (Mansell, 1996). The family comprises two subfamilies, Crocinae (thread-wings) and Nemopterinae (spoon- and ribbon-wings). Crocinae have filamentous hind wings that have a sensory function in the confined cavernicolous habitats they occupy (Mansell, 1996), while those of Nemopterinae vary from ribbon-shaped to extensive dilations that are pigmented and aerodynamically twisted to provide stability during flight as well as camouflage when at rest (Mansell, 1996). In some diurnal South African taxa additional functions of the hind wings include heat absorption when sitting on a substrate and semiotic functioning when the black and white bilobed hind wings of *Sicyoptera Navás* species are rapidly ratcheted dorsally and ventrally (Ball pers. obs.). Although these functions have been refuted (Leon and Picker, 1990b; Picker, 1984) for the species *Palmipenna aeoleoptera* Picker,

numerous field observations (Ball, Brinkman and Mansell, pers. obs) on other taxa: *Sicyoptera*, *Barbibucca* Tjeder and *Palmipenna pilicornis* Tjeder provide strong support for these functions. The ephemeral adults usually have elongated mouthparts that have evolved in response to their specialist pollenophagous diet. Larvae by contrast, are all predacious with the autapomorphy of piercing and sucking mouthparts that defines the order Neuroptera. Larval nemopterids occupy a variety of habitats ranging from small caves and rock overhangs, disused buildings and hollow tree trunks, to psammophiles and litter-dwellers, to inquilines in the nests of ants. The first crocine larva, with a bizarrely elongated prothorax, was discovered in tombs at the pyramids of Giza in Egypt (Roux, 1833), giving rise to the almost-mythological status of Nemopteridae.

Nemopterids are an ancient group of lacewings that are widespread in the drier regions of the world, with the exception of North America where the family is represented only by two fossil records (Carpenter, 1959). Nemopteridae occur in parts of Africa, particularly southern Africa, Socotra Island (1 Nemopterinae, 1 Crocinae), Australia (3 Nemopterinae, 6 Crocinae), South America (1 Nemopterinae, 6 Crocinae), Mediterranean Europe, the Middle East and India (1 Crocinae). The southern African fauna was originally monographed by Tjeder (1967), with several papers dealing with Crocinae (Mansell, 1976, 1977, 1980, 1981a,b, 1986, 1996)

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and the Nemopterinae (Leon and Picker, 1990a,b; Mansell, 1973; Picker, 1984, 1987a,b; Picker and Leon, 1990; Picker et al., 1991, 1992; Walker et al., 1994) having followed Tjeder's quintessential treatise.

There are currently 142 valid species worldwide, 43 Crocinae and 99 Nemopterinae, and at least a further 10 undescribed nemopterine species in southern Africa. The present distribution of the family appears to have been largely influenced by the vicariance events of plate tectonics, resulting in relic populations in some parts of the world and extensive evolutionary radiations in others, particularly southern Africa where 72 species, 48% of the world's Nemopteridae occur. The vast majority of these, 57 species (38% of the world fauna) are endemic to the Western and Northern Cape Provinces of South Africa (Fig. 1). The southern African Nemopterinae (excluding Crocinae) comprise 57% (62 species) of the global fauna, with 47% of the world's taxa (51 species) being endemic to these two provinces of South Africa.

The Cape nemopterines are consequently a unique and rich biological heritage that requires special research and conservation attention. While the subfamily Crocinae is comparatively well known, knowledge of the taxonomy, biology, phylogeny, local biogeography and conservation status of the Nemopterinae remains inadequate beyond that recorded by Tjeder (1967). While the conservation of Crocinae is reasonably assured owing to their arid and rocky habitats, unsuited to agriculture, the Nemopterinae are extremely vulnerable, as many of the habitats of the rare Cape endemics have already been destroyed by agricultural and urban expansion, with the remainder being severely threatened.

The South African Nemopterinae are characterised by numerous fragmented populations, with many species being known from a single locality only, and the almost clockwork precision of adult emergence at specific times of the year and, sometimes only in certain years. This has engendered the notion that they co-evolved

with the species-rich Cape flora, leading to the hypothesis that certain plant and nemopterine taxa may be interdependent (Mansell and Ball pers. obs.). Although the Crocinae are central to the evolutionary processes of the family Nemopteridae, there are no observations to indicate that their diversity in southern Africa has been influenced by flowering plant diversity to the same extent as that of the Nemopterinae. The habitats of larval crocines are also different from those of nemopterines, being confined to dusty recesses under rock overhangs, small caves and completely sheltered microhabitats, whereas most nemopterines are not confined by precise habitat requirements, where many species are psammophilous. This unrestricted habitat facilitates mass and synchronised emergence by many nemopterine taxa (Mansell and Ball pers. obs.). The focus of this paper is consequently confined to the subfamily Nemopterinae.

A detailed study of the taxonomy, with emphasis on molecular and morphological analysis, phylogeny, phylogeography, biogeography and biology of the subfamily Nemopterinae is consequently being undertaken. The first priority of these studies, and the main objective of this paper, was to establish the validity of the currently recognised 11 genera using molecular data, as this would provide the basis for investigations into their ecological role and objective criteria for the conservation of a unique South African biological heritage. The overall project is especially designed to validate the hypothesis that the Cape nemopterines co-evolved with certain taxa in the Cape flora, one of the world's six floral kingdoms (Galley and Linder, 2006; Goldblatt and Manning, 2002; Linder, 2003, 2005; Schulze et al., 2005). By combining the data from four gene regions (16S rDNA, 18S, 28S domain 2 and COI) in a total evidence approach we attempt to resolve the phylogenetic relationships of the 11 currently recognised genera. In addition, we estimate divergence times for the origin and diversification of the major lineages within the Nemopterinae.

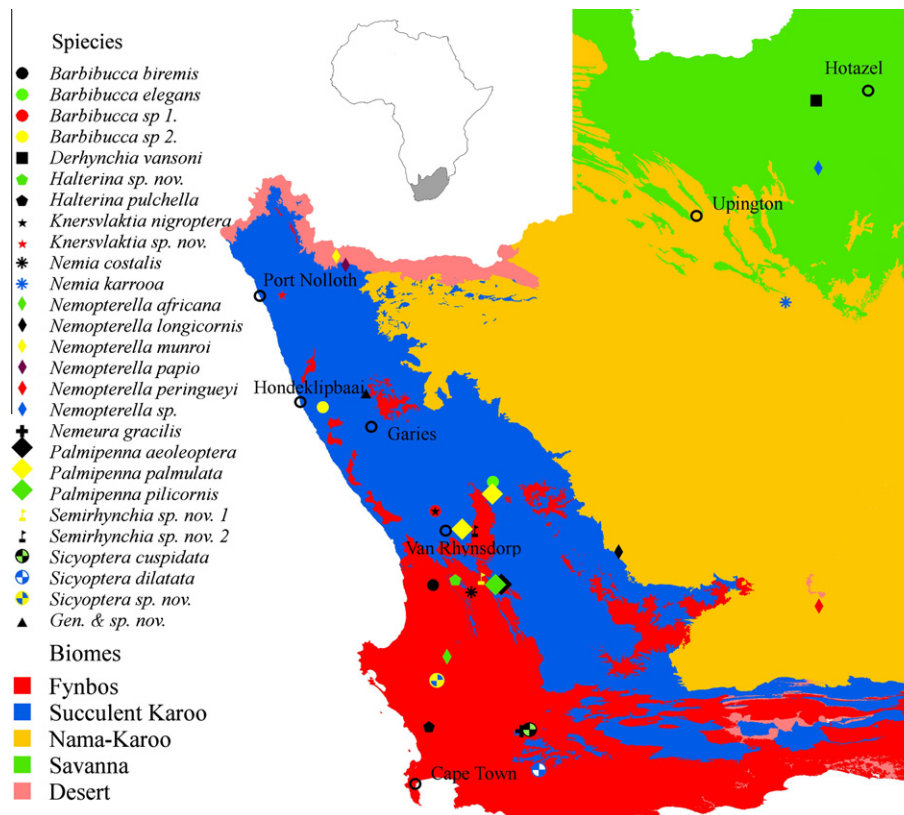


Fig. 1. Map indicating the genera and species localities of the Nemopterinae.

2. Material and methods

2.1. In-group taxa

Ten of the eleven genera South African Nemopterinae were included in this phylogenetic study: *Barbibucca*, *Derhynchia* Tjeder, *Halterina* Navás, *Knervlaktia* Picker, *Nemia* Navás, *Nemopterella* Banks, *Nemeura* Navás, *Palmipenna* Tjeder, *Semirhynchia* Tjeder and *Sicyoptera*. A single representative of the Australian Nemopterinae, the genus *Chasmoptera* Westwood, was also included. The

only genus not included in this study is *Nemopistha* Navás, a rare Savanna biome taxon.

2.2. Out-group taxa

Considerations for out-group comparisons were based on a recent phylogenetic study of the Neuropterida by Winterton et al. (2010). Based on this a species of Ascalaphidae (*Neomelambrotus molestus* Tjeder) and a representative of the subfamily Crocinae

Table 1

Sampled species for this study along with their collection locality. X – indicates amplification failed i.e. no sequence data available, Comb. – indicates the samples used in the combined analysis.

Taxon	Locality	Specimen ID	COI	16S	28S domain 2	18S	Comb.
<i>Barbibucca biremis</i>	Graafwater	BBGW01	✓	✓	✓	✓	✓
<i>Barbibucca elegans</i>	Klompbome, Loeriesfontein	BEL01	✓	✓	X	X	✓
	Klompbome, Loeriesfontein	BEL02	✓	✓	✓	✓	✓
<i>Barbibucca</i> sp.	Wallekraal	BWK01	✓	✓	✓	✓	✓
<i>Barbibucca</i> sp.	Knervlakte	BKP01	X	✓	✓	✓	✓
<i>Chasmoptera huitti</i>	Coorow, Western Australia	CS01	✓	✓	✓	✓	✓
<i>Derhynchia vansoni</i>	Tswalu–Gosa Dunes	DVT01	✓	✓	✓	✓	✓
	Tswalu–Gosa Dunes	DVT02	✓	✓	✓	✓	✓
	Tswalu–Gosa Dunes	DVT03	✓	✓	✓	✓	✓
<i>Halterina</i> sp. nov.	Zandrug 10 km north Clanwilliam	HBCZ03	✓	✓	✓	✓	✓
<i>Halterina pulchella</i>	Koeberg	HPCT01	✓	✓	✓	✓	✓
<i>Knervlaktia nigroptera</i>	Knervlakte	KNK01	✓	✓	✓	✓	✓
	Knervlakte	KNK02	✓	✓	✓	X	✓
<i>Knervlaktia</i> sp. nov.	Vyftienmylberg	BVB01	✓	✓	X	✓	✓
<i>Nemia costalis</i>	Clanwilliam	NCCD02	✓	✓	✓	✓	✓
	Clanwilliam	NCCD03	✓	✓	✓	✓	✓
	Clanwilliam	NCCD04	✓	✓	✓	✓	✓
<i>Nemia karrooa</i>	Marydale, Swartkopspan	NAMD01	✓	✓	✓	✓	✓
	Marydale, Swartkopspan	NAMD02	✓	✓	✓	✓	✓
<i>Nemopterella africana</i>	Doornfontein	NKDTK01	✓	✓	✓	✓	✓
	Doornfontein	NKDTK03	✓	✓	X	✓	✓
	Doornfontein	NKDTK04	✓	✓	✓	✓	✓
	Uitkyk, Piketberg	NAC02	✓	✓	✓	✓	✓
<i>Nemopterella longicornis</i>	Uitkyk, Piketberg	NAC03	✓	✓	✓	✓	✓
	Sterkfontein	NLTS01	✓	✓	✓	✓	✓
<i>Nemopterella munroi</i>	Sterkfontein	NLTS02	✓	✓	✓	✓	✓
	Vioolsdrif	NMSK01	✓	✓	✓	✓	✓
<i>Nemopterella papio</i>	Vioolsdrif	NMSK02	✓	✓	✓	✓	✓
	Vioolsdrif	NMSK03	✓	✓	✓	✓	✓
	Vioolsdrif	NPSK01	✓	✓	✓	✓	✓
<i>Nemopterella papio</i>	Vioolsdrif	NPSK03	✓	✓	✓	✓	✓
	Vioolsdrif	NPSK05	✓	✓	X	✓	✓
<i>Nemopterella papio</i>	Vioolsdrif	NPVD01	✓	✓	✓	✓	✓
<i>Nemopterella peringueyi</i>	Beaufort West	NBW01	✓	✓	✓	✓	✓
<i>Nemopterella</i> sp.	Tswalu Reserve	NSS01	✓	✓	✓	✓	✓
	Tswalu Reserve	NSS02	✓	✓	✓	✓	✓
<i>Nemeura gracilis</i>	Worcester	NGW01	✓	✓	✓	✓	✓
	Worcester	NGW02	✓	✓	✓	✓	✓
	Worcester	NGW03	✓	✓	✓	✓	✓
<i>Palmipenna aeoleoptera</i>	Biedou' Clanwilliam	PAC02	✓	✓	✓	✓	✓
	Biedou' Clanwilliam	PAC03	✓	✓	✓	✓	✓
	Biedou' Clanwilliam	PAC10	✓	✓	✓	✓	✓
<i>Palmipenna palmulata</i>	Brandkop	PBK01	✓	✓	✓	✓	✓
	Kobee Pass	PPKP02	✓	✓	X	✓	✓
<i>Palmipenna pilicornis</i>	Biedou' Clanwilliam	PPB02	✓	✓	X	✓	✓
	Biedou' Clanwilliam	PPB03	✓	✓	X	✓	✓
<i>Semirhynchia</i> sp. nov.	Clanwilliam	SMC02	✓	✓	X	✓	✓
<i>Semirhynchia</i> sp. nov.	Vanrhynsdorp - Kobee Pass	SV01	✓	✓	✓	✓	✓
<i>Sicyoptera cuspidata</i>	Worcester	SCBB01	✓	✓	✓	✓	✓
<i>Sicyoptera dilatata</i>	Galgeberg	SDGG01	✓	✓	X	✓	✓
	Galgeberg	SDGG02	✓	✓	✓	✓	✓
	Galgeberg	SDGG03	X	✓	✓	✓	✓
	Galgeberg	SDGB01	✓	✓	✓	✓	✓
<i>Sicyoptera</i> sp. nov.	Welbedacht	SWB01	✓	✓	✓	X	✓
	Welbedacht	SWB02	✓	✓	✓	X	✓
Gen. & sp. nov.	Kamieskroon	SSKK01	✓	✓	✓	✓	✓
Out-groups							
Ascalaphidae (<i>Neomelambrotus molestus</i>)	Klipvlei Farm	MA01	✓	✓	✓	✓	✓
Crocinae (<i>Laurhervasia setacea</i>)	Kelkiewyn Farm	LSKF01	✓	✓	✓	X	✓
Crocinae (<i>Laurhervasia setacea</i>)	Piketberg 'Uitkyk'	CPC01	✓	✓	✓	X	✓

Table 2
Summary of oligonucleotide primers used in this study.

Locus (length)	Primer name and sequence	Length	Reference
Cytochrome oxidase I	C1-J-2183 (5'CAACATTTATTTGATTTTTGG 3')	23 m	Simon et al. (1994)
	TL2-N-3014 (TCCAATGCACTAATCTGCCATATTA3')	25 m	Simon et al. (1994)
16S rRNA	LR-J-12961 (5'TTTAATCCAACATCGAGG3')	18 m	Cognato and Vogler (2001)
	LRN-N-13398 (5'CGCCTGTTTAAACAAAACAT 3')	20 m	Simon et al. (1994)
28S rRNA domain 2	D2-3551 (5'CGTGTGCTGTAGTAGCAGC3')	21 m	Gillespie et al. (2005)
	D2-4057 (5'TCAAGACGGGCTCTGAAAGT3')	20 m	Gillespie et al. (2005)
18S rRNA	18S-intfw-ST12 (5'ATCAAGAACGAAAGTTAGAG3')	20 m	Haring and Aspöck (2004)
	18S-rev1 (5'ATGGGGAACAATTGCAAGC3')	19 m	Haring and Aspöck (2004)

(Laurhervasia setacea (Klug)) were used as out-group taxa (See Table 1 for details of taxa used in this study).

2.3. Species identification

Morphological identifications were provided by a specialist on the group (M.W. Mansell) and based on material in the South African National Collection of Insects, Pretoria and in the J.B. Ball Collection, Cape Town. These two collections are the largest and most comprehensive holdings of southern African Nemopteridae currently available. Molecular analyses were based on freshly-collected specimens, authoritatively identified and further verified by comparison with material in these collections. Fresh material was collected by hand-netting and at mercury vapour light traps and preserved in absolute ethanol.

2.4. DNA extraction, cycling conditions and sequencing

Genomic DNA was extracted from a leg of at least three individuals per species representing each genus using the Roche High Pure PCR Template Preparation Kit (Roche, Penzberg, Germany) according to the manufacturer's specifications.

Sequence data were generated for four different gene regions: three ribosomal genes (16S rDNA, 18S rDNA and a portion of the nuclear rRNA large subunit 28S domain 2), along with a single protein coding gene region (cytochrome oxidase I – COI). Primer sequences used to amplify the four gene regions are listed in Table 2. Amplification using polymerase chain reaction (PCR) was performed using the following cycling conditions: a C. 456 base pair (bp) fragment of 16S rDNA was generated using the primer pair LR-N-13398 (Simon et al., 1994) and LR-J-12961 (Cognato and Vogler, 2001) and 1221 bp of COI using the primer pair C1-J-1718 and TL2-N-3014 (Simon et al., 1994) with the following protocol: initial denaturation at 95 °C (5 min); 33 cycles of 93 °C (20 s), 50 °C (40 s), 72 °C (20 s); final extension at 72 °C (5 min). A C. 836 bp of 18S rDNA was amplified using the primers 18S-intfw-ST12 and 18S-rev1 (Haring and Aspöck, 2004) with the following cycling conditions: initial denaturation at 95 °C (2 min); 30 cycles of 95 °C (10 s), 48 °C (10 s), 72 °C (1 min); final extension at 72 °C (5 min). A 'three-cycle' touchdown PCR program was used to amplify C. 735 bp stretch of 28S domain 2: initial denaturation for 20 s at 96 °C was followed by 3 cycles (15 s at 96 °C, 20 s at 54 °C, 1 min at 72 °C), thereafter 7 cycles (12 s at 96 °C, 18 s at

Table 3

Data characteristics and estimated model parameters for 16S, COI, 28S domain 2 and 18S datasets as applied to the MrBayes and *BEAST analyses (I = proportion of invariable sites).

	16S	COI	D2	18S
Best-fit model	GTR + I + G	HKY + I + G	GTR + I + G	GTR + I + G
A frequency	0.405	0.374	0.336	0.255
C frequency	0.154	0.152	0.123	0.199
G frequency	0.084	0.068	0.167	0.242
T frequency	0.358	0.406	0.373	0.305
Gamma	1.529	0.381	0.675	0.454
I	0.579	0.467	0	0.581

53 °C, 55 s at 72 °C) and 30 cycles (12 s at 96 °C, 15 s 52 °C, 50 s at 72 °C) with a final extension of 1 min at 72 °C. For all gene regions PCR was performed in a final volume of 50 µl containing approximately 50–100 ng genomic DNA template, 2.5 mM MgCl₂, 20 pmol of each primer, 10 mM dNTP's (0.25 mM of each of the four nucleotides (Promega)) and 1X buffer in the presence of one unit of *Taq* DNA polymerase (Super-Therm® DNA polymerase, Southern Cross Biotechnology).

2.5. Processing and alignment of sequences

All sequences were viewed, edited and assembled in CLC Bio 5.6 (<http://www.clcbio.com/>). Sequences for 16S, 18S and 28S domain two were subsequently aligned using the algorithm described by Löytynoja and Goldman (2005) as implemented in the Probabilistic Alignment Kit (PRANK: <http://www.ebi.ac.uk/goldMyan-srv/web-PRANK>) (Löytynoja and Goldman, 2005, 2008). Once aligned these alignments were checked manually. Alignment results showed areas of these gene regions that are conserved while others have significantly large amounts of inferred indels. Gblocks (Castresana, 2000) was used to select confidently aligned areas by eliminating the poorly aligned positions and divergent regions. The resulting alignment from Gblocks was used in subsequent analyses. The protein-coding gene COI could unambiguously be aligned displaying no stop codons when translated in MacClade version 4.03 (Maddison and Maddison, 1992). Ambiguous sites were coded using the appropriate IUB symbols after double-checking the electropherograms for recognisable sequencing artefacts. All sequences were submitted to GenBank under accession numbers JX294077–JX294294.

2.6. Phylogenetic analysis

Parsimony and Maximum Likelihood analyses were conducted using PAUP* 4.0b10 (Swofford, 2003). We used a heuristic tree search protocol with 10 random addition sequences and tree bisection and reconnection (TBR). For Parsimony we excluded all uninformative characters, gaps were treated as 5th state characters and bootstrap support values (Felsenstein, 1985) were calculated based on 1000 replicates. Starting trees for the Maximum likelihood (ML) analysis were obtained through the Neighbor-Joining (NJ) method and bootstrap support values were calculated based on 100 replicates. For ML inference a model for the entire dataset as favoured by the Akaike Information Criteria (AIC) was estimated in MrModelTest version 2.2 (Nylander, 2004). Bayesian analyses were performed in MrBayes version 3.1.2 (Huelsenbeck and Ronquist, 2001). All Bayesian analyses used the model favoured by the AIC implemented in MrModelTest see Table 3 for models of the respective gene regions. All analyses were initiated from random starting trees using one cold and three incrementally heated metropolis coupled chains (0.01) run for 10 million iterations with trees being sampled every 1000th iteration, of which 20% were discarded dur-

ing the burn-in, with the posterior probabilities being calculated from the remaining saved majority rule consensus trees. Two independently repeated Monte Carlo Markov Chain (MCMC) approximation runs were performed.

2.7. Divergence time estimates

Relaxed molecular clock estimates of divergence time were estimated using Beast version 1.6.2 (Drummond and Rambaut, 2007), a Bayesian coalescent analysis with the MCMC approximation. Nemopterinae were constrained to be monophyletic, reflecting our phylogenetic analysis, and the dataset was partitioned by gene region, with the respective substitution models (Table 3) applied to each partition. The fossil record for the Myrmeleontiforms is relatively diverse with the oldest fossils appearing to represent the stem groups of the Nymphidae, Myrmeleontidae, Ascalaphidae, Psychopsidae and Nemopteridae dating back to the Jurassic (Grimaldi and Engel, 2005). To reflect this a normal prior was applied to the root using the mid-point of the Jurassic (172 million years ago (Mya) with a standard deviation of 11) to allow for soft minimum and maximum bounds of 144 and 200 Mya, representing the upper and lower bounds of the Jurassic epoch, respectively. Two fossils from the genus *Marquetia*, morphologically considered the most primitive of the Nemopteridae, were described by Carpenter (1959), from the Eocene–Oligocene boundary (33.9 Mya) of the Florissant Shale in Colorado (Grimaldi and Engel, 2005). Based on hind wing shape the fossil genus *Marquetia*

appears morphologically similar to the extant genus *Sicyoptera* we therefore used the midpoint of 34 million years (my) as a hard minimum age constraint in an exponential prior for the node containing the genera *Nemeura*, *Sicyoptera* and *Semirhynchia*. A soft maximum constraint was applied such that 97.5% of the prior probability density would fall prior to the 200 Mya soft upper bound of the root of the tree. Priors on the ages of unconstrained nodes were derived from a birth–death tree model. Two independent Markov chains were run for 20 million iterations using a random starting tree. The program TRACER version 1.5 (Drummond and Rambaut, 2007) was used to assess the convergence between runs and posterior probabilities of the estimates.

3. Results

3.1. Dataset properties and phylogeny

The combined aligned molecular matrix consisted of 2681 base pairs (bp): 16S \approx 434 bp; 18S \approx 798 bp; domain 2 \approx 663 bp and COI = 786 bp and included 1053 parsimony informative characters. As with most arthropod genomes the A/T bias is reflected here across the four gene regions (Table 3). Parsimony analysis recovered 957 trees with a length of 3431, CI of 0.505, RI of 0.853 and RC of 0.430. A single ML tree was obtained assuming the GTR model with a gamma distribution shape parameter of 0.801 and proportion of invariable sites 0.620. The phylogram depicted in Fig. 2 is the Bayesian consensus tree with Bayesian posterior prob-

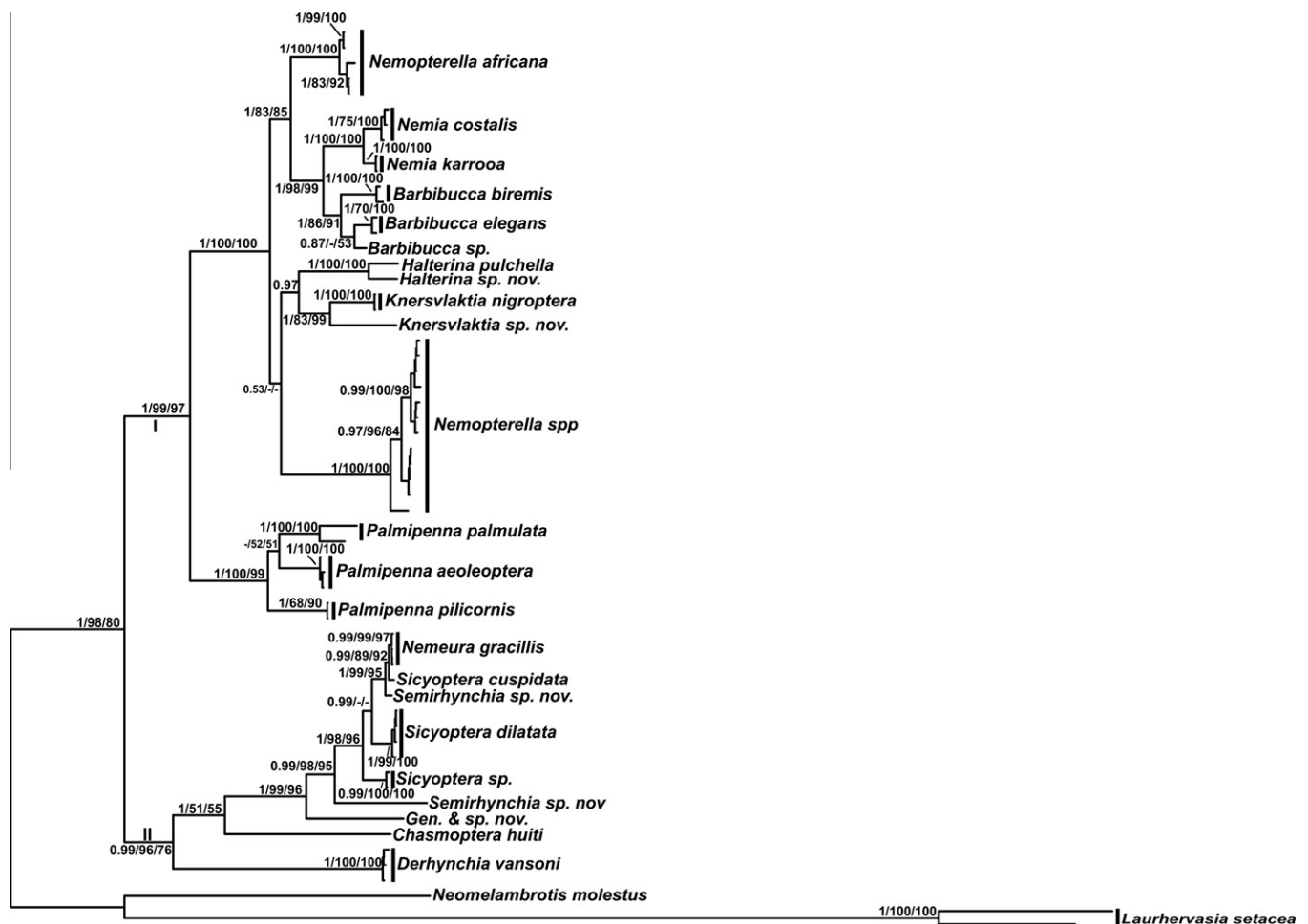


Fig. 2. Bayesian phylogram of combined dataset analysis (COI, 16S, 18S and 28S domain 2). Posterior probabilities, parsimony and ML bootstrap are given, respectively. Dashes (-) on nodes indicate weak/no support.

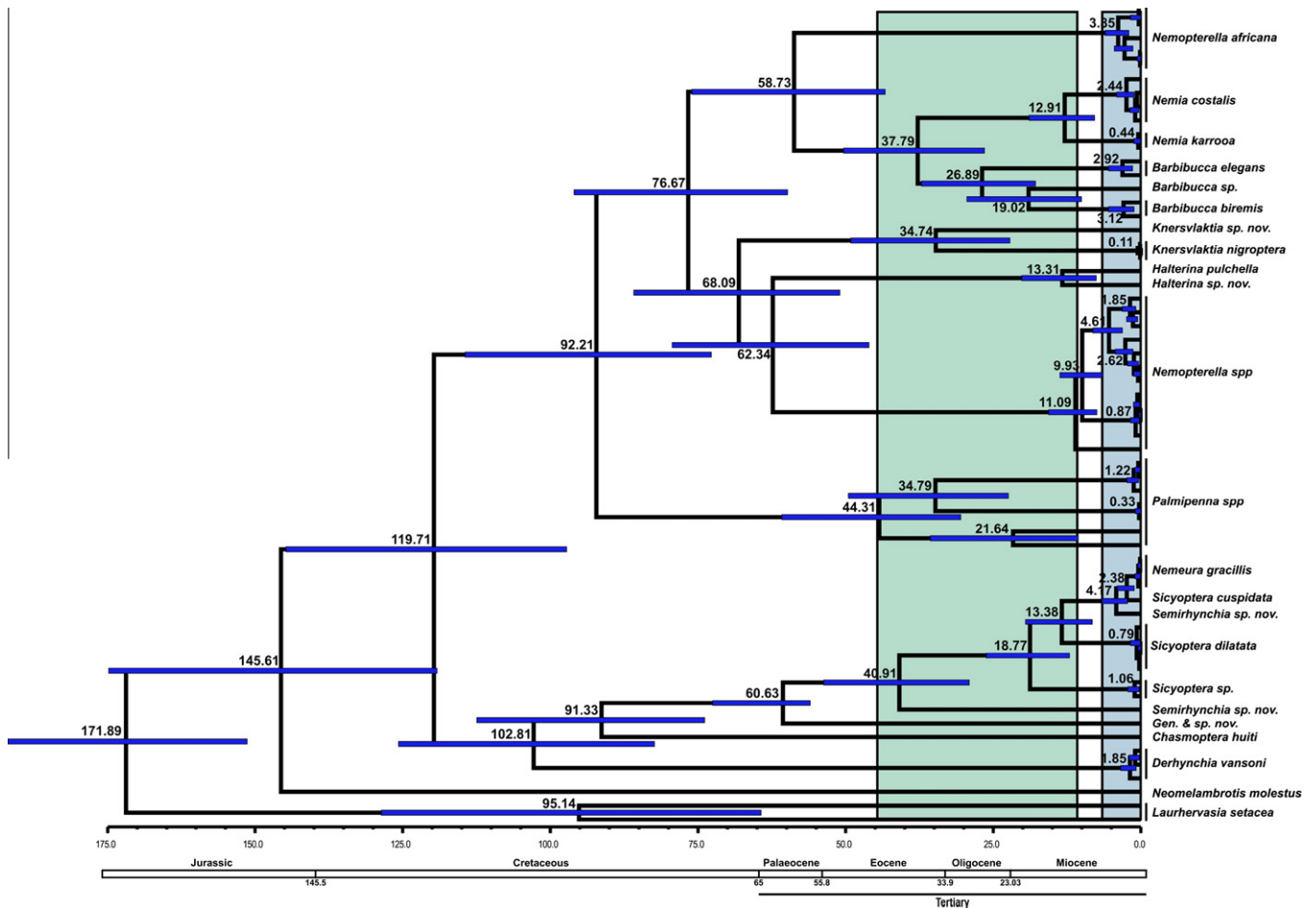


Fig. 3. Estimated times of divergence for the major lineages of the Nemopterinae. Nodes on the phylogram represent means of the probability distributions for node ages. The time intervals for the 95% probability of actual age are represented as blue bars. Green box represents the time period of slow accumulation of genera, blue box represents recent diversification. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

ability, parsimony and ML bootstrap values presented on nodes. Only values above 50% bootstrap and 0.5 posterior probability are indicated, and nodes with bootstrap support values above 70% and/or posterior probability above 0.95 are considered as strongly supported nodes.

The Nemopterinae were strongly supported as being monophyletic (Bayesian posterior probability (PP) 1.00, Parsimony bootstrap (PB) 98% and Maximum Likelihood Bootstrap (MLB) 80%). Two distinct lineages can be identified within the phylogenetic trees labelled I (1.00 PP; 99% PB; 97% MLB) and II (0.99 PP; 96% PB; 76% MLB) (Fig. 2), respectively. There are currently 11 recognised genera of Nemopterinae in southern Africa, based on morphological criteria. Within the phylogram seven of the 11 genera are well supported, based on their representative taxa: *Barbibucca* (1.00 PP; 86% PB; 91% MLB), *Derhynchia* (1.0 PP; 100% PB; 100% MLB), *Halterina* (1.0 PP; 100% PB; 100% MLB), *Knersvlaktia* (1.0 PP; 83% PB; 99% MLB), *Nemoptera* – excluding *Nemoptera africana* (1.0 PP; 100% PB; 100% MLB), *Nemia* (1.0 PP; 100% PB; 100% MLB) and *Palmipenna* (1.0 PP; 100% PB; 99% MLB). The genus *Palmipenna* appears sister to the genera *Knersvlaktia*, *Nemoptera*, *Barbibucca*, *Halterina*, and *Nemia* (1.00 PP; 99% PB; 97% MLB). The genera *Barbibucca* and *Nemia* are well supported as sister genera to each other (1.0 PP; 98% PB; 99% MLB) and are in turn sister to *Nemoptera africana* (1.0 PP; 83% PB; 85% MLB). A sister relationship between *Halterina* and *Knersvlaktia* is only supported by the Bayesian analysis (0.97 PP). Lineage II contains the genera *Nemeura*, *Sicyoptera*, *Semirhynchia* and *Chasmoptera*. Although this lineage is monophyletic and well

supported the genera *Nemeura*, *Sicyoptera* and *Semirhynchia* are polyphyletic. *Derhynchia* forms a well-supported genus sister to all the genera in lineage II, including the Australian genus *Chasmoptera*.

3.2. Divergence estimates

The blue bars in Fig. 3 indicate the 95% high posterior density (HPD) interval for each divergence. The mean estimated divergence of the Nemopterinae (including *Chasmoptera*) is ca. 145.61 million years ago (Mya), late Jurassic. The split between lineages I and II (as depicted in Fig. 2) appears to have occurred during the mid Cretaceous (ca. 119.71 Mya). The divergence of the southern African Nemopterinae appears to have occurred gradually with two distinct patterns being obvious. Most of the genera appear to have diversified gradually during the middle Eocene (ca. 44 Mya) to the middle Miocene (ca. 11 Mya) while other genera have diverged more recently during the last ca. 4.5 million years (Fig. 3).

4. Discussion

This study is the most comprehensive phylogenetic analysis yet undertaken on Nemopterinae. The overall phylogeny appeared well resolved, the Nemopterinae are monophyletic and a good indication is given as to which genera are well supported.

4.1. Phylogenetic considerations

Two major lineages are revealed by the molecular analysis, the first comprising the genera *Nemia*, *Barbibucca*, *Halterina*, *Nemopterella*, *Knervslaktia* and *Palmipenna* (lineage I – Fig. 2). The second lineage consisting of *Nemeura*, *Sicyoptera*, *Semirhynchia*, and including the exotic *Chasmoptera* shows two clear separations (*Gen. & sp. nov.*, *Semirhynchia sp. nov.* and *Chasmoptera*) and a polyphyletic complex of *Nemeura*, *Sicyoptera cuspidata* and *Semirhynchia sp. nov.*, which requires further investigation. The genera in lineages I and II are also distinguished morphologically in that the former have abdomens that are short and stout, whereas the latter have long and slender abdomens. Pleuritocavae occur sporadically in the male abdomen among the genera of lineage I, but are absent from all members of lineage II. Nocturnally active species are divided between the two lineages, *Nemopterella* and *Nemia* (lineage I) and *Semirhynchia* and *Nemeura* (lineage II). The remaining genera are diurnal a potential further adaptation to their polle-nophagous habits.

The close morphological similarity between *Nemia* and *Nemopterella* is not supported by molecular data in the phylogeny (Fig. 2). This raises the question of reliable morphological characters to distinguish these two genera. Most *Nemopterella* species, which are morphologically difficult to separate from one another, are superficially distinct from *Nemia*, although these morphological distinctions are tenuous – requiring further study. Navás (1915) divided the genus *Nemopterella* into two, *Nemeva*, with type species *Nemopteryx africana* Leach, 1815 and *Nemia*, with type species *Nemoptera costalis* Westwood, 1836. According to Tjeder (1967) this separation was based on inconsistent characters of no taxonomic significance, but he did discover an important feature, the presence of pleuritocavae in the male of *N. africana* that were absent from that of *N. costalis*, and he separated the two genera on that clear basis. Tjeder (1967) then synonymised *Nemeva* Navás with *Nemopterella*, as a valid existing name (*Nemopterella*) cannot be substituted by a new name with the same type species. However, further species recently discovered in South Africa that could be assigned to *Nemia* have pleuritocavae thereby casting doubt upon their value in distinguishing *Nemia* and *Nemopterella*. *Nemopterella africana* is indicated as distinct from both *Nemia* and *Nemopterella* although it shares the important distinguishing features of both genera (pleuritocavae as in *Nemopterella* and the characteristic body patterns of *Nemia*). It consequently suggests that *N. africana* may represent a previously undetected monotypic genus and also that the main distinguishing feature currently separating *Nemia* and *Nemopterella* (pleuritocavae) may not be supported by molecular data.

The genus *Barbibucca* is morphologically distinctive in that they are robust with uniformly broad hind wings. The genus *Knervslaktia* has been distinguished from other genera, and is clearly underpinned by molecular data. The morphologically distinctive diurnal genus *Palmipenna* has several distinguishing characters: antennae short and stout less than half forewing length, hind wings less than twice forewing length with broad apical dilations and very small eyes characteristic of diurnal taxa. Molecular data unequivocally support *Palmipenna* as a valid genus. The genus *Halterina* is also morphologically distinct, and its two species are supported by molecular data.

Derhynchia is a distinct monotypic genus supported by the autapomorphy of reduced mouthparts and rostrum, the only nemopterine with this unique feature. The biology of this genus is also unique in that it inhabits the dunes in the Kalahari ecosystem where its free-living psammophilous larva lives several centimetres under the sand surface near vegetation. The larva also has several unique features including vestigial eyes and peculiar burrowing behaviour, indicating that the entire larval life is spent

underground (Mansell, 1973). The most readily available food source for adults is pollen from dune grasses that do not require a long rostrum for harvesting, probably leading to the secondarily atrophied mouthparts.

In lineage II, *Nemeura*, *Semirhynchia* and *Sicyoptera* are morphologically distinct from one another. *Semirhynchia* has distinctive short mouthparts and ribbon-shaped wings, while *Sicyoptera* is easily distinguished from *Nemeura* by the broad double pre-apical expansions of the hind wings. The value of the hind wing shape in distinguishing genera is however, questionable as broad hind wings occur over a wide range of taxa, especially those in this study (lineages I and II). Furthermore, a newly discovered taxon (*Gen. & sp. nov.*) which also has broad double pre-apical hind wing expansion, and closely resembles species of *Sicyoptera*, separates out from other genera within this group, and is furthermore distinguished by forewing characteristics supporting its molecular distinction.

The molecular analysis applied in this study has led to three important conclusions. It has shown clear support for at least seven of the southern African genera currently based on morphological criteria, it has indicated that further studies, both morphological and molecular are required on the remaining three genera, and it has also indicated the correct generic placement for taxa that were previously doubtful. It has furthermore, revealed two distinct major lineages, with lineage II apparently more closely related to the Australian *Chasmoptera*, possibly suggesting that it may comprise an ancestral lineage of the Nemopterinae. The family is clearly a Gondwanan element with the sporadic relic distribution on the southern continents being due to plate tectonics. Only one species, *Stenorhachus walkeri* (McLachlan) remains in South America, while one genus, *Chasmoptera* with three described species (*C. hutti* (Westwood), *C. mathewsi* Koch and *C. superba* Tillyard all confined to Western Australia near Perth, is the only representative of the Nemopterinae in the Australasian region. The fact that genera in lineage II are most closely related to *Chasmoptera* suggests that they are part of the lineage that emerged before South America and Australia became separated, and before the major radiation took place on the southern African fragment. *Chasmoptera* is also morphologically very similar to *Sicyoptera*, and to the two fossil Nemopterinae, *Marquettia americana* (Cockerell) and *M. metzeli* (Pierce and Kirkby), from North America, as well as to some species of the Palaearctic genus, *Lertha* Navás, suggesting that the double pre-apical dilation of the hind wing may be part of the “ground-plan” of the Nemopterinae that has been retained by widely disparate genera, including the enigmatic *Parasicyoptera guichardi* Tjeder, known only from Socotra Island.

4.2. Biogeographic inferences and speciation events

The Cape Floristic Region (CFR), one of six floral kingdoms (Galley and Linder, 2006; Goldblatt and Manning, 2002; Linder, 2003, 2005; Schulze et al., 2005) and its extent encompasses the bulk of the distribution of the South African nemopterines (Fig. 1). The biomes in which South African Nemopterinae occur are Fynbos, Succulent Karoo, Nama Karoo, Albany Thicket and Savanna biomes (Mucina and Rutherford, 2006). Recent detailed paleoclimatic records indicate that large fluctuations in the African climate have caused changes, among others, in the topology, ecology and soil which, in turn, has had a major effect on the speciation, extinction and dispersal of the flora and fauna over the last 5–6 Mya (deMenocal, 2004). These landscape type changes result in the island-like fragmentation of a habitat, which in turn may act as speciation ‘hotspots’ (Linder, 2003). One of the major driving forces behind the floral diversification within CFR is thought to be the Plio-Pleistocene glacial and interglacial cycles and the onset of the winter rainfall regime, having occurred

ca. 5 Mya (deMenocal, 1995, 2004; Linder, 2003) with African bovids, birds and hominids having diversified over the last ca. 2–3 Mya (deMenocal, 2004; Linder, 2003; Potts, 1998). In particular, the increase in aridity that is accentuated by the rain shadow created by the Great Escarpment and reinforced by the cold upwelling of the Benguela current system and the subtropical anticyclone (Late Miocene) is thought to have played a major role in the CFR diversification (deMenocal, 2004; Linder, 2003; van Zinderen Bakker, 1975; Ward et al., 1983). It has been suggested that allopatric faunal distribution may have therefore been driven by distributional changes and fragmentation of the flora as a result of climatic oscillations (Price et al., 2007; Tolley et al., 2006).

The flora of the CFR *sensu lato* can be divided into three elements: (1) a Succulent Karoo element that has its main diversity in the northern part of the CFR e.g. *Mesembryanthemum sensu stricto* (2) a Tropical African element in which the main diversity occurs to the east of the CFR e.g. *Rhus*, *Aloe* and many genera typical of ticket and forest vegetation and, (3) the so-called 'Cape clades' occurring in the South Western Region (Bolus, 1886) that all have their species richness centred within the CFR e.g. *Erica*, Proteaceae, Bruniaceae, Restionaceae *p.p.*, and *Phyllica*.

It is hypothesised here that the Nemopterinae have co-evolved mainly with Ruschioideae (Aizoaceae). The Aizoaceae have their greatest diversity in the summer-dry west and reach into Namaqualand up to the Orange River (Linder, 2003). The sub-family Ruschioideae contains approximately 1600 species almost exclusively endemic to southern Africa, are the most speciose sub-family within the Aizoaceae and dominate the Namaqualand in terms of species numbers and density. The core Ruschioideae radiation driven by the onset of the winter rainfall regime was estimated to have occurred c. 3.8–7.8 Ma (Klak et al., 2004).

Two hypotheses exist regarding the high species diversity of the CFR, either the flora and fauna of the CFR have undergone rapid diversification or there has been a slow accumulation of species over time (Galley et al., 2007; Linder, 2003). When looking more closely at the climatic changes since the early Oligocene, the west appears to have undergone a gradual change in climate (Linder, 2005), arid to semi-arid conditions interspersed with wetter periods (Klak et al., 2004), indicating that the two hypotheses are not necessarily mutually exclusive. A wide range of postulated dates exists for the radiation of the Cape Flora, those dating back to the arid phase in the early Oligocene indicate a slow accumulation of diversity, while others show typical recent rapid radiations (Linder, 2005). The crown age of the Nemopterinae is estimated to be at ca. 119.7 Mya (144.8–97.2; upper and lower estimates, respectively), indicating that the group has been present since the Cretaceous. Most of the genera appear to have diversified during the middle Eocene and into the middle Miocene (ca. 44–11 Mya, Fig. 3) with recent rapid divergence of several of the genera, *Barbibucca*, *Nemia*, *Nemoptera*, *Nemeura*, *Palmipenna* and *Sicyoptera/Semirhynchia*, occurring during the late Miocene (ca. 4.5 Mya), alluding to the fact that the diversification of the Nemopterinae appears to support both these hypotheses. The timing of the recent diversification events seems to follow the recent speciation of the Ruschioideae, indicating that an adaptive shift of the nemopterines may have occurred in response to the Ruschioideae distribution shifts linked to climatic oscillations. Further empirical evidence however is needed to test whether this is an adaptive radiation. Present observations, noted on radially symmetrical flowers, of adult nemopterine pollen feeding include the plant families Aizoaceae, Asteraceae and Molluginaceae (Ball pers. obs.). Feeding behaviour of *Nemoptera sinuata* Olivier in the Balkan-Anatolian peninsula was noted on flowers from the families Asteraceae and Brassicaceae (Krenn et al., 2008). It is therefore possible that the initial diversification of Nemopterinae was influenced by the radiation of the ruschoids, and that they subsequently adapted to feeding on other plant species as well.

This is a hypothesis that will be tested in subsequent studies on pollen composition in the alimentary canals of preserved and freshly-collected specimens, as well as field observations. The greatest extant concentration of nemopterines is in the more arid portions of the Western and Northern Cape Provinces, where a variety of families of ephemeral spring flowers are visited by a large number of different orders of insects. Present local observations (Ball pers. obs.) confirm that the community patterns of plant-pollinator interactions reveal that angiosperm species are typically visited by many taxa of potential pollinators and that the majority of flower visitors are noted on multiple plant species (Geber and Moeller, 2006; Waser and Ollerton, 2006).

Apart from the radiation in South Africa, there has also been a similar, albeit smaller, radiation of a single genus, *Chasmoptera* in Western Australia where three species occur. In the southern Palaeoartic there has been a radiation of two genera, *Nemoptera* Latrielle and *Lertha*. Both of these areas share a Mediterranean climate similar to that of the Western Cape Province of South Africa. The South African genus, *Sicyoptera*, most closely resembles *Chasmoptera* and *Lertha* and is restricted to the fynbos component in the south western part of the Western Cape Province, whereas most of the recent radiation of genera has taken place along the central and northern parts of the west coast area, where the largest simultaneous radiation of Ruschioideae has also taken place (Klak et al., 2004).

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