

Molecular phylogeny and systematics of the Pieridae (Lepidoptera: Papilionoidea): higher classification and biogeography

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The systematic relationships of the butterfly family Pieridae are poorly understood. Much of our current understanding is based primarily on detailed morphological observations made 50–70 years ago. However, the family and its putative four subfamilies and two tribes, have rarely been subjected to rigorous phylogenetic analysis. Here we present results based on an analysis of molecular characters used to reconstruct the phylogeny of the Pieridae in order to infer higher-level classification above the generic level and patterns of historical biogeography. Our sample contained 90 taxa representing 74 genera and six subgenera, or 89% of all genera recognized in the family. Three complementary approaches were employed: (1) a combined analysis of a 30 taxon subset for sequences from four gene regions, including elongation factor-1 alpha (*EF-1α*), *wingless*, cytochrome oxidase subunit I (*COI*), and *28S* (3675 bp, 1031 parsimony-informative characters), mainly to establish higher-level relationships, (2) a single-gene analysis of the 90 taxon data set for sequences from *EF-1α* (1066 bp, 364 parsimony-informative characters), mainly to establish lower-level relationships, and (3) an all available data analysis of the entire data set for sequences from the four genes, to recover both deep and shallow nodes. Analyses using maximum parsimony, maximum likelihood and Bayesian inference provided similar results. All supported monophyly for the four subfamilies but not for the two tribes, with the Anthocharidini polyphyletic and the Pierini paraphyletic. The combined and all available data analyses support the following relationships among the subfamilies: ((Pseudopontiinae + Dismorphiinae) + (Coliadinae + Pierinae)), corroborating Ehrlich's 1958 phenetic hypothesis. On the basis of these analyses, and additional morphological and life history evidence, we propose a reclassification of the subfamily Pierinae into two tribes (Anthocharidini s.s., Pierini s.s.) and two informal groups (*Colotis* group, *Leptosia*), with the tribe Pierini s.s. subdivided into three subtribes (Appiadina, Pierina, Aporiina) and three genera (*Elodina*, *Dixeia*, *Belenois*) of uncertain status (*incertae sedis*). The combined and all available data analyses support the following relationships among the Pierinae: (*Colotis* group + Anthocharidini s.s. + *Leptosia* + (*Elodina* + ((*Dixeia* + *Belenois*) + Appiadina + Pierina + Aporiina))). Application of a molecular clock calibrated using fossil evidence and semiparametric rate smoothing suggests that divergence between the Pierina and Aporiina occurred no later than the Palaeocene (> 60 Myr). The minimum estimate for the age of the crown-group of the Pieridae was 112–82 Myr, with a mean of 95 Myr. A historical biogeographical hypothesis is proposed to explain the present-day distribution of the clade Pseudopontiinae + Dismorphiinae, which argues for an origin of the two subfamilies in western Gondwana (Africa + South America) during the Late Cretaceous. © 2006 The Linnean Society of London, *Zoological Journal of the Linnean Society*, 2006, 147, 239–275.

ADDITIONAL KEYWORDS: Coliadinae – Cretaceous – Dismorphiinae – Dispersal – Gondwana – Pierinae – Pseudopontiinae – Vicariance.

INTRODUCTION

The Pieridae are among the most poorly understood butterfly families within the Papilionoidea in terms of their higher-level systematics and classification. Indeed, almost 20 years ago Robbins & Henson (1986)

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emphasized that ‘... there is glaring need for a worldwide treatment of the pierines’. Yet pierids have played an important role in evolutionary studies (e.g. Courtney, 1986; Watt, Donohue & Carter, 1996; Brunton, 1998; Stavenga *et al.*, 2004; Kemp, Rutowski & Mendoza, 2005) and include species of major economic significance, such as the cabbage whites (*Pieris*) and sulphurs (*Colias*). Unlike several other families, or subfamilies/tribes within other families, such as the Papilionidae (Miller, 1987; Caterino *et al.*, 2001; Braby, Trueman & Eastwood, 2005), Nymphalidae (Ackery & Vane-Wright, 1984; Brower, 2000; Penz & Peggie, 2003; Wahlberg, Weingartner & Nylin, 2003; Freitas & Brown, 2004) and Riodinidae (Harvey, 1987; Campbell, Brower & Pierce, 2000), the Pieridae and its putative subfamilies and tribes have rarely been subjected to rigorous phylogenetic analyses of morphological or molecular data. Although Janz & Nylin (1998; pers. comm) published a phylogeny of the Pieridae based on 39 terminal taxa (with five taxa each comprising the combination of two genera), their cladistic analysis of previously published morphological data dealt with only two subfamilies (Coliadinae and Pierinae) and provided little resolution within the major clades.

Although the monophyly of the Pieridae is well established (Kristensen, 1976; de Jong, Vane-Wright & Ackery, 1996; Ackery, de Jong & Vane-Wright, 1999; Wahlberg *et al.*, 2005), the phylogenetic position of pierids in relation to the other butterfly families is uncertain (Robbins, 1988; Vane-Wright, 2003; Wahlberg *et al.*, 2005). The Pieridae are considered to be either the sister family to the Papilionidae (Ehrlich, 1958; Scott, 1985), or, more probably, sister to Nymphalidae + (Riodinidae + Lycaenidae) (Kristensen, 1976; de Jong *et al.*, 1996; Weller, Pashley & Martin, 1996; Ackery *et al.*, 1999; Wahlberg *et al.*, 2005). However, unlike some other families (e.g. Nymphalidae, Lycaenidae), the integrity of the Pieridae as a natural group has never been in dispute. Synapomorphies supporting monophyly include wing scales with pterin pigments, foretarsus with distinctly bifid claws, outer edge of forewing third axillary with tooth, and lateral plates of pronotum not fused medially (Ackery *et al.*, 1999; Vane-Wright, 2003). The family is worldwide in distribution, and contains approximately 1100 species (Robbins, 1982; Ackery *et al.*, 1999; Vane-Wright, 2003) currently arranged in 98 lower taxa (83 genera, 15 subgenera) (Braby, 2005). The two most speciose genera are *Delias* Hübner and *Catasticta* Butler, but putative radiations have also occurred in *Colias* Fabricius, *Eurema* Hübner, *Colotis* Hübner, *Mylothris* Hübner and the *Tatochila* Butler group of genera. The adult butterflies are of medium size, typically white, orange or yellow in colour, and the pupal morphology is highly distinctive (Chapman,

1895; Talbot, 1939; Mosher, 1969). Many species migrate and/or exhibit seasonal phenotypic variation.

Much of our current understanding of the higher classification and interrelationships of the Pieridae has been based on detailed morphological work conducted 50–70 years ago (Klots, 1933; Ehrlich, 1958). The family is currently arranged in four subfamilies (Pseudopontiinae, Dismorphiinae, Coliadinae, Pierinae), with the Pierinae usually divided into two tribes (Pierini, Anthocharidini) (Ackery, 1984; Bridges, 1988; de Jong *et al.*, 1996; Ackery *et al.*, 1999; Vane-Wright, 2003). The Pseudopontiinae are monotypic, containing the single monobasic genus *Pseudopontia* Plötz from central and western Africa. The Dismorphiinae are relatively small, comprising approximately 60 species in seven genera and, with the exception of the single disjunct genus *Leptidea* Billberg in the Palaearctic, are found predominantly in South America, with a smaller representation in Central America. The Coliadinae comprise approximately 220 species in 18 genera, and are cosmopolitan, although the greater proportion of species occurs in tropical latitudes. The Pierinae, also cosmopolitan, are by far the largest subfamily, containing approximately 840 species in 57 genera (Ackery *et al.*, 1999; Braby, 2005), and thus make up between two-thirds and three-quarters of the total species and generic diversity of the family.

Although the four subfamilies have remained relatively stable in terms of their composition, considerable uncertainties exist in the systematics and phylogenetic relationships among the higher taxa (Fig. 1). Klots (1933), building on his own earlier work (Klots, 1929) as well as that of Butler (1870), Scudder (1875b), Dixey (1894, 1932), Grote (1900), Röber (1908–09), and Aurivillius (1910) among others, recognized three subfamilies, with one of these, the Pierinae, consisting of three tribes (Euchloini, Rhodocerini, Pierini). The Rhodocerini and Euchloini have since proven to be subjective synonyms of the Coliadini (Talbot, 1935) and Anthocharidini (Bridges, 1988), respectively. Klots’ intuitive phylogeny (Fig. 1A) showed that the Dismorphiinae and Pseudopontiinae were closely related and formed the sister group to the Pierinae. Clench (1955) followed Klots and recognized the same three subfamilies, but noted that the Pseudopontiinae were ‘intermediate’ between the two other subfamilies.

Ehrlich’s (1958) phenetic tree (Fig. 1B) was similar to that of Klots, except that the tribe Coliadini was treated as a distinct subfamily, the Coliadinae, in accordance with Talbot (1935), and phylogenetically removed from, and sister to, the Pierinae. Scott (1985) reached the same conclusion as Ehrlich (1958) with regard to the classification and relationships of the pierid subfamilies. In both Ehrlich’s and Scott’s classifications, the Pierinae were not further subdivided

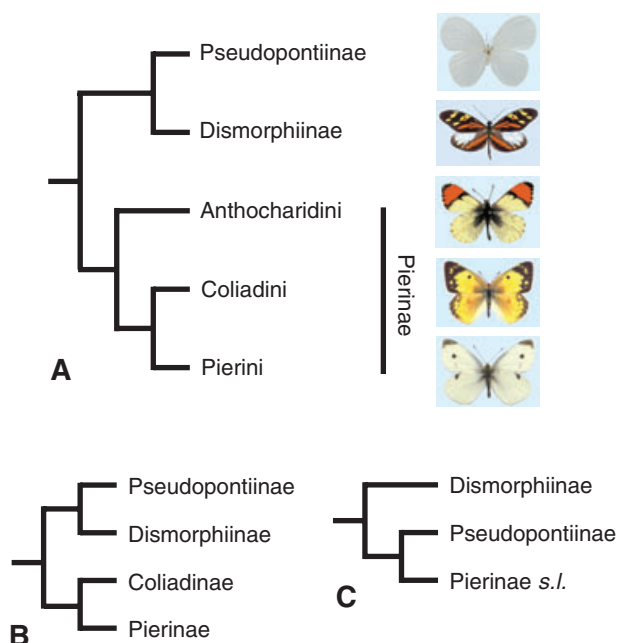


Figure 1. Three different phylogenetic hypotheses of the higher classification of the Pieridae. A, Klots (1933). B, Ehrlich (1958). C, Venables (1993).

into tribes. The only other broad-based study of the higher classification of the Pieridae is the work of Venables (1993), who made a preliminary cladistic analysis of a larger data set (43 genera) and incorporated Klots' morphological characters. Her cladogram showed that the Coliadinae were paraphyletic, whereas the Pierinae were largely monophyletic except they contained the coliadine taxon *Nathalis* Boisduval. On this basis, Venables (1993) tentatively subsumed the Coliadinae within the Pierinae so that her classification of higher taxa was essentially similar to that of Klots (1933) and Clench (1955), except that the Dismorphiinae were sister to Pseudopontiinae + Pierinae *s.l.* (Fig. 1C).

Several other studies have dealt with the higher systematics of the Pieridae, but these are more limited in scope, as only small numbers of taxa or characters were analysed. Ehrlich & Ehrlich (1967) analysed five taxa within their broader phenetic study of the Papilionoidea. The relationships among pierid subfamilies were found to be variable, and the family grouped inconsistently with the Papilionidae. Geiger (1981) studied phenetic relationships among 24 European taxa using enzyme electrophoretic data. His results showed a clear biochemical distinction between the subfamilies Dismorphiinae, Coliadinae, and Pierinae. However, separation of the tribes Pierini and Anthocharidini within the Pierinae was much weaker. In a study of butterflies and their host plants, Janz &

Nylin (1998) published a simplified version of the phylogeny of the Papilionoidea, based on the data sets of Ehrlich & Ehrlich (1967) and Geiger (1981) for the Pieridae. Their cladistic analysis of 39 terminal taxa in the Coliadinae and Pierinae recovered the two subfamilies as reciprocally monophyletic, although their study did not include *Pseudopontia*, and the Dismorphiinae were used as a single outgroup taxon. Cheong (1990) studied the female genitalia from 90 species representing 23 genera. However, too few independent morphological characters (a total of 15) were available to infer phylogenetic relationships. Lukhtanov (1991) studied chromosome relationships and noted that the Dismorphiinae, Coliadinae, and Anthocharidini, but not Pierini, all had the same basic number ($n = 31$). de Jong *et al.* (1996) and Ackery *et al.* (1999) included seven exemplar pierid species in their higher-level cladistic analysis of morphological characters of the butterflies. They provisionally maintained the four subfamilies but noted that relationships between them were uncertain. The Dismorphiinae (*Dismorphia*) were sister to the six other species, but the Pierinae appeared to be paraphyletic as the exemplar genera (*Pieris*, *Delias*, *Euchloe*) rarely grouped together and the subfamily included both the Coliadinae (*Eurema* + *Colias*) and Pseudopontiinae as subordinate taxa. Moreover, they were unable to find any convincing synapomorphies for either the Coliadinae or the Pierinae. Pollock *et al.* (1998) sequenced a small fragment of mitochondrial cytochrome oxidase subunit I (*COI*) and ribosomal *12S* + *16S* genes for 21 taxa (mainly *Colias*) representing eight pierid genera. In contrast to de Jong *et al.* (1996), their molecular phylogenetic analysis recovered the Coliadinae (four genera) as a strongly supported monophyletic group and sister to the four other genera (*Anthocharis*, *Euchloe*, *Pieris*, *Pontia*). Similarly, T. Yamauchi, O. Yata & A. Venables (unpubl. data), conducted a phylogenetic analysis of adult morphological characters representing all genera and also recovered the Coliadinae as a monophyletic group.

Part of the uncertainty and lack of agreement among workers in the interpretation of phylogenetic relationships and systematic status of higher taxa of the Pieridae may lie in the fact that Klots' (1933) original systematic classification and ideas of evolutionary relatedness were incongruent with one another. In Figure 2, we have attempted to reconstruct Klots' intuitive phylogeny as a cladogram, according to his generic revision and systematic framework, and hypothetical chart of evolution of the higher taxa (subfamilies, tribes, and generic groups). Klots proposed several 'natural' groupings based on the examination of numerous morphological characters of the male genitalia and wing venation. However, he expressed considerable uncertainty about the placement of 12

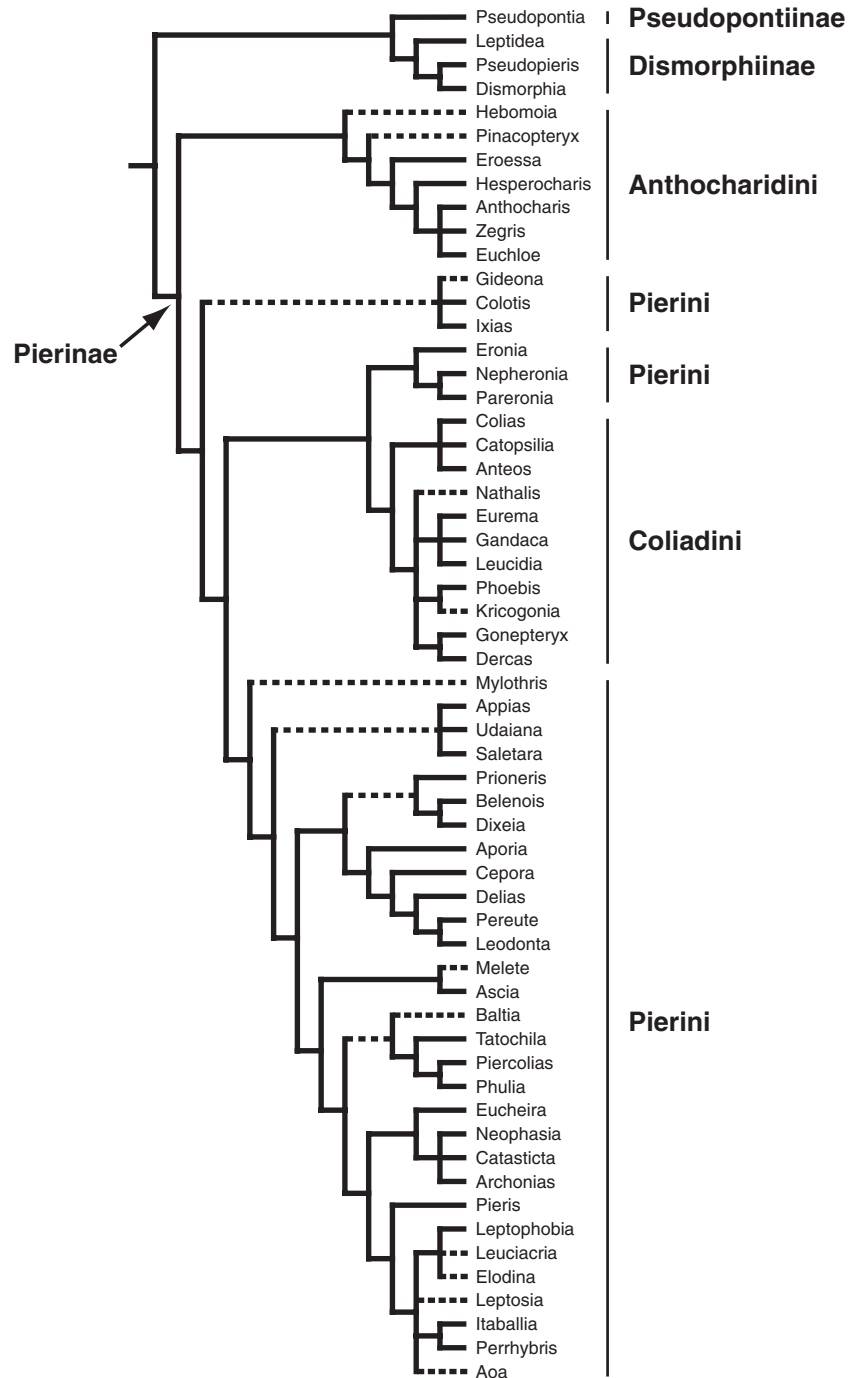


Figure 2. Klots' (1933) intuitive phylogeny of the Pieridae, reconstructed from his generic revision and systematic classification, and hypothetical chart of evolution of the subfamilies and main stock of the Pierinae. Dashed lines indicate uncertainty in the phylogenetic position of genera or groups of genera.

genera, namely *Hebomoia* Hübner and *Pinacopteryx* Wallengren in the Anthocharidini; *Nathalis* and *Kricogonia* Reakirt in the Coliadini; and *Gideona* Klots, *Mylothris*, *Melete* Swainson, *Baltia* Moore, *Leuciactria* Rothschild & Jordan, *Elodina* Felder & Felder,

Leptosia Hübner and *Aoa* de Nicéville in the Pierini. Moreover, the Pierini were not envisaged as a monophyletic entity. Klots (1933) regarded the genera *Colotis* and *Ixias* Hübner to be 'derived' from the Anthocharidini in an evolutionary sense, but nonethe-

less classified them with the Pierini; he also considered the *Eronia* group of genera (*Eronia*, *Nepheronia*, *Pareronia*) to be more closely related to the Coliadini than to the Pierini, in which he placed them.

The goal of this study was to reconstruct the phylogeny of the Pieridae using molecular characters and exemplar species representing nearly all of the currently recognized lower taxa (genera, subgenera). We employed three complementary approaches to investigate the monophyly and relationships of the extant taxa in the family: (1) a combined analysis of fragments of four genes, namely nuclear elongation factor-1 alpha (*EF-1 α*), nuclear *wingless*, mitochondrial *COI*, and ribosomal 28S (*28S*), of a 30 taxon data set to establish higher-level phylogenetic patterns at deeper nodes (subfamilies, tribes); (2) a single-gene (*EF-1 α*) analysis of a 90 taxon data set to infer lower-level phylogenetic patterns at more shallow nodes (genera, subgenera); and (3) an all available data analysis of the entire 90 taxon data set using all four genes to recover both deep and shallow nodes. The phylogenetic hypothesis based on the combined and all available data analyses was then used as a framework to revise the higher classification of the family and to explore patterns of historical biogeography. We also estimated the age of divergence events calibrated with fossil evidence for the *EF-1 α* analysis.

MATERIAL AND METHODS

MOLECULAR MARKERS

Currently, there are relatively few genes available for reconstructing arthropod divergences of Mesozoic age (Caterino, Cho & Sperling, 2000). *EF-1 α* is a nuclear protein-encoding gene involved in the translation of mRNA to protein, specifically the binding of aminoacyl-tRNA to the ribosome (Kamie *et al.*, 1993; Palumbi, 1996). It evolves relatively slowly, insertion/deletions are absent and it provides relatively unambiguous alignment (Cho *et al.*, 1995; Caterino *et al.*, 2000; Sperling, 2003). For Lepidoptera that have been studied, most of the phylogenetic information lies in the third codon position, most substitutions are synonymous, pairwise differences between closely related taxa are small, and saturation levels (transversion/transition ratios) tend to be low (Cho *et al.*, 1995; Mitchell *et al.*, 1997; Roger *et al.*, 1999). Because first and second positions are highly conserved, nonsynonymous changes are rare, but third positions frequently show saturation. These properties render the gene as a useful marker for resolving the more recent divergence events (e.g. mid-Tertiary) of insects, especially Lepidoptera, which have lost all introns and which have only a single copy of the gene (i.e. there are no paralogous copies) (Cho *et al.*, 1995; Danforth & Shu-

qing, 1998). In this group, *EF-1 α* has proven useful in reconstructing phylogenies at the 'intermediate' systematic levels, such as genus and tribe (Cho *et al.*, 1995; Mitchell *et al.*, 1997; Mitchell, Mitter & Regier, 2000; Friedlander *et al.*, 1998; Reed & Sperling, 1999; Caterino *et al.*, 2001; Monteiro & Pierce, 2001; Morinaka, Miyata & Tanaka, 2002; Wahlberg *et al.*, 2003).

Wingless is another protein-encoding gene involved in wing pattern formation. It belongs to the *wnt* gene family, whose paralogs are easily distinguishable, and shows a relatively rapid rate of substitution. In Lepidoptera, it has been used successfully for resolving phylogenetic relationships at both higher and lower systematic levels (Brower & Egan, 1997; Brower & DeSalle, 1998; Brower, 2000; Campbell *et al.*, 2000; Wahlberg *et al.*, 2003).

COI is a widely used mitochondrial protein-encoding gene. Because it is faster evolving, third codon positions quickly become saturated at the deeper levels of divergence, but it is relatively conserved compared with other mitochondrial genes (e.g. Simon *et al.*, 1994; Hillis *et al.*, 1996a; Palumbi, 1996). In molecular phylogenetic studies of Lepidoptera it has shown great utility for resolving shallow (recent) divergence events (Caterino *et al.*, 2000; Sperling, 2003).

The rDNA 28S marker has been used successfully for reconstructing phylogenetic relationships among many invertebrate taxa (Caterino *et al.*, 2000). In Lepidoptera, the unambiguously aligned regions are highly conserved, rendering the gene especially useful for recovering deeper (old) divergence levels (Weller *et al.*, 1992, 1994, 1996; Weller & Pashley, 1995).

Because the four genes evolve at different rates, combining all of them will probably increase the phylogenetic estimation and resolution of most, if not all, nodes, provided the data partitions are congruent (Caterino *et al.*, 2000). In Lepidoptera, several recent studies have demonstrated improved resolution based on standard measures of nodal support at both deep and shallow levels of divergence in a combined analysis of nuclear, mitochondrial or ribosomal genes (Monteiro & Pierce, 2001; Caterino *et al.*, 2001; Wahlberg *et al.*, 2003; Kandul *et al.*, 2004; Zakharov, Caterino & Sperling, 2004).

TAXON SAMPLING

For the combined analysis, 26 exemplars (Table 1) were sampled, representing the entire systematic and phylogenetic diversity of the family, based on previous systematic hypotheses and the results from the *EF-1 α* larger taxon data set. This data set also included four taxa (*Leptosia*, *Elodina*, *Dixeia*, *Belenois*) of uncertain status. For the single-gene (*EF-1 α*) analysis, 86 exemplar species of Pieridae (Table 1) were sampled from

Table 1. Exemplar taxa used in this study, with collection localities and GenBank accession numbers. Voucher numbers refer to those specimens deposited in the Museum of Comparative Zoology (MCZ), Harvard University, USA. The nomenclature of pierid taxa follows Braby (2005)

Taxon	MCZ voucher no.	Locality	GenBank accession no.			
			<i>EF-1α</i>	<i>wingless</i>	<i>COI</i>	28S
Pieridae						
<i>Anteos clorinde</i>	MFB-00-P188	COSTA RICA. La Guacima	AY870570			
<i>Anthocharis belia</i>	RV-00-T884	SPAIN. Vallgrassa, Parc del Garraf, Barcelona	AY870560	AY954604	AY954574	AY954544
<i>Aoa affinis</i>	MFB-00-P469	INDONESIA. Karaentha, S. Sulawesi	AY870585	AY954608	AY954578	AY954548
<i>Aphrissa statira</i>	MFB-00-P199	COSTA RICA. Río Poas, La Garita	AY870572			
<i>Aporia (Aporia) crataegi</i>	RV-00-T758	SPAIN. Universitat Autònoma de Barcelona, Cerdanyola del Vallés, Barcelona	AY870529			
<i>Aporia (Metaporia) agathon</i>	MFB-00-P481	CHINA. Near Lijiang, Yunnan Province	AY870590			
<i>Appias (Catophaga) paulina</i>	MFB-00-P159	AUSTRALIA. Buderim Forest Park, Buderim, Qld	AY870588			
<i>Appias (Glutophrissa) drusilla</i>	MFB-00-P195	COSTA RICA. Río Poas, La Garita	AY870525	AY954609	AY954579	AY954549
<i>Appias (Phrissura) aegis</i>	MFB-00-P491	PHILIPPINES. Camiguin Island, near Luzon.	AY870589			
<i>Archonias brassolis</i>	MFB-00-P263	COSTA RICA. Río Orosi, Cartago Province	AY870534			
<i>Ascia monuste</i>	MFB-00-P170	COSTA RICA. Colón	AY870582			
<i>Baltia butleri</i>	MFB-00-P492	INDIA. Tagalanga-la, Kashmir	AY870583			
<i>Belenois java</i>	DL-00-Q546	AUSTRALIA. Waikeri, SA	AY870593	AY954617	AY954587	AY954557
<i>Catanticta teutila</i>	MFB-00-P247	COSTA RICA. Copey	AY870532	AY954614	AY954584	AY954554
<i>Catanticta cerberus</i>	MFB-00-P266	COSTA RICA. Cerro de la Muerte, Villa Mills	AY870533			
<i>Catopsilia pomona</i>	MFB-00-P059	PAPUA NEW GUINEA. WEI, Wau, Morobe Province	AY870569			
<i>Cepora primale</i>	MFB-00-P131	AUSTRALIA. Brisbane Forest Park, Qld	AY870524			
<i>Charonias eurytele</i>	MFB-00-P395	PERU. 9 km south of Tingo María	AY870531			
<i>Colias eurytheme</i>		CANADA. Ontario	AF173400*			
<i>Colias philodice</i>	NP-98-U268	USA. Harvard Forest, MA	AY870565	AY954600	AY954570	AY954540
<i>Colotis hetaera</i>	SC-01-T434	KENYA. Kibwezi	AY870520	AY954601	AY954571	AY954541
<i>Cunizza hirlanda</i>	MFB-00-P398	PERU. 9 km south of Tingo María	AY870556			
<i>Delias aganippe</i>	MFB-97-U344	AUSTRALIA. Mt. Ainslie, ACT	AY870511			
<i>Delias belladonna</i>	DL-01-N104	THAILAND. Doi Inthanon NP, Chiang Mai	AY870510			
<i>Dercas gobrias</i>	MFB-00-P475	MALAYSIA. Mt. G Serapi, Sarawak	AY870512			
<i>Dismorphia zathoe</i>	MFB-00-P231	COSTA RICA. Monteverde	AY870578	AY954596	AY954566	AY954536
<i>Dixeta charina</i>	SC-01-T400	KENYA. Kibwezi	AY870523	AY954619	AY954589	AY954559
<i>Elodina angulipennis</i>	MFB-00-P133	AUSTRALIA. Brisbane Forest Park, Qld	AY870522	AY954618	AY954588	AY954558
<i>Enantia lina</i>	MFB-00-P405	PERU. 9 km south of Tingo María	AY870576			
<i>Eroessa chiliensis</i>	MFB-00-P437	CHILE. Parque Nacional Puyehue, Los Lagos Región	AY870554	AY954605	AY954575	AY954545
<i>Eronia cleodora</i>	SC-01-T424	KENYA. Kibwezi	AY870551			
<i>Euheira socialis</i>	MFB-00-P284	MEXICO. Kilometer 64 on Highway 40, town of El Madrono, Durango	AY870535			
<i>Euclloe ausonides</i>	AF-94-A022	USA. Naked Hills, Gothic, CO	AY870558			

<i>Eurema (Eurema) mexicana</i>	DW-92-Z084	USA. South fork of Cave Creek, Chiricahua Mtns., Cochise Co., AZ	AY870563	AY954598	AY954568	AY954538
<i>Eurema (Terias) hecabe</i>	MFB-00-P036	PAPUA NEW GUINEA. WEI, Wau, Morobe Province	AY870587			
<i>Gandaca harina</i>	MFB-00-R059	BORNEO. Sungai Turan, north-east of Kalimantan	AY870514			
<i>Ganyra josephina</i>	MFB-00-P201	COSTA RICA. Río Poas, La Garita	AY870539			
<i>Gideona lucasi</i>	GA-93-Z024	MADAGASCAR. Kirindy Forest	AY870521			
<i>Gonpteryx rhamnii</i>	RV-00-R023	SPAIN. Cánoves (Vallés Oriental), Barcelona	AY870568			
<i>Hebomoia glaucippe</i>	NP-95-Y258	MALAYSIA. Fraser's Hill, Pahang	AY870581	AY954606	AY954576	AY954546
<i>Hesperocharis crocea</i>	MFB-00-P268	COSTA RICA. Alajuela	AY870555			
<i>Hypsochila wagenknechti</i>	MFB-00-P432	CHILE. Farellones, Región Metropolitana	AY870544			
<i>Infraophulia ilyodes</i>	MFB-00-P416	CHILE. Parinacota, Parque Nacional Lauca, Tarapaca	AY870542			
<i>Itaballia demophile</i>	MFB-00-P277	COSTA RICA. Near Río Virilla, 6 km north-west of Colón	AY870584			
<i>Ixias pyrene</i>	DL-00-Q581	THAILAND. Bang Khantak, Samut Songkram	AY870552	AY954602	AY954572	AY954542
<i>Kricogonia lyside</i>	MFB-00-P215	COSTA RICA. Parque Nacional Santa Rosa	AY870566			
<i>Leodonta tellane</i>	MFB-00-P265	COSTA RICA. Río Oroso	AY870537			
<i>Leptidea sinapis</i>	RV-00-T760	SPAIN. Vallgrassa, Parc del Garraf, Barcelona	AY870573	AY954595	AY954565	AY954535
<i>Leptophobia aripa</i>	MFB-00-P189	COSTA RICA. La Guacima	AY870546			
<i>Leptosia nina</i>	NP-95-Y248	MALAYSIA. Fraser's Hill, Pahang	AY870519	AY954616	AY954586	AY954556
<i>Leuciacria acuta</i>	MFB-00-P468	INDONESIA. Pass Valley Wamena, Papua.	AY870591			
<i>Leuciacria olivei</i>	MFB-00-S095	PAPUA NEW GUINEA. Schleinitz Mts, New Ireland	AY870592			
<i>Leucidia brephos</i>	PDV-94-B004	ECUADOR. Prov. Sucumbios, Garza Cocha – Anangu, 175 km east-south-east of Coca	AY870561			
<i>Lieinix nemesis</i>	MFB-00-P284	COSTA RICA. Monteverde	AY870579			
<i>Mathania leucothea</i>	MFB-00-P428	CHILE. Farellones, Región Metropolitana	AY870557			
<i>Melete lycimnia</i>	MFB-00-P316	PERU. 10 km south-west of San Ramón, Chanchamayo	AY870530			
<i>Moschoneura pinthous</i>	MFB-00-P403	PERU. 9 km south of Tingo María	AY870575			
<i>Mylothris agathina</i>	NP-99-T486	SOUTH AFRICA. Muizenberg, Western Cape	AY870528	AY954615	AY954585	AY954555
<i>Mylothris bernice</i>	SC-01-M002	TANZANIA. Mufindi, Kihansi	AY870513			
<i>Nathalis iole</i>	DC-98-U711	USA. Ft. Huadrúa field, Garden Canyon, AZ	AY870562	AY954599	AY954569	AY954539
<i>Neophasia menapia</i>	AMS-00-R052	USA. Ice Spring Road, Glenn Co, CA	AY870536			
<i>Nepheronia thalassina</i>	SC-01-T403	KENYA. Kibwezi	AY870518			
<i>Pareronia valeria</i>	MFB-00-P442	INDIA. Madras, Tamil Nadu	AY870517	AY954603	AY954573	AY954543
<i>Patia orize</i>	MFB-00-P353	PERU. 10 km south-west of San Ramón, Chanchamayo	AY870577			
<i>Perete charops</i>	MFB-00-P283	COSTA RICA. Río Macho, Cartago Province	AY870538			
<i>Perrhybris pamela</i>	MFB-00-P202	COSTA RICA. Río Poas, La Garita	AY870545			
<i>Phoebis sennae</i>	MFB-00-P197	COSTA RICA. Río Poas, La Garita	AY870571			
<i>Phulia nymphula</i>	MFB-00-P375	PERU. 22 km south-south-east of Junín	AY870541	AY954610	AY954580	AY954550
<i>Pieriballia viardi</i>	MFB-00-P221	COSTA RICA. Río Alondra, San Luís	AY870547	AY954612	AY954582	AY954552
<i>Pieris napi</i>		USA. CA	AF173401*			
<i>Pieris rapae</i>	NM-95-Y381	USA. Nora Murphy culture	AY870550	AY954611	AY954581	AY95451

Table 1. Continued

Taxon	MCZ voucher no.	Locality	GenBank accession no.		
			<i>EF-1α</i>	<i>wingless</i>	<i>COI</i>
<i>Pierphulia rosea</i>	MFB-00-P410	CHILE. Parque Nacional Lauca, Tarapaca Region	AY870543		28S
<i>Pinacopteryx eriphia</i>	GA-93-Z028	MADAGASCAR. Kirindy Forest	AY870553		
<i>Pontia (Pontia) helice</i>	NP-99-T476	SOUTH AFRICA. Swartberg Pass, north of Oudtshoorn, Western Cape	AY870549		
<i>Pontia (Synchloe) callidice</i>	AF-94-A003	USA. Gold Basin, Gunnison, CO	AY870548		
<i>Prioneris philonome</i>	DL-00-Q610	THAILAND. Gaw Chan Waterfall, Ratchaburi	AY870527	AY954613	AY954583
<i>Pseudopieris nehemia</i>	MFB-00-P312	PERU. 10 km south-west of San Ramón, Chanchamayo	AY870574	AY954597	AY954537
<i>Pseudopontia paradoxo</i>	SC-01-T380	ZAMBIA. Zambesi Bridge, Ikelenge	AY870580	AY954594	AY954534
<i>Pyrisitia proterpia</i>	MFB-00-P211	COSTA RICA. Parque Nacional Santa Rosa	AY870586		
<i>Saletara liberia</i>	NP-95-Y239	MALAYSIA. Fraser's Hill, Pahang	AY870526	AY954607	AY954577
<i>Talbotia naganum</i>	MFB-00-P467	LAOS. Laksao	AY870516		
<i>Tatochila autodice</i>	MFB-00-P306	PERU. San Mateo	AY870540		
<i>Teriocolas zelia</i>	MFB-00-P419	CHILE. Socoroma, Tarapaca Region	AY870564		
<i>Theochila maenacte</i>	MFB-00-P449	ARGENTINA. Escobar, Buenos Aires Province	AY870515		
<i>Zegris (Zegris) eupheme</i>	RV-00-Q026	SPAIN. Mequinenza, N.211 Km 293, Aragón	AY870559		
<i>Zerene cesonia</i>	MFB-00-P204	COSTA RICA. Parque Nacional Santa Rosa	AY870567		
Papilionidae					
<i>Papilio rutulus</i>	AF-94-A002	USA. Gold Basin, Gunnison, CO	AY954620	AF233563†	AY954530
Nymphalidae					
<i>Vanessa virginiensis</i>	CA-94-N006	USA. Charles River, Middlesex County, MA	AY954621	AY954591	AY954561
Lycaenidae					
<i>Lycaena helleoides</i>	NP-99-W131	USA. Lost Man Creek, Pitkin County, CO	AY954622	AY954592	AY954562
Riodimidae					
<i>Uraneis hyalina</i>	PDV-94-T013	ECUADOR. 175 km east-south-east of Coca, Prov. Sucumbios Garza Cocha – Anangu	AY954623	AY954593	AY954563

EF-1 α , elongation factor-1 alpha; *COI*, cytochrome oxidase subunit I.

*Sequences for these taxa are those published by Caterino *et al.* (2001).

†The sequence for this taxon is that published by Campbell *et al.* (2000).

74 genera plus six subgenera (i.e. a total of 80 lower taxa) representing all the higher systematic groups (four subfamilies, two tribes). This sample represents 89% of all genera and 82% of all lower taxa (genera and subgenera) currently recognized within the Pieridae (Braby, 2005).

Four species from the butterfly families Papilionidae, Nymphalidae, Riodinidae, and Lycaenidae were chosen as outgroup taxa (Table 1). The final data set for the combined analysis thus comprised 30 taxa (26 Pieridae, four outgroups), whereas that for the *EF-1 α* analysis comprised 90 taxa (86 Pieridae, four outgroups). The Pieridae are considered to be either the sister group to the Papilionidae (Ehrlich, 1958; Scott, 1985) or the sister group to Nymphalidae + (Riodinidae + Lycaenidae) (Kristensen, 1976; de Jong *et al.*, 1996; Weller *et al.*, 1996; Ackery *et al.*, 1999). A recent combined molecular and morphological study of all the butterfly families and superfamilies by Wahlberg *et al.* (2005) suggested that the latter hypothesis is more probable.

Nine pierid genera [*Abaeis*, *Prestonia*, *Rhabdodryas*, *Glennia*, *Reliquia*, *Piercolias*, *Calopieris*, *Udaiana*, *Appias* (*Appias*)] were not sampled, in some cases because of their rarity or occurrence in inaccessible/remote locations. The relationships of most of these taxa have been reasonably well hypothesized based on morphology and their absence was assumed probably not to affect overall tree topology and hence the higher-level systematic relationships at the tribal and subfamily level, although their inclusion in future research will help elucidate relationships among the lower levels (e.g. genera). A further three subgenera [*Zegris* (*Microzegris*), *Aporia* (*Mesapia*), *Appias* (*Hiposcritia*)], and the putative subgenera of *Colias*, were not sampled in the present study, although the genera to which all of these taxa belong were included in our study, in some cases by more than one species.

To improve our sampling, and to test for potential nonmonophyly, two exemplar species (sometimes representing different subgenera) were included in each of the following ten genera: *Eurema*, *Colias*, *Pieris*, *Pontia*, *Aporia*, *Delias*, *Leuciacria*, *Catasticta*, *Mylothris*, and *Appias* Hübner. Previous molecular phylogenetic studies have confirmed the monophyly of at least three genera: *Colias* (Brunton, 1998; Pollock *et al.*, 1998), *Gonepteryx* [Leach] (Brunton & Hurst, 1998), and *Delias* (Morinaka *et al.*, 2002; Braby & Pierce, 2006).

MOLECULAR TECHNIQUES

The following protocol was adopted to obtain DNA sequences of *EF-1 α* , *wingless*, *COI*, and *28S*. Three additional sequences were obtained from GenBank

based on the published work of Campbell *et al.* (2000) and Caterino *et al.* (2001) (see Table 1).

Specimen preparation

Specimens were collected as fresh adults from the field using a hand net and killed by pinching the thorax. Wings were immediately excised and stored in paper envelopes as vouchers for identification and the bodies were preserved in plastic vials containing 100% ethyl alcohol. The specimens were temporarily stored at -20°C for laboratory use and then ultimately transferred to -80°C for permanent storage. A few of the specimens were collected and preserved (from a few months to several years) as dried adults before the wings and body were stored and preserved as for the fresh specimens. All specimens are deposited in the DNA and tissues collection at the Museum of Comparative Zoology, Harvard University, USA.

DNA extraction

For the freshly preserved specimens, gDNA was extracted from the metathorax, homogenized manually in a 1.5 ml microcentrifuge tube containing 200–400 μl buffer solution [2% sodium dodecyl sulphate, 50 mM Tris-HCl, 20 mM ethylene diamine tetra acetic acid (EDTA) at pH 8.0], digested with Proteinase K (Gibco BRL/Life Technologies) 20 g l^{-1} for 2–3 h at 55°C , and then purified to separate the nucleic acids from the cellular debris through successive salt solution and ethanol precipitation at low temperature. The purified gDNA was dried and then resuspended in 110 μl of TE buffer (10 mM Tris, 0.1 M EDTA at pH 8) and stored at -20°C . For dried specimens, gDNA was extracted from a leg; the tissue was first rehydrated in 200 μl of buffer solution in a 1.5 ml Eppendorf tube for approximately 1 week at 4°C before homogenization, digestion, and precipitation. The precipitation steps were similar to the method used for fresh material but adjusted to maximize extraction of the degraded DNA fragments.

DNA amplification

The primers used for the amplification of the four genes in this study are given in Table 2. Approximately 1.1 kb of the *EF-1 α* gene was amplified in one or two fragments using different sets of primers. We used both published (Cho *et al.*, 1995; Monteiro & Pierce, 2001) and original primers for polymerase chain reaction (PCR) amplifications and sequencing. For *wingless*, an approximately 420 bp fragment was amplified using the single set of primers published in Brower & DeSalle (1998). For mitochondrial *COI*, an approximately 1.2 kb fragment was amplified using

Table 2. Primers used for the amplification and sequencing of the four genes. Position numbers correspond to the following reference sequences: elongation factor-1 alpha (*EF-1 α*), *Bombyx mori* (GenBank D13338); *wingless*, *Drosophila melanogaster* (GenBank M17230); cytochrome oxidase subunit 1 (*COI*), *Drosophila yakuba* (GenBank X03240); *28S*, *Drosophila melanogaster* (GenBank M21017)

Gene	Primer name (forward or reverse)	Positions (5'→3')	Sequence (5'→3')
<i>EF-1α</i>	EF44 (fwd)	240–262	GCYGARC GYGARCGTGGTATYAC
	EF46.1I (fwd)	549–567	GAGGAAATYAARAAGGAAG
	EF46.1III (fwd)	548–567	CGAGGAAATCAARAARGAAG
	EF46.1IV (fwd)	549–567	GAAGAAATCAAAAARGAAG
	EF51.9 (fwd)	798–817	CARGACGTATACAAAATCGG
	EF77I (fwd)	816–835	GGTGGTATTGGAACAGTRCC
	EF77II (fwd)	816–835	GGTGGTATTGGAACAGTSCC
	EF51r (rev)	650–631	CATGTTGTCGCCGTGCCAAC
	EF52.6r (rev)	940–921	GCTTCGTGGTGCATTTCAAC
	EFrcM4 (rev)	1351–1329	ACAGCVACKGTYTGYCTCATRTC
	<i>wingless</i>	LepWG1 (fwd)	1111–1136
LepWG2 (rev)		1775–1750	ACTICGRCACCARTGGAATGTRCA
<i>COI</i>	LCO1490 (fwd)	1490–1514	GGTCAACAATAATCATAAAGATATTGG
	Ron (fwd)	1729–1751	GGATCACCTGATATAGCATTCCC
	Tonya (fwd)	2191–2216	GAAGTTTATATTTTAAATTTTACCGGG
	Nancy (rev)	2216–2191	CCCCGTAATAATTAATAAATAAACTTC
	Hobbes (rev)	2756–2735	AAATGTTGNGGRAAAAATGTTA
<i>28S</i>	Mo6 (fwd)	3318–3337	CCCCCTGAATTTAAGCATAT
	D2B (fwd)	3549–3568	GTCGGGTTGCTTGAGAGTGC
	S3660 (fwd)	3668–3690	GAGAGTTMAASAGTACGTGAAAC
	D3A (fwd)	4046–4065	GACCCGCTTGAAACACGGA
	D2B-r (rev)	3568–3549	GCACTCTCAAGCAACCCGAC
	D3A-r (rev)	4065–4046	TCCGTGTTTCAAGACGGGTC
	A335 (rev)	4394–4375	TCGGARGGAACCAGCTACTA
	D3B (rev)	4414–4395	TCGGAAGGAACCAGCTACTA

standard primers (Folmer *et al.*, 1994; Monteiro & Pierce, 2001). For ribosomal *28S*, approximately 1.2 kb was amplified according to the primers published in Schmitz & Moritz (1994), Sequeira, Normark & Farrell (2000), and Saux, Fisher & Spicer (2004), although for approximately half the taxa only the 'downstream' 800 bp was amplified and sequenced.

Fragments were amplified according to standard PCR techniques using a thermal cycler and Qiagen PCR kit. For *EF-1 α* , *wingless*, and *COI*, standard PCR reactions, with a total volume of 25 μ l, were prepared using 0.5 μ l of gDNA template at various dilutions, with 2.5 μ l of buffer (100 mM Tris-HCl solution with 50 mM KCl), 0.5 μ l MgCl₂ (25 mM), 0.125 μ l of each dNTP (2.5 mM), 1.25 μ l of each primer (10 μ M), and 0.125 μ l of *Taq* polymerase (5 units μ l⁻¹). For *28S*, 25 μ l reactions were prepared using 0.25 μ l of gDNA template at various dilutions, with 2.5 μ l of buffer (100 mM Tris-HCl solution with 50 mM KCl), 2 μ l MgCl₂ (25 mM), 1 μ l dimethyl sulphoxide, 0.25 μ l of each dNTP (2.5 mM), 1.2 μ l of each primer (10 μ M), and 0.2 μ l of *Taq* polymerase (5 units μ l⁻¹).

For *EF-1 α* , samples were initially denatured at 95 °C for 2 min followed by 30 cycles of amplification (denaturation at 95 °C for 60 s, annealing at 55–51 °C for 60 s, extension at 72 °C for 2 min) with a final extension at 72 °C for 10 min; three or four cycles were used at each successive annealing temperature. If faint or no DNA bands were detected in the gel, PCRs were repeated and the concentrations of the template and/or the magnesium optimized. For dried specimens, a second amplification of the PCR product was necessary. The conditions for the amplification of *wingless* and *COI* followed the protocols reported in Campbell *et al.* (2000), and Rand *et al.* (2000) and Monteiro & Pierce (2001), respectively. For *28S*, samples were initially denatured at 95 °C for 2 min followed by 35 cycles of amplification (denaturation at 95 °C for 60 s, annealing at 52 °C for 60 s, extension at 72 °C for 2 min) with a final extension at 72 °C for 4 min. Negative controls were included in all PCRs to check for possible contamination. The PCR products of each template were combined and separated by electrophoresis on a 1 or 2% low-melting temperature

agarose gel. The portion of the gel containing DNA fragments was excised and the gel-extracted PCR products then purified using QIAquick gel extraction kit columns.

DNA sequencing and alignment

Both strands of purified DNA fragments for each gene were reamplified and sequenced with a range of forward and reverse primers (see Table 2) using ABI Dye Terminator or Big Dye cycle sequencing kits. Half cycle sequence reactions (10 µl) were prepared and denatured at 96 °C for 3 min followed by 25 cycles (Dye Terminator: 96 °C for 30 s, 50 °C for 15 s, 60 °C for 4 min; Big Dye: 96 °C for 10 s, 50 °C for 10 s, 60 °C for 4 min). Samples were loaded on to a polyacrylamide gel and sequenced on an ABI 370 or 377 automated sequencer, or loaded into a 3100 ABI genetic analyser capillary sequencer. Sequence contigs generated from each reaction were edited manually and then aligned for each sample using SEQUENCHER version 3.0 (Sequencher, 1995) or version 4.1.2 (Sequencher, 2000) software. Ambiguities and gaps (typically at the ends of a sequence) were treated as missing data.

For *EF-1α*, the consensus sequence of each sample was aligned against the published sequence for *Bombyx mori* (Kamie *et al.*, 1993) and primer ends were removed, resulting in 1066 bp (corresponding to positions 263–1328). For *wingless*, sequences (403 bp after the removal of primer ends) were aligned against other published Lepidoptera sequences (Brower & DeSalle, 1998; Campbell *et al.*, 2000). For *COI*, the consensus sequences were aligned against the published reference sequence for *Drosophila yakuba* (Clary & Wolstenholme, 1985) and/or other Lepidoptera sequences on GenBank; the final fragment was 1220 bp (corresponding to positions 1515–2734). Aligning *EF-1α* and *COI* sequences did not require any indels, but in *wingless* one sample (*Mylothris agathina*) had a one-codon deletion, and another sample (*Phulia nymphula*) had three-codon deletions. Codon positions were either analysed in SEQUENCHER 3.0 or exported into MacClade version 3.08a (Maddison & Maddison, 1999) or version 4.03 (Maddison & Maddison, 2001) and translated to amino acids. For *28S*, sequences were initially aligned against the published reference sequence for *Drosophila melanogaster* (Tautz *et al.*, 1988); improved alignment was obtained using CLUSTALX version 1.81 (Thompson *et al.*, 1997) and then manually using MacClade 4.03 (Maddison & Maddison, 2001). Ambiguous regions were removed, resulting in a final character set of 986 bp, which included internal gaps as well as nonsequenced terminal regions for some taxa. GenBank accession numbers for all sequences are given in Table 1.

PHYLOGENETIC ANALYSIS

Maximum (cladistic) parsimony (MP), maximum likelihood (ML) (using PAUP), ML (using PHYML), and Bayesian inference (BI) were carried out for the smaller taxon data set of the four genes combined, as well as for the larger taxon data set of the *EF-1α* gene. We also ran an 'all available data' analysis, using MP and ML, of the entire data by combining the 30 taxon data set of the four genes with the 90 taxon data set of *EF-1α*. The final data matrix of this data set thus consisted of 90 taxa, that is, 30 taxa with sequences from *EF-1α*, *wingless*, *COI*, and *28S*, plus 60 taxa with sequences from *EF-1α* only, with the remaining three genes coded as 'missing' data.

MP

Phylogenetic trees were reconstructed using unweighted and weighted MP as the optimality criterion, as implemented in PAUP* version 4.0b10 (Swofford, 2002). Tree estimation involved heuristic searches with the tree-bisection-reconnection (TBR) branch-swapping algorithm, stepwise addition with up to 1000 random starts to check for islands of trees, and 'MulTrees' option in effect. Searches of large data sets that still recovered numerous islands of trees after approximately 100 random additions were repeated using PAUPRat (Sikes & Lewis, 2001). Strict consensus trees were computed where there was more than one equally parsimonious tree. Results based on MP analyses of each codon position, as well as those obtained from other methods (e.g. neighbour joining), were compared to establish that there was no conflict of signal within each data set. Bootstrap analyses (Felsenstein, 1985, 1988), based on a full heuristic search of 1000 pseudoreplicates using TBR branch swapping and simple stepwise addition, were carried out for each analysis to determine the level of support of each node (clades with bootstrap values < 50% were collapsed). In order to ascertain the extent of saturation, transition:transversion ratios were plotted against the observed or uncorrected pairwise 'p' distance for each codon position. Various weighting schemes were also explored, including weighting transversions over transitions (2 : 1 or 3 : 1).

For the smaller taxon data set of the four genes combined, each gene partition was first analysed separately using unweighted MP and the topology of the resulting trees compared for congruence before combining the data. Clade robustness was also evaluated using Bremer support (decay index) (Bremer, 1988, 1994) using the program TreeRot version 2 (Sorenson, 1999). Partitioned Bremer support was calculated to assess the contribution of each data partition to the total Bremer support values in the combined analysis.

ML

Phylogenetic trees were estimated using ML tree-building methods, as implemented in PAUP* version 4.0b10. Analyses based on the ML optimality criterion were performed according to the general time reversible substitution model (Lanave *et al.*, 1984; Rodríguez *et al.*, 1990) with among-site rate variation (invariable sites and gamma distribution) (i.e. GTR + I + Γ). Model selection was determined according to the hierarchical likelihood ratio test (hLRT) as implemented in ModelTest 3.06 (Posada & Crandall, 1998) with the starting trees obtained by MP. Models that best fitted the observed data were then used to generate an ML tree under a heuristic search using the TBR branch-swapping algorithm with as-is stepwise addition. Minor variations in estimates of model parameters were found not to affect the final tree topology. ML trees were also reconstructed using PHYML version 2.4.3 (Guindon & Gascuel, 2003). The model used was GTR + I + Γ , according to hLRT, with model parameters optimized automatically. Starting trees were distance based (BIONJ) according to the default option. Nonparametric bootstrap analyses based on 2000 pseudoreplicates were carried out to determine the approximate level of support for all branching events, with support percentages computed by majority rule consensus.

BI

Finally, we ran BI partitioned by codon position (first and second; third) in MrBayes 3.0b4 (Ronquist & Huelsenbeck, 2003), with the HKY85 + I + Γ model of sequence evolution for first and second positions, and GTR + I + Γ for third positions. Unlinked model parameters were preset as starting values for all partitioned analyses. Three independent Bayesian runs

at temperature settings from 0.2 to 0.4 were performed on the data using metropolis-coupled Markov chain Monte Carlo simulations, from one to 10 million generations each, and tree sampling every 100 generations. Bayesian topology and branch posterior probabilities were computed by majority rule consensus after deleting as 'burn in' all preasymptotic tree scores.

AGE OF DIVERGENCE ESTIMATIONS

In order to estimate the approximate age of divergence events within the Pieridae, the evolutionary rate of substitution for the molecular data set was calibrated using dated fossils, rather than ages of vicariance events inferred from biogeography and geological data, to estimate minimum divergence times of lineages within the framework of our phylogenetic hypothesis (Hillis, Mable & Moritz, 1996b; Arbogast *et al.*, 2002; Hedges & Kumar, 2003; Magallón, 2004). Although butterflies are rarely preserved as fossils, several have been discovered and described from the Pieridae (Scudder, 1875a, 1889; Zeuner, 1942; Brown, 1976; Shields, 1976; Carpenter, 1992; Emmel, Minno & Drummond, 1992). Four of these fossils are recorded from the Tertiary (Table 3), the oldest being two species from the Florissant Formation, Colorado, dated Late Eocene (34.07 ± 0.10 Myr) (Evanoff, McIntosh & Murphey, 2001). Because the nearest relatives of these fossils have been determined with some degree of certainty, the fossils served as useful calibration points.

To calibrate the rate of substitution, we first assessed if the rate was constant (i.e. clock-like) by comparing the likelihood scores of our best ML model of the *EF-1 α* data set with and without enforcing a molecular clock, using a LRT in PAUP. The LRT test rejected the null hypothesis that the data were clock-

Table 3. Summary of known pre-Quaternary fossils recorded for the Pieridae*. Data collated from Scudder (1875a), Zeuner (1942), Shields (1976), Brown (1976), Emmel *et al.* (1992), and Carpenter (1992)

Taxon	Closest relative(s)	Locality	Deposit	Epoch (Myr)
<i>Stolopsyche libytheoides</i> Scudder, 1889	<i>Pieris</i>	USA (Colorado)	Lacustrine shales	Upper Eocene (34)
<i>Oligodonta florissantensis</i> F.M. Brown, 1976	<i>Catasticta</i> group (possibly <i>Leodonta</i>)	USA (Colorado)	Lacustrine shales	Upper Eocene (34)
<i>Coliates proserpina</i> Scudder, 1875	<i>Delias-Prioneris</i> group (possibly <i>Aporia</i>)	France (Aix-en-Provence)	Calcareous marls of gypsum quarries	Lower Oligocene (33.5–30)
<i>Miopieris talboti</i> Zeuner, 1942	<i>Pontia</i> group (possibly <i>Pontia</i>)	Germany (Randecker Maar)	Dysodil shales	Upper Miocene (10.5–5.5)

**Mylothrites pluto* (Heer, 1849), recorded from marls of lacustrine beds of Lower Miocene age (23.5–16.5) from Radoboj, Yugoslavia, is excluded from the list because of doubt over its correct identity. Zeuner (1942) considered that it belonged to the Nymphalidae, but Carpenter (1992) treated it as a pierid, noting that its wings were similar in venation to *Mylothris* but similar in shape to the distantly related *Hebomoia*.

like ($\delta = 196$, d.f. = 88, $P < 0.0001$). However, inspection of the topology and branch lengths of our phylogram showed that rates of change within and between two major clades of interest were reasonably homogeneous (i.e. clock-like), relative to the rest of the clades in the tree. We therefore applied two methods: (1) Sanderson's semiparametric rate smoothing using a penalized likelihood method, as implemented in the r8s program (Sanderson, 2002), to correct rate heterogeneity across the entire tree; and (2) the quartet method (Cooper & Penny, 1997), which assumes that the data are ultrametric and clock-like, but does allow for rate variation among lineages (i.e. nonclock-like subsets of the data). The latter method involves determining the average genetic divergence between a pair of related taxa (i.e. between the lineages of two fossils), and calculating the rate of substitution using the age of the oldest fossil. The calculation is then repeated for another pair of related taxa. The two distantly related pairs of taxa are then combined into a quartet, and the two substitution rates averaged to give a calibrated rate for the gene. The minimum divergence time of the two pairs forming the quartet is then estimated based on the average corrected pairwise distance between the two pairs of fossils.

RESULTS

COMBINED ANALYSIS

The smaller (30 taxon) data set was assembled primarily to investigate patterns of higher-level relatedness within the Pieridae (e.g. subfamilies, tribes) by combining four independent markers. The final data set comprised a total of 3675 bp, of which 1031 bp (28%) were parsimony informative (Table 4).

The results of MP, ML (PHYML), and BI of the combined data set of the four genes are summarized in Figure 3. The results of the ML (PAUP) analysis were identical to those for ML (PHYML) (tree not shown). Tree topologies generated by each method of analysis were broadly similar, the major differences being in

the placement of *Pareronia*, *Leptosia*, and *Elodina*. Eleven major clades within the Pieridae (labelled A–K) were identified. With the exception of clade E, these clades were consistently recovered under the different methods of analysis and with a high level of support (Fig. 3, Table 5). MP and ML analyses of the individual data partitions showed no deep phylogenetic structure, and hence little conflict, among the basal nodes (trees not shown). Partitioned Bremer support of the combined data set under MP revealed a high level of congruence between the four genes for most nodes (Table 5). The only substantial source of conflict was with the mitochondrial gene *COI*, which contributed negatively to the total support value in two major clades (C and G) (Table 5), probably as a result of saturation due to high A–T bias.

In terms of the current higher-level classification, relationships within the Pieridae showed good agreement with those based on morphology (Klots, 1933; Ehrlich, 1958; Bridges, 1988). There was strong support for the monophyly of the three larger subfamilies in the combined analyses: Dismorphiinae (clade B: bootstrap 100% MP, ML, and BI), Coliadinae (clade C: bootstrap 73% MP, 100% ML and BI), and Pierinae (clade D: bootstrap 76% MP, 97% ML, 100% BI) (Fig. 3). A sister relationship between the Dismorphiinae and the Pseudopontiinae was evident in two analyses with reasonable support (bootstrap 76% MP, 80% ML), and in the two ML trees this clade formed the sister group to the rest of the Pieridae (Fig. 3B). The Coliadinae and Pierinae were sister taxa in two analyses (bootstrap 85% ML, 97% BI). The Pierinae was found to consist of three major clades (E–G), with clade G composed of four smaller subclades (H–K). The tribe Pierini, however, was paraphyletic as it included the three exemplars from the Anthocharidini (clade F).

EF-1 α ANALYSIS

The *EF-1 α* analysis of the larger (90 taxon) data set was used primarily to investigate patterns of lower-level relatedness within the Pieridae (e.g. genera,

Table 4. Character summary for the combined data set, with numbers of sites for each codon position for each gene partition

Gene partition	<i>EF-1α</i>				<i>wingless</i>				<i>COI</i>				28S	Total
	1st	2nd	3rd	All	1st	2nd	3rd	All	1st	2nd	3rd	All		
Number parsimony informative	22	10	271	303	40	11	127	178	74	15	320	409	141	1031
Number variable but parsimony uninformative	19	6	42	67	21	26	3	50	34	15	47	96	87	300
Number constant	314	339	43	696	73	97	5	175	299	376	40	715	758	2344
Total characters	355	355	356	1066	134	134	135	403	407	406	407	1220	986	3675

EF-1 α , elongation factor-1 alpha; *COI*, cytochrome oxidase subunit I.

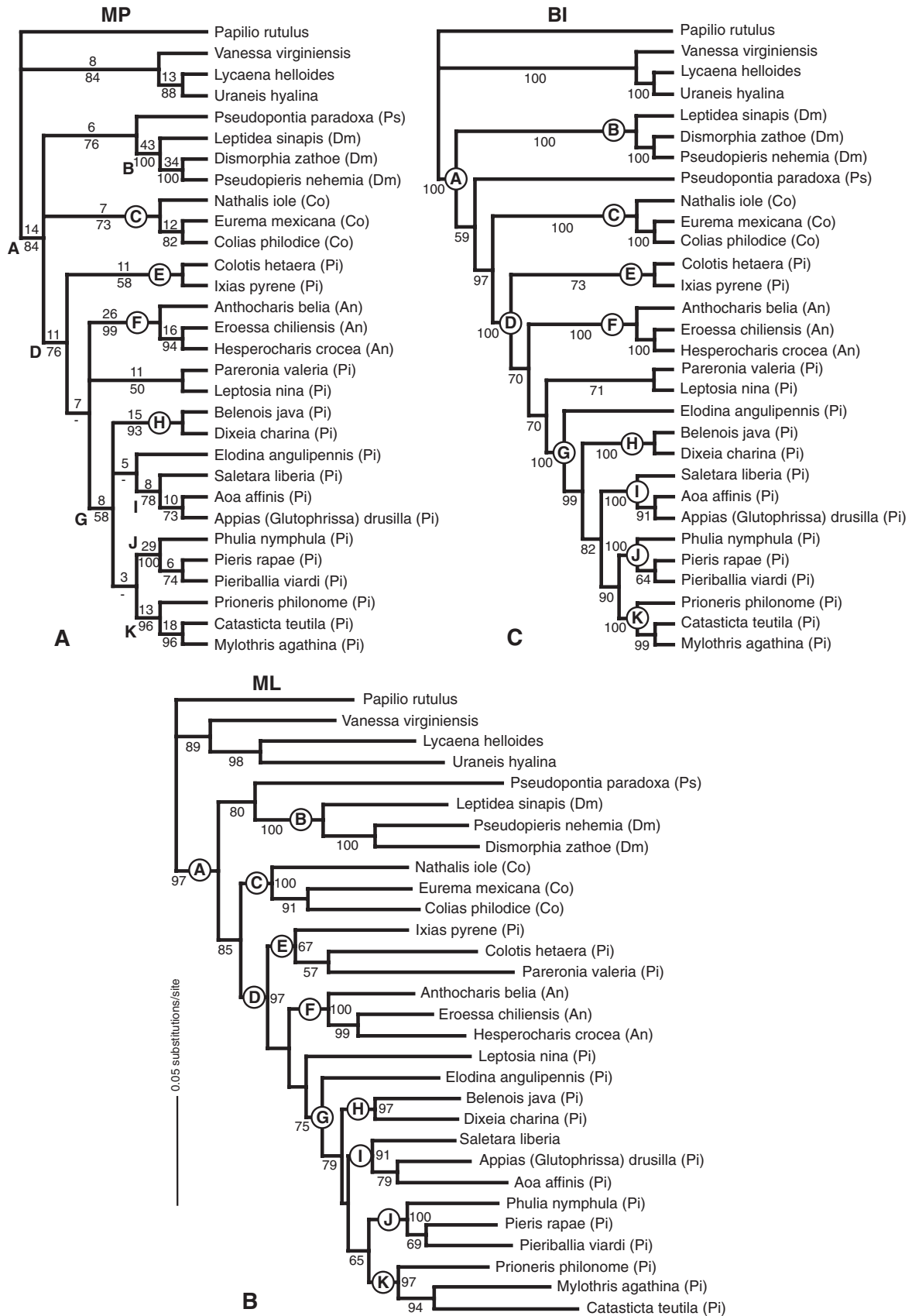


Figure 3. Phylogenetic trees for 26 pierid taxa based on the combined data set of four genes (3675 bp, 1031 informative characters). A, strict consensus of three equally most parsimonious trees [length = 6572, consistency index (CI) = 0.321, retention index (RI) = 0.316]; values below the branches are bootstraps (1000 full heuristic search replicates) for nodes with ≥50% support, those above the branches are total Bremer support indices. B, maximum likelihood (ML–PHYML) tree according to the GTR + I + Γ substitution model [log likelihood score = -32254.152; relative rate matrix A↔C 2.2381, A↔G 7.7859, A↔T 7.5951, C↔G 2.3345, C↔T 14.0782, G↔T 1.0; base frequencies A = 0.2482, C = 0.2307, G = 0.2095, T = 0.3116; proportion of invariable sites (I) = 0.5423; shape parameter (α) of gamma distribution (Γ) = 0.8036], with bootstrap values (2000 pseudoreplicates) shown below the branches or adjacent to nodes with ≥50% support. C, Bayesian inference (BI) tree partitioned by gene and codon position; the unlinked partition model is HKY85 + I + Γ for first and second codon positions for elongation factor-1 alpha (*EF-1α*), *wingless*, and cytochrome oxidase subunit I (*COI*) partitions and for all positions for *28S* partition, GTR + I + Γ for third codon positions for each gene partition, at a sampling temperature of 0.4; values below the nodes are posterior branch supports estimated from majority rule consensus of 90 000 trees (10⁷ generations, 10⁶ burned). The two letters in parentheses after each taxon name are the currently recognized subfamily/tribe, abbreviated as follows: Ps, Pseudopontiinae; Dm, Dismorphiinae; Co, Coliadinae; Pi, Pierini (Pierinae); An, Anthocharidini (Pierinae). Bold capitalized letters (A–K) denote major clades evident in the analysis. *Papilio*, *Vanessa*, *Lycaena*, and *Ura-neis* are outgroup taxa.

Table 5. Total Bremer support and partitioned Bremer support for each gene for nodes in the strict consensus maximum parsimony (MP) cladogram of the combined data set (Fig. 3A). Rows in bold refer to nodes/clades that comprise the higher taxa recognized in this work (see Discussion)

Node	Clade	Higher taxon	Total	<i>EF-1α</i>	<i>wingless</i>	<i>COI</i>	<i>28S</i>
1			8	-0.7	3.0	-1.3	7.0
2			13	3.3	1.0	3.7	5.0
3	A	Pieridae	14	3.3	3.0	0.7	7.0
4		Pseudopontiinae + Dismorphiinae	6	0.3	3.0	-2.3	5.0
5	B	Dismorphiinae	43	11.3	16.0	3.7	12.0
6			34	8.3	11.0	4.7	10.0
7	C	Coliadinae	7	3.3	2.0	-3.3	5.0
8			12	11.3	-5.0	-3.3	9.0
9	D	Pierinae	11	5.3	3.0	0.7	2.0
10	E		11	2.3	2.0	3.7	3.0
11			7	2.3	-2.0	7.7	-1.0
12	F	Anthocharidini s.s.	26	3.7	6.2	7.7	8.4
13			16	6.8	6.5	3.7	-1.0
14			11	0.3	5.0	2.7	3.0
15	G	Pierini s.s.	8	1.3	10.5	-3.3	-0.5
16	H	<i>Dixeia</i> + <i>Belenois</i>	15	6.3	-2.0	9.2	1.5
17			5	-0.7	-2.0	5.7	2.0
18	I	Appiadina	8	3.3	4.0	-0.3	1.0
19			10	3.3	5.0	-0.3	2.0
20			3	2.3	0.0	1.2	-0.5
21	J	Pierina	29	10.3	10.3	2.0	6.4
22			6	-2.7	0.0	7.7	1.0
23	K	Aporiina	13	1.3	3.0	8.7	0.0
24			18	5.8	1.5	8.7	2.0

EF-1α, elongation factor-1 alpha; *COI*, cytochrome oxidase subunit I.

subgenera), that is, relationships among the shallow nodes or tips of the tree. Of the 1066 bp sequenced for the gene, 364 sites (34%) were parsimony informative. As expected, most of the variable sites were in the third codon position. First and second positions were

highly conserved, with a total of only 46 sites (4%) parsimony informative. Mean base frequencies were similar and not significantly different across bases (A = 0.26263, C = 0.25572, G = 0.23161, T = 0.25004). A plot of the transition/transversion ratio against

uncorrected pairwise 'p' distance for all codon positions, third codon positions, and first and second positions inferred under MP, revealed that the gene was significantly saturated after approximately 10% divergence (not shown). Saturation was limited to third positions, being most pronounced among the deeper-level divergences (> 30%), and was not evident in first and second positions.

Figure 4 shows the strict consensus of nine equally MP trees based on unweighted analysis. The analysis recovered the Dismorphiinae (clade B), Coliadinae (clade C), and two clades within the Pierinae (clades J and K), with moderate to high support (bootstrap 64–100%), plus a number of smaller, less well-supported groups (clades E–I) that were evident in the combined analysis (Fig. 3). BI gave a similar tree to MP (tree not shown). ML (PHYML) yielded a single tree with similar topology in terms of shallow relationships among the exemplar taxa (Fig. 5). The same four well-supported clades were again evident (bootstrap 75–100%), and there was increased support for the monophyly of another large clade (F). However, the deeper nodes lacked support and, although the supported clades were concordant with those obtained in the combined smaller data set, their relationships were not resolved.

Of the ten genera where multiple species were examined, seven (70%) were monophyletic and three (*Eurema*, *Catantacta*, *Appias*) were not (Figs 4, 5). However, only in *Eurema* was there significant evidence (bootstrap 75% MP, 87% ML) in support of paraphyly: the *Eurema* clade included the genera *Leucidia*, *Teriocolias*, and *Pyrisitia*. Relationships within the two other genera, *Catantacta* and *Appias*, were essentially unresolved.

Dismorphiinae

The exemplars of the subfamily Dismorphiinae formed a tightly structured, well-supported monophyletic group (clade B: bootstrap 98% MP, 99% ML) (Figs 4, 5). Phylogenetic relationships within the Dismorphiinae were extremely well resolved. The Palaearctic *Leptidea* was sister to the remaining genera, all from the Neotropical region. In the latter clade, both MP and ML analyses yielded the following topology: *Pseudopieris* + (*Moschoneura* + ((*Enantia* + *Patia*) + (*Dismorphia* + *Lieinix*))).

Coliadinae

Members of the subfamily Coliadinae (the sulphurs) formed a well-supported monophyletic group (clade C: bootstrap 70% MP, 90% ML) (Figs 4, 5). Deeper-level splits within the Coliadinae were not well supported and provided only a polytomy. However, tree topologies generated by MP and ML methods were in general agreement, with both analyses suggesting that *Nathalis* and *Kricogonia* from the New World were sister taxa to all other genera in the Coliadinae. In all analyses, *Zerene* and *Colias* were recovered as sister taxa and were well supported by MP and ML (bootstrap 71–76%).

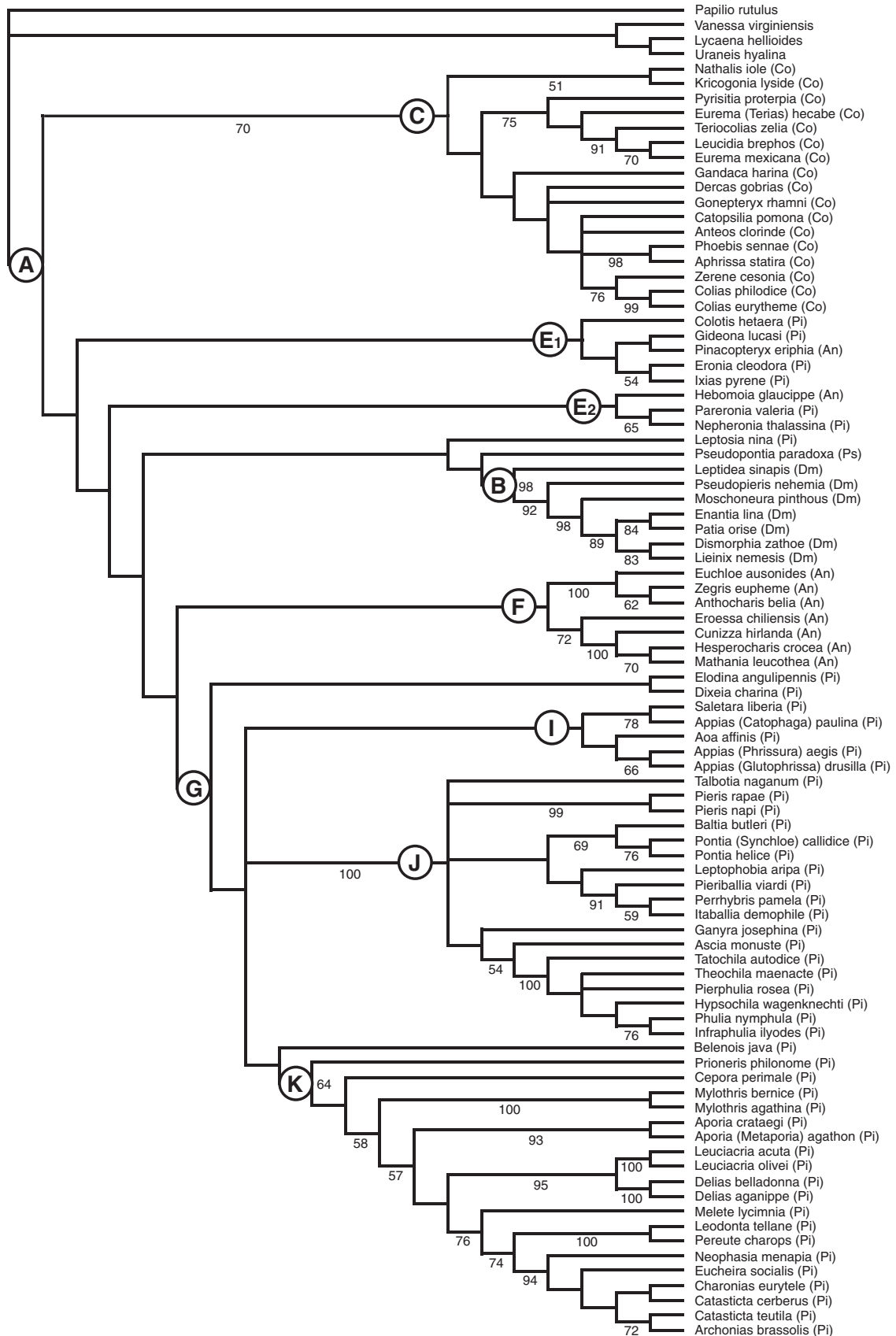
Pierinae

The Pierinae (clade D) were recovered as a monophyletic group under ML but without support. The Anthocharidini and Pierini were both nonmonophyletic, with two genera (*Hebomoia*, *Pinacopteryx*) traditionally placed in the Anthocharidini (clade F) and seven genera (*Colotis*, *Gideona*, *Ixias*, *Eronia*, *Pareonia*, *Nepheronia*, *Leptosia*) normally associated with the Pierini (clade G) comprising three separate groups (clades E₁ and E₂, *Leptosia*) outside these two tribes (Figs 4, 5).

The seven other genera from the Anthocharidini [*Euchloe*, *Anthocharis*, *Zegris* (*Zegris*), *Eroessa*, *Cunizza*, *Hesperocharis*, *Mathania*] appeared to form a clade, but support for their monophyly was weak (clade F: bootstrap < 50% MP, 68% ML) (Figs 4, 5). This clade, however, consisted of two well-supported subclades: the Holarctic *Anthocharis* group, commonly known as the 'orange tips', and the Neotropical *Hesperocharis* group. The *Anthocharis* group (bootstrap 100% MP, 99% ML) comprised the genera *Euchloe*, *Anthocharis*, and *Zegris* (*Zegris*) in an unresolved trichotomy; whereas the *Hesperocharis* group (bootstrap 72% MP, 82% ML) consisted of *Eroessa*, *Cunizza*, *Hesperocharis*, and *Mathania*. The latter three taxa formed a well-supported monophyletic group (bootstrap 100% MP and ML) sister to *Eroessa* with the following topology: *Cunizza* + (*Hesperocharis* + *Mathania*).

Of the other taxa within the Pierini (clade G), one group of taxa, comprising the genera *Appias* (*Catopha*), *Appias* (*Glutophrissa*), *Appias* (*Phrissura*), *Saltara*, and *Aoa*, appeared to form a cluster (clade I),

Figure 4. Strict consensus of nine equally maximum parsimony (MP) trees for the family Pieridae based on 1066 bp elongation factor-1 alpha (*EF-1 α*) [364 informative characters: length = 3777, consistency index (CI) = 0.190, retention index (RI) = 0.548]. Bootstrap values (1000 full heuristic search replicates) are shown below the branches or adjacent to nodes with $\geq 50\%$ support. Letters in parentheses after each taxon name are as per Figure 3. Bold capitalized letters (A–K) denote the major clades evident in the combined analysis of Figure 3. *Papilio*, *Vanessa*, *Lycaena*, and *Uraneis* are outgroup taxa.



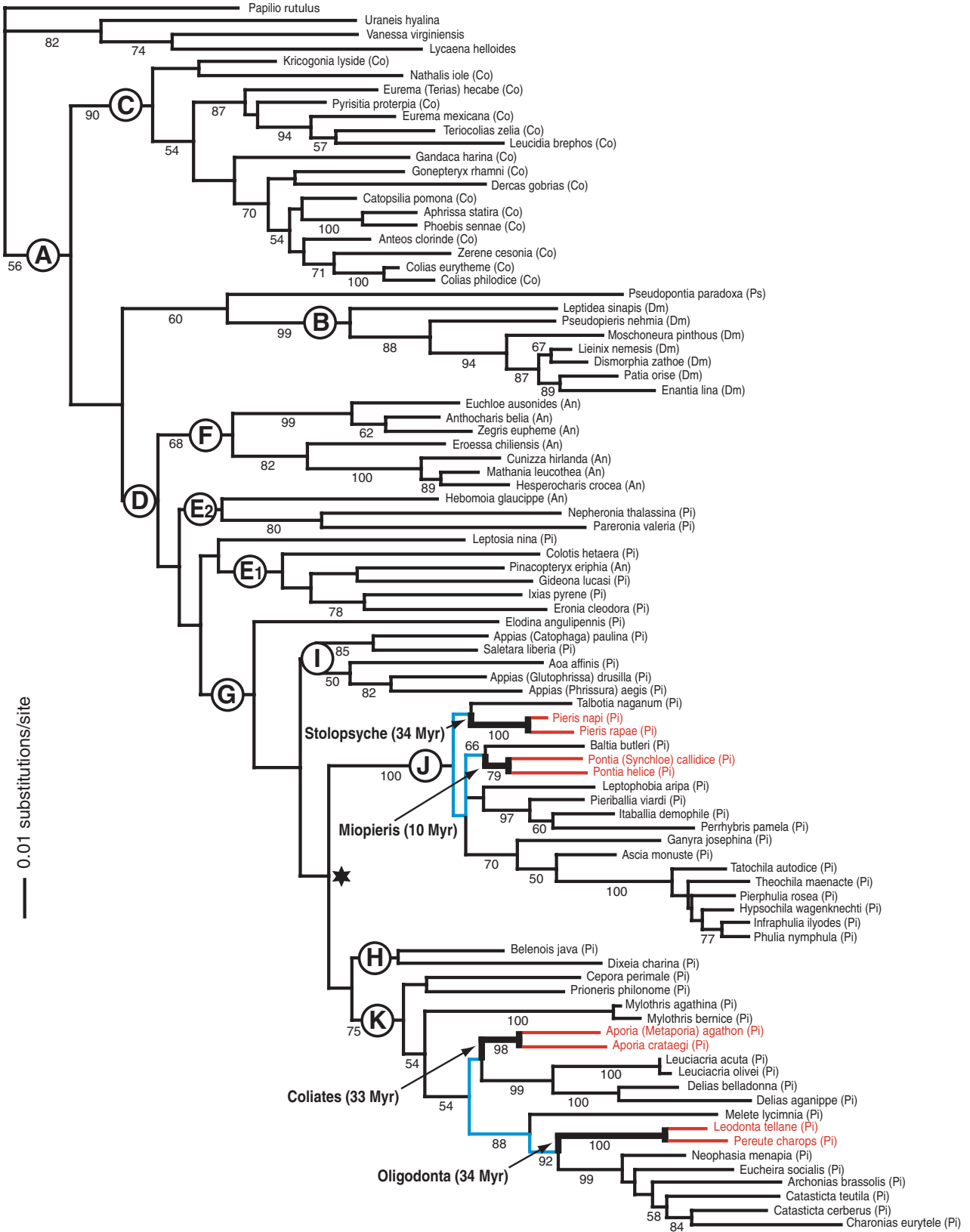


Figure 5. Maximum likelihood (ML–PHYML) tree for the family Pieridae based on 1066 bp elongation factor-1 alpha (*EF-1 α*) according to the GTR + I + Γ substitution model [log likelihood score = -17727.75: relative rate matrix A \leftrightarrow C 2.1063, A \leftrightarrow G 9.3033, A \leftrightarrow T 3.9410, C \leftrightarrow G 1.4829, C \leftrightarrow T 13.0754, G \leftrightarrow T 1.0; base frequencies A = 0.2729, C = 0.2222, G = 0.1946, T = 0.3102; proportion of invariable sites (I) = 0.5903; shape parameter (α) of gamma distribution (Γ) = 1.0686]. Bootstrap values are given below the branches or adjacent to nodes with $\geq 50\%$ support (2000 pseudoreplicates). Letters in parentheses after each taxon name are as per Figure 3. Bold capitalized letters (A–K) denote the major clades evident in the combined analysis of Figure 3. *Papilio*, *Vanessa*, *Lycaena*, and *Uraneis* are outgroup taxa. Extinct taxa based on fossils are indicated by thick lines at various nodes and along internal branches according to their putative relative(s) (taxa and external branches are highlighted in red) (see Table 3). Internal branches for each set of pairwise comparisons are shown in blue. The minimum age of divergence between clades J and K (node indicated by an asterisk) is estimated to be approximately 60 Myr.

but support for monophyly was not evident (Figs 4, 5).

Another 17 genera/subgenera within the Pierini (clade G) comprised an extremely well-supported monophyletic group in our analysis (clade J: bootstrap 100% MP and ML) (Figs 4, 5). This clade included the familiar *Pieris* and allied taxa often referred to as the typical ‘whites’. MP and ML analyses revealed three well-resolved subclades within clade J: the Neotropical *Itaballia* group, the largely Holarctic *Pontia* group, and the Neotropical *Tatochila* group. Relationships among these groups, however, were unresolved and the positions of five taxa (*Ascia*, *Ganyra*, *Leptophobia*, *Pieris*, *Talbotia*) were uncertain. The *Itaballia* group (bootstrap 91% MP, 97% ML) included three taxa: *Itaballia*, *Pieriballia*, and *Perrhybris*. The *Pontia* group (bootstrap 69% MP, 66% ML) included the taxa *Pontia* (*Pontia*), *Pontia* (*Synchloe*), and *Baltia*. The *Tatochila* group comprised a well-supported monophyletic group (bootstrap 100% MP and ML) of six genera from South America (*Tatochila*, *Hypsochila*, *Theochila*, *Pierphulia*, *Phulia*, *Infraphulia*) but not the Palaearctic *Baltia*. *Ascia* and *Ganyra* from the New World appeared to comprise sister taxa to the *Tatochila* group, although evidence for the associations were weak.

A further 15 genera/subgenera within the Pierini (clade G) formed a second major monophyletic group in our analysis (clade K: bootstrap 64% MP, 75% ML) (Figs 4, 5). This clade included the large and speciose genera *Delias* and *Catasticta*, as well as the Palaearctic *Aporia*, Afrotropical *Mylothris*, and the predominantly Oriental *Cepora* and *Prioneris*, both of which were sister to the remaining taxa. Both MP and ML analyses revealed strikingly similar topologies and significant structure within clade K, with several major subclades evident, including the Australian–Oriental *Delias* group, and the predominantly Neotropical *Catasticta* group. The *Delias* group comprised two Old World genera, *Delias* and the Australian *Leuciactria*, the monophyly of which was extremely well

supported (bootstrap 95% MP, 99% ML). The *Catasticta* group comprised a well-supported monophyletic group of eight genera (bootstrap 76% MP, 88% ML), all from the New World but predominantly from Central and South America (i.e. *Melete*, *Leodonta*, *Pereute*, *Neophasia*, *Eucheira*, *Catasticta*, *Archonias*, *Charonias*).

The remaining 12 taxa currently recognized from the Pierini did not belong to the four clades (F, I, J, K) outlined above (Figs 4, 5). Eight genera (*Colotis*, *Eronia*, *Ixias*, *Gideona*, *Pinacopteryx*, *Hebomoia*, *Pareronia*, *Nepheronia*), predominantly from the Afrotropical–Oriental regions, appeared to form two closely related clades (E₁, E₂), but there was no support for their monophyly. Four genera (*Leptosia*, *Elodina*, *Dixeia*, *Belenois*) were scattered across the topology of the trees, but their phylogenetic positions were inconsistent and hence their systematic relationships unresolved.

ALL AVAILABLE DATA ANALYSIS

The all available data analysis of the entire 90 taxon data set (i.e. 30 taxa for *EF-1 α* , *wingless*, *COI*, and *28S*, plus 60 taxa for *EF-1 α*) summarized well the topologies generated by the smaller and larger data sets, with high support for both shallow and deep nodes (Fig. 6). The deeper branching events were concordant with those generated by most trees in the combined analysis (Fig. 3), whereas the tips of the tree showed the same basic structure as that generated by the *EF-1 α* analysis (Figs 4, 5). However, the level of support for the basal nodes and for the monophyly of some clades was not as high as that recovered in the smaller combined data set.

AGE OF DIVERGENCE ESTIMATIONS

The wings, or parts thereof, of the four fossils used as calibration points were sufficiently well preserved to determine their broad systematic relationships within

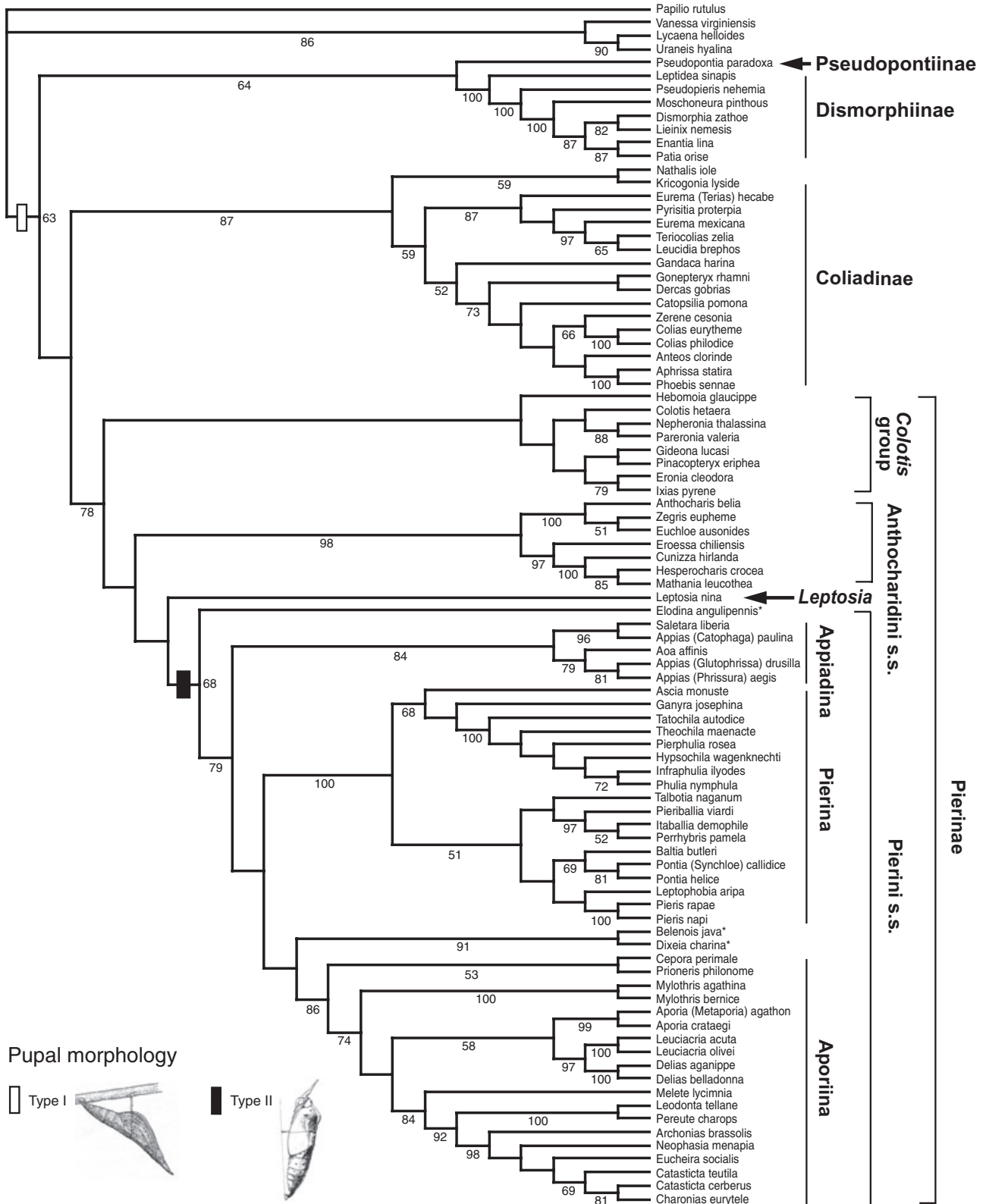


Figure 6. Cladogram for the Pieridae based on the all available data analysis of the 90 taxon data set of the four genes [30 taxa for elongation factor-1 alpha (*EF-1 α*), *wingless*, and cytochrome oxidase subunit I (*COI*), and 28*S*, plus 60 taxa for *EF-1 α*] (3675 bp, 1091 informative characters). Tree topology according to the GTR + I + Γ substitution model inferred under maximum likelihood (ML–PHYML) [log likelihood score = -41028.64: relative rate matrix A \leftrightarrow C 2.6276, A \leftrightarrow G 8.7266, A \leftrightarrow T 6.7753, C \leftrightarrow G 2.3732, C \leftrightarrow T 14.7215, G \leftrightarrow T 1.0; base frequencies A = 0.2518, C = 0.2191, G = 0.2023, T = 0.3265; proportion of invariable sites (I) = 0.5663; shape parameter (α) of gamma distribution (Γ) = 0.2260]. Bootstrap values (500 pseudoreplicates) are shown below the branches for nodes with $\geq 50\%$ support. *Papilio*, *Vanessa*, *Lycaena*, and *Uraneis* are outgroup taxa. To the right of the tree are the formal and informal names of the higher taxa recognized in this work (see Discussion and Appendix). Asterisks denote taxa of uncertain status. The two main pupal forms (types I, II) are mapped on the tree (see Discussion).

the family with some degree of confidence (Table 3). *Stolopsyche libytheoides* is considered to be the ancestor or sister taxon of *Pieris* (Scudder, 1889; Carpenter, 1992), whereas the venation of *Miopieris talboti* is very similar to *Pontia* and its allies (Zeuner, 1942), especially *Baltia*. *Oligodonta florissantensis* shows features reminiscent of the *Catasticta* group (Brown, 1976), particularly *Leodonta* and *Catasticta*: according to our estimate of the phylogeny it could be the ancestor of either *Leodonta* + *Pereute* or *Neophasia* + *Eucheira* + *Catasticta* + *Archonias* + *Charonias*. The forewing of *Coliatus proserpina* is similar to *Delias* in shape, the form of the anterior end of the discal cell, and in having vein R_2 absent, but the venation is unusual with veins R_3 and R_4 stalked, R_3 + R_4 long-stalked with R_5 (in *Delias* and allied genera, R_4 is fused with R_5 into a single vein), and vein M_2 forming a straight line with the discocellular vein at its point of origin. Scudder (1875a) placed *Coliatus* in the *Prioneris–Delias* group; according to our phylogeny, it could belong either with *Prioneris*, *Cepora*, *Mylothris*, *Aporia* or *Delias* + *Leuciacria*. We provisionally placed it with *Aporia* on biogeographical grounds, although we acknowledge that the forewing discal cell (which has the anterior end neatly truncated) and radial venation of *Coliatus* are quite distinct from *Aporia*. According to our phylogeny, two fossil genera (*Miopieris*, *Stolopsyche*) belong to clade J, whereas the two other genera (*Oligodonta*, *Coliatus*) belong to clade K. We mapped the approximate positions of the four fossils on the nodes and internal branches of the phylogram of our ML model that best fitted the observed data according to their nearest sister taxa (Fig. 5). From the phylogenetic distribution of these fossils, and their known age, it should be possible to estimate the approximate minimum ages of clades J and K, and their immediate common ancestor, to which the four extinct taxa belong. Although there was little support for the basal nodes in the tree generated by the *EF-1 α* analysis, the topology did not contradict that estimated in both the combined and all available data analyses for the nodes of interest. That is, clades J and

K are either sister taxa (Fig. 3) or comprise a monophyletic group with clade H (Fig. 6).

Penalized likelihood method

From the distribution of the four fossils (Fig. 5), it is clear that, for each fossil, two calibration points can be made depending upon which node is selected along the internal branches. In order to provide a conservative estimate of the substitution rate (i.e. fastest rate) and hence minimum age, we selected the most basal node for each fossil. From these four calibration points, the ages of various nodes were estimated in r8s with the value of the smoothing parameter λ set to 1000 and 3000 (i.e. two reconstructions were performed). The smoothing parameter was optimized using the cross-validation method, which minimizes the square and chi-square error terms. The high estimates of the smoothing parameter suggest that the data were in fact behaving in a clock-like manner. Confidence intervals were calculated for each estimate in r8s based on two (95%) and four (99.9%) standard deviations (SD) of the mean, with the four calibration points fixed and not free to vary.

The average rate of substitution for the *EF-1 α* gene was estimated to be $0.1277 \pm 0.0024\%$ (SD) per site per million years, which is equivalent to a divergence rate of 1% in 7.83 Myr. This substitution rate seems reasonable given that the average substitution rate for mtDNA (*COI*), which is much faster evolving than nuclear *EF-1 α* , is approximately $1.5\% \text{ Myr}^{-1}$ (i.e. 1% in 0.667 Myr) for arthropods (Quek *et al.*, 2004). In other words, our estimate of substitution for the nuclear gene is approximately 12 times slower than that for mitochondrial *COI* of other arthropods. The estimated minimum age of divergence for the putative split between clades J and K varied from 62.3 Myr ($\lambda = 1000$) to 60.7 Myr ($\lambda = 3000$). Errors in these estimates were small, with the confidence interval varying from 66.4–55.8 Myr (2 SD) to 68.8–54.1 Myr (4 SD) for the latter estimate. The minimum mean estimate for the crown-group of clade J was 40.6 Myr

$[\lambda = 3000$; confidence interval (4 SD) = 46.2–36.9 Myr], whereas that for the crown-group of clade K was 50.1 Myr [$\lambda = 3000$; confidence interval (4 SD) = 56.7–44.9 Myr]. The minimum mean estimate for the crown-group of the Pieridae was 95.5 Myr [$\lambda = 3000$; confidence interval (4 SD) = 111.6–82.5 Myr], although we are cautious about the wisdom of extrapolating too far beyond the calibration points to nodes deeper in the tree.

Quartet method

The mean corrected pairwise distance for each pair of fossil taxa, based on their nearest extant relatives, and their respective evolutionary rates are given in Table 6. The substitution rates for each pair varied greatly, with the *Coliatus–Oligodonta* split (0.327% Myr⁻¹) being almost twice that of the *Miopieris–Stolopsyche* split (0.178% Myr⁻¹). Averaging the two rates gave an overall mean rate of evolution within clades J–K of $0.252 \pm 0.106\%$ Myr⁻¹ (SD) (i.e. 1% = 4.0 Myr). The mean corrected pairwise distance between the *Miopieris–Stolopsyche* lineage and the *Coliatus–Oligodonta* lineage, that is, between (*Pontia (Pontia) callidice + Pontia (Synchloe) helice*) and (*Pieris rapae + P. napi*), and (*Aporia (Aporia) crataegi + Aporia (Metaporia) agathon*) and (*Leodonta tellane + Pereute charops*), was calculated to be 14.67%. According to the average rate of substitution (0.252% Myr⁻¹), and assuming a molecular clock, this level of genetic divergence between the two pairs of fossil lineages extrapolates to an average divergence time of 58 Myr (Table 6). In other words, the minimum age of the common ancestor of clades J and K, to which the fossil taxa belong, is 58 Myr.

DISCUSSION

HIGHER CLASSIFICATION

Our study represents the first rigorous phylogenetic analysis of the Pieridae, and indeed the first comprehensive phylogenetic study of a higher butterfly taxon at the familial level to date. The monophyly of the four currently recognized subfamilies is well supported (Table 5), corroborating several previous studies that recognize these taxa as distinct lineages (Ehrlich, 1958; Geiger, 1981; Scott, 1985; Janz & Nylin, 1998; Wahlberg *et al.*, 2005). However, it is clear that the largest subfamily, the Pierinae, contains far greater within-phylogenetic diversity than hitherto recognized. We therefore use these results in combination with previously published hypotheses to propose a revised higher-level systematic classification of the Pieridae (see the Appendix; also shown to the right of terminal taxa in Fig. 6 and in Table 5). In this classification, we provisionally place the 83 genera in the conventional four subfamilies in order to maintain the nomenclatural stability of the present classification (Knapp *et al.*, 2004), but divide the Pierinae into two tribes (Anthocharidini *s.s.*, Pierini *s.s.*) and two informal groups (*Colotis* group, *Leptosia*). The Pierini *s.s.* are further subdivided into three subtribes (Appiina, Pierina, Aporiina) and two informal groups comprising three genera (*Elodina*, *Dixeia*, *Belenois*) of uncertain status (*incertae sedis*). We discuss the status, monophyly and phylogenetic relationships of these higher taxa in more detail below.

A revised estimate of the phylogeny for the Pieridae, summarizing the interrelationships of the higher taxa and major clades recovered in the smaller combined

Table 6. Estimated time of divergence between clades J and K according to the phylogeny of Figure 5 using the quartet method (see Cooper & Penny, 1997). The estimate is based on the minimum divergence time for each pair of related taxa (lineages) according to their oldest known fossils. The fossils *Miopieris talboti* and *Stolopsyche libytheoides* belong to clade J, and *Oligodonta florissantensis* and *Coliatus proserpina* belong to clade K. Pairwise distances are the average corrected pairwise distances according to a GTR + I + Γ substitution model (see Fig. 5). The nearest relative of *Coliatus* is provisionally placed with *Aporia* on biogeographical grounds

Fossil taxon	Nearest related taxa (lineage)	Pairwise distance (%)	Minimum divergence time (Myr)	Average substitution rate (% Myr ⁻¹)	Split between clades J and K	
					Pairwise distance (%)	Estimated age (Myr)
<i>Miopieris</i>	<i>Pontia (Synchloe) callidice + Pontia helice</i>	6.0415	34	0.1777	0.2523	14.668
<i>Stolopsyche</i>	<i>Pieris rapae + P. napi</i>					
<i>Coliatus</i>	<i>Aporia crataegi + A. (Metaporia) agathon</i>	11.1148	34	0.3269		
<i>Oligodonta</i>	<i>Leodonta + Pereute</i>					

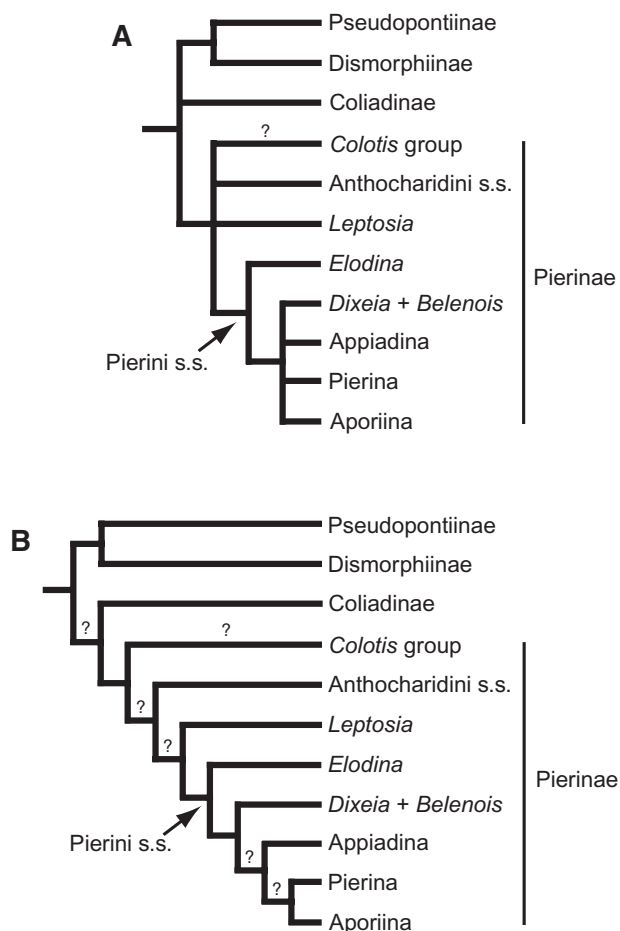


Figure 7. Higher classification of the Pieridae, showing two possible phylogenetic hypotheses according to the combined and all available data analyses of this study (Figs 3, 6). A, consensus tree summarizing nodes that are well supported or that are consistently recovered under different methods of analysis (maximum parsimony, maximum likelihood, Bayesian inference), with a question mark denoting uncertainty in the monophyly of the *Colotis* group. B, fully resolved tree, with question marks denoting uncertainty among nodes and in the monophyly of the *Colotis* group. Four subfamilies are recognized, with the subfamily Pierinae comprising four major lineages (two tribes, two informal groups); the tribe Pierini is subdivided into five lineages (three subtribes, two subclades of uncertain status).

analysis (Fig. 3) and the larger all available data analysis (Fig. 6), is presented in Figure 7A. In this tree, only nodes that are well supported or consistently recovered under different analytical methods (Giribet, 2003) are shown. Figure 7B portrays a fully resolved hypothesis and higher classification of the Pieridae according to the branching order of our ML tree (Fig. 3B), with question marks denoting uncertainty of nodes that are not well supported. Although there

is uncertainty in the phylogenetic position of the Coliadinae, the sister relationship between the Pseudopontiinae and the Dismorphiinae is well supported, and this lineage is almost certainly the sister group to the rest of the Pieridae. The most probable hypothesis for the subfamilial relationships is therefore (Pseudopontiinae + Dismorphiinae) + (Coliadinae + Pierinae) (Fig. 7B). This topology is identical to that originally proposed by Ehrlich (1958) (Fig. 1B) and will serve as our best estimate until further evidence is obtained to indicate a contrary pattern. Resolution of some of the uncertainties among the deeper-level divergences may only be overcome by addition of further genetic markers and/or integration of morphological characters from both adult and early stages (Wahlberg & Nylin, 2003).

An interesting feature of our all available data analysis (Fig. 6) is the close agreement with Klots' (1933) intuitive phylogeny based on morphology (Fig. 2). With the exception of the phylogenetic placement of the Anthocharidini and the Coliadinae, there is remarkable concordance of the two trees in terms of the higher-level structure. In relation to the lower-level structure, there is also strong concordance between groups of genera at the tips of the trees. Indeed, most of the 'shallow' differences lie among the 12 genera whose relationships Klots himself was uncertain about. Quite striking are parallels in the evolutionary relationships of the Pierini *s.l.* In a phylogenetic sense, Klots' concept of the Pierini was paraphyletic as it included the Coliadinae. Moreover, he envisaged two clades comprising six genera (*Gideona* + *Colotis* + *Ixias*) and (*Eronia* + (*Nepheronia* + *Pareronia*)), as each having separate origins from the main stock of Pierini *s.l.* (Fig. 2). These taxa show a similar pattern in our analyses, and comprise an assembly, termed the *Colotis* group, phylogenetically removed from the remaining Pierini *s.s.* (Fig. 6).

Our results also show a strong association between the morphology of the pupal stage and higher taxa. Pierid pupae approximately fall into two major groups according to differences in their morphology and habits (Talbot, 1939). In the Pseudopontiinae, Dismorphiinae, Coliadinae, and some groups in the Pierinae (i.e. *Colotis* group, Anthocharidini, *Leptosia*), the pupa (type I) is smooth, the wings are often strongly curved ventrally to form a prominent 'keel', and the head is tapered apically, often forming a prominent point or spine. The pupa is suspended loosely by the central girdle, usually horizontally or sometimes slightly upwards or downwards, but always with the ventral surface facing uppermost, similar to that of many Papilionidae. In contrast, the pupae of all members of the Pierini *s.s.* (i.e. Appiadina, Pierina, Aporiina, *Elodina*, *Dixeia*, *Belenois*) are characterized by having markedly different morphology and habits. In these

taxa, the pupa (type II) is more elongate with the ventral surface flat, the head has a horn or spine-like process anteriorly, which may be very prominent, the thorax has a pronounced dorsal ridge, the anterior abdominal segments (usually two to four) frequently have a series of dorsolateral spines or projections, and sometimes the abdomen has a series of dorsal projections on each segment. A central girdle secures the pupa close to the substrate, usually vertically or horizontally, but nearly always with the dorsal surface facing uppermost. These two pupal forms are mapped on the topology of our cladogram for the all available data analysis (Fig. 6). From the phylogeny it is clear that pupal form type I is the ancestral (plesiomorphic) form and that pupal form type II is a derived trait, having evolved once within the family and after the origin of the Pierinae. Pupal form type II is thus a synapomorphy for the Pierini *s.s.*

Pseudopontiinae and Dismorphiinae

The close relationship between the Pseudopontiinae and the Dismorphiinae (Table 5, Figs 3–6) supports previous conclusions drawn by Klots (1933) and Ehrlich (1958) (Fig. 1A, B), and Ackery *et al.* (1999) based on morphological evidence, particularly the male genitalia. *Pseudopontia* was placed in a separate subfamily because of its peculiar venation and other features. The lineage is long-branched (Figs 3B, 5) and comprises a single terminal taxon, indicating that either substantial evolutionary change has occurred since it diverged from the ancestor of Dismorphiinae + Pseudopontiinae (possibly due to a population bottleneck in the past), or that there have been considerable extinction events in the lineage. Yoshimoto (2000) suggested that *Leptidea* was unrelated to the *Dismorphia* group, but our analysis refutes this and recovers the Dismorphiinae as a well-supported monophyletic group, with *Leptidea* sister to the remaining six genera.

Coliadinae

In contrast to the preliminary analysis based on morphology by Venables (1993), our results (Table 5, Figs 3–6) support the conclusion of a number of other studies (Klots, 1933; Ehrlich, 1958; Geiger, 1981; Janz & Nylin, 1998; Pollock *et al.*, 1998; T. Yamauchi, O. Yata & A. Venables, unpubl. data) that recognize the Coliadinae as a natural or monophyletic group. de Jong *et al.* (1996) and Ackery *et al.* (1999) were unable to identify clear synapomorphies for their exemplar taxa, although T. Yamauchi, O. Yata & A. Venables (unpubl. data) noted two characters (patagia sclerotized; valvenansatz a short, narrow lobe) that appear to be diagnostic. The subfamily is almost certainly the sister

group to the Pierinae. Of the 18 genera currently recognized in the subfamily (Braby, 2005), only three (*Abaeis*, *Prestonia*, *Rhabdodryas*) were not included in our study. *Abaeis* Hübner contains two species restricted to North and Central America (Lamas, 2004); it was previously considered to be a subgenus of *Eurema* and may well belong to the New World *Eurema* and allied taxa (Klots, 1933). *Prestonia* Schaus is a monotypic genus, with type species *clarki* Schaus, from Mexico (Lamas, 2004); it was previously treated as a synonym of *Phoebis* Hübner (Klots, 1933). *Rhabdodryas* Godman & Salvin includes the single species *trite* (Linnaeus) from Central and South America (Lamas, 2004); Klots (1933) treated it as a subgenus of *Phoebis*.

Pierinae

The analyses of the combined and all available data provide strong support for the monophyly of the Pierinae (Table 5, Fig. 6). Of the 57 genera currently recognized in the subfamily (Braby, 2005), 51 were included in our study. On the basis of our analyses, together with morphological evidence (Klots, 1933), we propose a reclassification of the Pierinae and divide the subfamily into four rather than two main lineages. These lineages comprise two tribes (Anthocharidini *s.s.*, Pierini *s.s.*) and two informal groups (*Colotis* group, *Leptosia*), the interrelationships of which are unresolved (Fig. 7A).

Colotis group

We place eight genera, previously included in the Pierini *s.l.* (i.e. *Colotis*, *Eronia*, *Ixias*, *Gideona*, *Parenonia*, *Nepheronia*) or Anthocharidini *s.l.* (i.e. *Hebomoia*, *Pinacopteryx*), into an informal group termed the *Colotis* group. *Calopieris* Aurivillius, although not included in this study, presumably belongs here. It is a monotypic genus, with type species *eulimene* (Klug), restricted to areas adjacent to the Red Sea of Africa (Chad, Sudan, Arabia) (Ackery, Smith & Vane-Wright, 1995); it was previously regarded as a subgenus of *Colotis*, to which it is probably closely related. These nine taxa may well comprise a separate lineage sister to the rest of the Pierinae, but evidence for their monophyly is currently lacking (Fig. 6). As noted earlier, Klots (1933) regarded the first six genera to be phylogenetically unrelated to the other Pierini *s.l.* (Fig. 2), and this pattern is also evident in our analysis. The higher taxon name Teracolini, introduced by Reuter (1896) for the genus *Colotis*, is available (Bridges, 1988), but for the present we prefer not to recognize the group as a formal tribe without further evidence.

Klots (1933) described the monotypic genus *Gideona* endemic to Madagascar on the basis of its distinct genitalia, noting that it was probably related to *Colotis* or

possibly *Eronia* and *Nepheronia* + *Pareronia*. However, Lees, Kremen & Raharitsimba (2003), following Bernardi (1954), treated *Gideona* as a subgenus of *Colotis*. In our all available data analysis, *Gideona* appears to be more closely related to *Pinacopteryx* and *Eronia* + *Ixias*, although evidence supporting this arrangement is not convincing. Although the monophyly of *Nepheronia* + *Pareronia* is well supported, reinforcing Klots' view (Fig. 2) of a close relationship between these taxa, the systematic relationship of *Colotis* is not resolved.

Anthocharidini s.s.

In our *EF-1 α* (Figs 4, 5) and all available data (Fig. 6) analyses, the Anthocharidini, as delimited and classified by Klots (1933), are polyphyletic, with two Old World genera, the Afrotropical *Pinacopteryx* and the predominantly Oriental *Hebomoia*, falling outside the remaining genera. Indeed, Klots (1933: 174–175) stated that he 'does not regard the Euchloini as here delineated as being an entirely natural group' by inclusion of *Pinacopteryx* and *Hebomoia*. We thus narrow the concept of the Anthocharidini to include only seven genera (*Euchloe*, *Anthocharis*, *Zegris*, *Eroessa*, *Cunizza*, *Hesperocharis*, *Mathania*), the monophyly of which is extremely well supported in the combined and all available data analyses (Table 5, Fig. 6). Interestingly, the broad, straight subapical orange band present on the forewing in the Holarctic *Anthocharis* and *Zegris* (*Zegris*) of the *Anthocharis* group, also occurs in *Eroessa* of the Neotropical *Hesperocharis* group, and may be a synapomorphy for the tribe, with independent losses in *Euchloe* and the *Hesperocharis* subclade.

Leptosia group

Klots (1933) was uncertain about the phylogenetic position of *Leptosia*, noting that 'In none of its characters does it show any close relationship to any other modern Pieridae, but stands alone.' He went on further to state that '*Leptosia* appears to have no close relatives. It probably represents a derivative of a stock that split off far back on the Pierine line of development.' Klots' sentiments are clearly borne out in our all available data analysis in which the genus shows no close relatives other than belonging somewhere in the subfamily Pierinae (Fig. 6). The taxon possibly comprises a distinct lineage and may well warrant formal tribal status. However, for the moment we recognize it as an informal group within the Pierinae, pending further study of its exact relationships.

Pierini s.s.

We remove ten genera (in the *Colotis* group, and *Leptosia*) from the *Pierini* s.l.; otherwise the tribe is non-

monophyletic in the broad sense. The monophyletic *Pierini* s.s. (Table 5, Fig. 6) is distinguished from all other pierids by possession of pupal type II morphology. Five lineages are recognized in the tribe: three subtribes (Appiadina, Pierina, Aporiina) and two groups of uncertain status (*Elodina*, *Dixeia* + *Belenois*). The interrelationships of these lineages are largely unresolved, although *Elodina* is almost certainly sister to the remaining taxa (Fig. 7A).

Appiadina

We provisionally place four genera (*Saletara*, *Appias*, *Udaiana*, *Aoa*) in the subtribe Appiadina, introduced by Kusnezov (1921), based on strong evidence of monophyly in the combined and all available data analyses (Table 5, Fig. 6). Klots (1933) considered *Saletara*, *Appias* and *Udaiana* to be very closely related on morphological grounds (Fig. 2), and Yata (1985: 359) noted that *Saletara* 'should be phylogenetically included in the comprehensive genus *Appias*'. *Udaiana* Distant, not included in this study, contains a single species, *cynis* (Hewitson), known only from a restricted area in south-east Asia. *Aoa* was thought to be unrelated to these genera, although Klots (1933: 223) commented that 'Its exact relationships are... very obscure' and Yata (1985) noted that the butterfly is similar to *Appias* and *Cepora*. The genitalia of *Aoa* are remarkably similar to *Appias*, and our molecular evidence suggests a close relationship between these two taxa. The genus *Appias*, which includes five subgenera (*Appias*, *Hiposcritia*, *Catophaga*, *Phrissura*, *Glutophrissa*), is almost certainly paraphyletic. *Appias* (*Appias*), not included in this study, includes seven species restricted to the Oriental and Australian regions (Yata, 1985; Vane-Wright & de Jong, 2003). Although not well supported, the Appiadina may be sister to Pierina + Aporiina.

Pierina

We place 19 genera in the subtribe Pierina. Of these, 16 genera were included in our *EF-1 α* and all available data analyses, all of which formed an extremely well-supported monophyletic group (Figs 4–6). The three genera (*Glennia*, *Reliquia*, *Piercolias*) not included in our study presumably belong here according to morphological evidence. The monotypic genus *Glennia* Klots contains the species *pylotis* Godart from southern Brazil, but its systematic position is problematic, having affinities with either *Pieris* Schrank or *Pontia* Fabricius (Robbins & Henson, 1986). *Reliquia* Ackery is another monotypic genus, containing the species *santamarta* Ackery restricted to Sierra Nevada de Santa Marta of north-eastern Colombia (above 3000 m) (Ackery, 1975; Shapiro, 1978b). It has

affinities with the *Pieris* and *Tatochila* groups of genera, and is considered to have phylogenetic and biogeographical importance in understanding the evolution and radiation of the high Andean and Patagonian pierine fauna. *Piercolias* Staudinger contains three rare species from the high Andes of southern Peru and Bolivia, and belongs to the *Tatochila* group of genera; it is probably the sister genus of *Pierphulia* Field (Field, 1958; Field & Herrera, 1977).

Klots (1933: 218–219) was uncertain about the position of *Baltia* from the Himalayas, stating that, on the one hand it ‘probably represents a group, originally derived from *Synchloe* or some closely related stock’, but on the other hand ‘Whether there is a real relationship between *Baltia* and *Phulia* or whether the resemblances are merely to be regarded as similar developments, in the same type of environment is a matter of doubt’. Field (1958) placed *Baltia* in the *Tatochila* group of genera, the members of which predominantly inhabit the high Andes of South America. Although our analysis strongly supports the monophyly of the South American genera of the *Tatochila* group, it does not support a close relationship between these otherwise disjunct taxa, with *Baltia* being more closely related to *Pontia* (*Pontia*) + *Pontia* (*Synchloe*) from the Holarctic, as originally suggested by Klots. We tentatively conclude that the striking similarities in morphology between some members of *Baltia* (Central Asia) and the *Tatochila* group s.s. (South America) are due to convergence of living at extreme altitudes and not to common ancestry.

Aporiina

We place 14 genera (embracing around 480 species) in the subtribe *Aporiina*, first introduced by Chapman (1895) as a subfamily to distinguish genera such as *Aporia* and *Delias* from *Pieris*, but expand the concept of the taxon to include a larger number of genera. The monophyly of the group is well supported (Table 5, Fig. 6). Most of the genera are morphologically distinct (especially the early stages; M.F. Braby & K. Nishida, unpubl. data), biologically peculiar with the larvae of the vast majority of species feeding gregariously and producing considerable quantities of silk, and phylogenetically removed from the *Pierina*. Indeed, more than 25 years ago, the late John Eliot (in Corbet & Pendlebury, 1978, 1992) suggested that *Delias* together with the African *Mylothris* and South American *Catasticta*, *Archonias*, *Pereute*, and *Leodonta* probably form a distinct tribe. Chapman’s (1895) higher divisions of the Pieridae were based primarily on fundamental differences in pupal morphology, including the structure, shape (especially wing cases), and motility of segments. In the *Aporiina*, he noted that both abdominal segments 5 and 6 are movable

(when molested the abdomen twitches violently), whereas only abdominal segment 5 is movable in the *Coliadinae* and the *Pierina*, but no segments are movable in the *Anthocharidini*. Our combined analysis suggests that the well-supported *Aporiina* (Table 5, Fig. 6) may be the sister taxon to the *Pierina*.

Six major lineages are evident in the *Aporiina*: *Cepora*, *Prioneris*, *Mylothris*, *Aporia*, *Delias* group, and the *Catasticta* group. The last four taxa/subclades are very closely related (Braby, Pierce & Vila, 2006); they share a number of larval and adult morphological features, and the majority of species for which life histories are known feed as larvae on mistletoes in the order Santalales. It is not clear whether *Cepora* and *Prioneris* form a monophyletic group sister to (*Mylothris* + *Aporia* + *Delias* group + *Catasticta* group), or represent two independent lineages that diverged early in the evolution of the *Aporiina*.

Incertae sedis

The phylogenetic positions of three genera (*Elodina*, *Dixeia*, *Beleinois*) are uncertain in our combined and all available data analyses (Figs 3, 6) in that they do not belong to any of the higher taxa recognized above. Klots (1933) was equally uncertain about the relationships of the Australian *Elodina*, which appears to be the sister lineage to the rest of the *Pierini*. Klots (1933) suggested that the African *Dixeia* and *Beleinois* were closely related and probably allied to *Prioneris* (Fig. 2). The monophyly of *Dixeia* and *Beleinois* is corroborated in our combined analysis (Table 5), and these two genera probably constitute a separate subtribe within the *Pierini* (Fig. 7A).

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The age of divergence estimates generated by the two different methods based on fossils were approximate. Nevertheless, both estimates for the age of divergence between the *Pierina* (clade J) and *Aporiina* (clade K) were similar and around 60 Myr (Palaeocene): 61 Myr for penalized likelihood and 58 Myr for the quartet method. Moreover, the 99.9% confidence interval for the mean estimate under penalized likelihood was relatively small and precise (69–54 Myr). These observations indicate that the *EF-1 α* data are robust, clock-like and, if potential sources of error are assumed to be small, the age estimates may be accurate. The main assumptions and potential sources of error in the estimates are that the topology of the phylogram accurately represents evolutionary relationships of the extant taxa, the age of the fossil deposits have been dated accurately, and that the fossils have been identified and placed on the tree correctly. Our all available data analysis suggests that our phyloge-

netic hypothesis of the Pieridae is reasonably robust (Fig. 6), although additional or independent data (e.g. from morphological characters) would be desirable. The age of the deposit from which the two oldest fossils were described (Florissant Formation) has been accurately dated using the ^{40}Ar – ^{39}Ar decay method (Evanoff *et al.*, 2001). The identity of the fossils has been determined with a high degree of certainty at the subclade level (i.e. generic groups) (Table 3), but with less certainty among the extant genera within those groups. Given that fossils provide only minimum estimates of age, and that our estimates are conservative in that we calculated the fastest possible rate under penalized likelihood, the common ancestor of the Pierina + Aporiina is more likely to have originated before than during the Palaeocene. Clearly the ancestor of the Pieridae must be older than 60 Myr. Extrapolation of our phylogram from the node uniting the Pierina and Aporiina (Fig. 5) indicates that the stem-group of the family must have arisen well before the Tertiary. Indeed, our extrapolated mean estimate of 95 Myr (99.9% confidence interval: 112–82 Myr) for the crown-group of the Pieridae under penalized likelihood is in close agreement with the maximum age of 94–91 Myr estimated from the analysis of larval host plant associations and reconstruction of the ancestral host in relation to the maximum age of the plants (Braby & Trueman, 2006). These findings suggest a possible maximum origin of the Pieridae in the Cenomanian-Turonian of the Late Cretaceous.

Pierids occur worldwide but are not evenly distributed throughout the major zoogeographical regions (Table 7). In terms of taxonomic richness at the generic and subgeneric level, the Neotropical region has by far the highest diversity (46 taxa), whereas the Australian region has the lowest diversity (13 taxa). The Afrotropical and Palaearctic regions have similar totals in richness, but considerably less than the Oriental and Nearctic regions. More than two-thirds of the Neotropical fauna (70%) is endemic to the region at the generic/subgeneric level, and more than one-

third of the fauna in the Afrotropical region (42%) is endemic. In contrast, the large Holarctic region (Palaearctic and Nearctic) has a low level of endemism, with only five endemic taxa, of which two are restricted to the Himalaya (*Baltia*, *Aporia* (*Mesapia*)) and two to North America (*Eucheira*, *Neophasia*). The Australian region likewise has a very low level of endemism (15%), with *Leuciactria* and *Elodina* being the only taxa endemic to the region, with the latter genus extending as far west as Sulawesi in Wallacea. Although the Oriental region has a relatively high richness (second to the Neotropics), the level of endemism is comparatively low (29%), but much higher than that of the Australian, Nearctic, and Palaearctic. The Oriental and Australian regions, however, often share taxa because of frequent dispersal across Wallacea: combining the faunas for the two regions revealed a high level of endemism (55%), most of which is centred near the Old World tropics, although overall richness (29 taxa) is still low compared with the Neotropics.

Although the generic/subgeneric framework is incomplete for the Pieridae (Braby, 2005), it is unlikely that further improvements to the higher classification will affect the broad patterns enumerated in Table 7. South America clearly stands out as an area that has a highly distinctive fauna in terms of its composition, richness and endemism, whereas the Australian region has the most impoverished fauna. Moreover, four groups (Pseudopontiinae, Dismorphiinae, Anthocharidini, Pierina) are notably absent from Australia.

In terms of the higher taxa recognized in this work, the subfamilies, tribes, subtribes, and informal groups have markedly different distribution patterns among the six major zoogeographical regions. The Pseudopontiinae are endemic to Africa, whereas the Dismorphiinae are restricted mainly to the Neotropical region, with a disjunct occurrence in the Palaearctic. The Coliadinae, although especially well represented in the Oriental and New World faunas, are cosmopolitan. Indeed, two genera (*Eurema s.l.*,

Table 7. Comparison of taxonomic richness between the zoogeographical regions at the generic and subgeneric level (compiled from data listed in the Appendix)

Zoogeographical region	Number of taxa			No. of endemic taxa (%)
	Major occurrence	Minor occurrence	Total	
Neotropical	46	0	46	32 (70)
Oriental	20	8	28	8 (29)
Nearctic	15	8	23	2 (9)
Afrotropical	14	5	19	8 (42)
Palaearctic	14	4	18	3 (17)
Australian	8	5	13	2 (15)

Colias) are almost cosmopolitan. Within the Pierinae, the *Colotis* group and *Leptosia* occur in the Old World, mainly in the Afrotropical and Oriental regions, with weaker representations in the Australian and/or Palaearctic. The Anthocharidini s.s. are restricted geographically to the Neotropical and Holarctic regions (with a weak representation in Africa) and contain two monophyletic subclades, the *Hesperocharis* and *Anthocharis* groups, confined to each of these zoogeographical regions, respectively. Within the Pierini s.s., the Appiadina are pantropical, but the higher taxa (genera, subgenera) are concentrated in the Oriental region and few species occur in the other regions. The Aporiina, the putative sister group of the Pierina, are more strongly represented in the southern hemisphere (Neotropical, Australian, and to a lesser extent Afrotropical) and south-east Asia (Oriental) than in the Holarctic. In contrast, the Pierina are strongly represented in the Neotropical and Holarctic regions, with a weaker representation in the Afrotropical and Oriental. *Elodina* is restricted to the Australian region, with a very small representation in submontane eastern Indonesia of Wallacea (Vane-Wright & de Jong, 2003), whereas *Dixeia* and *Belenois* are predominantly African.

Phylogenetic relationships within most of these higher taxa are, in general, too poorly resolved to interpret broad historical biogeographical patterns. Also, given the strong migratory tendencies in the family, ancient vicariant patterns have probably long been obscured by subsequent dispersal events, and levels of differentiation may only be sought among closely related genera or species within genera. Moreover, few higher taxa are restricted to areas of endemism. The Pseudopontiinae and Dismorphiinae, however, are an exception. The phylogenetic hypothesis concerning the generic relationships of the clade Pseudopontiinae + Dismorphiinae is well supported and the terminal taxa are restricted either to the Afrotropical, Neotropical, or Palaearctic regions. These two subfamilies are therefore the most amenable of the higher taxa within the Pieridae to historical biogeographical analysis.

Pseudopontiinae–Dismorphiinae

Scott (1985: 261) hypothesized that the ancestor of Pseudopontiinae + Dismorphiinae evolved in Western Gondwana before Africa and South America finally split apart (100–90 Myr); the two subfamilies then diverged vicariantly following the break-up of the two continents. However, the disjunct geographical distribution of the Dismorphiinae has remained an outstanding biogeographical enigma in the Pieridae. To explain the presence of the Dismorphiinae in the Palaearctic, Scott suggested that *Leptidea* dispersed

across the Bering Strait to reach Eurasia. However, this scenario assumes at least four major biogeographical steps or costs [in the sense of Ronquist (1997)]: (1) dispersal from South America to North America, (2) extinction in South America, (3) dispersal from North America to Eurasia, and (4) extinction in North America. Dispersal is relatively easy to envisage in most pierid butterflies, but the extinction or range contraction of a whole genus from both North and South America is much harder to comprehend. Even if the ancestor of the Dismorphiinae expanded its range to the northern hemisphere before the split between *Leptidea* and the Neotropical Dismorphiinae, such that *Leptidea* and Neotropical Dismorphiinae diverged allopatrically in Eurasia and South America, respectively, this three-step hypothesis implies that the ancestor became extinct in North America but not in Europe.

An alternative vicariance hypothesis is that the ancestor of *Leptidea* reached Europe via northern Africa rather than via North America. According to our phylogeny, there are two major speciation events: (a) Pseudopontiinae (Africa) and Dismorphiinae (South America), and (b) Neotropical Dismorphiinae (northern South America) and *Leptidea* (northern Africa). The first speciation event may have occurred by vicariance, and the second event possibly through long-distance postspeciation dispersal. This hypothesis thus requires a minimum of three biogeographical steps: (1) long-distance dispersal of the ancestor of Dismorphiinae from northern South America to northern Africa followed by allopatric speciation of *Leptidea* in northern Africa; (2) dispersal (range expansion) of the ancestor of *Leptidea* from northern Africa to Eurasia; and (3) extinction (range contraction) of *Leptidea* in northern Africa (Fig. 8). *Leptidea* is currently not known from northern Africa, although Tennent (1996: 102) drew attention to the possibility that the genus may occur in the coastal regions. Step (2) probably involved simple range expansion following collision of the African plate with Eurasia during the early Tertiary (60 Myr), rather than long-distance dispersal. Step (3) probably involved range contraction following aridification of northern Africa with formation of the Sahara Desert after the Miocene. Subsequent differentiation of the Neotropical Dismorphiinae (at the generic level) in South America presumably represents a duplication event (sympatric speciation) within this area of endemism.

Scott's and our biogeographical hypotheses rest on the assumption that the ancestor of the Pseudopontiinae + Dismorphiinae originated in Western Gondwana, that is, when Africa and South America were still connected. Plate tectonic models show that the opening of the South Atlantic Ocean between Africa and South America started in the south in the

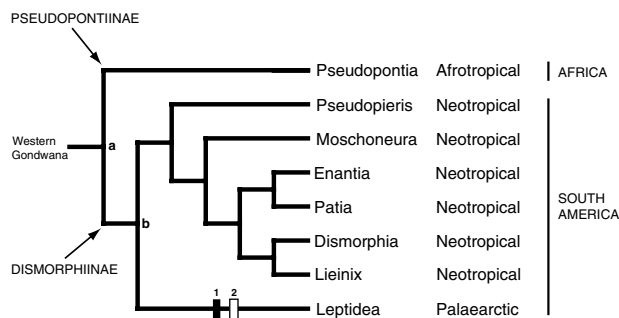


Figure 8. Historical biogeographical hypothesis of the Pseudopontiinae + Dismorphiinae, with dispersal and extinction events optimized to reconcile the area cladogram. Letters designate speciation events: a, vicariance between Pseudopontiinae (Africa) and Dismorphiinae (South America), following the final break-up of Western Gondwana (Late Cretaceous); b, long-distance dispersal of the ancestor of Dismorphiinae from northern South America to northern Africa (Late Cretaceous), followed by allopatric speciation of *Leptidea* in northern Africa (Late Cretaceous). Numbers designate major biogeographical events: 1, dispersal (range expansion) of the ancestor of *Leptidea* from northern Africa to Eurasia, following contact of Africa with Eurasia (early Tertiary); 2, extinction (range contraction) of *Leptidea* in northern Africa following formation of the Sahara Desert (Quaternary). Once *Leptidea* reached Eurasia it colonized much of the Palearctic, the Neotropical Dismorphiinae subsequently spread into Central America, whereas the Pseudopontiinae contracted to central western Africa.

Early Cretaceous (from *c.* 135 to 130 Myr) and propagated northwards until the mid- to Late Cretaceous (*c.* 110–90 Myr) when a transform fault opened between Guinea and Brazil, so that northern Western Gondwana separated much later than southern Western Gondwana (Smith, Smith & Funnell, 1994; White, 1994; Cox & Moore, 2000; Scotese, 2001; Sanmartín & Ronquist, 2004). Thus, if the ancestor of Pseudopontiinae + Dismorphiinae occurred in Western Gondwana such that the two subfamilies evolved under a process of vicariance, then the two lineages would have diverged sometime in the Late Cretaceous (> 90 Myr) at the very latest. This implies an origin of the Pieridae around the Late Cretaceous, which agrees well with our approximate estimates based on fossils (95 Myr) and larval host plant associations (94–91 Myr). The alternative scenario is that the speciation event occurred more recently through long-distance dispersal from South America to Africa, or vice versa. Although possible, this hypothesis is less parsimonious, as it requires an extra biogeographical step.

Whatever the true sequence of events and mode of speciation, the molecular and morphological diver-

gence between the Pseudopontiinae and Dismorphiinae is substantial (average corrected pairwise distance for *EF-1 α* = 30.5%), and no doubt reflects a long period of isolation between the two subfamilies. Moreover, a recent molecular phylogeny of *Leptidea* (Martin, Gilles & Descimon, 2003) suggests that *L. duponcheli* (endemic to the Mediterranean) is the sister taxon to the remaining species, most of which occur widely in the Palearctic, including Siberia (e.g. *L. sinapis*, *L. morsei*, *L. amurensis*). Such a biogeographical pattern is consistent with our hypothesis that *Leptidea* reached Europe from Africa and not from North America. We tentatively conclude that the Pseudopontiinae + Dismorphiinae originated in Western Gondwana, and that divergence of the two groups occurred by vicariance between South America and Africa, probably sometime during the Late Cretaceous.

Other taxa

Three other groups (Coliadae, Anthocharidini *s.s.*, *Tatochila* group) show interesting biogeographical patterns that also point towards an origin in South America/southern hemisphere. Although relationships within the Coliadae are not well resolved, it is curious that both *Nathalis* and *Kricogonia*, relictual taxa sister to the rest of the subfamily, are found only in the New World, especially Central and South America. Indeed, A. Shapiro (pers. comm.) has suggested that *Nathalis*, which has its main occurrence in the high altitudes of the Andes, probably originated in South America and colonized North America recently.

The Anthocharidini *s.s.* are restricted largely to the Neotropical and Holarctic regions. An origin of the tribe in South America would involve two major biogeographical steps: (1) long-distance dispersal or range expansion of the ancestor of the stem-group to North America, followed by differentiation of the *Anthocharis* and *Hesperocharis* groups in North and South America, respectively, and (2) dispersal (range expansion) of the *Anthocharis* group to Europe/northern Africa. However, a northern hemisphere origin in North America is equally parsimonious. The relictual distribution of *Eroessa*, which is limited to cool temperate rainforest (valdivian forest) of southern Chile–western Argentina (Shapiro, 1991; M.F. Braby & K. Nishida, unpubl. data), provides circumstantial evidence in favour of the first hypothesis. If correct, the timing of such events may date back to the early Tertiary (50–40 Myr) when North and South America joined and then separated again following formation of the Greater Antilles.

Within the subtribe Pierina (Pierini), the *Tatochila* group of genera (*Tatochila*, *Hypsochila*, *Theochila*, *Pierphulia*, *Phulia*, *Infraphulia*, and probably *Pierco-*

lias and *Reliquia*) comprises a well-supported monophyletic group. Theories concerning its origin and evolution in the high Andes of South America have long attracted attention (summarized by Shapiro, 1978a, 1994). It has generally been assumed that the ancestor of the *Tatochila* group dispersed from the Palaearctic/Holarctic to South America during the Great American Interchange 3–2 Myr, and then radiated explosively once it colonized the high Andes, which are young geologically. However, this hypothesis rests on the presumption that the *Tatochila* group is closely related to *Baltia* [which is limited to high altitudes (> 5000 m) in Central Asia] and/or *Pontia* from the northern hemisphere. Our molecular phylogeny shows that the nearest relatives of the *Tatochila* group are probably *Ascia* and *Ganyra*, and that the subclade *Baltia* + *Pontia* is more distantly related. *Ascia* and *Ganyra* are restricted to the New World, but have their major centre of distribution (i.e. in terms of both diversity and area of occurrence) in Central and South America. Hence, it is probable that lowland tropical Amazonia may have been the source for high-altitude colonization by the *Tatochila* group rather than stock from the northern hemisphere. Our phylogram (Fig. 5) supports the contention that the *Tatochila* group represents an example of rapid and probably recent radiation in the high Andes: the stem-group is subtended by a long branch and the crown-group shows little resolution, with the terminal taxa (six genera in our analysis) having very short branches, giving a long 'broom handle' pattern typical of explosive radiations (Crisp, Cook & Steane, 2004). The minimum mean estimate for the node (crown-group) of the *Tatochila* group under penalized likelihood was 10.4 Myr [$\lambda = 3000$; confidence interval (4 SD) = 14.5–7.3 Myr], which coincides with the time of initial uplift of the Andes.

DIRECTIONS FOR FUTURE WORK

Our study represents the first thorough phylogenetic study of the Pieridae and has provided a preliminary framework for higher-level classification of the family. However, many issues still require attention, and future systematic studies of the Pieridae might concentrate in the following three areas. (1) Among the higher taxa of the Pieridae, deep-level relationships are still poorly resolved, especially the relationship of the Coliadinae to the three other subfamilies. The Coliadinae are assumed to be sister to the Pierinae. (2) Within the subfamily Pierinae, the relationships of the four major lineages are not well understood, particularly the two informal groups (*Colotis* group, *Leptosia*), which may prove to constitute separate tribes. The *Colotis* group is envisaged to comprise nine genera (*Colotis*, *Calopieris*, *Eronia*, *Ixias*, *Gideona*,

Pareronia, *Nepheronia*, *Hebomoia*, *Pinacopteryx*), but further investigation is required to establish monophyly. (3) Within the tribe Pierini *s.s.*, the relationships of the five major lineages are also poorly resolved, particularly the phylogenetic positions of the genus *Elodina* and the subclade *Dixeia* + *Belenois*.

Reconstructing deep-level relationships is often fraught with difficulty because ancient divergence times inevitably result in considerable noise (homoplasy) among characters so that the phylogenetic signal is weak. Some of these issues may only be resolved by inclusion of data from other gene regions that are able to recover deeper-level splits. The analysis and integration of morphological characters (e.g. Wahlberg & Nylin, 2003; Wahlberg *et al.*, 2005), especially from immature stages, may also improve resolution and aid in the recognition of synapomorphies to help diagnose clades and further define the higher taxa proposed in this work.

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REFERENCES

- Ackery PR. 1975.** A new pierine genus and species with notes on the genus *Tatochila* (Lepidoptera: Pieridae). *Bulletin of the Allyn Museum* **30**: 1–9.
- Ackery PR. 1984.** Systematics and faunistic studies on butterflies. In: Vane-Wright RI, Ackery PR, eds. *The biology of butterflies. Symposium of the Royal Entomological Society of London, No. 11*. London: Academic Press, 11–21.
- Ackery PR, de Jong R, Vane-Wright RI. 1999.** The butterflies: Hedyloidea, Hesperioidea and Papilionoidea. In: Kristensen NP, ed. *Lepidoptera, moths and butterflies*, Vol. 1. *Evolution, systematics and biogeography*. Berlin: de Gruyter, 263–300.
- Ackery PR, Smith CR, Vane-Wright RI. 1995.** *Carcasson's African butterflies. An annotated catalogue of the Papilionoidea and Hesperioidea of the Afrotropical Region*. Melbourne: CSIRO Publishing.
- Ackery PR, Vane-Wright RI. 1984.** *Milkweed butterflies: their cladistics and biology*. London: British Museum (Natural History).
- Arbogast BS, Edwards SV, Wakeley J, Beerli P, Slowinski JB. 2002.** Estimating divergence times from molecular data on phylogenetic and population genetic timescales. *Annual Review of Ecology and Systematics* **33**: 707–740.
- Aurivillius C. 1910.** Pieridae. In: Seitz A, ed. *Macrolepidoptera of the world*, Vol. 13. Stuttgart: Alfred Kern, 29–69.
- Bernardi G. 1954.** Révision des Pierinae de la faune Malgache (Lepid. Pieridae). *Mémoires de l'Institut Scientifique de Madagascar Série E Entomologie* **5**: 239–375.
- Braby MF. 2005.** Provisional checklist of genera of the Pieridae (Lepidoptera: Papilionidae). *Zootaxa* **832**: 1–16.
- Braby MF, Pierce NE. 2006.** Systematics, biogeography and diversification of the Indo-Australian genus *Delias* Hübner (Lepidoptera: Pieridae): phylogenetic evidence supports an 'out-of-Australia' origin. *Systematic Entomology* (in press).
- Braby MF, Pierce NE, Vila R. 2006.** Phylogeny and historical biogeography of the subtribe Aporiina (Lepidoptera: Pieridae): implications for the origin of Australian butterflies. *Biological Journal of the Linnean Society* (in press).
- Braby MF, Trueman JWH. 2006.** Evolution of larval host plant associations and adaptive radiation in pierid butterflies. *Journal of Evolutionary Biology* (in press).
- Braby MF, Trueman JWH, Eastwood RG. 2005.** When and where did troidine butterflies (Lepidoptera: Papilionidae) evolve? Phylogenetic and biogeographic evidence suggests an origin in remnant Gondwanan in the Late Cretaceous. *Invertebrate Systematics* **19**: 113–143.
- Bremer K. 1988.** The limits of amino acid sequence data in angiosperm phylogenetic reconstruction. *Evolution* **42**: 795–803.
- Bremer K. 1994.** Branch support and tree stability. *Cladistics* **10**: 295–304.
- Bridges CA. 1988.** *Catalogue of Papilionidae and Pieridae (Lepidoptera: Rhopalocera)*. Urbana, Illinois: Charles A. Bridges.
- Brower AVZ. 2000.** Phylogenetic relationships among the Nymphalidae (Lepidoptera) inferred from partial sequences of the wingless gene. *Proceedings of the Royal Society of London B* **267**: 1201–1211.
- Brower AVZ, DeSalle R. 1998.** Patterns of mitochondrial versus nuclear DNA sequence divergence among nymphalid butterflies: the utility of wingless as a source of characters for phylogenetic inference. *Insect Molecular Biology* **7**: 73–82.
- Brower AVZ, Egan MG. 1997.** Cladistic analysis of *Heliconius* butterflies and relatives (Nymphalidae: Heliconiini): a revised phylogenetic position for *Eueides* based on sequences from mtDNA and nuclear gene. *Proceedings of the Royal Society of London B* **264**: 969–977.
- Brown FM. 1976.** *Oligodonta florissantensis*, gen. n., sp. nov. (Lepidoptera: Pieridae). *Bulletin of the Allyn Museum* **37**: 1–4.
- Brunton CFA. 1998.** The evolution of ultraviolet patterns in European *Colias* butterflies (Lepidoptera, Pieridae): a phylogeny using mitochondrial DNA. *Heredity* **80**: 611–616.
- Brunton CFA, Hurst GDD. 1998.** Mitochondrial DNA phylogeny of brimstone butterflies (genus *Gonepteryx*) from the Canary Islands and Madeira. *Biological Journal of the Linnean Society* **63**: 69–79.
- Butler AG. 1870.** A revision of the genera of the subfamily Pierinae. *Cistula Entomologica* **1**: 33–58.
- Campbell DL, Brower AVZ, Pierce NE. 2000.** Molecular evolution of the Wingless gene and its implications for the phylogenetic placement of the butterfly family Riodinidae (Lepidoptera: Papilionoidea). *Molecular Biology and Evolution* **17**: 684–696.
- Carpenter FM. 1992.** *Treatise on invertebrate paleontology, Part R. Arthropoda 4*, Vol. 4: *Superclass Hexapoda*. Boulder/Lawrence: Geological Society of America and University of Kansas.
- Caterino MS, Cho S, Sperling FAH. 2000.** The current state of insect molecular systematics: a thriving tower of babel. *Annual Review of Entomology* **45**: 1–54.
- Caterino MS, Reed RD, Kuo MM, Sperling FAH. 2001.** A partitioned likelihood analysis of swallowtail butterfly phylogeny (Lepidoptera: Papilionidae). *Systematic Biology* **50**: 106–127.
- Chapman TA. 1895.** Notes on butterfly pupae, with some remarks on the phylogenesis of the Rhopalocera. *Entomologist's Record and Journal of Variation* **6**: 101–107, 125–131, 147–152.
- Cheong SW. 1990.** *Comparative morphology and phylogeny on the female internal genitalia of the Pieridae (Lepidoptera)*. Taegu: Department of Biology, Kyungpook National University, 80.

- Cho S, Mitchell A, Regier JC, Mitter C, Poole RW, Friedlander TP, Zhao S. 1995.** A highly conserved nuclear gene for low-level phylogenetics: elongation factor-1 α recovers morphology-based tree for heliothine moths. *Molecular Biology and Evolution* **12**: 650–656.
- Clary DO, Wolstenholme DR. 1985.** The mitochondrial DNA molecule of *Drosophila yakuba*: nucleotide sequence, gene organization, and genetic code. *Journal of Molecular Evolution* **22**: 252–271.
- Clench HK. 1955.** Revised classification of the butterfly family Lycaenidae and its allies. *Annals of the Carnegie Museum* **33**: 261–274.
- Cooper A, Penny D. 1997.** Mass survival of birds across the Cretaceous–Tertiary boundary: molecular evidence. *Science* **275**: 1109–1113.
- Corbet AS, Pendlebury HM. 1978.** *The butterflies of the Malay Peninsula*. Kuala Lumpur: Malayan Nature Society.
- Corbet AS, Pendlebury HM. 1992.** *The butterflies of the Malay Peninsula*. Kuala Lumpur: Malayan Nature Society.
- Courtney SP. 1986.** The ecology of pierid butterflies: dynamics and interactions. *Advances in Ecological Research* **15**: 51–131.
- Cox CB, Moore PD. 2000.** *Biogeography: an ecological and evolutionary approach*. Oxford: Blackwell Science.
- Crisp MD, Cook LG, Steane D. 2004.** Radiation of the Australian flora: what can comparisons of molecular phylogenies across multiple taxa tell us about the evolution of diversity in present-day communities? *Philosophical Transactions of the Royal Society of London B* **359**: 1551–1571.
- D’Abrera B. 1971.** *Butterflies of the Australian region*. Melbourne: Lansdowne Press.
- D’Abrera B. 1980.** *Butterflies of the Afrotropical region*. East Melbourne: Lansdowne Editions.
- D’Abrera B. 1981.** *Butterflies of the Neotropical region, Part 1: Papilionidae and Pieridae*. Melbourne: Lansdowne Editions.
- D’Abrera B. 1982.** *Butterflies of the Oriental region, Part 1: Papilionidae, Pieridae and Danaidae*. Ferny Creek, Victoria: Hill House.
- D’Abrera B. 1990.** *Butterflies of the Holarctic region, Part 1: Papilionidae, Pieridae, Danaidae and Satyridae (Partim)*. Black Rock, Melbourne: Hill House.
- Danforth BN, Shuqing J. 1998.** Elongation factor-1 α occurs as two copies in bees: implications for phylogenetic analysis of EF-1 α sequences in insects. *Molecular Biology and Evolution* **15**: 225–235.
- Dixey FA. 1894.** On the phylogeny of the Pierinae, as illustrated by their wing-markings and geographical distribution. *Transactions of the Entomological Society of London Part 2*: 249–334.
- Dixey FA. 1932.** The plume-scales of the Pierinae. *Transactions of the Entomological Society of London* **80**: 57–75.
- Ehrlich PR. 1958.** The comparative morphology, phylogeny and higher classification of the butterflies (Lepidoptera: Papilionoidea). *University of Kansas Science Bulletin* **39**: 305–364.
- Ehrlich PR, Ehrlich AH. 1967.** The phenetic relationships of the butterflies. I. Adult taxonomy and the nonspecificity hypothesis. *Systematic Zoology* **16**: 301–317.
- Emmel TC, Minno MC, Drummond BA. 1992.** *Florissant butterflies. A guide to the fossil and present-day species of central Colorado*. Stanford: Stanford University Press.
- Evanoff E, McIntosh WC, Murphey PC. 2001.** Stratigraphic summary and 40Ar/39Ar geochronology of the Florissant Formation, Colorado. In: Evanoff E, Gregory-Wodzicki KM, Johnson KR, eds. *Fossil flora and stratigraphy of the Florissant Formation, Colorado*. Denver, Colorado: Denver Museum of Nature and Science, series 4, 1–16.
- Felsenstein J. 1985.** Confidence limits on phylogenies: an approach using the boot strap. *Evolution* **39**: 783–791.
- Felsenstein J. 1988.** Phylogenies from molecular sequences: inference and reliability. *Annual Review of Genetics* **22**: 521–565.
- Field WD. 1958.** A redefinition of the butterfly genera *Tatochila*, *Phulia*, *Piercolias*, and *Baltia*, with descriptions of related genera and subgenera. *Proceedings of the United States National Museum* **108**: 103–131.
- Field WD, Herrera JV. 1977.** The pierid butterflies of the genera *Hypsochila* Ureta, *Phulia* Herrich-Schaffer, *Infra-phulia* Field, *Pierphulia* Field, and *Piercolias* Staudinger. *Smithsonian Contributions to Zoology* **232**: 1–64.
- Folmer O, Black M, Hoeh W, Lutz R, Vrijenhoek R. 1994.** DNA primers for amplification of mitochondrial cytochrome c oxidase subunit I from diverse metazoan invertebrates. *Molecular Marine Biology and Biotechnology* **3**: 294–299.
- Freitas AVL, Brown KSJ. 2004.** Phylogeny of the Nymphalidae (Lepidoptera). *Systematic Biology* **53**: 363–383.
- Friedlander TP, Horst KR, Regier JC, Mitter C, Peigler RS, Fang QQ. 1998.** Two nuclear genes yield concordant relationships within Attacini (Lepidoptera: Saturniidae). *Molecular Phylogenetics and Evolution* **9**: 131–140.
- Geiger HJ. 1981.** Enzyme electrophoretic studies on the genetic relationships of pierid butterflies (Lepidoptera: Pieridae) I. European taxa. *Journal of Research on the Lepidoptera* **19**: 181–195.
- Giribet G. 2003.** Stability in phylogenetic formulations and its relationship to nodal support. *Systematic Biology* **52**: 554–564.
- Grote AR. 1900.** The descent of the pierids. *Proceedings of the American Philosophical Society* **39**: 4–67.
- Guindon S, Gascuel O. 2003.** A simple, fast, and accurate algorithm to estimate large phylogenies by maximum likelihood. *Systematic Biology* **52**: 696–704.
- Harvey DJ. 1987.** The higher classification of the Riodinidae. PhD Thesis, University of Texas, Austin.
- Hedges SB, Kumar S. 2003.** Genomic clocks and evolutionary timescales. *Trends in Genetics* **19**: 200–206.
- Hillis DM, Mable BK, Larson A, Davis SK, Zimmer EA. 1996a.** Nucleic acids IV. Sequencing and cloning. In: Hillis DM, Moritz C, Mable BK, eds. *Molecular systematics*. Sunderland: Sinauer Associates, 321–384.
- Hillis DM, Mable BK, Moritz C. 1996b.** Applications of molecular systematics: the state of the field and a look to the future. In: Hillis DM, Moritz C, Mable BK, eds. *Molecular systematics*. Sunderland: Sinauer Associates, 515–543.
- Janz N, Nylin S. 1998.** Butterflies and plants: a phylogenetic study. *Evolution* **52**: 486–502.

- de Jong R, Vane-Wright RI, Ackery PR. 1996.** The higher classification of butterflies (Lepidoptera): problems and prospects. *Entomologica Scandinavica* **27**: 65–101.
- Kamie K, Taira H, Ooura H, Kakuta A, Matsumoto S, Ejiri S-i, Katsumata T. 1993.** Nucleotide sequence of the cDNA encoding silk gland elongation factor 1 α . *Nucleic Acids Research* **21**: 742.
- Kandul NP, Lukhtanov VA, Dantchenko AV, Coleman JWS, Sekercioglu CH, Haig D, Pierce NE. 2004.** Phylogeny of *Agrodiaetus* Hübner 1822 (Lepidoptera: Lycaenidae) inferred from mtDNA sequences of *COI* and *COII* and nuclear sequences of *EF-1 α* : karyotype diversification and species radiation. *Systematic Biology* **53**: 278–298.
- Kemp DJ, Rutowski RL, Mendoza M. 2005.** Colour pattern evolution in butterflies: a phylogenetic analysis of structural ultraviolet and melanic markings in North American sulphurs. *Evolutionary Ecology Research* **7**: 133–141.
- Klots AB. 1929.** The generic status of *Catopsilia* Hübner and *Phoebus* Hübner, with a discussion of the relationships of the species and the homologies of the male genitalia (Lepidoptera, Pieridae). *Bulletin of the Brooklyn Entomological Society* **24**: 203–214.
- Klots AB. 1933.** A generic classification of the Pieridae (Lepidoptera) together with a study of the male genitalia. *Entomologica America* **12**: 139–242.
- Knapp S, Lamas G, Lughadha EN, Novarino G. 2004.** Stability or stasis in the names of organisms: the evolving codes of nomenclature. *Philosophical Transactions of the Royal Society of London B* **359**: 611–622.
- Kristensen NP. 1976.** Remarks on the family-level phylogeny of butterflies (Insecta, Lepidoptera, Rhopalocera). *Zeitschrift für Zoologische Systematik und Evolutionsforschung* **14**: 25–33.
- Kusnezov NJ. 1921.** On 'taxonomic conceptions' and some attempts of their foundation on morphological data. *Revue Russe d'Entomologie* **17**: 53–80.
- Lamas G. 2004.** Pieridae. In: Lamas G, ed. *Checklist: Part 4a Hesperioidea–Papilionoidea*. In: Heppner JB, ed. *Atlas of Neotropical Lepidoptera*, Vol. 5A. Gainesville: Association for Tropical Lepidoptera/Scientific Publishers, 99–117.
- Lanave C, Preparata G, Saccone C, Serio G. 1984.** A new method for calculating evolutionary substitution rates. *Journal of Molecular Evolution* **20**: 86–93.
- Lees DC, Kremen C, Raharitsimba T. 2003.** Classification, diversity, and endemism of the butterflies (Papilionoidea and Hesperioidea): a revised species checklist. In: Goodman SM, Benstead JP, eds. *The natural history of Madagascar*. Chicago: University of Chicago Press, 762–793.
- Lukhtanov VA. 1991.** Evolution of the karyotype and system of higher taxa of the Pieridae (Lepidoptera) of the world fauna. *Entomologicheskoe Obozrenie* **70**: 619–641.
- Maddison WP, Maddison DR. 1999.** *MacClade: analysis of phylogeny and character evolution*, Version 3.08a for Macintosh. Sunderland: Sinauer Associates.
- Maddison WP, Maddison DR. 2001.** *MacClade: analysis of phylogeny and character evolution*, Version 4.03. Sunderland: Sinauer Associates.
- Magallón SA. 2004.** Dating lineages: molecular and paleontological approaches to the temporal framework of clades. *International Journal of Plant Sciences* **165**: S7–S21.
- Martin J-F, Gilles A, Descimon H. 2003.** Species concepts and sibling species: the case of *Leptidea sinapis* and *Leptidea reali* (Lepidoptera). In: Boggs CL, Watt WB, Ehrlich PR, eds. *Butterflies: ecology and evolution taking flight*. Chicago: University of Chicago Press, 459–476.
- Miller JS. 1987.** Phylogenetic studies in the Papilioninae. *Bulletin of the American Museum of Natural History* **186**: 365–512.
- Mitchell A, Cho S, Regier JC, Mitter C, Poole RW, Matthews M. 1997.** Phylogenetic utility of elongation factor-1 α in Noctuoidea (Insecta: Lepidoptera): the limits of synonymous substitution. *Molecular Biology and Evolution* **14**: 381–390.
- Mitchell A, Mitter C, Regier JC. 2000.** More taxa or more characters revisited: combining data from nuclear protein-encoding genes for phylogenetic analysis of Noctuoidea (Insecta: Lepidoptera). *Systematic Biology* **49**: 202–224.
- Monteiro A, Pierce NE. 2001.** Phylogeny of *Bicyclus* (Lepidoptera: Nymphalidae) inferred from *COI*, *COII*, and *EF-1 α* gene sequences. *Molecular Phylogenetics and Evolution* **18**: 264–281.
- Morinaka S, Miyata T, Tanaka K. 2002.** Molecular phylogeny of the *Eichhorni* group of *Delias* Hübner, 1819 (Lepidoptera: Pieridae). *Molecular Phylogenetics and Evolution* **23**: 276–287.
- Mosher E. 1969.** *Lepidoptera pupae. Five collected works on the pupae of North American Lepidoptera*. East Lansing: Entomological Reprint Specialists.
- Palumbi SR. 1996.** Nucleic acids II: the polymerase chain reaction. In: Hillis DM, Moritz C, Mable BK, eds. *Molecular systematics*, 2nd edn. Sunderland: Sinauer Associates, 205–247.
- Penz CM, Peggie D. 2003.** Phylogenetic relationships among Heliconiinae genera based on morphology (Lepidoptera: Nymphalidae). *Systematic Entomology* **28**: 451–479.
- Pollock DD, Watt WB, Rashbrook VK, Iyengar EV. 1998.** Molecular phylogeny for *Colias* butterflies and their relatives (Lepidoptera: Pieridae). *Annals of the Entomological Society of America* **91**: 524–531.
- Posada D, Crandall KA. 1998.** MODELTEST: testing the model of DNA substitution. *Bioinformatics* **14**: 817–818.
- Quek S-P, Davies SJ, Itino T, Pierce NE. 2004.** Codiversification in an ant-plant mutualism: stem texture and the evolution of host use in *Crematogaster* (Formicidae: Myrmicinae) inhabitants of *Macaranga* (Euphorbiaceae). *Evolution* **58**: 554–570.
- Rand DB, Heath A, Suderman T, Pierce NE. 2000.** Phylogeny and life history evolution of the genus *Chrysoritis* within the Aphaeini (Lepidoptera: Lycaenidae), inferred from mitochondrial cytochrome oxidase I sequences. *Molecular Phylogenetics and Evolution* **17**: 85–96.
- Reed RD, Sperling FAH. 1999.** Interaction of process partitions in phylogenetic analysis: an example from the swallowtail butterfly genus *Papilio*. *Molecular Biology and Evolution* **16**: 286–297.

- Reuter E. 1896.** Ueber die Palpen der Rhopaloceren. Ein Beitrag zur Erkenntnis der verwandtschaftlichen Beziehungen unter den Tagfaltern. *Acta Societatis Scientiarum Fennicae* **22**: 1–577.
- Robbins RK. 1982.** How many butterfly species? *News of the Lepidopterists' Society* **3**: 40–41.
- Robbins RK. 1988.** Comparative morphology of the butterfly foreleg coxa and trochanter (Lepidoptera) and its systematic implications. *Proceedings of the Entomological Society of Washington* **90**: 133–154.
- Robbins RK, Henson PM. 1986.** Why *Pieris rapae* is a better name than *Artogeia rapae* (Pieridae). *Journal of the Lepidopterists' Society* **40**: 79–92.
- Röber JKM. 1908–09.** Pieridae section. In: Seitz A, ed. *Macrolepidoptera of the world*, Vol. 5. Stuttgart: Alfred Kernen, 53–111.
- Rodríguez F, Oliver JL, Marín A, Medina JR. 1990.** The general stochastic model of nucleotide substitution. *Journal of Theoretical Biology* **142**: 485–501.
- Roger AJ, Sandblom O, Doolittle WF, Philippe H. 1999.** An evaluation of Elongation Factor 1 α as a phylogenetic marker for eukaryotes. *Molecular Biology and Evolution* **16**: 218–233.
- Ronquist F. 1997.** Dispersal-vicariance analysis: a new approach to the quantification of historical biogeography. *Systematic Biology* **46**: 195–203.
- Ronquist F, Huelsenbeck JP. 2003.** MrBayes 3: Bayesian phylogenetic inference under mixed models. *Bioinformatics* **19**: 1572–1574.
- Sanderson MJ. 2002.** Estimating absolute rates of molecular evolution and divergence times: a penalized likelihood approach. *Molecular Biology and Evolution* **19**: 101–109.
- Sanmartín I, Ronquist F. 2004.** Southern hemisphere biogeography inferred by event-based models: plant versus animal patterns. *Systematic Biology* **53**: 216–243.
- Saux C, Fisher BL, Spicer GS. 2004.** Dracula ant phylogeny as inferred by nuclear 28S rDNA sequences and implications for ant systematics (Hymenoptera: Formicidae: Amblyoponinae). *Molecular Phylogenetics and Evolution* **33**: 457–468.
- Schmitz J, Moritz RFA. 1994.** Sequence analysis of the D1 and D2 regions of 28S rDNA in the hornet (*Vespa crabro*) (Hymenoptera, Vespinae). *Insect Molecular Biology* **3**: 273–277.
- Scotese CR. 2001.** *Atlas of Earth history*. Arlington: University of Texas.
- Scott JA. 1985.** The phylogeny of butterflies (Papilionoidea and Hesperioidea). *Journal of Research on the Lepidoptera* **23**: 241–281.
- Scudder SH. 1875a.** Fossil butterflies. *Memoirs of the American Association for the Advancement of Science* **1**: 1–99.
- Scudder SH. 1875b.** Historical sketch of the generic names proposed for butterflies. *Proceedings of the American Academy of Arts and Sciences* **10(2d Series, Vol. 2)**: 89–294.
- Scudder SH. 1889.** The fossil butterflies of Florissant. *US Geological Survey, 8th Annual Report Part I*, 433–474.
- Sequeira AS, Normark BB, Farrell BD. 2000.** Evolutionary assembly of the conifer fauna: distinguishing ancient from recent associations in bark beetles. *Proceedings of the Royal Society of London Series B* **267**: 2359–2366.
- Sequencher. 1995.** *Sequencher*, Version 3.0 for Macintosh. Ann Arbor, Michigan: Gene Codes Corporation.
- Sequencher. 2000.** *Sequencher*, Version 4.1.2 for Macintosh. Ann Arbor, Michigan: Gene Codes Corporation.
- Shapiro AM. 1978a.** The life history of an equatorial montane butterfly, *Tatochila xanthodice* (Lepidoptera: Pieridae). *New York Entomological Society* **86**: 51–55.
- Shapiro AM. 1978b.** The life history of *Reliquia santamarta*, a Neotropical alpine pierine butterfly (Lepidoptera: Pieridae). *New York Entomological Society* **86**: 45–50.
- Shapiro AM. 1991.** The zoogeography and systematics of the Argentine Andean and Patagonian Pierid fauna. *Journal of Research on the Lepidoptera* **28**: 137–238.
- Shapiro AM. 1994.** Why are there so few butterflies in the high Andes? *Journal of Research on the Lepidoptera* **31**: 25–56.
- Shields O. 1976.** Fossil butterflies and the evolution of Lepidoptera. *Journal of Research on the Lepidoptera* **15**: 132–143.
- Sikes DS, Lewis PO. 2001.** *Pauprat: Paup* implementation of the parsimony ratchet*. Storrs: Department of Ecology and Evolutionary Biology, University of Connecticut.
- Simon C, Frati F, Beckenbach A, Crespi BJ, Liu H, Flook P. 1994.** Evolution, weighting, and phylogenetic utility of mitochondrial gene sequences and a compilation of conserved polymerase chain reaction primers. *Annals of the Entomological Society of America* **87**: 651–701.
- Smith AG, Smith DG, Funnell BM. 1994.** *Atlas of Mesozoic and Cenozoic coastlines*. Cambridge: Cambridge University Press.
- Sorenson MD. 1999.** *Treerot*, Version 2. Boston: Boston University.
- Sperling FAH. 2003.** Butterfly molecular systematics: from species definitions to higher-level phylogenies. In: Boggs CL, Watt WB, Ehrlich PR, eds. *Butterflies: ecology and evolution taking flight*. Chicago: University of Chicago Press, 431–458.
- Stavenga DG, Stowe S, Siebke K, Zeil J, Arikawa K. 2004.** Butterfly wing colours: scale beads make white pierid wings brighter. *Proceedings of the Royal Society of London B* **271**: 1577–1584.
- Swofford DL. 2002.** *PAUP*. Phylogenetic analysis using parsimony (*and other methods)*, Version 4.0b10 for Macintosh. Sunderland: Sinauer Associates.
- Talbot G. 1935.** Pieridae III. In: Aurivillius C, Wagner H, Strand E, eds. *Lepidopterorum catalogus*. Berlin: Dr W. Junk, 385–697.
- Talbot G. 1939.** *The fauna of British India, including Ceylon and Burma butterflies*, Vol. I. London: Taylor & Francis.
- Tautz D, Hancock JM, Webb DA, Tautz C, Dover GA. 1988.** Complete sequences of the rRNA genes of *Drosophila melanogaster*. *Molecular Biology and Evolution* **5**: 366–376.
- Tennent WJ. 1996.** *The butterflies of Morocco, Algeria and Tunisia*. Wallingford: Gem Publishing.
- Thompson JD, Gibson TJ, Plewniak F, Jeanmougin F, Higgins DG. 1997.** The ClustalX windows interface: flexible

- strategies for multiple sequence alignment aided by quality analysis tools. *Nucleic Acids Research* **24**: 4876–4882.
- Vane-Wright RI. 2003.** Evidence and identity in butterfly systematics. In: Boggs CL, Watt WB, Ehrlich PR, eds. *Butterflies: ecology and evolution taking flight*. Chicago: University of Chicago Press, 477–513.
- Vane-Wright RI, de Jong R. 2003.** The butterflies of Sulawesi: annotated checklist for a critical island fauna. *Zoologische Verhandelingen, Leiden* **343**: 3–267.
- Venables BAB. 1993.** Phylogeny of the white and sulphur butterflies (Pieridae). Dissertation project proposal. Made available by Dr R. K. Robbins, National Museum of Natural History, Smithsonian Institution.
- Wahlberg N, Braby MF, Brower AZ, de Jong RM-ML, Nylin S, Pierce NE, Sperling FAH, Vila RU, Warren AD, Zakharov E. 2005.** Synergistic effects of combining morphological and molecular data in resolving the phylogeny of butterflies and skippers. *Proceedings of the Royal Society of London B* **272**(1572): 1577–1586.
- Wahlberg N, Nylin S. 2003.** Morphology versus molecules: resolution of the positions of *Nymphalis*, *Polygonia*, and related genera (Lepidoptera: Nymphalidae). *Cladistics* **19**: 213–223.
- Wahlberg N, Weingartner E, Nylin S. 2003.** Towards a better understanding of the higher systematics of Nymphalidae (Lepidoptera: Papilionoidea). *Molecular Phylogenetics and Evolution* **28**: 473–484.
- Watt WB, Donohue K, Carter PA. 1996.** Adaptation at specific loci. VI. Divergence vs. parallelism of polymorphic allozymes in molecular function and fitness-component effects among *Colias* species (Lepidoptera, Pieridae). *Molecular Biology and Evolution* **13**: 699–709.
- Weller SJ, Friedlander TP, Martin JA, Pashley DP. 1992.** Phylogenetic studies of ribosomal RNA variation in higher moths and butterflies (Lepidoptera: Ditrysia). *Molecular Phylogenetics and Evolution* **1**: 312–337.
- Weller SJ, Pashley DP. 1995.** In search of butterfly origins. *Molecular Phylogenetics and Evolution* **4**: 235–246.
- Weller SJ, Pashley DP, Martin JA. 1996.** Reassessment of butterfly family relationships using independent genes and morphology. *Annals of the Entomological Society of America* **89**: 184–192.
- Weller SJ, Pashley DP, Martin JA, Constable JL. 1994.** Phylogeny of noctuid moths and the utility of combining independent nuclear and mitochondrial genes. *Systematic Biology* **43**: 194–211.
- White ME. 1994.** *After the greening, the browning of Australia*. East Roseville, NSW: Kangaroo Press.
- Yata O. 1985.** Part 1: Pieridae. In: Tsukada E, ed. *Butterflies of the South East Asian islands. II: Pieridae, Danaidae*. Tokyo: Plapac, 33–120, 205–438.
- Yoshimoto H. 2000.** Wood whites (Pieridae), the most primitive pierids? *Butterflies* **26**: 52–59.
- Zakharov EV, Caterino MS, Sperling FAH. 2004.** Molecular phylogeny, historical biogeography, and divergence time estimates for swallowtail butterflies of the genus *Papilio* (Lepidoptera: Papilionidae). *Systematic Biology* **53**: 193–215.
- Zeuner FE. 1942.** Two new fossil butterflies of the family Pieridae. *Annals and Magazine of Natural History* **9**: 409–416.

APPENDIX

Zoogeographical distributions of the Pieridae [higher classification according to this study; lower classification according to Braby (2005)]. Zoogeographical distributions are divided into two categories: ‘major’ and ‘minor’, the latter category refers to taxa that are poorly represented in the region in terms of numbers of species relative to other ‘major’ region(s) and/or that have a very small area of occurrence relative to other region(s). The eastern boundary of the Oriental region is set along Wallace’s Line, whereas the western boundary of the Australian region is set along Lydekker’s Line. *Aoa*, the only taxon endemic to Wallacea, the intervening area between Wallace’s Line and Lydekker’s Line, was categorized as ‘Oriental’ because of its restricted occurrence in Sulawesi, close to Wallace’s Line. Distributions are based primarily on D’Abrera (1971, 1980, 1981, 1982, 1990), supplemented with other regional faunistic works for specific continents.

Taxon	Zoogeographical distribution	
	Major	Minor
Subfamily Pseudopontiinae		
<i>Pseudopontia</i>	Afrotropical	
Subfamily Dismorphiinae		
<i>Leptidea</i>	Palearctic	
<i>Pseudopieris</i>	Neotropical	
<i>Moschoneura</i>	Neotropical	
<i>Dismorphia</i>	Neotropical	
<i>Lieinix</i>	Neotropical	
<i>Enantia</i>	Neotropical	Nearctic
<i>Patia</i>	Neotropical	

APPENDIX *Continued*

Taxon	Zoogeographical distribution	
	Major	Minor
Subfamily Coliadinae		
<i>Nathalis</i>	Neotropical, Nearctic	
<i>Kricogonia</i>	Neotropical, Nearctic	
<i>Eurema</i>	Neotropical, Nearctic, Oriental, Australian, Afrotropical	
Subgen. <i>Terias</i>	Oriental, Australian	Afrotropical
<i>Abaeis</i>	Neotropical, Nearctic	
<i>Pyrisitia</i>	Neotropical	Nearctic
<i>Teriocolias</i>	Neotropical	
<i>Leucidia</i>	Neotropical	
<i>Gandaca</i>	Oriental, Australian	
<i>Gonepteryx</i>	Palaeartic	Oriental
<i>Dercas</i>	Oriental	
<i>Phoebis</i>	Neotropical, Nearctic	
<i>Prestonia</i>	Neotropical	
<i>Rhabdodryas</i>	Neotropical	
<i>Aphrissa</i>	Neotropical	Nearctic
<i>Catopsilia</i>	Oriental, Australian	Afrotropical
<i>Anteos</i>	Neotropical	Nearctic
<i>Colias</i>	Palaeartic, Nearctic, Neotropical	Afrotropical, Oriental
<i>Zerene</i>	Nearctic, Neotropical	
Subfamily Pierinae		
<i>Colotis</i> group		
<i>Colotis</i>	Afrotropical	Palaeartic, Oriental
<i>Calopieris</i>	Afrotropical	
<i>Eronia</i>	Afrotropical	
<i>Ixias</i>	Oriental	
<i>Pinacopteryx</i>	Afrotropical	
<i>Gideona</i>	Afrotropical	
<i>Hebomoia</i>	Oriental	Palaeartic
<i>Nepheronia</i>	Afrotropical	
<i>Pareronia</i>	Oriental	Australian
Tribe Anthocharidini s.s.		
<i>Euchloe</i>	Palaeartic, Nearctic	Afrotropical
<i>Anthocharis</i>	Palaeartic, Nearctic	
<i>Zegris</i>	Palaeartic	Nearctic
Subgen. <i>Microzegris</i>	Palaeartic	
<i>Eroessa</i>	Neotropical	
<i>Cunizza</i>	Neotropical	
<i>Hesperocharis</i>	Neotropical	
<i>Mathania</i>	Neotropical	
<i>Leptosia</i> group		
<i>Leptosia</i>	Afrotropical, Oriental	Australian
Tribe Pierini s.s.		
Subtribe Appiadina		
<i>Saletara</i>	Oriental	Australian
<i>Appias</i>	Oriental	Australian
Subgen. <i>Catophaga</i>	Oriental, Australian	
Subgen. <i>Hiposcritia</i>	Oriental	
Subgen. <i>Glutophrissa</i>	Afrotropical, Neotropical	Nearctic
Subgen. <i>Phrissura</i>	Oriental	
<i>Udaiana</i>	Oriental	
<i>Aoa</i>	Oriental	

APPENDIX *Continued*

Taxon	Zoogeographical distribution	
	Major	Minor
Subtribe Pierina		
<i>Pieris</i>	Palaeartic, Nearctic	Oriental, Afrotropical
<i>Talbotia</i>	Oriental	
<i>Glennia</i>	Neotropical	
<i>Leptophobia</i>	Neotropical	
<i>Itaballia</i>	Neotropical	
<i>Pieriballia</i>	Neotropical	
<i>Perrhybris</i>	Neotropical	
<i>Pontia</i>	Palaeartic, Nearctic, Afrotropical	
Subgen. <i>Synchloe</i>	Nearctic, Palaeartic	
<i>Baltia</i>	Palaeartic	
<i>Ganyra</i>	Neotropical	Nearctic
<i>Ascia</i>	Neotropical	Nearctic
<i>Reliquia</i>	Neotropical	
<i>Tatochila</i>	Neotropical	
<i>Hypsochila</i>	Neotropical	
<i>Theochila</i>	Neotropical	
<i>Piercolias</i>	Neotropical	
<i>Pierphulia</i>	Neotropical	
<i>Phulia</i>	Neotropical	
<i>Infraphulia</i>	Neotropical	
Subtribe Aporiina		
<i>Cepora</i>	Oriental	Australian
<i>Prioneris</i>	Oriental	
<i>Mylothris</i>	Afrotropical	
<i>Aporia</i>	Palaeartic	Oriental
Subgen. <i>Metaporis</i>	Palaeartic	Oriental
Subgen. <i>Mesapia</i>	Palaeartic	
<i>Delias</i>	Australian, Oriental	Palaeartic
<i>Leuciactria</i>	Australian	
<i>Melete</i>	Neotropical	
<i>Pereute</i>	Neotropical	
<i>Leodonta</i>	Neotropical	
<i>Neophasia</i>	Nearctic	
<i>Eucheira</i>	Nearctic	
<i>Catasticta</i>	Neotropical	
<i>Archonias</i>	Neotropical	
<i>Charonias</i>	Neotropical	
incertae sedis		
<i>Elodina</i>	Australian	
<i>Dixeia</i>	Afrotropical	
<i>Belenois</i>	Afrotropical	Oriental, Australian, Palaeartic