Morphology, molecules and fritillaries: approaching a stable phylogeny for Argynnini (Lepidoptera: Nymphalidae)

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> We examine the phylogenetic relationships among 29 species of Argynnini based on 141 previously published morphological characters and new data from the mitochondrial gene COI and the two nuclear genes EF-1a and wingless. We investigate the stability and robustness of the resulting phylogenetic hypotheses through various combinations of the 4 functionally separate datasets. Increasing the number of datasets in combined analyses led to increased support for clades, sometimes substantially. We find that the tribe Argynnini is a well-supported, robust, monophyletic clade with the following internal subtribal structure: (Euptoietina (Yrameina (Boloriina Argynnina))). Our analyses support the classification of argynnine species into six robust and stable genera: Euptoieta, Yramea, Boloria, Issoria, Brenthis and Argynnis. We suggest that for moderate amounts of data, a total evidence approach is always

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Introduction

Researchers working on the evolutionary history of a group of species are interested in generating robust phylogenetic hypotheses for their taxon. A robust phylogenetic hypothesis can be seen as one that does not change when new characters are added to the data matrix. When inferring phylogenetic hypotheses, the combined analysis of data from various sources is commonly considered to lead to the most robust hypothesis (Kluge 1989). Since the molecular revolution in systematics, large amounts of new data have been generated by finding and sequencing new gene regions, although positive effects of combining morphological data with molecular data have been reported (Miller et al. 1997, Baker & Gatesy 2002, Wahlberg & Nylin 2003, Wahlberg et al. 2005). Combining different kinds of data into a single analysis is still debated, but it is now understood that separate analysis of biologically delimited data sets is a way of investigating the strength of the phylogenetic signal apparent in the total evidence analysis (DeSalle & Brower 1997; Gatesy et al. 1999).

Here we investigate the relative contributions of morphological characters and DNA sequences from three gene regions to the pattern of phylogenetic relationships among butterflies in the tribe Argynnini of the subfamily Heliconiinae. We are specifically interested in the effects of adding new molecular data to an already published morphological dataset (Simonsen 2006a). Do we really need to add data from 20 gene regions, as suggested by Rokas et al. (2003), to arrive at a robust phylogenetic hypothesis for this tribe of butterflies?

Phylogenetic assessments of relationships among various clades in the nymphalid subfamily Heliconiinae have been increasing over the last few years. The work of Penz & Peggie (2003) established a clear hypothesis for the higher level diversification and relationships of the major lineages in the subfamily, although problems still exist. Based on a large morphological data set, Penz & Peggie (2003) divided Heliconiinae into four tribes: Acraeini, Heliconiini, Vagrantini and Argynnini. Although these clades did not receive strong bootstrap support, they do conform to the intuitive groupings used by many authors prior to that study.

Most recent studies of relationships of taxa in Heliconiinae have concentrated on the tribe Heliconiini (Brower 1994; Brower 1997; Brower & Egan 1997; Penz 1999), but lately some attention has been given to the mainly Holarctic tribe Argynnini. Since the precladistic works of Warren (1944, 1955), Dos Passos & Grey (1945) and Warren et al. (1946) and the early cladistic or systematic works of Shirôzu & Saigusa (1973, 1975) and Higgins (1975), only Aubert et al. (1996) have dealt with the group in a modern cladistic way, using an outgroup and computer analyses. However, two morphology based phylogenetic studies of the tribe Argynnini and its subtribes have recently been concluded (Simonsen 2005, 2006a), and both morphology and DNA-based phylogenetic studies of the Nearctic subgenus Speveria are in preparation (J. Dunford, pers. comm.). The only molecule based phylogenetic study of the group so far (Aubert et al. 1996) was limited to western Palaearctic taxa, and the results should be considered preliminary (H. Descimon, pers. comm.). No attempt to combine morphological and molecular data has hitherto been published for the tribe Argynnini.

The Argynnini comprise 100+ species, historically placed in up to 19 genera, although currently placed in 6 genera (Simonsen 2006a). Almost all species are found in temperate, arctic and/or alpine areas, mainly in the Palaearctic and Nearctic biomes. A few species are found in the high Andes of South America, or the mountains of East Africa, and a single species (*Argynnis hyperbius*) occurs widely from Japan to Australia to Eastern Africa. There are two previously published studies with taxon sampling relevant to our study, both based on morphological data. Penz & Peggie (2003) sampled 18 species of Argynnini which they clas-

sified in 12 genera, and found that the enigmatic genus *Euptoieta* was nested within the tribe Argynnini. Simonsen (2006a) found *Euptoieta* to be the sister-group to the rest of the Argynnini, and that there were additionally two distinct clades, which were termed the subtribes Yramiina and Argynnina.

In this contribution we present a phylogenetic hypothesis of the Argynnini at the species group level, based on the adult genitalia and wing morphology from Simonsen (2006a) as well as the mitochondrial gene *cytochrome oxidase subunit I* (COI) and the nuclear genes *elongation factor-1* α (EF-1 α) and *wingless*.

Material and methods

Taxon sampling

29 ingroup taxa representing all major groups within the tribe (17 "genera") and 7 heliconiine outgroup taxa used by Simonsen (2006a) are included in the analysis of the morphological dataset. A complete list of the species including specimen data is given in Table 1. The morphological character matrix and character list can be found in Simonsen (2006a).

Preparation and terminology

Morphological characters. – Preparation techniques for morphological characters are found in Simonsen (2006a).

Taxonomy. – We initially adopt the argynnine classification proposed by Simonsen (2006a) but propose some changes based upon the current results.

Molecular characters. – We extracted DNA either from one or two legs or from the thorax musculature of freshly frozen, dried or alcohol conserved butterflies using QIAgen's DNEasy extraction kit.

For each of the 36 species we sequenced COI, EF-1 α and *wingless*. Primers for COI were taken from Wahlberg & Zimmermann (2000), for EF-1 α from Peña *et al.* (2006), and for *wingless* from Brower and DeSalle (1998). We performed all PCRs in a 20-ml reaction volume. The cycling profile for both COI and *wingless* was 95°C for 5 min, 35 cycles of 94°C for 30 s, 47°C for 30 s, 72°C for 1 min 30 s, and a final extension period of 72°C for 10 min. For EF-1 α the cycling profile was 95°C for 7 min, 35 cycles 95°C for 1 min,

55°C for 1 min, 72°C for 2 min and a final extension period of 72°C for 10 min. For all three generegions the PCR primers were also used for sequencing. Sequencing was done with a Beckman-Coulter CEQ8000 capillary sequencer (Stockholm University) or an ABI Prism 377 DNA sequencer (University of Leiden). The resulting chromatograms were verified using the program BioEdit (Hall 1999) and the sequences were aligned by eye. The sequences are available on GenBank (Accession numbers in Table 1).

Phylogenetic analyses

Phylogenetic analyses were carried out in TNT 1.0 (Goloboff et al. 2003) using maximum parsimony and a heuristic search algorithm. Molecular data were equally weighted and unordered; morphological data were coded as in Simonsen (2006a). Heuristic searches were run with 1000 randomaddition replicates using TBR branch swapping. The four datasets were analysed separately and in combination. The effects of adding new data to a published phylogenetic dataset (Simonsen 2006a) was analysed by sequentially adding the molecular datasets to the morphological dataset in all combinations, including a total evidence analysis with all four datasets combined in a single analysis. The molecular data were also combined and analysed without the morphological data.

Robustness of the clades in the resulting cladograms was evaluated with Bremer support values (BS) (Bremer 1988; Bremer 1994). The scripting feature of TNT was used to calculate BS values (see Peña et al. 2006). The contribution of each data partition to the BS values of the combined analyses was assessed using partitioned Bremer support (Baker & DeSalle 1997; Baker et al. 1998; Gatesy et al. 1999) using another script in TNT (scripts available from N. Wahlberg). Basic sequence statistics and genetic distances were calculated in MEGA2 (Kumar et al., 2001). The distribution of homoplasy within and among partitions was examined with the ILD statistic (Mickevich & Farris 1981). The Genbank accession codes for the gene sequences from the different species are shown in Table 1.

Results

General

Of the 141 morphological characters from Simon-

sen (2006a), 125 were parsimony informative with the current taxon sampling.

1450 bp from COI, 1240 bp from EF-1 α and 400 bp from wingless were sequenced for all species. The nucleotide base frequencies for each gene are given in Table 2. The AT bias for COI is in accordance with the general AT bias for insect mt genes (DeSalle et al., 1987; Liu & Beckenbach 1992, Simon et al. 1994). The tree statistics for each individual data partition and the combined sets (including tree lengths, number of trees and total PBS contribution) are given in Table 3. Almost all homoplasy is intrinsic to individual data partitions, rather than due to incongruence among partitions when they are combined. Between 48% and 67% of tree lengths for the analyses of individual partitions are due to homoplastic character state transformations, while the combination of the three gene regions adds 1%, and the addition of the morphological data to the combined DNA data adds another 1%.

Phylogenetic patterns in Argynnini

The analyses of each data partition both separately and combined in various ways revealed stable patterns of relationships (Figs 1-4). All analyses recover the following clades: Euptoieta, Yramea, Boloria, Brenthis and the clade comprising all Argynnini except *Euptoieta*. The only analysis in which the monophyly of the tribe Argynnini is not resolved is when COI is analysed on its own (Fig. 1B). The monophyly of the subtribe Argynnina is not resolved when morphology and the wingless are analysed separately (Figs 1A, D), nor when these two are combined (Fig. 2C), but the subtribe is found to be monophyletic in all other analyses. Monophyly of *Issoria* is likewise unresolved in the separate analyses of morphology and wingless (Figs 1A, D), but is found to be monophyletic in the combined analysis of these two partitions (Fig. 2C), as well as in all other analyses. The genus Argynnis (sensu Simonsen, 2006a) is not found to be monophyletic in the analysis of the EF-1 α partition (Fig. 1C), and its monophyly is unresolved in the analysis of wingless (Fig. 1 D), and in the combined analyses of morphology+EF-1α and morphology+EF- 1α +wingless (Figs 2B, F).

The combined molecular dataset (1 tree, 3318 steps) recovers the Argynnini as monophyletic (Fig. 3) and only differs from the total evidence analysis in the internal phylogeny of a subordinate

Table 1. A taxonomic list of the species and origin of the specimens used in the present analysis. Ingroup taxa listed to subgenera following Simonsen (2006a) with generic names in bold. BMNH: Natural History Museum, London, Great Britain. RMNH: National Museum of Natural History, Leiden, The Netherlands. ZMUC: The Natural History Museum of Denmark (Zoology), Copenhagen, Denmark.

Species	Locality DNA specimen	Locality Morphology specimen	Museum	Genbank Accession numbers COI EF-1α	sion numbers EF-1α	wingless
Outgroup Cupha erymanthis (Drury, 1773) Cupha prosope (Fabricius, 1775) Phalanta phalama (Drury, 1773) Cethosia cyalippe (Linnaeus, 1767) Cethosia cyane (Drury, 1773) Vindula arsinoe (Cramer, 1777) Agraulis vanillae (Linnaeus, 1758) Dryas iulia (Fabricius, 1775) Heliconius charitonia (Linnaeus, 1767) Heliconius hecale (Fabricius, 1775)	Australia Tanzania Bangladesh Australia USA Costa Rica	Java Nyasaland, Nigeria Java Philippines, Malaysia Cuba, Brazil Brazil Panama, Venezuela	o+\$-ZMUC o+\$-ZMUC o+\$-ZMUC o+\$-ZMUC o+\$-ZMUC o+\$-ZMUC	DQ922839 DQ922840 DQ922870 AY090204 DQ922841 DQ922842 AY090202	DQ922871 DQ922872 DQ922902 AY090170 DQ922873 DQ922874 AY090168	DQ922808 DQ922809 DQ922838 AY090137 AF169921 DQ922810
Ingroup						
Subtribe Argynnina Argynnis s.l. Argynnis paphia (Linnaeus, 1758) Argyreus hyperbius (Linnaeus, 1763) Argyronome laodice (Pallas, 1771) Argyronome ruslana, Motschulsky, 1866 Childrena childreni (Gray, 1831) Damora sagana (Doubleday, 1847) Fabriciana adippe (Denis & Schiff, 1775) Fabriciana niobe (Linnaeus, 1758) Fabriciana kamala (More, 1875)	Sweden Japan Russia Japan China Japan Finland Denmark	Denmark China, India Estonia Japan Sikkim, Assam Russia, Japan Denamrk Mashmir	O+9-ZMUC O+9	AY090200 DQ922843 DQ922844 DQ922845 DQ922849 DQ922850 DQ922851 DQ922853	AY090166 DQ922875 DQ922876 DQ922877 DQ922881 DQ922882 DQ922883 DQ922883	AY090133 DQ922811 DQ922812 DQ922813 DQ922813 DQ922817 DQ922819 DQ922819 DQ922819
Messacidatia agidja (Linnaeus, 1738) Nephargynnis anadyomene (Feld. & Feld., 1862) Spandoriana pandora (Denis & Schiff., 1775) Speedic cybele (Fabricius, 1775) Brenthis	Sweden Japan Russia USA	Denmark Japan France, Spain USA	o+o-ZMUC o+o-ZMUC o+o-ZMUC	DQ922860 DQ922861 DQ922862 DQ922863	DQ922892 DQ922893 DQ922894 DQ922895	DQ922828 DQ922829 DQ922830 DQ922831
Brenthis daphne (Denis & Schiff., 1775) Brenthis ino (Rottemburg, 1775) Brenthis hecate (Denis & Schiff., 1775) Isonia	Spain Finland Kirgisia	France, Romania Denmark Hungary	°+9-ZMUC °+9-ZMUC °+9-ZMUC	DQ922848 DQ922847 DQ922846	DQ922880 DQ922879 DQ922878	DQ922816 DQ922815 DQ922814
Issoria lathonia (Linnaeus, 1758) Issoria eugenia Eversmann, 1847 Issoria smaragdifera (Butler, 1895) Issoria hamningtoni Elwes, 1889	France Russia Tanzania Tanzania	Denmark Nepal Nysaland, Nigeria British East Africa	σ+φ-ZMUC σ+φ-RMNH σ+φ-BMNH σ+φ-BMNH	DQ922854 DQ922857 DQ922855 DQ922856	DQ922886 DQ922889 DQ922887 DQ922888	DQ922822 DQ922825 DQ922823 DQ922824

92

32

Subtribe Boloriina						
Boloria s.l.						
Boloria aquilonaris (Stichel, 1908)	Denmark	Denmark	♂+ç-ZMUC	DQ922867	DQ922899	DQ922835
Boloria pales (Denis & Schiff., 1775)	Italy	Switzerland	♂+ç-ZMUC	DQ922866	DQ922898	DQ922834
Clossiana euphrosyne (Linnaeus, 1758)	Spain	Denmark	o'+o-ZMUC	DQ922868	DQ922900	DQ922836
Clossiana selene (Denis & Schiff., 1775)	Sweden	Denmark	o'+o-ZMUC	AY090201	AY090167	AY090134
Proclossiana eunomia (Esper, 1799)	Spain	Norway, Switzerland	♂+ç-ZMUC	DQ922869	DQ922901	DQ922837
Subtribe Yrameiina						
Yramea						
Yramea cytheris (Drury, 1773)	Chile	Argentina	♂+ç-ZMUC	DQ922858	DQ922890	DQ922826
Yramea inca (Staudinger, 1894)	Peru	Peru	♂+ç-ZMUC	DQ922859	DQ922891	DQ922827
Subtribe Euptoietina						
Euptoieta						
Euptoieta claudia (Cramer, 1775)	USA	USA	♂+ç-ZMUC	DQ922864	DQ922896	DQ922832
Euptoieta hegesia (Cramer, 1779)	Brazil	Costa Rica, Cuba	♂+ç-ZMUC	DQ922865	DQ922897	DQ922833

Table 2. Empirical base frequencies (%) for the three molecular data sets in 36 species of Heliconiinae included in the analysis.

	Empiri	Empirical base frequencies (%)					
Gene	A	G	С	T			
COI	31.2	13.6	14.5	40.6			
EF-1 α	26.8	23.5	26.1	23.6			
wingless	25.1	28.0	25.2	21.7			

group within *Argynnis*, where *A. pandora* is found to be the sister taxon of a group comprising *A. paphia*, *A. sagana*, *A. anadyomene*, *A. laodice*, *A. ruslana*, *A. hyperbius* and *A. childrena*, in which (*A. paphia A. sagana*) in turn are found to comprise the sister group of (*A. anadyomene* (*A. laodice A. ruslana*)).

The total evidence analysis yielded one most parsimonious tree of 3724 steps (Fig. 4). Compared to the molecules only analysis, the support for almost all nodes in the total evidence tree is increased, sometimes substantially (Fig. 5). Adding morphological data decreases the BS at only four nodes: BS for the two nodes within the *Issoria* clade is decreased by 3, support for the sister relationship of *A. hyberbius* and *A. childreni* is decreased by 1, and the support for the sister relationship of Argynnina and Bolorina is decreased by 1. There is a clear trend of increasing BS values for the major clades (genera and subtribes) as additional characters are added to the combined analyses (Fig. 5).

In the total evidence analysis (Fig. 4), the Argynnini are monophyletic with high Bremer support (BS 15). The first split within the Argynnini is between *Euptoieta* and the remaining in-group genera. The latter clade is well supported (BS 20). *Euptoieta* is very strongly supported with a very high BS (50). *Yramea* is also very strongly supported with a very high BS (87). The sister group of *Yramea*, a clade comprising *Boloria* and Argynnina, is fairly well supported with a BS of 7. *Boloria* appears very strongly supported with a BS of 18. Within *Boloria* s.l., the position of *B. eunomia* is unstable to method of analysis, being sister to *Boloria* s. str. (BS 5). *Boloria* s. str. is strongly supported by a BS value of 37.

The Argynnina is monophyletic and well supported with a high BS of 10. Within Argynnina, *Issoria* s.l. is the sister group of a clade comprising *Brenthis* and *Argynnis*. *Issoria* s.l. is well sup-

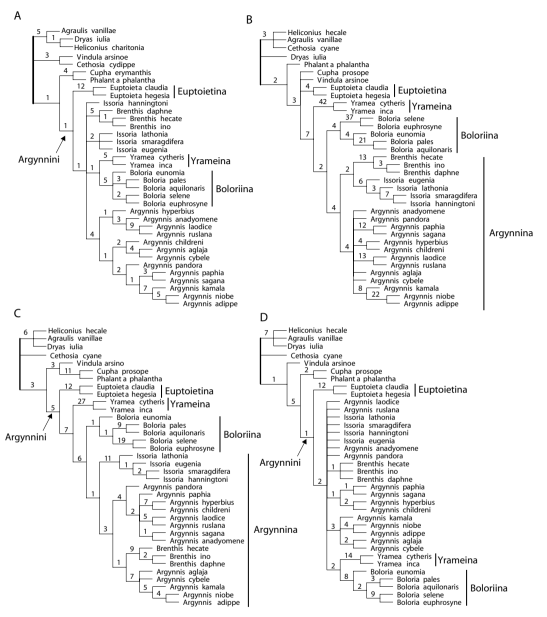


Fig. 1. Strict consensus trees from the separate analyses of the four datasets. **A** Morphology (8 trees, each 376 steps with CI = 0.41 and RI = 0.69), **B** COI (26 trees, each 2127 steps, CI = 0.36, RI = 0.43), **C** EF-1 α (4 trees, each 772 steps, CI = 0.52, RI = 0.68), **D** wingless (6 trees, each 388 steps, CI = 0.51, RI = 0.63). The numbers above the nodes are Bremer support values.

ported by a high BS of 26. Within *Issoria*, *I. eugenia* is the sister group of the remainder of the genus, which is modestly supported with a BS of 4. *I. lathonia* is the sister group of the two African

species *I. smaragdifera* and *I. hanningtoni*, a clade modestly supported (BS 5). The clade comprising *Brenthis* and *Argynnis* has a fairly high BS of 8. *Brenthis* is well supported (BS 26).

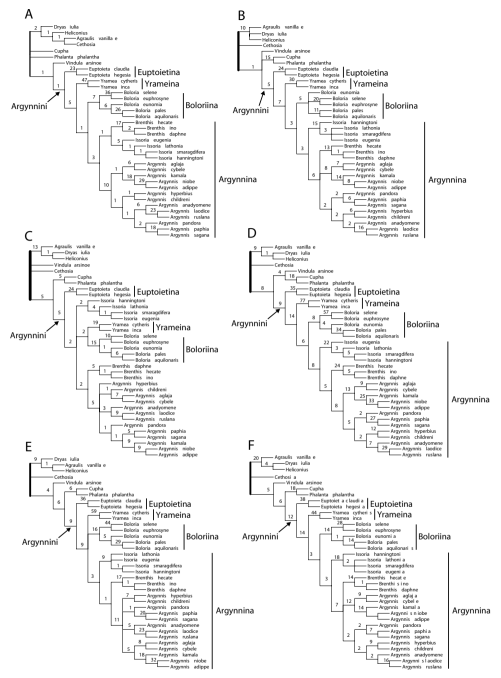


Fig. 2. Strict consensus trees from various combinations of the four datasets. A Morphology and COI (2 trees, each 2544 steps, CI = 0.36 and RI = 0.48), **B** Morphology and EF-1 α (12 trees, each 1179 steps, CI = 0.47 and RI = 0.67), **C** Morphology and *wingless* (8 trees, each 778 steps, CI = 0.46 and RI = 0.66), **D** Morphology, COI and EF-1 α (1 tree, 3327 steps, CI = 0.40 and RI = 0.53), **E** Morphology, COI and *wingless* (6 trees, each 2940 steps, CI = 0.38 and RI = 0.50), **F** Morphology, EF-1 α and *wingless* (3 trees, each 1573 steps, CI = 0.48 and RI = 0.66). The numbers above the nodes are Bremer support values.

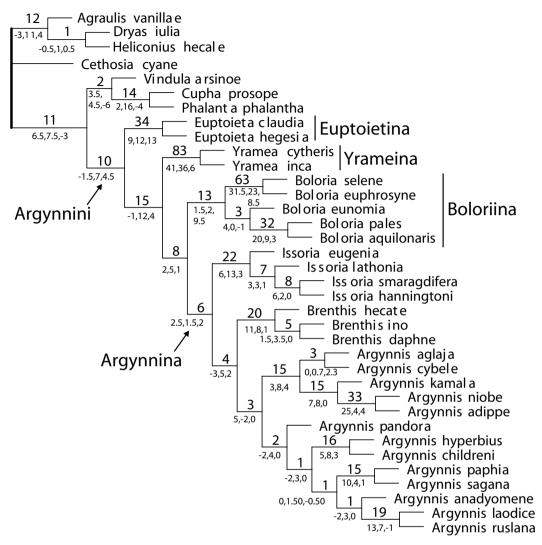


Fig. 3. The combined analysis of the three molecular datasets. The single most parsimonious tree (3318 steps, CI = 0.41, RI = 0.52). The numbers above the nodes are Bremer support values, whereas the numbers below the nodes are partitioned Bremer support values yielded by COI, EF- 1α and wingless respectively.

Argynnis s. l. as defined by Simonsen (2006a) is reasonably well supported and has a fairly high BS (8). The first split within Argynnis is between a clade comprising the two subgenera Fabriciana (represented by A. kamala, A. niobe and A. adippe) and Speyeria (represented by A. aglaja and A. cybele) and a clade comprising the remaining Argynnis. The Fabriciana+Speyeria clade is well supported with a very high BS (17). The clade comprising the two representatives of subgenus

Speyeria is well supported and the BS (11) is high. The subgenus Fabriciana is well supported and the BS (25) is very high. The clade comprising the remaining Argynnis is only moderately supported (BS 5). The basal split within this clade is between a clade comprising A. pandora, A. sagana and A. paphia and a clade comprising A. hyperbius, A. childreni, A. anadyomene, A. laodice and A. ruslana. The latter is poorly supported with a low BS (2). Within this clade, A. hyperbius and A. chil-

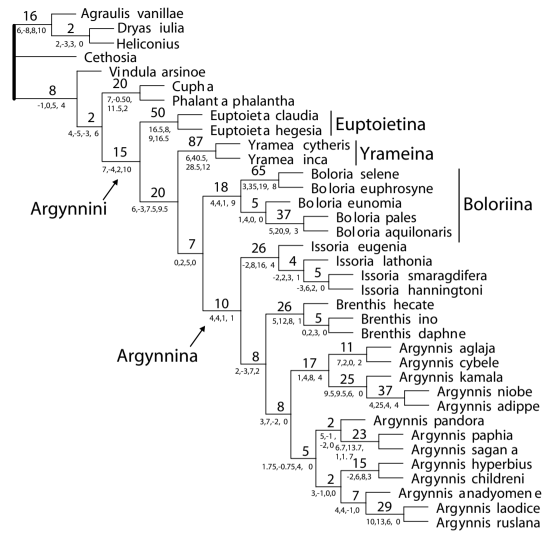


Fig. 4. The combined analysis of all four datasets. The single most parsimonious tree (3724 steps, CI = 0.41, RI = 0.54). The numbers above the nodes are Bremer support values, whereas the numbers below the nodes are partitioned Bremer support values yielded by morphology, COI, EF-1 α and *wingless* respectively.

dreni comprise the sister group of A. anadyomene, A. laodice and A. ruslana. A. hyperbius and A. childreni form a well supported clade with a high BS (15). The clade comprising A. anadyomene, A. laodice and A. ruslana is well supported and the BS (7) is fairly high. A. laodice and A. ruslana (subgenus Argyronome) form a strongly supported clade (BS 29). The clade comprising A. pandora, A. sagana and A. paphia is weakly supported, with a BS of 2, while the clade comprising A. sagana

and A. paphia is well supported with a BS of 24.

The positive partitioned total support values show that all datasets contributed significantly to the final result (Table 3). However, the datasets were conflicting or ambiguous at some nodes (Fig. 4). Of 28 nodes, 12 were unanimously supported by all datasets. In 4 cases the three molecular datasets unanimously supported the node, whereas the morphology was in conflict (3 of these nodes are in the *Issoria* clade). There was no node for

increasing support with sample size

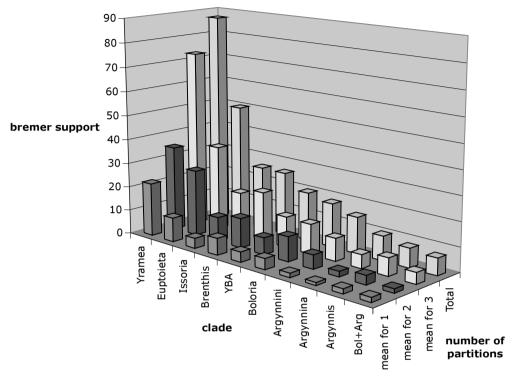


Fig. 5. The development of Bremer support values at major nodes as the number of partitions increase. Bol+Arg = Boloriina+Argynnina, YBA = Yrameina+Boloriina+Argynnina.

which the three molecular datasets unanimously yielded negative PBS. In one case (the position of *A. pandora* as sister to *A. paphia* and *A. sagana*) only morphology supported a node (the molecular datasets were either conflicting or ambiguous). The extremely small number of extra steps entailed by combining the molecular data partitions, and subsequently the molecular and morphological partitions (Table 3), also gives a clear indication of the high degree of congruence among these different sources of evidence.

Discussion

Combining data to increase support

The question of how much data is needed to arrive at a robust phylogenetic hypothesis is still debated (Rokas et al. 2003, Gatesy & Baker 2005). In our

study we have shown that increasing the number of functionally separate datasets also increases the stability and robustness of the resulting phylogenetic hypothesis. All four datasets have considerable positive impact on the results of the combined analysis. By sequentially adding the new molecular datasets to the already published morphological dataset, we have been able to discover which clades are very stable (i.e. unlikely to change no matter how much new data we add), which clades tend to stabilise as new data are added (Bremer supports increase substantially in combined analyses) and which clades are still relatively uncertain.

When analysed separately, each of the datasets show incongruence, both among one another and with the combined result. However, this is likely to be caused by the intrinsic properties of each finite dataset, i.e. the number of variable characters is

Table 3. Parameters for the individual and combined data partitions. D homoplasy is the additional steps due to incongruence among data partitions when they are combined. The total support is the sum of PBS values across all branches of the combined tree for a given partition.

Data partition	No. Char.	Variable Char.	Informative Chars	No. of Trees	Total Tree length	Intrinsic homo- plasy	D homo- plasy	Total support in TE tree
Morphology COI Ef-1a Wingless Comb. DNA Tot. evidence	141 1450 1240 400 3090 3231	140 550 305 136 991 1131	125 421 225 112 233 248	8 26 4 6 1	376 2127 772 388 3318 3724	222 (59%) 1361 (64%) 371 (48%) 190 (49%)	31 (1%) 30 (1%)	123.4 202.4 177.5 113.7 493.6 617

limited and homoplasy is high (as in all of our datasets). Combining the various datasets allows the underlying phylogenetic signal to come out (Gatesy et al. 1999, Baker & Gatesy 2002). In our analyses, this phenomenon is striking in the total evidence analysis (Fig. 5). Recent studies have suggested that incongruence between molecular datasets may be due to real differences in evolutionary histories of genes (e.g. Holland et al. 2004, Jeffroy et al. 2006). Such studies have taken advantage of very large datasets that are not yet tractable for many groups of organisms. Our study is typical of most studies today (about 3000 characters) and we suggest that in such cases a total evidence approach is not only appropriate but also necessary. It is likely that instances of real incongruence will only be reliably inferred with larger numbers of independent datasets.

In our case, by combining morphological characters and sequences from three genes, a more resolved, well-supported phylogenetic hypothesis is inferred than when any of the data partitions are analyzed alone. As shown in Fig. 5, the support for all higher clades increased as character sampling increased. Our study emphasizes the desirability of analyzing large datasets to arrive at robust phylogenetic hypotheses. Single gene studies, particularly using short mitochondrial gene regions, should be viewed with circumspection (Brower 2006).

Phylogeny

Although the present results are largely in agreement with those of Simonsen (2006a), some very important differences are obvious and deserve attention. There is agreement that Argynnini are monophyletic and that the subtribe Euptoietina

comprises the sister group of the remaining taxa. The placement of *Yramea* as the sister group of *Boloria* + Argynnina, however, conflicts with the earlier results. In Simonsen (2006a), *Yramea* was sister of *Boloria*, and these two genera were placed in a subtribe named Yrameina. However, according to our current results this name should be reserved only for the genus *Yramea*. The internal phylogeny of *Boloria*, where *B. eunomia* is the sister group of (*B. pales B. aquilonaris*) is in conflict with Simonsen (2006a) where *B. eunomia* was placed as the sister group of (*B. selene B. euphrosyne*), but is congruent with the morphology based analysis focused on *Boloria* by Simonsen (2005).

This analysis corroborates the hypothesis that the subtribe Argynnina is monophyletic. However, in contrast to Simonsen (2006a), we found that the genus *Issoria* s.l. is monophyletic. In part due to the absence of the 'rectal plate' in two African species (I. baumanni and I. hanningtoni), Simonsen (2006a) found that these two species formed the sister group to the rest of Argynnina and they were thus placed in the genus Prokuekenthaliella. Our results here suggest that the two African species are nested well within Issoria. The rectal plate is a very complex structure associated with the tegumen and uncus in Palaearctic Issoria and the African species I. smaragdifera (Simonsen 2006b), but absent in other Heliconiinae. Given the simple structure of the uncus and tegumen in *I*. hanningtoni and I. baumanni the present results seem to indicate that these structures (and hence the rectal plate) have been secondarily reduced in these two species compared to other Issoria.

There is agreement that *Brenthis* form the sister group of a monophyletic *Argynnis* s.l.. The genus *Brenthis* is well supported here though the internal

relationships of the genus differ from Simonsen (2006a).

Monophyly of Argynnis s.l. is moderately well supported, but the internal phylogeny of the genus contradicts the results of Simonsen (2006a), where a clade comprising (A. hyperbius (A. anadyomne (A. laodice A. ruslana))) is the sister group of the remaining species in the genus. In our total evidence analysis the first split in Argynnis is between a clade comprising the two subgenera Fabriciana and Speveria (including Mesoacidalia) and a clade comprising the remaining subgenera. Though contradicting Simonsen (2006a), a close relationship between Fabriciana and Speveria was suggested by Penz & Peggie (2003) and here the clade is well supported by all datasets. Although not found by Simonsen (2006a), the clade comprising the remaining Argynnis bears some similarities to the previous results, but also conflicts with these. The clade comprising A. pandora, A. sagana and A. paphia is supported in both analyses and the sister group relationship between the latter two suggested by Simonsen (2006a) is strongly supported here. The sister clade that here comprises A. hyperbius, A. childreni, A. anadyomene, A. laodice and A. ruslana contradicts Simonsen (2006a) where A. childreni (and its sister species A. zenobia) is placed with the subgenus Speyeria. However, Penz & Peggie (2003) placed A. childreni with A. hyperbius. We agree with Simonsen (2006a) that A. ruslana and A. laodice form a strongly supported clade and that their sister group is A. anadyomene.

Classification

The present results necessitate three changes in the classification proposed by Simonsen (2006a). Since the clade comprising *Boloria* and *Yramea* is not supported here we suggest that the name Yrameina should be reserved for a subtribe comprising only *Yramea*. Given its phylogenetic position, *Boloria* should be placed in its own subtribe. The name Boloriina (Warren 1944, Warren et al. 1946) is available for this genus and should be assigned to it. The inclusion of the re-established genus *Prokuekenthaliella* (Simonsen 2006a) in *Issoria* removes the need for retaining *Prokuekenthaliella* as a separate genus.

The fairly well supported clade comprising the *Argynnis* s.l. species supports the unification of all the "larger fritillaries" in one genus. As argued by

Simonsen (2006a) there is little justification for the large number of generic names traditionally applied to various members of that group, and one large, unified genus Argynnis seems to be the only stable and "natural" solution for this problem. Not only is Argynnis as defined here well delimited and easily recognized based on the highly specialized male alar androconials (Barth 1944), it is also supported in the morphological analysis, the combined molecular analysis and the total evidence analysis. Additionally, no alternative division of Argynnis into two or three genera seems ideal. The combined analyses do split Argynnis into two reciprocally monophyletic groups. However, these two groups are contradicted by morphology and not easily and immediately recognizable as a unit.

In summary, the tribe Argynnini is a well supported monophyletic group comprising of the subtribes Euptoietina, Yrameina, Boloriina and Argynnina. Six monophyletic, well-supported, robust and morphologically well-defined clades are termed genera in this study. These are *Euptoieta*, *Yramea*, *Boloria*, *Issoria*, *Brenthis* and *Argynnis*. We feel that this classification of Argynnini will be stable to the addition of new data based on our analyses in this paper.

Correction added in proof

Just as the paper was being typeset, the authors became aware that the *Boloria euphrosyne* specimen from which the DNA sequences were obtained was in fact a misidentified, slightly aberrant, *Boloria selene* (identified by T. J. Simonsen). This does not affect the results and conclusions, since the internal relationships of *Boloria* were outside the scope of the study.

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