# MOLECULAR PHYLOGENY OF COLEOID CEPHALOPODS (MOLLUSCA: CEPHALOPODA) INFERRED FROM THREE MITOCHONDRIAL AND SIX NUCLEAR LOCI: A COMPARISON OF ALIGNMENT, IMPLIED ALIGNMENT AND ANALYSIS METHODS

### JAN STRUGNELL<sup>1,2</sup> AND MICHELE K. NISHIGUCHI<sup>3</sup>

<sup>1</sup>School of Biology and Biochemistry, Queen's University Belfast, 97 Lisburn Road, Belfast BT9 7BL, UK;

<sup>2</sup>British Antarctic Survey, Natural Environmental Research Council, High Cross, Madingley Road, Cambridge CB3 0ET, UK;

<sup>3</sup>Department of Biology, New Mexico State University, Box 30001, MSC 3AF, Las Cruces, NM 88003-8001, USA

(Received 15 December 2006; accepted 1 September 2007)

### ABSTRACT

Recent molecular studies investigating higher-level phylogenetics of coleoid cephalopods (octopuses, squids and cuttlefishes) have produced conflicting results. A wide range of sequence alignment and analysis methods are used in cephalopod phylogenetic studies. The present study investigated the effect of commonly used alignment and analysis methods on higher-level cephalopod phylogenetics. Two sequence homology methods: (1) eye alignment, (2) implied alignment, and three analysis methods: (1) parsimony, (2) maximum likelihood, (3) Bayesian methodologies, were employed on the longest sequence dataset available for the coleoid cephalopods, comprising three mitochondrial and six nuclear loci. The data were also tested for base composition heterogeneity, which was detected in three genes and resolved using RY coding. The Octopoda, Argonautoidea, Oegopsida and Ommastrephidae are monophyletic in the phylogenies resulting from each of the alignment and analysis combinations. Furthermore, the Bathyteuthidae are the sister taxon of the Oegopsida in each case. However many relationships within the Coleoidea differed depending upon the alignment and analysis method used. This study demonstrates how differences in alignment and analysis methods commonly used in cephalopod phylogenetics can lead to different, but often highly supported, relationships.

### INTRODUCTION

The class Cephalopoda comprises two extant subclasses, Nautiloidea (*Nautilus* and *Allonautilus*) and the Coleoidea. The Coleoidea contains two subdivisions, the Belemnoidea, which became extinct at the end of the Cretaceous, and the Neocoleoidea, which contains the octopuses, squids and cuttlefishes. Neocoleoid cephalopods are characterized by the reduction and internalization, or complete loss, of the shell and, as a result, they very rarely fossilize well. Therefore, very little information regarding the origins and relationships of extant coleoid cephalopods can be gleaned from the fossil record (Nishiguchi & Mapes, 2007).

Morphological studies have proved to be useful in classifying species within subfamilies and/or genera (e.g. Berthold & Engeser, 1987; Clarke, 1988; Khromov, 1990; Voight, 1993a,b; Young & Vecchione, 1996; Voss, 1988), but less so in determining higher-level relationships. Morphological studies attempting to resolve these relationships have been constrained by the number of characters used with confidence due to "problems primarily involving character independence, apomorphic 'loss', or assessment of homology/homoplasy" (Young & Vecchione, 1996).

Presently, extant coleoids are divided into two superorders, Decapodiformes and Octopodiformes (Berthold & Engeser, 1987). In his website 'The Fossil Coleoidea Page' (http://userpage.fu-berlin.de/~palaeont/fossilcoleoidea/welcome.html), Engeser draws attention to the fact that the term Octopodiformes is in use elsewhere and suggests the use of Vampyropoda (Boletzky, 1992) instead. The Decapodiformes (Decembrachiata Winckworth sensu Engeser, loc. cit.) contains the orders Teuthoidea [suborders Myopsida (closed-eye squids) and Oegopsida (open-eye

Correspondence: J. Strugnell; e-mail: jmst@bas.ac.uk

squids) and Sepioidea (families Idiosepiidae (pygmy squid), Sepiidae (cuttlefishes), Spirulidae (ram's horn squid), Sepiolidae (bobtail squids) and Sepiadariidae (bottletail squids)]. Current debate exists on the validity of the ordinal level of classification (Naef, 1921–1923; Voss, 1977; Berthold & Engeser, 1987; Young & Vecchione, 1996). Furthermore, Lindgren, Giribet & Nishiguchi (2004) question whether the suborder Oegopsida is monophyletic.

The Octopodiformes contains the orders Vampyromorpha (vampire 'squid') and Octopoda (pelagic and benthic octopuses), hence the name Vampyropoda (Boletzky, 1992). A sister-taxon relationship between these two orders is accepted primarily based on morphology (Pickford, 1939; Boletzky, 1992; Young & Vecchione, 1996; Engeser, 1997; Young, Vecchione & Donovan, 1998; Carlini, Reece & Graves, 2000), but combined analysis using molecular and morphological data suggests a sister-taxon relationship between the Decapodiformes and Vampyromorpha (Lindgren et al., 2004). The Octopoda comprises the suborders Cirrata (deep-sea finned octopuses) and Incirrata (benthic octopuses and pelagic octopuses, including the argonautoids and blanket octopuses. A sister-taxon relationship between these suborders is also widely accepted (Grimpe, 1921; Naef, 1921–1923; Young & Vecchione, 1996; Voight, 1997). Phylogenetic relationships between the nine Incirrata families remain unresolved and have been debated in the literature (Naef, 1921-1923; Robson, 1929, 1931; Voss, 1977; Young & Vecchione, 1996; Voight, 1997).

In the mid 1990s the first studies using DNA sequence data to estimate phylogenetic relationships within cephalopods were reported (Bonnaud, Boucher-Rodoni & Monnerot, 1994, 1996, 1997; Boucher-Rodoni & Bonnaud, 1996). These studies sequenced portions of 16S rDNA, COII and COIII from 8 to 28 cephalopod taxa. These authors aligned their sequences by

eye (with the aid of the secondary structure where possible) and analysed the data using neighbour-joining (NJ) and parsimony methods. Although the genes proved useful in helping resolve intrafamilial relationships, little resolution of higher-level relationships was recovered. Subsequently, molecular studies investigating higher-level phylogenetic relationships of cephalopods have sequenced additional mitochondrial genes (Carlini & Graves, 1999; Piertney et al., 2003; Nishiguchi, Lopez & Boletzky, 2004; Zheng et al., 2004; Guzik et al., 2005) including whole mitochondrial genomes (Yokobori et al., 2004; Akasaki et al., 2006) and also nuclear genes (Carlini et al., 2000; Warnke et al., 2003; Strugnell et al., 2004; Guzik et al., 2005; Strugnell et al., 2005) often from a greater number of taxa (Carlini & Graves, 1999; Anderson, 2000a, b; Carlini et al., 2000; Lindgren et al., 2004; Strugnell et al., 2005) (Table 1).

Furthermore, since these first studies of cephalopod molecular phylogenetics, the range of sequence alignment and analysis methods available to phylogeneticists has increased (Table 1), and debate concerning the best methods to use has flourished (e.g. Wheeler, 1995, Kjer, Gillespie & Ober, 2007). Studies investigating cephalopod phylogenetics have aligned sequences by eye (Carlini & Graves, 1999; Carlini et al., 2000; Strugnell et al., 2004, 2005) or with the aid of alignment packages (Piertney et al., 2003; Yokobori et al., 2004; Zheng et al., 2004; Guzik et al., 2005) and have employed a variety of methods of analysis, including neighbour-joining (Allcock & Piertney, 2002; Warnke et al., 2003; Yokobori et al., 2004; Zheng et al., 2004), parsimony (Carlini & Graves, 1999; Anderson, 2000a,b; Carlini et al., 2000; Carlini, Young & Vecchione,

2001; Allcock & Piertney, 2002; Warnke et al., 2003; Lindgren et al., 2004, 2005; Nishiguchi et al., 2004; Zheng et al., 2004; Guzik et al., 2005), maximum likelihood (ML) (Anderson, 2000a,b; Carlini et al., 2000, 2001; Allcock & Piertney, 2002; Warnke et al., 2003; Strugnell et al., 2004; Yokobori et al., 2004; Guzik et al., 2005), Bayesian (Strugnell et al., 2004, 2005; Guzik et al., 2005) and LogDet (Anderson, 2000b; Strugnell et al., 2005). Recently, some studies have employed direct optimization where alignment is coupled with tree estimation in a dynamic procedure (Nishiguchi et al., 2004; Lindgren et al., 2004, 2005) (Table 1).

Although providing some insights [e.g. sister taxon relationships between the suborder Oegospida and family Bathyteuthidae (Strugnell *et al.*, 2005)] none of these studies have conclusively resolved all higher-level cephalopod phylogenetic relationships and in many cases the results have been conflicting (see Akasaki *et al.*, 2006; Nishiguchi & Mapes, 2007 for review of conflicting decapodiform relationships).

A number of reasons have been suggested for these varying and unresolved relationships. These include the early divergence of taxa, saturated sequence data, insufficient data, insufficient taxa and gene duplication (see Bonnaud *et al.*, 1994, 1996; Carlini & Graves, 1999; Carlini *et al.*, 2000; Lindgren *et al.*, 2004; Strugnell *et al.*, 2005 for discussion).

The large molecular data sets generated by Lindgren et al. (2004) (four genes) and Strugnell et al. (2004, 2005) (six genes) contained 18 of the same species (including 6 Octopodiformes and 11 Decapodiformes). Together, these provide the single largest dataset (with regard to sequence length) available for investigating higher-level phylogenetic relationships within

**Table 1.** Summary of studies of the molecular phylogenetics of coleoid cephalopods.

Reference	Focal taxa	Genes used	No. of species	Sequence alignment method	Analysis method(s)	
Bonnaud et al. (1994)	Decapodiformes	16S	28	eye (2° structure)	NJ, P	
Bonnaud et al. (1996)	Decapodiformes	16S, COIII	8	eye	NJ, P	
Boucher-Rodoni & Bonnaud (1996)*	Coleoidea	16S	10		NJ, P	
Bonnaud et al. (1997)	Coleoidea	COIII, COII	17	eye	NJ, P	
Bonnaud et al. (1998)	Onychoteuthidae	16	14	eye	NJ, P	
Carlini & Graves (1999)	Coleoidea	COI	48	eye	Р	
Anderson (2000)	Loliginidae	16S, COI	$\sim$ 30	Clustal and eye	P, ML, LogDet	
Anderson (2000)*	Loliginidae	16S, COI	53	Clustal and eye	P, ML	
Carlini et al. (2000)	Coleoidea	actin	44	eye	P, ML	
Carlini et al. (2001)	Octopoda	COI	29	eye	P, ML	
Allcock & Piertney (2002)	Octopodidae	16S	9	Clustal X and eye	NJ, P, ML	
Piertney et al. (2003)	Cirrata	16S	27	Clustal X and eye	NJ, P, ML	
Warnke et al. (2003)	Decapodiformes	complete 18S	8	Clustal V, MegAlign,	NJ, P, ML	
				checked by eye		
Bonnaud et al. (2004)	Nautilus	complete 18S	3	eye	2° structure	
Lindgren et al. (2004)*	Coleoidea	complete 18S, 28S, hist. COI	60	POY	Р	
Nishiguchi et al. (2004)	Sepiolidae	12S, 16S, COI, 28S	30	POY	Р	
Strugnell et al. (2004)	Octopodiformes	16S, 12S, COI, rhod, pax-6, ODH		eye	ML, Bayesian	
Yokobori et al. (2004)	Coleoidea	whole mitochondrial genome	3	ClustalX	NJ, ML	
Zheng et al. (2004)	Decapodiformes	COI, 16S	13	ClustalX v1.8	NJ, P	
Guzik et al. (2005)	Octopodinae	COIII, cyt b, ef-1 $\alpha$	30	Sequencher 3.1	P, ML, Bayesian	
Lindgren et al. (2005)	Gonatidae	12S, 16S, COI	39	POY	Р	
Strugnell et al. (2005)	Coleoidea	16S, 12S, COI, rhod, pax-6, ODH	35	eye	Bayesian, LogDet	
Takumiya et al. (2005)	Coleoidea	12S, 16S, COI	36	SeqPup v. 0.9, ClustalX	NJ, P, ML	
				ver1.83		
Akasaki et al. (2006)	Coleoidea	whole mitochondrial genome	5	-	ML	

<sup>\*</sup>note these studies also used further information in some analyses in addition to gene sequences, e.g. morphology, allozymes, immunology etc. Abbreviations: cyt b, *cytochrome b apoenzyme*, COI, *cytochrome c oxidase subunit l*; 16S, 16S rDNA; 12S, 12S rDNA; 28S, 28S rDNA; 18S, 18S rDNA; ODH, *octopine dehydrogenase*; rhod, *rhodopsin*; hist, *histone* H3; ef-1α, elongation factor-1α; All sequences were of partial fragments unless otherwise stated. NJ, neighbour-joining; P, parsimony; ML, maximum likelihood.

the subclass Coleoidea. In the present study we used two methods to align these data: by eye and implied alignment using POY; and also three methods of analysis: parsimony, ML and Bayesian, to investigate the effect of these analyses on the resulting phylogeny. The effect of base composition heterogeneity upon coleoid phylogenetic relationships was also investigated.

### MATERIAL AND METHODS

Eighteen species were used in the present study, including representatives from each higher-level taxon within the subclass Coleoidea (Table 2). Portions of nine genes were included, three mitochondrial genes (12S rDNA, 16S, rDNA, COI) and six nuclear genes (28S rDNA, 18S rDNA, histone, octopine

**Table 2.** Accession numbers of each of the genes used in this study.

	Mitochondrial genes			Nuclear genes					
	12S rDNA	16S rDNA	COI	28S rDNA	18S rDNA	hist.	ODH	pax-6	rhod.
Nautiloidea									
Nautilida									
Nautilidae									
Nautilus pompilius	AY616965	AY377628	AY557514	AF311688	AY557452			AY617039	
Coleoidea									
Octopodiformes									
Vampyromorpha									
Vamyroteuthidae									
Vampyroteuthis infernalis	AY545077	AY545101	AF000071	AY557548	AY557459	AY557408	AY545114	AY545139	AY545163
Octopoda									
Allopsidae									
Haliphron atlanticus	AY616942	AY616971	AY557516	AY557549	AY557460	AY557409	AY616910	AY617016	AY617040
Argonautidae									
Argonauta nodosa	AY545080	AY545104	AY557517	AY557551	AY557462	AY557411	AY545117	AY545142	AY545166
Bolitaenidae	71.0.0000	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	71.007017	711007001	711007102	71.007 111	71.0.01.7	71.0.01.2	711010100
Japetella diaphana	AY545093	A252766	AY545192	AY557552	AY557463		AY545130	AY545155	AY545179
Octopodidae	711010000	71202700	711010102	711007002	711007100		711010100	711010100	711010170
Eledone cirrhosa	AY616946	AY616973	AY557520	AY557556	AY557467		AY616992	AY617020	AY617043
Graneledone verrucosa	AY545091	AY545111	AF000042	AY557557	AY557468	AY557413	AY545129	AY545153	AY545177
Decapodiformes	A 1 54509 1	A1343111	AI 000042	A1337337	A1337400	A1557415	A1545129	A1343133	A1545177
•									
Sepiolida									
Sepiolidae	4)/040070	AV040004	A F000044	AV000700	AVEE7470	AV/557440	A)/040000	4)/040007	4)/040000
Heteroteuthis hawaiiensis	AY616873	AY616884	AF000044	AY293703	AY557472	AY557416	AY616906	AY616937	AY616922
Sepiida									
Sepiidae	43/545000	\\0==0		****	*>//	*>/	43/5/5/5	***	. =
Sepia officinalis	AY545098	X9570	AJ583491	AY557560	AY557471	AY557415	AY545135	AY545160	AF000947
Idiosepiida									
Idiosepiidae									
Idiosepius pygmaeus	AY545095	AJ001647	AY545193	AY293684	AY557477	AY557421	AY545132	AY5157	AY545181
Spirulida									
Spirulidae									
Spirula spirula	AY545097	AY293659	AY293709	AY557563	AY557476	AY557420	AY545134	AY545159	AY545183
Teuthida Myopsdia									
Loliginidae									
Sepioteuthis lessoniana	AY616869	AJ001649	AY131036	AY557566	AY557480	AY557424	AY616902	AY616933	AY616918
Teuthida Myopsdia									
Bathyteuthidae									
Bathyteuthis abyssicola	AY616958	AJ000104	AF000030	AY557568	AY557483	AY557427	AY617002	AY617032	AY617057
Octopoteuthidae									
Octopoteuthis nielseni	AY616957	AY616983	AF000055	AY557591	AY557507		AY617011-	AY617031	AY617056
							AY617013		
Cranchiidae									
Cranchia scabra	AY616962	DQ280046	AF000035	AY557571	AY557487	AY557430	AY617014	AY617036	AY617061
							AY617015		
Ommastrephidae									
Illex coindetii	AY616963	AY616985	AY617065	AY557593	AY557509	AY557450	AY617008	AY617037	AY617062
							AY617015		
Sthenoteuthis oualaniensis	AY545100	X79582	AF000069	AY557595	AY557511	AY557452	AY545137	AY545162	AY545185
Ommastrephes batramii	AY616866	AY616880	AF000057	AY557594	AY557510	AY557451	AY616899	AY616930	AY616915

dehydrogenase [ODH], pax-6 and rhodopsin). Sample details and methodologies used to obtain DNA sequences from these species are outlined in Lindgren et al. (2004) and Strugnell et al. (2005). Accession numbers for these sequences are listed in Table 2.

### Sequence alignment and homology assessment

Two methods of sequence alignment were used within this study (1) by eye, and (2) using implied alignment using the homology scheme via POY (Wheeler, 2003; Giribet, 2005).

### Aligned by eye

DNA sequences were compiled and aligned by eye in Se-Al v2.0a11 Carbon (Rambaut, 2002). Gaps were inserted where necessary to allow sequences to be aligned. Sequence data that were not alignable using this method were removed prior to analyses. Sequence alignment files are available on request. The total concatenated sequence length was 5,651 bp, of which 2,219 bp were variable.

### Dynamic homology and implied alignments

Sequence data were analysed by using the direct optimization method described by Wheeler (1996) and implemented in the computer program POY. This method directly assesses the number of DNA sequence transformations (evolutionary events) required by a phylogenetic topology without the use of multiple sequence alignment. This is accomplished by generalization of existing character optimization procedures, including insertion and deletion events (indels) in addition to base substitutions. This method treats indels as processes, as opposed to the patterns implied by multiple sequence alignment (Wheeler, 1995). It is claimed that this method generates more efficient (and therefore simpler) explanations of sequence variation than multiple sequence alignment (Wheeler, 1996). Direct optimization, although computationally intense, is much less demanding than parsimony-based multiple sequence alignments when congruence among partitions is used as a criterion (Wheeler & Hayashi, 1998). The implied alignments produced via POY were used for both ML and Bayesian analyses. These sequences were concatenated for ML and Bayesian analysis (6,377 bp, of which 2,330 bp were variable).

### Base composition heterogeneity

PAUP\*4.0b10 (Swofford, 1998) was used for  $\chi^2$  tests of composition homogeneity of the sequence data aligned by eye. Tests of base homogeneity were based on variable sites only. Where base composition heterogeneity was detected it was RY coded to remove base composition heterogeneity.

The three sequence data sets, (1) implied alignments, (2) aligned by eye, (3) aligned by eye and RY coded were analysed using three methods, (a) parsimony, (b) maximum likelihood, (c) Bayesian analysis. It is important to note that the sequence data aligned by eye were analysed using parsimony analyses in PAUP rather than POY.

### Dynamic homology under parsimony

Molecular data were analysed with the computer program POY (Wheeler *et al.*, 1996–2003) using the direct optimization method (Wheeler, 1996), with parsimony as the optimality criterion. Nodal support was calculated in POY using Farris's parsimony jackknifing procedure (Farris *et al.*, 1996) for 100 replicates (using the commands: jackboot; replicates 100). Tree searches were conducted in parallel at Harvard University

on a 19 dual-processor cluster (Darwin.oeb.harvard.edu) using pvm (parallel virtual machine). Commands for lad balancing of spawned jobs were used to optimize parallelization procedures (-parallel-dpm-jobspernode 2). Trees were built via a randomaddition sequence procedure (10 replicates) followed by a combination of branch-swapping steps [SPR (subtree pruning and regrafting) and TBR (tree bisection and reconnection)] and tree fusing (Goloboff, 1999) in order to further improve on tree length minimization. Discrepancies between heuristic and actual tree length calculations were addressed by adjusting slop values (-slop5-checkslop10). Phylogenetic trees were obtained using parsimony with a gap/ts/tv cost of various weighting. Several analyses were implemented with character transformations weighted differently to determine how various phylogenetic hypotheses were affected (sensitivity analysis sensu Wheeler, 1995). Each gene was analysed separately, using character transformations (indels/ts/tv) of equal weighting (111), and unequal weighting (121, 141, 211, 221, 241, 411, 421, 441). The parameter set that optimized the least amount of character incongruence was the equal weighted transformation (111) for all genes. Histone H3 and pax-6 were the two exceptions that also had similar character incongruence values for the 211 and 411 transformations. The final tree was drawn with Tree View (Win32) and consensus trees were analysed in PAUP version 4.02b (Swofford, 1998). To determine nodal support all jackknife calculations were performed in POY using the procedure described in Nishiguchi et al. (2004).

### Implied alignment under parsimony

PAUP\*4.0b10 (Swofford, 1998) was used to perform maximum parsimony analyses on the sequence data that were aligned by eye. All parsimony searches were performed with 1,000 random sequence-addition replicated and TBR (tree bisection-reconnection) branch swapping. All characters were unordered and equally weighted. One thousand bootstrap replicates were performed to measure the support for each clade on the phylogenetic trees.

### Alignment by eye and implied alignment under maximum likelihood

PAUP\*4.0b10 (Swofford, 1998) was used to perform 100 full heuristic searches. Starting trees were generated by the neighbour-joining method (NJ) (Saitou & Nei, 1987). A GTR + I +  $\Gamma$  likelihood model incorporating rate heterogeneity was used. Branch swapping was performed using TBR (tree-bisection-reconnection). Parameters were then re-estimated, and final branch swapping was performed using NNI (nearest-neighbour-interchange). ML bootstrap values of clade support were generated using the parameters estimated in the analysis, but with starting trees generated by the neighbour-joining method.

## Alignment by eye and implied alignment under Bayesian analyses

MrBayes 3.1.2 (Ronquist & Huelsenbeck, 2003) was used to calculate marginal posterior probabilities using the GTR + I +  $\Gamma$  model of nucleotide substitution. Model parameter values were treated as unknown and were estimated in each analysis. Random starting trees were used for the analyses and were run between 1 and 500,000 generations, sampling the Markov chain every 100 generations.

Three strategies were used to ensure that analyses were not trapped in local optima: (1) analysis was performed twice, starting with a different random tree and log-likelihood values at stationarity were compared for convergence

(Huelsenbeck & Bolback, 2001); (2) the topologies and clade posterior probabilities from each of the two analyses were compared for congruence (Huelsenbeck & Imennov, 2002); and (3) Metropolis-coupled Markov chain Monte Carlo (MCMCMC) was used with one cold and three incrementally heated Markov chains run simultaneously (default Mr Bayes heating values) to allow a more extensive exploration of parameter space (Huelsenbeck & Ronquist, 2001).

Stationarity was deemed to be reached when the average standard deviation of split frequencies, shown in MrBayes 3.1.2 was less than 0.01 (Ronquist & Huelsenbeck, 2003).

Tracer v1.3 (Rambaut & Drummond, 2003) was used to determine the correct 'burnin-in' for the analysis (i.e. the number of initial generations that must be discarded before stationarity is reached).

### RESULTS

### Sequence alignment

Alignment of the ODH, pax-6, COI and histone sequences required no insertion/deletion events (indels). Indels were introduced into aligned sequences of 12S rDNA, 16S rDNA, 28S rDNA, 18S rDNA and *rhodopsin* both by eye and 'dynamically' during the analysis using POY. The alignments of these genes where indels were required differed notably depending upon the alignment method (Table 3) (alignments available on request). A greater number of gaps were inserted using POY than by eye for the 12S rDNA, 16S rDNA and 28S rDNA genes (Table 3), whereas a greater number of insertions was used aligning by eye than by using POY for rhodopsin and 18S rDNA (Table 3). For each of these five genes requiring indels, regions that were deemed to be unalignable with confidence by eye were removed prior to analysis. In contrast, no sequence was removed from the POY analysis due to the fact that sequences are aligned simultaneously during analysis.

### Base composition heterogeneity

Chi-squared homogeneity tests of each of the genes shows that third positions of ODH, *rhodopsin* and COI have significant base frequency heterogeneity (Table 4). RY-coding the third positions of these genes was used to resolve base composition heterogeneity (Table 4). RY coding pools purines (adenine and guanine:R) and pyrimidines (cytosine and thymine:Y) into two-state categories (R,Y), and helps resolve bias resulting from differences in the relative frequency of either the two purines or pyrimidines (Phillips *et al.*, 2001).

A number of taxonomic groupings are robust to the different methods of coding, alignment and analysis. The following taxa are always monophyletic: Octopoda, Argonautoidea, Ommastrephidae and Oegopsida (Figs 1–9). Furthermore, in each topology the Bathyteuthoida is the sister taxon to the Oegopsida (Figs 1–9). Bayesian posterior probabilities provide the highest support for each of these clades (Figs 3, 6, 9).

Vampyromorpha and the Decapodiformes are sister taxa in the phylogenies resulting from parsimony, ML and Bayesian analyses of the sequences aligned using POY and also by eye (no RY) with variable levels of support (Figs 1–6). This relationship is also recovered from parsimony analysis of RY coded sequence (BS = 99) (Fig. 7). In contrast, ML and Bayesian analysis of RY coded sequence aligned by eye recovered a sister-taxon relationship between Vampyromorpha and Octopoda, i.e. the Octopodiformes (Figs 8, 9). However, these relationships are not highly supported by bootstraps or posterior probabilities (Figs 8, 9).

The placement of *Eledone* within the Octopoda differs depending upon alignment and analysis method. *Eledone* is the sister

**Table 4.** Chi-squared homogeneity test for base composition across all genes and codon positions.

Gene	Codon position	$\chi^2(P)$		
12S rDNA	-	0.998		
16S rDNA	_	0.997		
18S rDNA	_	0.963		
28S rDNA	_	1.000		
COI	1st	1.000		
COI	2nd	1.000		
COI	3rd	0.000		
COI (RY)	3rd	0.938		
histone H3	1st	1.000		
histone H3	2nd	1.000		
histone H3	3rd	0.560		
ODH	1st	1.000		
ODH	2nd	1.000		
ODH	3rd	0.000		
ODH (RY)	3rd	0.999		
pax-6	1st	1.000		
pax-6	2nd	1.000		
pax-6	3rd	0.945		
rhodopsin	1st	0.994		
rhodopsin	2nd	0.962		
rhodopsin	3rd	0.003		
rhodopsin (RY)	3rd	0.721		

Tests were performed on variable sites only.  $\chi^2(P) < 0.05$  are in bold.

**Table 3.** Comparison of alignment length of genes.

	Gene	Total base pairs in gene sequenced (no gaps) (bp)	Alignment method						
		σοφασίποσα (πο θαφο) (οφ)	POY (bp)	Eye (total alignment length) (bp)	Eye (unalignables remove in analysis) (bp)				
fitochondrial 12S rDNA 417	417	573 (417)*	486 (417)*	283					
	16S rDNA	528	627 (528)*	554 (528)*	427				
Nuclear	18S rDNA	2,845	1,893 (1,842)*	3,202 (2,845)*	1,943				
	28S rDNA	661	198 (191)*	166 (166)*	166				
	rhodopsin	1,040	1,022 (991)*	1,032 (954)*	765				

<sup>\*</sup>Number in brackets indicates the starting sequence length without gaps. The portion of available sequence able to be aligned by eye was less for 28S and *rhodopsin* than by POY. A larger sequence fragment of 18S was attempted for alignment by eye, however a large proportion was unalignable and was removed prior to analysis.

**Table 5.** Phylogenetic relationships recovered by two alignment methods (by eye, dynamic homology/implied alignment using POY) and three analysis methods (P, parsimony; ML, maximum likelihood; Bayes, Bayesian).

Alignment method	POY			By eye						
				No RY		RY				
Analysis method	POY	ML	Bayes	POY	ML	Bayes	POY	ML	Bayes	
Vampyromorpha(Decapodiformes)	$\checkmark$	X	Χ							
Vampyromorpha(Octopoda)	Χ	Χ	X	Χ	Χ	Χ	Χ	$\checkmark$	$\checkmark$	
Octopoda	$\checkmark$	$\checkmark$	$\checkmark$	$\checkmark$	$\sqrt{}$	$\checkmark$	$\checkmark$	$\checkmark$	$\checkmark$	
Argonautoidea	$\checkmark$									
((Japetella, Graneledone)Eledone)	$\checkmark$	$\checkmark$	$\checkmark$	$\checkmark$	Χ	Χ	$\checkmark$	X	Χ	
(Eledone((Japetella,Graneledone)(Haliphron,Argonauta))	X	X	Χ	X	$\checkmark$	$\checkmark$	Χ	$\checkmark$	$\checkmark$	
Decapodiformes	$\checkmark$									
(Oegopsida)(remaining Decapodiformes)	Χ	X	Χ	X	$\checkmark$	$\checkmark$	Χ	$\checkmark$	$\checkmark$	
Polyphyletic Sepioidea	$\checkmark$									
Ommastrephidae	$\checkmark$									
Oegopsida	$\checkmark$									
Bathyteuthoida(Oegopisda)	$\checkmark$									
Spirulida(Bathyteuthoida(Oegopisda))	$\checkmark$	$\checkmark$	$\checkmark$	$\checkmark$	Χ	Χ	Χ	X	Χ	
Idiosepiidae(Sepioidea(Myopsida(Spirulida(Bathyteuthoida(Oegopisda)))))	$\checkmark$	X	Χ	X	Χ	Χ	Χ	X	Χ	
Sepioidea(Myopsida(Spirulida(Bathyteuthoida(Oegopisda))))	$\checkmark$	$\checkmark$	$\checkmark$	Χ	Χ	Χ	Χ	Χ	Χ	
Myopsida(Spirulida(Bathyteuthoida(Oegopisda)))	$\checkmark$	$\checkmark$	$\checkmark$	X	Χ	Χ	Χ	X	Χ	
(Heteroteuthis, Idiosepius)	X	$\checkmark$	$\checkmark$	X	Χ	Χ	Χ	X	Χ	
(Sepioteuthis, Idiosepius)	X	X	Χ	$\checkmark$	Χ	Χ	Χ	X	Χ	
(Sepia, Idiosepius)	Χ	Χ	Χ	Χ	$\checkmark$	$\checkmark$	$\checkmark$	$\checkmark$	$\checkmark$	
((Sepia, Idiosepius)Sepioeuthis)	X	X	Χ	X	$\checkmark$	$\checkmark$	$\checkmark$	X	$\checkmark$	
(((Sepia, Idiosepius)Sepioeuthis)Spirula)	Χ	Χ	X	Χ	$\checkmark$	$\checkmark$	$\checkmark$	Χ	$\checkmark$	
((((Sepia, Idiosepius)Sepioeuthis)Spirula)Heteroteuthis)	Χ	Χ	Χ	Χ	$\checkmark$	$\checkmark$	$\checkmark$	Χ	$\checkmark$	
(Oegopsida,Bathyteuthoidea)(Sepioidea, Myopsida*)	Χ	X	Χ	Χ	$\checkmark$	√	X	$\checkmark$	$\checkmark$	

<sup>\*</sup>Myopsida falls within Sepioidea in this topology.

taxon to a clade containing Japetella and Graneledone in each of the phylogenies resulting from the POY alignment, and also parsimony analysis of the sequence data aligned by eye, both RY coded and not RY coded (Figs 1–4,7). High support for this

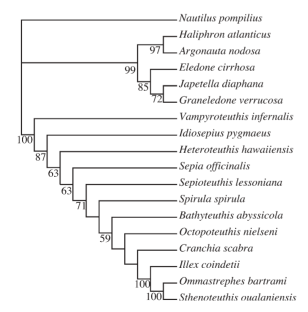


Figure 1. Parsimony topology of coleoid cephalopod relationships obtained using direct optimization using POY. Jackknife support values are indicated beneath each node.

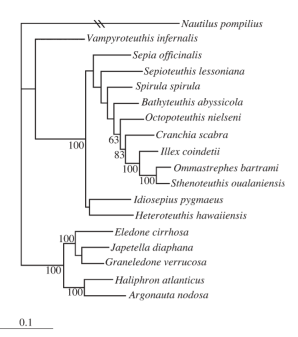
relationship is evident on each of these phylogenies. However, *Eledone* is highly supported as being basal within the Octopoda in analysis of sequence data aligned by eye (both RY coded and not RY coded) and analysed using ML (no RY, BS = 100; RY, BS = 100) and Bayesian analyses (no RY, PP = 1.00; RY, PP = 1.00) (Figs 5, 6, 8, 9).

Higher-level decapodiform relationships differ markedly between the various methods of alignment, coding, and analysis (Figs 1–9). Phylogenies generated from ML and Bayesian analyses of sequences aligned by eye (both RY coded and not RY coded) demonstrate Decapodiformes to be divided into two monophyletic groups, one containing the Oegopsida, and the second containing the remaining decapodiforms (i.e. Myopsida, Spirulidae, Sepiidae, Sepiolidae and Idiosepiidae) (Figs 5, 6,8,9). This division is highly supported by bootstrap support (no RY, BS = 98; RY, BS = 98) and posterior probabilities (no RY, PP = 0.99; RY, PP = 0.97) (Figs 5, 6,8,9). Within these topologies *Sepia* and *Idiosepius* are sister taxa, thereby rendering 'Sepioidea' (including Sepiidae, Sepiadariidae and Sepiolidae) polyphyletic (Figs 5, 6, 8, 9).

In contrast, a clade containing *Heteroteuthis* and *Idiosepius* is basal within decapodiforms in ML and Bayesian analysis (PP = 0.90) of sequence data aligned using POY (Figs 2, 3). *Heteroteuthis* alone is basal in phylogenies resulting from parsimony analysis of sequences aligned by eye, both RY coded (BS = 100) and not RY coded (BS = 100) (Figs 4, 7).

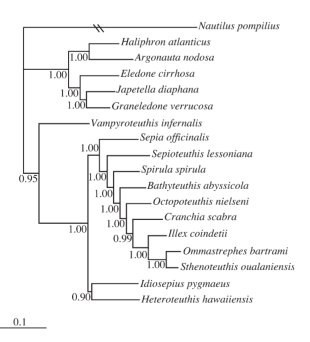
The position of Spirulidae within the Decapodiformes is highly dependent upon the method of alignment and analysis. Spirulidae are the sister taxon to a clade containing the Oegopsida and Bathyteuthoidea in all three analyses where sequences were aligned using POY, although support was only obtained

The data aligned by eye have been analysed for both nucleotide data and RY coded data.  $\sqrt{\ }$ , the relationship is supported, X, the relationship is not supported.

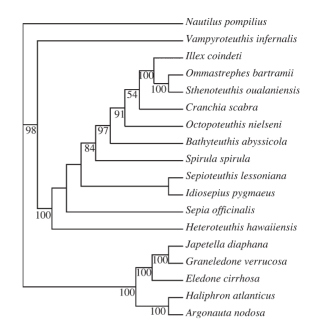


**Figure 2.** ML topology of coleoid cephalopod relationships obtained using  $GTR + I + \Gamma$ . Sequences were obtained from implied alignments using POY. Bootstrap support values are indicated beneath each node.

for this relationship from the Bayesian analysis (PP = 1.00) (Figs 1–3). Interestingly, this same arrangement results from parsimony analysis of sequence data aligned by eye, not RY coded (BS = 84) (Fig. 4). In contrast, Spirulidae are the sister taxon to a clade containing Idiosepiidae, Sepiidae and Myopsida in the topologies resulting from ML and Bayesian analysis (PP = 0.99) of data aligned by eye (not RY coded) (Figs 5, 6) and in parsimony and Bayesian analysis (PP = 0.92) of RY coded data aligned by eye (Figs 8, 9).



**Figure 3.** Bayesian topology of coleoid cephalopod relationships obtained using  $GTR + I + \Gamma$ . Sequences were obtained from implied alignments using POY. Bayesian posterior probabilities are indicated beneath each node.

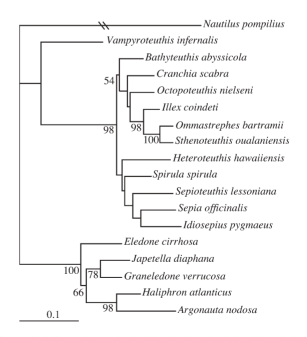


**Figure 4.** Parsimony topology of coleoid cephalopod relationships. Sequences were aligned by eye. Boostrap support values are indicated beneath each node.

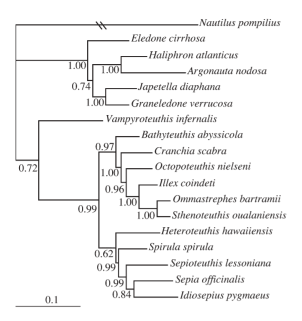
### DISCUSSION

The present study is the largest molecular analysis of cephalopod phylogeny to date, with regard to sequence length, and provides a thorough comparison of the effect of commonly used alignment and analysis methodologies on the resulting higher-level phylogenetic relationships.

The different alignment, analysis and coding methods used within this study produced a range of considerably different topologies. Only the clades Octopoda, Argonautoidea, Decapodiformes, Oegopsida, Ommastrephidae and a sister-taxon



**Figure 5.** ML topology of coleoid cephalopod relationships obtained using  $GTR + I + \Gamma$ . Sequences were aligned by eye. Bootstrap support values are indicated beneath each node.

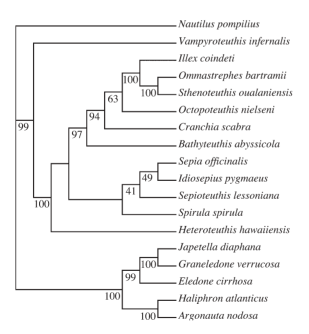


**Figure 6.** Bayesian topology of coleoid cephalopod relationships obtained using  $GTR + I + \Gamma$ . Sequences were aligned by eye. Bayesian posterior probabilities are indicated beneath each node.

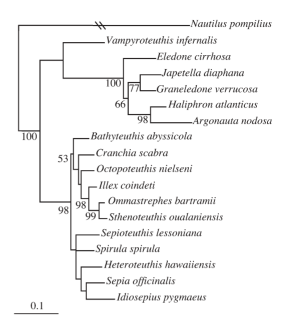
relationship between Bathyteuthidae and Oegopsida are robust to the alignment and analysis methods used.

### Alignment methods

It is not surprising that different alignments can affect the resulting phylogeny, as the process of alignment aims to recover the evolutionary history of the sequences and therefore provides the very data upon which the algorithm performs (Giribet,



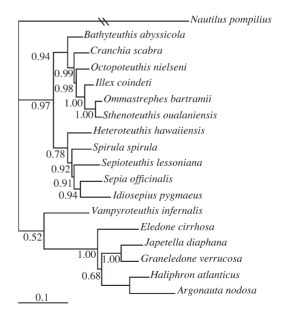
**Figure 7.** Parsimony topology of coleoid cephalopod relationships. Sequences were aligned by eye, and third positions of *rhodopsin*, COI and ODH were RY coded. Bootstrap support values are indicated beneath each node.



**Figure 8.** ML topology of coleoid cephalopod relationships obtained using  $GTR + I + \Gamma$ . Sequences were aligned by eye, and third positions of *rhodopsin*, COI and ODH were RY coded. Boostrap support values are indicated beneath each node.

Desalle & Wheller, 2002). For protein coding genes, the method of sequence alignment is usually insignificant, since in theory they should all produce the same alignment, i.e. an alignment without indels. However, as we demonstrate here, alignment methods for rDNAs, and both coding (i.e. *rhodopsin*) and non-coding genes can differ in their resulting sequence alignment and phylogenies.

There is debate in the literature regarding the best method of sequence alignment. Proponents of aligning sequences by eye (using secondary structural information) claim that



**Figure 9.** Bayesian topology of coleoid cephalopod relationships obtained using  $GTR + I + \Gamma$ . Sequences were aligned by eye, and third positions of *rhodopsin*, COI and ODH were RY coded. Bayesian posterior probabilities are indicated beneath each node.

they are 'both philosophically and operationally superior' (Kjer *et al.*, 2007), whereas proponents of computational methods claim that alignments performed by eye are subjective and therefore not repeatable (Giribet & Wheeler, 1999).

Proponents of the method of direct optimization using POV claim that it avoids the problem of alignment by generalizing phylogenetic character analysis to include insertion/deletion events (indels), with the sequence data proceeding directly to phylogenetic reconstruction, obviating the necessity to create gap characters. Indels do not appear as states, but as transformations linking ancestral to descendent nucleotide sequences (Giribet & Wheeler, 1999; Giribet et al., 2002). POY assumes that shorter trees are better trees and that aligning nucleotides together based on state is parsimonious and algorithmically less costly. Kjer et al. (2007) argues that this is not justified in structurally conserved molecules such as rDNAs, where conserved structures in the molecules are more important than the states of the nucleotides.

There has been intense disagreement over the relative merits of manual alignment and direct optimization (Kjer, 1995; Wheeler, 1995; Shull et al., 2001; Belshaw & Quicke, 2002; Gillespie, Yoder & Wharton, 2005) but few rigorous comparisons of these methods. Recently, Kjer et al. (2007) compared the phylogenies obtained by three phylogeneticists who independently aligned and analysed the same 16S rDNA datset by eye (using rDNA secondary structure and analysed by parsimony) and using direct optimization within POY. Interestingly, although all three alignments by eye differed at some positions, each alignment produced nearly identical topologies. In contrast, when using POY, none of the three phylogeneticists converged on the same parameters or the same tree. Kjer et al. (2007) suggest that the reason for this is that gap cost to change ratios (used within POY) are arbitrary, and this allows different researchers to obtain different results.

Sequence alignments resulting from POY have been reported to be 'gappy' in some studies (Pons & Vogler, 2006) with the program inserting a greater number of indels than other methods when utilizing indel costs of 1. Similarly, in the present study POY inserted a greater number of gaps in the 12S rDNA, 16S rDNA and 28S rDNA alignments, even though several parameters were explored using POY (costs of 1, 2 and 4 for indels, transitions and transversions, respectively). In addition, no sequence data were removed from the POY alignments used in the analysis. In contrast, a notable proportion of the sequence alignments of 12S rDNA, 16S rDNA, 18S rDNA and rhodopsin was removed after alignment by eye as it was deemed to be unalignable and would contribute noisy signal to the analysis. Therefore the starting information present in both datasets differed. The sequence data that were deemed 'unalignable' when aligning by eye are by their nature 'variable' and would therefore have an important contribution in the POY alignments in determining the resulting phylogenetic relationships. Differences in phylogenetic relationships observed in this study between the two alignment methods is largely due to the deleted sequences. Over 60% of the sequence information in both the datasets was constant (i.e. not variable) further demonstrating the significance of these variable sites. Despite these obvious differences in output, both of these methods of sequence alignment are widely accepted and appear in the cephalopod (Table 1) and wider literature today. It is likely that debate will continue regarding the best method of sequence alignment and while this continues to be the case, it may be beneficial to employ more than one method of alignment in phylogenetic studies.

Analysis methods

There is considerable debate in the literature regarding methods of phylogenetic analysis (e.g. Giribet, 2003). Parsimony methods have the benefit of being relatively easy to understand and require few assumptions about the evolutionary process (Page & Holmes, 1998). However they have been shown to produce the wrong topology under the most realistic models of evolution (e.g. long branch attraction; Huelsenbeck & Hillis, 1993).

ML methods allow the incorporation of sophisticated models of sequence evolution and allow statistical tests of different evolutionary hypotheses (i.e. likelihood ratio testing Felsenstein, 1981) yet require very large computational resources. Furthermore ML methods have been shown to be susceptible to long branch repulsion and long branch attraction under some circumstances (Pol & Siddall, 2001).

Bayesian methodologies (differing from likelihood methods only in the use of a prior distribution of the quantity being inferred, which is typically the tree) have the advantage over ML methods of being computationally efficient. They allow very complex models of sequence evolution to be implemented and also can efficiently analyse large datasets. Bayesian methods have been criticized however, for producing unrealistically high posterior probability support (Suzuki, Glazko & Nei, 2002; Simmons, Pickett & Miya, 2004).

In the present study, the majority of topologies resulting from the three analysis methods on the implied aligned data (from POY) are very similar. The exception to this is the position of Idiosepius. In contrast, the method of analysis had a greater effect on the data aligned by eye. In many cases ML and Bayesian methods of analysis produced the same or very similar topology for both RY coded and non-RY coded data, while the parsimony analysis produced a different topology. This is the case for the relationships of octopod taxa, and the relationship between the Oegopsida and the rest of the decapodiforms. It is unsurprising that ML and Bayesian analysis methods produce more similar topologies than parsimony analysis, because both are based on the same probabilistic model of evolution. In contrast, parsimony analysis is based on the idea that the preferred phylogenetic tree is the one that requires the fewest evolutionary changes.

### Discussion of phylogenetic relationships

Order Vampyromorpha: Vampyroteuthis infernalis is the only species within the order Vampyromorpha. It possesses a number of unusual characteristics including two pairs of fins in juveniles (one pair in adults) and a second pair of arms modified into retractile filaments. Traditionally Vampyromorpha and Octopoda have been suggested to be sister taxa due to embryological, developmental (Naef, 1928; Young & Vecchione, 1996; Boletzky, 2003) and morphological similarities, such as sperm morphology (Healy, 1989) and the presence of radial sucker symmetry (Lindgren et al., 2004). However, the vampyromorph gladius is known to be morphologically similar to that of decapodiforms (Toll, 1982, 1998). Previous molecular studies have found support for both a sister taxon relationship between Vampyromorpha and Octopoda (Bonnaud et al., 1997; Carlini & Graves, 1999; Lindgren et al., 2004; Strugnell et al., 2004,2005) and Vampyromorpha and the Decapodiformes (Bonnaud et al., 1997; Lindgren et al., 2004). This present study found support for both of these relationships. The majority of alignment and analysis combinations support a sister-taxon relationship between Vampyromorpha and Decapodiformes. Only ML and Bayesian analysis of the 'by eye' alignment of RY coded data support a sister-taxon relationship between Vampyromorpha and Octopoda. RY coding rectified the base composition heterogeneity identified in the third positions of

COI, rhodopsin and ODH and thus is possible that this contributed to the Vampyromorpha and Octopoda sister-taxon relationship. RY coding also would have aided in reducing the effect of saturation (Phillips & Penny, 2003). However, parsimony analysis of the same dataset recovered a vampyromorph and decapodiform sister-taxon relationship. These results suggest that this relationship is unstable. The lineage Vampyromorpha is supposed to be at least 162 Myr from fossil evidence (Fischer & Riou, 2002) and has been estimated from fossil and molecular data to be potentially 252 Myr (Strugnell et al., 2006). The ancient diversification of this lineage provides support for the supposition that the molecular data used within this study are likely to be saturated at this level (Strugnell et al., 2005). Furthermore, the numerous extinction events throughout the Coleoidea during this time may contribute to the obscuring of affinities of Vampyromorpha (Lindgren et al., 2004).

Order Octopoda: Eledone was traditionally placed within the subfamily Eledoninae because it possesses an ink sac, a single row of suckers and large eggs (Robson, 1929). The taxonomic value of these characters has been debated; the presence of an ink sac has been suggested to be a function of depth (Robson, 1931; Voss, 1988; Allcock & Piertney, 2002) and sucker arrangement has been suggested to be a plastic character (Naef, 1921-1923; Voight, 1993a; Allcock & Piertney, 2002). Allcock & Piertney, (2002) suggested that sub-familial level assignment within the Octopodidae is 'a totally artificial classification with no evolutionary basis.' Eledone has been included in relatively few molecular studies (Bonnaud et al., 1997; Lindgren et al., 2004; Warnke et al., 2004). The present study recovered two differing placements for *Eledone*. All parsimony analyses, and also ML and Bayesian analyses of the POY alignment, show a sister-taxon relationship between Eledone and a clade containing Japetella and Graneledone, thus grouping together all species with a single row of suckers. In contrast ML and Bayesian analyses of data aligned by eye show Eledone to be basal within the Octopoda. This relationship was also recovered by Strugnell (2004), using a subset of the genes used within the present study, but with additional octopod species. Eledone possesses a number of morphological features supporting a basal position within the Octopoda, including the absence of a ligula (Naef, 1921–1923). It must be noted that there are relatively few octopod taxa included within the present study. The inclusion of additional taxa such as Benthoctobus, Bathybolybus and members of the suborder Cirrata would likely improve stability and resolution of octopod relationships.

Suborder Oegopsida and the family Bathyteuthidae: The suborder Oegospida contains squids that possess a gladius and lack a cornea. Molecular studies by Bonnaud et al. (1994, 1997), Carlini & Graves (1999), Carlini et al. (2000) and Lindgren et al. (2004) have suggested that the suborder may be polyphyletic, the later three studies reporting Spirula to fall within the Oegopsida. In contrast, Strugnell et al. (2005) supported a monophyletic Oegopsida. The present study also strongly supports a monophyletic Oegospida, since all alignment and analysis combinations supported this grouping. It is possible that the datasets in the previous studies that suggested a polyphyletic Oegopsida have been too small, and thus contained insufficient information to recover this relationship. All alignment and analysis combinations also support a sister-taxon relationship between the Oegopsida and the family Bathyteuthidae. This supports previous molecular studies by Carlini et al. (2000) and Strugnell et al. (2005) and also agrees with Naef's (1921-1923) suggestion that the Bathyteuthidae possess 'primitive characters for all Oegopsida'.

Suborder Myopsida and Sepioidea: Traditionally Spirulidae, Sepiidae, Idiosepiidae and Sepiadariiae/Sepiolidae have been grouped together in the suborder Sepioidea (Naef, 1921–1923), while the suborder Myopsida was grouped with the suborder Oegopsida in the order Teuthoidea on the basis of similar gladii and tentacular clubs (Naef, 1916, 1921-1923). However, the Myopsida has also been suggested to be derived from the 'Sepioidea' line based on a number of characteristics including possession of a cornea, suckers with circularis muscle, beak without angle point and a vena cava ventral to the intestine (d'Orbigny, 1845; Berthold & Engeser, 1987; Engeser, 1997; Haas, 1997, see Young et al., 1998, for a more detailed discussion). Molecular studies have suggested a close relationship between the Myopsida and some or all members of the Sepioidea (Carlini et al., 2000; Lindgren et al., 2004; Strugnell et al., 2005), although the precise relationship has varied depending upon the genes and analyses used. The present study also suggests a closer relationship between the Myopsida and the Sepioidea than the Myopsida and the Oegopsida, although the exact configuration of this is dependent upon the alignment method and analysis employed. In the phylogenies resulting from data aligned using POY, Myopsida was consistently the sister taxon to a clade containing Spirulida, Bathyteuthoidea and Oegopsida, with the remaining Sepioidea taxa falling outside this clade. However, in the phylogenies resulting from ML and Bayesian analyses of data aligned by eve (RY coded and not RY coded) the Myopsida fell within Sepioidea, together forming a sister taxon to a clade containing the Oegopsida and Bathyteuthidae.

These results clearly show that differing alignment and analysis strategies commonly used in coleoid cephalopod phylogenetics can produce notably different phylogenetic relationships. Researchers are far from agreeing on a single 'best' strategy of phylogenetic analysis, because the advantages and disadvantages of competing strategies are not yet clear. Until such a time, we advocate the use of a variety of different alignment and analysis strategies in phylogenetic analysis.

### ACKNOWLEDGEMENTS

We thanks the following people for their generosity in donating tissue samples: S. von Boletzky, David Carlini, Martin Collins, Stephen Craig, Eileen Dillane, F.G. Hochberg, T. Kubodera, W.K. Macy, Mark Norman, S. Piertney, Richard Stride, Michael Vecchione, Kerstin Warnke and Richard Young. We are also grateful to G. Giribet for running the POY analysis on the Darwin cluster. J.S. is supported by a Natural Environment Research Council Antarctic Funding Initiative grant (NE/C506321/1). M.K.N. is partially supported by NIH SO6 GM008136-32S2 and NSF DEB-0316516.

#### REFERENCES

- AKASAKI, T., NIKAIDO, M., TSUCHIYA, K., SEGAWA, S., HASEGAW, M. & OKADA, N. 2006. Extensive mitochondrial gene arrangements in coleoid Cephalopoda and their phylogenetic implications. *Molecular Phylogenetics and Evolution*, **38**: 648–658.
- ALLCOCK, A.L. & PIERTNEY, S.B. 2002. Evolutionary relationships of Southern Ocean Octopodidae (Cephalopoda: Octopda) and a new diagnosis of *Pareledone. Marine Biology*, **140**: 129–135.
- ANDERSON, F.E. 2000a. Phylogenetic relationships among loliginid squids (Cephalopoda: Myopsida) based on analyses of multiple data sets. *Zoological Journal of the Linnean Society*, **130**: 603–633.
- ANDERSON, F.E. 2000b. Phylogeny and historical biogeography of the Loliginid squids (Mollusca: Cephalopoda) based on mitochondrial DNA sequence data. *Molecular Phylogenetics and Evolution*, 15: 191–214.

- BELSHAW, R. & QUICKIE, D.L.J. 2002. Robustness of ancestral state estimates: Evolution of life strategy in ichneumonoid parasitoids. *Systematic Biology*, **51**: 450–477.
- BERTHOLD, T. & ENGESER, T. 1987. Phylogenetic analysis and systematization of the Cephalopoda (Mollusca). Verhandlungen des naturwissenschaftlichen Vereins Hamburg, 29: 187–220.
- BOLETZKY, S.von 1992. Evolutionary aspects of development, life style, and reproductive mode in incirrate octopods (Mollusca, Cephalopoda). *Revue suisse de Zoologie*, **99**: 755–770.
- BOLETZKY, S.von 2003. Biology of early life stages in cephalopod molluscs. Advances in Marine Biology, 44: 144-184.
- BONNAUD, L., BOUCHER-RODONI, R. & MONNEROT, M. 1994. Phylogeny of decapod Cephalopods based on partial 16S rDNA nucleotide sequences. *Comptes Rendus de l'Académie des Sciences*, **317**: 575–580.
- BONNAUD, L., BOUCHER-RODONI, R. & MONNEROT, M. 1996. Relationship of some coleoid cephalopods established by 3' end of the 16S rRNA and cytochrome oxidase III gene sequence comparison. American Malacological Bulletin, 12: 87–90.
- BONNAUD, L., BOUCHER-RODONI, R. & MONNEROTT, M. 1997. Phylogeny of cephalopods inferred from mitochondrial DNA sequences. Molecular Phylogenetics and Evolution, 7: 44–54.
- BONNAUD, L., RODHOUSE, P.G. & BOUCHER-RODONI, R. 1998. A phylogenetic study of the squid family Onychoteuthidae (Cephalopoda: Oegopsida). Proceedings of the Royal Society of London. Series B, 265: 1761–1770.
- BOUCHER-RODONI, R. & BONNAUD, L. 1996. Biochemical and molecular approach to cephalopod phylogeny. American Malacological Bulletin, 12: 79–85.
- CARLINI, D.B. & GRAVES, J.E. 1999. Phylogenetic analysis of cytochrome c oxidase I sequences to determine higher-level relationships within the coleoid cephalopods. *Bulletin of Marine Science*, **64**: 57–76.
- CARLINI, D.B., REECE, K.S. & GRAVES, J.E. 2000. Actin gene family evolution and the phylogeny of coleoid cephalopods (Mollusca: Cephalopoda). *Molecular Biology and Evolution*, 17: 1353–1370.
- CARLINI, D.B., YOUNG, R.E. & VECCHIONE, M., 2001. A molecular phylogeny of the octopoda (Mollusca:Cephalopoda) evaluated in light of morphological evidence. *Molecular Phylogenetics* and Evolution 21: 388–397.
- CLARKE, M.R. 1988. Evolution of recent cephalopods-A brief review.
  In: The Mollusca. Paleontology and neontology of cephalopods. Vol. 12
  (M.R. Clarke & E.R. Trueman, eds), 1–355. Academic Press, San Diego.
- D'ORBIGNY, A. 1845. Mollusques vivants et fossiles, 2. Paris.
- ENGESER, T. 1997. The fossil Coleoidea page. http://userpage.fuberlin.de/ $\sim$ palaeont/fosscol.html.
- FARRIS, J.S., ALBERT, V.A., KÄLLERSJÖ, M., LIPSCOMB, D. & KLUGE, A.G. 1996. Parsimony jacknifing outperforms neighborjoining. Cladistics, 12: 99–124.
- FELSENSTEIN, J. 1981. Evolutionary trees from DNA sequences: a maximum likelihood approach. Journal of Molecular Evolution, 17: 368-376
- FISCHER, J.C. & RIOU, B. 2002. Vampyronassa rhodanica nov. gen. nov sp., vampyromorphe (Cephalopoda, Coleoidea) du Callovien inférieur de la Voulte-sur-Rhône (Ardèche, France). Annales de Paléontologie, **88**: 1–17.
- GILLESPIE, J.J., YODER, M.J. & WHARTON, R.A. 2005. Predicted secondary structures for 28S and 18S rRNA from Ichneumonoidea (Insecta: Hymenoptera: Apocrita): impact on sequence alignment and phylogeny estimation. Journal of Molecular Evolution, 61:114–137.
- GIRIBET, G. 2003. Stability in phylogenetic formulations, and its relationship to nodal support. Systematic Biology, 52: 554–564.
- GIRIBET, G. 2005. Generating implied alignments under direct optimization using POY. Cladistics, 21: 396–402.
- GIRIBET, G., DESALLE, R. & WHEELER, W.C. 2002. 'Pluralism' and the aims of phylogenetic research. In: Molecular systematics and

- evolution: theory and practice (R. DeSalle, G. Giribet, & W.C. Wheeler, eds), 141–146. Birkhauser, Basel.
- GIRIBET, G. & WHEELER, W.C. 1999. On gaps. Molecular Phylogenetic and Evolution, 13: 132–143.
- GOLOBOFF, P.A. 1999. Analyzing large data sets in reasonable times: solutions for composite optima. *Cladistics*, 15: S26–S34.
- GRIMPE, G. 1921. Teuthologische Mitteilungen. VII:Systematiche Ubersicht der Nordsee-Cephalopoden. Zoologischer Anzeiger. 52: 297–305.
- GUZIK, M.T., NORMAN, M.D. & CROZIER, R.H. 2005. Molecular phylogeny of the benthic shallow-water octopuses (Cephalopoda: Octopodinae). Molecular Phylogenetic and Evolution, 37: 235–248.
- HAAS, W. 1997. Der Ablauf der Entwicklungsgeschichte der Decabrachia (Cephalopoda, Coleoidea). *Palaeontographica*, **245A**: 63–81.
- HEALY, J.M. 1989. Spermatozoa of the deep-sea cephalopod *Vampyroteuthis infernalis* Chun: ultrastructure and possible phylogenetic significance. *Philosophical Transactions of the Royal Society, Series B*, **323**: 589–600.
- HUELSENBECK, J.P. & BOLBACK, J.P. 2001. Empirical and hierchical Bayesian estimation of ancestral states. Systematic Biology, 50: 351–366.
- HUELSENBECK, J.P. & HILLIS, D.M. 1993. Success of phylogenetic methods in the four-taxon case. Systematic Biology, 40: 247–264.
- HUELSENBECK, J.P. & IMENNOV, N.S. 2002. Geographic origin of mitochondrial DNA: accommodating phylogenetic uncertainty and model comparison. Systematic Biology, 51: 155–165.
- HUELSENBECK, J.P. & RONQUIST, F. 2001. MrBayes:Bayesian inference of phylogeny. *Bioinformatics*, **17**: 754–755.
- KHROMOV, D.N. 1990. Cuttlefishes in the systematics and phylogeny of Cephalopoda. *Zoologicheskii Zhurnal*, **69**: 12–20.
- KJER, K.M. 1995. Use of rRNA secondary structure in phylogenetic studies to identify homologous positions: an example of alignment and data presentation from the frogs. *Molecular Phylogenetics and Evolution*, **4**: 314–330.
- KJER, K.M., GILLESPIE, J.J., & OBER, K.A. 2007. Opinions on multiple sequence alignment, and an empirical comparison of repeatability and accuracy between POY and structural alignment. Systematic Biology, 56: 133–146.
- LINDGREN, A.R., GIRIBET, G. & NISHIGUCHI, M.K. 2004. A combined approach to the phylogeny of Cephalopoda (Mollusca). *Cladistics*, 20: 454–486.
- LINDGREN, A.R., KATUGIN, O.N., AMEZQUITA, E. & NISHIGUCHI, M.K. 2005. Evolutionary relationships among squids of the family Gonatidae (Mollusca: Cephalopoda) inferred from three mitochondrial loci. Molecular Phylogenetics and Evolution, 26: 101–111.
- NAEF, A. 1916. Uber neue Sepioliden aus dem Golf von Neapel. Pubblicazioni della Stazione Zooloica di Napoli, 1: 1-10.
- NAEF, A. 1921–1923. Cephalopoda. Fauna e flora del Golfo di Napoli, Monograph. (translated from German by the Israel program for Scientific translations, Jerusalem 1972).
- NAEF, A. 1928. *Die Cephalopoden. Vol. 2. Embryologie* Fauna e Flora del Golfo di Napoli, Monographie 35.
- NISHIGUCHI, M.K. & MAPES, R. 2007. Cephalopoda. In: *Molluscan evolution* (W. Ponder & D. Lindberg, eds), 161–197. University of California Press, Berkeley, California.
- NISHIGUCHI, M.K., LOPEZ, J.E. & BOLETZKY, S.von 2004. Enlightenment of old ideas from new investigations: more questions regarding the evolution of bacteriogenic light organs in squids. *Evolution and Development*, **6**: 41–49.
- PAGE, R.D.M. & HOLMES, E.C. 1998. Molecular evolution. A phylogenetic approach. Blackwell Science, Oxford.
- PHILLIPS, M.J., LIN, Y.-H., HARRISON, G.L. & PENNY, D. 2001. Mitochondrial genomes of a bandicoot and a brushtail possum confirm the monophyly of australidelphian marsupials. *Proceedings* of the Royal Society of London, Series B, 268: 1533–1538.
- PHILLIPS, M.J. & PENNY, D. 2003. The root of the mammalian tree inferred from whole mitochondrial genomes. *Molecular Phylogenetics and Evolution*, **28**: 171–185.

- PICKFORD, G.E. 1939. The Vampyromorpha. A new order of dibranchiate Cephalopoda. Věstník Československé zoologické společnosti, 6-7: 346-358.
- PIERTNEY, S.B., HUDELOT, C., HOCHBERG, F.G. & COLLINS, M.A. 2003. Phylogenetic relationships among cirrate octopods (Mollusca: Cephalopoda) resolved using mitochondrial 16S ribosomal DNA sequences. Molecular Phylogenetics and Evolution, 27: 348–353
- POL, D. & SIDDALL, M.E. 2001. Biases in maximum likelihood and parsimony: a simulation approach to a 10-taxon case. *Cladistics*, 17: 266-281.
- PONS, J. & VOGLER, A.P. 2006. Size, frequency, and phylogenetic signal of multiple-residue indels in sequence alignment of introns. *Cladistics*, **22**: 144–156.
- RAMBAUT, A. 2002. Se-Al. Ver. v2.0a11 Carbon. Oxford University. RAMBAUT, A. & DRUMMOND, A.J. 2003. Tracer. v1.0.1. Oxford University.
- ROBSON, G.C. 1929. A monograph of Recent Cephalopoda. Part I. Octopodinae. British Museum (Natural History), London.
- ROBSON, G.C. 1931. A monograph of the Recent Cephalopoda. Part II. The Octopoda (excluding the Octopodinae). British Museum (Natural History), London.
- RONQUIST, F. & HUELSENBECK, J.P. 2003. MRBAYES 3: Bayesian phylogenetic inference under mixed models. *Bioinformatics*, **19**: 1572–1574.
- SAITOU, N. & NEI, M. 1987. The neighbor-joining method a new method for reconstructing phylogenetic trees. *Molecular Biology and Explusion*. 4: 406–425.
- SHULL, V.L., VOGLER, A.P., BAKER, M.D., MADDISON, D.R. & HAMMOND, P.M. 2001. Sequence alignment of 18S ribosomal RNA and the basal relationships of adephagan beetles: evidence for monophyly of aquatic families and the placement of Trachypachidae. *Systematic Biology*, **50**: 945–969.
- SIMMONS, M.P., PICKETT, K.M. & MIYA, M. 2004. How meaningful are Bayesian support values? *Molecular Biology and Evolution*, **21**: 188–199.
- STRUGNELL, J. 2004. The molecular evolutionary history of the Class Cephalopoda (Phylum Mollusca). DPhil thesis, Department of Zoology, University of Oxford.
- STRUGNELL, J., JACKSON, J., DRUMMOND, A.J. & COOPER, A. 2006. Divergence time estimates for major cephalopod groups: evidence from multiple genes. *Cladistics*, **22**: 89–96.
- STRUGNELL, J., NORMAN, M., DRUMMOND, A.J., JACKSON, J. & COOPER, A. 2005. Molecular phylogeny of coleoid cephalopods (Mollusca: Cephalopoda) using a multigene approach; the effect of data partitioning on resolving phylogenies in a Bayesian framework. *Molecular Phylogenetics and Evolution*, 37: 426–441.
- STRUGNELL, J.M., NORMAN, M.D., DRUMMOND, A.J. & COOPER, A. 2004. The octopuses that never came back to earth: neotenous origins for pelagic octopuses. *Current Biology*, **18**: R300–R301.
- SUZUKI, Y., GLAZKO, G.V. & NEI, M. 2002. Overcredibility of molecular phylogenies obtained by Bayesian phylogenetics. *Proceedings of the National Academy of Science of the USA*, **99**: 16138–16143.
- SWOFFORD, D.L. 1998. PAUP\*4.0-Phylogenetic Analysis Using Parsimony (\*and other methods). Sinauer, Sunderland, MA.
- TAKUMIYA, M., KOBAYASHI, M., TSUNEKI, K. & FURUYA, H. 2005. Phylogenetic relationships among major species

- of Japanese coleoid cephalopods (Mollusca: Cephalopoda) using three mitochondrial DNA sequences. *Zoological Science*, **22**: 147–155.
- TOLL, R.B. 1982. The comparative morphology of the gladius in the order Teuthodea (Mollusca: Cephalopoda) in relation to systematics and phylogeny. PhD thesis, University of Miami.
- TOLL, R.B. 1998. The gladius in teuthoid systematics. In: *International workshop on the systematics and biogeography of cephalopods*, 65–68. Smithsonian Institution Press, Washington D.C.
- VOIGHT, J.R. 1993a. The arrangement of suckers on octopodid arms as a continuous character. *Malacologia*, **35**: 351–359.
- VOIGHT, J.R. 1993b. A cladistic reassessment of Octopodid classification. *Malacologia*, 35: 343–349.
- VOIGHT, J.R. 1997. Cladistic analysis of the octopods based on anatomical characters. Journal of Molluscan Studies, 63: 311–325.
- VOSS, G.L. 1977. Present status and new trends in cephalopod systematics. In: *The biology of cephalopods* (M. Nixon & J.B. Messenger, eds), 49–60. Academic Press, London.
- VOSS, G.L. 1988. Evolution and phylogenetic relationships of deep-sea octopods (Cirrata and Incirrata). In: The Mollusca, 12: Paleontology and neontology of cephalopods (M.R. Clarke & E.R. Trueman, eds), 253–276. Academic Press, San Diego.
- WARNKE, K., PLÖTNER, J., SANTANA, J.I., RUEDA, M.J. & LLINAS, O. 2003. Reflections on the phylogenetic position of *Spirula* (Cephalopoda): preliminary evidence from the 18S ribosomal RNA gene. *Berliner Paläobiologisch Abhandlungen*, **3**: 253–260.
- WARNKE, K., SOLLER, R., BLOHM, D. & SAINT-PAUL, U. 2004. A new look at geographic and phylogenetic relationships within the species group surrounding Octopus vulgaris (Mollusca, Cephalopoda): indications of very wide distribution from mitochondrial DNA sequences. Journal of Zoological Systematics and Evolutionary Research, 42: 306–312.
- WHEELER, W.C. 1995. Sequence alignment, parameter sensitivity, and the phylogenetic analysis of molecular data. *Systematic Biology*, **44**: 321–331.
- WHEELER, W.C. 1996. Optimization alignment: the end of multiple sequence alignment in phylogenetics? *Cladistics*, **12**: 1–9.
- WHEELER, W.C. 2003. Implied alignment: a synapomorphy-based multiple-sequence alignment method and its use in cladogram search. *Cladistics*, **19**: 261–268.
- WHEELER, W.C. & HAYASHI, C.Y. 1998. The phylogeny of extant chelicerate orders. *Cladistics*, 14: 173–192.
- WHEELER, W.C., GLADSTEIN, D.S. & LAET, J.D. 1996–2003. POY, v.3.0.11 ftp.amnh.org/pub/molecular/poy Wheeler.
- YOKOBORI, S.-I., FUKUDA, N., NAKAMURA, M., AOYAMA, T. & OSHIMA, T. 2004. Long-term conservation of six duplicated structural genes in cephalopod mitochondrial genomes. *Molecular Biology and Evolution*, 22: 2034–2046.
- YOUNG, R.E. & VECCHIONE, M. 1996. Analysis of morphology to determine primary sister-taxon relationships within celeoid cephalopods. *American Malacological Bulletin*, **12**: 91–112.
- YOUNG, R.E., VECCHIONE, M. & DONOVAN, D.T. 1998. The evolution of coleoid cephalopods and their present biodiversity and ecology. South African Journal of Marine Science, 20: 393–420.
- ZHENG, X., YANG, J., LIN., X. & WANG, R. 2004. Phylogenetic relationships among the decabrachia cephalopods inferred from mitochondrial DNA sequences. *Journal of Shellfish Research*, 23: 881–886.