BMC Evolutionary Biology



Research article Open Access

Phylogenetic placement of the enigmatic parasite, *Polypodium* hydriforme, within the Phylum Cnidaria

Nathaniel M Evans¹, Alberto Lindner², Ekaterina V Raikova³, Allen G Collins⁴ and Paulyn Cartwright*¹

Address: ¹Department of Ecology and Evolutionary Biology, University of Kansas, Lawrence, Kansas 66045, USA, ²CEBIMar, University of São Paulo, São Sebastião, Brazil, ³Institute of Cytology of the Russian Academy of Sciences, St. Petersburg, Russia and ⁴National Systematics Laboratory of NOAA Fisheries Service, National Museum of Natural History, Smithsonian Institution, Washington, DC 20013-7012, USA

Email: Nathaniel M Evans - evansnat@ku.edu; Alberto Lindner - alindner@usp.br; Ekaterina V Raikova - raikova@swipnet.se; Allen G Collins - COLLINSA@si.edu; Paulyn Cartwright* - pcart@ku.edu

* Corresponding author

Published: 9 May 2008

Received: 4 December 2007 Accepted: 9 May 2008

BMC Evolutionary Biology 2008, 8:139 doi:10.1186/1471-2148-8-139

This article is available from: http://www.biomedcentral.com/1471-2148/8/139

© 2008 Evans et al; licensee BioMed Central Ltd.

This is an Open Access article distributed under the terms of the Creative Commons Attribution License (http://creativecommons.org/licenses/by/2.0), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

Abstract

Background: Polypodium hydriforme is a parasite with an unusual life cycle and peculiar morphology, both of which have made its systematic position uncertain. Polypodium has traditionally been considered a cnidarian because it possesses nematocysts, the stinging structures characteristic of this phylum. However, recent molecular phylogenetic studies using 18S rDNA sequence data have challenged this interpretation, and have shown that Polypodium is a close relative to myxozoans and together they share a closer affinity to bilaterians than cnidarians. Due to the variable rates of 18S rDNA sequences, these results have been suggested to be an artifact of long-branch attraction (LBA). A recent study, using multiple protein coding markers, shows that the myxozoan Buddenbrockia, is nested within cnidarians. Polypodium was not included in this study. To further investigate the phylogenetic placement of Polypodium, we have performed phylogenetic analyses of metazoans with 18S and partial 28S rDNA sequences in a large dataset that includes Polypodium and a comprehensive sampling of cnidarian taxa.

Results: Analyses of a combined dataset of 18S and partial 28S sequences, and partial 28S alone, support the placement of *Polypodium* within Cnidaria. Removal of the long-branched myxozoans from the 18S dataset also results in *Polypodium* being nested within Cnidaria. These results suggest that previous reports showing that *Polypodium* and Myxozoa form a sister group to Bilateria were an artifact of long-branch attraction.

Conclusion: By including 28S rDNA sequences and a comprehensive sampling of cnidarian taxa, we demonstrate that previously conflicting hypotheses concerning the phylogenetic placement of *Polypodium* can be reconciled. Specifically, the data presented provide evidence that *Polypodium* is indeed a cnidarian and is either the sister taxon to Hydrozoa, or part of the hydrozoan clade, Leptothecata. The former hypothesis is consistent with the traditional view that *Polypodium* should be placed in its own cnidarian class, *Polypodiozoa*.

Background

Polypodium hydriforme is an endocellular parasite whose unusual life cycle, peculiar morphology, and high rates of DNA evolution, have led to much controversy regarding its phylogenetic position within metazoans [1-5]. Polypodium spends most of its life inside the oocytes of acipenseriform fishes (sturgeons and paddlefish). During this time, Polypodium develops from a binucleate cell into an inside-out planuliform larva and then into an elongate inside-out stolon; the epidermal cell layer is located internal to the body and the gastrodermis is located externally [6-8]. The embryo, larva and stolon are surrounded by a protective polyploid cell, which also functions in digestion [7]. Just prior to host spawning, Polypodium everts to the normal position of cell layers, revealing tentacles scattered along the stolon. During eversion, the yolk of the host oocyte fills the gastral cavities of the parasite, supplying the future free-living stage with nutrients [6,7]. Finally, upon emerging from the host egg in fresh water, the freeliving stolon (Figure 1A) fragments into individual medusoid-like forms (Figure 1B) that go on to multiply by means of longitudinal fission, form sexual organs, and ultimately infect host fish with their gametophores [6-9].

Two conflicting hypotheses have been proposed regarding the phylogenetic placement of *Polypodium*. The first, more

traditional hypothesis is that *Polypodium* is a cnidarian. Some have suggested it is nested within a derived group of hydrozoans, the Narcomedusae [10-13] or the cnidarian class Scyphozoa [14]; while others have suggested it belongs to a separate cnidarian class, Polypodiozoa [1,15,16]. The assignment of Polypodium to Cnidaria is based primarily on morphological evidence, most notably the fact that Polypodium possesses nematocysts [17,18], the stinging structures characteristic of all cnidarians. In addition, the presence of tentacles and overall body-plan organization of Polypodium are reminiscent of cnidarians, although it is unclear if the adult free-living stage is homologous to a polyp or medusa stage. This hypothesis is supported by a cladistic analysis of small subunit nuclear ribosomal DNA (18S rDNA) sequences in conjunction with morphological characters (including nematocysts) [2]. In this study, Polypodium falls within the medusozoan clade of cnidarians, although the non-cnidarian placozoan, Trichoplax [19,20], also fell within this clade, rendering Cnidaria paraphyletic.

The second hypothesis is that *Polypodium* is the sister taxon to Myxozoa, a diverse group of parasites in aquatic animals, and that *Polypodium* + Myxozoa is the sister group to Bilateria [2-4]. This hypothesis is derived from cladistic analyses utilizing 18S rDNA sequences [2-4]. However,

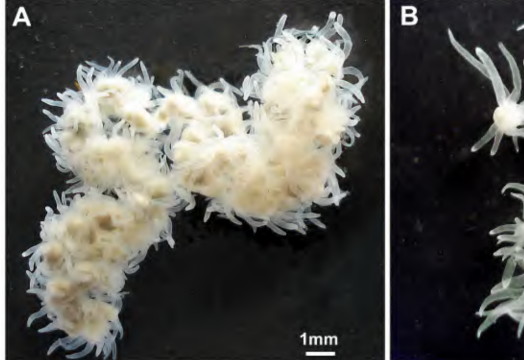




Figure 1

Polypodium hydriforme. A) Stolon stage just after emerging from the host oocyte. B) Four specimens of free-living Polypodium with 12 tentacles. Photos by E. Raikova.

because Polypodium and myxozoans have unusually high divergence rates in their 18S rDNA sequences, these cladistic analyses have been criticized by a number of authors who suggest that the data might be unduly affected by long-branch attraction (LBA) [5,21,22]. Despite some attempts to overcome the effects of LBA through the use of a maximum likelihood (ML) approach [21-23] and pruning long branches [5,22], these results have been largely silent on the placement of Polypodium. For instance, Kim et al. [22] applied a maximum likelihood approach to 18S rDNA sequence data and found that myxozoans and Polypodium did not group together. Instead, *Polypodium* was part of an unresolved polytomy that included several cnidarian lineages and Trichoplax, as well as myxozoans + Bilateria. Most recently, Jimenez-Guri et al. [24] utilized multiple protein-coding gene sequences in a ML analysis and found the myxozoan, Buddenbrockia plumatellae nested within cnidarians. Unfortunately, this study had relatively limited sampling of cnidarians and did not include Polypodium.

In an attempt to resolve this controversy, we sequenced an additional marker in *Polypodium*, a partial gene sequence of the large nuclear ribosomal unit (28S rDNA), and greatly expanded the taxonomic sampling of cnidarian sequences. Using this approach, we provide evidence that *Polypodium* is nested within Cnidaria and does not group with myxozoans.

Results

Sampled taxo

All taxa used in this study are arranged taxonomically in Table 1. 155 sequences were obtained from GenBank. 45 new cnidarian sequences for 18S and 59 for 28S (including 2 new 18S and 2 new partial 28S from Polypodium taxa) were generated for this study and deposited in Gen-Bank (see Table 1 for accession numbers). Polypodium hydriforme sequences were obtained from both North American and Eurasian hosts. Eurasian samples were collected from two individuals of Acipenser ruthenus. North American samples were collected from *Polyodon spathula* and Scaphirhynchus platorynchus. This is the first reported presence of Polypodium infection in Scaphirhynchinae. While *Polypodium* was recovered from the oocytes of *S. pla*torynchus, the sample from which we extracted sequence data was found externally attached to its presumed host. More specific collection data for *Polypodium* specimens are associated with each sequence submitted to GenBank (see Table 1 for accession numbers).

All *Polypodium* sequences were newly generated for this study. We did not include the previously published 18S *Polypodium* sequence (GenBank accession number <u>II37526</u>) because of concern over the quality of the sequence which included a number of ambiguities. Fur-

thermore, while the two new *Polypodium* 18S sequences (from hosts *Acipenser ruthensus* and *Polyodon spathula*) differed from each other by a total of 8 sites they differed from #U37526 by 77 and 83 sites respectively. These differences included a large number of insertions and deletions. The two new 28S sequences (from hosts *Acipenser ruthensus* and *Scaphirhynchus platorynchus*) only differed from each other by 2 sites.

Position of Polypodium

The complete combined dataset of 18S rDNA and partial 28S rDNA contains 4842 characters, 2901 of which are variable and 2124 parsimony informative. Both the ML and parsimony topologies reconstructed from the combined dataset suggest that Polypodium is nested within a monophyletic Cnidaria, and myxozoans are the sister taxon to bilaterians (Figure 2). The ML bootstrap values supporting a monophyletic Cnidaria (including Polypodium), a monophyletic Medusozoa (including Polypodium) and the Polypodium + hydrozoan clade are 73, 67 and 73 respectively (Figure 2A, and Additional file 1). Parsimony analysis of the combined dataset differs from that of ML in that Polypodium is nested within a group of hydrozoans, the leptothecates (Figure 2B). The parsimony bootstrap values supporting a monophyletic Cnidaria and Hydrozoa, with *Polypodium* nested within these clades are 50 and 51 respectively (Figure 2B). The clade nested within hydrozoans, that includes Polypodium + leptothecates is weakly supported in the sub-sampling tests with a bootstrap value of less than 50.

The analyses using partial 28S rDNA sequences alone (129 sampled taxa) contains 1756 characters, 1196 of which are parsimony informative. The ML topology using this dataset reveals Polypodium nested within Cnidaria, specifically within leptothecate hydrozoans, (Additional file 2). This analysis however fails to recover a monophyletic Cnidaria, as the anthozoans are placed outside the Cnidaria + Bilateria clade. Analysis of the 18S rDNA dataset alone (132 taxa, 3038 characters, 1469 parsimony informative) under both optimality criteria conflicts with the combined and partial 28S topologies. The 18S rDNA topology for both criteria place Polypodium at the base of Bilateria (Figure 3A, Additional files 3, 4 and 5). However, the ML topology also reflects a sister relationship between Polypodium and myxozoans (Figure 3A and Additional file 3A) while the parsimony topology does not (Additional files 4 and 5). Moreover, under parsimony criteria the position of myxozoans is dependent upon how gaps are coded: if gaps are coded as a fifth character state, myxozoans are placed as a highly derived clade of bilaterians (Additional file 4); if gaps are coded as missing, myxozoans are placed as sister to all metazoans (Additional file 5). The 18S analysis showing placement of Polypodium with Bilateria, and more specifically as sister to myxo-

Table I: Taxon and sequence list

Higher classification	Taxon ID	Accessio		
		285	185	Voucher
Bilateria				
Annelida	Proceraea cornuta	AF212165	AF212179	
Annelida	Urechis caupo	AF342804	AF342805	
Arthropoda	Limulus polyphemus	AF212167	<u>U91490</u>	
Arthropoda	Tenebrio sp./Tenebrio molitor	AY210843	X07801	
Brachiopoda	Phoronis vancouverensis	AF342797	<u>U12648</u>	
Chordata	Oncorhynchus sp./O. kisutch	<u>U34341</u>	AF030250	
Chordata	Petromyzon marinus	AF061798	M97575.1	
Chordata	Raja schmidti	AF278683	AF278682	
Chordata	Triakis semifasciata	AF212182	AF212180	
chinodermata	Strongylocentrotus purpuratus	AF212171	L28056.	
lemichordata	Cephalodiscus gracilis	AF212172	AF236798	
lemichordata	Harrimania sp.	AF212173	AF236799	
lemichordata	Ptychodera flava	AF212176	AF278681	
lemichordata	Ptychoderidae	AF278684	D14359	
Hemichordata	Saccoglossus kowalevskii	AF212175	L28054	
Kinorhyncha	Pycnophyes sp.Tjarno	AY859597	AY859598	
10llusca	Parvicardium minimum	DQ279966	DO279942	
10llusca			X53899	
	Placopecten magellanicus	AF342798		
Nematoda	Caenorhabditis elegans	X03680	X03680	
Nematomorpha	Chordodes morgani	AF342787	AF036639	
Nemertea	Amphiporus sp.	AF342786	AF119077	
Nemertodermatida	Meara stichopi	AY157605	AF119085	
Onychophora	Peripatus sp.	AY210836	AY210837	
Platyhelminthes	Diclidophora denticulata	<u>AY157169</u>	<u>AJ228779</u>	
Platyhelminthes	Stenostomum leucops	<u>AYI57151</u>	D85095	
Platyhelminthes	Stylochus zebra	<u>AF342800</u>	<u>AF342801</u>	
Priapulida	Priapulus caudatus	<u>AY210840</u>	<u>Z38009</u>	
ipuncula	Phascolopsis gouldii	<u> AF342795</u>	<u> AF342796</u>	
⁻ ardigrada	Milnesium.sp.\M. tardigradum	<u>AY210826</u>	<u>U49909</u>	
Jrochordata	Styela plicata	<u> AFI58724</u>	<u>L12444</u>	
Jrochordata	Thalia democratica	<u>AFI58725</u>	D14366	
Cnidaria				
olypodiozoa	Polypodium (Host: Acipenser ruthenus)	EU272585	EU272630	
Polypodiozoa	Polypodium (Host: Polyodon spathula)		EU272629	
Polypodiozoa	Polypodium (Host:Scaphirhynchus platorynchus)	EU272586		
Anthozoa, Antipatharia	Antipathes galapagensis	AY026365	<u>AF100943</u>	
Anthozoa, 5cleractinia	Montastraea franksi	AY026375	AY026382	
Cubozoa, Carybdeidae	Carybdea rastonii	AY920787	AF358108	
Cubozoa, Carybdeidae	Darwin carybdeid sp.	AY920788	AF358105	
Cubozoa, Carybdeidae	Tripedalia cystophora	EU272595	EU272637	
Cubozoa, Chirodropidae	Chironex fleckeri	AY920785	AF358104	
Cubozoa, Chirodropidae	Chiropsalmus sp.	AY920786	AF358103	
lydrozoa, Capitata	Dipurena ophiogaster	EU272560	EU272615	KUNHM 2803
lydrozoa, Capitata	Ectopleura dumortieri	EU272561	EU272616	
lydrozoa, Capitata	Euphysora bigelowi	EU272563	EU272618	KUNHM 2829
lydrozoa, Capitata	Moerisia sp.	AY920801	AF358083	
lydrozoa, Capitata	Pennaria disticha	EU272581	AY920762	
lydrozoa, Capitata	Polyorchis penicillatus		AF358090	
lydrozoa, Capitata Iydrozoa, Capitata	Porpita sp.	AY920803	AF358086	
lydrozoa, Capitata Iydrozoa, Capitata	Ralpharia gorgoniae	EU272590	EU272633	KUNHM 2778
lydrozoa, Capitata lydrozoa, Capitata	Scrippsia pacifica	AY920804	AF358091	NOINI II Z//0
lydrozoa, Capitata Iydrozoa, Capitata	Solanderia ericopsis	EU272593	EU272636	MHNG INVE29593
	•			TIMING INVEZ7373
lydrozoa, Capitata	Velella sp.	EU272597	AF358087	VIINIUM 2702
Hydrozoa, Capitata	Zanclea prolifera	EU272598	EU272639	KUNHM 2793
Hydrozoa, Capitata	Zyzzyzus warreni	EU272599	EU272640	KUNHM 2777
Hydrozoa, Capitata	Candelabrum cocksii	<u>AY920796</u>	<u>AY920758</u>	MHNG INVE2953 I

Table I: Taxon and sequence list (Continued)

Hydrozoa Capitata	Cladaranne florrasa	EU272551	EU272608	
Hydrozoa, Capitata Hydrozoa, Filifera	Cladacaryne flaccasa Bimerio vestita	EU272551 EU272548	EU272608 EU272605	
Hydrozoa, Filifera	Baugainvillia carolinensis	EU272549	EU272606	
Hydrozoa, Filifera	Brinckmannia hexoctinellidaphila	EU272550	EU272607	MHNG INVE38148
Hydrozoa, Filifera	Clava multicarnis	EU272552	EU272609	1 11 11 10 11 11 2501 10
Hydrozoa, Filifera	Clovactinia gallensis	EU272553	EU272610	MHNG INVE33470
Hydrozoa, Filifera	Cardylaphara caspia	EU272556	EU272612	1 11 11 10 11 11 233 11 0
Hydrozoa, Filifera	Corydendrium sp.	EU272557	EU272613	KUNHM 2764
Hydrozoa, Filifera	Dicaryne canybearei	EU272559	EU272614	MHNG INVE32949
Hydrozoa, Filifera	Eudendrium.rocemosum	EU272562	EU272617	
Hydrozoa, Filifera	Fabienna sphaerica	AY920797	AY920767	
Hydrozoa, Filifera	Gorveio annulata/Gorveia sp.	EU272564	AY920766	KUNHM 2860
Hydrozoa, Filifera	Hydro circumcincto	AY026371	AF358080	
Hydrozoa, Filifera	Hydroctinio symbialangicarpus	EU272568	EU272621	
Hydrozoa, Filifera	Hydrichthella epigorgia	EU272569	EU272622	KUNHM 2665
Hydrozoa, Filifera	Hydrichthys boycei	EU272570		MHNG INVE37417
Hydrozoa, Filifera	Kaellikerina fasciculata	EU272571	EU272623	
Hydrozoa, Filifera	Leuckartiara octana	EU272573	EU272624	
Hydrozoa, Filifera	Lizzio blandina	EU272574	EU272625	
Hydrozoa, Filifera	Pochycordyle pusillo	EU272579	EU272627	MHNG INVE32953
Hydrozoa, Filifera	Pondeo sp.	EU272580	AY920765	
Hydrozoa, Filifera	Padacoryne cornea	AY920802	AF358092	
Hydrozoa, Filifera	Prabascidactyla ornata	EU272587	EU272631	KUNHM 2767
Hydrozoa, Filifera	Pruvatella grisea	EU272588	EU272632	MHNG INVE34436
Hydrozoa, Filifera	Rothkea octapunctata	EU272591	EU272634	KUMIP 314321
Hydrozoa, Filifera	Rhizagetan nudus	EU272592	EU272635	MHNG INVE35757
Hydrozoa, Filifera	Turritopsis dohrnii	EU272596	EU272638	MHNG INVE29753
Hydrozoa, Leptothecata	Abietinoria filicula	EU272540	EU272600	MHNG INVE29947
Hydrozoa, Leptothecata	Aglaaphenia tubifarmis	EU272543	EU272601	MHNG INVE29967
Hydrozoa, Leptothecata	Amphisbetia minima	EU272544	EU272602	MHNG INVE25071
Hydrozoa, Leptothecata	Anthahebella parasitica	EU272545	EU272603	MHNG INVE29762
Hydrozoa, Leptothecata	Clytia naliformis	EU272554	EU272611	
Hydrozoa, Leptothecata	Holecium muricotum	EU272565	EU272619	MHNG INVE29028
Hydrozoa, Leptothecata	Halopteris minuta	EU272567	EU272620	MHNG INVE25073
Hydrozoa, Leptothecata	Melicertum octocostatum	EU272575	AY920757	USNM 1073342
Hydrozoa, Leptothecata	Octophialucium indicum	EU272577	EU272626	MHNG INVE29970
Hydrozoa, Leptothecata	Plumulorio setoceo	EU272583	EU272628	MHNG INVE36298
Hydrozoa, 5iphonophorae	Agolmo elegans	EU272542	AY937313	YPM 35029
Hydrozoa, 5iphonophorae	Apalemia sp.	EU272546	AY937331	YPM 35090
Hydrozoa, 5iphonophorae	Cardagalma cardiforme	EU272555	AY937317	YPM 35032
Hydrozoa, 5iphonophorae	Halistemma rubrum	EU272566	AY937358	YPM 35359
Hydrozoa, 5iphonophorae	Nonomia bijugo	EU272576	AY937338	YPM 35043
Hydrozoa, 5iphonophorae	Nectopyromis sp.	AY026377	AF358068	
Hydrozoa, Siphonophorae	Physophara hydrastatica	<u>EU272582</u>	AY937342	YPM 35046
Hydrozoa, 5iphonophorae	Sulculeoloria quodrivalvis	<u>EU272594</u>	AY937353	YPM 35357
Hydrozoa, 5tylasteridae	Crypthelia cryptatrema	<u>EU272558</u>	<u>EU272641</u>	USNM1027758
Hydrozoa, Stylasteridae	Lepidopora micrastylus	<u>EU272572</u>	EU272644	USNM1027724
Hydrozoa, 5tylasteridae	Pseudocrypthelio pachypama	EU272589	EU272643	U5NM1027728
Hydrozoa, 5tylasteridae	Adelapara crassilabrum	EU272541	EU272642	U5NM1027760
Hydrozoa, Trachylina	Limnocnido tongonyicoe	AY920795	AY920755	
Hydrozoa, Trachylina	Maeatias marginata	<u>EU247810</u>		
Hydrozoa, Trachylina	Olindias phospharica	EU247808	AY920753	MHNG INVE29811
5cyphozoa, Coronatae	Atollo vanhaeffeni	AY026368	AF100942	
Scyphozoa, Coronatae	Nousithoe rubro	<u>AY920776</u>	AF358095	
5cyphozoa, Rhizostomea	Cotostylus sp.	<u>AY920777</u>	AF358100	
5cyphozoa, 5emaeostomeae	Chrysoora melonoster	AY920780	AF358099	
Scyphozoa, Semaeostomeae	Aurelio sp.	<u>EU272547</u>	EU272604	
5cyphozoa, 5emaeostomeae	Phocellophara camtschatica	<u>AY920778</u>	AF358096	
	Croteralophus canvolvulus	<u>AY920781</u>	AY845344	
Staurozoa, Stauromedusae				
	Haliclystus actorodiatus	<u>AH014894</u>	<u>AY845346</u>	
Staurozoa, Stauromedusae Staurozoa, Stauromedusae Staurozoa, Stauromedusae	•	<u>AH014894</u> <u>AY920782</u>	AY845346 AF358102	
Staurozoa, Stauromedusae	Haliclystus actorodiatus			

Table I: Taxon and sequence list (Continued)

Myxosporea	Henneguyo salminicala	AY302726		
Myxosporea	Kudoo trifolio	AM490336	AM183300	
Myxosporea	Kudoo unicopsulo	AM490335	AM490334	
Myxosporea	Myxobalus cerebralis		EF370481	
Myxosporea	Myxabalus dagieli		EU003978	
Myxosporea	Parvicapsula limondae		EF429096	
Outgroups	· ····			
Choanoflagellida				
Codonosigidae	Manasiga brevicollis	AY026374	AF084618	
Salpingoecidae	Solpingoeco infusionum	AY026380	<u>AF100941</u>	
Ctenophora,	,			
Cyclocoela	Beroe ovota	AY026369	AF293694	
Cyclocoela	Mnemiopsis leidyi	AY026373	AF293700	
Typhlocoela	Pleurabrachia bachei	AY026378	AF293677	
Fungi				
Ascomycota	Condido albicans	<u>×70659</u>	X53497	
Ascomycota	Saccharomyces cerevisiae	101355	M27607	
Basidiomycota	Tricholomo matsutake	<u>U62964</u>	<u>U62538.1</u>	
Mucoromycotina	Mucor rocemosus	<u> </u>	A]271061	
Porifera,				
Calcarea	Leucosolenia sp.	AY026372	AF100945	
Demospongia	Mycole fibrexilis	AY026376	AF100946	
Demospongia	Suberites ficus	AY026381	AF100947	

A complete list of sequences used in the analyses with GenBank accession numbers and museum voucher numbers. Bold numbers indicate new sequences generated for this study. KUMIP = University of Kansas Museum of Invertebrate Paleontology, KUNHM = University of Kansas Natural History Museum, MHNG = Muséum d'histoire naturelle de Genève, YPM = Yale Peabody Museum, USNM = U5 National Museum of Natural History.

zoans, is consistent with previously reported studies using the same marker [2-4], but raises similar concerns of longbranch attraction [5].

Test of long-branch attraction

Myxozoans and *Polypodium* have unusually high rates of evolution in their 18S and 28S rDNA sequences relative to the other sampled taxa. To investigate the influence of myxozoans on the placement of *Polypodium*, we removed the myxozoans from our three datasets and re-ran each analysis. Under the ML analysis of 18S rDNA, the removal of myxozoans results in the placement of *Polypodium* nested within Cnidaria (Figure 3B and Additional file 3B). This result suggests that the placement of *Polypodium* at the base of bilaterians in the 18S analysis (Figure 3A) was indeed an artifact of LBA. The placement of *Polypodium* within Cnidaria was not effected by the removal of myxozoans in the 28S (Additional file 6) and combined datasets (Additional file 7).

To investigate the possible role of LBA on myxozoan placement, we removed *Polypodium* from the combined ML analyses and found that it did not affect the position of Myxozoa at the base of the Bilateria (not shown). Given that bilaterians also form long branches, we tried removing all bilaterian sequences in the combined ML analysis. This resulted in a Myxozoa + *Polypodium* clade nested within Cnidaria (not shown). However, when *Polypodium*

and bilaterians were removed, myxozoans fell outside the cnidarians (not shown). Similar effects of myxozoan placement to long-branches were also found in parsimony analyses of the combined dataset (not shown).

Discussion

Polypodium is a cnidarian

Our metazoan dataset of 18S and partial 28S rDNA sequences, with a large taxonomic sample of cnidarians, places Polypodium within a monophyletic Cnidaria. This accords with the fact that Polypodium possesses nematocysts [17,18] and a cnidarian-like body plan [7-9,12]. The precise placement of Polypodium within Cnidaria is less certain. The ML combined analysis places *Polypodium* as sister to Hydrozoa (Figure 2A), a hypothesis consistent with the suggestion that Polypodium be considered a separate class of cnidarians, Polypodiozoa [1]. By contrast, the combined parsimony analysis (Figure 2B) and the ML analyses of 28S alone (Additional file 2 and 6) place Polypodium within the hydrozoan clade Leptothecata. Given that leptothecates have relatively high rates of evolution within hydrozoans, one possible explanation for the conflicting hypotheses is that the placement of Polypodium within leptothecates is an artifact of LBA and that the combined data, in conjunction with the ML approach (Figure 2A), overcame this localized LBA artifact.



Figure 2
Phylogenetic hypotheses of relationships among 126 metazoan taxa, based on a combined analysis of nearly complete 18S and partial 28S rDNA sequences. Arrow indicates *Polypodium* taxa. A) Maximum likelihood topology. The assumed model (GTR+I+G) has six substitutions rates estimated from the data (A-C, 1.1786; A-G, 3.3654; A-T, 1.7283; C-G, 0.7403; C-T, 4.7803; G-T, 1.0000), an assumed proportion of invariant sites (0.1692) and a gamma shaped parameter or (0.5584). The length of the bar indicates 0.1 substitutions per site. Bootstrap values for this topology are indicated on the cladogram in Additional file 1. B) Strict consensus of 32 trees of length 25141 from a parsimony analyses. Bootstrap values of 50 or greater are indicated.

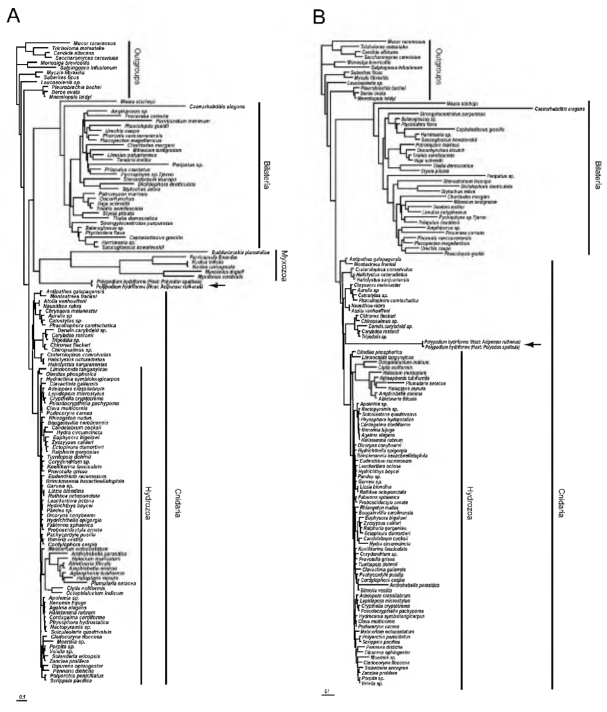


Figure 3 ML topologies of metazoan relationships of nearly complete 18S rDNA sequences. Arrow indicates *Polypodium* taxa. Bootstrap values for both topologies are indicated on the cladograms in Additional file 3. A) 132 taxa including 6 myxozoan taxa and two *Polypodium* taxa. The assumed model (GTR+I + G) has six substitutions rates estimated from the data (A-C, 1.4071; A-G, 3.3470; A-T, 1.6901; C-G, 0.84888; C-T, 4.7638; G-T, 1.0000), an assumed proportion of invariant sites (0.1757) and a gamma shaped parameter or (0.5837). B) Same dataset as (A) but with the 6 myxozoan taxa removed. The assumed model (GTR+I + G) has six substitutions rates estimated from the data (A-C, 1.4115; A-G, 3.3559; A-T, 1.7502; C-G, 0.8342; C-T, 4.8554; G-T, 1.0000), an assumed proportion of invariant sites (0.2464) and a gamma shaped parameter or (0.6326). The length of the bar indicates 0.1 substitutions per site.

Evolution of Polypodium life-history characters

Although the fresh water habitat of *Polypodium* is unusual for cnidarians, it is not unheard of, especially within hydrozoans. For instance, the model organism *Hydra* and the jellyfish *Craspedacusta* are both exclusively fresh-water hydrozoans. *Hydra* and *Craspedacusta* are distantly related [25] and our analyses do not indicate a close phylogenetic affinity of *Polypodium* to either of the clades containing these taxa. Thus, it appears that in the evolution of cnidarians, invasion to fresh-water habitats has happened at least three separate times.

Although *Polypodium* is the only known intracellular cnidarian parasite, other cnidarians have adopted parasitic life-styles [11,26-29]. For example, parasites belonging to the Narcomedusae (Hydrozoa) have been reported to live in the stomach cavities of other narcomedusae [11,27] and anthomedusae [27]. In addition, the sea anemone *Edwardsiella lineata* parasitizes the stomach cavity of the ctenophore *Mnemiopsis leidyi* [28] and the anemone *Peachia quinquecapitata* is reported to parasitize the stomachs of hydromedusa [29].

Effects of long-branch attraction

The well-documented effects of long-branch attraction artifacts (reviewed in Bergsten [30]) are particularly concerning when investigating relationships amongst earlydiverging metazoans, where rates between lineages vary greatly [22]. Suggestions for avoiding LBA artifacts include choice of appropriate markers [31,32], increased taxonomic sampling to effectively break up long branches [33,34] and utilization of best-fit models that incorporate rate variation [21-23]. Previous conflicting reports that show Polypodium and myxozoans form a sister taxon to Bilateria [2-4] can be explained by limited taxon sampling and an inadequate number of informative characters in their analyses, both of which confound long-branch problems. In this study, the increased taxonomic sampling of cnidarians and the addition of 28S rDNA sequence data proved critical to placing the highly divergent Polypodium taxon within Cnidaria. The choice of optimality criteria (ML vs. parsimony) both supported Polypodium as a cnidarian but did affect the placement within Cnidaria.

Polypodium and Myxozoa

Our analyses are inconclusive in the placement of Myxozoa within metazoans. We found that myxozoans consistently grouped with long-branched taxa and that removal of long-branches resulted in myxozoans being placed to the next longest branch. For example myxozoans group with *Polypodium* in the absence of Bilateria and group with Bilateria in the absence of *Polypodium* (not shown).

Jimenez-Guri et al. [24] sampled the myxozoan, Budden-brockia, and found it to fall within Cnidaria, as the sister

group to two hydrozoan representatives and a single scyphozoan. Previous studies have suggested a sister group relationship between cnidarians and myxozoans [2-4], and some morphological evidence has been used to support this view [35]. Although our present study does not support this relationship, further investigation is merited. Myxozoans are a highly diverse group (reviewed in Kent et al. [36]) that comprise two clades, the Myxosporea and the Malacosporea [37]. We were only able to include 28S rDNA sequences from myxosporeans, although the malacosporean Buddenbrockia was included in our 18S analysis and found to group with other myxozoans and outside of Cnidaria. Future studies with a comprehensive sampling of myxozoans together with Polypodium, in a dataset that includes a large taxonomic sampling of cnidarians, should shed further light on the relationships between myxozoans and Polypodium.

Conclusion

Although previous molecular phylogenetic hypotheses conflicted with the traditional interpretation of cnidarian affinity for *Polypodium*, the molecular evidence we present, using an augmented dataset, ultimately confirms and reconciles this traditional hypothesis and suggests that *Polypodium* is indeed a cnidarian. This study also reaffirms the importance to large taxonomic sampling and inclusion of additional informative characters for avoiding long-branch attraction artifacts.

Methods

DNA isolation, amplification and sequencing

Genomic DNA was extracted using Qiagen DNeasy kits according to manufacturer's protocol (QIAGEN Inc., Mississauga, ON) or a standard phenol/chloroform protocol. The latter method involved tissue digestion with proteinase K (20 mg/ml) in a lysis buffer (20 mM Tris-CL pH 8.0, 5 mM EDTA pH 8.0, 400 mM NaCl, 2%SDS), extraction with phenol/chloroform (1:1), precipitation with 2.5 vol. 95% EtOH, and elution in TE or $\rm H_2O$.

An approximately 1.8 kb portion of the gene coding for 18S was amplified and sequenced with universal eukaryotic primers as described by Medlin et al. [38], with the annealing temperature modified to 57°C. With the exception of Polypodium samples, a nearly complete, roughly 3 kb portion of the gene coding for 28S was amplified and sequenced with an approach modified from that reported in Collins et al. [25]. 28S was directly amplified in two combinations fragments with of primers F63mod+R2077sq and F1379+R3264 from Medina et al. [39] or newly developed medusozoan specific primers F97+R2084 and F1383+R3238 (F97: CCYYAGTAACG-GCGAGT, R2084: AGAGCCAATCCTTTTCC, F1383: GGACGGTGGCCATGGAAGT, and R3238: SWACAGAT-GGTAGCTTCG). Amplifications of 28S were conducted with the following thermal profile: 4 minutes at 94°C; 30 cycles of 30 seconds at 94°C, 1 minute at 45°C, and 3 minutes at 72°C; and 10 minutes at 72°C. For *Polypodium*, a portion of the 5' end of 28S (approx. 0.8–1.0 kbps) was amplified using two universal metazoan primers (fw1and rev2) as reported by Sonnenberg et al. [40]. Sequencing was carried out using amplification primers and F635sq and R635sq from Medina et al. [39].

All gene fragments were purified and sequenced by Cogenics, Inc. (Houston, TX) and assembled and edited using Sequencher v4.5 (Gene Code Co., 2005). Sequences for each marker were aligned using the program MUSCLE [41]. The 28S sequence alignment was then trimmed to reflect only that region which included sequence data for *Polypodium*. This trimmed 28S dataset was analyzed separately and used in conjunction with the complete 18S sequences to create the combined dataset.

Phylogenetic analyses

Phylogenetic analyses were performed using both maximum likelihood (ML) and parsimony criteria. ML searches were performed using GARLI v0.951.OsX-GUI [42] under an assumed GTR model with rates estimated from the data. The assumed model of nucleotide substitution was selected by using the Akaike Information Criterion (AIC) as implemented in ModelTest [43]. Each run was repeated 10 times from random starting trees using default termination conditions. Each run gave identical topologies and similar likelihood scores. 100 bootstrap replications were run in GARLI v0.951.0sX-GUI [42] under the same parameters.

To assess the effect that omitting length-variable regions has on topology, we removed these regions from the combined dataset, using the less stringent settings of Gblocks [44]. This dataset contained 126 metazoan taxa, 2415 characters, 1391 of which are parsimony informative. We found that removal of length-variable regions had no effect on the placement of *Polypodium* and minimal effect on overall topology in our combined ML analyses (Additional file 8). Therefore we performed all other analyses with the complete datasets, including the more variable regions.

Parsimony analyses were performed using TNTv.1.1 [45]. Separate tree searches were performed with gaps coded as missing and gaps coded as a fifth state. However, with one exception (see results for myxozoan placement with 18S data) there was no significant difference in topology. Numerous search methods available in TNT were utilized to search the tree space but the following approach was found to consistently recover trees with minimum lengths

from our datasets. The implemented search was a driven new technology search with a random seed of 0 (where 0 = time). Default settings for sectorial searches (RSS and CSS) and tree fusing were used [46], with 5 replicates per repetition, and a requirement that the global optimum be found 20 times. TBR branch swapping was performed on the resulting trees and a strict consensus was calculated. TNT was used to calculate standard bootstrap values (1000 replicates). Alignments and trees for 18S, 28S and combined datasets have been submitted to TreeBASE http://www.treebase.org/treebase/index.html.

Authors' contributions

This study was inspired by the work of EVR and originally conceived by EVR, AL and AGC. NME performed most of the data collection and submission of the new sequences. NME and PC performed the analyses. PC took the lead in organizing the study and drafting the manuscript with contributions from NME. All other authors provided comments and suggestions and approved the final manuscript.

Additional material

Additional file 1

ML topology of relationships based on combined data. ML topology identical to Figure 1A but as a cladogram showing bootstrap values. Click here for file

[http://www.biomedcentral.com/content/supplementary/1471-2148-8-139-S1.pdf]

Additional file 2

ML topology of relationships based on partial 28S rDNA sequences. Relationships of 128 metazoan taxa were analyzed with partial 28S rDNA sequences.

Click here for file

[http://www.biomedcentral.com/content/supplementary/1471-2148-8-139-S2.pdf]

Additional file 3

ML topologies of relationships based on 18S data with and without myxozoans. ML topologies identical to Figure 3 but as cladograms showing bootstrap values.

Click here for file

[http://www.biomedcentral.com/content/supplementary/1471-2148-8-139-S3.pdf]

Additional file 4

Parsimony topology of relationships based on 18S rDNA sequences. This parsimony analysis of 18S rDNA sequences included 132 taxa with gaps coded as a fifth state.

Click here for file

[http://www.biomedcentral.com/content/supplementary/1471-2148-8-139-S4.pdf]

Additional file 5

Parsimony topology of relationships based on 18S rDNA sequences. This parsimony analysis of 18S rDNA sequences included 132 taxa with gaps coded as missing.

Click here for file

[http://www.biomedcentral.com/content/supplementary/1471-2148-8-139-S5.pdf]

Additional file 6

ML topology of relationships excluding myxozoans, based on partial 28S rDNA data. This ML analysis of partial 28S rDNA sequences excluded myxozoan taxa.

Click here for file

[http://www.biomedcentral.com/content/supplementary/1471-2148-8-139-S6.pdf]

Additional file 7

ML topology of relationships excluding myxozoans, based on combined data. This ML analysis of partial 28S rDNA and 18S sequences excluded myxozoan taxa.

Click here for file

[http://www.biomedcentral.com/content/supplementary/1471-2148-8-139-S7.pdf]

Additional file 8

ML topology of relationships based on combined data. This analysis of 126 metazoan taxa was based on combined 18S and partial 28S rDNA with length variable sequences removed.

Click here for file

[http://www.biomedcentral.com/content/supplementary/1471-2148-8-139-S8.pdf]

Acknowledgements

We thank S. Ash Bullard, Jan Dean, Bobby Reed and Ron Bruch for contributions of *Polypodium* specimens. We also thank Peter Schuchert, Casey Dunn and Michael Dawson for contributions of other cnidarian specimens. NME acknowledges the instructors at the 2007 Ohio State Cladistics workshop, especially P. Goloboff, for help with TNT. We thank M. Holder for discussions, B. Bentlage, B. Lieberman, A. Nawrocki, and five anonymous reviewers for helpful comments on earlier versions of this manuscript. This work was supported by grants from NSF ATOL EF-0531779 (to PC and AGC) and support for AL from NSF (PEET DEB-9978086) and FAPESP (06/02960-8/05821-9/60327-0).

References

- Raikova EV: On the systematic position of Polypodium hydriforme Ussov, (Coelenterata). In Porifera and Cnidaria Contemporary state and perspectives of investigations Edited by: Koltum VM, Stepanjants SD. Leningrad: Zoological Institute of Academy of Sciences of USSR; 1988:116-122.
- Siddall ME, Martin DS, Bridge D, Cone DM, Desser SS: The demise
 of a phylum of protests: Phylogeny of Myxozoa and other
 parasitic Cnidaria. J Parasitol 1995, 81:961-967.
- Siddall ME, Whiting MF: Long-branch abstractions. Cladistics 1999, 15:9-24.
- Zrzavý J, Hypša V: Myxozoa, Polypadium, and the origin of the Bilateria: The phylogenetic position of "Endocnidozoa" in the light of the rediscovery of Buddenbrockio. Cladistics 2003, 19(3):164-169.
- 5. Hanelt B, Van Schyndel D, Adema CM, Lewis L, Loker ES: The phylogenetic position of *Rhopalura ophiocomae* (Orthonectida)

- based on 18S ribosomal DNA sequence analysis. Mol Biol Evol 1996, 13:1187-1191.
- Raikova EV: Life cycle and systematic position of Polypodium hydriforme Ussov (Coelenterata), a cnidarian parasite of the eggs of Acipenseridae. Publ Seto Mar Biol Lab 1973, 20:165-173.
- Raikova EV: Morphology, ultrastructure and development of the parasitic larva and its surrounding trophamnion of Polypodium hydriforme Ussov (Coelenterata). Cell Tissue Res 1980, 206(3):487-500.
- Raikova EV: Life cycle, cytology, and morphology of Polypodium hydriforme, a coelenterate parasite of the eggs of acipenseriform fishes. J Parasitol 1994, 80(1):1-22.
- Raikova EV, Suppes VC, Hoffmann GL: The parasitic coelenterate, Polypodium hydriforme Ussov, from the eggs of the American acipenseriform Polyadon spathula. J Parasitol 1979, 65(5):804-810.
- Berrill NJ: Development and medusa-bud formation in the Hydromedusae. Quart Rev Biol 1950, 25:292-316.
- Bouillon J: Considérations sur les dévéloppement des narcoméduses et sur leur position phylogenétique. Indo-Malayan Zool 1987, 4:189-278.
- 12. Hyman L: **The invertebrates**. *I. Protozoa through Ctenophara*. New York and London: McGraw-Hill. 1940
- Schimkevitch VM: Essay on contemporary state of the problem of the development of Hydrozoa. Vestnik Estestvoznanija 1890, 4:171-176.
- Lipin A: Geschlechtliche Form, Phylogenie und systematische Stellung von Polypodium hydriforme Ussov. Zool Jahrb Anat 1925, 47:541-635.
- Bouillon J, Medel MD, Pagès F, Gili JM, Boero F, Gravili C: Fauna of the Mediterranean Hydrozoa. Scientia Marina 2004, 68(Suppl 2):5-438.
- Bouillon J, Gravili c, Pagès F, Gili JM, Boero F: An introduction to Hydrozoa. Paris: Publications Scientifiques du Muséum, Paris; 2006.
- Raikova EV: Fine structure of the nematocytes of Polypodium hydriforme Ussov (Cnidaria). Zoologica Scripta 1990, 19(1):1-11.
- Ibragimov A, Raikova E: Nematocysts of Polypodium hydriforme, a cnidarian parasite of acipenseriform fishes. Hydrobiologia 2004, 531:165-171.
- Dellaporta SL, Xu A, Sagasser S, Jakob W, Moreno MA, Buss LW, Schierwater B: Mitochondrial genome of Trichoplax adhaerens supports Placozoa as the basal lower metazoan phylum. Proc Natl Acad Sci USA 2006, 103(23):8751-8756.
- Collins AG: Phylogeny of Medusozoa and the evolution of cnidarian life cycles. | Evol Biol 2002, 15(3):418-432.
- Huelsenbeck JP: Is the Felsenstein zone a fly trap? Syst 8iol 1997, 46:247-264.
- Kim JH, Kim W, Cunningham CW: A new perspective on lower metazoan relationships from 18S rDNA sequences. Mol Biol Evol 1999, 16:423-427.
- Cunningham CW, Zhu H, Hillis DM: Best-fit maximum likelihood models for phylogenetic inference: Empirical tests with known phylogenies. Evolution 1998, 52:978-987.
- 24. Jimenez-Guri E, Philippe H, Okamura B, Holland PWH: **Buddenbrockio** is a cnidarian worm. Science 2007, 317(5834):116-118.
- Collins AG, Schuchert P, Marques AC, Jankowski T, Medina M, Schierwater B: Medusozoan phylogeny and character evolution clarified by new large and small subunit rDNA data and an assessment of the utility of phylogenetic mixture models. Syst Biol 2006, 55(1):97-115.
- Ósborn DA: Cnidárian "parasites" on Solmissus incisa, a Narcomedusa. Sci Mar 2000, 64:157-163.
- Pagès F, Corbera J, Lindsay D: Piggybacking pycongonids and parasitic narcomedusae on Pandea rubra (Anthomedusae, Pandeidae). Plankton 8enthos Res 2007, 2(2):83-90.
- 28. Bumann D, Puls G: Infestation with larvae of the sea anemone Edwardsia lineata affects nutrition and growth of the ctenophore Mnemiopsis leidyi. Parasitology 1996, 113:123-128.
- Spaulding JG: The life cycle of Peachia quinquecapitata, an anemone parasitic on medusae during its larval development. 8iol Bull 1972, 143(2):440-453.
- Bergsten J: A review of long-branch attraction. Cladistics 2005, 21:163-193.
- Rokas A, King N, Finnerty J, Carroll SB: Conflicting phylogenetic signals at the base of the metazoan tree. Evol Dev 2003, 5(4):346-359.

- 32. Xiang QY, Moody ML, Soltis DE, Fan C, Soltis PS: Relationships within Cornales and circumscription of Cornaceae matK and rbcL sequence data and effects of outgroups and long branches. Mol Phyl Evol 2002, 24:3S-S7.
- Hillis DM: Taxonomic sampling, phylogenetic accuracy, and investigator bias. Syst 8iol 1998, 47:3-8.
- Zwickl DJ, Hillis DM: Increased taxon sampling greatly reduces phylogenetic error. Syst Biol 2002, 51:588-598.
- Raikova EV: Cytomorphological characters of Polypodium hydriforme and problems of myxozoan and cnidarian phylogeny. Tsitologiio 200S, 47(10):933-939.
- Kent ML, Andree KB, Bartholomew JL, El-Matbouli M, Desser SS, Devlin RH, Feist SW, Hedrick RP, Hoffmann RW, Khattra J, Hallett SL, Lester RJG, Longshaw M, Oswaldo P, Siddall ME, Xiao C: Recent advances in our knowledge of the Myxozoa. J Eukoryot Microbiol 2001, 48(4):395-413.
- Canning EÜ, Curry A, Feist SW, Longshaw M, Okamura B: A new class and order of myxozoans to accommodate parasites of bryozoans with ultrastructural observations on Tetrocopsulo bryosolmonoe (PKX Organism). J Eukoryot Microbiol 2000, 47(5):4S6-468.
- Medlin LH, Elwood HJ, Stickel S, Sogin ML: The characterization of enzymatically amplified eukaryotic 16S-like rRNA-coding regions. Gene 1988, 71:491-499.
- Medina M, Collins AG, Silberman JD, Sogin ML: Evaluating hypotheses of basal animal phylogeny using complete sequences of large and small subunit rRNA. Proc Natl Acad Sci USA 2001, 98(18):9707-9712.
- Sonnenberg R, Nolte A, Tautz D: An evaluation of LSU rDNA D1-D2 sequences for their use in species identification. Front Zool 2007, 4(1):6.
- 41. Edgar RC: MUSCLE: multiple sequence alignment with high accuracy and high throughput. Nucleic Acids Res 2004, 32(5):1792-1797.
- Zwickl DJ: Genetic algorithm approaches for the phylogenetic analysis of large biological sequence datasets under the maximum likelihood criterion. Austin: The University of Texas; 2006.
- Posada D, Crandall KA: Modeltest: testing the model of DNA substitution. 8ioinformatics 2000, 14:817-818.
- 44. Castresana J: Selection of conserved blocks from multiple alignments for their use in phylogenetic analysis. Mol Biol Evol 2000, 17:S40-SS2.
- Goloboff P, Farris J, Nixon K: TNT: Tree Analysis Using New Technology. 1.1th edition. Tucuman, Argentina: Published by the authors; 2003.
- Goloboff P: Analyzing large data sets in reasonable times: Solutions for composite optima. Clodistics 1999, 14:415-428.

Publish with **Bio Med Central** and every scientist can read your work free of charge

"BioMed Central will be the most significant development for disseminating the results of biomedical research in our lifetime."

Sir Paul Nurse, Cancer Research UK

Your research papers will be:

- · available free of charge to the entire biomedical community
- peer reviewed and published immediately upon acceptance
- cited in PubMed and archived on PubMed Central
- ullet yours you keep the copyright

Submit your manuscript here: http://www.biomedcentral.com/info/publishing_adv.asp

