

RESEARCH ARTICLES

Testing the Molecular Clock: Molecular and Paleontological Estimates of Divergence Times in the Echinoidea (Echinodermata)

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The phylogenetic relationships of 46 echinoids, with representatives from 13 of the 14 ordinal-level clades and about 70% of extant families commonly recognized, have been established from 3 genes (3,226 alignable bases) and 119 morphological characters. Morphological and molecular estimates are similar enough to be considered suboptimal estimates of one another, and the combined data provide a tree that, when calibrated against the fossil record, provides paleontological estimates of divergence times and completeness of their fossil record. The order of branching on the cladogram largely agrees with the stratigraphic order of first occurrences and implies that their fossil record is more than 85% complete at family level and at a resolution of 5-Myr time intervals.

Molecular estimates of divergence times derived from applying both molecular clock and relaxed molecular clock models are concordant with estimates based on the fossil record in up to 70% of cases, with most concordant results obtained using Sanderson's semiparametric penalized likelihood method and a logarithmic-penalty function. There are 3 regions of the tree where molecular and fossil estimates of divergence time consistently disagree. Comparison with results obtained when molecular divergence dates are estimated from the combined (morphology + gene) tree suggests that errors in phylogenetic reconstruction explain only one of these. In another region the error most likely lies with the paleontological estimates because taxa in this region are demonstrated to have a very poor fossil record. In the third case, morphological and paleontological evidence is much stronger, and the topology for this part of the molecular tree differs from that derived from the combined data. Here the cause of the mismatch is unclear but could be methodological, arising from marked inequality of molecular rates. Overall, the level of agreement reached between these different data and methodological approaches leads us to believe that careful application of likelihood and Bayesian methods to molecular data provides realistic divergence time estimates in the majority of cases (almost 80% in this specific example), thus providing a remarkably well-calibrated phylogeny of a character-rich clade of ubiquitous marine benthic invertebrates.

Introduction

Congruence between independent sources of data is one of the most persuasive arguments when assessing the reliability of a phylogenetic hypothesis (De Queiroz et al. 1995; Cunnigham 1997). Although some striking differences initially existed between morphological and molecular-based phylogenies, many of these are gradually being resolved as characters are added or reassessed, denser sampling is carried out, new fossils are found, and better methods of analysis adopted. For example, morphological and molecular data initially pointed to different sister-group relationships for the whales (Cetacea) within mammals. This conflict has now been resolved by the discovery of new fossils (Gingerich 2005), which demonstrated that previous morphological hypotheses were in error. Conversely, Xia et al. (2003) have shown that a bird–mammal linkage in 18S ribosomal RNA (rRNA) (which conflicted with morphological evidence) could be explained by misalignment of sequences, inappropriate generalization of base-frequency parameters over the whole tree, and poor sequence quality. The comparison of morphological and molecular data

sets provides an important cross-check on the reliability of results.

In recent years, attention has shifted to another aspect where concordance between morphological and molecular estimates has been hard to achieve, namely, the dating of divergence times. Here workers seem to be more polarized in their outlook, either critical of the reliability and accuracy of molecular clock methods to date divergence times (e.g., Benton 1999; Rodriguez-Trelles et al. 2002; Benton and Ayala 2003) or dismissive of the quality of the fossil record (e.g., Eastal 1999). Until now, debate has logically focused on areas where the two approaches give the most divergent results, specifically the radiation of the metazoan phyla and on the Mesozoic origins of birds and of mammals (see Smith and Peterson 2002; Aris-Brosou and Yang 2003; Donoghue and Smith 2003; Peterson et al. 2004; Pisani et al. 2004; Blair and Hedges 2005; Ho et al. 2005; Peterson and Butterfield 2005; Welch et al. 2005). In all 3 cases, however, the fossil record is far from satisfactory. The fossil record of birds, for example, is miserable by comparison to almost all other fossil groups, with approximately two-thirds of fossil species from the Mesozoic still represented only by a single specimen (Fountaine et al. 2005), implying that the fossil bird record is still very poorly sampled. Similarly, the pre-Cambrian fossil record of bilaterian metazoans (Eumetazoans) is scant or nonexistent. In order to advance this debate, molecular clock methods need to be more widely tested in groups where the fossil record is considerably better and the phylogeny unequivocally established. Apart from the pioneering work of Pérez-Losada et al. (2004)

¹ A.B.S. led the research for this paper and is responsible for the paleontological and morphological data; D.P. undertook all molecular dating analyses; J.A.M.D., B.S., and B.L.W. carried out the sequencing; and D.T.J.L. was responsible for phylogenetic alignment and analysis.

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on barnacles (Thoracica), no studies have looked at the reliability and consistency of molecular clock methods using marine invertebrate groups with a good fossil record.

Sea urchins (Echinoidea) are a diverse group of marine invertebrate deuterostomes (Schultz 2005; Smith 2006; Smith et al. 2004). Their multielement skeleton is complex, providing a large number of phylogenetically informative characters. This skeleton also preserves well, creating a rich fossil record that has been the focus for much paleontological research. Importantly, echinoid taxonomy, which was largely established in the great monographic works of Mortensen (1928, 1935, 1940, 1943a, 1943b, 1948a, 1948b, 1950, 1951), is based almost exclusively on skeletal characters and so is equally applicable to living and fossil taxa.

Today there are some 900 extant species distributed in about 50 families and 14 orders. Previous phylogenetic studies (e.g., Smith 1988; Smith et al. 1992; Littlewood and Smith 1995; Lessios et al. 1999, 2001; Jeffery et al. 2003; Lee 2003; Stockley et al. 2005) have generally found good levels of congruence between morphological and molecular estimates of relationship. With their good fossil record and well-established phylogenetic relationships, echinoids should provide a model system against which to examine the performance of molecular methods of dating.

Here we compare molecular and paleontological estimates of divergence times for Echinoidea as an empirical cross-check on the reliability of the methods and assumptions. We do this by 1) constructing phylogenies from both molecular data and traditional morphological data to arrive at the best-supported tree, 2) estimating the quality of the echinoid fossil record by calibrating this tree against the observed record of first occurrences, 3) estimating divergence times using a molecular clock method and a variety of relaxed molecular clock models applied to the molecular data, and 4) quantifying the match between the observed times of appearance of clades in the fossil record and the results derived from molecular data.

Materials and Methods

Taxa Included

To construct our phylogeny, we compiled morphological information and gene sequence data for 46 genera (listed in table 1) with representatives from 28 families and 13 of the 14 orders of living echinoid. Where sequence data for multiple species of the same genus existed, a strict consensus sequence was constructed. Because we restricted our analyses to only those regions of the genes that could be unambiguously aligned across all genera, sequences of congeneric species were effectively identical. However, the 2 species of *Araeosoma* were retained separately as these showed modest amounts of sequence divergence.

To compare divergence times based on the fossil record and molecular data, we selected 1 taxon from each family (asterisked in table 1). These were chosen, after examining our initial molecular phylogenetic analysis (fig. 2), to avoid taxa showing anomalously long or short branches, thereby creating a “partially linearized tree” (sensu Takezaki et al. 1995).

Morphological data were rooted by outgroup comparison using the fossil *Archaeocidaris*, a late stem-group echinoid. Data on this taxon were taken from a remarkably

well-preserved Carboniferous species (Lewis and Ensom 1982). Molecular data were rooted using a combination of representatives from each of the other 4 classes of echinoderm as outgroup: the starfish *Asterias*, the ophiuroid *Ophiocanops*, the crinoid *Antedon*, and 2 holothurians, *Cucumaria* and *Psychopetes*. In regions where the ingroup and outgroup sequences were too divergent for meaningful alignment, outgroup sequences were scored as unknown to avoid spurious rooting.

Morphological Characters

Morphological characters were compiled from published analyses, notably from Smith (1988), Littlewood and Smith (1995), Jeffery et al. (2003), and Stockley et al. (2005). In total, 119 characters were scored, 29 of which are multistate (see Supplementary Material online). Two multistate characters (characters 6 and 8 in supplementary table 1, Supplementary Material online) reflect a clear and unambiguous ontogenetic sequence of character states and were, therefore, treated as ordered; the remainder was left unordered. The great majority of characters relate to skeletal features of the adult, which forms the basis for the classification of the group. Only 3 characters relate to the larval skeleton as larval characteristics have been shown to be more homoplastic than characters based on adult skeletal morphology (Smith et al. 1996; Smith and Littlewood 1997). Characters were obtained from direct observation or from modern descriptions in the literature and scored on the basis of the states shown in the species for which sequence data were available. The complete character listing and data matrix are provided as supplementary data (Supplementary Material online).

Molecular Characters

Three rRNA genes, 2 nuclear (18S small subunit [18S] and 28S large subunit [28S]) and 1 mitochondrial (16S large subunit [16S]), were sequenced. These were selected to encompass a range of different evolutionary rates and have been used successfully in a wide range of metazoan phylogeny studies aimed at resolving divergences over the past 250 Myr, including echinoderms (e.g., Littlewood and Smith 1995; Littlewood et al. 1997; Jeffery et al. 2003; Winchell et al. 2004; Stockley et al. 2005). Approximately 630 bp from the 3' end of the 16S gene, 1,250 bp from the 5' end of the 28S gene, and the entire 18S gene (ca. 1,800 bp) were sequenced.

For details of tissue selection and DNA extraction and amplification methods see Littlewood and Smith (1995) and Stockley et al. (2005). Sequence data were obtained for both forward and reverse reads. An initial multiple sequence alignment was made using MacClade (Maddison DR and Maddison WP 2001) followed by alignment of sequences by eye. Areas of high variability, for which no reliable alignment across the different orders could be made, were excluded from further data analysis. In total, out of 3,990 bp, 3,226 were alignable and 449 were phylogenetically informative. All sequences are lodged with GenBank/European Bioinformatics Institute under accession numbers listed in table 1. The aligned data matrix is available as supplementary data (Supplementary Material online).

Table 1
List of Taxa Included in This Study, Their Higher Taxonomic Placement, and GenBank Accession Numbers for Their Gene Sequences

Order	Family	Genus	Species	18S rRNA	28S rRNA	16S rRNA
Cidaroida	Cidaridae	<i>Calocidaris</i>	<i>micans</i> (Mortensen, 1903)	DQ073782	DQ073756	DQ073737
		* <i>Prionocidaris</i>	<i>bispinosa</i> (Lamarck, 1816)	DQ073792	DQ073767	DQ073747
		<i>Stereocidaris</i>	<i>excavatus</i> Mortensen, 1936	DQ073795	DQ073772	DQ073740
Diadematoida	Aspidodiadematidae	* <i>Aspidodiadema</i>	<i>jacobi</i> Agassiz, 1880	DQ073780	DQ073754	DQ073734
	Diadematidae	<i>Diadema</i>	<i>setosum</i> (Leske, 1778)	Z37122	DQ073760	DQ073741
		<i>Centrostephanus</i>	<i>coronatus</i> (Verrill, 1867)	Z37120	—	—
Echinothurioida	Echinothuriidae	* <i>Araeosoma</i>	<i>fenestratum</i> Wyville Thomson, 1872	DQ073777	DQ073752	DQ073732§
		<i>Araeosoma</i>	<i>owstoni</i> Mortensen, 1904	Z37118	Z37507	DQ073735
		* <i>Caenopedina</i>	<i>cubensis</i> Agassiz, 1869	DQ073781	DQ073755	DQ073736
Pedinoida	Pedinidae	* <i>Arbacia</i>	<i>lixula</i> (Linnaeus, 1758)	Z37514	DQ073753	X80396
Arbacioida	Arbaciidae	<i>Arbacia</i>	<i>punctulata</i> (Lamarck, 1816)	DQ073778	AY26367	DQ073733
		<i>Coelopleurus</i>	<i>floridanus</i> Agassiz, 1871	DQ073784	DQ073758	DQ073739
Phymosomatoida	Stomopneustidae	* <i>Stomopneustes</i>	<i>variolaris</i> (Lamarck, 1816)	AF279214, Z37133	DQ073773	AF279169
Temnopleuroida	Temnopleuridae	<i>Temnopleurus</i>	<i>reevesii</i> (Gray, 1855)	AF279200	—	AF279149
		<i>Temnopleurus</i>	<i>hardwickii</i> (Gray, 1855)	Z37135	—	—
		<i>Temnopleurus</i>	<i>toreumaticus</i> (Leske, 1778)	—	DQ073774	AF279164
		<i>Temnopleurus</i>	<i>alexandrei</i> (Bell, 1880)	AF279206	—	AF279156
		<i>Temnotrema</i>	<i>sculptum</i> Agassiz, 1863	AF279201	—	AF279150
		* <i>Salmacis</i>	<i>sphaeroides</i> (Linnaeus, 1758)	AF279210, Z37131	DQ073770	AF279162
		<i>Salmacis</i>	<i>belli</i> Doderlein, 1902	AF279213	—	AF279167
		<i>Salmaciella</i>	<i>oligopora</i> (Clark, 1916)	AF279211	—	AF279163
		<i>Mespilia</i>	<i>globulus</i> (Linnaeus, 1758)	AF279203, Z37130	—	AF279152
		<i>Microcyphus</i>	<i>annulatus</i> Mortensen, 1904	AF279216	—	AF279172
		<i>Microcyphus</i>	<i>olivaceus</i> (Doderlein, 1885)	AF279202	—	AF279151
		<i>Amblypneustes</i>	<i>ovum</i> (Lamarck, 1816)	AF279207	—	AF279157
		<i>Amblypneustes</i>	<i>formosus</i> Valenciennes, 1846	AF279212	—	AF279165
		<i>Holopneustes</i>	<i>porosissimus</i> Agassiz, 1846	AF279208	—	AF179159
		<i>Holopneustes</i>	<i>inflatus</i> Lutken, in Agassiz, 1872	AF279209	—	AF279161
		<i>Pseudechinus</i>	<i>albocinctus</i> (Hutton, 1872)	AF279204	—	AF279153
		<i>Pseudechinus</i>	<i>novaezealandiae</i> Mortensen, 1921	AF279205	—	AF279155
	Trigonocidaridae	* <i>Maculata</i>	<i>maculata</i> Agassiz, 1869	AF279199	—	AF279148
Echinoida	Strongylocentrotidae	* <i>Strongylocentrotus</i>	<i>purpuratus</i> (Stimpson, 1857)	L28056	AF212171	—
		<i>Strongylocentrotus</i>	<i>intermedius</i> (Agassiz, 1863)	—	—	AB154276
	Echinometridae	* <i>Anthocidaris</i>	<i>crassispinus</i> (Agassiz, 1863)	DQ073776	DQ073751	AB154278
	Echinidae	<i>Psammechinus</i>	<i>miliaris</i> (Muller, 1771)	AF279215, Z37149	DQ073768	AF279170
		* <i>Paracentrotus</i>	<i>lividus</i> (Lamarck, 1816)	AY428816	DQ073766	J04815
	Toxopneustidae	<i>Tripneustes</i>	<i>gratilla</i> (Linnaeus, 1758)	Z37134	—	AB154279
		* <i>Cyrtechinus</i>	<i>verruculatus</i> (Lutken, 1864)	DQ073786	DQ073759	DQ073740
		<i>Sphaerechinus</i>	<i>granulosus</i> (Lamarck, 1816)	Z37132	DQ073771	DQ073749
		<i>Lytechinus</i>	<i>variegatus</i> (Lamarck, 1816)	DQ073790	AJ225816	DQ073746
Echinoneoida	Echinoneidae	* <i>Echinoneus</i>	<i>cyclostomus</i> Leske, 1778	DQ073789	AJ639778	AJ639801
Cassiduloida	Cassidulidae	* <i>Cassidulus</i>	<i>mitis</i> Krau, 1954	Z37148	—	—
	Echinolampadidae	* <i>Echinolampas</i>	<i>crassus</i> (Bell, 1880)	DQ073788	DQ073764	DQ073744
		<i>Conolampas</i>	<i>sigsbei</i> (Agassiz, 1878)	DQ073785	AJ639777	AJ639800
Clypeasteroida	Arachnoididae	* <i>Fellaster</i>	<i>zealandiae</i> (Gray, 1855)	Z37128	DQ073765	DQ073745
	Echinocyamiidae	* <i>Echinocyamus</i>	<i>pusillus</i> (Muller, 1776)	DQ073787	DQ073762	DQ073743
	Laganidae	* <i>Rumphia</i>	<i>orbicularis</i> (Leske, 1778)	DQ073793	DQ073769	DQ073748
	Astriclypeidae	* <i>Echinodiscus</i>	<i>bisperforatus</i> Leske, 1778	Z37124	DQ073763	—
	Mellitidae	* <i>Encope</i>	<i>abberans</i> Martens, 1867	Z37126	Z37117	—
Holasteroida	Plexechinidae	* <i>Plexechinus</i>	<i>planus</i> Mironov, 1978	AY957468	AY957469	AY957467
Spatangoida	Schizasteridae	* <i>Abatus</i>	<i>cavernosus</i> Philippi, 1845	DQ073775	AJ639776	AJ639803
	Paleopneustidae	* <i>Paleopneustes</i>	<i>cristatus</i> Agassiz, 1873	DQ073791	AJ639784	AJ639808
	Archaeopneustidae	* <i>Archaeopneustes</i>	<i>hystrix</i> Agassiz, 1880	DQ073779	AJ639785	AJ639809
	Brissidae	<i>Brissopsis</i>	<i>atlantica</i> Mortensen, 1907	—	AJ639794	AJ639818
		<i>Brissopsis</i>	<i>lyrifera</i> (Forbes, 1841)	Z37119	—	—
		* <i>Meoma</i>	<i>ventricosa</i> Lamarck, 1816	Z37129	AJ639796	AJ639820
		<i>Spatangus</i>	<i>multispinus</i> Mortensen, 1925	—	AJ639786	AJ639810
	Spatangidae	* <i>Spatangus</i>	<i>raschi</i> Loven, 1869	DQ073794	AJ639787	AJ639811
	Loveniidae	* <i>Echinocardium</i>	<i>laevigaster</i> Agassiz, 1869	—	AJ639789	AJ639813
		<i>Echinocardium</i>	<i>cordatum</i> (Pennant, 1777)	Z37123	DQ073761	DQ073742

NOTE.—Taxa with asterisks were used as representatives of their family in molecular analyses.

Analytical Methods

Phylogenetic analysis of the morphological data was carried out with maximum parsimony as the optimality criterion, using the Macintosh version of PAUP* (4.0b10

[Altevec] (Swofford 2002). Because of the large number of taxa included, we used a heuristic search method, with 1,000 random additional replicates and tree bisection reconnection branch swapping. Node support was tested

by bootstrapping with 250 replicates and by clade decay analysis (Bremer 1994).

All 3 genes were combined for phylogenetic and molecular clock analyses. The program Modeltest (version 3.06) (Posada and Crandall 1998) was used to analyze each data set and produce an appropriate nucleotide substitution model. We used the general time reversible (GTR) + Γ + I model (rates set to gamma, with 6 rate categories). Bayesian inference analyses were conducted using a separate GTR + Γ + I model for each data partition independently and also for the combined 3-gene analysis, thus allowing separate estimates for each model parameter per data set. Bayesian analyses were performed using MrBayes (Huelsenbeck and Ronquist 2003). The number of generations permitted was 5,000,000 with 4 chains, and the 50% majority rule consensus tree was constructed from the non-burn-in trees. Maximum likelihood (ML) analyses were implemented using subtree pruning regrafting (Hordijk and Gascuel 2005) under the best-fitting model (GTR + Γ + I). Support for the nodes in the ML tree was estimated using the bootstrap (100 replicates).

Combined morphological and molecular data were analyzed using both parsimony and Bayesian methods. The likelihood model developed for morphological data by Lewis (2001) was used for the morphological partition, and separate, unlinked GTR + Γ + I models were used for the molecular partitions.

Data Congruence Tests

The appropriateness of combining the morphological and molecular data sets was tested using the partition homogeneity test (Farris et al. 1994), as implemented in PAUP*. A Templeton (1983) test of data heterogeneity was performed on the trees that were obtained from morphological and molecular data, to determine whether they could be considered suboptimal estimates of the same underlying topology. We also used the recommended approach of Wiens (1998) of comparing support levels for nodes in the morphological and molecular analyses. We identified problematic areas as nodes where morphological and molecular data pointed to incongruent groupings with strong support, as indicated by bootstrap proportions of >70% or posterior probabilities of >95%.

Stratigraphic Congruence Tests

Benton's (1995, 2001) Relative Completeness Index (RCI) was used to measure the fit of stratigraphic data to cladogram topology. This measures the amount of missing range that must be added to make stratigraphic record fit the phylogeny and has the advantage over Huelsenbeck's (1994) Stratigraphic Consistency Index or the Spearman rank correlation proposed by Norrell and Novacek (1992) in that it takes account of the relative size of the mismatch over the entire tree (Hitchin and Benton 1997). We calculated RCI for the trees derived from the combined morphological and molecular data (fig. 3), using both parsimony and Bayesian analysis. We divided the fossil record into 5-Myr intervals and, for each sister-group pairing, used the earliest occurrence of either to establish the minimum

time of origin of both sister groups. The geological ages of the earliest fossil representative of each clade at the family level or above included in this analysis are listed in table 2. Supporting evidence for the paleontological dating of nodes is provided in Appendix.

Where a family is the only representative of a larger clade, the oldest member of the more inclusive clade is given. For example, the Cidaridae is the only included family-level representative of the subclass Cidaroidea, sister group to all other living echinoids (fig. 1). Whereas the fossil record of Cidaridae is not much older than 150 Myr, stem-group members of the Cidaroidea extend back to 255 Myr. In such cases, we always use the earliest representative of a member of the most inclusive taxonomic group to date the divergence.

Molecular Estimates of Divergence Times

All molecular estimates of divergence times were calculated for the 28 family representatives selected (asterisked in table 1) plus outgroups, as explained in Materials and Methods. The likelihood ratio test (LRT) was used to test the null hypothesis that the data evolved under a molecular clock. This was done using Modeltest in "LRT calculator mode," after having estimated the likelihood for the molecular tree (fig. 2) under the GTR + Γ + I model, both imposing a molecular clock and after having removed this assumption. As the molecular clock assumption could be rejected ($P < 0.00001$ with 29 degrees of freedom), divergence time estimates were calculated from the combined 16S, 18S, and 28S rRNA sequences using a variety of relaxed molecular clock approaches. We used the fully parametric Bayesian method of Thorne et al. (1998), the nonparametric rate smoothing (NPRS), and the semiparametric penalized likelihood (PL) methods of Sanderson (2002). However, we also used a dating method imposing a strict molecular clock, the Langley-Fitch method (LF; Langley and Fitch 1974), to evaluate its performance for comparison. The Bayesian method of Thorne et al. (1998) was implemented using the software Multidivtime (Thorne and Kishino 2002), whereas NPRS, PL, and LF analyses were performed using r8s version 1.70 (Sanderson 2004). The r8s version 1.70 allows divergence times under PL to be estimated using either a logarithmic- or an additive-penalty function, and divergence times for the internal nodes were obtained using both types of penalty function (see Sanderson 2004). Optimal smoothing parameters for the PL and log-PL analyses were obtained by cross-validation (Sanderson 2002, 2004).

Both r8s and Multidivtime need, as input information, a tree topology with associated branch lengths. For the r8s analyses, branch lengths for the ML molecular tree (fig. 2) were calculated via likelihood using PAUP 4b10, under the best-fitting substitution model (GTR + Γ + I), which had been reestimated using Modeltest after removing all the taxa that were not selected for the molecular clock analyses. For comparison, in the Bayesian analyses branch lengths were estimated with the software Estbranches (which is part of the Multidivtime package) under the F84 + Γ substitution model (as suggested in the Estbranches manual). Software limitations did not allow the implementation of more complex substitution model (i.e., GTR + Γ + I), and the

Table 2
Paleontological Age Estimates Based on Oldest Fossil Occurrence for Each Branch in the Tree Generated from Parsimony Analysis of the Combined Morphological and Molecular Data

Branch	Higher Taxon	First Record	Date (Myr)	Observed Record (Myr)	Missing Record (Myr)
1	Euechinoidea	<i>Serpianotiaris coeava</i> (Quenstedt)	235–240	45	15
2	Acroechinoidea	<i>Diademopsis serialis</i> Agassiz	205–210	0	0
3		<i>Acrosalenia chartroni</i> Lambert	200–205	0	0
4	Diadematoidea	<i>Gymnotiara varusense</i> Cotteau	190–195	10	5
5		<i>Plesiechinus hawkinsi</i> Jesionek Szymanska	195–200	5	5
6	Irregularia	<i>Plesiechinus hawkinsi</i> Jesionek Szymanska	195–200	15	0
7	Microstomata	<i>Galeropygus sublaevis</i> (McCoy)	180–185	0	0
8	Neognathostomata	<i>Galeropygus sublaevis</i> (McCoy)	180–185	80	10
9	Cassiduloida	<i>Hungaresia ovum</i> (Grateloup)	85–90	90	15
10	Clypeasteroidea	<i>Nucleopygus angustatus</i> (Clark)	100–105	50	0
11	Scutellina	<i>Eoscutum doncieuxi</i> (Lambert)	50–55	0	0
12	Laganiformes	<i>Sismondia logotheti</i> Fraas	50–55	0	0
13	Scutelliformes	<i>Eoscutum doncieuxi</i> (Lambert)	50–55	25	0
14	Atelostomata	<i>Hyboclypus ovalis</i> (Leske)	175–180	25	0
15	Spatangoida	<i>Disaster moeschi</i> Desor	160–165	65	5
16	Paleopneustina	<i>Polydesmaster fourtaui</i> Lambert	90–95	0	5
17	Brissidea	<i>Micraster distinctus</i> Agassiz & Desor	95–100	45	0
18		<i>Meoma antiqua</i> Arnold & Clark	40–45	0	0
19		<i>Eupatagus haburiensis</i> Khanna	50–55	15	0
20	Stirodonta + Camarodonta	<i>Atlasaster jeanneti</i> Lambert	195–200	30	0
21		?	—	0	0
22	Camarodonta	<i>Glyptocyphus difficilis</i> (Agassiz)	115–120	0	65
23	Echinoidea	<i>Pseudarbacia archaici</i> (Cotteau)	90–95	65	5
24		?	?	0	0
25		<i>Lytechinus axiologus</i> (Arnold & Clark)	45–50	0	45
26	Cidaroidea	<i>Eotiaris keyserlingi</i> (Geinitz)	250–255	255	
27	Echinothurioida	<i>Pelanechinus oolithicum</i> (Hess)	170–175	175	35
28	Pedinoida	<i>Hemipedina hudsoni</i> Kier	205–210	210	0
29	Aspidodiadematae	<i>Gymnotiara varusense</i> Cotteau	190–195	195	5
30	Diadematae	<i>Farquharsonia crenulata</i> Kier	165–170	170	30
31	Echinoneoidea	<i>Pygopyrina icaunensis</i> Cotteau	160–165	165	35
32	Cassidulidae	<i>Rhyncholampas macari</i> (Smiser)	65–70	70	20
33	Echinolampadidae	<i>Hungaresia ovum</i> (Grateloup)	85–90	90	0
34	Clypeasterina	<i>Clypeaster calzadai</i> Via & Padreny	40–45	45	10
35	Fibularidae	<i>Echinocyamus gurnahensis</i> Roman & Strugo	50–55	55	0
36	Laganidae	<i>Sismondia logotheti</i> Fraas	50–55	55	0
37	Mellitidae	<i>Encope ciae</i> Cortazar	20–25	25	5
38	Astriclypeidae	<i>Amphiope duffi</i> Gregory	25–30	30	0
39	Holasteroidea	<i>Collyrites ellipticus</i> (Lamarck)	165–170	170	0
40	Schizasteridae	<i>Periaster elatus</i> d'Orbigny	90–95	95	0
41	Paleopneustidae	<i>Polydesmaster fourtaui</i> Lambert	90–95	95	0
42	Archaeopneustids	<i>Heterobrissus salvae</i> (Cotteau)	40–45	45	0
43	Brissidae	<i>Meoma antiqua</i> Arnold & Clark	40–45	45	0
44	Spatangidae	<i>Granopatagus lonchophorus</i> Meneghini	35–40	40	0
45	Loveniidae	<i>Hemimareta subrostrata</i> Clark	35–40	40	0
46	Arbacioidea	<i>Atopechinus cellensis</i> Thiery	165–170	170	0
47	Somopneustids	<i>Phymechinus mirabilis</i>	155–160	160	0
48	Temnopleuridae	<i>Zeuglopleurus costulatus</i> Gregory	95–100	100	0
49	Echinidae	<i>Psammechinus dubius</i> (Agassiz)	15–20	20	10
50	Strongylocentrotidae	<i>Strongylocentrotus antiquus</i> Philip	20–25	25	5
51	Echinometridae	<i>Plagiechinus priscus</i> Cotteau	25–30	30	0
52	Toxopneustidae	<i>Lytechinus axiologus</i> (Arnold & Clark)	45–50	50	0
53	Trigonocidaridae	<i>Arbacina monilis</i> (Desmarest)	15–20	20	30
				3210	360

NOTE.—Branches numbered 1–53 in figure 4 are listed in the table along with the higher taxonomic group they represent, the oldest fossil representative that can be assigned to that group, and the paleontological age of that fossil (date) in millions of years. The last 2 columns list, for each branch, the length of time (in millions of years) represented by the observed fossil record and the length of time predicted to exist from the calibrated cladogram but for which no fossil representatives have been found (missing record). The sum of each is given at the foot of each column.

specific parameters for the F84 + Γ model were estimated using phylogenetic analysis by maximum likelihood (PAML) (Yang 1997). Each Bayesian analysis was run 4 times, comparing the divergence times obtained in order to estimate whether convergence was reached. For each

analysis, the Markov Chain Monte Carlo chain was run for 1,000,000 cycles, sampling every 100 cycles. The first 200,000 cycles were considered burn-in.

For all Bayesian divergence time estimates, 95% credibility intervals were calculated. For PL, NPRS, and the LF

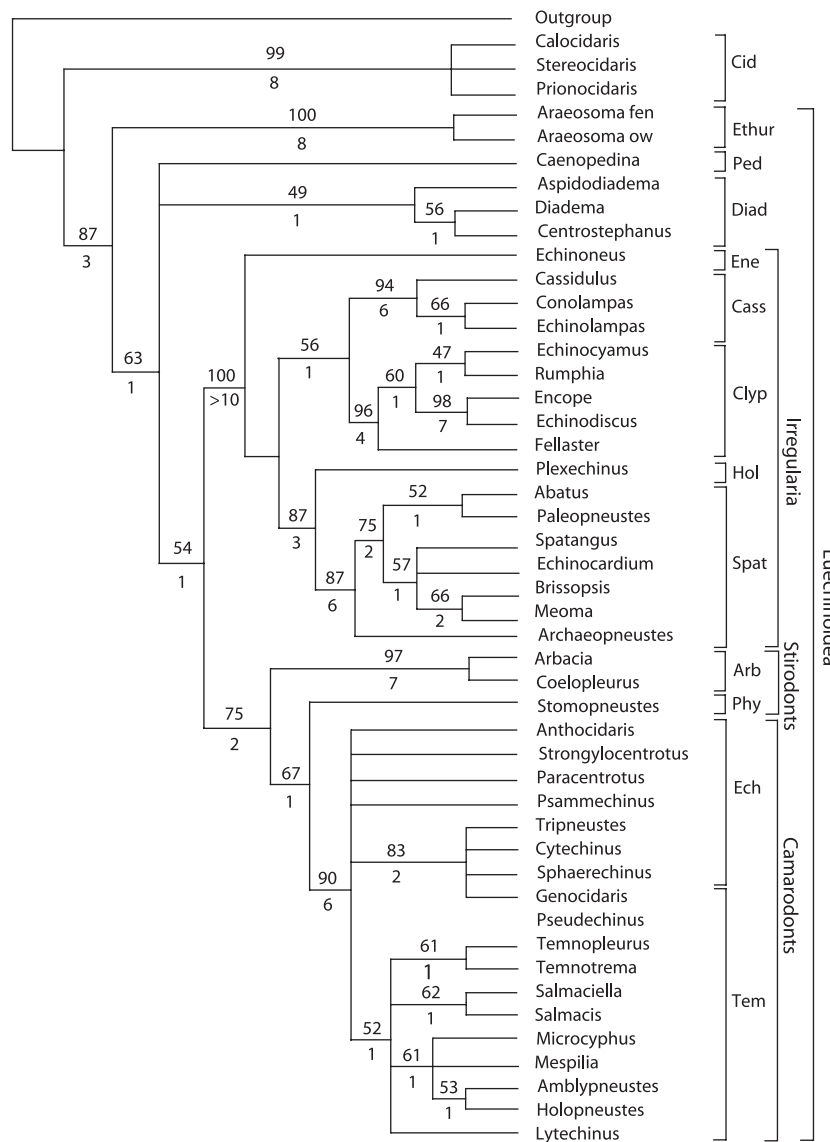


FIG. 1.—Semistrict consensus tree of 4,605 equally parsimonious solutions of TL 249 steps with a CI of 0.60 and retention index of 0.88 derived from the morphological data matrix of 119 characters (see supplementary tables 1 and 3, Supplementary Material online). Bootstrap values for each node are given above each internal branch, and Bremer support values are given beneath. Abbreviations for higher taxonomic groupings as follows: Arb = Arbacioida; Cass = Cassiduloidea; Cid = Cidaroida; Clyp = Clypeasteroidea; Diad = Diadematoidea; Ech = Echinoidea; Ene = Echinoneoidea; Ethur = Echinothurioida; Hol = Holasteroidea; Ped = Pedinoida; Phy = Phymosomatoidea; Spat = Spatangoida; and Tem = Temnopleuroidea.

analyses, confidence intervals around the estimated divergence times were obtained by bootstrapping (Sanderson 2004). Thousand bootstrapped data sets were generated using Seqboot (Felsenstein 2004), and for each bootstrapped data set, the branch lengths of the input tree were reestimated using likelihood (see also above). The results of these analyses were sets of 1,000 bootstrap trees with associated branch lengths, with the trees in each set having the same topology and bootstrapped branch lengths. Divergence times were then obtained, and, taking each of these 1,000 bootstrap trees and each node in the input tree, the standard deviation (SD) of its estimated age was calculated. The 95% confidence intervals around each clade's estimated age were approximated as $(X \pm 2SD)$, where X is the estimated age of a given node. All the computational steps in the estimation of the bootstrap confidence intervals

for the penalized likelihood analyses were automated using several PERL scripts written by D.P.

In the PL, NPRS, and LF analyses the ingroup node was fixed at 265 Myr, which is 10 Myr prior to the first appearance of the oldest recognizable member of the crown group in the fossil record. For the Bayesian analyses of divergence times, we set a date for this node of 265 Myr with a SD of 1, whereas the prior date of the root node was set to 480 Myr, which represents the earliest occurrence of members of 2 of our outgroups, the Asterozoa and Ophiurozoa (Dean 2005). Two sets of analyses were then run, the first without any internal constraints (with the exclusion of the ingroup node) and a second in which 4 local calibration points were enforced. These 4 points were set as minimal divergence estimates, and the basal dichotomy (ingroup node) was taken as a fixed point. The 4 internal calibrations

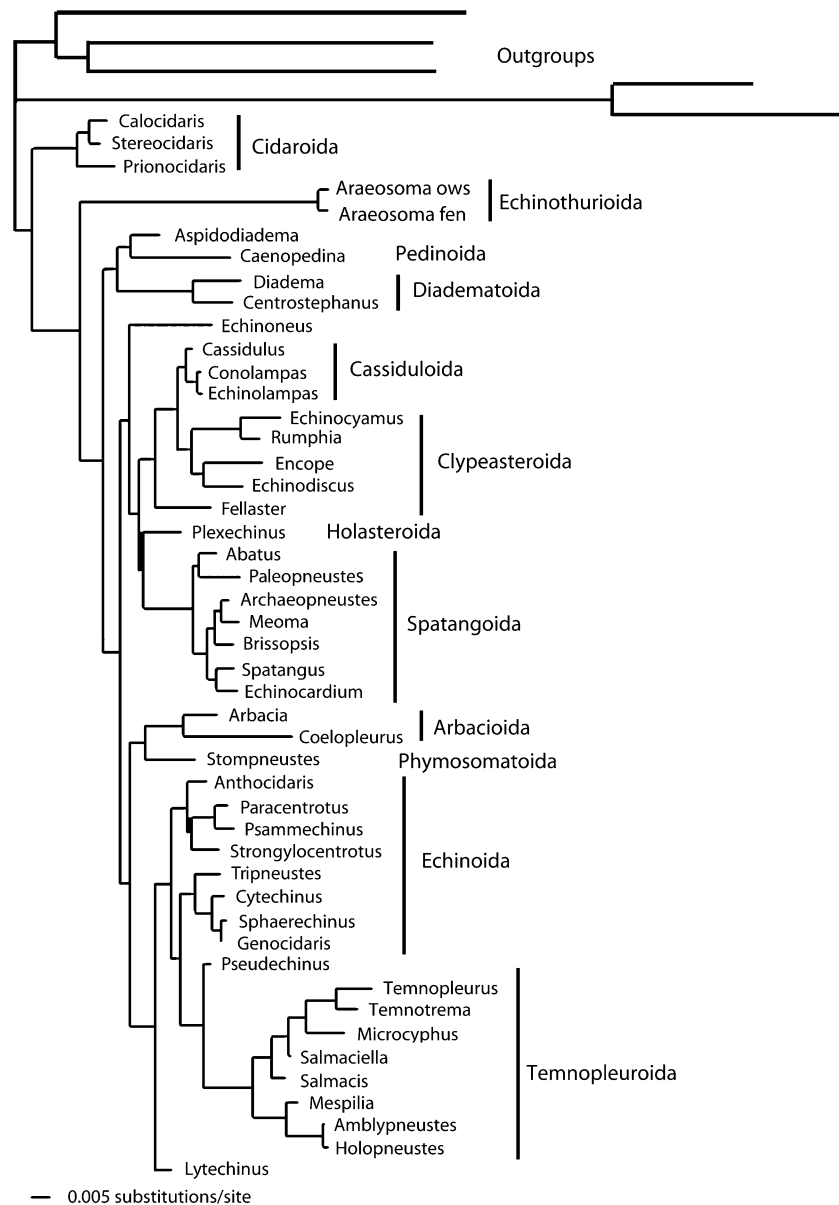


FIG. 2.—Phylogram derived from analysis of the combined molecular sequence data (see supplementary table 3, Supplementary Material online) by ML. Numbers on branches are posterior probabilities.

were selected from across the tree topology to provide constraints on local rate variation. The following local calibration points were employed: the first occurrence of Pedinoida (fig. 5, node 4) at 210 Myr, the first occurrence of Scutellina (fig. 5, node 11) at 55 Myr, the first occurrence of paleopneustid spatangoids (fig. 5, node 15) at 95 Myr, and the first occurrence of Temnopleuridae (fig. 5, node 25) at 45 Myr (all dates from Smith 2006).

Measuring Congruence of Divergence Estimates

For comparing the different methods of estimating divergence times, we use the following measure. Each node where we have an independent molecular and paleontological estimate of divergence time was given a score between 0 and 2: 2 if the fossil date lies within 1 SD of the mean of the molecular estimate, 1 if it lies between 1 and

2 SD of the molecular estimate, and 0 when the paleontological estimate falls outside 2 SDs of the molecular estimate. Our overall measure of congruence is then simply the score summed over all nodes divided by the total number of nodes \times 2; the higher the score (maximum 1 and minimum 0), the better the agreement achieved between molecular and paleontological estimates of divergence on our tree.

Results Phylogenetic Relationships

Parsimony analysis of morphological data found 4,605 equally parsimonious solutions of tree length (TL), 249 steps with a consistency index (CI) of 0.60 and retention index 0.88. A semistrict consensus of these trees (fig. 1) shows good resolution in all parts of the tree except among

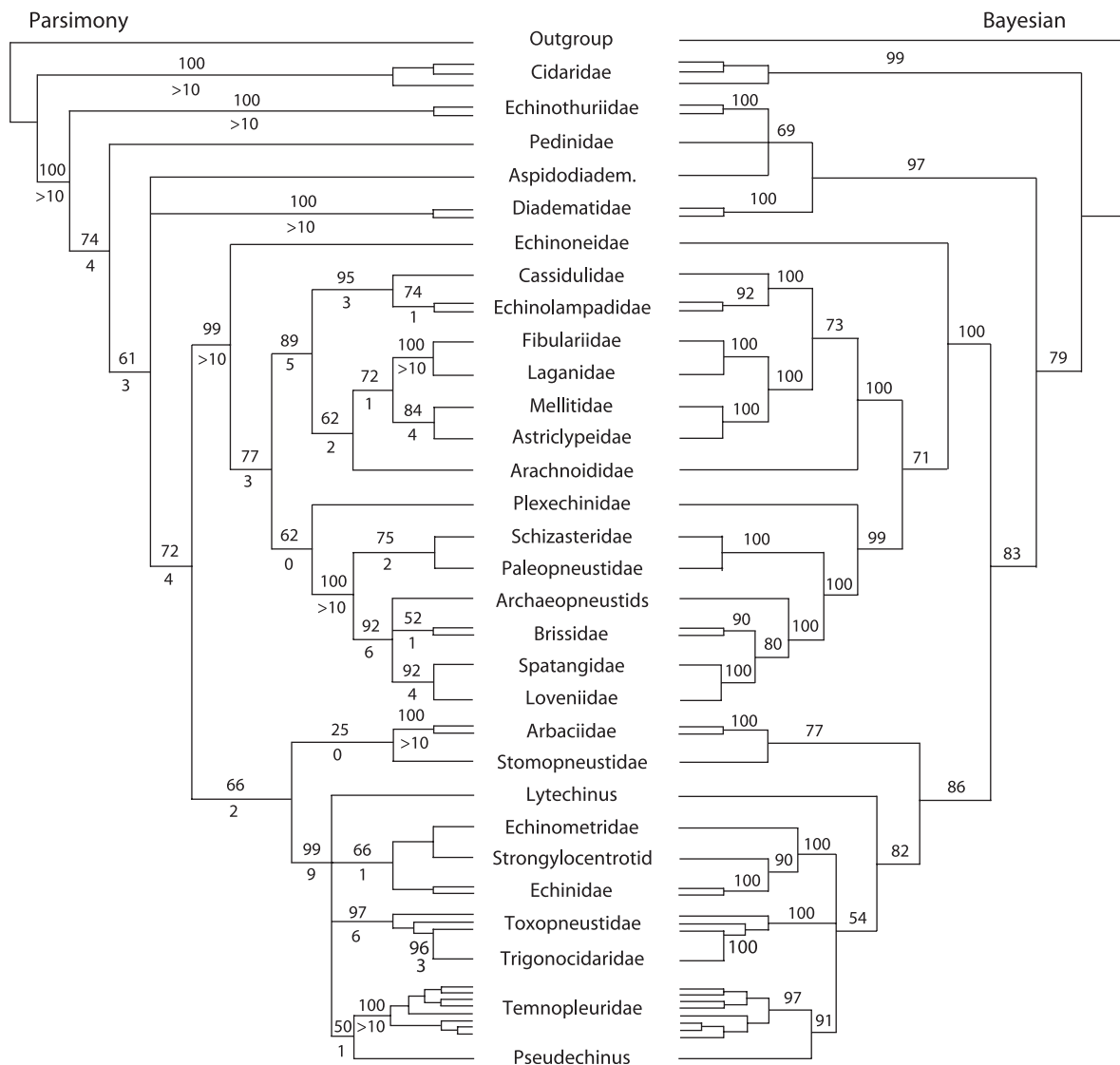


FIG. 3.—Trees derived from analysis of the combined morphological and molecular data. (Left hand side) Semistrict consensus of 6 equally most parsimonious trees derived from parsimony analysis. Bootstrap values (above) and Bremer support values (below) are given for each internal branch. (Right hand side) Tree derived from Bayesian analysis with posterior probabilities for each branch.

members of the Camarodonta. Some nodes are well supported, especially within the irregular echinoids, but there are a number of weakly supported areas.

The ML analyses of molecular data identified a topology very similar, but not identical, to that supported by morphological data (fig. 2). Of the few differences, only 1 taxon, *Fellaster*, is placed with strong support at different positions in the 2 rival trees. Other conflicting placements are weakly supported in either one or both of the trees. The morphological and molecular trees passed Templeton's test. Although the trees based on molecular data could not have been produced by the morphological data, the consensus tree of equally parsimonious trees from the morphological analysis was not significantly different in terms of TL from the molecular ML tree at $P > 0.05$ (number of differences = 31, rank sums = 331.0, $P = 0.07$ for Templeton's test, and $P = 0.15$ for Winning sites test). Therefore, the 2 sources of data, morphological and molec-

ular, can be considered to be suboptimal estimates of the same underlying topology.

Because different nodes are strongly supported in the 2 trees, a combined data set arguably provides the most appropriate way to combine the strengths of the 2 data sets. Parsimony analysis of the combined data found 6 trees whose strict consensus is shown (fig. 3, left-hand side). Bootstrap and Bremer support values were moderate to high for most branches. The Bayesian analysis of the combined data produces a tree that is very similar to the parsimony tree, except for its placement of the echinothurioids (*Araeosoma*) and the pedinoid *Caenopedina* (fig. 3, right-hand side). The topology in this part of the tree is closer to that derived from analysis of molecular data alone.

Stratigraphic Completeness Estimate

Both Bayesian and parsimony trees derived from the combined morphological and molecular data (fig. 3) were

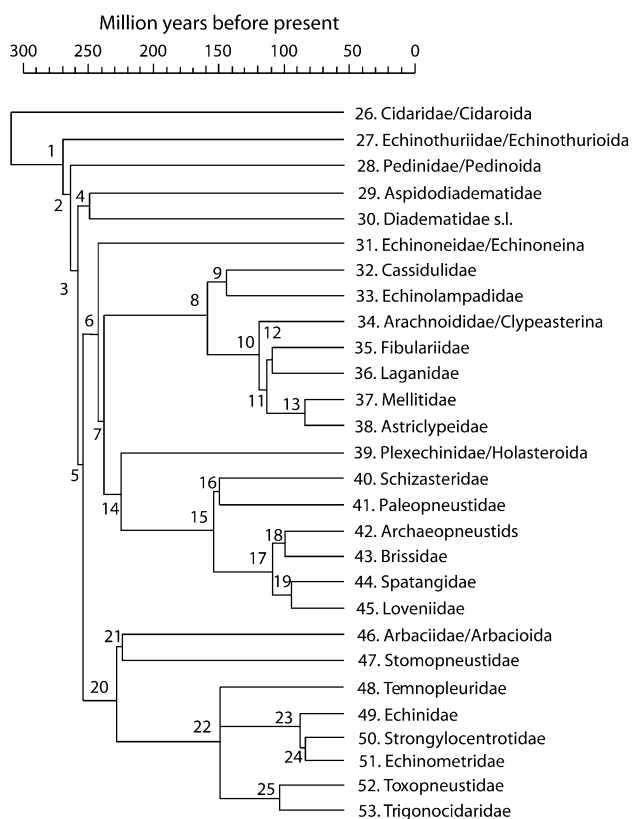


FIG. 4.—Calibrated tree constructed by optimizing the maximum parsimony cladogram derived from combined morphological and molecular data (fig. 3A) against the known fossil record of the group. Thick black lines = existing fossil record; thin lines = missing record that must be inferred from the cladogram structure; and scale at top = millions of years before present. Branches are numbered 1–53, with 1–25 being internal branches and 26–53 terminal branches. The oldest fossil member belonging to each of the branches numbered 1–53 are listed in table 2.

calibrated against the known fossil record to calculate the minimum amount of fossil record that must be missing. For the parsimony tree (table 2, fig. 4), the duration of all branch lengths (observed and inferred) implied by the combined morphological and molecular tree is 3,530 Myr, of which missing (ghost) lineages implied at family level constitute approximately 10% of the total duration (360 Myr inferred minimal time missing). Furthermore, almost half the perceived mismatch arises from relationships within the Camarodonta, where relationships are least well resolved. Almost identical results were obtained using the Bayesian combined-data tree (not shown).

Molecular Divergence Estimates

An ML analysis of the molecular data matrix for the 28 family representatives (plus outgroups) generated the tree shown in figure 5, and this was used to estimate molecular divergence times. When only the basal node of the ingroup was fixed, estimated divergence times varied considerably according to the specific method applied (table 3, fig. 6). The LF method failed to estimate the great majority (>80%) of paleontological nodes correctly, for the most part greatly underestimating divergence dates. The PL ap-

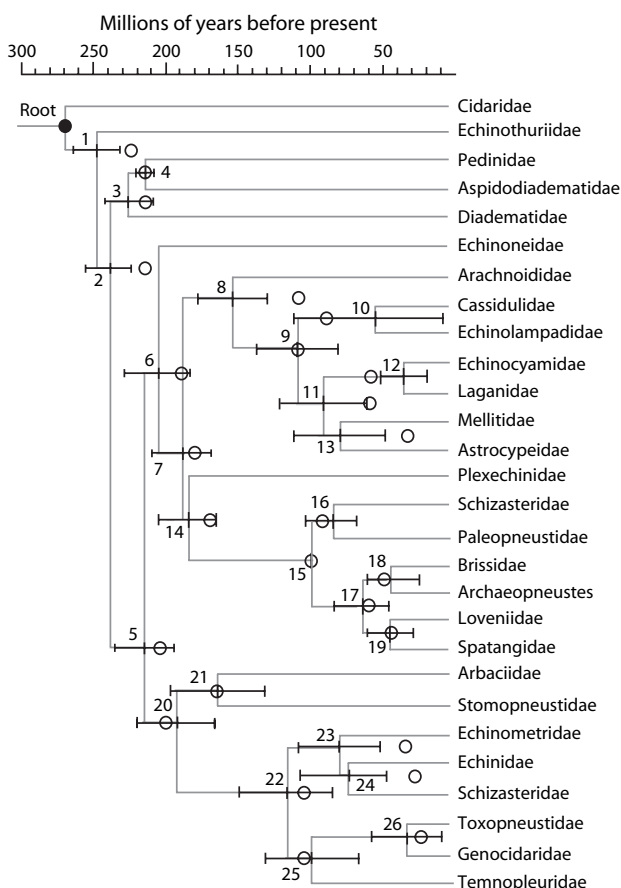


FIG. 5.—ML tree constructed from molecular data only and calibrated against time using divergence dates estimated by Sanderson's penalized likelihood method with logarithmic-penalty function. Black bars and error bars indicate estimate of divergence dates with 2 standard errors (see text for details). Circles indicate divergence times for internal nodes based on the fossil record. Numbers 1–26 refer to dated nodes listed in tables 3 and 4.

proach with additive-penalty function also performed poorly. However, the other 3 methods (NPRS, Bayesian, and log-PL) performed more or less equally well, having 17–18 of the 26 internal nodes congruent with paleontological estimates (i.e., the paleontological estimate lies within 2 SDs of the molecular estimate). Overall, the PL approach using logarithmic-penalty function gave the closest match to paleontological estimates.

When the additional 4 local calibration points were enforced, all molecular estimation methods performed reasonably well (table 4), having 17–18 of the 26 internal nodes congruent with paleontological estimates (figs. 5 and 7). However, in these conditions PL with additive-penalty function and LF greatly improved their rates of success, whereas the performance of NPRS was distinctly poorer compared with when only the ingroup node was constrained. Focusing on the number of nodes where the molecular estimate encompasses the paleontological estimate within 1 SD of the mean, the log-PL method outperforms other methods with NPRS performing the poorest.

Surprisingly, under these conditions LF performed better than all the relaxed methods we applied except for log-PL. This implies that relaxed methods should not be considered a priori better than methods for dating

Table 3

Divergence Dates for the 26 Internal Nodes Identified on the Tree Resulting from ML of Molecular Data (fig. 7, nodes 1–26) as Estimated by the LF, NPRS, Penalized Likelihood with Additive-Penalty Function (PL), Penalized Likelihood with Logarithmic-Penalty Function (log-PL), and Bayesian Methods (see text for details). In All Cases, the Basal Node Is Fixed at 265 Myr and No Internal Calibration Points Were Used

Node	Paleo Dates	LF			NPRS			PL (additional)			PL (logarithmic)			Bayes			
	Age (Myr)	Age (Myr)	95% CI (Myr)	C	Age (Myr)	2 × SD (Myr)	C	Age (Myr)	2 × SD (Myr)	C	Age (Myr)	2 × SD (Myr)	C	Age (Myr)	95% CI (Myr)	C	
Root	255																
1	220	222	193–251	●●	227	190–264	●●	216	177–253	●●	225	197–253	●●	226	180–254	●●	
2	210	177	147–207		213	185–241	●●	196	158–234	●●	212	182–242	●●	210	164–243	●●	
3	210	151	112–190		192	157–227	●	173	129–217	●	193	161–224	●	189	138–229	●	
4	210	122	84–158		167	131–203		146	106–186		169	132–206		160	109–205		
5	200	146	114–178		196	161–231	●●	173	134–212	●	197	165–229	●●	186	138–226	●●	
6	185	136	107–165		188	161–215	●●	164	125–203	●	190	161–219	●●	173	126–214	●●	
7	175	121	95–147		173	146–200	●●	146	111–181	●	177	145–209	●●	157	112–198	●●	
8	105	103	78–128	●●	146	115–177		118	84–152	●●	151	118–184		131	87–175	●	
9	105	83	62–104		110	81–139	●●	81	47–115	●	114	83–145	●●	98	60–141	●●	
10	85	29	0–59		68	9–127	●●	43	5–86	●	77	21–132	●●	52	7–104	●	
11	55	74	55–93	●	94	67–121		69	38–100	●●	97	68–126		78	44–121	●	
12	55	31	19–43		35	20–50		24	0–38		36	20–52		28	12–54		
13	30	65	4–86		81	50–112		58	26–90	●	83	51–115		65	35–105		
14	170	116	91–141		169	140–198	●●	142	106–178	●	174	142–206	●●	149	104–190	●	
15	95	52	38–66		78	56–100	●	61	42–80		82	61–104	●	70	39–110	●	
16	80	43	30–56		66	45–87	●	51	33–69		69	48–90	●	57	29–94	●	
17	55	35	24–46		52	32–72	●●	39	24–54		54	35–73	●●	47	23–83	●●	
18	45	22	17–27		35	17–53	●	26	13–39		37	21–53	●●	28	10–57	●	
19	40	24	15–33		35	19–51	●●	27	16–38		36	19–53	●●	32	13–62	●●	
20	195	130	101–159		180	150–210	●●	154	117–191		182	150–214	●●	167	119–210	●	
21	160	105	76–134		152	116–188	●●	129	94–164	●	155	119–191	●●	144	95–190	●●	
22	100	86	63–109	●	120	84–156	●	91	59–123	●●	125	97–153	●●	108	67–155	●●	
23	30	50	33–67		86	55–117		63	35–91		93	63–123		78	42–122		
24	25	47	29–65		82	50–114		59	31–88		88	57–119		70	36–112		
25	100	75	54–96		102	75–129	●●	76	45–107	●	108	81–135	●●	92	53–139	●●	
26	20	22	8–34	●●	36	5–67	●●	25	4–46	●●	43	12–74	●	37	11–76	●	
				0.15			0.63			0.39			0.65			0.62	

NOTE.—Paleo dates = estimated divergence times based on the first occurrence of fossils in the geological record; Age = divergence date estimated by method; 2 × SD = 2 SDs given as maximum and minimum age around the estimate in millions of years; C = congruence statistic measuring the fit of paleontological and molecular estimates of divergence time: ●● = paleontological date lies within 1 SD of the mean of the molecular date; ● = paleontological date lies between 1 and 2 SDs of the mean of the molecular date. The congruence statistic averaged over all 26 nodes is given at the foot of each column.

Table 4
Divergence Dates for the 26 Internal Nodes Identified on the Tree Resulting from ML of Molecular Data (fig. 7, nodes 1–26) as Estimated by the LF, NPRS, Penalized Likelihood with Additive-Penalty Function (PL), Penalized Likelihood with Logarithmic-Penalty Function (log-PL) and Bayesian Methods (see text for details). In All Cases the Basal Node Is Fixed at 265 Myr and 4 Internal Calibration Points Were Set as Minimal Times of Divergence (nodes in brackets)

Node	Paleo dates		LF				NPRS			PL (Additional)			PL (Logarithmic)			Bayes		
	Age (Myr)	Age (Myr)	95% CI (Myr)	C	Age (Myr)	2 × SD (Myr)	C	Age (Myr)	2 × SD (Myr)	C	Age (Myr)	2 × SD (Myr)	C	Age (Myr)	95% CI (Myr)	C		
Root	255	265	0		265	0		265	0		265	0						
1	220	245	227–263		243	226–260		244	227–261		243	226–260		257	240–265			
2	210	232	212–252		234	216–252		235	218–252		234	220–248		248	231–261			
3	210	221	196–246	●●	223	202–244	●	223	201–244	●	222	206–238	●	236	217–254			
[4]	210	210	210	●●	210	205–215	●●	210	205–215	●●	210	206–214	●●	218	210–234	●		
5	200	193	167–219	●●	220	197–243	●	218	196–240	●	211	189–233	●●	223	192–248	●		
6	185	181	157–205	●●	212	190–234		210	189–231		202	179–225	●	211	180–238	●		
7	175	164	144–184	●	196	174–218	●	194	173–215	●	185	164–206	●●	195	162–225	●		
8	105	138	115–161		166	137–195		164	136–192		149	123–175		165	125–203			
9	105	109	87–131	●●	126	97–155	●	124	96–152	●	104	72–136	●●	124	86–166	●●		
10	85	37	0–74	—	39	0–99	●	75	10–140	●●	54	6–102	●	67	8–128	●●		
[11]	55	98	78–118	—	107	79–135		106	80–132		88	58–118		99	62–145			
12	55	41	27–55	●	39	23–55	●	39	23–55	●	32	17–47		36	16–67	●		
13	30	86	63–109		92	61–123		91	60–122		75	43–107		83	48–127			
14	170	160	141–179	●	192	170–214	●	190	169–211	●	181	161–201	●	187	154–217	●		
[15]	95	95	95	●●	95	95–107	●●	95	95–104	●●	95	95	●●	110	95–144	●		
16	80	72	57–87	●	81	67–95	●●	80	66–94	●●	79	65–93	●●	91	65–124	●●		
17	55	54	39–69	●●	64	45–83	●●	63	43–83	●●	61	42–80	●●	76	48–112	●		
18	45	33	21–45	●	44	26–62	●●	42	23–61	●●	39	21–57	●●	46	19–81	●●		
19	40	36	24–48	●●	43	25–62	●●	42	25–59	●●	41	26–56	●●	52	25–88	●●		
20	195	171	143–199	●	204	180–228	●●	199	174–224	●●	189	162–216	●●	202	164–234	●●		
[21]	160	137	109–165	●	177	146–208	●●	168	135–201	●●	159	127–191	●●	175	130–215	●●		
22	100	111	87–135	●●	139	112–166		133	106–160		113	81–145	●●	133	88–180	●		
23	30	65	44–86		105	73–137		94	61–127		77	49–105		97	56–145			
24	25	61	39–83		100	68–132		89	55–123		73	43–103		87	48–135			
25	100	97	74–120	●●	118	90–146	●	113	86–140	●●	94	61–127	●●	113	68–162	●●		
26	20	28	11–45	●●	47	12–82	●	38	5–71	●	31	6–56	●●	47	14–96	●		
				0.56			0.48			0.52			0.62				0.50	

NOTE.—Paleo dates = estimated divergence times based on the first occurrence of fossils in the geological record; Age = divergence date estimated by method; 2 × SD = 2 SDs given as maximum and minimum age around the estimate in millions of years; C = congruence statistic measuring the fit of paleontological and molecular estimates of divergence time: ●● = paleontological date lies within 1 SD of the mean of the molecular date; ● = paleontological date lies between 1 and 2 SDs of the mean of the molecular date. The congruence statistic averaged over all 26 nodes is given at the foot of each column.

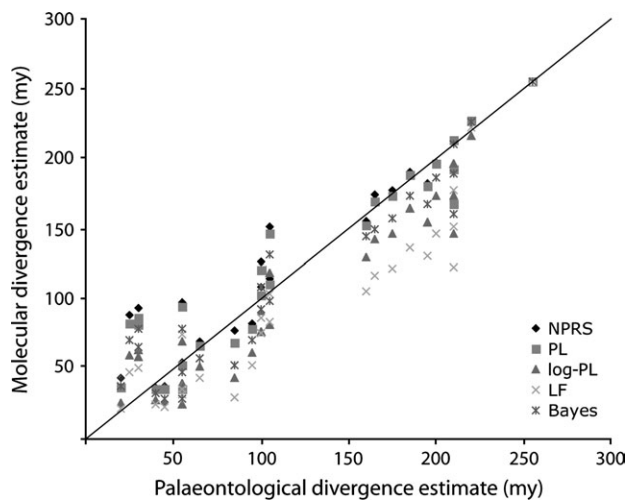


FIG. 6.—Plot of estimated divergence times based on molecular data (using 5 different methods as indicated) against paleontological estimate for the 26 nodes identified in table 3. Molecular divergence estimates were calculated using only fixed basal date without local calibration points. PL = penalized likelihood method with additive-penalty function; log-PL = penalized likelihood method with logarithmic-penalty function; and Bayes = Bayesian analysis.

divergence times imposing a molecular clock. Methods imposing a single, global, rate of evolution could return biased results, but the same is true for relaxed models as correctly noted, for example, by Welch et al. (2005) and Ho et al. (2005).

It is also interesting to note that log-PL and the Bayesian method of Thorne et al. (1998) were the most consistent, correctly estimating a high proportion of divergence times, no matter whether internal constraints different from the ingroup node were enforced. However, log-PL always performed better than the method of Thorne et al. (1998). It is, however, difficult to evaluate what causes this difference.

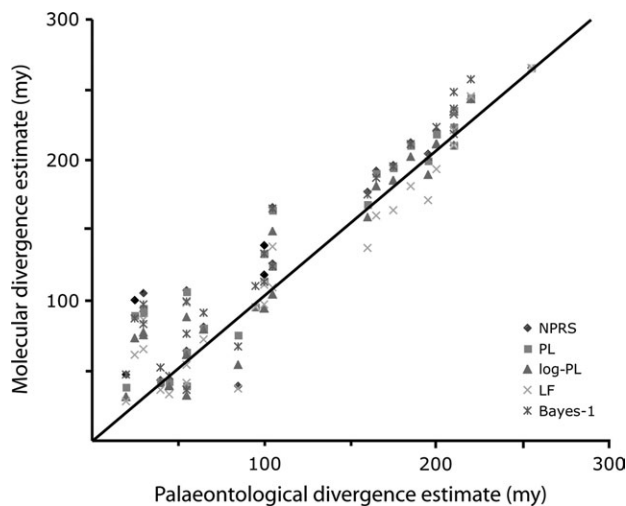


FIG. 7.—Plot of estimated divergence times based on molecular data (using 5 different methods as indicated) against paleontological estimate for the 26 nodes identified in table 3. Molecular divergence estimates were calculated using a fixed basal date plus 4 local calibration points, as described in the text. PL = penalized likelihood method with additive-penalty function; log-PL = penalized likelihood method with logarithmic-penalty function; and Bayes = Bayesian analysis.

The method of Thorne et al. (1998), as it is currently implemented, can only use simple models (e.g., $F84 + \Gamma$), but it build these models into the likelihood calculations, whereas Sanderson's (2004) software (r8s) relies on trees with branch lengths that must be previously estimated using other software (e.g., PAUP) and does not build the models into the likelihood calculations. However, in this way, it allows using more complex models and hence more accurate branch length estimations. If the use on simple and perhaps suboptimal model will be discovered to be the cause of the difference in performance between the 2 methods, then we should expect that the Bayesian method of Thorne et al. (1998) would improve as more models are integrated into it.

In any case, when the molecular estimates and paleontological estimates for each node are regressed, there is a strong and highly significant correlation with $r^2 = 0.91$ (for log-PL estimates). For the 8 cases where there is significant mismatch, paleontological data mostly underestimate the date of divergence.

Irrespective of which specific method was applied, mismatch between molecular and morphological estimates of divergence time is confined to the same 3 areas of the cladogram, 2 nodes near the base of the tree (the divergence of Euechinoidea; fig. 5, nodes 1 and 2), 4 nodes within the Clypeasteroidea (fig. 5, nodes 8 and 11–13), and 2 within the Camarodonta (fig. 5, nodes 23 and 24).

When the tree derived from the combined-data analysis (fig. 3) was used as the base for calculating divergences instead of the molecular tree, the overall congruence between molecular and paleontological estimates decreases (table 5), and the camarodont and clypeasteroid nodes that were incongruent in the previous analyses are also incongruent in this analysis. However, the basal nodes now show a better fit, with molecular and paleontological estimates for nodes 1 and 2 now congruent and in close agreement.

Discussion

There has been much recent discussion about the validity of the different methods used to estimate divergence times from molecular data but few empirical tests in groups with a good fossil record. The study of Pérez-Losada et al. (2004) tested a variety of different methods on the phylogeny of barnacles and their relatives but examined only a small number of nodes. In this study, we have adopted a similar approach but applied across a much larger number of paleontologically well-dated nodes. The fossil record and molecular-based divergence dates provide effectively independent estimates for the 26 internal nodes on our family-level tree. Furthermore, we specifically used the topology derived from only molecular data (rather than the combined morphological and molecular data tree) in order to set a more stringent test of the accuracy of the molecular approach.

It has been suggested that approaches that can accommodate rate variation in their inference procedures generate significantly more realistic results (e.g., Kishino et al. 2001; Sanderson 2002; Thorne and Kishino 2002; Pérez-Losada et al. 2004). However, whether these methods truly return better results seem strongly dependent on the model used to estimate rate variation, on the calibration points used

Table 5
Divergence Dates for the 23 Internal Nodes Identified on the Tree Resulting from ML Analysis of the Combined Morphological and Genetic Molecular Data as Estimated by the LF, NPRS, Penalized Likelihood with Additive-Penalty Function (PL), Penalized Likelihood with Logarithmic-Penalty Function (log-PL), and Bayesian Methods (see text for details). In All Cases, the Basal Node Is Fixed at 265 Myr and 4 Internal Calibration Points Were Used (nodes in brackets; see text for details)

Node	Paleo dates		LF		NPRS			PL (additional)			PL (logarithmic)			Bayes			
	Age (Myr)	Age (Myr)	95% CI (Myr)	C	Age (Myr)	2 × SD(Myrr)	C	Age (Myr)	2 × SD (Myr)	C	Age (Myr)	2 × SD (Myr)	C	Age (Myr)	95% CI (Myr)	C	
Root	255	265	0		265	0		265	0		265	0					
1	220	236	214–258	•	232	208–256	••	238	216–260	•	230	203–257	••	250	224–264		
[2]	210	210	195–225	••	216	193–239	••	223	223	••	214	194–234	••	237	213–258		
3	195	201	178–224	••	211	184–238	•	219	192–246	•	208	181–235	••	221	193–245	•	
4	200	167	145–189		180	156–204	•	189	163–215	••	175	150–200	•	193	160–226	••	
5	185	161	142–180		173	148–198	••	183	161–205	••	169	148–190	•	182	149–215	••	
6	175	149	131–167		161	139–183	•	170	147–193	••	156	136–176	•	168	135–201	••	
7	105	111	89–133	••	114	85–143	••	118	89–147	••	97	64–130	••	122	88–161	••	
8	90	32	0–64		67	16–118	••	55	0–115	•	44	0–88		61	9–116	•	
[9]	55	103	82–124		101	73–129		106	79–133		85	54–116	•	104	71–145		
10	55	92	74–110		87	63–111		92	68–116		73	48–97	•	84	57–123		
11	50	39	26–52	•	33	19–47		35	21–49		26	14–38		33	15–61	•	
12	30	81	60–102		76	50–102		80	53–107		63	38–88		72	46–109		
13	165	146	130–162		157	137–177	••	167	145–189	••	154	138–170	•	162	130–195	••	
[14]	95	95	95	••	95	94–96	••	95	94–96	••	95	95	••	105	95–131	•	
15	65	71	55–87	••	85	73–97		82	68–96	•	82	69–95		87	64–114	•	
16	55	45	33–57	•	58	40–76	••	54	37–71	••	51	33–69	••	65	41–95	••	
17	40	27	18–36		32	18–46	•	32	20–44	•	29	17–41	•	39	17–70	••	
18	195	146	123–169		162	136–188		168	139–197	•	153	125–181		170	133–207	•	
19	160	122	97–147		142	114–170	•	144	112–176	••	131	103–159		150	109–189	••	
[20]	100	99	78–120	••	108	82–134	••	112	84–140	••	95	69–128	••	111	72–154	••	
21	30	59	40–78		80	51–109		77	45–109		66	37–95		80	45–123		
22	30	48	30–66	•	68	50–86		64	33–95		55	29–81	•	66	34–107		
23	20	25	10–40	••	34	8–60	•	31	7–55	••	27	5–49	••	42	11–85	•	
				0.39			0.52			0.61			0.52				0.54

NOTE.—Paleo dates = estimated divergence times based on the first occurrence of fossils in the geological record; Age = divergence date estimated by method; 2 × SD = 2 SDs given as maximum and minimum age around the estimate in millions of years; C = congruence statistic measuring the fit of paleontological and molecular estimates of divergence time: •• = paleontological date lies within 1 SD of the mean of the molecular date; • = paleontological date lies between 1 and 2 SD of the mean of the molecular date. The congruence statistic averaged over all 25 nodes is given at the foot of each column.

(Welch et al. 2005), and, in the case of the Bayesian methods, on the prior on-divergence times chosen. For example, Aris-Brosou and Yang (2003) obtained high concordance with the fossil record from their molecular clock analysis; however, the major determinant of their divergence times was not their data, it was their prior (Ho et al. 2005; Welch et al. 2005). This also seems to be the case in our study, which found that although the LF method performed poorly when only the root is fixed, the inclusion of internal calibration points surprisingly increased its performance, whereas this same operation negatively affected NPRS. Furthermore, it is interesting to note that with our data, when LF behaved poorly, it underestimated most of the divergence time, contradicting the claim of Benton and Ayala (2003) that molecular clock methods by default overestimate divergence times. The correspondence we obtain between paleontological and molecular estimates of divergence using both the fully parametric Bayesian method of Thorne et al. (1998) and the semiparametric PL methods of Sanderson (2002) is reassuringly strong. In approximately 70% of cases, morphological and molecular estimates are congruent, that is, the paleontological estimate falls within the 95% confidence error bars of the molecular estimate. The use of internal fossil calibration points made little difference for log-PL and the Bayesian method of Thorne et al. (1998), and this consistency of performance is encouraging, suggesting that these methods should probably be favored over others. Overall, for our data, log-PL was best at recovering known divergence times, although employing better substitution models to estimate branch lengths in the Bayesian approach of Thorne et al. (1998) might be expected to improve the success rate of this method.

The few nodes that consistently show a mismatch between paleontological and molecular divergence estimates are restricted to 3 parts of the tree (fig. 5). These affect basal nodes 1 and 2 and nodes within the Echinoida clade (nodes 23 and 24) and clypeasteroid clade (nodes 8 and 11–13).

There are 3 reasons why disagreement between paleontological and molecular estimates of divergence might arise: inaccuracy of the phylogenetic reconstruction being used, incompleteness of the fossil record, or methodological problems in the way molecular estimates are derived. In order to test the first possibility, we calculated molecular divergence dates using the tree constructed from the combined morphological and molecular data (fig. 3), which we take as our best-supported estimate of phylogenetic relationships. This tree differs most obviously from the molecular tree (fig. 2) in the relationships of the basal nodes and the monophyly of clypeasteroids. Using the combined-data tree reduces the correspondence between molecular and paleontological estimates of divergence in some parts of the tree but improves the fit of basal nodes 1 and 2 considerably. Consequently, the poor match between paleontological and molecular estimates of divergence times seen in the 2 basal nodes in the molecular analyses could be the result of there being a suboptimal arrangement of basal taxa in the molecular tree. Significantly, however, mismatch between paleontological and molecular divergence estimates in the 2 other regions of the tree remains when using the best overall supported tree. In these cases, therefore, it is unlikely that

the mismatch is simply a problem of inaccurate phylogenetic reconstruction.

To check whether inadequacies in the fossil record could explain the mismatch, we calibrated the combined morphology and gene tree against the observed fossil record (fig. 4). Although this implies that the fossil record of echinoids is relatively well sampled overall, almost half the missing record identified is concentrated in the Camarodonta, suggesting that this group is seriously under-represented in the fossil record. The reason for this is obvious. It is notoriously difficult to assign fossil echinoids to specific families within the Camarodonta because Mortensen (1943a, 1943b) established his taxonomy on the basis of structures (details of pedicellariae anatomy) that are rarely preserved in fossils. Consequently, it is likely that the fossil record in this part of the tree is better than that suggested by our data and that it is simply a lack of diagnostic morphological characters that creates the apparent mismatch. The fossil record may well be there, it is just that we cannot currently place many fossil taxa with any degree of confidence. On this part of the tree, molecular estimates of divergence are always deeper than paleontological estimates (figs. 5–7). Consequently, the observed mismatch of molecular and paleontological estimates of divergence at nodes 23 and 24 is probably due to inadequacies of the fossil record, rather than due to errors in the molecular-based estimate.

For the remaining 4 nodes (nodes 8 and 11–13), the mismatch between paleontological and molecular estimates of divergence times is less easily explained. All these nodes refer to divergences within clypeasteroids, a morphologically complex and intensively studied clade (Durham 1955; Kier 1982; Mooi 1987, 1990). Furthermore, they have a highly distinctive synapomorphy (large numbers of pores perforating all ambulacral plates) that makes them immediately recognizable in the fossil record, even from small fragments. The group primarily inhabits shallow water, living in environments that are well represented in the fossil record. Consequently, the chance that crown-group clypeasteroids have a deep, hidden fossil record stretching back into the Early Cretaceous seems improbable from what we know of the echinoid fossil record.

Clypeasteroid divergence times estimated from molecular data may be afflicted by two potentially significant problems. First, the topology of this part of the molecular tree differs from that supported by the combined analysis of morphological and molecular data that we take as our best estimate of the phylogeny (cf. figs. 3 and 5). Specifically, the 2 cassiduloid families group within the clypeasteroids in the molecular tree (fig. 5) rather than as their sister group (fig. 3). However, even when the combined-data tree is used as the model to estimate divergence times (log-PL analysis, table 5), the paleontological and molecular-based estimates of divergence time remain strongly incongruent. Errors in the topology of the tree cannot, therefore, explain why molecular and paleontological estimates of divergence times are so incongruent in this part of the tree.

A second potential problem arises from the very uneven rate of molecular evolution shown by the cassiduloids and clypeasteroids on the combined-data tree. Whereas cassiduloids show 0.0051–0.0057 substitutions per site, all 5

clypeasteroids have rates about 4 times faster (0.018–0.026 substitutions per site). This may be a large amount of rate heterogeneity for molecular methods to accommodate. We predict that denser taxon sampling together with two or more local tie points (minima) might help generate a closer match between paleontology and molecular data in this part of the tree.

In conclusion, because morphological and molecular data both point to closely similar phylogenetic relationships among echinoid clades (with the exception of the clypeasteroid *Fellaster*) and we estimate that their fossil record is relatively complete, we have the first real opportunity to compare the accuracy of molecular and paleontological methods of estimating divergence times empirically. In approximately 70% of nodes tested, paleontological and molecular methods give congruent estimates of divergence dates using methods that allow for rate variation over the tree. Although this is pleasantly reassuring, it is not as good as we were hoping for. For those nodes where paleontological and molecular estimates of divergence time are incongruent, paleontological data mostly underestimate divergence times. In some cases, the problem clearly lies with the paleontological data because of the poor preservation potential of key diagnostic characteristics by which paleontologists recognize those clades. A second problem may be that the molecular tree being used to establish divergence times is suboptimal in its arrangement of certain branches. When these problems are discounted, only a small proportion of nodes (ca. 15%) show a significant mismatch between molecular and paleontological estimates where the error may be the fault of our molecular techniques. Although care is still needed in selecting calibration points when using molecular data to estimate divergence times, our study demonstrates that, so long as a realistic model of rate variation is applied (see Welch et al. 2005), modern parametric and semiparametric approaches that assume rate heterogeneity can and do generate realistic divergence time estimates in the great majority of cases.

Supplementary Material

Figures 6 and 7, in color, and the full description of morphological characters and character states together with the data matrix and the aligned sequences used in our analyses given as supplementary tables 1, 2, and 3 are available at *Molecular Biology and Evolution* online (<http://www.mbe.oxfordjournals.org/>).

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Appendix

Supporting evidence for the paleontological dating of nodes in this study is given in this section.

Paleontological dates for nodes 1–26 in figures 5–7 are fixed as follows.

Root: Cidaroida–Euechinoidea Divergence. These 2 clades have fundamentally different lantern and perignathic girdle structures, both derived compared with the arrangement seen in Paleozoic late stem-group archaocidarids. Kier (1984a) recognized that this divergence probably occurred in the Permian, and Smith and Hollingworth (1990) showed that the well-dated Late Permian (Kazanian) “*Miocidaris*” *keyserlingi* possessed cidaroid synapomorphies in both the structure of its lantern and perignathic girdle. As the oldest demonstrable member of the Cidaroida, *M. keyserlingi* sets the minimum time of crown-group divergence at 255 Myr.

- 1: Echinothurioida–Acroechinoidea Divergence. Echinothurioida are the only extant clade of echinoids whose tests have remained imbricate, and, consequently, they have the poorest fossil record of any of the groups considered here (see Smith and Wright 1990). This divergence is, therefore, dated by the first occurrence of a member of the Acroechinoidea, which has, as its primary synapomorphies, a fully tessellate test, the lack of multiple peristomial ambulacral plates, and the presence of true ambulacral plate compounding. “*Hemipedina*” *hudsoni* (Kier 1977), though relatively poorly known, has a solid, nonimbricate test plating and compound ambulacral plating characteristic of an Acroechinoidea. It comes from the Norian of Oman and dates the origin of Acroechinoidea at around 220 Myr.
- 2: Aulodont–[Echinacea + Irregularia] Divergence. The major synapomorphy distinguishing the Echinacea and Irregularia from aulodonts, is their possession of keeled, as opposed to grooved, teeth. Isolated teeth are not uncommon from the Middle Triassic (Carnian) St Cassian beds, but all are grooved (Vadet 1999). The lantern of the Late Triassic (Rhaetian) *Diademopsis serialis* is known in detail (Smith 1981) and is aulodont in structure. The earliest known keeled teeth come from the Lower Jurassic (Pliensbachian; Smith 1981; Markel 1978). However, Kier (1977, p. 33) argued that the Late Triassic (Rhaetian) *Pseudodiadema silbinense* (which is more correctly placed in the genus *Stereopyga*) represented the oldest echinacean on the basis of general test characteristics. These occurrences suggest that this divergence had definitely occurred by 195 Myr and probably by 210 Myr.
- 3: Diadematidae–[Aspidodiadematidae + Pedinidae] Divergence. The pairing of Aspidodiadematidae and Pedinidae as sister groups to the exclusion of Diadematidae in the molecular analysis is unexpected. Traditionally, the Diadematidae and Aspidodiadematidae have been grouped together in the order Diadematoida and distinguished from members of the order Pedinoida by their crenulated tubercles (e.g., Mortensen 1940; Smith 1981).

- No morphological synapomorphy exists uniting the molecular clade Aspidodiadematidae + Pedinidae, and so the date of this divergence is set by the oldest member showing synapomorphies of any 1 of the 3 constituent families. The Upper Triassic (Rhaetian) *Diademopsis serialis* is generally placed in the Pedinidae (Smith 1981, 2006) as is the Lower Jurassic (Pliensbachian) *Hemipedina*, which displays the derived dicyclic pedinid apical disc. Synapomorphies of the Aspidodiadematidae include their distinctive apical disc plating, which is only very loosely connected to the corona, and their distinctive sphaeridial pit arrangement. Both occur in the Lower Jurassic (Pliensbachian) *Gymnotiara varusense* (Smith 2006). Divergence must have occurred before 195 Myr and probably by 210 Myr, the date used here.
- 4: Pedinoida–Aspidodiadematidae Divergence. See Discussion for node 4. The oldest member of the Aspidodiadematidae is *G. varusense*, from the Pliensbachian at 195 Myr (Smith 2006). The oldest member of the Pedinoida is *Diademopsis serialis* from the Rhaetian, at 210 Myr.
 - 5: Echinacea–Irregularia Divergence. Both these groups have keeled teeth but differ in their apical disc structure: Echinacea having a normal endocyclic apical disc with 5 gonopores in contrast to the more derived apical disc arrangement of primitive irregular echinoids in which the periproct is displaced to the posterior behind the posterior genital plates and genital plate 5 has lost its gonopore. The oldest putative echinacean is the Rhaetian *Stereopyga silbinense* (redescribed by Kier 1977). However, there is nothing to definitely place this as a derived Echinacea rather than a stem-group Echinacea + Irregularia. Cladistic analysis consistently identifies the Sinemurian *Jesionekchinus hawkinsi* as the earliest member of the Irregularia (Smith and Anzalone 2000; Barras forthcoming), establishing divergence at 200 Myr.
 - 6: Echinoneoida–Microstomata Divergence. Among irregular echinoids, Microstomata is more derived than its sister group Echinoneoida in having evolved specialized ambulacral zones in the form of aboral petals and adoral phyllodes (Barras forthcoming). The Lower Jurassic (Toarcian) *Galeropygus sublaevis* has distinct phyllodes and a subanal groove and establishes the date of this split as no younger than 185 Myr.
 - 7: Atelostomata–Neognathostomata Divergence. Both atelostomates and neognathostomates have long, well-documented stem groups. The synapomorphies that distinguish basal atelostomates are mostly associated with apical disc arrangement, atelostomates having a “stretched” apical disc with ocular plates 2 and 4 intercalated between the anterior and posterior pair of genital plates, as opposed to the more compact ethmophract disc structure of early neognathostomates (Smith 1981; Barras forthcoming). The earliest echinoid to show the derived atelostomate disc plating is *Aulacopygus caudatus* from the Late Bajocian (Smith 2006), setting this divergence as no later than 175 Myr.
 - 8: *Fellaster*–[Cassiduloida + Scutellina] Divergence. As discussed in the text, the position taken by *Fellaster* in the molecular tree is in strong contradiction to that in traditional taxonomies based on morphology (Mortensen 1948b; Smith 1981; Mooi 1987, 1990). Traditional taxonomies place *Fellaster* as a member of the Clypeasteroidea along with its sister group the Scutellina, whereas the molecular topology implies that Clypeasteroidea are diphyletic, with the 2 cassiduloids as closer to members of the Scutellina than *Fellaster*. A further complication is that the Cassiduloida, as currently recognized, is a paraphyletic group (Smith 2001). Cladistic analysis of morphological data suggests that the cassiduloid families Cassidulidae and Echinolampadidae form a clade, whereas a third cassiduloid family, Apatopygidae (not included in the molecular analysis), represents the closest living sister group to *Fellaster* and the Scutellina (Smith 2001). This latter grouping is based on a derived pattern of ambulacral plating seen in apatopygids and undoubted stem-group Clypeasteroidea and implies that the Clypeasteroid–[Cassidulidae + Echinolampadidae] split can be traced back in the Late Cretaceous (*Nucleopygus angustatus* being the earliest recognizable representative of the Apatopygidae + Clypeasteroidea total group). Given the molecular topology, divergence at this node is thus placed at 105 Myr.
 - 9: Cassiduloida–Scutellina Divergence. As for node 8, above.
 - 10: Cassidulidae–Echinolampadidae Divergence. The phylogenetic relationships of cassiduloids remain problematic because of the small number of convincing synapomorphies available from morphology (Smith 2001). Convincing cassidulids (*Rhyncholampas*) and echinolampadids (*Vologesia*) are, however, present and clearly differentiated by the Maastrichtian (70 Myr; Smith and Jeffery 2000) with echinolampadids extending back to the Santonian (*Hungaresia ovum*; Smith 2006). This places their divergence at no younger than 85 Myr.
 - 11: Scutelliformes–Laganiformes Divergence. The most paleontologically useful synapomorphies characterizing these 2 clades are 1) the single terminal interambulacral plate unique to laganines and 2) the complex arrangement of buttressing internal meshwork in the Scutellina (e.g., Mooi 1987). Both *Sismondia* and *Echinocyamus* have interambulacra that terminate in a single plate, and both extend back to the Eocene. However, the very oldest members of each genus (the Lower Eocene *Echinocyamus gurnahensis* and *Sismondia logotheti*) are not sufficiently known to confirm that their interambulacra show the laganiform arrangement (e.g., Roman and Struogo 1994). The oldest scutelliform, *Eoscutum doncieuxi*, is also of Lower Eocene and has the apomorphic dense internal buttressing of a scutelliform (Roman 1990; Smith 2006). Divergence is thus taken as having occurred around 50–55 Myr.
 - 12: Echinocyamidae–Laganidae Divergence. Echinocyamidae are small laganiforms with interambulacral zones ending adapically in a single plate and with internal buttressing composed of radial buttresses only. Unlike other laganiform families, there is no certainty that this is a monophyletic clade, and it could include pedomorphic forms of other laganiforms. Small laganids might be extremely difficult to distinguish from taxa currently placed in the Echinocyamidae. Forms attributed to *Echinocyamus* extend back into the Eocene (Kier 1968), with *E. gurnahensis* being the oldest (Roman

- and Struogo 1994). Laganidae are larger forms with a well-developed internal skeleton of both concentric partitions and adradial bars. The highly stellate nature of their basicoronal plates, with their radially directed points, is also a synapomorphy. One stem-group branch of the Laganidae shows an additional synapomorphy of having pseudocompound plating in their petals and is found as far back as the Middle Eocene (Kier 1968, 1980). The 2 groups were thus clearly separated by the Middle Eocene with a divergence probably in the Early Eocene, at 50–55 Myr.
- 13: Mellitidae–Astricylpeidae Divergence. Mellitidae and Astricylpeidae are sister taxa, united by having a well-developed microcanal system internally, spines and tubercles on the oral surface clearly differentiated into food-gathering and locomotory areas, and the periproct opening in the first postbasicoronal interambulacral plate. Mellitidae is the more derived taxa in possessing an anal lunule (Smith 2006). Although both clades possess ambulacral notches or lunules, these are constructed differently, as noted by Seilacher (1979), and are presumably independently evolved in each. *Amphiope duffi*, from the Upper Oligocene of North Africa, is the oldest fossil with astricylpeid-style lunules, whereas *Encopociae*, from the Lower Miocene, is the oldest fossil with an anal lunule (Durham 1955). This places their divergence at around 30 Myr.
- 14: Holasteroidea–Spatangoida Divergence. Traditionally, holasteroids and spatangoids have both been thought as originating at the base of the Cretaceous (e.g., Durham 1966). However, this is because both are apomorphy based rather than total group definitions. Mintz (1968) was first to realize that both originated from among the Jurassic “disasteroids” and might have a long pre-Cretaceous record. Recent cladistic analysis of this paraphyletic grade by Barras (forthcoming) confirms this and places the split at 170 Myr, based on the occurrence of *Collyrites ellipticus*, the oldest stem-group holasteroid, and *Disaster moeschi*, the oldest stem-group spatangoid.
- 15: Schizasterina–Micrasterina Divergence. Micrasterina and Schizasterina display fundamentally different derived fasciole patterns (Smith and Stockley 2005): micrasterines having a subanal fasciole and schizasterines a lateroanal fasciole. *Periaster elatus*, from the Cenomanian, is the earliest spatangoid to have a lateroanal fasciole, whereas the slightly younger *Micraster leskei*, from the Lower Turonian, is the oldest micrasterine with a subanal fasciole (Smith 2006), although the Cenomanian “*Micraster*” *distinctus* may be an early micrasterine, lacking a subanal fasciole. Divergence is set at 95 Myr.
- 16: Schizasteridae–Paleopneustidae Divergence. These 2 families share a similar fasciole pattern, but in Paleopneustidae the lateroanal fasciole is discrete from the peripetalous fasciole, whereas in Schizasteridae the 2 fascioles are coalesced around the anterior. Markov and Solovjev (2001) have argued that paleopneustids (here defined to include pericosmids) are derived and that their fasciole pattern is the synapomorphic state. The oldest echinoid with paleopneustid fascioles and apical disc according to Markov and Solovjev is *Eopericosmus typicus* from the Early Palaeocene, establishing divergence at 65 Myr.
- 17: [Brissidae + Archaeopneustid]–[Spatangidae + Loveniidae] Divergence. Cladistic analysis of the Spatangoida (Stockley et al. 2005) suggested that the composition of all these families needs revision. Nevertheless, considering just those taxa discussed in this paper, a dichotomy between *Brissus*, *Meoma*, and *Archaeopneustes* on the one hand and *Spatangus*, *Echinocardium*, and *Lovenia* on the other hand is clearly evident. The occurrence of *Granopatagus* (a Spatangidae) and *Brissus* (a Brissidae) by the late Middle Eocene demonstrates that this split had already occurred by 45 Myr. If the cladogram in Stockley et al. (2005) is correct, then it suggests that the divergence was even earlier, at around 55 Myr, because of the occurrence of *Eupatagus* in the Early Eocene (Roman and Struogo 1994).
- 18: Brissidae–Archaeopneustid Divergence. The brissid *Meoma* and the deep-sea Archaeopneustid *Archaeopneustes* are sister groups in both morphological and molecular analyses of spatangoid genera by Stockley et al. (2005). *Archaeopneustes* extends back in the fossil record to the Oligocene (*Heteropneustes elegans* (Jackson)), whereas *Meoma* extends back to the Middle Eocene (*Meoma caobaensis* Sanchez Roig) (Kier 1984b; Smith 2006). Their divergence is therefore placed at 45 Myr.
- 19: Spatangidae–Loveniidae Divergence. Loveniidae are more derived than Spatangidae, having as synapomorphies an inner fasciole and deeply sunken aboral tubercles. The latter are found in *Hemimaretia*, with *Hemimaretia subrostrata* dated from the late Middle Eocene of the United States representing the oldest Loveniidae (A. Kroh, personal communication). This, together with the occurrence of the spatangid *Granopatagus* by the late Middle Eocene, demonstrates that this split had already occurred by 40 Myr.
- 20: Stirodonta–Camarodonta Divergence. Camarodonta are a monophyletic clade united by their characteristic and derived lantern structure (Mortensen 1943a; Smith 1981). Unfortunately, lanterns are rarely preserved in fossils, making distinction between camarodonts and stirodonta difficult. Echinoid-style plate compounding is a reliable synapomorphy for crown-group Camarodonta but is absent from the earliest forms (Zeuglopleuridae). A further problem is that the major clades within the Stirodonta are possibly not a monophyletic group. Smith and Wright (1996) argued that the sister group to camarodonts was the stirodont-group Salenioida as both shared the derived character of possessing a suranal plate. The split between stirodonta with and without suranal plates forming an integral part of the apical disc occurred very early, with the Hettangian *Acrosalenia chartoni* being the oldest member of the former group. Divergence is thus set at 195 Myr.
- 21: Arbacioida–Stomopneustidae Divergence. Stomopneustidae are a small, well-defined group with characteristic ambulacral plating and test morphology that can be traced back to the Upper Jurassic (Oxfordian) *Phymechinus mirabilis* (Smith 2006). However, no cladistic analysis has been carried out to investigate the relationships of this clade to potential Middle Jurassic sister taxa and

its divergence with arbacioids. Tentatively, therefore, divergence is set at 160 Myr, though it may be anywhere up to 30 Myr older.

- 22: Echinoida–Temnopleuroida Divergence. These 2 clades share the synapomorphy of having echinoid-style ambulacral plate compounding, with temnopleuroids further derived in having a strongly developed ornament over test plates. The Cenomanian *Zeuglopleurus* has a primitive forerunner to echinoid-style ambulacral plate compounding and a strong ornament of pits (Smith and Wright 1996) and is generally accepted to be the oldest stem-group temnopleuroid. Divergence is thus estimated at 100 Myr.
- 23: Echinometridae–[Strongylocentrotidae + Echinidae] Divergence. Neither Echinidae nor Strongylocentrotidae are defined on synapomorphies that are likely to be preserved in fossils (see below) and so are difficult to recognize in the rock record. However, several of the Echinometridae have a very unusual ovate test, and this distinctive morphology is first seen in the Upper Oligocene *Plagiechinus priscus* (see Smith 2006), placing their divergence at 30 Myr.
- 24: Echinidae–Strongylocentrotidae Divergence. These families are distinguished on pedicellarial features, which are rarely preserved in fossils. Although Echinidae usually have trigeminate ambulacral plating and Strongylocentrotidae polygeminate plating, there are exceptions and based on test features alone no reliable distinction can be drawn. Dating is therefore based on finding taxa that are close enough in morphology to the extant species to be considered congeneric. The oldest putative strongylocentrotid *Strongylocentrotus antiquus* from the Lower Miocene of Australia was described by Philip (1965), whereas *Psammechinus dubius* from the Early Miocene (see Kroh 2005) is probably a valid Echinidae. No earlier species can be reliably placed in these families (see also Smith 1988). A divergence date is therefore tentatively set at 25 Myr.
- 25: Temnopleuridae–[Toxopneustidae + Trigonocidaridae] Divergence. Temnopleuridae have, as their most distinctive apomorphy, a test ornamentation of deep sutural pits, a character first seen in the Cenomanian *Zeuglopleurus* (Smith and Wright 1996). Divergence is thus estimated at 100 Myr. However, Cretaceous temnopleuroids such as *Zeuglopleurus* lack echinoid-style ambulacral plate compounding, a character that unites all extant Temnopleuridae, Trigonocidaridae, Toxopneustidae as well as Echinoida (Smith 1988). It is therefore possible that this divergence is younger than commonly estimated. As a minimum date of divergence, we used the first appearance of a member of the Toxopneustidae, *Lytechinus* [*Scoliechinus*] *axiologus* Arnold & Clark, at 45 Myr, as our local calibration point.
- 26: Toxopneustidae–Trigonocidaridae Divergence. Molecular data unexpectedly placed the trigonocidarid *Genocidaris* within the Toxopneustidae, despite its lack of deep buccal notches, and there are no clear morphological synapomorphies uniting *Genocidaris* and its molecular sister taxon *Cyrtechinus*. *Genocidaris*, however, has an almost identical in-test morphology to the fossil *Arbacina*, with *Arbacina monilis* (Desmarest) from the Burdigalian being

the oldest representative known. This establishes a putative divergence date of 20 Myr for *Genocidaris*.

Literature Cited

- Aris-Brosou S, Yang Z. 2003. Bayesian models of episodic evolution support a late Precambrian diversification of the Metazoa. *Mol Biol Evol* 20:1947–54.
- Barras C. 2007. Phylogeny of the Jurassic to early Cretaceous ‘disasteroid’ echinoids (Echinoidea; Echinodermata), and the origins of spatangoids and holasteroids. *J Syst Palaeontol*. Forthcoming.
- Benton MJ. 1995. Testing the time axis of phylogenies. *Phil Trans R Soc Lond B* 349:5–10.
- Benton MJ. 1999. Early origins of modern birds and mammals: molecules vs. morphology. *BioEssays* 21:1043–51.
- Benton MJ. 2001. Finding the tree of life: matching phylogenetic trees to the fossil record through the 20th century. *Proc R Soc Lond B* 268:2123–30.
- Benton MJ, Ayala FJ. 2003. Dating the tree of life. *Science* 300:1698–700.
- Blair JE, Hedges SB. 2005. Molecular clocks do not support the Cambrian explosion. *Mol Biol Evol* 22:387–90.
- Bremer K. 1994. Branch support and tree stability. *Cladistics* 10:295–304.
- Cunningham CW. 1997. Can three incongruence tests predict when data should be combined? *Mol Biol Evol* 14:733–40.
- Dean J. 2005. Skeletal homologies, phylogeny and classification of the earliest asterozoan echinoderms. *J Syst Palaeontol* 3: 29–114.
- De Queiroz A, Donoghue MJ, Kim J. 1995. Separate versus combined analysis of phylogenetic evidence. *Annu Rev Ecol Syst* 26:657–81.
- Donoghue PCJ, Smith MP. 2003. Telling the evolutionary time: molecular clocks and the fossil record. *Syst Assoc Spec Pub* 66:296.
- Durham JW. 1955. Classification of clypeasteroid echinoids. *Univ Calif Publ Geol Sci* 31:73–198.
- Durham JW. 1966. Phylogeny and evolution. In: Moore RC, editor. *Treatise on invertebrate paleontology Part U. Echinodermata 3*. Boulder, CO: University of Kansas Press and The Geological Society of America. p U266–9.
- Easteal S. 1999. Molecular evidence for the early divergence of placental mammals. *Bioessays* 21:1052–8.
- Farris JS, Källersjö M, Kluge AG, Bult C. 1994. Testing significance of incongruence. *Cladistics* 10:315–9.
- Felsenstein J. 2004. PHYLIP (phylogeny inference package). Version 3.6. Distributed by the author. Seattle: Department of Genome Sciences, University of Washington.
- Fountaine TMR, Benton MJ, Dyke GJ, Nudds RL. 2005. The quality of the fossil record of Mesozoic birds. *Proc R Soc B* 272:289–94.
- Gingerich PD. 2005. Cetacea. In: Rose KD, Archibald JD, editors. *Placental mammals: origin, timing, and relationships of the major extant clades*. Baltimore, MD: Johns Hopkins University Press. p 234–52.
- Hitchin R, Benton MJ. 1997. Congruence between parsimony and stratigraphy: comparisons of three indices. *Paleobiology* 23:20–32.
- Ho SYW, Phillips MJ, Drummond AJ, Cooper A. 2005. Accuracy of rate estimation using relaxed-clock models with a critical focus on the early metazoan radiation. *Mol Biol Evol* 22:1355–63.
- Hordijk W, Gascuel O. 2005. Improving the efficiency of SPR moves in phylogenetic tree search methods based on maximum likelihood. *Bioinformatics* 21:4338–47.

- Huelsenbeck JP. 1994. Comparing the stratigraphic record to estimates of phylogeny. *Paleobiology* 20:470–83.
- Jeffery CH, Emler RB, Littlewood DTJ. 2003. Phylogeny and evolution of developmental mode in temnopleurid echinoids. *Mol Phylogenet Evol* 28:99–118.
- Kier PM. 1968. Echinoids from the Middle Eocene Lake City Formation of Georgia. *Smithson Misc Collect* 153:1–45, pls 1–10.
- Kier PM. 1977. Triassic echinoids. *Smithson Contrib Paleobiol* 30:1–88.
- Kier PM. 1980. The echinoids of the Middle Eocene Warley Hill Formation, Santee Limestone, and Castle Hayne Limestone of North and South Carolina. *Smithson Contrib Paleobiol* 39:1–102.
- Kier PM. 1982. Rapid evolution in echinoids. *Palaeontology* 25:1–9.
- Kier PM. 1984a. Echinoids from the Triassic (St Cassian) of Italy, their lantern supports, and a revised phylogeny of Triassic echinoids. *Smithson Contrib Paleobiol* 56:1–41.
- Kier PM. 1984b. Fossil spatangoids of Cuba. *Smithson Contrib Paleobiol* 55:1–336.
- Kishino H, Thorne JL, Bruno WJ. 2001. Performance of a divergence time estimation method under a probabilistic model of rate evolution. *Mol Biol Evol* 18:352–61.
- Kroh A. 2005. *Catalogus fossilium Austriae. Band 2 Echinoidea neogenica*. Wien, Austria: Osterreichischen Akademie der Wissenschaften. 210 p, 82 pls.
- Langley CH, Fitch W. 1974. An estimation of the constancy of the rate of molecular evolution. *J Mol Evol* 3:161–77.
- Lee Y-H. 2003. Molecular phylogenies and divergence time of sea urchin species of Strongylocentrotidae, Echinoidea. *Mol Biol Evol* 20:1211–21.
- Lessios HA, Kessing BD, Pearse JS. 2001. Population structure and speciation in tropical seas: global phylogeography of the sea urchin *Diadema*. *Evolution* 55:955–75.
- Lessios HA, Kessing BD, Robertson DR, Pauley G. 1999. Phylogeography of the pantropical sea urchin *Eucidaris* in relation to land barriers and ocean currents. *Evolution* 53:806–17.
- Lewis DN, Ensom PC. 1982. *Archaeocidaris whatleyensis* sp. nov. (Echinoidea) from the carboniferous limestone of Somerset and notes on echinoid phylogeny. *Bull Brit Mus (Nat Hist) Geol Ser* 36:77–104.
- Lewis PO. 2001. A likelihood approach to estimating phylogeny from discrete morphological character data. *Syst Biol* 50:913–25.
- Littlewood DTJ, Smith AB. 1995. A combined morphological and molecular phylogeny for echinoids. *Phil Trans R Soc Lond B* 347:213–34.
- Littlewood DTJ, Smith AB, Clough KA, Emson RH. 1997. The interrelationships of the echinoderm classes: morphological and molecular evidence. *Biol J Linn Soc* 61:409–38.
- Maddison DR, Maddison WP. 2001. *MacClade 4. Analysis of phylogeny and character evolution. Version 4.03*. Sunderland, MA: Sinauer Associates.
- Markel K. 1978. On the teeth of the recent cassiduloid *Echinolampas depressa* Gray, and on some liassic fossil teeth nearly identical in structure (Echinodermata, Echinoidea). *Zoomorphology* 89:125–44.
- Markov AV, Solovjev AN. 2001. Echinoids of the family Paleopneustidae (Echinoidea, Spatangoida): morphology, taxonomy, phylogeny. *Trans Pal Inst Moscow* 280:1–109.
- Mintz LW. 1968. Echinoids of the Mesozoic families Collyritidae d'Orbigny, 1853, and Disasteridae Gras, 1848. *J Paleontol* 42:1272–88.
- Mooi R. 1987. A cladistic analysis of the sand dollars (Clypeasteroidea: Scutellina) and the interpretation of heterochronic phenomena. [PhD thesis]. Toronto, Canada: University of Toronto; 208 p.
- Mooi R. 1990. Paedomorphosis, Aristotle's Lantern, and the origin of the sand dollars (Echinodermata; Clypeasteroidea). *Paleobiology* 16:25–48.
- Mortensen T. 1928. A monograph of the Echinoidea. I. Cidaroida. Copenhagen, Denmark: C. A. Reitzel. p 551, 88 pls.
- Mortensen T. 1935. A monograph of the Echinoidea. II. Bothriocidaroida to Stirodonta. Copenhagen, Denmark: C. A. Reitzel. p 647, 89 pls.
- Mortensen T. 1940. A monograph of the Echinoidea. III(i). Aulodonta. Copenhagen, Denmark: C. A. Reitzel. p 369, 79 pls.
- Mortensen T. 1943a. A monograph of the Echinoidea. III(ii). Camarodonta 1. Copenhagen, Denmark: C. A. Reitzel. p 553, 56 pls.
- Mortensen T. 1943b. A monograph of the Echinoidea. III(ii). Camarodonta 2. Copenhagen, Denmark: C. A. Reitzel. p 466, 66 pls.
- Mortensen T. 1948a. A monograph of the Echinoidea. IV(1). Holoctypoida, Cassiduloida. Copenhagen, Denmark: C. A. Reitzel. p 363, 14 pls.
- Mortensen T. 1948b. A monograph of the Echinoidea. IV(2). Clypeasteroidea. Copenhagen, Denmark: C. A. Reitzel. p 471, 72 pls.
- Mortensen T. 1950. A monograph of the Echinoidea. V(1). Spatangoida. Copenhagen, Denmark: C. A. Reitzel. p 432, 25 pls.
- Mortensen T. 1951. A monograph of the Echinoidea. V(2). Spatangoida 2. Copenhagen, Denmark: C. A. Reitzel. p 593, 64 pls.
- Norrell MA, Novacek MJ. 1992. Congruence between superpositional and phylogenetic patterns: comparing cladistic patterns with fossil records. *Cladistics* 8:319–37.
- Pérez-Losada M, Hoeg JT, Crandall KA. 2004. Unravelling the evolutionary radiation of the Thoracican barnacles using molecular and morphological evidence: a comparison of several divergence time estimation approaches. *Syst Biol* 53:244–64.
- Peterson KJ, Butterfield NJ. 2005. Origin of the Eumetazoa: testing ecological predictions of molecular clocks against the Proterozoic fossil record. *PNAS* 102:9547–52.
- Peterson KJ, Lyons JB, Nowak KS, Takacs CM, Wargo MJ, McPeck MA. 2004. Estimating metazoan divergence times with a molecular clock. *PNAS* 101:6536–41.
- Philip GM. 1965. The Tertiary echinoids of South-eastern Australia III Stirodonta, Aulodonta, and Camarodonta (1). *Proc R Soc Victoria* 78:181–196, pls 26–29.
- Pisani D, Poling LL, Lyons-Weiler M, Hedges SB. 2004. The colonization of land by animals: molecular phylogeny and divergence times among arthropods. *BMC Biology* 2:1.
- Posada D, Crandall KA. 1998. MODELTEST: testing the model of DNA substitution. *Bioinformatics* 14:817–8.
- Rodriguez-Trelles F, Tarrío R, Ayala FJ. 2002. A methodological bias towards overestimation of molecular evolutionary time scales. *Proc Natl Acad Sci USA* 99:8112–5.
- Roman J. 1990. l'ancetre Eocene des Scutellidae (Echinoidea; Clypeasteroidea). In: De Ridder C, Dubois P, Lahaye M-C, Jangoux M, editors. *Echinoderm research*. Rotterdam, the Netherlands: A. A. Balkema. p 41–47.
- Roman J, Struogo A. 1994. Echinoides du Libyen (Eocene inférieur) d'Égypte. *Rev Paleobiol* 13:29–57.
- Ronquist F, Huelsenbeck JP. 2003. MRBAYES 3: Bayesian phylogenetic inference under mixed models. *Bioinformatics* 19:1572–4.
- Sanderson MJ. 2002. Estimating absolute rates of molecular evolution and divergence times: a penalized likelihood approach. *Mol Biol Evol* 19:101–9.

- Sanderson MJ. 2004. r8s, version 1.70: user's manual. Section of evolution and ecology. Davis, CA: University of California.
- Schultz H. 2005. Sea urchins. Hemdingen, Germany: Heinke & Peter Schulz. p 484.
- Seilacher A. 1979. Constructional morphology of sand dollars. *Paleobiology* 5:191–221.
- Smith AB. 1981. Implications of lantern morphology for the phylogeny of post-Palaeozoic echinoids. *Palaeontology* 24: 779–801.
- Smith AB. 1988. Phylogenetic relationship, divergence times, and rates of molecular evolution for camarodont sea urchins. *Mol Biol Evol* 5:345–65.
- Smith AB. 2001. Optimizing phylogenetic analysis by the inclusion of fossils: cassiduloid paraphyly and the origin of clypeasteroid echinoids. *Paleobiology* 27:392–404.
- Smith AB. 2006. The echinoid directory. London: The Natural History Museum. [Electronic publication]. Available at: <http://www.nhm.ac.uk/research-curation/projects/echinoid-directory>. Accessed 24 Apr 2006.
- Smith AB, Anzalone L. 2000. *Loriolella*, a key taxon for understanding the early evolution of irregular echinoids. *Palaeontology* 43:303–24.
- Smith AB, Hollingworth NTJ. 1990. Tooth structure and phylogeny of the Upper Permian echinoid *Miocidaris keyserlingi*. *Proc Yorkshire Geol Soc* 48:47–60.
- Smith AB, Jeffery CH. 2000. Maastrichtian and Palaeocene echinoids: a key to world faunas. *Spec Pap Palaeontol* 63:1–406.
- Smith AB, Lafay B, Christen R. 1992. Comparative variation in morphological and molecular evolution through time: 28S ribosomal RNA versus morphology in echinoids. *Phil Trans R Soc Lond B* 338:365–82.
- Smith AB, Littlewood DTJ. 1997. Molecular and morphological evolution during the post-Palaeozoic diversification of echinoids. In: Givnish TJ, Systma KJ, editors. *Molecular evolution and adaptive radiation*. Cambridge: Cambridge University Press. p 559–83.
- Smith AB, Littlewood DTJ, Wray GA. 1996. Comparing patterns of evolution: larval and adult life-history stages and small ribosomal RNA of post-Palaeozoic echinoids. *Phil Trans R Soc Lond B* 349:11–8.
- Smith AB, Peterson KJ. 2002. Dating the time of origin of major clades: molecular clocks and the fossil record. *Annu Rev Earth Planet Sci* 30:65–88.
- Smith AB, Peterson KJ, Wray GA, Littlewood DTJ. 2004. From bilateral symmetry to pentaradiality. The phylogeny of hemichordates and echinoderms. In: Cracraft J, Donoghue MJ, editors. *Assembling the tree of life*. Oxford: Oxford University Press. p 365–83.
- Smith AB, Stockley B. 2005. Fasciole pathways in spatangoid echinoids: a new source of phylogenetically informative characters. *Zool J Linn Soc* 144:15–35.
- Smith AB, Wright CW. 1990. British Cretaceous echinoids. Part 2, Echinothurioida, Diadematoidea and Stirodonta (1, Calycina). *Palaeontogr Soc Monogr* 101–198, pls 33–72.
- Smith AB, Wright CW. 1996. British Cretaceous echinoids. Part 4, Stirodonta 3 (Phymosomatoidea 2) and Camarodonta. *Palaeontogr Soc Monogr* 268–341, pls 93–114.
- Stockley B, Smith AB, Littlewood DTJ, Lessios HA, MacKenzie-dodds JA. 2005. Phylogenetic relationships of spatangoid sea urchins (Echinoidea): taxon sampling density and congruence between morphological and molecular estimates. *Zool Scr* 34:447–68.
- Swofford DL. 2002. PAUP*. Phylogenetic analysis using parsimony (and other methods). Version 4. Sunderland, MA: Sinauer Associates.
- Takezaki N, Rzhetsky A, Nei M. 1995. Phylogenetic test of the molecular clock and linearized trees. *Mol Biol Evol* 12:823–33.
- Templeton A. 1983. Phylogenetic inference from restricted endonuclease cleavage site maps with particular reference to the evolution of humans and apes. *Evolution* 37:221–44.
- Thorne JL, Kishino H. 2002. Divergence time and evolutionary rate estimation with multilocus data. *Syst Biol* 51:689–702.
- Thorne JL, Kishino H, Painter IS. 1998. Estimating the rate of evolution of the rate of evolution. *Mol Biol Evol* 15: 1647–57.
- Vadet A. 1999. Revision des échinides de Saint Cassian et évolution des échinides post-carbonifères. *Mem Soc Acad Boulonnais Hist Nat* 20:1–59.
- Welch JJ, Fontanillas E, Bromham L. 2005. Molecular dates for the “Cambrian explosion”: the influence of prior assumptions. *Syst Biol* 54:672–8.
- Wiens JJ. 1998. Combining data sets with different phylogenetic histories. *Syst Biol* 47:568–81.
- Winchell CJ, Martin AP, Mallatt J. 2004. Phylogeny of elasmobranchs based on LSU and SSU ribosomal RNA genes. *Mol Phylogenet Evol* 31:214–24.
- Xia X, Xie Z, Kjer KM. 2003. 18S ribosomal RNA and tetrapod phylogeny. *Syst Biol* 52:283–95.
- Yang, Z. 1997. PAML: a program package for phylogenetic analysis by maximum likelihood. *Comput Appl Biosci* 13: 555–6.

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