Pancrustacean Phylogeny in the Light of New Phylogenomic Data: Support for Remipedia as the Possible Sister Group of Hexapoda

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Abstract

Remipedes are a small and enigmatic group of crustaceans, first described only 30 years ago. Analyses of both morphological and molecular data have recently suggested a close relationship between Remipedia and Hexapoda. If true, the remipedes occupy an important position in pancrustacean evolution and may be pivotal for understanding the evolutionary history of crustaceans and hexapods. However, it is important to test this hypothesis using new data and new types of analytical approaches. Here, we assembled a phylogenomic data set of 131 taxa, incorporating newly generated 454 expressed sequence tag (EST) data from six species of crustaceans, representing five lineages (Remipedia, Laevicaudata, Spinicaudata, Ostracoda, and Malacostraca). This data set includes all crustacean species for which EST data are available (46 species), and our largest alignment encompasses 866,479 amino acid positions and 1,886 genes. A series of phylogenomic analyses was performed to evaluate pancrustacean relationships. We significantly improved the quality of our data for predicting putative orthologous genes and for generating data subsets by matrix reduction procedures, thereby improving the signal to noise ratio in the data. Eight different data sets were constructed, representing various combinations of orthologous genes, data subsets, and taxa. Our results demonstrate that the different ways to compile an initial data set of core orthologs and the selection of data subsets by matrix reduction can have marked effects on the reconstructed phylogenetic trees. Nonetheless, all eight data sets strongly support Pancrustacea with Remipedia as the sister group to Hexapoda. This is the first time that a sister group relationship of Remipedia and Hexapoda has been inferred using a comprehensive phylogenomic data set that is based on EST data. We also show that selecting data subsets with increased overall signal can help to identify and prevent artifacts in phylogenetic analyses.

Key words: phylogenomics, EST, matrix reduction, orthology prediction, Crustacea, Remipedia.

Introduction

A monophyletic taxon Pancrustacea is supported by phylogenies that are based on mitochondrial, single nuclear gene, multigene, and large phylogenomic analyses (Friedrich and Tautz 1995, 2001; Shultz and Regier 2000; Giribet et al. 2001; Hwang et al. 2001; Regier and Shultz 2001; Nardi et al. 2003; Carapelli et al. 2005, 2007). These results all support the hypothesis that hexapods are more closely related to crustaceans than to myriapods and thus contradict the Atelocerata (a.k.a. Tracheata) hypothesis, which assumes a sister group relationship of hexapods and myriapods (Pocock 1893; Heymons 1901). If the Pancrustacea hypothesis (Zrzavý and Stys 1997) is accepted, it still remains unclear which among the major crustacean groups represents the sister group of Hexapoda. Many studies based on large molecular data sets have proposed Branchiopoda as the sister group of Hexapoda (Roeding et al. 2007, 2009; Dunn et al. 2008; Timmermans et al. 2008; Meusemann et al. 2010; Rota-Stabelli et al. 2011). However, these studies are characterized by a relatively poor sampling of crustacean taxa. A recent comprehensively sampled molecular phylogenetic analysis of arthropods instead suggests that hexapods are

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the sister group to a clade "Xenocarida," which comprises Remipedia and Cephalocarida (Regier et al. 2010). A close relationship between hexapods and Remipedia was previously suggested by a phylogenetic analysis of hemocyanin sequences (Ertas et al. 2009) as well as by several morphological studies (Moura and Christoffersen 1996; Fanenbruck et al. 2004; Fanenbruck and Harzsch 2005; Bäcker et al. 2008). By contrast, other morphological analyses inferred Remipedia and Malacostraca as being sister taxa (Koenemann et al. 2007, 2009).

To shed light on the higher level pancrustacean phylogeny and the closest crustacean relatives of hexapods, we performed a series of phylogenetic analyses on the most exhaustive crustacean phylogenomic data set derived from ESTs compiled to date. This includes 454 expressed sequence tag (EST) data from six hitherto unsampled crustacean species, namely *Lynceus brachyurus* (Laevicaudata, Branchiopoda), Spinicaudata sp. (Branchiopoda), Cypridininae sp. (Ostracoda), *Sarsinebalia urgorrii*, *Nebalia bipes* (Leptostraca, Malacostraca), and *Speleonectes* cf. *tulumensis* (Remipedia). Data from Cephalocarida, however, were not available for inclusion in this study, despite a tremendous effort and several field trips to collect sufficient specimens of this taxon.

Nonphylogenetic signal (Felsenstein 1988; Philippe et al. 2005, 2011) can seriously mislead phylogenomic studies. The greatest challenges are therefore to optimize the quality of the data, to separate signal from noise, and to handle efficiently missing data (Driskell et al. 2004; Philippe et al. 2005, 2011; Dunn et al. 2008; Hartmann and Vision 2008; Wiens and Moen 2008; Meusemann et al. 2010). Here, these issues are addressed by using the HaMStR approach (Hidden Markov Model based Search for Orthologs using Reciprocity) for orthology prediction (Ebersberger et al. 2009), automated alignment evaluation and masking (Misof B and Misof K 2009; Kück et al. 2010), and a recently developed approach to matrix reduction that selects optimal data subsets featuring increased signal (see Meusemann et al. 2010; Meyer and Misof 2010, http://mare.zfmk.de).

In summary, this study has three goals. 1) To address pancrustacean phylogeny with the largest phylogenomic data set derived from ESTs compiled to date, including data from hitherto unsampled key taxa, such as Remipedia. 2) To assess the likely sister group of Hexapoda based on phylogenomic EST data. 3) To evaluate the impact of matrix reduction procedure on inferred trees by selecting optimal data subsets derived from two different orthologous gene sets.

Materials and Methods

Molecular Techniques

454-pyrosequencing (ROCHE) was used to generate EST sequences from six crustacean species (see supplementary file 1, Supplementary Material online). Fresh tissue was preserved in RNAlater and stored at -20 or -80 °C. Total RNA of Cypridininae sp. (Ostracoda), Speleonectes cf. tulumensis (Remipedia), and S. urgorrii (Leptostraca) was extracted (Absolutely RNA kit; Stratagene) and its corresponding cDNA

synthesized (Mint kit; Evrogen) at the Max Planck Institute for Molecular Genetics, Berlin, Germany. Subsequently, cDNA fragments were size selected with the Chromaspin 1000 kit (Clontech), and the cDNA library was normalized with the Trimmer kit (Evrogen). cDNA was digested with the restriction enzyme SfiI (NEB). The digested cDNA was purified with the Qiagen polymerase chain reaction kit and subsequently ligated with 454 pyrosequencing adaptors (Roche). A total of 1,000,000 reads per species were sequenced on a Titanium FLX sequencer (Roche). Total RNA of N. bipes (Leptostraca), L. brachyurus (Laevicaudata), and Spinicaudata sp. (a new species that is currently being described by Nicolas Rabet, Université Pierre et Marie Curie, Paris) was extracted with the Qiagen RNAeasy Kit by R.A.J. at the University of Bath. Synthesis of cDNA, construction of nonnormalized cDNA libraries, 454-pyrosequencing (100,000-140,000 reads per species), and sequence assembly were performed at the GenePool genomics facility, University of Edinburgh, United Kingdom.

Sequence Processing, Orthology Prediction, and Alignment Masking

Vector sequences of the obtained reads were removed with CrossMatch (Green 1993-1996, 0.990329) using UNIVEC (build 5.2, December 2009; http://www.ncbi.nlm.nih.gov/ VecScreen/UniVec.html) after lowercase nucleotides were clipped with the aid of a custom-made PERL script written by Sascha Strauss (CIBIV, Vienna, Austria). Additionally, vector sequences and poly-A tails were removed with Seq-Clean (http://compbio.dfci.harvard.edu/tgi/software/) using UNIVEC (build 5.2, December 2009). Subsequently, sequences were masked with RepeatMasker (Smit et al. 1996-2010, open-3.1.6) and RepBase (20061006; http:// www.girinst.org/server/RepBase). Clustering and assembly were performed using MIRA 3.0.3 (Chevreux et al. 2004). EST sequences of other taxa (see supplementary file 1, Supplementary Material online) were retrieved from GenBank. All crustaceans, for which EST sequences are available (39 species), were added to our data set. The data set comprised a total of 46 crustaceans, 46 hexapods, 32 chelicerates, and 3 myriapods as well as 3 onychophorans and 1 polychaete (Capitella sp.) (see supplementary file 1, Supplementary Material online). Onychophorans and the polychaete were included as outgroup taxa. All EST sequences were quality checked and assembled in the processing pipeline described above. Assembled sequences of our own 454 projects were submitted to the Transcriptome Sequences Assembly database at the National Center for Biotechnology Information (NCBI) (accession numbers are summarized in supplementary file 1, Supplementary Material online).

Our strategies for orthology prediction and for alignment masking followed the procedures described in Meusemann et al. (2010). Two sets of orthologous genes were constructed using the InParanoid transitive closure approach described by Ebersberger et al. (2009). Selection of orthologs in these two sets was guided by protein sequences available in proteome data sets of the so-called

"primer taxa." Sequences of primer taxa were aligned within each set of orthologs and used to generate profile hidden Markov models (pHMMs). Subsequently, the pHMMs were used to search for putative orthologous sequences among the translated ESTs from all the taxa in our data set. Ortholog set 1 included the amino acid sequences of those genes for which the algorithm 4.1s from InParanoid (Ostlund et al. 2010) inferred orthologous sequences based on the following five primer taxa: Ixodes scapularis (Chelicerata), Daphnia pulex (Crustacea), Apis mellifera, Aedes aegypti (Hexapoda), and Capitella sp. (Polychaeta). Ortholog set 2 included genes for which InParanoid 7 inferred orthologous sequences based on the following six primer taxa: I. scapularis (Chelicerata), D. pulex (Crustacea), A. mellifera, Tribolium castaneum, Bombyx mori (Hexapoda), and Capitella sp. (Polychaeta). HaMStR then assigned ESTs to the core ortholog groups (Ebersberger et al. 2009) (options-representative, -strict, and -eval limit = 0.01). Each group of orthologous amino acid sequences was aligned separately with MAFFT L-INS-I (Katoh and Toh 2008). Randomly, similar aligned positions were identified with ALI-SCORE. We applied the default sliding window size, the maximal number of pairwise comparisons (-r), and a special scoring for gappy amino acid data (-e) (Misof B and Misof K 2009; Kück et al. 2010). Randomly aligned positions were subsequently removed with ALICUT (Kück 2009; http:// www.utilities.zfmk.de). All masked gene alignments were finally concatenated with FASconCAT (Kück and Meusemann 2010).

Orthology prediction resulted in two data sets: ortholog set 1 (hereafter, set 1A_{unred}) encompasses 1,886 genes and ortholog set 2 (set 2A_{unred}) contains 1,579 genes (see supplementary files 2-4, Supplementary Material online). Each set consists of 131 taxa. The reference species for the reciprocal Blast procedure are given in supplementary file 1 (Supplementary Material online). To generate additional data sets, the number of hexapod and chelicerate species was reduced. This a priori exclusion of taxa allowed the quartet mapping and subsequent gene selection procedures (see next paragraph) to preferentially retain genes that are proportionally more represented in crustaceans and eventually more informative for resolving relationships among crustaceans. In addition, we removed several hexapod and chelicerate taxa with long terminal branches in the trees inferred from set 1 and 2 (e.g., Glycophagus domesticus) in order to reduce the impact of possible long-branch attraction artifacts (see supplementary file 1, Supplementary Material online). This yielded the additional data sets 1B_{unred} and 2B_{unred}, each with 105 species.

We assessed the overlap of our putatively orthologous genes with those presented in Meusemann et al. (2010) and with the sequences analyzed by Regier et al. (2010). Of the data presented in Regier et al. (2010), all mRNA sequences for nine representatives of the major crustacean taxa present in and complementary to our data set (including Remipedia and Cephalocarida, supplementary file 5a, Supplementary Material online) were downloaded from NCBI (September 2010). Sequences of these nine crustaceans

were also analyzed with HaMStR (same settings as before) to search for orthologous genes that correspond to those in our data sets (supplementary file 5a, Supplementary Material online).

Matrix Reduction and Selection of Data Subsets

There are various strategies to handle highly incomplete matrices (i.e., data sets with a large proportion of missing entries or gaps). Most often, concatenated "supermatrices" are filtered using predefined thresholds of data availability (Dunn et al. 2008; Philippe et al. 2009). We utilized an alternative approach to data reduction here, selecting a subset of genes and taxa from each supermatrix based on information content in addition to data availability (MARE v 0.1.2-alpha; Meyer and Misof 2010, http://mare.zfmk.de). The approach yields a condensed and more informative data set by maximizing the ratio of signal to noise and by reducing the number of uninformative genes and poorly sampled taxa. MARE first evaluates the "tree likeness" of each single gene. Tree-likeness reflects the fraction of all possible (but not more than 20,000, due to computational limitations) quartets that are resolved for a given sequence alignment. The process is based on geometry-weighted quartet mapping (Nieselt-Struwe and von Haeseler 2001), extended to amino acid data. For further details on the procedure and the algorithm, see Meyer and Misof (2010; http://mare.zfmk.de).

Matrix reduction was performed on the four data sets $(1A_{unred}, 1B_{unred}, 2A_{unred}, and 2B_{unred})$ defined above (for an overview, see supplementary file 4, Supplementary Material online), using Acerentomon franzi (Protura, Hexapoda) and Balanus amphritite (Cirripedia, Crustacea) as constraint taxa and applying a taxon weighting parameter (-t) of 1.5 to keep more taxa. The constraints aim to maximize the retention of entognathous hexapods and cirripede crustacean taxa, respectively. The matrix reduction resulted in the reduced data sets $1A_{red}$, $1B_{red}$, $2A_{red}$, and $2B_{red}$.

Phylogenetic Analyses

Phylogenetic relationships were inferred by analyzing data sets 1A_{red}, 1B_{red}, 2A_{red}, and 2B_{red} under the maximum likelihood optimality criterion in RAxML v7.2.6 (Stamatakis 2006; Ott et al. 2007) (see table 1). Tree searching and bootstrapping were conducted simultaneously (PROTCATWAG, -f a, 1,000 bootstrap replicates). In all analyses, the "bootstopping" criterion (Pattengale et al. 2010) was used (default settings) a posteriori to assess whether or not a sufficient number of bootstrap replicates had been computed for evaluating tree robustness. Additionally, the unreduced data sets (i.e., 1A_{unred}, 1B_{unred}, 2A_{unred}, and 2B_{unred}) were analyzed using the same procedures, except that we used the "on-the-fly" bootstopping criterion (to avoid unnecessary computations and save computational resources) with the SSE-3-vectorized Pthreads-parallelized version 7.2.8 of RAxML. All analyses were done on the high performance computing clusters at the ZFMK Bonn (Zoologisches Forschungsmuseum Alexander Koenig, Bonn), the RRZK in Cologne (Regionales Rechenzentrum Köln:

Table 1. Comparison of the Unreduced and Reduced Data Sets and Resulting Support Values for Major Taxa in Both Approaches.

Data set (matrix)	Set 1				Set 2			
	Set 1A _{unred}	Reduced Set 1A _{red}	Set 1B _{unred}	Reduced Set 1B _{red}	Set 2A _{unred}	Reduced set 2A _{red}	Set 2B _{unred}	Reduced set 2B _{red}
Number of included genes	1,886	316	1,886	351	1,579	272	1,579	280
Alignment lengths (aa)	831,013	62,638	866,479	74,941	711,430	54,209	736,307	56,481
Information content	0.276	0.617	0.26	0.62	0.276	0.62	0.26	0.614
Clade support								
Malacostraca	100	100	100	100	100	100	100	100
(Leptostraca, Eumalacostraca)	100	100	99	100	100	100	100	77
Decapoda	99	100	99	100	99	100	100	100
(Eucarida, Decapoda)	65	99	61	_	62	100	58	100
Cirripedia	100	100	100	100	100	100	100	100
(Cirripedia, Malacostraca)	100	88	99	94	99	_	100	_
(Cirripedia, Copepoda)	_	_	_	_	_	96	_	94
Copepoda	100	100	100	100	100	100	100	100
(Ostracoda, Copepoda)			22		_	_	69	_
(Ostracoda, (remaining								
Pancrustacea))	_	100 (IS)	_	100 (IS)	_	_	_	100 (IS)
(Ostracoda, (Malacostraca,								
(Cirripedia, Copepoda)))	_	_	_	_	_	79 (HS)	_	_
(Ostracoda, (Malacostraca, Cirripedia))	3	_	_	_	6	_	_	_
Branchiopoda	100	100	100	100	100	100	100	100
(Branchiopoda, (Remipedia, Hexapoda))	100	78	100	43	_	83	100	100
(Remipedia, Hexapoda)	100	98	94	100	100	96	100	100
Hexapoda	100	99	100	100	100	96	100	100
Pancrustacea	100	100	100	100	100	100	99	100
Mandibulata	91	_	96	_	99	_	83	_

NOTE.—The numbers of taxa and genes, the alignment length, and information content of all constructed matrices for both ortholog sets are given. Selected major taxa in all resulting topologies are listed with statistical support (bootstrap values). Dashes indicate low clade support (under 50%). Leaf stability values above 95% represent highly stable taxa. HS denotes high stability; IS, an instable position; see figures.

SUGI—Sustainable Grid Infrastructure project—and CHEOPS—Cologne High Efficient Operating Platform for Science), and the HITS gGmbH in Heidelberg (Heidelberg Institute for Theoretical Studies). Leaf stability indices were computed with Phyutility (Smith and Dunn 2008) on the respective sets of bootstrap trees from each data set. The indices are a measure for the consistency of the position of each terminal taxon (leaf) relative to remaining taxa across replicates. Potentially unstable positions of taxa can be detected in the reconstructed topologies using this method. The lower the leaf stability index for a given terminal taxon is, the less stable is its phylogenetic position.

Results

Sets of Orthologous Genes

Set 1A_{unred} of orthologous genes comprises sequences of 131 taxa, 1,886 genes, and 831,013 aligned amino acid positions (supplementary files 2 and 6, Supplementary Material online). Set 2A_{unred} includes sequences of 131 taxa, 1,579 genes, and 711,430 aligned amino acid positions (see supplementary files 3 and 7, Supplementary Material online). The two sets have 1,410 genes in common (see supplementary files 2, 3, and 5, Supplementary Material online). After applying MARE, the information content in each data subset was approximately doubled (see table 1). MARE removed nearly the same species from each data set such that the two a priori reduced data sets (1A_{red} and 2A_{red}) had very similar taxon samples (supplementary

file 1, Supplementary Material online). Four hundred and ninety six of these genes are present in the unreduced data set analyzed by Meusemann et al. (2010). Of the 129 genes present in the reduced data set (selected optimal data subset) of Meusemann et al. (2010), 75 were found in the reduced data sets 1A_{red} and 2A_{red} and 74 genes in the reduced data sets 1B_{red} and 2B_{red} (see supplementary file 5, Supplementary Material online).

Of the sequences of Regier et al. (2010), 42 sequences were assigned to our groups of orthologous sequences in data set 1A_{unred} and 37 to our groups of orthologous sequences in data set 2A_{unred}. However, only 19 sequences of Remipedia and Cephalocarida overlap with set 1A_{unred}, and 18 overlap with set 2A_{unred}. Only four remipede and cephalocarid genes used in Regier et al. 2010 were present in our data sets 1A_{red} and 2A_{red}; five genes were shared with our data sets 1B_{red} and 2B_{red} (see supplementary file 5, Supplementary Material online).

Pancrustacean Relationships in the Trees Inferred from Reduced Data Sets

The monophyly of Pancrustacea received 99–100% bootstrap support (BS) in all of our trees. Likewise, many major crustacean groups (i.e., Malacostraca, Branchiopoda, Copepoda, and Cirripedia) have high (BS = 99%) or maximal support in all trees (see table 1).

Two large clades are found in the trees inferred from the reduced data sets: A clade composed of Malacostraca, Cirripedia, and Copepoda and another comprising Branchiopoda, Remipedia, and Hexapoda. Support for the first clade is much higher in the trees that we derived from the submatrices of set 2 (fig. 2: data set 2A_{red} and fig. 4: data set $2B_{red}$; BS = 75% and 100%, respectively) than in the trees derived from the submatrices of set 1. However, the relationships of the major lineages within this clade (i.e., Malacostraca, Cirripedia, and Copepoda) differ between the trees inferred from submatrices of sets 1 and 2. The reduced data sets of set 1 suggest a sister group relationship of cirripedes and malacostracans (fig. 1: data set 1A_{red} and fig. 3: 1B_{red}). In contrast, the reduced data sets of set 2 imply that cirripedes and copepods are sister groups (fig. 2: data sets 2A_{red} and fig. 4: 2B_{red}). Similarly, the clade comprising branchiopods, remipedes, and hexapods receives stronger support in the trees (83% and 100%) that were inferred from submatrices of set 2. Trees based upon set 2 also show higher average leaf stability indices (see figs. 1-4) than those based upon set 1. Importantly, all trees inferred from the reduced data sets support the relationship (Branchiopoda (Remipedia, Hexapoda)). Data sets 1A_{red}, 1B_{red}, and 2B_{red} maximally support a clade (Ostracoda, remaining pancrustaceans), whereas data set 2A_{red} suggests a clade (Ostracoda ((Malacostraca (Copepoda, Cirripedia)))) (BS = 79%).

Inferred phylogenetic relationships within the monophyletic higher level crustacean taxa are consistent between our data sets. Within Malacostraca, both the unreduced and the reduced data sets suggest a sister group relationship of Leptostraca and Eumalacostraca. Eucarida (Euphausiacea, Decapoda) are supported in three of the four reduced trees (figs. 1, 2, and 4). In the fourth tree (fig. 3), Euphausia superba (Euphausiacea) was not present because this taxon had been excluded from the data set during matrix reduction. In all inferred trees, Eucarida and Peracarida (represented by Amphipoda) are sister taxa. All trees, except one of the unreduced trees (supplementary file 6, Supplementary Material online, data set 1Aunred), support the same phylogenetic relationships within Decapoda. Decapoda is divided into two sister clades. The first unites Caridea and Dendrobranchiata as sister taxa. The second clade supports the relationships ((Anomura, Brachyura) (Astacidea, Achelata)). Within branchiopods, all our analyses suggest the same topology: (Anostraca (Notostraca (Laevicaudata (Spinicaudata, Cladocera)))). Finally, in Hexapoda, a split between Ectognatha and Entognatha (Insecta) is recovered consistently. Within Entognatha, Collembola is inferred invariably as the sister group to Protura (together constituting the clade Ellipura).

Comparison of Trees Inferred from Unreduced and Reduced Data Sets

All of the trees inferred from the unreduced data sets suggest consistently the monophyly of Mandibulata (BS =83–99%). In all of these, Myriapoda is the sister group of Pancrustacea. In contrast, Mandibulata is not supported by any of the trees derived from the reduced data sets. Rather, a clade (Chelicerata, Pancrustacea) is recovered with weak to maximal support (BS =51–100%). Within Pancrustacea, the

trees based upon the unreduced data sets strongly support a clade of cirripedes and malacostracans (BS = 99–100%). The same relationships are obtained when analyzing the reduced subsets of set 1 (i.e., data sets $1A_{\rm red}$ and $1B_{\rm red}$). Three of four phylogenetic trees inferred from the reduced data sets show a sister group relationship of Ostracoda plus the remaining pancrustaceans (figs. 1, 3, and 4), although the precise phylogenetic position of Ostracoda remains uncertain.

Discussion

Pancrustaceans

The monophyly of Pancrustacea (Zrzavý and Stys 1997) has been suggested by several studies that investigated nuclear and/or mitochondrial sequences (Friedrich and Tautz 1995, 2001; Shultz and Regier 2000; Giribet et al. 2001; Hwang et al. 2001; Regier and Shultz 2001; Nardi et al. 2003; Carapelli et al. 2005, 2007;). This clade, sometimes also referred to as Tetraconata (Dohle 2001), has also been advocated because of conspicuous similarities in the ommatidia of the compound eyes shared between hexapods and crustaceans (but see also Harzsch and Hafner 2006) and because of similarities in their neuroanatomy and neuroembryology (Harzsch et al. 2005; Harzsch 2006; Ungerer and Scholtz 2008). Recent phylogenomic analyses (Roeding et al. 2009; Meusemann et al. 2010; Rota-Stabelli et al. 2011) also strongly support the monophyly of Pancrustacea. Our results corroborate strongly a clade Pancrustacea, which is maximally or highly supported in all trees inferred from of our data sets.

Malacostraca

Malacostraca was consistently recovered as a clade in our analyses. Nonetheless, the phylogenetic relationships of the major lineages within Malacostraca as well as the phylogenetic position of the Malacostraca within the Pancrustacea are still unclear (Jenner 2010). Our data support a split of the Malacostraca into the lineages Leptostraca and Eumalacostraca. This is consistent with morphological data (Wills et al. 1995, 2009; Wills 1998a, 1998b; Richter and Scholtz 2001; Jenner et al. 2009). Although our phylogenomic data are unable to completely resolve the relationships within Eumalacostraca, they do suggest a common origin of Anomura, Brachyura, Astacidea, and Achelata by exclusion of Dendrobranchiata and Caridea. This last result is largely consistent with recently published molecular phylogenetic investigations including these taxa (Bracken et al. 2009, 2010; Toon et al. 2009). Most contentious of all is the position of the Malacostraca within crustaceans (Jenner 2010). Even if we only focus on recently published molecular phylogenetic (von Reumont et al. 2009; Koenemann et al. 2010; Regier et al. 2010) and phylogenomic studies with reasonable sampling of crustacean taxa (Meusemann et al. 2010; Andrew 2011), no consistent pattern emerges. Our current results support two alternative sister groups for Malacostraca: Cirripedia (representing Thecostraca) or (Cirripedia, Copepoda). In Meusemann et al.

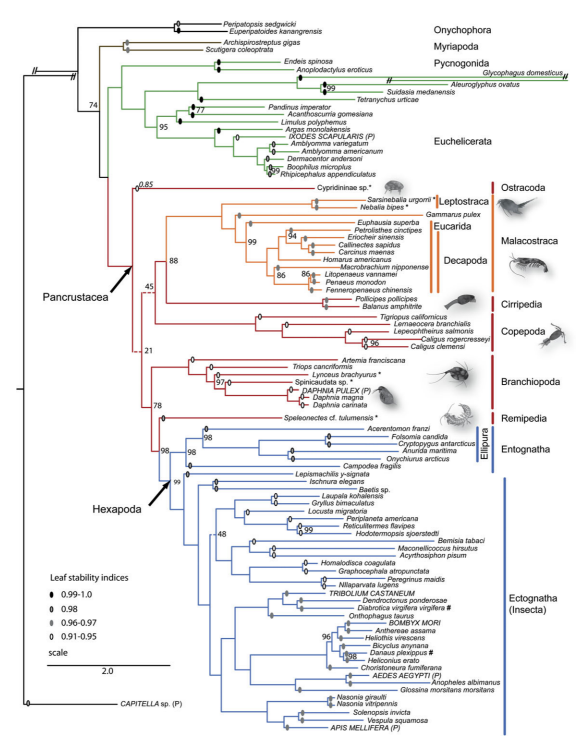


Fig. 1. Phylogram derived from data matrix 1A_{red} (91 taxa, 316 genes) in RaxML Topology inferred from set 1A_{red} in RAxML (PROTCATWAG, 1,000 BS replicates, -f a). Bootstrap values are given only for nodes that lack maximum support. Ellipses on the branches (as explained in the bottom left corner) summarize the leaf stability values calculated with Phyutility (Smith and Dunn 2008), the value for the highly unstable Ostracoda is shown in italic for this branch. One thousand sampled bootstrap trees converged after 50 replicates applying the a posteriori bootstop function (Pattengale et al. 2010). Color code: crustaceans red and orange, hexapods blue, chelicerates green, myriapods brown, and outgroup taxa black. Species that are marked by an asterisk (*) are newly sequenced in this study, species marked by a pound sign (#) are only present in the respective data set, species written in CAPITALS represent proteome taxa, a (P) indicates the used primer taxa.

(2010), these two alternatives were inferred from the same data set using Bayesian and likelihood methods, respectively. In our study, the results of six of eight analyses support (Malacostraca, Cirripedia), with only the reduced data sets based on ortholog set 2 supporting (Malacostraca

(Cirripedia, Copepoda)). Since matrix reduction is shown to increase the signal to noise ratio (table 1), we speculate that the clade (Malacostraca, Cirripedia), which was also found by Regier et al. (2010) and Andrew (2011), might be an artifact, a hypothesis at least consistent with the

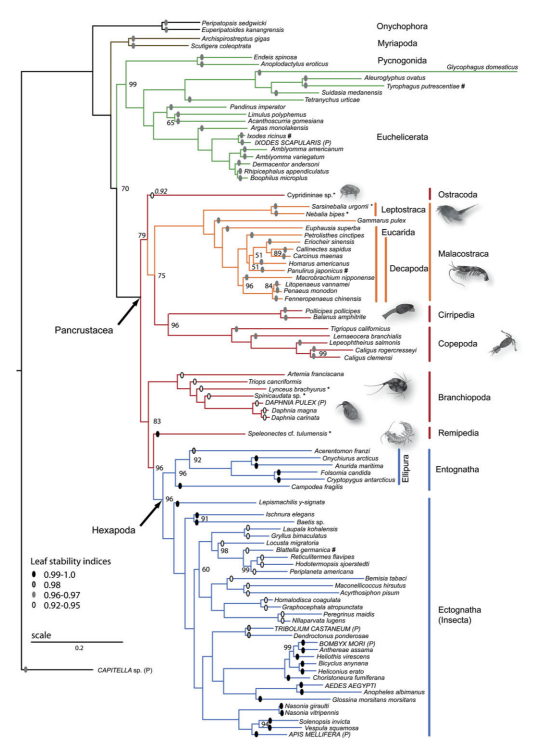


Fig. 2. RAxML topology derived from data matrix 2A_{red} (92 taxa, 272 genes). Topology inferred from set 2A_{red} in RAxML (PROTCATWAG, 1,000 BS replicates, -f a). Taxa are represented with the same colors as described in figure 1. Bootstrap values are given only for nodes that lack maximal support. Thousand sampled bootstrap trees converged after 50 replicates. For color codes and Phyutility usage, see figure 1.

slight drop in support value for this clade in the reduced data sets based on ortholog set 1. More importantly perhaps, support for this clade was also significantly reduced in the analysis of Regier et al. (2010) that was corrected for heterogeneity in base composition. A closer affinity of copepods and cirripedes would also be more congruent with some analyses of morphological data (e.g., Wills 1998a, 1998b; Martin and Davis 2001).

Branchiopoda

Our results strongly support monophyly of Branchiopoda, in line with earlier molecular and morphological studies (Wills 1998a, 1998b; Stenderup et al. 2006; Olesen 2007; Richter et al. 2007). Furthermore, we found the conchostracans to be paraphyletic, in agreement with recent studies (Braband et al. 2002; Olesen 2007; Richter et al. 2007). Unfortunately, there is still no agreement on the position of

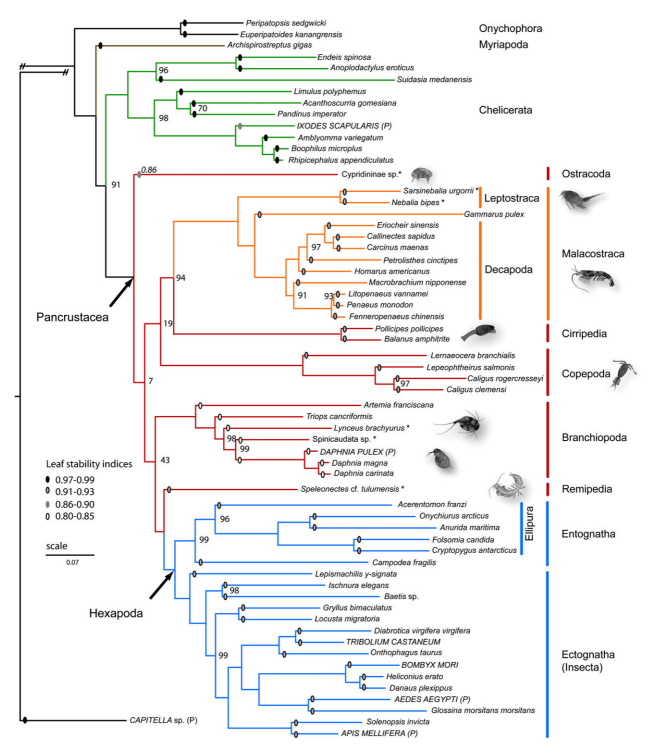


Fig. 3. RAxML topology derived from data matrix 1B_{red} with a priori taxa exclusion (62 taxa, 351 genes). Topology inferred from set 1B_{red} in RAxML (PROTCATWAG, 1,000 BS replicates, -f a). Taxa are colored as described in figure 1. Bootstrap values are given only for nodes that lack maximal support. Thousand sampled bootstrap trees converged after 50 replicates. For color codes and Phyutility usage, see figure 1.

Branchiopoda within the crustaceans. In terms of the number of recently proposed alternative hypotheses, the placement of Branchiopoda remains one of the most intriguing challenges in higher level pancrustacean phylogenetics (Jenner 2010). One recent well-supported hypothesis that has attracted considerable interest is the possible sister group relationship of branchiopods and hexapods (Glenner et al. 2006; Roeding et al. 2009; Meusemann et al. 2010;

Andrew 2011; Rota-Stabelli et al. 2011). Indeed, this hypothesis underpins a seductive scenario, in which hexapods are conjectured to have evolved from marine ancestors via some Late Silurian, freshwater, branchiopod-like intermediate (Glenner et al. 2006). However, if the marine fossil Rehbachiella kinnekullensis (Walossek 1993) represents a stem group branchiopod (Schram and Koenemann 2001), then branchiopods themselves are also likely to be ancestrally

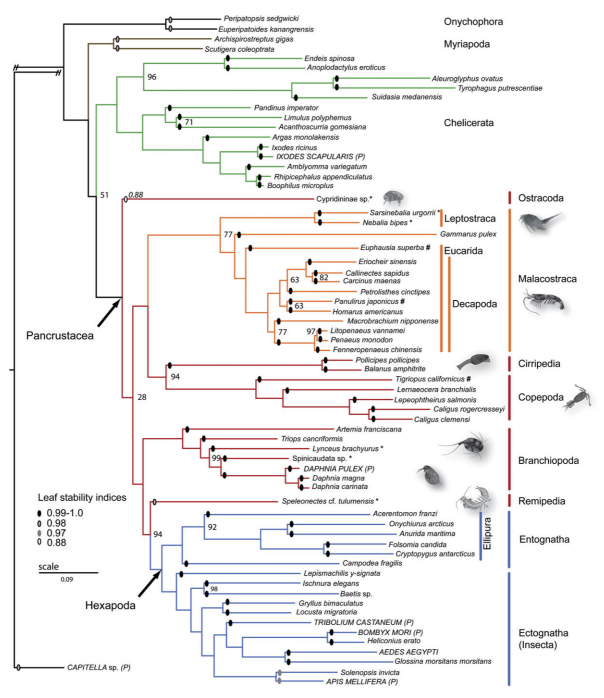


Fig. 4. RAXML topology derived from data matrix 2B_{red} (67 taxa, 280 genes). Topology inferred from set 2B_{red} in RAXML (PROTCATWAG, 1,000 BS replicates, -f a). Taxa are colored as seen in figure 1. Bootstrap values are given only for nodes that lack maximal support. Thousand sampled bootstrap trees converged after 100 replicates. For color codes and Phyutility usage, see figure 1.

marine (Olesen 2007), contrary to Glenner et al. (2006): see also figure 5.

Importantly, no previous phylogenomic analyses of EST data have included the enigmatic remipedes. Our new EST data strongly suggest that Branchiopoda is the sister group of Remipedia plus Hexapoda (with the single exception of our unreduced set 2A_{unred}, supplementary file 7, Supplementary Material online). Our data thus challenge the monophyly of Vericrustacea (= (Branchiopoda (Copepoda (Malacostraca, Thecostraca)))) found by Regier et al. (2010).

The conflict between molecular and morphological data regarding the evolutionary history of Branchiopoda, Malacostraca, and Remipedia is illustrated in figure 5. Our data, in common with most molecular studies (Regier et al. 2005, 2008; Mallatt and Giribet 2006; Roeding et al. 2009; von Reumont et al. 2009; Meusemann et al. 2010; Andrew 2011; Rota-Stabelli et al. 2011), imply that Branchiopoda is more closely related to Hexapoda and Remipedia than is Malacostraca. In conflict with these molecular results are morphological and neuroanatomical studies that support

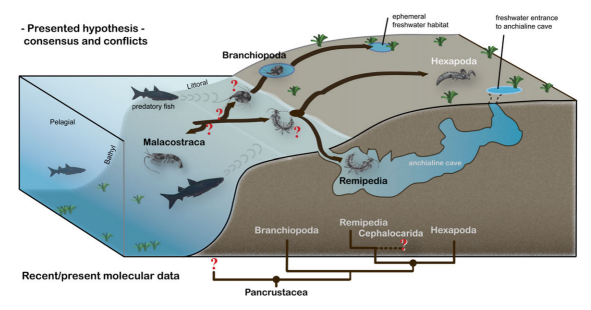


Fig. 5. Schematic illustrating the proposed evolutionary scenario highlighting conflicts between morphological and molecular data of pancrustaceans. Brown arrows and lines represent evolutionary lineages. The impact of predatory fishes as a possible evolutionary driver is illustrated by the gray waves. Circles represent nodes that are strongly supported by morphological and molecular data. Dashed lines indicate more weakly supported relationships. Red question marks indicate branches whose position is uncertain: variously because of ambiguity in the molecular data, conflict with morphological data, or a large gap in the fossil record. Molecular and morphological evidence suggest conflicting positions for Branchiopoda and Malacostraca. Molecular analyses generally place Branchiopoda closer to Hexapoda, whereas selected morphological, neuroanatomical, and larval development data suggest a closer relationship of Malacostraca to Remipedia and Hexapoda. The figure illustrates the close relationship of Remipedia and Hexapoda, which is strongly supported by the present study.

a clade of Malacostraca, Remipedia, and Hexapoda (Fanenbruck et al. 2004; Fanenbruck and Harzsch 2005).

Is Remipedia the Sister Group to Hexapoda?

Remipedes have been considered crucial for understanding the origin of crustaceans ever since they were first described in the 1980s (Yager 1981; Yager and Schram 1986). Their homonymous trunks and the presence of a pair of biramous appendages on each segment have usually been interpreted as crustacean plesiomorphies (Schram 1986; Emerson and Schram 1991; Schram and Hof 1998; Ax 1999; Wheeler et al. 2004). However, new and substantially more comprehensive molecular, morphological, neuroanatomical, and developmental data have started to challenge the idea that remipedes diverged early during crustacean evolution. Similarities in neuroanatomy suggest a close relationship of remipedes, malacostracans, cephalocarids, and hexapods, which has been used to argue for a less basal position of remipedes. These taxa possess highly complex brains with a markedly different construction from those of other crustaceans (Fanenbruck et al. 2004; Fanenbruck and Harzsch 2005). Intriguingly, remipede larvae show many similarities with those of some malacostracans (Koenemann et al. 2007, 2009).

Until recently, molecular phylogenetic analyses provided evidence for conflicting hypotheses with respect to the position of remipedes within pancrustaceans (see Jenner 2010). For example, mitochondrial and nuclear ribosomal RNA sequences suggested a sister group relationship of remipedes to cirripedes (Carapelli 2000; Lavrov et al. 2004; Hassanin 2006; Lim and Hwang 2006), to ostracods (Cook

et al. 2005), to collembolans (Cook et al. 2005; Hassanin 2006), to diplurans (Carapelli et al. 2007), and to various "maxillopodan" taxa (Lavrov et al. 2004; von Reumont et al. 2009). The set of possible crustacean sister groups proposed for hexapods has been equally diverse, including branchiopods (Babbit and Patel 2005; Glenner et al. 2006; Roeding et al. 2009; Meusemann et al. 2010; Andrew 2011), malacostracans (Lim and Hwang 2006; Strausfeld et al. 2009), and copepods (Mallatt and Giribet 2006; Dell'Ampio et al. 2009; von Reumont et al. 2009). However, the taxonomic coverage in these studies was often sparse and usually did not include remipedes.

In ribosomal DNA (rDNA)-based phylogenies, Remipedia and Cephalocarida show long branches, and at least the cephalocarids are affected by nonstationary substitution processes (von Reumont et al. 2009). Spears and Abele (1998) interpreted a sister group relationship of Cephalocarida and Remipedia inferred from 18S rDNA sequence data with caution and suggested the possibility of pseudogenes in addition to nonstationary substitution processes. The putative sister group relationship of these two taxa must therefore be regarded with caution (von Reumont et al. 2009; Koenemann et al. 2010). Nonetheless, remipedes and cephalocarids have also emerged as close relatives from analyses of nuclear coding genes (Shultz and Regier 2000; Regier et al. 2005, 2008) but mostly without strong support. However, in the study by Regier et al. (2010), support for this clade was higher when models were applied that exclude the degenerated third codon positions on nucleotide level. Yet, support was again weak when studying the phylogenetic relationships at the amino acid level. Testing this hypothesis by means of analyzing exhaustive phylogenomic data must await the generation of EST data for cephalocarids.

Ertas et al. (2009) provided the first molecular phylogenetic support for a close relationship of remipedes and hexapods. This result was soon corroborated by a multigene analysis at the nucleotide level by Regier et al. (2010), which recovered the clade Xenocarida = (Remipedia, Cephalocarida) as a sister group of Hexapoda. Our results provide strong support for a close relationship of remipedes and hexapods and on the basis of significantly more nuclear protein-coding genes than analyzed before. Given the minimal overlap between our data and those by Regier et al., our results offer a largely independent test of this hypothesis. We therefore propose an evolutionary scenario, in which the last common ancestor of remipedes and hexapods lived in a shallow marine environment, from which crown group remipedes and hexapods colonized their respective anchialine and terrestrial habitats (fig. 5).

Impact of Ortholog Sets and Matrix Reduction

This study shows that the size and precise composition of phylogenomic data sets can have marked effects on the results of phylogenetic inference. Large size alone clearly does not make a data set reliable (Philippe et al. 2011). However, understanding the relative contributions of the size and composition of data sets on the results requires more studies in the future. Using the HaMStR approach, the set of orthologous genes selected for analysis (both the total number and identity) is strongly dependent upon the choice of primer taxa (supplementary files 2, 3, and 5, Supplementary Material online). Of course one expects a smaller set or orthologs when using more primer taxa. The exclusion of the dipteran Aedes and inclusion of Tribolium and Bombyx consequently result in a smaller number of orthologous genes in data sets derived from ortholog set 2 (see supplementary file 4 and 5, Supplementary Material online). The percentage of present genes that overlap between the two ortholog sets is significantly higher in data sets derived from ortholog set 2 (90% 2A_{unred}, 89% 2A_{red}, and 92% 2B_{red},) compared with data sets from ortholog set 1 (75% $1A_{unred}$, 77% $1A_{red}$, and 73% $1B_{red}$): see table 2. Overlapping genes between the unreduced and reduced data sets within ortholog set 1 and 2 is nearly identical, see table 2 and supplementary file 5b (Supplementary Material online).

Nonetheless, it remains difficult to determine which ortholog set should be considered as the most "reliable." Not only the contribution of each gene to the inferred relationships is unknown, the interactions of signals present in all genes also remain wholly unexplored. The software MARE attempts to address the first of these issues by excluding genes with low tree-likeness in order to reduce noise. However, more studies are needed to fully explore the efficiency and performance of this approach. For example, the clade Mandibulata is replaced with a clade Chelicerata + Pancrustacea in the topologies of the reduced data sets. This could conceivably be an artifact of matrix reduction. During the random substitution process, one expects that older phylogenetic signal is more likely to be

Table 2. Comparison of Gene Overlap and Exclusive Gene Occurrence in the Data Sets.

Come Occasion of Data Cota	Numbers	Percentage	C (0/)
Gene Overlap of Data Sets	of Genes	of Genes (%)	Sum (%)
Ortholog set 1: set 1A _{unred}	Total: 1,886		
Set 1 specific only	442	23	_
Set 1 and SOS	34	2	∑ 25
Set 1, set 2, and SOS	496	26	
Set 1 and set 2 only	914	49	∑ 75
Ortholog set 2: set 2A _{unred}	Total: 1,579		
Set 2 specific only	131	8	
Set 2 and SOS	38	2	∑ 10
Set 2, set 1, and SOS	496	32	
Set 2 and set 1 only	914	58	∑ 90
Ortholog set 1: set 1A _{red}	Total: 316		
Set 1 specific only	68	21	
Set 1 and SOS	6	2	∑ 23
Set 1, set 2, and SOS	65	21	
Set 1 and set 2 only	177	56	\sum 77
Ortholog set 2: set 2A _{red}	Total: 272		
Set 2 specific only	26	10	
Set 2 and SOS	4	1	\sum 11
Set 2, set 1, and SOS	65	24	
Set 2 and set 1 only	177	65	∑ 89
Ortholog set 1: set 1B _{red}	Total: 351		
Set 1 specific only	90	26	
Set 1 and SOS	4	1	∑ 27
Set 1, set 2, and SOS	66	19	_
Set 1 and set 2 only	191	54	\sum 73
Ortholog set 2: set 2B _{red}	Total: 280		_
Set 2 specific only	19	7	
Set 2 and SOS	4	1	∑ 8
Set 2, set 1, and SOS	66	24	_
Set 2 and set 1 only	191	68	\sum 92

NOTE.—The total numbers and the percentage of genes that are found in each data set derived from the two ortholog sets are given. Overlapping genes and exclusively represented genes for each data set are highlighted. Additionally, the overlap with the reduced data set (SOS) from Meusemann et al. (2010) with each of our data sets is included. The sum-column shows the percentages of genes unique to each particular data set, and those shared with the corresponding data set derived from the other ortholog set (for a graphical comparison, see supplementary file 5b, Supplementary Material online).

substituted by multiple hits (noise) than younger phylogenetic signal. Since MARE excludes genes that have lower tree-likeness scores, it could be that it disproportionally removes genes that contain older and distorted phylogenetic signal. This could lead to a loss of support for deeper nodes in the tree. However, because MARE does not distinguish between such secondarily noisy genes and pure noise, the potential loss of some phylogenetic signal is an inescapable side effect of trying to increase the overall signal to noise ratio of the data.

An important methodological issue may be illustrated by considering the variable placement of Cirripedia. Data sets based on set 1 support a clade Cirripedia and Malacostraca, independent of matrix reduction (albeit with decreased support in the reduced data sets). In contrast, when data sets based on set 2 are reduced with MARE (sets 2A_{red} and 2B_{red}), Cirripedia are inferred as the sister group to Copepoda (figs. 2 and 4). The latter hypothesis is in line with results from morphological and several molecular analyses (see Martin and Davis 2001; Jenner 2010). This indicates that some genes that are found exclusively in both reduced

matrices of set 1 (supplementary file 5, Supplementary Material online) apparently obscure the signal for a clade (Cirripedia, Copepoda). Interestingly, the clade (Cirripedia, Malacostraca) collapses in the study by Regier et al. (2010) when these authors tried to reduce the effects of sequence saturation corroborating the suggestion that conflicting signal is present in some genes.

Conclusions

- 1) This is the first phylogenomic analysis (including new EST data), which supports a sister group relationship of Remipedia and Hexapoda (Fanenbruck et al. 2004; Fanenbruck and Harzsch 2005; Ertas et al. 2009). This particular conclusion is unaffected by the precise procedures used for identifying orthologous genes or for reducing the data sets.
- 2) Our results suggest that Pancrustacea is divided into two clades: i) Malacostraca, Copepoda, and Cirripedia and ii) Branchiopoda, Remipedia, and Hexapoda.
- 3) The methods used for selection of putative orthologous genes, namely the primer taxa choice for the HaMStR approach and matrix reduction by selecting optimal data subsets can markedly influence the inferred relationships. For example, matrix reduction indicates that the clade Communostraca (Malacostraca, Thecostraca), with Cirripedia representing Thecostraca in our study that was supported by Regier et al. (2010) and by the phylogenomic analysis of Andrew (2011) might be artificial. This underlines the importance of implementing the most appropriate methods for compiling data sets and for controlling their quality.
- 4) By increasing the information content of data sets via matrix reduction, some conflicts in the data become visible and can be removed like (Malacostraca, Cirripedia). However, this study serves in parallel as a test case to explore the idea that MARE might introduce potential artifacts such as a collapse of Mandibulata in the reduced topologies.
- 5) High-level pancrustacean phylogeny remains a challenging area of research. The recent study by Regier et al. (2010) sampled significantly more genes and taxa than its forebears and represented a major advance. In view of the limited overlap between the genes included in that study and ours, our results allow an ostensibly independent test of some of the more surprising relationships reported by Regier et al. (2010). Future work should aim to incorporate hitherto unsampled taxa in phylogenomic data sets, most notably Cephalocarida.
- 6) An alternative approach to the one employed here is to assemble genomic data for more pancrustacean taxa to infer more pancrustacean-typical putative orthologous genes that might carry a less noisy signal. Critically, the prediction of orthologous genes could then be based on a larger sample of completely sequenced genomes. HaMStR could represent one possible strategy to identify the ortholog genes. In an additional second step after the HaMStR approach, gene subsets could be selected with MARE targeting in general only those genes that show a high tree-likeness and chance to be informative. Subsequently, the sequences of the identified genes can be used to reconstruct primer toolboxes to amplify genes

for specific taxa groups. This method will allow us additionally to include species that can be collected for DNA-based work, but which are difficult to collect fresh, and in sufficient quantity for mRNA-based EST sequencing.

Supplementary Material

Supplementary files 1-7 are available at Molecular Biology and Evolution online (http://www.mbe.oxfordjournals.org/).

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