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# Decapod Crustacean Phylogenetics 

edited by

Joel W. Martin, Keith A. Crandall, and Darryl L. Felder

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Edited by<br>Joel W. Martin<br>Natural History Museum of L. A. County<br>Los Angeles, California, U. S. A.

Keith A. Crandall<br>Brigham Young University<br>Provo, Utah, U. S. A.

Darryl L. Felder<br>University of Louisiana<br>Lafayette, Louisiana, U. S. A.

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# Decapod Phylogenetics and Molecular Evolution 

ALICIA TOON, MAEGAN FINLEY, JEFFREY STAPLES \& KEITH A. CRANDALL<br>Department of Biology, Brigham Young University, Provo, Utah, U.S.A.


#### Abstract

Decapoda is the most species-rich group of crustaceans, with numerous economically important and morphologically diverse species leading to a large amount of research. Our research groups are attempting to estimate a robust phylogeny of the Decapoda based on molecular and morphological data to resolve the relationships among the major decapod lineages and then to test a variety of hypotheses associated with the diversity of decapod morphological evolution. Thus, we have developed a database of molecular markers for use at different scales of the evolutionary spectrum in decapod crustaceans. We present potential mitochondrial and nuclear markers with an estimation of variation at the genus level, family level, and among infraorders for Decapoda. We provide a methodological framework for molecular studies of decapod crustaceans that is useful at different taxonomic levels.


## 1 MOLECULAR TAXONOMY

There are several competing hypotheses concerning the relationships of the major lineages of Decapoda based on morphological estimates of phylogeny. Early taxonomy of the decapods was largely based on the mode of locomotion; taxa were divided into the swimming lineages (Natantia) and the crawling lineages (Reptantia) (Boas 1880). Morphological and molecular studies suggest Natantia is paraphyletic; it is presently classified based on gill structure (Burkenroad 1963, 1981) dividing Decapoda into the suborders Dendrobranchiata (penaeoid and sergestoid shrimps) and Pleocyemata (all other decapod crustaceans). Relationships within Pleocyemata are still controversial and remain unresolved. As morphological data, both recent and fossil, and genetic data continue to accumulate, we are moving towards phylogenetic resolution of these controversial relationships. Here we present a progress report for the Decapoda Tree of Life effort and the tools with which we will continue our analysis of decapod crustacean phylogenetic relationships.

Several recent hypotheses based on combined analysis of morphological and molecular data or molecular data alone suggest that resolving the systematics of this group is a difficult task (see Fig. 1). There is agreement among these studies that Dendrobranchiata represents a basal lineage within the decapod crustaceans and that within Pleocyemata the Caridea and Stenopodidea are basal infraorders (Porter et al. 2005; Tsang et al. 2008). Molecular research also supports the removal of polychelids from Palinura following Scholtz and Richter (1995) and its establishment as a separate infraorder (Polychelida) (Tsang et al. 2008; Ahyong this volume). Relationships among reptant decapods remain unresolved by the addition of molecular data. Several recent phylogenetic analyses incorporating mitochondrial and nuclear data (Robles et al. this volume) or nuclear data alone (Tsang et al. 2008; Chu et al. this volume) suggest Thalassinidea are not monophyletic but rather may represent several infraorders. The timeline of diversification among the reptant decapods or specifically whether Astacidea (Porter et al. 2005) or the Anomura/Brachyura lineages (Ahyong \& O'Meally 2004; Tsang et al. 2008) are the most recently derived lineages remains a question of interest.


Crandall et al. 2000


Ahyong \& O'Meally 2004


Tsang et al. 2008

Figure 1. Hypotheses of decapod evolutionary relationships based on molecular data. R shows the position of the reptant decapods.

## 2 DEVELOPING GENETIC MARKERS FOR MOLECULAR PHYLOGENY

The order Decapoda includes roughly 175 families (extant and extinct) and more than 15,000 described species. Complicating things further are the estimated 437 million years since the origin of the Decapoda with the major lineages estimated to have been established by 325 million years ago (Porter et al. 2007). Constructing a molecular phylogeny across such breadth of taxa and depth of timescale requires serious consideration of markers that have enough variation to reconstruct relationships at the fine scale (at and within the family level) as well as being conservative enough to be used across infraorders representing these deeper timescales. Our approach is to accumulate molecular sequence data for different gene regions including both mitochondrial and nuclear genes, coding and non-coding. In this way, we will be able to maximize data at deeper nodes where alignment of sequence data is most difficult while retaining information among families and between the most recently diverged taxa.

There are two molecular approaches to amplifying sequence data for use in phylogenetic studies. (1) Isolation of RNA from tissues, coupled with reverse transcription-polymerase chain reaction (RT-PCR) to amplify target genes or gene fragments, reduces problems associated with amplification of pseudogenes (non-coding duplicated gene segments) and sequencing through large introns. The main limitation of RNA work is that fresh tissues, or at least tissues collected in an RNA preserving agent such as RNAlater, require rapid transfer to $-80^{\circ} \mathrm{C}$ storage. (2) Phylogenetic work using genomic tissue extractions and amplifications is still favored over RNA techniques due to lower costs, ease of field sampling, and the ability to use previously collected specimens in ethanol. To reduce the risk of sequencing multiple copy genes or pseudogenes, gene fragments are first cloned to identify the number of copies that a primer set amplifies. Although this is not the focus of this paper, in the course of looking for useful phylogenetic markers, we have sequenced a number of multigene families such as hemocyanin, actin, and opsins. These markers may be phylogenetically useful if a single gene is isolated and amplified. They also have many uses when looking at genome evolution and the expression of these genes in Decapoda (e.g., Porter et al. 2007; Scholtz this volume). However, one must be certain that the same copy is being amplified across taxa for useful phylogenetic results.

Introns or highly variable regions need to be considered when sequencing as they can be large (greater than 1000 base pairs in length) and include repeat regions in some taxa, making amplification and sequencing difficult. Often there is too much variation in the intron among taxa to be aligned and included in the analysis. Introns can be avoided by first identifying their position and then designing primer sets within the exon to remove the introns. Here we redesigned primers for elongation factor 2 (EF-2) and transmembrane protein (TM9sf4) to exclude regions of high variability of approximately 300 base pairs in EF2 and 500-1000 base pairs in TM9sf4. Although this reduced the total length of sequence amplified, the highly variable regions produce a greater noise-to-signal ratio at the higher phylogenetic relationships, our principal focus. Of course, these more variable introns might become very useful for population genetic and species level phylogenetic work, and we continue to explore their utility at these lower levels of diversity.

## 3 THE GENES AND THEIR DIVERSITY

### 3.1 Mitochondrial genes: $12 S, 16 \mathrm{~S}$, and COI

Mitochondrial ribosomal genes 12 S and 16 S and coding genes such as COI have been extremely useful in population genetic and systematic studies. Mitochondrial markers have been favored in studies for several reasons (see Schubart, this volume, for details and proposed primer sets for decapod mtDNA amplification). The high copy number of mitochondria in tissues makes them relatively easy to isolate. They are haploid and maternally inherited and consequently are one quarter the effective population size of nuclear genes (Moritz et al. 1987), thus allowing population level studies and systematic studies among recently diverged taxa. Possibly the most important reason to use mitochondrial genes is the availability of universal mtDNA primer sets that have minimized laboratory time in the initial setting up of a project. Finally, there is already an extensive set of nucleotide sequences from these genes in GenBank, as they have been the staple for crustacean molecular phylogenetic work since its inception.

To provide a comparison of gene utility, we have included uncorrected divergence estimates between pairs of taxa: between species, between genera, between families, and between infraorders/ suborders for a number of genes. We also included COI on each graph as a reference (see Figs. 2-5). The ribosomal mitochondrial genes show similar levels of divergence to each other across all comparisons. In 12S, divergence estimates range from 3.9\% among Euastacus species,


Figure 2. Pairwise divergence estimates between species of Euastacus (Astacidea) for mitochondrial and nuclear genes. Species are A: E. eungella and E. spinichelatus, B: E. robertsi and E. eungella, C: E. robertsi and E. spinichelatus.


Figure 3. Pairwise divergence estimates between species of Parastacidae (Astacidea) for mitochondrial and nuclear genes. For genes COI, 12S, 16S, 18S, 28S, H3, EF-2, TM9SF4, EPRS the species are A: Euastacus robertsi and Astacoides betsileoensis, B: E. robertsi and Parastacus defossus, C: A. betsileoensis and P. defossus. Species for genes PEPCK and NaK are A: Homarus gammarus and Nephropides caribaeus, B: H. gammarus and Nephropsis stewarti, C: N. caribaeus and N. stewarti.
$18 \%$ among genera within Parastacidae, $18.6 \%$ among families of Astacidea, and up to $24.2 \%$ among infraorders of Pleocyemata. Divergence of 16 S ranges from $3.5 \%$ among species, $17.6 \%$ among genera, $23.5 \%$ among families, and up to $26.2 \%$ among infraorders of Pleocyemata. The coding mitochondrial gene COI is highly variable among species, thus making it a good candidate at lower levels. High divergence estimates were found above and including the family level, suggesting that this gene may have problems of nucleotide saturation above this level. This gene may still be useful for phylogenetic inference for resolving deeper nodes; however, it is important to test for


Figure 4. Pairwise divergence estimates among family representatives of Astacidea for mitochondrial and nuclear genes. For genes COI, 12S, $16 \mathrm{~S}, 18 \mathrm{~S}, 28 \mathrm{~S}, \mathrm{H} 3, \mathrm{EF}-2$, TM9SF4, EPRS the species are A: E. robertsi and Procambarus clarkii (TM9SF4: Orconectes virilis), B: E. robertsi and Nephropsis aculeata (COI: Homarus americanus), C: P. clarkii (TM9SF4: Orconectes virilis) and N. aculeate (COI: Homarus americanus). Species for genes PEPCK and NaK are A: H. gammarus and Cherax quadricarinatus, B: H. gammarus and P. clarkii, C: C. quadricarinatus and P. clarkii.


Figure 5. Pairwise divergence estimates among representatives of Decapoda for mitochondrial and nuclear genes. For genes COI, 12S, 16S, 18S, 28S, H3, EF-2, TM9SF4, EPRS the species are A: E. robertsi and Calappa gallus (COI: Praebebalia longidactyla), B: C. gallus (COI: P. longidactyla) and Penaeus sp., C: E. robertsi and Penaeus sp. Species for genes PEPCK and NaK are A: H. gammarus and Calappa philargius, B: C. philargius and Penaeus monodon, C: H. gammarus and P. monodon.
saturation and consider this in the analysis (i.e., use a model of evolution that incorporates multiple mutations at the same site - see Palero \& Crandall this volume). A disadvantage of mitochondrial markers is that they are effectively a single locus, and, when used alone, they may not represent the true species tree.

Another problem of some mitochondrial genes such as COI is the presence of pseudogenes (nuclear copies of mitochondrial genes) in some species of decapods (Song et al. 2008).

### 3.2 Nuclear genes

Use of nuclear genes in addition to mitochondrial genes adds to the number of independent markers in a dataset, thus increasing the chances of reconstructing the true species phylogeny. In addition, a larger effective population size, and, on average, a lower substitution rate (Moriyama \& Powell 1997), results in nuclear genes evolving slower than mitochondrial genes. Consequently, they may be better at resolving deeper phylogenetic nodes (see Chu et al. this volume). There are several considerations when choosing nuclear markers. There are at least two copies of each gene, although this is not usually a problem for phylogenetic studies as variation within an individual is less than between species. However, as mentioned previously, many genes belong to multigene families where duplications have resulted in genes or domains with a similar nucleotide sequence. In order to establish a single copy or at least the amplification of one dominant copy for new primer sets (EF-2, EPRS, TM9sf4) presented here, we analyzed 16-24 clones in several taxa representing Pleocyemata (Astacidea (Homarus americanus), Brachyura (Cancer sp.)) and Dendrobranchiata (Penaeus sp.). Low variation among some of the clones was observed. This could be attributed to taq polymerase error assuming an error rate of $1.6 \times 10^{-6}$ to $2.1 \times 10^{-4}$ per nucleotide per cycle (Hengen 1995) or to very low variation of a diploid gene.

The ribosomal nuclear genes 18 S rDNA and 28 S rDNA have been extensively used in arthropod systematics including several decapod studies (e.g., Ahyong \& O'Meally 2004; Porter et al. 2005; Mitsuhashi et al. 2007; Ahyong et al. 2007). Rates of evolution vary among and within these genes, making them valuable phylogenetic tools at different taxonomic levels (Hillis \& Dixon 1991). We found divergence rates for 18 S were consistently moderate among species (5.8-7.2\%) and
Table 1. Gene regions and primer sets selected for reconstructing the phylogeny of decapod crustaceans. For each primer, details of position (3) and a reference sequence are given. NR (nested reaction) refers to the primers used in the first reaction (1) and subsequent hemi-nested reaction (2).

| Gene Region | Primer Name | Primer Sequence ( $\mathbf{5}^{\prime}$ - ${ }^{\prime}$ ) | NR | Position | Reference Sequence | Primer Reference |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Mitochondrial |  |  |  |  |  |  |
| Genes |  |  |  |  |  |  |
| 12 S rRNA | 12sf | GAA ACC AGG ATT AGA TAC CC |  | 390 | AY659990 | Mokady et al. 1994 |
|  | 12sr | TTT CCC GCG AGC GAC GGG CG |  | 778 | AY659990 | Mokady et al. 1994 |
| 16S rRNA | 16s-1472 | AGA TAG AAA CCA ACC TGG |  | 99 | AF200829 | Crandall \& Fitzpatrick 1996 |
|  | 16sf-cray | GAC CGT GCK AAG GTA GCA TAA TC |  | 552 | AF200829 | Crandall \& Fitzpatrick 1996 |
| COI | LCO1-1490 | GGT CAA CAA ATC ATA AAG ATA TTG |  | * |  | Folmer et al. 1994 |
|  | HCO1-2198 | TAA ACT TCA GGG TGA CCA AAA AAT |  | * |  | Folmer et al. 1994 |
|  |  | CA |  |  |  |  |
| Nuclear Genes |  |  |  |  |  |  |
| 18S rRNA | 18 s 1 f | TAC CTG GTT GAT CCT GCC AGT AG |  | * |  | Whiting et al. 1997, Whiting 2002 |
|  | 18 s b3.0 | GAC GGT CCA ACA ATT TCA CC |  | * |  | Whiting et al. 1997, Whiting 2002 |
|  | 18s a 0.79 | TTA GAG TGC TYA AAG C |  | * |  | Whiting et al. 1997, Whiting 2002 |
|  | 18 s bi | GAG TCT CGT TCG TTA TCG GA |  | * |  | Whiting et al. 1997, Whiting 2002 |
|  | 18 s a 2.0 | ATG GTT GCA AAG CTG AAA C |  | * |  | Whiting et al. 1997, Whiting 2002 |
|  | 18s 9R | GAT CCT TCC GCA GGT TCA CCT AC |  | * |  | Whiting et al. 1997, Whiting 2002 |
| 28 S rRNA | $28 \mathrm{~s}-\mathrm{rD} 1.2 \mathrm{a}$ | CCC SSG TAA TTT AAG CAT ATT A |  | * |  | Whiting et al. 1997, Whiting 2002 |
|  | 28s-rD3a | AGT ACG TGA AAC CGT TCA GG |  | * |  | Whiting et al. 1997, Whiting 2002 |
|  | 28s-rd3.3f | GAA GAG AGA GTT CAA GAG TAC G |  | * |  | Whiting et al. 1997, Whiting 2002 |
|  | 28sA | GAC CCG TCT TGA AGC ACG |  | * |  | Whiting et al. 1997, Whiting 2002 |
|  | 28s-rD4.5a | AAG TTT CCC TCA GGA TAG CTG |  | * |  | Whiting et al. 1997, Whiting 2002 |
|  | 28S rD5a | GGY GTT GGT TGC TTA AGA CAG |  | * |  | Whiting et al. 1997, Whiting 2002 |
|  | 28s-rD4b | CCT TGG TCC GTG TTT CAA GAC |  | * |  | Whiting et al. 1997, Whiting 2002 |
|  | 28S B | TCG GAA GGA ACC AGC TAC |  | * |  | Whiting et al. 1997, Whiting 2002 |
|  | 28s-rD5b | CCA CAG CGC Cag TTC TGC TTA C |  | * |  | Whiting et al. 1997, Whiting 2002 |
|  | 28s-rD6b | AAC CRG ATT CCC TTT CGC C |  | * |  | Whiting et al. 1997, Whiting 2002 |
|  | 28S rD7bl | GAC TTC CCT TAC CTA CAT |  | * |  | Whiting et al. 1997, Whiting 2002 |
|  | 28s3.25a | CAG GTG GTA AAC TCC ATC AAG G |  | 602 | AY210833 | this study |
|  | 28s4.4b | GCT ATC CTG AGG GAA ACT TCG |  | 1594 | AY210833 | this study |

Table 1. continued.

| Gene Region | Primer Name | Primer Sequence ( $\mathbf{5}^{\prime}-3^{\prime}$ ) | NR | Position | Reference <br> Sequence | Primer Reference |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| H3 | H3 AF | ATG GCT CGT ACC AAG CAG ACV GC |  | 321 | AB044542 | Colgan et al. 1998 |
|  | H3 AR | ATA TCC TTR GGC ATR ATR GTG AC |  | 694 | AB044542. | Colgan et al. 1998 |
| EF-2 | EF2a IF2 | TGG GGW GAR AAC TTC TTY AAC |  | 824 | EF426560 | Porter ML pers. comm. |
|  | EF2a 1R2 | ACC ATY TTK GAG ATG TAC ATC AT |  | 1236 | EF426560 | Porter ML pers. comm. |
|  | EF2a-F978 | TGG ANA CBC TGA ARA TCA A | 1,2 | 978 | EF426560 | this study |
|  | EF2-R1435 | GTT ACC HGC TGG VAC RTC TTC | 2 | 1435 | EF426560 | this study |
|  | EF2-R1536 | GAC ACG NWG AAC TTC ATC ACC | 1 | 1536 | EF426560 | this study |
| EPRS | 192finlf | +GAR AAR GAR AAR TTY GC |  | 6874 | U59923 | www.umbi.umd.edu/users/jcrlab/ |
|  | 192fin 2 r | +TCC CAR TGR TTR AAY TTC CA |  | 7316 | U59923 | www.umbi.umd.edu/users/jcrlab/ |
| TM9SF4 | 3064fin6f | CAR GAR GAR TTY GGN TGG AA | 1 | 1198 | $\begin{aligned} & \mathrm{NM}_{-} \\ & 014742 \end{aligned}$ | www.umbi.umd.edu/users/jcrlab/ |
|  | 3064fin7r | AAN CCR AAC ATR TAR TA |  | 1841 | $\begin{aligned} & \text { NM_ } \\ & 014742 \end{aligned}$ | www.umbi.umd.edu/users/jcrlab/ |
|  | 3064-F1204 | +GAA TTT GGR TGG AAG CTG GT | 2 | 1204 | $\begin{aligned} & \text { NM- }_{-} \\ & 014742 \end{aligned}$ | this study |
|  | 3064-R1697 | +CTG GGN ATY TGG TTG GTT CG | 1,2 | 1697 | $\begin{aligned} & \text { NM- }_{-} \\ & 014742 \end{aligned}$ | this study |

"*" see primer reference for primer positions. " + " addition of M13 primers to the 5 ' end improves PCR amplification (Regier \& Shi 2005).
among infraorders ( $5.6 \%$ ) within Pleocyemata but were higher among the suborders Pleocyemata and Dendrobranchiata ( $12.8 \%$ and $14.1 \%$ ). Two hypervariable regions of 28 S were identified and removed to avoid inflated estimates of divergence among poorly aligned repeat regions. 28 S divergence estimates were higher than 18 S among species ( $9.1-11.6 \%$ ), within Pleocyemata ( $11.3 \%$ ), and among the suborders (20.8-21.8\%). Levels of divergence were lower for the intermediate taxon levels, among genera ( $3.4-8.0 \%$ ), and among families ( $7.3-9.9 \%$ ), and possibly represented a shorter nucleotide alignment due to indels (insertions or deletions) that are absent among species (within a genus).

Two nuclear protein coding genes that are currently used in arthropod systematics are histone 3 (H3) (e.g., Porter et al. 2005) and elongation factor 2 (EF-2) (e.g., Regier \& Shultz 2001). Primer sets already developed for H 3 (Colgan et al. 1998) amplify the target fragment across a range of decapod crustaceans and show moderate levels of divergence among species ( $2.2-8.4 \%$ ), suggesting they are useful nuclear protein coding markers for relationships within a genus. It should be noted that Euastacus is relatively older than some decapod genera (see Breinholt et al. this volume) and consequently H 3 may not be appropriate for phylogenetic analyses among recently diverged species. Divergence within and among families is also moderate ( $8.9-12.4 \%$ ), with a higher level of divergence between Euastacus robertsi and Calappa gallus within Pleocyemata ( $17 \%$ ).

Although we were able to amplify genomic fragments of the EF-2 gene with currently designed primer sets (see Table 1), an intron was located at base pair position 860 relative to mRNA in Libinia emarginata (GenBank accession AY305506). The intron may be useful for species/genera level studies, although preliminary analysis suggests it is fewer than 300 base pairs in caridean (Hippolytidae) and brachyuran (Calappidae, Leucosiidae, Goneplacidae, Majidae, Cyclodorippidae) decapods. A new forward primer was designed to exclude the intron, and GenBank sequences were downloaded and aligned to design reverse primers 400-500 base pairs downstream of the forward primer. Using different primer sets, we were able to isolate two copies of EF-2. The two copies were more similar within an individual than between species of Euastacus crayfish. Two similar copies of EF-2 are present in Drosophila melanogaster (Lasko 2000). The divergence estimates for the longer fragment are presented in figure 2 and were low among species of Euastacus $(1.3 \%)$. Percent divergence within Parastacidae ( $6.7-9.3 \%$ ) and between families of Astacidea ( $13.6 \%$ ) was moderate. High divergences were noted within Pleocyemata between E. robertsi and C. gallus ( $18.7 \%$ ).

The EPRS locus is a potentially useful nuclear gene for reconstructing phylogenetic relationships among the deeper nodes of decapod crustaceans. The EPRS locus encodes a multifunctional aminoacyl tRNA synthetase, glutamyl-prolyl-tRNA synthetase (Cerini et al. 1991). The two proteins are involved in the aminoacylation of glutamic acid and praline tRNA in Drosophila (Cerini et al. 1991; Cerini et al. 1997). Few phylogenetic studies have used EPRS, although a recent study of Paramysis (Crustacea: Mysida) demonstrates its usefulness in reconstructing relationships among genera of mysids (Audzijonyte et al. 2008). We found divergence levels were low among species of Euastacus ( $0.8-1.5 \%$ ) but moderate for within the family Parastacidae ( $5.2-8.6 \%$ ) and high between some families of Astacidea (11.3-20.5\%). This locus showed high divergences within Pleocyemata between E. robertsi and C. gallus ( $33.9 \%$ ) and between E. robertsi and Penaeus sp. (15.5-30.1\%). The different levels of divergence at different taxonmic levels suggest this marker may be useful among genera up to order level for phylogenetic estimation.

Transmembrane 9 superfamily protein member 4, or TM9sf4, is a small molecule carrier or transporter. Our study is the first to present divergence estimates and phylogenetic results using this gene. Uncorrected pairwise divergence results suggest it has potential as a valuable gene for reconstructing family to order level relationships. Divergence among species within Euastacus was low ( $0.7-1.5 \%$ ), suggesting this marker may be less informative than other nuclear protein coding markers such as Histone 3 when reconstructing relationships among species. As with EPRS, this marker shows greater divergences ( $18.8-23 \%$ ) at the deeper level (among infraorders/suborders)
than Histone 3. High levels of divergence are often considered indicative of saturation; however, we found increasing divergence with increasing evolutionary distance, suggesting saturation may not have been reached even among the deeper nodes, indicating the utility of this gene to infer phylogenetic relationships at these higher levels of divergence.

## 4 PHYLOGENY BASED SYSTEMATICS

Reconstructing the evolutionary relationships among decapod crustaceans using molecular data has taken two directions: using only protein coding genes, which are phylogenetically informative at deeper nodes, or incorporating as much molecular information available including both ribosomal RNA and protein coding genes in a family level supertree. We have taken the latter approach and reconstructed Decapoda relationships using a total of eight genes and 46 taxa (see Table 2) including representatives of seven infraorders of Pleocyemata and a representative of Dendrobranchiata ( Pe naeus sp.) as an outgroup. Pleocyemata representatives include Astacidea, Achelata, Polychelida, Thalassinidea, Brachyura, Anomura and Caridea. Non-decapod crustaceans, Lysiosquillina maculata (Lysiosquillidae: Stomatopoda), were also included in the analysis as outgroups to all the decapods. Rather than focus on representing all lineages equally, we were interested in reconstructing relationships at many levels from among species within genera, among families, and among infraorders within decapod crustaceans. Therefore, we focused on sampling the Astacidea to demonstrate the usefulness of these genes for reconstructing phylogenies at these various taxonomic levels.

The genes included in our analyses were 12S, 16S, 18S, 28S, H3, EF-2, EPRS, and TM9sf4. A second analysis was run on the four nuclear protein-coding genes. Use of nuclear rRNA 18 S and 28S data has been criticized for ambiguities noted in alignments (Tsang et al. 2008). The difficulties in aligning highly variable data may be overcome by using sophisticated methods of alignment employed in recently developed programs such as DIALIGN-T (Subramanian et al. 2005) and MAFFT (Katoh et al. 2002; Katoh et al. 2005). These programs produce more accurate alignments than ClustalW with increasing evolutionary distance (e.g., MAFFT, Nuin et al. 2006) or when gaps are present (indels) in the resulting alignment of sequence data (e.g., DIALIGN-T and MAFFT, Golubchik et al. 2007). To further improve the alignment, GBlocks can be used to identify and exclude ambiguous regions of sequence data (Castresana 2000; Talavera \& Castresana 2007). We used MAFFT to align all gene fragments and subsequently ran each dataset through GBlocks (retaining half gap positions) to recover the most useful sequence data. As an example, this reduced the 28 S MAFFT alignment from 4489 to 1254 base pairs. Our resulting alignment for the eight-gene dataset was 5104 nucleotides.

Maximum likelihood phylogenies were constructed with RAxML (Stamatakis 2006; Stamatakis et al. 2008) at the CIPRES portal assuming a GTR $+G+I$ model and estimation and optimization of $\alpha$-shape parameters, GTR-rates, and empirical base frequencies for each gene. We allowed the program to choose the number of bootstrap replicates, and for the eight-gene dataset, 150 bootstrap replicates were run before termination. For the smaller nuclear protein coding alignment, 250 bootstrap replicates were run before the program terminated. The estimated parameters are presented in Table 3.

The relationships within Astacidea were well resolved, with bootstrap support in 11 of 14 nodes supported by $95 \%$ or greater and all nodes supported greater than $80 \%$ (see Fig. 6). As a comparison, the ML phylogeny based on the four-gene dataset (nuclear protein coding) constructed a similar topology within Astacidea although the nodes were not as strongly supported. Only six nodes were supported greater than $95 \%$, with an additional five nodes supported greater than $70 \%$. This result suggests that although the nuclear coding genes have the power to resolve relationships within an infraorder, additional data from ribosomal genes adds to the information available for reconstructing relationships across the whole of decapod diversity. Our group continues to add genes and taxa to achieve our goal of reconstructing a robust phylogenetic estimate for the decapod crustaceans.
Table 2. Taxonomy and accession numbers of decapod samples and outgroup included in this study. Accession numbers in bold were obtained from GenBank.

| Taxon | $\begin{aligned} & \text { Voucher } \\ & \text { ID } \end{aligned}$ | $\begin{aligned} & \text { 12S } \\ & \text { rRNA } \end{aligned}$ | $\begin{aligned} & \text { 16S } \\ & \text { rRNA } \end{aligned}$ | $\begin{aligned} & 18 \mathrm{~S} \\ & \text { rRNA } \end{aligned}$ | $\begin{aligned} & \text { 28S } \\ & \text { rRNA } \end{aligned}$ | H3 | EF-2 | EPRS | TM9SF4 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Decapoda Latreille, 1802 |  |  |  |  |  |  |  |  |  |
| Dendrobranchiata Bate, 1888 |  |  |  |  |  |  |  |  |  |
| Penaeoidea Rafinesque, 1815 |  |  |  |  |  |  |  |  |  |
| Penaeus sp. Fabricius, 1798 | KCpen | EU920908 | EU920934 | EU920969 | $\begin{aligned} & \text { EU921005- } \\ & \text { EU921006 } \end{aligned}$ | EU921075 | - | - | EU921109 |
| Pleocyemata Burkenroad, 1963 |  |  |  |  |  |  |  |  |  |
| Anomura MacLeay, 1838 |  |  |  |  |  |  |  |  |  |
| Galatheoidea Samouelle, 1819 |  |  |  |  |  |  |  |  |  |
| Aegla alacaluf (Jara \& López, 1981) | KAC798 | AY050012 | AY050058 | EU920958 | AY595958 | EU921042 | EU921009 | EU910098 | EU921077 |
| Eumunida funambulus (Miyake, 1982) | KC3100 | EU920892 | EU920922 | EU920957 | EU920984 | EU921056 | EU921032 | EU910124 | EU921089 |
| Kiwa hirsute (Jones \& Segonzac, 2005) | KC3116 | - | - | EU920942 | EU920987 | EU921065 | EU921035 | EU910128 | EU921097 |
| Munidopsis rostrata (Milne-Edwards, 1880) | KC3102 | EU920898 | EU920928 | EU920961 | EU920985 | EU921066 | EU921034 | EU910126 | EU921100 |
| Lomisoidea Bouvier, 1895 |  |  |  |  |  |  |  |  |  |
| Lomis hirta (Lamarck, 1810) | KAClohi | AY595547 | AY595928 | AF436013 | AY596101 | DQ079680 | EU921040 | EU910131 | EU921098 |
| Paguroidea Latreille, 1802 |  |  |  |  |  |  |  |  |  |
| Pomatocheles jeffreysii (Miers, 1879) | KC3097 | EU920903 | EU920930 | EU920965 | EU920983 | EU921070 | EU921031 | EU910123 | EU921105 |
| Astacidea Latreille, 1802 |  |  |  |  |  |  |  |  |  |
| Astacoidea Latreille, 1802 |  |  |  |  |  |  |  |  |  |
| Astacus astacus (Linnaeus, 1758) | KC702 | EU920881 | AF235983 | AF235959 | DQ079773 | DQ079660 | EU921008 | - | EU921078 |
| Barbicambarus cornutus (Faxon, 1884) | KC1941 | EU920883 | EU920913 | EU920951 | EU920993 | EU921045 | EU921017 | EU910106 | EU921080 |
| Orconectes virilis (Hagen, 1870) | KC709 | EU920900 | AF235989 | AF235965 | DQ079804 | DQ079693 | EU921041 | - | EU921102 |
| Procambarus clarkii (Girard, 1852) | KC1497 | EU920901 | AF235990 | EU920952 | EU920970 | EU921067 | EU921011 | EU910100 | - |
| Parastacoidea Huxley, 1879 |  |  |  |  |  |  |  |  |  |
| Astacoides betsileoensis (Petit, 1923) | KC1822 | EU920882 | EU920912 | EU920955 | EU920992 | EU921044 | EU921014 | EU910103 | EU921079 |
| Cherax cuspidatus (Riek, 1969) | KC1175 | DQ006421 | DQ006550 | EU920960 | EU920996 | EU921048 | EU921010 | EU910099 | EU921083 |
| Euastacus eungella (Morgan, 1988) | KC2671 | DQ006464 | DQ006593 | EU920964 | $\begin{aligned} & \text { EU92 100- } \\ & \text { EU921002 } \end{aligned}$ | EU921055 | EU921018 | EU910109 | EU921088 |
| Euastacus robertsi (Monroe, 1977) | KC2781 | DQ006507 | DQ006633 | EU920962 | EU920988 | EU921058 | EU921019 | EU910110 | EU921091 |
| Euastacus spinichelatus (Morgan, 1997) | KC2631 | DQ006512 | DQ006638 | EU920963 | EU920989 | EU921059 | - | EU910108 | EU921092 |
| Gramastacus insolitus (Riek, 1972) | KC640 | EU920895 | EU920926 | EU920968 | EU920994 | EU921062 | EU921007 | EU910097 | EU921094 |
| Ombrastacoides huonensis (Riek, 1967) | KC611 | EU920905 | AF135997 | EU920956 | EU920995 | EU921072 | - | EU910096 | EU921106 |
| Parastacus defossus (Faxon, 1898) | KC1515 | EU920902 | AF175243 | EU920953 | EU920991 | EU921068 | EU921012 | EU910101 | EU921103 |
| Parastacus varicosus (Faxon, 1898) | KC1529 | EU920907 | EU920933 | EU920954 | EU920990 | EU921074 | EU921013 | EU910102 | EU921108 |

Table 2. continued

| Taxon | Voucher ID | $\begin{aligned} & \text { 12S } \\ & \text { rRNA } \end{aligned}$ | $\begin{aligned} & \text { 16S } \\ & \text { rRNA } \end{aligned}$ | $\begin{aligned} & \text { 18S } \\ & \text { rRNA } \end{aligned}$ | $28 S$ <br> rRNA | H3 | EF-2 | EPRS | TM9SF4 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Nephropoidea Dana, 1852 |  |  |  |  |  |  |  |  |  |
| Homarus americanus (Milne-Edwards, 1837) | KAChoam | DQ298427 | HAU11238 | AF235971 | DQ079788 | DQ079675 | - | - | EU921095 |
| Nephropsis aculeate (Smith, 1881) | KC2117 | EU920899 | DQ079727 | DQ079761 | DQ079802 | DQ079691 | - | EU910107 | EU921101 |
| Brachyura Latreille, 1802 |  |  |  |  |  |  |  |  |  |
| Calappoidea Milne-Edwards, $1837{ }^{\text {- }}$ |  |  |  |  |  |  |  |  |  |
| Cycloes granulose (de Haan, 1837) | KC3082 | EU920887 | EU920917 | EU920943 | EU920976 | EU921050 | EU921025 | EU910116 | EU921085 |
| Calappa gallus (Herbst, 1803) | KC3083 | EU920886 | EU920916 | EU920947 | EU920977 | EU921049 | EU921026 | EU910117 | EU921084 |
| Dorippoidea MacLeay, 1838 |  |  |  |  |  |  |  |  |  |
| Ethusa sp. (Roux, 1830) | KC3088 | - | EU920925 | EU920966 | EU920980 | EU921061 | EU921029 | EU910120 | EU921093 |
| Grapsoidea MacLeay, 1838 |  |  |  |  |  |  |  |  |  |
| Cyclograpsus cinereus (Dana, 1851) | KC3417 | EU920884 | EU920914 | EU920945 | EU920997 | EU921046 | EU921038 | EU910130 | EU921081 |
| Leucosioidea Samouelle, 1819 |  |  |  |  |  |  |  |  |  |
| Ebalia tuberculosa (Milne-Edwards, 1873) | KC3085 | EU920894 | EU920924 | EU920944 | EU920978 | EU921060 | EU921027 | EU910118 | - |
| Praebebalia longidactyla (Yokoya, 1933) | KC3086 | EU920904 | EU920931 | EU920946 | EU920979 | EU921071 | EU921028 | EU910119 | - |
| Majoidea Samouelle, 1819 |  |  |  |  |  |  |  |  |  |
| Chorilia longipes (Dana, 1852) | KC3089 | EU920889 | EU920919 | EU920948 | EU920981 | EU921052 | EU921039 | EU910121 | EU921087 |
| Raninoidea de Haan, 1839 |  |  |  |  |  |  |  |  |  |
| Cosmonotus grayi (White, 1848) | KC3092 | EU920888 | EU920918 | EU920949 | EU920982 | EU921051 | EU921030 | EU910122 | EU921086 |
| Caridea Dana, 1852 |  |  |  |  |  |  |  |  |  |
| Palaemonoidea Rafinesque, 1815 |  |  |  |  |  |  |  |  |  |
| Anchistioides antiguensis (Schmitt, 1924) | KC3051 | EU920880 | EU920911 | EU920936 | EU920971 | EU921043 | EU921020 | EU910111 | - |
| Coutierella tonkinensis (Sollaud, 1914) | KC3068 | EU920890 | EU920920 | EU920937 | EU920975 | EU921053 | EU921024 | EU910115 | - |
| Crangonoidea Haworth, 1825 |  |  |  |  |  |  |  |  |  |
| Crangon crangon (Linnaeus, 1758) | KC3052 | EU920885 | EU920915 | EU920938 | EU920972 | EU921047 | EU921021 | EU910112 | EU921082 |
| Bresilioidea Calman, 1896 |  |  |  |  |  |  |  |  |  |
| Discias sp. (Rathbun, 1902) | KC3108 | EU920891 | EU920921 | EU920941 | EU920986 | EU921054 | - | EU910127 | - |
| Alpheoidea Rafinesque, 1815 |  |  |  |  |  |  |  |  |  |
| Hippolyte bifidirostris (Miers, 1876) | KC3059 | EU920896 | EU920927 | EU920939 | EU920974 | EU921063 | EU921023 | EU910114 | - |
| Eualus gaimardii (Milne-Edwards, 1837) | KC3056 | EU920893 | EU920923 | EU920940 | EU920973 | EU921057 | EU921022 | EU910113 | EU921090 |

Table 2. continued.

| Taxon | Voucher ID | $\begin{aligned} & \text { 12S } \\ & \text { rRNA } \end{aligned}$ | $\begin{aligned} & \text { 16S } \\ & \text { rRNA } \end{aligned}$ | $\begin{aligned} & \text { 18S } \\ & \text { rRNA } \end{aligned}$ | $\begin{aligned} & \text { 28S } \\ & \text { rRNA } \end{aligned}$ | H3 | EF-2 | EPRS | TM9SF4 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Achelata Scholtz \& Richter, 1995 |  |  |  |  |  |  |  |  |  |
| Palinuroidea Latreille, 1802 |  |  |  |  |  |  |  |  |  |
| Jasus edwardsii (Hutton, 1875) | KC3209 | - | DQ079716 | AF235972 | DQ079791 | EU921064 | EU921036 | EU910129 | EU921096 |
| Palinurus elephas (Fabricius, 1787) | KC3210 | - | EU920929 | EU920959 | $\begin{aligned} & \text { EU920999- } \\ & \text { EU921000 } \end{aligned}$ | EU921069 | EU921037 | - | EU921104 |
| Polychelida de Haan, 1941 |  |  |  |  |  |  |  |  |  |
| Polycheles typhlops (Heller, 1862) | KC3101 | EU920906 | EU920932 | EU920950 | $\begin{aligned} & \text { EU921003- } \\ & \text { EU921004 } \end{aligned}$ | EU921073 | EU921033 | EU910125 | EU921107 |
| Thalassinidea Latreille, 1831 |  |  |  |  |  |  |  |  |  |
| Callianassoidea Dana, 1852 |  |  |  |  |  |  |  |  |  |
| 1935) <br> Lepidophthalmus louisianensis (Schmitt, | KAC1852 | EU920897 | DQ079717 | DQ079751 | DQ079792 | DQ079678 | EU921015 | EU910104 | EU921099 |
| Sergio mericeae (Manning \& Felder, 1995) | KAC1865 | EU920909 | DQ079733 | DQ079768 | DQ079811 | DQ079700 | EU921016 | EU910105 | EU921110 |
| Outgroup |  |  |  |  |  |  |  |  |  |
| Stomatopoda Latreille, 1817 |  |  |  |  |  |  |  |  |  |
| Lysiosquilloidea Giesbrecht, 1910 |  |  |  |  |  |  |  |  |  |
| Lysiosquillina maculata (Fabricius, 1793) | KC3832 | EU920910 | EU920935 | EU920967 | EU920998 | EU921076 | - | - | EU921111 |

Table 3. Empirical base frequencies for each gene region and associated model parameters estimated from the sequence data in RAxML.

|  | A | C | G | T | alpha | pinvar |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: |
| 12S rRNA | 0.3670 | 0.0981 | 0.1726 | 0.3622 | 0.6030 | 0.1934 |
| 16S rRNA | 0.3399 | 0.1116 | 0.2027 | 0.3458 | 0.6235 | 0.2879 |
| 18S rRNA | 0.2502 | 0.2342 | 0.2780 | 0.2377 | 0.9231 | 0.4940 |
| 28S rRNA | 0.2501 | 0.2357 | 0.3161 | 0.1981 | 0.7772 | 0.2735 |
| H3 | 0.2152 | 0.3172 | 0.2654 | 0.2022 | 1.0618 | 0.5882 |
| EF-2 | 0.2364 | 0.2469 | 0.2655 | 0.2512 | 1.4067 | 0.4872 |
| EPRS | 0.2857 | 0.2159 | 0.2523 | 0.2460 | 1.6197 | 0.3690 |
| TM9SF4 | 0.1587 | 0.2784 | 0.2455 | 0.3174 | 0.9592 | 0.4982 |



Figure 6. Maximum likelihood phylogeny based on two mitochondrial and six nuclear genes constructed in RAxML. Values at nodes represent bootstrap support greater than $70 \%$.

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