

CRUSTACEAN ISSUES 18



# Decapod Crustacean Phylogenetics

edited by

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



CRC Press  
Taylor & Francis Group

# Decapod Crustacean Phylogenetics

Edited by

**Joel W. Martin**

Natural History Museum of L. A. County  
Los Angeles, California, U. S. A.

**Keith A. Crandall**

Brigham Young University  
Provo, Utah, U. S. A.

**Darryl L. Felder**

University of Louisiana  
Lafayette, Louisiana, U. S. A.



**CRC Press**

Taylor & Francis Group

Boca Raton London New York

---

CRC Press is an imprint of the  
Taylor & Francis Group, an **informa** business

CRC Press  
Taylor & Francis Group  
6000 Broken Sound Parkway NW, Suite 300  
Boca Raton, FL 33487-2742

© 2009 by Taylor & Francis Group, LLC  
CRC Press is an imprint of Taylor & Francis Group, an Informa business

No claim to original U.S. Government works  
Printed in the United States of America on acid-free paper  
10 9 8 7 6 5 4 3 2 1

International Standard Book Number-13: 978-1-4200-9258-5 (Hardcover)

This book contains information obtained from authentic and highly regarded sources. Reasonable efforts have been made to publish reliable data and information, but the author and publisher cannot assume responsibility for the validity of all materials or the consequences of their use. The authors and publishers have attempted to trace the copyright holders of all material reproduced in this publication and apologize to copyright holders if permission to publish in this form has not been obtained. If any copyright material has not been acknowledged please write and let us know so we may rectify in any future reprint.

Except as permitted under U.S. Copyright Law, no part of this book may be reprinted, reproduced, transmitted, or utilized in any form by any electronic, mechanical, or other means, now known or hereafter invented, including photocopying, microfilming, and recording, or in any information storage or retrieval system, without written permission from the publishers.

For permission to photocopy or use material electronically from this work, please access [www.copyright.com](http://www.copyright.com) (<http://www.copyright.com/>) or contact the Copyright Clearance Center, Inc. (CCC), 222 Rosewood Drive, Danvers, MA 01923, 978-750-8400. CCC is a not-for-profit organization that provides licenses and registration for a variety of users. For organizations that have been granted a photocopy license by the CCC, a separate system of payment has been arranged.

**Trademark Notice:** Product or corporate names may be trademarks or registered trademarks, and are used only for identification and explanation without intent to infringe.

---

**Library of Congress Cataloging-in-Publication Data**

---

Decapod crustacean phylogenetics / editors, Joel W. Martin, Keith A. Crandall, Darryl L. Felder.  
p. cm. -- (Crustacean issues)

Includes bibliographical references and index.

ISBN 978-1-4200-9258-5 (hardcover : alk. paper)

1. Decapoda (Crustacea) 2. Phylogeny. I. Martin, Joel W. II. Crandall, Keith A. III. Felder, Darryl L.  
IV. Title. V. Series.

QL444.M33D44 2009

595.3'8138--dc22

2009001091

---

Visit the Taylor & Francis Web site at  
<http://www.taylorandfrancis.com>

and the CRC Press Web site at  
<http://www.crcpress.com>

# Contents

|  |     |
|--|-----|
| Preface  | ix  |
| JOEL W. MARTIN, KEITH A. CRANDALL & DARRYL L. FELDER   |     |
| <br>   |     |
| <b>I</b> <i>Overviews of Decapod Phylogeny</i>   |     |
| On the Origin of Decapoda  | 3   |
| FREDERICK R. SCHRAM  |     |
| Decapod Phylogenetics and Molecular Evolution  | 15  |
| ALICIA TOON, MAEGAN FINLEY, JEFFREY STAPLES & KEITH A. CRANDALL                                  |     |
| Development, Genes, and Decapod Evolution  | 31  |
| GERHARD SCHOLTZ, ARKHAT ABZHANOV, FREDERIKE ALWES, CATERINA BIFFIS & JULIA PINT                  |     |
| Mitochondrial DNA and Decapod Phylogenies: The Importance of Pseudogenes and Primer Optimization | 47  |
| CHRISTOPH D. SCHUBART  |     |
| Phylogenetic Inference Using Molecular Data  | 67  |
| FERRAN PALERO & KEITH A. CRANDALL  |     |
| Decapod Phylogeny: What Can Protein-Coding Genes Tell Us?  | 89  |
| K.H. CHU, L.M. TSANG, K.Y. MA, T.Y. CHAN & P.K.L. NG   |     |
| Spermatozoal Morphology and Its Bearing on Decapod Phylogeny                                     | 101 |
| CHRISTOPHER TUDGE  |     |
| The Evolution of Mating Systems in Decapod Crustaceans   | 121 |
| AKIRA ASAKURA  |     |
| A Shrimp's Eye View of Evolution: How Useful Are Visual Characters in Decapod Phylogenetics?     | 183 |
| MEGAN L. PORTER & THOMAS W. CRONIN   |     |
| Crustacean Parasites as Phylogenetic Indicators in Decapod Evolution                             | 197 |
| CHRISTOPHER B. BOYKO & JASON D. WILLIAMS   |     |
| The Bearing of Larval Morphology on Brachyuran Phylogeny   | 221 |
| PAUL F. CLARK  |     |

## II *Advances in Our Knowledge of Shrimp-Like Decapods*

- Evolution and Radiation of Shrimp-Like Decapods: An Overview 245  
CHARLES H.J.M. FRANSEN & SAMMY DE GRAVE
- A Preliminary Phylogenetic Analysis of the Dendrobranchiata Based on Morphological Characters 261  
CAROLINA TAVARES, CRISTIANA SEREJO & JOEL W. MARTIN
- Phylogeny of the Infraorder Caridea Based on Mitochondrial and Nuclear Genes (Crustacea: Decapoda) 281  
HEATHER D. BRACKEN, SAMMY DE GRAVE & DARRYL L. FELDER

## III *Advances in Our Knowledge of the Thalassinidean and Lobster-Like Groups*

- Molecular Phylogeny of the Thalassinidea Based on Nuclear and Mitochondrial Genes 309  
RAFAEL ROBLES, CHRISTOPHER C. TUDGE, PETER C. DWORSCHAK, GARY C.B. POORE & DARRYL L. FELDER
- Molecular Phylogeny of the Family Callinassidae Based on Preliminary Analyses of Two Mitochondrial Genes 327  
DARRYL L. FELDER & RAFAEL ROBLES
- The Timing of the Diversification of the Freshwater Crayfishes 343  
JESSE BREINHOLT, MARCOS PÉREZ-LOSADA & KEITH A. CRANDALL
- Phylogeny of Marine Clawed Lobster Families Nephropidae Dana, 1852, and Thaumastocheilidae Bate, 1888, Based on Mitochondrial Genes 357  
DALE TSHUDY, RAFAEL ROBLES, TIN-YAM CHAN, KA CHAI HO, KA HOU CHU, SHANE T. AHYONG & DARRYL L. FELDER
- The Polychelidan Lobsters: Phylogeny and Systematics (Polychelida: Polychelidae) 369  
SHANE T. AHYONG

## IV *Advances in Our Knowledge of the Anomura*

- Anomuran Phylogeny: New Insights from Molecular Data 399  
SHANE T. AHYONG, KAREEN E. SCHNABEL & ELIZABETH W. MAAS

## V *Advances in Our Knowledge of the Brachyura*

- Is the Brachyura Podotremata a Monophyletic Group? 417  
GERHARD SCHOLTZ & COLIN L. MCLAY

|   |     |
|---|-----|
| Assessing the Contribution of Molecular and Larval Morphological Characters in a Combined Phylogenetic Analysis of the Superfamily Majoidea   | 437 |
| KRISTIN M. HULTGREN, GUILLERMO GUERAO, FERNANDO P.L. MARQUES & FERRAN P. PALERO   |     |
| Molecular Genetic Re-Examination of Subfamilies and Polyphyly in the Family Pinnotheridae (Crustacea: Decapoda)   | 457 |
| EMMA PALACIOS-THEIL, JOSÉ A. CUESTA, ERNESTO CAMPOS & DARRYL L. FELDER  |     |
| Evolutionary Origin of the Gall Crabs (Family Cryptochiridae) Based on 16S rDNA Sequence Data   | 475 |
| REGINA WETZER, JOEL W. MARTIN & SARAH L. BOYCE  |     |
| Systematics, Evolution, and Biogeography of Freshwater Crabs  | 491 |
| NEIL CUMBERLIDGE & PETER K.L. NG  |     |
| Phylogeny and Biogeography of Asian Freshwater Crabs of the Family Gecarcinucidae (Brachyura: Potamoidea)   | 509 |
| SEBASTIAN KLAUS, DIRK BRANDIS, PETER K.L. NG, DARREN C.J. YEO & CHRISTOPH D. SCHUBART   |     |
| A Proposal for a New Classification of Portunoidea and Cancroidea (Brachyura: Heterotremata) Based on Two Independent Molecular Phylogenies   | 533 |
| CHRISTOPH D. SCHUBART & SILKE REUSCHEL  |     |
| Molecular Phylogeny of Western Atlantic Representatives of the Genus <i>Hexapanopeus</i> (Decapoda: Brachyura: Panopeidae)  | 551 |
| BRENT P. THOMA, CHRISTOPH D. SCHUBART & DARRYL L. FELDER  |     |
| Molecular Phylogeny of the Genus <i>Cronius</i> Stimpson, 1860, with Reassignment of <i>C. tumidulus</i> and Several American Species of <i>Portunus</i> to the Genus <i>Achelous</i> De Haan, 1833 (Brachyura: Portunidae) | 567 |
| FERNANDO L. MANTELATTO, RAFAEL ROBLES, CHRISTOPH D. SCHUBART & DARRYL L. FELDER   |     |
| Index   | 581 |
| Color Insert  |     |

# The Timing of the Diversification of the Freshwater Crayfishes

JESSE BREINHOLT<sup>1</sup>, MARCOS PÉREZ-LOSADA<sup>2</sup> & KEITH A. CRANDALL<sup>1</sup>

<sup>1</sup>Department of Biology, Brigham Young University, Provo, Utah, U.S.A.

<sup>2</sup>CIBIO, Universidade do Porto, Vairão, Portugal

## ABSTRACT

Freshwater crayfish (Astacoidea) serve as model organisms for many diverse disciplines, from neurology to toxicology, and have been the focus of many physiological, ecological, and molecular-based studies. Although much of the recent work has focused on the evolutionary history, phylogeography, and conservation biology of freshwater crayfishes, estimations of their divergence times and radiations have never been made. Recently, divergence time estimations for decapods provided the first proposed molecular-timing hypothesis involving freshwater crayfish. In this study we focus specifically on estimating divergence among Astacoidea. We employ a Bayesian method implemented in multidivtime for timing estimation, calibrated with multiple fossils including a Parastacoidea fossil newly discovered in Australia. With our narrow taxonomic focus, we increase the accuracy and provide divergence estimations more specific to freshwater crayfish. Our molecular time estimation supports a late Permian to early Triassic divergence from Nephropoidea with radiation and dispersal before the breakup of Pangaea, as well as subsequent speciation and radiation prior to or directly associated with Gondwana and Laurasia disassembly. The breakup of Gondwana and Laurasia resulted in the separation of Parastacoidea and Astacoidea during the Jurassic. The hypothesized divergence and radiation of these two superfamilies are also supported by our molecular time estimations. For the three families of crayfish, we estimate the Astacidae radiation at ~153 million years ago (MYA), the Cambaridae radiation at ~90 MYA, and diversification of Parastacidae at ~161 MYA.

## 1 INTRODUCTION

Freshwater crayfish have a worldwide distribution, occurring on all continents except Antarctica and Africa excluding Madagascar. They are placed in the infraorder Astacoidea, which includes three superfamilies: 1) Astacoidea—Northern Hemisphere crayfish, 2) Parastacoidea—Southern Hemisphere crayfish, and 3) Nephropoidea—the clawed lobsters. The crayfish form a monophyletic group (Crandall et al. 2000b) and have ~640 described species (Crandall et al. 2008) with Nephropoidea, the clawed lobsters, hypothesized as their sister group (Crandall et al. 2000a). Parastacoidea contains one family, Parastacidae, with 15 genera (*Astacoides*, *Astacopsis*, *Cherax*, *Engaeus*, *Engaewa*, *Euastacus*, *Geocharax*, *Gramastacus*, *Ombrastacoides*, *Paranephrops*, *Parastacus*, *Samastacus*, *Spinastacoides*, *Tenuibranchiurus*, and *Virilastacus*) and 176 species. Astacoidea contains two families, Astacidae and Cambaridae. Astacidae has three genera (*Pacifastacus*, *Astacus*, *Austropotamobius*) (Hobbs 1974) to six genera (Starobogatov 1995), depending on whose taxonomy one prefers, and 16–39 species. Cambaridae has 2 subfamilies (Cambarellinae and Cambarinae) containing 11 genera (*Barbicambarus*, *Bouchardina*, *Cambarellus*, *Cambarus*, *Distocambarus*, *Fallicambarus*, *Faxonella*, *Hobbseus*, *Orconectes*, *Procambarus*, *Troglocambarus*), plus a distinct genus *Cambaroides* that appears to be more distantly related to these two subfamilies; Cambaridae has a total of 445 species (see Crandall & Buhay 2008 for a recent summary).

Freshwater crayfish relationships at higher taxonomic levels are well understood. The two superfamilies are monophyletic sister clades, and Parastacidae and Astacidae are monophyletic (Crandall et al. 2000b; Rode & Babcock 2003). Cambaridae is paraphyletic, as one of its genera, *Cambaroides*, is in a basal lineage to Astacidae and the rest of the Cambaridae genera (Braband et al. 2006; Crandall et al. 2000b). Most of the taxonomic relationships within Cambaridae are currently best explained by Hobbs' (1989) taxonomic revision. The following taxonomic groups within *Cambarinae* have been evaluated since Hobbs' (1989) revision: the genus *Orconectes* (Taylor and Knouft 2006); subgenus *Crockerinus* within *Orconectes* (Taylor and Hardman 2002); the subgenus *Scapulicambarus* within *Procambarus* (Busack 1989); and the subgenus *Aviticambarus* within *Cambarus* (Buhay et al. 2007). Within Astacidae, the taxonomy within *Astacus* and *Pacifastacus* is based on Hobbs' (1989) morphological taxonomic revision. The taxonomy within *Austropotamobius* was recently examined by Grandjean et al. (2000), Zaccara et al. (2004), and Fratini et al. (2004), all of whom reported multiple cryptic subspecies. However, Starobogatov (1995) provided a comprehensive overview of the Astacidae that resulted in 6 genera and 36 species, but his proposed taxonomy has not yet taken hold in the literature. The Astacidae in general is in need of a detailed examination to unify the diversity of ideas concerning its taxonomy.

The first comprehensive phylogenetic hypothesis of the Parastacoidea was morphologically based on male genitalia, cephalothorax, chelae, and body shape (Riek 1969). Studies that followed addressed the relations within this family using morphological, protein, and molecular data (Austin 1995; Crandall et al. 1995; Patak & Baldwin 1984; Patak et al. 1989; Riek 1972). These studies included limited sampling of genera and had conflicting results. The study by Crandall et al. (2000a) established well-supported relations within this family by analyzing 13 of the then 14 genera using mitochondrial DNA. Out of the now 15 genera in Parastacoidea, eight have been recently evaluated taxonomically and/or phylogenetically: *Engaewa* (by Horwitz and Adams 2000), *Cherax* (by Austin 1996), *Euastacus* (by Schull et al. 2005), two new genera *Spinastacoides* and *Ombrastacoides* (by Hansen and Richardson 2006), and *Engaeus*, *Geocharax*, and *Gramastacus* (by Schultz et al. 2007).

Through these recent studies, the problems of determining relationships among the freshwater crayfish become very apparent. Studies have not been fully comprehensive and have been limited in taxonomic sampling, due in part to the large number of freshwater crayfish taxa and their global distribution. The genetic and protein studies have shown high morphological and habitat variation within species and have demonstrated that convergent evolution is common (Braband et al. 2006; Crandall & Fitzpatrick 1996; Taylor & Hardman 2002). Additionally, these studies have revealed multiple cases of parphyly, discovery of cryptic species, and even some unsupported described species (e.g., Austin 1996; Grandjean et al. 2000; Hansen & Richardson 2006; Schull et al. 2005; Schultz et al. 2007; Crandall et al. 2008). As a result, Sinclair et al. (2004) proposed the completion of a worldwide phylogeny based on multiple mitochondrial and nuclear genes. Because of the group's extensive convergent evolutionary history, only through molecular analysis and full taxonomic coverage will it be possible to infer the relationships within this group. While this goal is yet to be achieved, here we report on a phylogenetic status of the major genera of freshwater crayfish and the associated divergence times to put such a phylogeny into a temporal perspective.

Recently, Porter et al. (2005) published a phylogeny and associated divergence time estimates for the decapods as a whole. This study was the first molecular-based time hypothesis that included the freshwater crayfish. The goal of that study was to estimate decapod divergences; hence, only two of their fossil calibrations came from within the infraorder Astacidea. Multiple studies have shown that the most important factor affecting molecular divergence time estimation is the number and distribution of the calibration points throughout the tree (Lee 1999; Porter et al. 2005; Thorne & Kishino 2002; Yang & Yoder 2003; Yoder & Yang 2000). In this study we focus specifically on estimating divergence among Astacidea. By including multiple fossil calibrations and a specific taxonomic focus we increase the accuracy and can provide divergence estimations more specific to freshwater crayfish events. The use of molecular-based divergence time estimates has improved the



understanding of the timing of evolutionary processes and events. A molecular time estimate for crayfish is particularly interesting because the current hypotheses of the divergence times correlates with estimates of the timing of the breakup of Pangaea and disassembly of Gondwana and Laurasia (Ahyong & O'Meally 2004; Crandall et al. 2000b; Porter et al. 2005; Rode & Babcock 2003). We test the hypotheses that freshwater crayfish diverged from *Nephropoidea* (clawed lobsters) during the Permian or Triassic, and that Parastacoidea (Southern Hemisphere) and Astacoidea (Northern Hemisphere) divergence occurred during the Jurassic (Ahyong & O'Meally 2004; Crandall et al. 2000b; Porter et al. 2005; Rode & Babcock 2003), using a comprehensive phylogeny at the genus level for the major lineages of freshwater crayfish.

## 2 METHODS

### 2.1 Taxon sampling, DNA extraction, PCR, and sequencing

Crayfish species were chosen to represent major crayfish lineages in order to date the divergence times of these major groups (Table 1). Multiple sequences were obtained from GenBank, and the remaining sequences were generated by Toon et al. (in prep.), as indicated by an asterisk in Table 1. Although specifics can be obtained from Toon et al. (this volume), crayfish collection, preservation, DNA extraction, and amplification were completed following protocols and methods described in Crandall & Buhay (2004) and Crandall & Fitzpatrick (1996) for 16S rDNA (~500 bp; Crandall & Fitzpatrick, 1996), 12S rDNA (~400; Mokady et al. 1999) and COI (~700 bp; Folmer et al. 1994), and two nuclear genes: 18S (~2,000 bp; Whiting et al. 1997) and 28S (~3,000 bp; Whiting et al. 1997).

### 2.2 Phylogenetic analyses

Astacoidea and Parastacoidea were aligned separately using MAFFT (Katoh et al. 2002; Katoh et al. 2005) implementing the G-INS-I alignment algorithm and then combined using the MAFFT profile alignment option with default parameters for each gene. *Homarus americanus* and *Sergio mericeae* were then aligned to the ingroup using MAFFT profile alignment for each gene. This multiple sequence alignment program has been shown to provide quick and accurate results by Notredame et al. (2000) and Katoh et al. (2005). The iterative algorithms used by MAFFT allow for repeatability of alignment. GBLOCKS 0.91b (Castresana 2000) was used to objectively trim sections of the alignment with questionable homology using the default parameter with the exception of the allowed gap positions parameter. The latter was set to allow gaps that are present in at least half of the sequences (Talavera & Castresana 2007). Models of evolution for each alignment were estimated in ModelTest (Posada & Crandall 1998) using the AIC criteria (Akaike 1973) to compare and choose best-fit models for the different gene partitions.

Phylogenies were estimated using maximum likelihood (ML) and Bayesian optimality criteria, with RAxML (Stamatakis 2006) and MrBayes (Ronquist & Huelsenbeck 2003), respectively (see Palero & Crandall, this volume, for a general description of these approaches). RAxML is a unique ML program in that it allows the use of multiple models, therefore giving better ML estimates. We partitioned the data set by gene and applied the model GTR+I+G to each gene allowing independent parameters to be estimated during analysis. We selected the tree with the best ML score after multiple independent runs with random starting positions and assessed confidence in nodal support through 1000 bootstrap pseudoreplications. Bayesian analysis was performed in MrBayes, in which four independent runs starting from random trees were run using the default flat priors for  $5 \times 10^6$  generations sampling every 100 generations. We also ran two independent MrBayes runs with the same settings using the best RAxML tree as a start tree. The negative log likelihood posterior distribution was checked for convergence and length needed for burn-in using the program Tracer

**Table 1.** Taxa and GenBank accession numbers associated with each sample. Asterisks (\*) indicate sequences from Toon et al. (submitted).

| Taxon   | Gene      |           |           |           |           |
|---|-----------|-----------|-----------|-----------|-----------|
|   | 12S       | 16S       | 18S       | 28S       | CO1       |
| <b>Astacidea</b> Latreille 1802                           |           |           |           |           |           |
| Astacoidea Latreille 1802                                 |           |           |           |           |           |
| <i>Astacus astacus</i> (Linnaeus 1758)                    | EU920881* | AF235983  | AF235959  | DQ079773  | AF517104  |
| <i>Cambarellus shufeldtii</i> (Faxon 1884)                | EU921117* | AF235986  | AF235962  | DQ079778  | EU921149* |
| <i>Cambaroides japonicus</i> (de Haan 1841)               | EU921118* | AF235987  | DQ079742  | DQ079779  | no seq    |
| <i>Cambarus maculatus</i> (Hobbs & Pflieger 1988)         | EU921119* | AF235988  | AF235964  | DQ079780  | no seq    |
| <i>Orconectes virilis</i> (Hagen 1870)                    | EU920900* | AF235989  | AF235965  | DQ079804  | AF474365  |
| <i>Pacifastacus leniusculus</i> (Dana 1852)               | EU921116* | AF235985  | AF235961  | DQ079806  | EU921148* |
| <i>Procambarus clarkii</i> (Girard 1852)                  | EU920901* | AF235990  | EU920952* | EU920970* | AY701195  |
| Parastacoidea (Huxley 1879)                               |           |           |           |           |           |
| <i>Astacoides betsileoensis</i> (Petit 1923)              | EU920882* | EU920912* | EU920955* | EU920992* | EU921146* |
| <i>Astacoides crosnieri</i> (Hobbs 1987)                  | EU921112* | EU921122* | EU921129* | EU921136* | EU921147* |
| <i>Astacopsis tricornis</i> (Clark 1936)                  | DQ006419  | DQ006548  | EU921123* | EU921135* | DQ006290  |
| <i>Cherax cairnsensis</i> (Riek 1969)                     | EU921113* | EU921120* | EU921124* | EU921132* | EU921113* |
| <i>Cherax quadricarinatus</i> (von Martens 1868)          | DQ006423  | DQ006552  | EU921125* | EU921139* | DQ006294  |
| <i>Engaeus fossor</i> (Erichson 1846)                     | EU921114* | EU921121* | EU921126* | EU921134* | EU921144* |
| <i>Euastacus sulcatus</i> (Riek 1951)                     | DQ006525  | DQ006651  | EU921127* | EU921133* | DQ006396  |
| <i>Geocharax gracilis</i> (Clark 1936)                    | EU921115* | AF235992  | AF235968  | EU921140* | EU921145* |
| <i>Paranephrops planifrons</i> (White 1842)               | DQ006544  | AF135995  | EU921128* | EU921141* | DQ006415  |
| <i>Omrastacoides huonensis</i> (Hansen & Richardson 2006) | EU920905* | AF135997  | EU920956* | EU920995* | EU921143* |
| <i>Parastacus brasiliensis</i> (von Martens 1869)         | EF599134  | AF175244  | EU921130* | EU921138* | EF599158  |
| <i>Samastacus spinifrons</i> (Phillipi 1882)              | EF599136  | AF175241  | EU921131  | EU921137  | EF599159  |
| Nephropoidea (Dana, 1852)                                 |           |           |           |           |           |
| <i>Homarus americanus</i> (H. Milne-Edwards 1837)         | DQ298427  | HAU11238  | AF235971  | DQ079788  | DQ889104  |
| Outgroup  |           |           |           |           |           |
| Thalassinidea   |           |           |           |           |           |
| Callianassoidea (Dana 1852)                               |           |           |           |           |           |
| <i>Sergio mericeae</i> (Manning & Felder 1995)            | EU920909* | DQ079733  | DQ079768  | DQ079811  | no seq    |

v1.4 (Rambaut & Drummond 2007) across all Bayesian runs. Converging MrBayes runs were combined after independent analysis and deletion of burn-in. Nodal confidence for the Bayesian trees was assessed using posterior probabilities compiled from the set of trees post-burn-in. We compared the support indices from our RAxML and MrBayes hypothesis and chose the phylogeny with the highest number of well-supported nodes considering bootstrap values  $\geq 70$  and Bayesian posterior probabilities  $\geq 95$  as high support for use in our molecular clock estimation.

### 2.3 Fossil calibrations

The fossil record is being continually updated, and relationships based on it are constantly being reanalyzed. The recent discovery of a new Australian fossil *Palaeoechinastacus australianus* (Martin et al. 2008) doubles the previously recorded geological time range of the family *Parastacidae* (Hasiotis 2002; Rode & Babcock 2003; Sokol 1987, 1988). Because fossil calibrations are a major source of error in molecular timing estimation, it is imperative to use multiple calibrations to get the best possible estimation, thus reducing the inherent amount of error associated with the fossil record (Table 2). Along with fossil calibrations, many studies have incorporated time estimations of vicariate events associated with the split in major land masses such as Pangaea, Laurasia, and Gondwana (Bocxlare et al. 2006; Porter et al. 2005). Our choice of Bayesian molecular time

**Table 2.** Fossil calibrations used for divergence time estimations, with the node referring to placement of the fossil on the crayfish chronogram.

| Taxonomy          | Species                                | Reference            | Geologic (MYA)                                      | Node |
|-------------------|--|----------------------|---|------|
| <b>Infraorder</b> |  |                      |   |      |
| Astacidea         |  |                      |   |      |
| <b>Family</b>     |  |                      |   |      |
| Chimaerastacidae  | <i>Chimaerastacus pacifluvialis</i>    | Amati et al. 2004    | Mid Triassic (Upper Ladinian) 227–234               | C1   |
| <b>Family</b>     |  |                      |   |      |
| Parastacidae      | <i>Palaeoechinastacus australianus</i> | Martin et al. 2008   | Early Cretaceous 106                                | C3   |
|                   | <i>Paranephrops fordycei</i>           | Feldmann & Pole 1994 | early middle Miocene (Otaian-Lillburnian) 21.7–12.7 | C4   |
| <b>Family</b>     |  |                      |   |      |
| Astacidae         | <i>Astacus licenti</i>                 | Van Straelen 1928    | Late Jurassic 144–159                               | C5   |
|                   | <i>Astacus spinostris</i>              | Imaizumi 1938        | Late Jurassic 144–159                               | C5   |
| <b>Family</b>     |  |                      |   |      |
| Cambaridae        | <i>Procambarus primaevus</i>           | Feldmann et al. 1981 | Late early Eocene 52.6–53.4                         | C6   |

Calibration C2 is 185 MYA, based on the splitting of Pangaea used as an upper limit

estimation requires that we have an estimation of at least one upper time limit (i.e., maximum age). Following Porter et al. (2005), we used the split of Pangaea at 185 MYA as an upper limit calibration for the divergence of the superfamilies Astacoidea and Parastacoidea (Crandall et al. 2000b). All other calibrations are estimated as the mean date of the fossil and set as the lower limit calibration indicating the absolute minimum age of the calibrated group (Porter et al. 2005). Additionally, we incorporated fossil calibrations for the origin of the family Astacidae and the split between Astacidea and Thalassinidea as the root node for our phylogenetic and molecular time estimation (Amati et al. 2004; Imaizumi 1938; Van Straelen 1928). Finally, we included three additional fossil calibrations: one to calibrate the genus *Procambarus* in Cambaridae and two to represent the family Parastacidae (Feldmann 2003; Feldmann et al. 1998; Martin et al. 2008). We agree with Porter et al. (2005) and others that trace fossil burrows are difficult to associate with crayfish with any amount of certainty (Babcock et al. 1998; Hasiotis 2002). Therefore, we chose to include only fossil records from descriptions of preserved animals. Our choice not to use trace fossils and to set each fossil calibration as the lower limit makes our estimate more conservative, while still allowing us to account for the fossil species existing for an undetermined amount of time before the actual fossilization event.

#### 2.4 Divergence time estimation

Freshwater crayfish divergence times were estimated using the multi-locus Bayesian method of Thorne and Kishino (2002) as implemented in the Multidivtime package (<http://statgen.ncsu.edu/thorne/multidivtime.html>). This approach was built on the continual improvements of molecular clock theory and applications (Kishino et al. 2001; Thorne et al. 1998). This method allows the use of multiple genes while not requiring a full taxa set for all genes included, does not assume a molecular clock in branch estimation, and allows for multiple calibrations. The use of multiple genetic loci and multiple fossil calibrations improves divergence times and rate estimations (Pérez-Losada et al. 2004; Porter et al. 2005; Thorne & Kishino 2002; Yang 2004; Yang & Yoder 2003; Yoder & Yang 2000). Multidivtime estimates times and rates by minimizing the discrepancies in

branch lengths and by minimizing rate changes over branches. This Bayesian method employs the rate evolution model of Thorne et al. (1998) and Kishino et al. (2001), which averages rates using a Markov chain Monte Carlo (MCMC) process.

We used three different parameter settings for Multidivtime. First, *rttm* and *rttmsd* (distribution of time separating the ingroup root from the present and the standard deviation, respectively) were set to 2.5 (250 MYA), and *rtrate* and *rtratesd* (prior evolutionary rate and standard deviation, respectively) were set to 0.0136 substitutions per million years. Second, *rttm* and *rttmsd* were set at 2.38 (238 MYA), and *rtrate* and *rtratesd* were set to 0.015 substitutions per million years, to see the effect of placing it closer to the age of the fossil calibration. Third, the *rttm* and *rttmsd* were set at 3.5 (350 MYA), and the *rtrate* and *rtratesd* were set to 0.0102 substitutions per million years to explore the effects of perturbations to the *rttm* setting. For each parameter setting, we applied two different burn-in period settings,  $10^4$  and  $10^6$  steps, combined with  $5 \times 10^5$  samples collected at every 100th cycle. The default settings were used for the rest of the required parameters. A total of 12 runs were completed with three independent random starts for each parameter and burn-in period setting. The three runs for each burn-in and parameter setting were checked, and the set with the most consistent estimations was chosen for our time estimation.

### 3 RESULTS

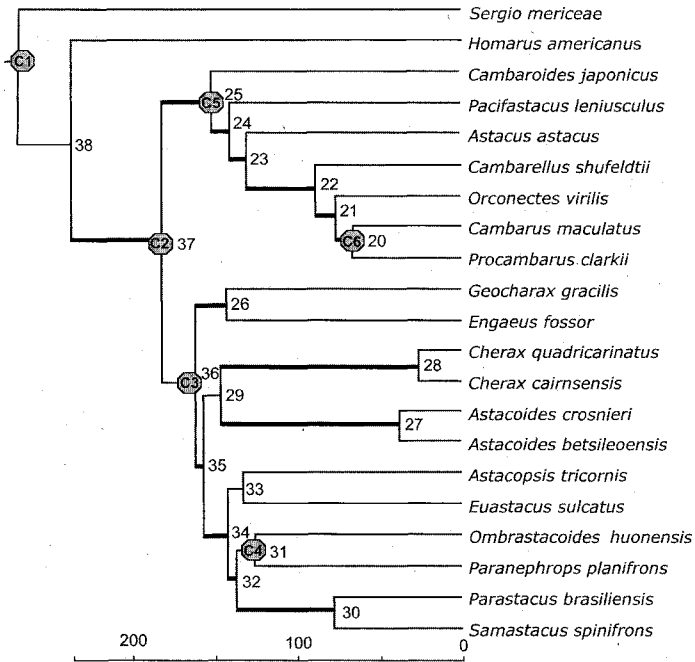
#### 3.1 Phylogenetics

All models selected by ModelTest were *nst*=6 with gamma and invariable sites (16S, 18S, and CO1=TVM+I+G; 28S=TrN+I+G; and 12S=GTR+I+G). There are a limited number of models in RAXML and MrBayes; therefore, the GTR+G+I model was chosen for each partition, allowing the respective programs to estimate the parameters during phylogenetic estimation. The RAXML best tree likelihood score was -24658.608503. Our RAXML tree compared to our Bayesian tree resulted in fewer nodes with high bootstrap support ( $\geq 70$ ) and Bayesian posterior probabilities ( $\geq 95$ ). Therefore, the Bayesian tree was used for the molecular divergent time estimation (Fig. 1). The relationships within Astacoidea are concordant with recent studies placing the genus *Cambariodes* basal to both Astacidae and Cambaridae. Although *Astacus* and *Pacifastacus* fall out independently, they both fall between the paraphyletic Cambaridae. Parastacids reflect the same relationships as in Crandall et al. (2000b), the most extensive study of the entire family to date.

#### 3.2 Divergence time estimations

Changing the *rttm* parameter, defined as the distribution of time separating the ingroup root from the present, to 2.386 and 3.5 hardly affected the results, with the largest difference in estimations being  $3 \times 10^5$  years (Table 3). Pérez-Losada et al. (2004) and Porter et al. (2005) found similar results using even larger perturbations and also reported a minimal effect on the overall time estimation. The burn-in period setting of  $10^6$  steps produced three nearly identical independent time estimations. From these three estimates, we chose the estimation with the smallest 95% posterior intervals for the chronogram (Fig. 1 & Table 3).

Our divergence time estimates between the crayfish lineages (Astacoidea and Parastacoidea) and Nephropoidea is  $\sim 239$  MYA (node 38). The divergence time estimates for the Northern Hemisphere families resulted in Astacidae divergence  $\sim 153$  MYA (node 25) being significantly older than Cambaridae divergence at  $\sim 90$  MYA (node 22). Parastacidae (the Southern Hemisphere crayfish) divergence time is estimated at  $\sim 161$  MYA (node 36) with the genera having much older divergence times than Northern Hemisphere crayfish.



**Figure 1.** Crayfish divergence time chronogram estimated with a Bayesian tree topology. Bolded branches indicate posterior probability of 1. Nodes labeled C1–C6 indicate locations of fossil calibration (Table 2). Node number refers to the estimated time and 95% posterior interval (Table 3).

**Table 3.** Node time estimations referring to crayfish chronogram (Fig. 1). Time is represented in MYA with 95% interval, standard deviation, and well-supported ML bootstrap and Bayesian posterior probability.

| Node | Time MYA | 95% Posterior Interval Lower | 95% Posterior Interval Upper | Standard Deviation | ML Bootstraps | Bayesian Posterior Probability |
|------|----------|------------------------------|------------------------------|--------------------|---------------|--------------------------------|
| 20   | 67.342   | 53.461                       | 96.797                       | 11.820             | 97            | 1                              |
| 21   | 77.593   | 56.790                       | 109.350                      | 13.966             | 100           | 1                              |
| 22   | 90.413   | 63.279                       | 125.150                      | 16.161             | 100           | 1                              |
| 23   | 132.263  | 100.796                      | 150.774                      | 13.184             | 82            | 1                              |
| 24   | 143.006  | 117.570                      | 154.648                      | 9.769              | -             | 1                              |
| 25   | 153.367  | 151.552                      | 157.798                      | 1.698              | 100           | 1                              |
| 26   | 144.531  | 128.907                      | 157.830                      | 7.363              | 99            | 1                              |
| 27   | 37.916   | 6.370                        | 73.685                       | 17.888             | 100           | 1                              |
| 28   | 25.915   | 12.882                       | 45.609                       | 8.481              | 100           | 1                              |
| 29   | 147.774  | 130.894                      | 161.587                      | 7.834              | -             | -                              |
| 30   | 78.473   | 40.3                         | 109.408                      | 17.520             | 100           | 1                              |
| 31   | 127.486  | 102.616                      | 149.049                      | 11.846             | -             | -                              |
| 32   | 138.331  | 115.897                      | 156.189                      | 10.326             | 87            | 1                              |
| 33   | 135.304  | 111.904                      | 153.525                      | 10.688             | 87            | -                              |
| 34   | 144.026  | 123.144                      | 160.854                      | 9.653              | 80            | 1                              |
| 35   | 158.120  | 143.756                      | 169.560                      | 6.56               | -             | 1                              |
| 36   | 161.875  | 150.093                      | 171.880                      | 5.542              | 100           | -                              |
| 37   | 183.459  | 179.650                      | 184.957                      | 1.446              | 100           | 1                              |
| 38   | 239.345  | 230.789                      | 258.697                      | 7.587              | -             | -                              |

## 4 DISCUSSION

### 4.1 *Phylogeny and divergence time estimations*

The phylogenetic results were consistent with the most recent molecular studies for freshwater crayfishes (Crandall et al. 2000b; Porter et al. 2005; Rode & Babcock 2003). The tree is generally well supported with the monophyly of the freshwater crayfish being recovered in 100% of the Bayesian posterior distributions. Most lineages within the Parastacidea are similarly supported, with a few of the deeper nodes having low support values. Our divergence time estimations support the divergent time hypotheses of Crandall et al. (2000b), Rode and Babcock (2003), Ahyong & O'Meally (2004), and Porter et al. (2005). In the most current divergence hypothesis, Porter et al. (2005) estimated the divergence between the crayfish lineages Astacoidea and Parastacoidea from Nephropoidea at ~278 MYA. Our estimation of ~239 MYA (node 38) differs probably because of the calibration of the node prior to this estimation in each study. Although both studies used *Chimaerastacus pacifluviialis* (C1) as a lower limit, we additionally used it as a guideline to estimate the time from the root to the tip, setting it at 250 MYA. Our estimation falls between their two estimations when they calibrated the previous node as a lower limit and when it was calibrated as an upper and lower bound. We estimate the Astacidae radiation at ~153 MYA (node 25), fitting within the range of the fossils used for calibration. We include *Cambaroides japonicus* in this estimation due to consistent placement of this genus within the Astacidae (Braband et al. 2006). Therefore, our estimate is significantly older than the Astacidae radiation estimate of Porter et al. (2005). Although their actual estimation is not reported, a visual inspection of the chronogram of Porter et al. (2005) reveals a similar estimation when including *Cambaroides japonicus*. The Cambaridae radiation was estimated at ~90 MYA (node 22), which coincides with Porter et al. (2005). These divergence estimates support the idea that Astacoidea diversified and was widespread before the split of Laurasia during the late Cretaceous (Owen 1976) ~65 MYA.

The diversification of Parastacidae was calibrated with a new fossil dated to 106 MYA (Martin et al. 2008), which resulted in our estimated divergence time of ~161 MYA. This divergence time suggests that older Parastacidae fossils are likely to be found in Australia. The first stages of Gondwana separation are estimated to have begun ~150 MYA with the separation of South America and Africa from Antarctica-India-Madagascar-Australia-New Zealand (Wit et al. 1999). Veevers (2006) estimates a later separation of Africa-India from Australia-Antarctica-South America at ~132 MYA. Regardless of the specific Gondwana breakup theory ascribed, the divergence time estimates between South America and Australia-New Zealand crayfish (node 32) and the Madagascar and Australian crayfish (node 29) can be explained by vicariance associated with the disassembly of Gondwana. The split between *Omrastacoides* (Australia) and *Paranephrops* (New Zealand) (node 32) ~127 MYA is also consistent in that vicariance may have happened before or in sync with this separation, which is commonly estimated at ~90 MYA, but rifting may have begun as early as ~110-115 (Stevens 1980, 1985).

### 4.2 *Interpreting results*

Molecular time estimations are prone to multiple errors, partially due to complete reliance on fossil calibration, in which there is an inherent amount of error, including incorrect assignment of fossils, error in chronological and date assignment, and introduced topological errors in the phylogenetic estimation (Graur & Martin 2004). With the amount of possible error, it is encouraging to get results that are consistent with the current fossil record and/or that are supported by theories of distribution and divergence. Although most time estimations were discussed as point estimation (the expected estimate of posterior distribution), readers should be aware of, and consider, the 95% posterior interval for all estimations. The Bayesian method employed is one of the few methods that allows the user to set minimum age fossil calibrations, but in doing so it results in a larger variance, increasing

the size of the posterior age interval. By setting fossil calibration intervals instead of minimum age estimates, you can effectively reduce the amount of variance resulting in a reduced size of the posterior age distribution. In the future, molecular clock estimates may consider using *Astacus licenti* and *Astacus spinirostris* fossil calibrations (C5) for Astacidae as an interval calibration instead of minimum age for two reasons. First, it is supported by two independent fossils. Second, our point estimation fits within the fossil estimated time interval. Including more upper limit calibrations or employing calibration intervals reduces the size of posterior interval estimates.

## 5 CONCLUSIONS

Our molecular clock estimation supports a late Permian to early Triassic divergence of freshwater crayfishes from Nephropoidea with radiation and dispersal before the breakup of Pangaea. Subsequent speciation and radiation prior to, or directly associated with, Gondwanan and Laurasian breakup resulted in the separation of the superfamilies Parastacoidea and Astacoidea during the Jurassic, thus supporting current divergent time estimations (Ahyong & O'Meally 2004; Crandall et al. 2000b; Porter et al. 2005; Rode & Babcock 2003). The hypothesized divergences and radiation of the two superfamilies attributed to the breakup of Laurasia and Gondwana are supported by our molecular time estimations. We do not expect this to be the last molecular divergence estimation for freshwater crayfishes, and we expect future estimates to improve in accuracy with the discovery of new fossils and new molecular dating techniques.

## ACKNOWLEDGEMENTS

We thank the wide variety of friends and colleagues who have helped us collect freshwater crayfish from around the world over the past 15 years. Likewise, this study was made possible by the exceptional undergraduates from Brigham Young University who have labored to collect DNA sequence data from freshwater crayfish. We thank Alicia Toon and two anonymous reviewers for helpful comments on this manuscript. Our work was supported by a grant from the US NSF EF-0531762 awarded to KAC.

## REFERENCES

- Ahyong, S.T. & O'Meally, D. 2004. Phylogeny of the Decapoda Reptantia: Resolution using three molecular loci and morphology. *Raffl. Bull. Zool.* 52: 673–693.
- Akaike, H. 1973. Information theory as an extension of the maximum-likelihood principle. In: Petrov, B. & Csake, F. (eds.), *Second International Symposium on Information Theory*. Budapest: Akademiai Kiado.
- Amati, L., Feldmann, R.M. & Zonneveld, J.P. 2004. A new family of Triassic lobsters (Decapoda: Astacidea) from British Columbia and its phylogenetic context. *J. Paleo.* 78: 150–168.
- Austin, C.M. 1995. Evolution in the genus *Cherax* (Decapoda: Parastacidae) in Australia: A numerical cladistic analysis of allozyme and morphological data. *Freshwater Crayfish* 8: 32–50.
- Austin, C.M. 1996. Systematics of the freshwater crayfish genus *Cherax* erichson (Decapoda: Parastacidae) in northern and eastern Australia: Electrophoretic and morphological variation. *Aust. J. Zool.* 44: 259–296.
- Babcock, L.E., Miller, M.F., Isbel, J.L., Collinson, J.W. & Hasiotis, S.T. 1998. Paleozoic-Mesozoic crayfish from Antarctica: Earliest evidence of freshwater Decapod crustaceans. *J. Geol.* 26: 539–542.
- Bocxlare, I.V., Roelants, K., Biju, S., Nagaraju, J. & Bossuyt, F. 2006. Late Cretaceous vicariance in Gondwanan amphibians. *PLoS ONE* 1: e74.

- Braband, A., Kawai, T. & Scholtz, G. 2006. The phylogenetic position of the east Asian freshwater crayfish *Cambaroides* within the northern hemisphere Astacoidea (Crustacea, Decapoda, Astacida) based on molecular data. *J. Zool. Syst. Evol. Res.* 44: 17–24.
- Buhay, J.E., Moni, G., Mann, N. & Crandall, K.A. 2007. Molecular taxonomy in the dark: Evolutionary history, phylogeography, and diversity of cave crayfish in the subgenus *Aviticambarus*, genus *Cambarus*. *Mol. Phylogenet. Evol.* 42: 435–488.
- Busack, C.A. 1989. Biochemical systematics of crayfishes of the genus *Procambarus*, subgenus *Scapulicambarus* (Decapoda: Cambaridae). *J. N. Amer. Benthol. Soc.* 8: 180–186.
- Castresana, J. 2000. Selection of conserved blocks from multiple alignments for their use in phylogenetic analysis. *Mol. Biol. Evol.* 17: 540–552.
- Crandall, K.A. & Buhay, J.E. 2004. Genomic databases and the tree of life. *Science* 306: 1144–1145.
- Crandall, K.A. & Buhay, J.E. 2008. Global diversity of crayfish (Astacidae, Cambaridae, and Parastacidae-Decapoda) in freshwater. *Hydrobiologia* 595: 295–301.
- Crandall, K.A., Fetzner, J.W., Jr., Jara, C.G. & Buckup, L. 2000a. On the phylogenetic positioning of the South American freshwater crayfish genera (Decapoda: Parastacidae). *J. Crust. Biol.* 20: 530–540.
- Crandall, K.A. & Fitzpatrick, J.F., Jr. 1996. Crayfish molecular systematics: Using a combination of procedures to estimate phylogeny. *Syst. Biol.* 45: 1–26.
- Crandall, K.A., Harris, D.J. & Fetzner, J.W. 2000b. The monophyletic origin of freshwater crayfishes estimated from nuclear and mitochondrial DNA sequences. *Proc. Roy. Soc. Lond. B. Biol. Sci.* 267: 1679–1686.
- Crandall, K.A., Lawler, S.H. & Austin, C. 1995. A preliminary examination of the molecular phylogenetic relationships of the crayfish genera of Australia (Decapoda: Parastacidae). *Freshwater Crayfish* 10: 18–30.
- Crandall, K.A., Robinson, H.W. & Buhay, J.E. 2008. Avoidance of extinction through nonexistence: The use of museum specimens and molecular genetics to determine the taxonomic status of an endangered freshwater crayfish. *Conservat. Genet.*: 10.1007/s10592-008-9546-9.
- Feldmann, R.M. 2003. The Decapoda: New initiatives and novel approaches. *J. Paleo.* 77: 1021–1038.
- Feldmann, R.M., Grande, L., Birkhimer, C.P., Hannibal, J.T. & McCoy, D.L. 1981. Decapod fauna of the Green River formation (Eocene) of Wyoming. *J. Paleo.* 55: 788–799.
- Feldmann, R.M. & Pole, M. 1994. A new species of *Paranephrops* White, 1842: A fossil freshwater crayfish (Decapoda: Parastacidae) from the Manuherikia group (Miocene), central Otago, New Zealand. *New Zeal. J. Geol. Geophys.* 37: 163–167.
- Feldmann, R.M., Vega, F.J., Applegate, S.P. & Bishop, G.A. 1998. Early Cretaceous arthropods from the Tlayua formation at Tepexi de Rodriguez, Puebla, Mexico. *J. Paleo.* 72: 79–90.
- Folmer, O., Black, M., Hoeh, W., Lutz, R. & Vrijenhoek, R. 1994. DNA primers for amplification of mitochondrial cytochrome c oxidase subunit i from diverse metazoan invertebrates. *Mol. Mar. Biol. Biotechnol.* 3: 294–299.
- Fratini, S., Zaccara, S., Barbaresi, S., Grandjean, F., Souty-Grosset, C., Crosa, G. & Gherardi, F. 2004. Phylogeography of the threatened crayfish (genus *Austropotamobius*) in Italy: Implications for its taxonomy and conservation. *Heredity* 94: 108–118.
- Grandjean, F., Harris, D.J., Souty-Grosset, C. & Crandall, K.A. 2000. Systematics of the European endangered crayfish species, *Austropotamobius pallipes* (Decapoda: Astacidae). *J. Crust. Biol.* 20: 522–529.
- Graur, D. & Martin, W. 2004. Reading the entrails of chickens: Molecular timescales of evolution and the illusion of precision. *Trends in Genetics* 20: 80–86.
- Hansen, B. & Richardson, A.M.M. 2006. A revision of the Taasmanian endemic freshwater crayfish genus *Parastacoides* (Crustacea: Decapoda: Parastacidae). *Invertebrate Systematics* 20: 713–769.



- Hasiotis, S.T. 2002. Where is the fossil evidence for Gondwanan crayfish? *Gondwana Res.* 5: 872–878.
- Hobbs, H.H., Jr. 1974. Synopsis of the families and genera of crayfishes (Crustacea:Decapoda). *Smithson. Contrib. Zoo.* 164: 1–32.
- Hobbs, H.H., Jr. 1989. An illustrated checklist of the American crayfishes (Decapoda: Astacidae, Cambaridae, and Parastacidae). *Smithson. Contrib. Zoo.* 480: 1–236.
- Horwitz, P. & Adams, M. 2000. The systematics, biogeography and conservation status of species in the freshwater crayfish genus *Engaewa* Riek (Decapoda: Parastacidae) from south-western Australia. *Invertebr. Taxon.* 14: 655–680.
- Imaizumi, R. 1938. Fossil crayfishes from Jehol. *Sci. Rep. Tokyo Imperial Univer., Sendai, Japan, Second Series* 19: 173–179.
- Katoh, K., Kuma, K., Toh, H. & Miyata, T. 2002. Mafft: A novel method for rapid multiple sequence alignment based on fast Fourier transform. *Nucleic Acids Res.* 30: 3059–3066.
- Katoh, K., Kuma, K., Toh, H. & Miyata, T. 2005. Mafft version 5: Improvement in accuracy of multiple sequence alignment. *Nucleic Acids Res.* 33: 511–518.
- Kishino, H., Thorne, J.L. & Bruno, W.J. 2001. Performance of a divergence time estimation method under a probabilistic model of rate evolution. *Mol. Biol. Evol.* 18: 352–361.
- Lee, M.S. 1999. Molecular clock calibrations and metazoan diversity. *J. Mol. Evol.* 49: 385–391.
- Martin, A.J., Rich, T.H., Poore, G.C.B., Schultz, M.B., Austin, C.M., Kool, L. & Vickers-Rich, P. 2008. Fossil evidence in Australia for oldest known freshwater crayfish of Gondwana. *Gondwana Res.* 14: 287–296.
- Mokady, M., Loya, Y., Achituv, Y., Geffen, E., Graur, D., Rozenblatt, S. & Brickner, I. 1999. Speciation versus phenotypic plasticity in coral inhabiting barnacles: Darwin's observations in an ecological context. *J. Mol. Evol.* 49: 367–375.
- Notredame, C., Higgins, D. & Heringa, J. 2000. T-coffee: A novel method for multiple sequence alignments. *J. Mol. Biol.* 302: 205–217.
- Owen, H.G. 1976. Continental displacement and expansion of the earth during the Mesozoic and Cenozoic. *Phil. Trans. Roy. Soc. Lond.* 281: 223–291.
- Palero, F. & Crandall, K.A. (this volume). Phylogenetic inference using molecular data. In: Martin, J.W., Crandall, K.A. & Felder, D.L. (eds.), *Crustacean Issues: Decapod Crustacean Phylogenetics*. Boca Raton, Florida: Taylor & Francis/CRC Press.
- Patak, A. & Baldwin, J. 1984. Electrophoretic and immunochemical comparisons of haemocyanins from Australian fresh-water crayfish (family parastacidae): Phylogenetic implications. *J. Crust. Biol.* 4: 528–535.
- Patak, A., Baldwin, J. & Lake, P.S. 1989. Immunochemical comparisons of haemocyanins of Australasian freshwater crayfish: Phylogenetic implications. *Biochem Systemat. Ecol.* 17: 249–252.
- Pérez-Losada, M., Høeg, J.T. & Crandall, K.A. 2004. Unraveling the evolutionary radiation of the thoracican barnacles using molecular and morphological evidence: A comparison of several divergence time estimation approaches. *Syst. Biol.* 53: 244–264.
- Porter, M.L., Pérez-Losada, M. & Crandall, K.A. 2005. Model based multi-locus estimation of decapod phylogeny and divergence times. *Mol. Phylogenet. Evol.* 37: 355–369.
- Posada, D. & Crandall, K.A. 1998. Modeltest: Testing the model of DNA substitution. *Bioinformatics* 14: 817–818.
- Rambaut, A. & Drummond, A. 2007. Tracer v1.4. Available from <http://slash/beast.bio.ed.ac.uk/Tracer>
- Riek, E.F. 1969. The Australian freshwater crayfish (Crustacea: Decapoda: Parastacidae), with descriptions of new species. *Aust. J. Zool.* 17: 855–918.
- Riek, E.F. 1972. The phylogeny of the Parastacidae (Crustacea: Astacoidea), and description of a new genus of Australian freshwater crayfishes. *Austral. J. Zool.* 20: 369–389.

- Rode, A.L. & Babcock, L.E. 2003. Phylogeny of fossil and extant freshwater crayfish and some closely related nephropid lobsters. *J. Crust. Biol.* 23: 418–435.
- Ronquist, F. & Huelsenbeck, J.P. 2003. MrBayes 3: Bayesian phylogenetic inference under mixed models. *Bioinformatics* 19: 1572–1574.
- Schull, H.C., Pérez-Losada, M., Blair, D., Sewell, K., Sinclair, E.A., Lawler, S., Ponniah, M. & Crandall, K.A. 2005. Phylogeny and biogeography of the freshwater crayfish *Euastacus* (Decapoda: Parastacidae) based on nuclear and mitochondrial DNA. *Mol. Phylogenet. Evol.* 37: 249–263.
- Schultz, M.B., Smith, S.A., Richardson, A.M.M., Horwitz, P., Crandall, K.A. & Austin, C.M. 2007. Cryptic diversity in *Engaeus* Erichson, *Geocharax* Clark and *Gramastacus* Riek (Decapoda: Parastacidae) revealed by mitochondrial 16S rDNA sequences. *Invertebrate Systematics* 21: 1–19.
- Sinclair, E.A., Fetzner, J.W., Jr., Buhay, J. & Crandall, K.A. 2004. Proposal to complete a phylogenetic taxonomy and systematic revision for freshwater crayfish (Astacida). *Freshwater Crayfish* 14: 1–9.
- Sokol, A. 1987. A note on the existence of Pre-Pleistocene fossils of parastacid crayfish. *Victorian Nat.* 104: 81–82.
- Sokol, A. 1988. Morphological variation in relation to the taxonomy of the *Destructor* group of the genus *Cherax*. *Invertebr. Taxon.* 2: 55–79.
- Stamatakis, A. 2006. Raxml-vi-hpc: Maximum likelihood-based phylogenetic analyses with thousands of taxa and mixed models. *Bioinformatics* 22: 2688–2690.
- Starobogatov, Y. 1995. Taxonomy and geographical distribution of crayfishes of Asia and East Europe (Crustacea Decapoda Astacoidei). *Arthropoda Selecta* 4: 3–25.
- Stevens, G.R. 1980. *New Zealand Adrift: The Theory of Continental Drift in a New Zealand Setting*. Wellington: A.H. & A.W. Reed.
- Stevens, G.R. 1985. *Lands in Collision: Discovering New Zealand's Past Geography*. Wellington: Science Information Publishing Centre, DSIR.
- Talavera, G. & Castresana, J. 2007. Improvement of phylogenies after removing divergent and ambiguously aligned blocks from protein sequence alignments. *Syst. Biol.* 56: 564–577.
- Taylor, C. & Hardman, M. 2002. Phylogenetics of the crayfish subgenus *Crockerinus*, genus, *Orconectes* (Decapoda: Cambaridae), based on cytochrome oxidase I. *J. Crust. Biol.* 22: 874–881.
- Taylor, C.A. & Knouft, J.H. 2006. Historical influences on genital morphology among sympatric species: Gonopod evolution and reproductive isolation in the crayfish genus *Orconectes* (Cambaridae). *Biol. J. Linn. Soc.* 89: 1–12.
- Thorne, J.L. & Kishino, H. 2002. Divergence time and evolutionary rate estimation with multilocus data. *Syst. Biol.* 51: 689–702.
- Thorne, J.L., Kishino, H. & Painter, I.S. 1998. Estimating the rate of evolution of the rate of molecular evolution. *Mol. Biol. Evol.* 15: 1647–1657.
- Toon, A., Finley, M., Staples, J. & Crandall, K.A. (this volume). Decapod phylogenetics and molecular evolution. In: Martin, J.W., Crandall, K.A. & Felder, D.L. (eds.), *Crustacean Issues: Decapod Crustacean Phylogenetics*. Boca Raton, Florida: Taylor & Francis/CRC Press.
- Van Straelen, V. 1928. On a fossil freshwater crayfish from eastern Mongolia. *Bulletin of the Geological Society of China* 7: 173–178.
- Veevers, J.J. 2006. Updated Gondwana (Permian–Cretaceous) earth history of Australia. *Gondwana Res.* 9: 231–260.
- Whiting, M.F., Carpenter, J.C., Wheeler, Q.D. & Wheeler, W.C. 1997. The Strepsiptera problem: Phylogeny of the holometabolous insect orders inferred from 18S and 28S ribosomal DNA sequences and morphology. *Syst. Biol.* 46: 1–68.

- Wit, M.D., Jeffrey, M., Bergh, H. & Nicolaysen, L. 1999. Gondawana reconstruction and dispersion. *Search and Discovery* Article #30001, <http://www.searchanddiscovery.net/documents/97019/index.htm>.
- Yang, Z. 2004. A heuristic rate smoothing procedure for maximum likelihood estimation of species divergence times. *Acta Zoolog. Sin.* 50: 645–656.
- Yang, Z. & Yoder, A.D. 2003. Comparison of likelihood and Bayesian methods for estimating divergence times using multiple gene loci and calibration points, with application to a radiation of cute-looking mouse lemur species. *Syst. Biol.* 52: 705–716.
- Yoder, A.D. & Yang, Z. 2000. Estimation of primate speciation dates using local molecular clocks. *Mol. Biol. Evol.* 17: 1081–1090.
- Zaccara, S., Stefani, F., Nardi, P.A. & Crosa, G. 2004. Taxonomic implications in conservation management of white-clawed crayfish *Austropotamobius pallipes* (Decapoda, Astacoidea). *Biol. Conservat.* 120: 1–10.