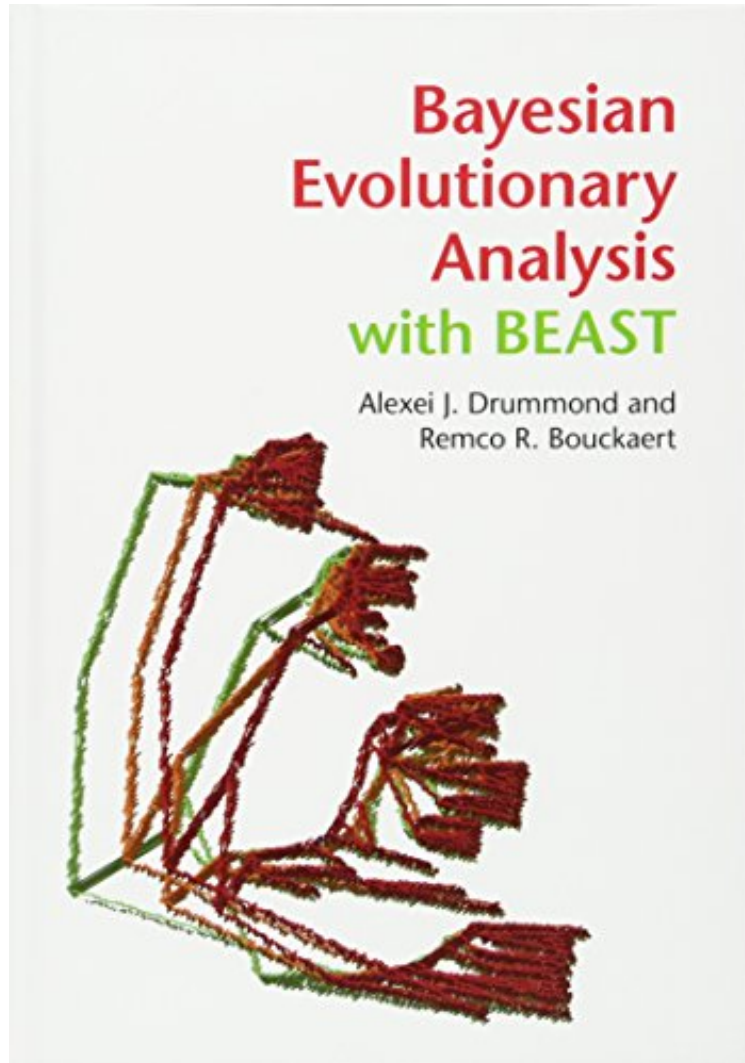


(Download pdf) Bayesian Evolutionary Analysis with BEAST

Bayesian Evolutionary Analysis with BEAST

Alexei J. Drummond, Remco R. Bouckaert
*ebooks | Download PDF | *ePub | DOC | audiobook*



 Download

 Read Online

#1212542 in Books 2015-10-08 Original language: English PDF # 1 9.72 x .63 x 6.851, 1.67 #File Name: 1107019656260 pages | File size: 42.Mb

Alexei J. Drummond, Remco R. Bouckaert : Bayesian Evolutionary Analysis with BEAST before purchasing it in order to gauge whether or not it would be worth my time, and all praised Bayesian Evolutionary Analysis with BEAST:

What are the models used in phylogenetic analysis and what exactly is involved in Bayesian evolutionary analysis using Markov chain Monte Carlo (MCMC) methods? How can you choose and apply these models, which parameterisations and priors make sense, and how can you diagnose Bayesian MCMC when things go wrong? These

are just a few of the questions answered in this comprehensive overview of Bayesian approaches to phylogenetics. This practical guide: Addresses the theoretical aspects of the field Advises on how to prepare and perform phylogenetic analysis Helps with interpreting analyses and visualisation of phylogenies Describes the software architecture Helps developing BEAST 2.2 extensions to allow these models to be extended further. With an accompanying website providing example files and tutorials (<http://beast2.org/>), this one-stop reference to applying the latest phylogenetic models in BEAST 2 will provide essential guidance for all users - from those using phylogenetic tools, to computational biologists and Bayesian statisticians.

"Want to construct a phylogeny, add in calibrated time points or work out the past history of an epidemic?The open source package BEAST has established itself as the industry standard for all this and more. This definitive book, explaining what is under the hood, how the user can customize extensions and most critically, a simple 'how to' users guide, is necessary reading for beginners and specialists alike." Laurence D. Hurst, University of Bath

"In concert with the dramatic improvements to DNA sequencing technology, Bayesian inference has revolutionized population genetics, phylogenetics, and divergence time estimation. A similar impact on epidemiology appears imminent via a suite of new Bayesian methods that incorporate host and pathogen DNA sequence data into established mathematical frameworks. This book is an accessible and thorough introduction to these Bayesian procedures. However, the book does far more than explain the theory. It also includes clear guides on how to use the BEAST 2 software for performing Bayesian analyses, and how to visualize the results. Because the software is designed to be extensible, the book instructs users to write their own code to supplement the diverse methods that are already implemented in BEAST 2. This book is timely and is written by two of the leaders of the field. I highly recommend it." Jeff Thorne, North Carolina State University

About the Author

Alexei J. Drummond is Professor of Computational Biology and Principal Investigator at the Allan Wilson Centre of Molecular Ecology and Evolution at the University of Auckland, New Zealand. He is the lead author of the BEAST software package and has gained a reputation in the field as one of the most knowledgeable experts for Bayesian evolutionary analyses.

Remco R. Bouckaert is a computer scientist with a strong background in Bayesian methods. He is the main architect of version 2 of BEAST and has been working on extensions to the BEAST software and other phylogenetics projects in Alexei Drummond's group at the University of Auckland.