Dear Editor,

Please consider our manuscript "babette: BEAUti 2, BEAST2 and Tracer for R" for publication in Methods in Ecology & Evolution.

BEAST2 is one of the most widely used phylogenetic software tools. One of the likely causes for its success is that it is accompanied with some programs that have a user-friendly graphical user interface. However, when doing an intensive theoretical exploration, with thousands of DNA alignments and/or model settings, a workflow of graphical user interfaces would be repetitive, plain dumb or impossible.

This manuscript describes babette, an R package that solves this problem. There have been many partial solutions to work with BEAST2 from R. Next to the partial solutions cited in this manuscript, there are many more (uncitable) code snippets that automate part of the BEAST2 workflow. babette unifies this scattered landscape of partial BEAST2 solutions, as it is the first R package that offers a complete end-to-end workflow to work with BEAST2 from R. This manuscript gives the reader an idea of the scope and ease of use of babette, thanks to the examples supplied.

We believe that our manuscript helps increase the joy and thoroughness of future studies within phylogenetics and should therefore appeal to a wide audience. Hence, we feel that Methods in Ecology & Evolution is the ideal platform for our manuscript.

Kind regards,

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