








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## Publisher Correction: Universal probabilistic programming offers a powerful approach to statistical phylogenetics

Fredrik Ronquist , Jan Kudlicka , Viktor Senderov , Johannes Borgström, Nicolas Lartillot, Daniel Lundén , Lawrence Murray, Thomas B. Schön  & David Broman

Correction to: *Communications Biology* <https://doi.org/10.1038/s42003-021-01753-7>, published online 24 February 2021.

Figures 1 and 2 and their associated captions were missing in the original PDF version of the Article. The online version of the Article was not affected. The errors have been corrected in the PDF version of the Article.

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