Systematic Biology -software for systematics and evolution

Running head PHYLODYNAMIC MODEL ADEQUACY **Title** Phylodynamic model adequacy using posterior predictive simulations **Authors** Sebastian Duchene^{1*}, Remco Bouckaert^{2,3}, David A. Duchene⁴, Tanja Stadler^{5,6}, Alexei J. Drummond² ¹Department of Biochemistry and Molecular Biology, Bio21 Molecular Science and Biotechnology Institute, University of Melbourne, Australia. ²Centre for Computational Evolution, University of Auckland, Auckland, New Zealand. ³Max Planck Institute for the Science of Human History, Jena, Germany. ⁴School of Life and Environmental Sciences, University of Sydney, Sydney, Australia. ⁵Department of Biosystems Science and Engineering, ETH Zurich, Basel, Switzerland. ⁶Swiss Institute of Bioinformatics, Lausanne, Switzerland. * To whom correspondence should be addressed: Sebastian Duchene

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Abstract

Rapidly evolving pathogens, such as viruses and bacteria, accumulate genetic

change at a similar timescale over which their epidemiological processes occur,

such that it is possible to make inferences about their infectious spread using

phylogenetic time-trees. For this purpose it is necessary to choose a

phylodynamic model. However, the resulting inferences are contingent on

whether the model adequately describes key features of the data. Model

adequacy methods allow formal rejection of a model if it cannot generate the

main features of the data. We present TreeModelAdequacy (TMA), a package for

the popular BEAST2 software, that allows assessing the adequacy of phylodynamic

models. We illustrate its utility by analysing phylogenetic trees from two viral

outbreaks of Ebola and H1N1 influenza. The main features of the Ebola data were

adequately described by the coalescent exponential-growth model, whereas the

H1N1 influenza data was best described by the birth-death SIR model.

Keywords

Model adequacy, Bayesian phylogenetics, Phylodynamics, Viral evolution, Posterior

predictive simulation, BEAST2

Phylogenetic trees depict the evolutionary relationships between groups of

organisms. In the context of infectious diseases, pathogen genetic data can be

used to infer such trees. By assuming a substitution model and including

independent information about time, one can calibrate the molecular clock to

obtain time-trees, where the branch lengths correspond to units of time, and

internal nodes of the tree represent the timing of divergence events. For rapidly

evolving viruses and bacteria it is possible to use the sampling times as time-

calibrations (Drummond et al. 2003; Rieux and Balloux 2016). In these organisms,

genetic change and epidemiological or ecological processes occur over a similar

timescale. Thus, time-trees can be informative about epidemiological dynamics, a

field of research known as phylodynamics (Holmes et al. 1993; Grenfell et al. 2004;

Kühnert et al. 2011; Volz et al. 2013).

Phylodynamic models describe the distribution of node times, branch lengths, and

sampling times. In full Bayesian phylogenetic analyses, the tree, parameters for

the molecular clock, the substitution model and the phylodynamic model can be

estimated simultaneously using molecular data. In this Bayesian framework, the

phylodynamic model is effectively a 'tree prior' (e.g. du Plessis and Stadler 2015).

The simplest phylodynamic models are the coalescent exponential-growth (CE)

and the coalescent constant-size (CC). These two models have very different

expectations about the shape of phylogenetic trees, with exponentially growing

populations tending to produce trees with longer external branches than those

evolving under constant population sizes (O'Meara, 2012; Volz et al., 2013).

Alternative phylodynamic models are the birth-death models, which include a

parameter to describe the sampling rate, so they have an expectation on the

number of taxa and their distribution over time (Stadler 2010; Stadler et al. 2012).

Choosing an appropriate model is important to draw reliable inferences from

parameters of interest. For instance, the CE and the constant birth-death (BD)

models can estimate the basic reproductive number, R_0 (Frost & Volz, 2010;

Stadler et al., 2012; Volz et al., 2013), which is defined as the average number of

new cases that a single case will generate over the course of its infection in a fully

susceptible population (Anderson and May 1979, 1992). Failing to account for

complex epidemiological dynamics can bias the estimate of this key parameter

(Stadler et al. 2014; Alkhamis et al. 2016; Ratmann et al. 2016). A Bayesian

approach to selecting a phylodynamic model is to estimate marginal likelihoods

for a pool of models and selecting that with the highest marginal likelihood

(Baele et al. 2012; Baele et al. 2016), but it is also possible to obtain weighted

averages of parameter estimates based on the support for each model (e.g. Baele,

Li, Drummond, Suchard, & Lemey, 2013; R. Bouckaert & Drummond, 2017;

Huelsenbeck, Larget, & Alfaro, 2004; Li & Drummond, 2012).

Model adequacy in phylogenetics

Model selection methods only allow a relative comparison of a set of models, but

they cannot determine whether any of the models in question could have

generated key features of the data at hand (i.e. absolute model fit). Such

information however is key to avoid unreliable inferences from a model and to

improve our understanding of the biological processes that produced the data.

Absolute model fit can be assessed via model adequacy methods, where a model

is considered 'adequate' if it is capable of generating the main features of the

empirical data. Consequently, model adequacy allows the user to formally reject a

model or to identify aspects of the data that are poorly described, instead of

ranking it with respect to other models, as is the case with model testing (e.g.

Goldman 1993; Bollback 2002; Ripplinger and Sullivan 2010; Brown 2014).

Model adequacy is typically conducted by fitting a model to the empirical data,

and generating synthetic data from the model in question, a procedure that is

similar to a parametric bootstrap (Goldman 1993). The adequacy of the model is determined depending on whether the synthetic data are similar to the empirical data, according to a descriptive test statistic (Gelman and Shalizi 2013; Gelman et al. 2014). The test statistics should summarise key aspects of the data or a combination of the data and parameter estimates (Gelman et al. 1996). Examples of test statistics that have been used to assess the substitution model include the multinomial likelihood or a measure of compositional homogeneity (Goldman 1993; Huelsenbeck et al. 2001; Foster 2004). The joint clock model and tree prior key can be assessed using the expected number of substitutions in individual branch lengths of the tree as test statistics (Duchêne et al. 2015). For DNA barcoding the number of OTUs and multinomial likelihood have been shown to be effective test statistics (Barley and Thomson 2016). Phylodynamic and diversification models are fitted to phylogenetic trees and their parameters depend on the distribution of nodes, such that some useful test statistics include the ratio of external to internal branch lengths, the tree height, and measures of phylogenetic tree imbalance (Revell et al. 2005, 2008; Drummond and Suchard 2008; Höhna et al. 2015). Clearly, designing test statistics is not trivial, but they should attempt to explicitly test some of the assumptions of the model. For

example, the CE and CC models have different expectations of the ratio of

external to internal branch lengths, such that this may be a useful test statistic.

Bayesian model adequacy

Bayesian model adequacy consists of a posterior predictive framework (Rubin

1981, 1984; Bollback 2002; Brown 2014a, 2014b; Lewis et al. 2014; Höhna et al.

2017). The posterior distribution of the model in question is approximated given

the empirical data, for example using Markov chain Monte Carlo (MCMC).

Samples from the MCMC are drawn to simulate data sets under the model used

for the empirical analysis. For example, the posterior distribution of the growth

rate and population size parameters of the CE model can be sampled to simulate

phylogenetic trees. Such simulations are known as posterior predictive

simulations. Test statistics are then calculated for every posterior predictive

simulation (i.e. for every simulated tree) to generate a distribution of values

according to the model. A posterior predictive probability, similar to the

frequentist ρ -value, can be calculated by determining where the value of the test

statistic for the empirical data (i.e. the empirical phylogenetic tree) falls with

respect to the posterior predictive distribution (Gelman et al. 2014). Following

Gelman et al. (2014) we refer to the posterior predictive probability as p_B to

differentiate it from the frequentist p-value. A useful quideline to determine

whether the model is adequate is to determine whether a test statistic within the

95% credible interval (Bollback 2002; Brown 2014a). This approach is sometimes

conservative, particularly when test statistics do not follow a Gaussian distribution,

and other methods of calculating posterior predictive probabilities are also

possible (Gelman et al. 2014; Höhna et al. 2017). Combining multiple test statistics

leads to multiple testing, which can be addressed by using a multivariate p_B

values (Drummond and Suchard 2008). However, Gelman et al. (2014) suggest

considering each test statistically separately to assess individual aspects of the

model and the data, which is the approach taken here.

Bayesian model adequacy is similar to Approximate Bayesian Computation (ABC)

techniques in that both methods use test statistics from simulated data. The aim

of ABC is to approximate the posterior by comparing test statistics from

simulations from the prior and the empirical data (Csilléry et al. 2010; Ratmann et

al. 2012; Poon 2015), which sometimes leads to biases for model testing (Robert

et al. 2011). In contrast, in model adequacy the simulations are generated from

the posterior distribution and they are not used to approximate the posterior. In

spite of these differences, test statistics developed for ABC can be useful to assess

model adequacy.

"TreeModelAdequacy" package in BEAST2

We implemented a computational framework to assess the adequacy of

phylodynamic models as a package for BEAST2 (Bouckaert et al., 2014). Analyses

as outlined in Fig 1 are easy to set up through BEAUti, the graphical user

interface for BEAST, to generate an xml file with the tree, the model, and test

statistics. Our package, TreeModelAdequacy (TMA), takes a tree with branch

lengths proportional to time. The tree can be a summary tree from BEAST2, or

estimated using a different method.

We fit phylodynamic models available in BEAST2 by approximating the posterior

distribution of the parameters of the model using MCMC. To generate the

posterior predictive simulations, we draw random samples from the posterior for

the parameters from the MCMC after removing the burn-in phase, and we

simulate phylogenetic trees using stochastic simulations and master equations

using MASTER (Vaughan and Drummond 2013) or the coalescent simulator in

BEAST2. The last step consists in calculating test statistics for the empirical data

and for the posterior predictive simulations, which depends on the TreeStat2

package (available at http://github.com/alexeid/TreeStat2). The user can select a

large number of test statistics (Supplementary material). At the end of the

analysis, p_B values and quantiles for the posterior predictive distribution are

shown, but they can also be visualised in Tracer (available at:

http://beast.bio.ed.ac.uk/tracer) or using an R script included in the package. At

present, the range of phylodynamic models that can be assessed includes the CC

and CE coalescent models, the constant BD with serial sampling (Stadler et al.

2012), and the birth-death susceptible-infected-recovered model (BDSIR; Kühnert

et al. 2014).

Our implementation allows parallelisation of the tree simulation step, which can

increase computational speed when the simulation conditions require extensive

calculations. This step can also be conducted independently on a computer

cluster. Our standalone application TreeModelAdequacyAnalyser can also

compute test statistics and p_B values, even for trees generated in a different

program than BEAST2, given that the posterior predictive trees are provided. TMA

is open source and freely available under a LGPL licence. It can be downloaded

from BEAUTi2 (part of BEAST2), and the documentation and example files are

available at: http://github.com/sebastianduchene/tree_model_adequacy.

To verify our implementation, we conducted a simple simulation experiment. We

simulated 100 trees under each of the four phylodynamic models (CC, CE, BD,

and BDSIR) using BEAST2 and MASTER and analysed them with the matching

model. We assessed their adequacy according to nine test statistics

(Supplementary material). The parameters for our simulations were based on

analyses of 72 whole genome sequences of Ebola virus (Gire et al. 2014). We

found that the p_B values for all test statistics were between 0.025 and 0.975 for

about 95% of each set of simulations, indicating that our implementation is

correct (Supplementary material).

Phylodynamic model adequacy in empirical virus data

West African Ebola virus

We obtained a phylogenetic tree inferred in a previous study from 72 Ebola virus

whole genome samples collected during the 2013-2016 epidemic (Gire et al.

2014). The samples were collected from May to July 2014 in Sierra Leone. These

data have been used in previous studies to estimate epidemiological parameters,

with estimates of R_0 ranging from 1.5 to 2.5, depending on the phylodynamic

model (Stadler et al., 2014; Volz & Pond, 2014). An important consideration about

our analysis is that we assume that the tree topology and divergence time

estimates are sufficiently accurate and that the data are informative.

We inferred phylodynamic parameters for the Ebola virus tree using four models;

CC, CE, the BD, and the BDSIR (Kühnert et al. 2014). The CE, BD, and BDSIR

models can estimate R_{θ} if information about the sampling process (for the birth-

death models) or present number of infected individuals (for the coalescent) is

available. In this case, we assumed that the sampling proportion was 0.7 (Gire et

al. 2014) by fixing this parameter in the BD and BDSIR models. For the remaining

parameters of these two models, we used the same prior distributions as in a

previous analysis of these data (Stadler et al. 2014). For the CE model we used a

Laplace distribution and a 1/x distribution as priors for the growth rate and for

the effective population size, respectively. We ran an MCMC of 10^7 steps and

generated 1,000 posterior predictive simulations, and we computed four test

statistics.

To compare the different models, we calculated p_{β} for four test statistics; the tree

height, the slope ratio of a lineages-through-time (LTT) plot, the ratio of external

to internal branch lengths, and the Colless index of phylogenetic imbalance (Fig.

2; Supplementary material). Importantly, the slope ratio of the LTT plot has been

found to be informative for inferring epidemiological parameters using ABC

(Saulnier et al. 2017).

In the CC model the p_B was < 0.05 for all test statistics, with the exception of the

tree height at 0.06 (Fig. 2 and Supplementary material Fig. S1). The CE and BD

models described these data better, with most p_{β} values between 0.11 and 0.56.

The ρ_p value for the Colless index in the CE model was the lowest for both of

these models, at 0.04. The BDSIR model had overall low p_B values, from to 0.01 to

0.18, with the lowest values found for the ratio of external to internal branch

lengths (0.03) and for the slope ratio of the LTT plot (0.01). The fact that most ρ_{β}

values for the CE and BD models frequently fall near the centre of the posterior

predictive distributions is consistent with the rapid spread that Ebola virus was

undergoing at the time the sequences were collected.

For the CE, BD, and BDISIR models we can estimate R_{θ} and the infectious period,

 $1/\delta$. The R_0 median estimates were: 1.6 (95% credible interval (CI): 1.12 – 2.2) for

the BD, 1.21 (95% CI: 1.1 – 1.5), for the CE, and 1.59 (95% CI: 1.22 – 1.94) for the

BDSIR. Estimates for the infectious period in calendar days were: 5.46 (95% CI:

4.14 – 7.24) for the BD, 2.90 (95% CI: 1.75 – 5.50) for the CE, and 5.21 (95% CI:

4.24 – 7.02) for the BDSIR. The estimates from these models were very similar and

overlapped with those from previous studies (Stadler et al. 2014). Although the

BDSIR did not capture some of the main features of these data, this model is

similar to the BD when the number of susceptible individuals is very large, which

probably explains the overlap in R_0 estimates between these models.

2009 H1N1 Influenza

We obtained a phylogenetic tree from a previous study (Hedge et al. 2013), which

was estimated from 328 whole genome samples from the 2009 H1N1 Influenza

pandemic. The samples were collected from April to December 2009, such that

they encompass a large portion of the duration of the pandemic. We used a

similar method as for the Ebola virus phylogenetic tree to fit the four

phylodynamic models (CC, CE, BD, and BDSIR). However, instead of fixing the

sampling proportion we used an informative prior distribution of the infectious

period via the becomeUnifectiousRate parameter, with a normal distribution of

mean 85 and standard deviation of 15 (corresponding to an infectious period of

about 4.45 days).

The CC and CE models had p_B values of 0.00 for all four test statistics, such that

they did not adequately describe any of these aspects of the tree. The BD model

had p_B values of 0.53 and 0.09 for the tree height and the slope ratio of the LTT

plot, and of 0.00 for the ratio of external to internal branch lengths (Fig. 2 and

Supplementary material Fig. S2). In contrast, the BDSIR model overall described

the H1N1 tree better overall than the other three models, with p_{β} values of

between 0.07 and 0.44. This result is consistent with the sampling time of the

data, which includes the start of the pandemic and the decline in the number of

infections towards the end of the year. We calculated an R_0 of mean 3.01 (95% CI:

2.5 – 3.7) at the start of the pandemic in January that declined to R_0 < 1 around

June, when infectious spread was lower. This estimate is similar to those made in

previous studies based on census data (Forsberg White et al. 2009), but in the

higher range of those based on the CE model for samples collected in early

stages of the pandemic (e.g. Hedge et al. 2013). For comparison, the R_0 estimate

from the BD model, which appeared inadequate, was substantially lower, with a

mean of 1.02 (95% CI: 1.00 – 1.03).

Conclusion

Model adequacy methods are useful to understand the biological processes that

generate the data, such as the evolutionary branching process. For example, our

approach reveals that in July of 2014 the West African Ebola outbreak was still

growing exponentially and that the 2009 H1N1 influenza virus pandemic had

evidence of a depletion of susceptible individuals in December. In some cases,

identifying models that indadequately describe key aspects of the data may

improve estimates of parameters of interest, such as R_0 in our H1N1 influenza

analyses. One consideration of our approach is that it requires an accurate

estimate of a single phylogenetic tree. Clearly, the phylogenetic tree should be

inferred using informative sequence data and the sensitivity to the prior should

be carefully examined (Ritchie et al. 2016; Boskova et al. 2018; Möller et al. 2018),

which is also the case for any Bayesian analysis. For example, a tree estimated

from uninformative sequence data will be driven by the prior and will necessarily

appear to be adequately described by the matching model, potentially leading to

increased rates of type 2 errors. A limitation of our method is that it does not

account for phylogenetic uncertainty, which can be addressed by comparing sets

of trees from the posterior with those from the posterior predictive distribution.

However, this approach will require the development of new test statistics and

model assessment criteria. Model adequacy in phylogenetics will benefit from

further development of methods to assess more sophisticated phylodynamic

models, such as those that account for population structure (Kühnert et al. 2016;

Müller et al. 2017a, 2017b; Volz and Siveroni 2018), and techniques to improve

the interpretation of ρ_B values for test statistics that are not normally distributed,

such at the Colless index. Model adequacy software, such as TMA, will be key to

address these questions.

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Conflict of interest

The authors gave final approval to the manuscript and have no conflict of interest

to declare.

Author contributions

SD, RB, and AJD wrote the computer code. SD analysed the data. TS, SD, and AJD

designed the experiments. SD wrote the manuscript with input from all the

authors.

Figure Legends

Fig 1. Posterior predictive simulation framework implemented in the

TreeModelAdequacy package. Step (1) consists in Bayesian analysis in BEAST

under model M to estimate the posterior distribution of parameters η , shown with

the arrow and posterior density in orange. In step (2) samples from the posterior

are drawn to simulate phylogenetic trees, known as posterior predictive

simulations, using MASTER as shown by the green arrow. In step (3) the posterior

predictive simulations are analysed in TreeStat to generate the posterior

predictive distribution of test statistic T_{a} , shown by the blue arrow and probability

density. Finally, T_a is also computed for the tree from the empirical data using

TreeStat, shown by the red arrow, to calculate a posterior predictive probability

 (p_B) . Test statistics and p_B values can also be computed for trees generated in

other programs using TreeModelAdequacyAnalyser, given that the tree from the

empirical data and the posterior predictive simulations are provided.

Fig 2. Model adequacy results for the two empirical data sets, West African Ebola

and 2009 H1N1 influenza. The histograms show the distribution of two test

statistics, the tree height and the slope ratio of the lineages-through-time (LTT)

plot, for the posterior predictive simulations. The black points also show the

distribution of the test statistics, and they have been jittered along the y-axis to

improve visualisation. The red lines denote the value for the tree estimated from

the empirical data. The posterior predictive p-value, p_B , is shown for each test

statistic. A feature of the empirical phylogenetic tree described by a test statistic,

such as the tree height, is considered adequately described by the model if the

empirical value falls within the 95% quantile range of the posterior predictive

distribution, such that the $p_B > 0.05$. Two more test statistics were computed, the

ratio of external to internal branch lengths and the Colless index, which are

shown in Supplementary material Fig. S1. Note that for the H1N1 influenza

analyses, the values are shown in a log₁₀ scale.

Supplementary figure captions

Fig S1. Model adequacy results for Ebola virus data with four test statistics. The

colours and symbols match those in Fig 2.

Fig S2. Model adequacy results for H1N1 influenza data with four test statistics.

The colours and symbols match those in Fig 2. Note that the values are shown in

log₁₀ scale.

References

- Alkhamis M.A., Perez A.M., Murtaugh M.P., Wang X., Morrison R.B. 2016.

 Applications of Bayesian phylodynamic methods in a recent US porcine reproductive and respiratory syndrome virus outbreak. Front. Microbiol. 7:67.
- Anderson R.M., May R.M. 1979. Population biology of infectious diseases: Part I. Nature. 280:361.
- Anderson R.M., May R.M. 1992. Infectious diseases of humans: dynamics and control. Oxford university press.
- Baele G., Lemey P., Bedford T., Rambaut A., Suchard M.A., Alekseyenko A. V. 2012. Improving the accuracy of demographic and molecular clock model comparison while accommodating phylogenetic uncertainty. Mol. Biol. Evol. 29:2157–2167.
- Baele G., Lemey P., Suchard M.A. 2016. Genealogical working distributions for Bayesian model testing with phylogenetic uncertainty. Syst. Biol. 65:250–264.
- Baele G., Li W.L.S., Drummond A.J., Suchard M.A., Lemey P. 2013. Accurate model selection of relaxed molecular clocks in bayesian phylogenetics. Mol. Biol. Evol. 30:239–243.
- Barley A.J., Thomson R.C. 2016. Assessing the performance of DNA barcoding using posterior predictive simulations. Mol. Ecol. 25:1944–1957.
- Bollback J.P. 2002. Bayesian model adequacy and choice in phylogenetics. Mol. Biol. Evol. 19:1171–1180.
- Boskova V., Stadler T., Magnus C. 2018. The influence of phylodynamic model specifications on parameter estimates of the Zika virus epidemic. Virus Evol. 4:vex044.

- Bouckaert R., Drummond A.J. 2017. bModelTest: Bayesian phylogenetic site model averaging and model comparison. BMC Evol. Biol. 17:42.
- Bouckaert R., Heled J., Kühnert D., Vaughan T., Wu C.-H., Xie D., Suchard M.A., Rambaut A., Drummond A.J. 2014. BEAST 2: a software platform for Bayesian evolutionary analysis. PLOS Comput. Biol. 10:e1003537.
- Brown J.M. 2014a. Detection of implausible phylogenetic inferences using posterior predictive assessment of model fit. Syst. Biol. 63:334–348.
- Brown J.M. 2014b. Predictive approaches to assessing the fit of evolutionary models. Syst. Biol. 63:289–292.
- Csilléry K., Blum M.G.B., Gaggiotti O.E., François O. 2010. Approximate Bayesian computation (ABC) in practice. Trends Ecol. Evol. 25:410–418.
- Drummond A.J., Pybus O.G., Rambaut A., Forsberg R., Rodrigo A.G. 2003. Measurably evolving populations. Trends Ecol. Evol. 18:481–488.
- Drummond A.J., Suchard M.A. 2008. Fully Bayesian tests of neutrality using genealogical summary statistics. BMC Genet. 9:68.
- Duchêne D.A., Duchêne S., Holmes E.C., Ho S.Y.W. 2015. Evaluating the adequacy of molecular clock models using posterior predictive simulations. Mol. Biol. Evol. 32:2896-2995.
- Forsberg White L., Wallinga J., Finelli L., Reed C., Riley S., Lipsitch M., Pagano M. 2009. Estimation of the reproductive number and the serial interval in early phase of the 2009 influenza A/H1N1 pandemic in the USA. Influenza Other Respi. Viruses. 3:267–276.
- Foster P.G. 2004. Modeling compositional heterogeneity. Syst. Biol. 53:485–495. Frost S.D.W., Volz E.M. 2010. Viral phylodynamics and the search for an "effective

- number of infections." Philos. Trans. R. Soc. London B Biol. Sci. 365:1879–1890.
- Gelman A., Carlin J.B., Stern H.S., Dunson D.B., Vehtari A., Rubin D.B. 2014. Model checking. Bayesian data analysis. Boca Raton, Florida: CRC press Boca Raton, FL. p. 141–163.
- Gelman A., Meng X.-L., Stern H. 1996. Posterior predictive assessment of model fitness via realized discrepancies. Stat. Sin.:733–760.
- Gelman A., Shalizi C.R. 2013. Philosophy and the practice of Bayesian statistics. Br. J. Math. Stat. Psychol. 66:8–38.
- Gire S.K., Goba A., Andersen K.G., Sealfon R.S.G., Park D.J., Kanneh L., Jalloh S., Momoh M., Fullah M., Dudas G. 2014. Genomic surveillance elucidates Ebola virus origin and transmission during the 2014 outbreak. Science (80-.). 345:1369–1372.
- Goldman N. 1993. Simple diagnostic statistical tests of models for DNA substitution. J. Mol. Evol. 37:650–661.
- Grenfell B.T., Pybus O.G., Gog J.R., Wood J.L.N., Daly J.M., Mumford J.A., Holmes E.C. 2004. Unifying the epidemiological and evolutionary dynamics of pathogens. Science. 303:327–332.
- Hedge J., Lycett S.J., Rambaut A. 2013. Real-time characterization of the molecular epidemiology of an influenza pandemic. Biol. Lett. 9:20130331.
- Höhna S., Coghill L.M., Mount G.G., Thomson R.C., Brown J.M. 2017. P3:

 Phylogenetic Posterior Prediction in RevBayes. Mol. Biol. Evol.
- Höhna S., May M.R., Moore B.R. 2015. TESS: an R package for efficiently simulating phylogenetic trees and performing Bayesian inference of lineage

- diversification rates. Bioinformatics. 32:789-791.
- Holmes E.C., Zhang L.Q., Simmonds P., Rogers A.S., Brown A.J.L. 1993. Molecular investigation of human immunodeficiency virus (HIV) infection in a patient of an HIV-infected surgeon. J. Infect. Dis. 167:1411–1414.
- Huelsenbeck J.P., Larget B., Alfaro M.E. 2004. Bayesian phylogenetic model selection using reversible jump Markov chain Monte Carlo. Mol. Biol. Evol. 21:1123–1133.
- Huelsenbeck J.P., Ronquist F., Nielsen R., Bollback J.P. 2001. Bayesian inference of phylogeny and its impact on evolutionary biology. Science (80-.). 294:2310–2314.
- Kühnert D., Stadler T., Vaughan T.G., Drummond A.J. 2014. Simultaneous reconstruction of evolutionary history and epidemiological dynamics from viral sequences with the birth–death SIR model. J. R. Soc. Interface. 11:20131106.
- Kühnert D., Stadler T., Vaughan T.G., Drummond A.J. 2016. Phylodynamics with migration: A computational framework to quantify population structure from genomic data. Mol. Biol. Evol. 33:2102–2116.
- Kühnert D., Wu C.H., Drummond A.J. 2011. Phylogenetic and epidemic modeling of rapidly evolving infectious diseases. Infect. Genet. Evol. 11:1825–1841.
- Lewis P.O., Xie W., Chen M.-H., Fan Y., Kuo L. 2014. Posterior predictive Bayesian phylogenetic model selection. Syst. Biol. 63:309–321.
- Li W.L.S., Drummond A.J. 2012. Model averaging and Bayes factor calculation of relaxed molecular clocks in Bayesian phylogenetics. Mol. Biol. Evol. 29:751–761.

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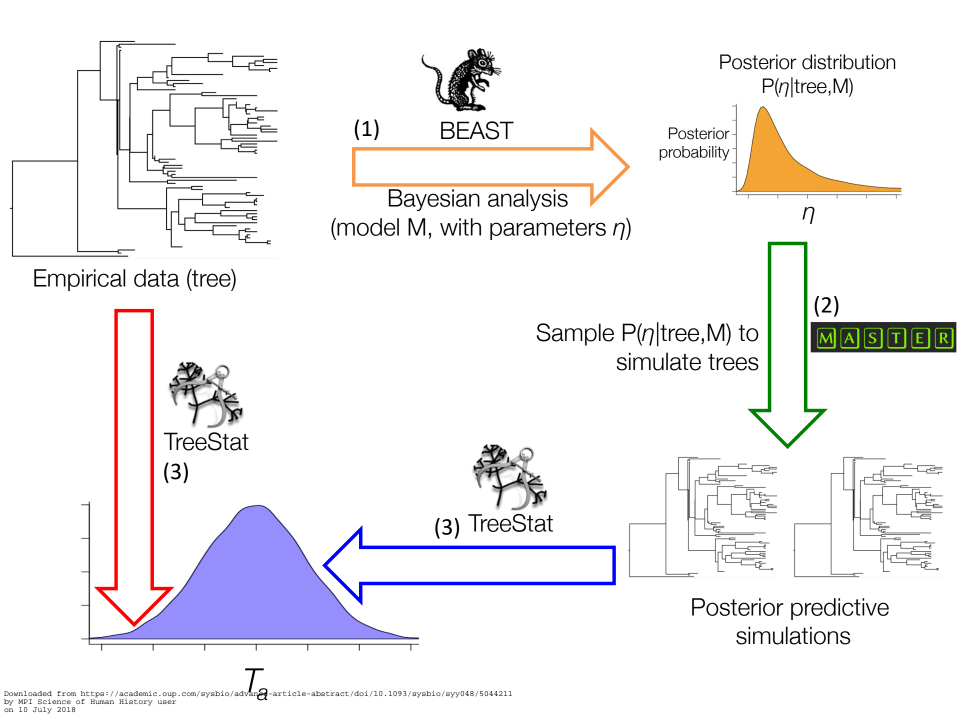
- Möller S., du Plessis L., Stadler T. 2018. Impact of the tree prior on estimating clock rates during epidemic outbreaks. Proc. Natl. Acad. Sci. USA. 115:4200–4205.
- Müller N.F., Rasmussen D.A., Stadler T. 2017a. MASCOT: Parameter and state inference under the marginal structured coalescent approximation. bioRxiv.:188516.
- Müller N.F., Rasmussen D.A., Stadler T. 2017b. The Structured Coalescent and Its Approximations. Mol. Biol. Evol. 34:2970–2981.
- O'Meara B.C. 2012. Evolutionary inferences from phylogenies: a review of methods. Annu. Rev. Ecol. Evol. Syst. 43:267–285.
- du Plessis L., Stadler T. 2015. Getting to the root of epidemic spread with phylodynamic analysis of genomic data. Trends Microbiol. 23:383–386.
- Poon A.F.Y. 2015. Phylodynamic inference with kernel ABC and its application to HIV epidemiology. Mol. Biol. Evol. 32:2483–2495.
- Ratmann O., Donker G., Meijer A., Fraser C., Koelle K. 2012. Phylodynamic inference and model assessment with approximate bayesian computation: influenza as a case study. PLoS Comput. Biol. 8:e1002835.
- Ratmann O., Hodcroft E.B., Pickles M., Cori A., Hall M., Lycett S., Colijn C., Dearlove B., Didelot X., Frost S. 2016. Phylogenetic Tools for Generalized HIV-1

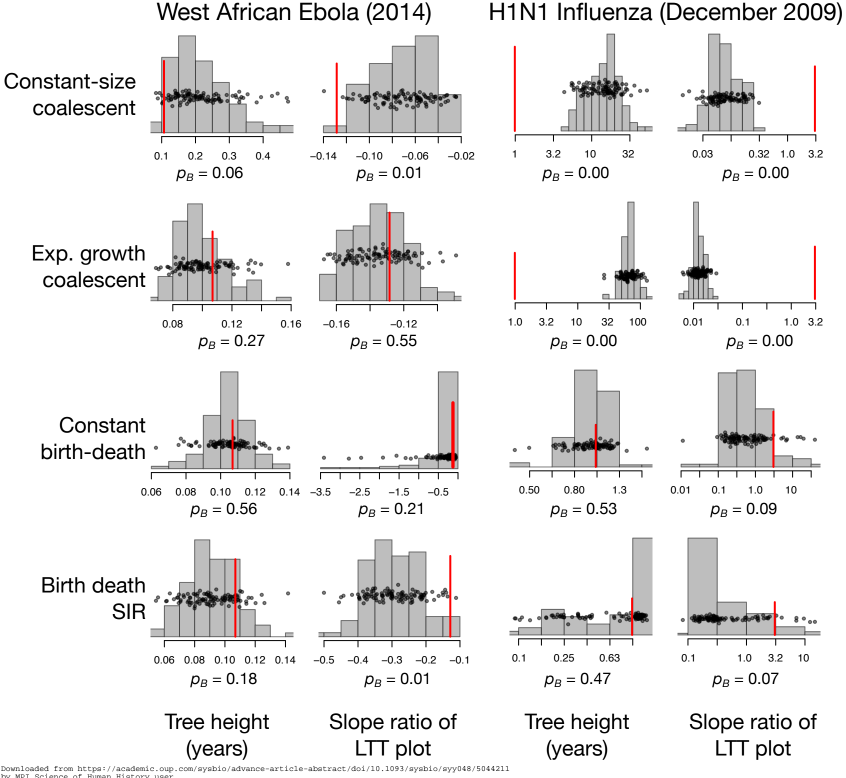
 Epidemics: Findings from the PANGEA-HIV Methods Comparison. Mol. Biol. Evol. 34:185–203.
- Revell L.J., Harmon L.J., Collar D.C. 2008. Phylogenetic signal, evolutionary process, and rate. Syst. Biol. 57:591–601.
- Revell L.J., Harmon L.J., Glor R.E. 2005. Under-parameterized model of sequence

- evolution leads to bias in the estimation of diversification rates from molecular phylogenies. Syst. Biol. 54:973–983.
- Rieux A., Balloux F. 2016. Inferences from tip-calibrated phylogenies: a review and a practical guide. Mol. Ecol. 25:1911–1924.
- Ripplinger J., Sullivan J. 2010. Assessment of substitution model adequacy using frequentist and Bayesian methods. Mol. Biol. Evol. 27:2790–2803.
- Ritchie A.M., Lo N., Ho S.Y.W. 2016. The impact of the tree prior on molecular dating of data sets containing a mixture of inter-and intraspecies sampling. Syst. Biol. 66:413–425.
- Robert C.P., Cornuet J.-M., Marin J.-M., Pillai N.S. 2011. Lack of confidence in approximate Bayesian computation model choice. Proc. Natl. Acad. Sci. USA. 108:15112–15117.
- Rubin D.B. 1981. Estimation in parallel randomized experiments. J. Educ. Stat. 6:377-401.
- Rubin D.B. 1984. Bayesianly justifiable and relevant frequency calculations for the applied statistician. Ann. Stat. 12:1151–1172.
- Saulnier E., Alizon S., Gascuel O. 2017. Assessing the accuracy of Approximate Bayesian Computation approaches to infer epidemiological parameters from phylogenies. PLOS Comput. Biol. 13:e1005416.
- Stadler T. 2010. Sampling-through-time in birth-death trees. J. Theor. Biol. 167:696-404.
- Stadler T., Kouyos R., von Wyl V., Yerly S., Böni J., Bürgisser P., Klimkait T., Joos B., Rieder P., Xie D. 2012. Estimating the basic reproductive number from viral sequence data. Mol. Biol. Evol. 29:347–357.

- Stadler T., Kühnert D., Rasmussen D.A., du Plessis L. 2014. Insights into the early epidemic spread of Ebola in Sierra Leone provided by viral sequence data.

 PLoS Curr. 6.
- Vaughan T.G., Drummond A.J. 2013. A stochastic simulator of birth–death master equations with application to phylodynamics. Mol. Biol. Evol. 30:1480–1493.
- Volz E., Pond S. 2014. Phylodynamic analysis of Ebola virus in the 2014 Sierra Leone epidemic. PLoS Curr. 6.
- Volz E., Siveroni I. 2018. Bayesian phylodynamic inference with complex models. bioRxiv.:268052.
- Volz E.M., Koelle K., Bedford T. 2013. Viral phylodynamics. PLOS Comput. Biol. 9:e1002947.





by MPI Science of Human History user on 10 July 2018