Simulating trajectories and phylogenies from population dynamics models with TiPS

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Abstract

We introduce TiPS, an R-based simulation software to generate time series and genealogies associated with a population dynamics model. The approach is flexible since it can capture any model defined with a set of ordinary differential equations (ODE), and allow parameter values to vary over time periods. Computational time is minimal thanks to the use of the Rcpp package to compile the ODEs into a program corresponding to an implementation of the Gillespie algorithm. This software is particularly suited for epidemiology and phylodynamics, where there is a need to generate numerous phylogenies for a variety of infections life cycles, and in population genetics as well.

Keywords: R package; simulation; phylogenies; compartment models; population dynamics
Introduction

Stochastic population dynamics simulations are routinely used in biology, ecology, or epidemiology [1–3]. These can be used to generate trajectories, also called time series, and genealogies that capture the relatedness between individuals. The increasing amount of genetic data is fuelling interest in linking population dynamics and genealogies because the way organisms spread can leave footprints in their genomes [4–6]. In particular, phylodynamics studies that rely on Approximate Bayesian Computing (ABC) require many simulated datasets [7–9]. Therefore, there is an increasing need for simulating trajectories as well as phylogenies rapidly and flexibly.

One of the most common methods to simulate population dynamics is Gillespie’s stochastic simulation algorithm (SSA) [10], which derives from the formal result by [11] and has been implemented in a variety of programming languages. In R, packages GillespieSSA [12], adaptivetau [13], and epimdr [14] allow to simulate trajectories but none of these allows to simulate genealogies. Conversely, the R packages geiger [15], phytools [16], ape [17], and TreeSim can simulate phylogenies using a birth-death model but they lack population dynamics dimension and the date of the tips of the tree are always drawn at random. One exception is the R package rcolgem (updated to phydynR) that allows simulating both trajectories and phylogenies using a coalescent process. It is also available as a software package PhyDyn [18] in the BEAST2 platform [19]. Another exception is the software package MASTER [20] in BEAST2. Both are not easy to use and phydynR package’s speed can be limited when simulating large phylogenies.

We introduce a flexible and easy-to-use R package to rapidly simulate population dynamics and phylogenies using a backward-in-time, i.e. coalescent, process. We performed a benchmarking analysis and found that our tool is comparable with the R package adaptivetau, but is the fastest when using our new algorithm to simulate trajectories, and is at least one order of magnitude faster than phydynR to simulate phylogenies. In Appendix, we provide additional details and the benchmarking analysis.

Methods

Structure overview

TiPS has two types of stochastic simulation outputs: population trajectories and phylogenies. These are obtained for a population-level model defined as a system of ordinary differential equations (ODE). This is performed by generating simulators in C++ and compiling them into R functions using the Rcpp package [21] for improved computational speed. The general structure of the pipeline is illustrated by the diagram in Figure 1.

Model description

We illustrate the functioning of TiPS using the SIR epidemiological compartmental model, where individuals can be susceptible (with density \(S\)), infected (\(I\)), and removed (\(R\)) [2]. The corresponding ODE system is

\[
\begin{align*}
\frac{dS}{dt} &= -\beta \ S \ I \\
\frac{dI}{dt} &= \beta \ S \ I - (\gamma + \alpha) \ I \\
\frac{dR}{dt} &= \gamma \ I
\end{align*}
\]

where \(\beta\) is the transmission rate, \(\gamma\) the recovery rate, and \(\alpha\) the virulence.
Figure 1: **Structure of the TiPS architecture.** The equations and outputs correspond to the *SIR* model. The functions of the R package are in blue. The simulator of trajectories built as a function is in orange. The variable *traj* in red is the output trajectory. The phylogeny is plotted using the ape R package. [17].
The model can be described as an individual-based model using a system of reactions:

\[ S + I \xrightarrow{\beta IS} I + I \]  
\[ I \xrightarrow{\gamma I} R \]  
\[ I \xrightarrow{\alpha I} \emptyset \]

where the rate of occurrence of each reaction is indicated above the transition arrow. More formally, we define three types of event reactions:

- ‘Birth’ reactions lead to a new individual in one of the compartments (Eq 2a),
- ‘Migration’ reactions correspond to the transition of an individual from one compartment to another (Eq 2b),
- ‘Death’ reactions correspond to the removal of an individual from the system (Eq 2c).

Simulating trajectories

TiPS uses Rcpp to write and compile into C++ a simulator that can then be used through R as a function.

Three simulation algorithms are implemented:

1. Gillespie’s Direct Algorithm (GDA) [10] simulates the time until the next event by assuming that waiting times are exponentially distributed. A limitation is that its computational complexity scales linearly with the number of events and the population size.

2. Gillespie’s Tau-Leap Algorithm (GTA) [22] introduces a fixed time-step during which the number of event of each type is assumed to be Poisson-distributed. This algorithm is limited when few events occur, e.g. early in an epidemic.

3. A Mixed Simulation Algorithm (MSA) that switches from GDA to GTA if over 10 iterations the time until the next event is below a threshold, and from GTA to GDA if the total number of realised event is lower than the number of possible events. For similar variations of the GDA see the next reaction method [23], the optimized direct method [24], the sorting direct method [25], or an adaptive explicit-implicit tau-leaping method [26].

The user can also input a vector of breaking time points, which allows parameter values to vary over time.

Simulating phylogenies

TiPS uses a coalescent approach [27] to simulate phylogenies based on simulated trajectories, i.e. a list of dated events (or ‘reactions’), and known sampling dates (typically corresponding to observed data in ABC approaches).

First, it incorporates the sampling dates into the events of the simulated trajectory. The user needs to define which compartments can be sampled. If more than one compartment can be sampled, the user needs to specify the proportion of external nodes (i.e. leaves) associated with each class. Sampling dates are then randomly associated with a compartment label.

The tree simulation starts from the last (i.e. most recent) sampling date and progresses through the simulated trajectory backwards-in-time. Each of the four types of reactions (birth, death, migration, and sampling) can result in a modification in the simulated tree.

Multiple events may occur at the same date with GTA or the MSA and multiple sampling events may also occur at the same date (e.g. sampling campaigns). If there are multiple occurrences of the same
type of event, we assume that the number of events that lead to a change in the phylogeny can be drawn without replacement from a hypergeometric distribution. Further details can be found in the Appendix.

The output simulated phylogeny is a "phylo" class R object.

Discussion

TiPS simulates trajectories and, from these, phylogenies using a coalescent approach. The analyses in the Appendix show that it outperforms existing R packages in terms of speed when generating numerous trajectories or phylogenies. Future extensions will consist in introducing non-Markovian dynamics, spatial structure, and simulating multifurcating phylogenies.
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