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Abstract

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Reference


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**Application notes**

**SPLATCHE3: simulation of serial genetic data under spatially explicit evolutionary scenarios including long-distance dispersal**

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**Supplementary information:** Details about the implemented evolutionary models and parameters are provided in the software documentation available at http://www.splatche.com/splatche3.

1 Introduction

The simulation of evolutionary scenarios using computer programs is based on the application of combined mathematical models defined by a series of parameters. It is a powerful tool for understanding the mechanisms that shape biodiversity. Compared to deterministic mathematical models, computer simulations have the advantage of considering stochastic processes that better mimic the evolution of species (e.g. Klopfstein, et al., 2006). In particular, they allow hypothesis testing by comparing empirical data to those simulated under models representing such hypotheses. When combined with approximate Bayesian computation (ABC) approaches, they also allow selecting among alternative evolutionary scenarios and estimating parameters of those scenarios (Bertorelle, et al., 2010; Csillery, et al., 2010). They can also be used for comparing and evaluating different analytical tools (e.g. Lopes, et al., 2014; Westesson and Holmes, 2009).

The spatial and temporal dynamics of populations, such as movements in space over time, demographic variation and interactions among populations, plays an important role in evolution. The genetic consequences of those various factors are difficult to disentangle and spatially explicit simulators have been developed for understanding the effects of these combined processes. By spatially explicit, we mean that the simulation takes into account the geographic position of populations.
To the best of our knowledge, the programs SPLATCHE (Currat, et al., 2004) and EASYPOP (Balloux, 2001) were the first spatially explicit computer simulators of genetic diversity available to the scientific community. Other spatially explicit simulation programs pre-dating SPLATCHE were not distributed (Barbujani, et al., 1995; Rendine, et al., 1986). It was initially designed to study the impact of spatial and ecological information on molecular diversity and it modeled migration as movements on a two dimensional stepping-stone lattice (Kimura, 1953). In a few words, SPLATCHE consists in a forward demographic simulation of population demography and migration, followed by a backward coalescent simulation step. In the coalescent step, the ancestry of a sample of gene lineages taken from one or several populations is simulated until the most recent common ancestor of these lineages. Then, genetic diversity of the sample is generated by adding mutations over the simulated coalescent tree. SPLATCHE can handle spatial and environmental complexity through the use of population carrying capacities (e.g. linked to available environmental parameters), migration rates (i.e. directional gene flow) and frictions (i.e. dispersal constraints in different environments) based on user-specified raster maps that can change over time. SPLATCHE and its sequel SPLATCHE2 (Ray, et al., 2010) have been used in many studies, i.e. those on human origin (Ray, et al., 2005) and evolution (Currat and Excoffier, 2005; Lao, et al., 2013; Mon, et al., 2013), on the genetic effects of past range shifts (Banks, et al., 2010; Currat, et al., 2006; Francois, et al., 2010; Gehara, et al., 2013; Ray, et al., 2003; Reid, et al., 2018; Schneider, et al., 2010; Sjodin and Francois, 2011) and climate changes (Brown and Knowles, 2012; Brown, et al., 2016), on admixture between species (Currat, et al., 2008; Durand, et al., 2009) including human and Neanderthals (Currat and Excoffier, 2004; Currat and Excoffier, 2011), as well as on other diverse species (Francois, et al., 2008; White, et al., 2013).

Since the publication of SPLATCHE2 in 2010, new capabilities have been implemented following suggestions from users and authors to provide more realistic simulations and to mimic additional evolutionary scenarios. As a result, we present here a new version of the program that brings together several innovative developments.

2 New capabilities implemented in SPLATCHE3
2.1 Long-distance dispersal
In its previous version, the program could only simulate migration between neighbouring demes under a stepping-stone migration model. SPLATCHE3 implements three new demographic models allowing the simulation of long-distance dispersal (LDD) events. The three LDD models differ by the arrival positions of the LDD events that can be set to target (i) any deme (occupied or empty), (ii) empty demes only, or (iii) occupied demes only. Other user-defined parameters are the proportion of LDD events, the shape of the Gamma distribution used to sample the distance of an LDD event, and the maximum distance of an LDD event. The effects of these models and associated parameters on genetic diversity during range expansion have been explored in Ray and Excoffier (2010). They have also been used to explore the human colonization of Eurasia (Alves, et al., 2016) and the Americas (Branco, et al., 2018).

2.2 Interspecific hybridization
SPLATCHE3 can simulate two interacting populations occupying the same area. Each population is modelled as a layer of interconnected demes spread over space and interactions between populations (competition and/or admixture) occur between demes located at the same position in their respective layers. The model of admixture implemented in the previous version of the program was criticized on the ground that introgression was assumed as asymmetric (Zhang, 2014). To address this point, we have implemented an alternative model of admixture that is more realistic to simulate hybridization between species. Indeed, the previous model was originally designed to represent the assimilation of individuals from one population to the other one, which affected the size of both populations. In the new model, gene flow is simulated in a more symmetric fashion and it does not modify population densities. We renamed the previous admixture model as “assimilation” and the new one as “hybridization”. Three variants for each model are provided in SPLATCHE3: (i) without inter-specific competition, (ii) with density-dependent inter-specific competition, and (iii) with density-independent inter-specific competition. The new hybridization model was described in detail in Excoffier, et al. (2014) and used to study hybridization between wildcats and domestic cats in Nussberger, et al. (2018).

2.3 Heterogeneous migration rates in space and time
In the previous version of the program, a single migration rate could be assigned to each population layer. In SPLATCHE3, it is now possible to assign different migration rates toward each direction (anisotropic migration) and individually for each deme. It is also possible to vary these individual migration rates over time. This feature has been used to simulate population range contractions caused by glacial periods (Arenas, et al., 2013; Arenas, et al., 2012; Mona, et al., 2014).

2.4 Full disappearance of a population within a deme
A reset of the population size when the carrying capacity of a deme is set to 0 was implemented in SPLATCHE3, allowing simulation of the full disappearance of a population (e.g. due to lack of resources caused by a climatic change). This capability was used in Arenas, et al. (2013); Arenas, et al. (2012); and Mona, et al. (2014).

2.5 Genetic data in serial sampling
Since the first successful retrieval of molecular material from ancient remains (Higuchi, et al., 1984; Paabo, 1985), the production of ancient DNA (aDNA) has increased at an almost exponential pace (e.g. Leonardi, et al., 2017) mainly thanks to rapid advances in sequencing techniques and bioinformatics data processing (e.g. Orlando, et al., 2015). SPLATCHE3 can now simulate genetic data for serial samples through the implementation of the serial coalescent algorithm (Rodrigo and Felsenstein, 1999). This capability of SPLATCHE3 may be used to study ancient DNA and has already been used in two studies investigating population continuity through time (Silva, et al., 2017; Silva, et al., 2018). If one wants to add aDNA post-mortem features, scripts (e.g. bash or R) need to be developed and applied to SPLATCHE3 DNA sequences outputted in ARLEQUIN format (Excoffier and Lischer, 2010).

2.6 Multiple mutation models for DNA evolution
The previous version of the program assumes a single rate of change among nucleotide states when simulating DNA sequence evolution (Jukes and Cantor, 1969). However, more complex patterns of change among nucleotide states are observed in empirical data (Arbiza, et al., 2011) and can be considered to provide more realistic evolutionary analysis (Lemmon and Moriarty, 2004). SPLATCHE3 implements models where 4 nucleotide frequencies and the 6 relative rates of change are allowed to vary independently among nucleotide states when simulating DNA sequence evolution (Excoffier and Lischer, 2010).
among nucleotide states are taken into account. These parameters allow the specification of the most commonly used DNA models of evolution, from the simple JC (Jukes and Cantor, 1969) to the complex GTR (Tavaré, 1986).

2.7 MacOs version

SPLATCHE3 includes a console executable for running on Mac OS, which was not available in previous versions of the program.

3 Conclusion

SPLATCHE3 simulates genetic data under a large combination of environmental and demographic features not available in other spatially explicit simulators. Its main strength is the combination of flexibility and efficiency (through the use of the backward coalescent algorithm), which allows huge gains in computing time when compared to alternative forward individual-based simulators (see Arenas, 2012 and references therein). In this respect, the great flexibility of forward individual-based simulations comes with a cost in computing power and time as they simulate all the individuals of all populations, whereas only a small number of individuals and populations are usually analysed. Forward simulations combined with the coalescent framework are much more efficient because they only simulate genetic information for the sample and its ancestral lineages. This approach is implemented in all versions of the program SPLATCHE, including SPLATCHE3. Its implementation in C++ offers a faster computation speed than the alternative program PHYLLOGEOSIM coded in JAVA (Dellicour, et al., 2014). A simple model of range expansion during 1,000 generations from a single deme in a layer composed of 25 to 1,600 demes (Carrying capacity = 300 to 1,000, one DNA sequence of 1,000 bp) results in running-times 2 to 40 faster with SPLATCHE3 than with PHYLLOGEOSIM using a 3.6GHz CPU station running on Linux Ubuntu. Other implementations of the coalescent, such as FASTSIMCOAL (Excoffier, et al., 2013; Excoffier and Foll, 2011) or ms (Hudson, 2002), potentially allow performing spatially explicit simulations by specifying an arbitrary migration matrix between all pairs of populations, but for models with complex geographic features the migration matrix becomes huge and is difficult to set up.

SPLATCHE3 is a flexible simulator allowing to investigate a large variety of evolutionary scenarios in a reasonable computational time. The simulation of full genomes would be too time consuming to explore complex evolutionary scenario with SPLATCHE3. However, it is possible to simulate thousands or tens of thousands of independent or partly linked loci and use them as a proxy for full genome data through forward individual-based simulators (see Arenas, 2012 and references therein). In this respect, the great flexibility of forward individual-based simulations comes with a cost in computing power and time as they simulate all the individuals of all populations, whereas only a small number of individuals and populations are usually analysed. Forward simulations combined with the coalescent framework are much more efficient because they only simulate genetic information for the sample and its ancestral lineages. This approach is implemented in all versions of the program SPLATCHE, including SPLATCHE3. Its implementation in C++ offers a faster computation speed than the alternative program PHYLLOGEOSIM coded in JAVA (Dellicour, et al., 2014). A simple model of range expansion during 1,000 generations from a single deme in a layer composed of 25 to 1,600 demes (Carrying capacity = 300 to 1,000, one DNA sequence of 1,000 bp) results in running-times 2 to 40 faster with SPLATCHE3 than with PHYLLOGEOSIM using a 3.6GHz CPU station running on Linux Ubuntu. Other implementations of the coalescent, such as FASTSIMCOAL (Excoffier, et al., 2013; Excoffier and Foll, 2011) or ms (Hudson, 2002), potentially allow performing spatially explicit simulations by specifying an arbitrary migration matrix between all pairs of populations, but for models with complex geographic features the migration matrix becomes huge and is difficult to set up.

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References


