

# Efficient ancestry and mutation simulation with msprime 1.0

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November 24, 2021 50

**Abstract** 51

Stochastic simulation is a key tool in population genetics, since the models involved are often analytically intractable and simulation is usually the only way of obtaining ground-truth data to evaluate inferences. Because of this necessity, a large number of specialised simulation programs have been developed, each filling a particular niche, but with largely overlapping functionality and a substantial duplication of effort. Here, we introduce `msprime` version 1.0, which efficiently implements ancestry and mutation simulations based on the succinct tree sequence data structure and `tskit` library. We summarise `msprime`'s many features, and show that its performance is excellent, often many times faster and more memory efficient than specialised alternatives. These high-performance features have been thoroughly tested and validated, and built using a collaborative, open source development model, which reduces duplication of effort and promotes software quality via community engagement. 52  
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**Keywords:** Simulation, Coalescent, Mutations, Ancestral Recombination Graphs 63

**Introduction** 64

The coalescent process (Kingman, 1982a,b; Hudson, 1983b; Tajima, 1983) models the ancestry of a set of sampled genomes, providing a mathematical description of the genealogical tree that relates the samples to one another. It has proved to be a powerful model, and is now central to population genetics (Hudson, 1990; Hein et al., 2004; Wakeley, 2008). The coalescent is an efficient framework for population genetic simulation, because it allows us to simulate the genetic ancestry for a sample from an idealised population model, without explicitly representing the population in memory or stepping through the generations. Indeed, Hudson (1983b) independently derived the coalescent *in order to* efficiently simulate data, and used these simulations to characterise an analytically intractable distribution. This inherent efficiency, and the great utility of simulations for a wide 65  
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range of purposes, has led to dozens of different tools being developed over the decades (Carvajal-Rodríguez, 2008; Liu et al., 2008; Arenas, 2012; Yuan et al., 2012; Hoban et al., 2012; Yang et al., 2014; Peng et al., 2015).

Two technological developments of recent years, however, pose major challenges to most existing simulation methods. Firstly, fourth-generation sequencing technologies have made complete chromosome-level assemblies possible (Miga et al., 2020), and high quality assemblies are now available for many species. Thus, modelling genetic variation data as a series of unlinked non-recombining loci is no longer a reasonable approximation, and we must fully account for recombination. However, while a genealogical tree relating  $n$  samples in the single-locus coalescent can be simulated in  $O(n)$  time (Hudson, 1990), the coalescent with recombination is far more complex, and programs such as Hudson’s classical `ms` (Hudson, 2002) can only simulate short segments under the influence of recombination. The second challenge facing simulation methods is that sample sizes in genetic studies have grown very quickly in recent years, enabled by the precipitous fall in genome sequencing costs. Human datasets like the UK Biobank (Bycroft et al., 2018) and gnomAD (Karczewski et al., 2020) now consist of hundreds of thousands of genomes and many other datasets on a similar scale are becoming available (Tanjo et al., 2021). Classical simulators such as `ms` and even fast approximate methods such as `scrm` (Staab et al., 2015) simply cannot cope with such a large number of samples.

The `msprime` simulator (Kelleher et al., 2016; Kelleher and Lohse, 2020) has greatly increased the scope of coalescent simulations, and it is now straightforward to simulate millions of whole chromosomes for a wide range of organisms. The “succinct tree sequence” data structure (Kelleher et al., 2016, 2018, 2019; Wohns et al., 2021), originally introduced as part of `msprime`, makes it possible to store such large simulations in a few gigabytes, several orders of magnitude smaller than commonly used formats. The succinct tree sequence has also led to major advances in forwards-time simulation (Kelleher et al., 2018; Haller et al., 2018), ancestry inference (Kelleher et al., 2019; Wohns et al., 2021) and calculation of population genetic statistics (Kelleher et al., 2016; Ralph et al., 2020). Through a rigorous open-source community development process, `msprime` has gained a large number of features since its introduction, making it a highly efficient and flexible platform for population genetic simulation. This paper marks the release of `msprime` 1.0. We provide an overview of its extensive features, demonstrate its performance advantages over alternative software, and discuss opportunities for ongoing open-source community-based development.

## Results

In the following sections we describe the main features of `msprime` 1.0, focusing on the aspects that are either new for this version, or in which our approach differs significantly from classical methods ([summarised in Table 1](#)). Where appropriate, we benchmark `msprime` against other simulators, but the comparisons are illustrative and not intended to be systematic or exhaustive. Please see Kelleher et al. (2016) for a performance comparison of `msprime` against simulators such as `ms`, `msms`, and `scrm`.

## User interface

The majority of simulation packages are controlled either through a command line interface (e.g. Hudson, 2002; Kern and Schrider, 2016), a text-based input file format (e.g. Guillaume and Rougemont, 2006; Excoffier and Foll, 2011; Shlyakhter et al., 2014), or a mixture of both. Command

<u>Interface</u>	<u>Separation of ancestry and mutation simulations. Ability to store arbitrary metadata along with simulation results, and automatic recording of provenance information for reproducibility. Jupyter notebook (Kluyver et al., 2016) integration. Rich suite of analytical and visualisation methods via <code>tskit</code> library.</u>
<u>Ancestry</u>	<u>SMC, SMC', Beta- and Dirac-coalescent, discrete time Wright-Fisher, and selective sweep models. Instantaneous bottlenecks. Discrete or continuous genomic coordinates, arbitrary ploidy, gene conversion. Output full ARG with recombination nodes, ARG likelihood calculations. Record full migration history and census events. Improved performance for large numbers of populations. Integration with forward simulators such as SLiM and fwdpy11 ("recapitation").</u>
<u>Demography</u>	<u>Improved interface with integrated metadata and referencing populations by name. Import from Newick species tree, *BEAST (Heled and Drummond, 2009), and Demes (Gower et al., 2022). Numerical methods to compute coalescence rates.</u>
<u>Mutations</u>	<u>JC69, HKY, F84, GTR, BLOSUM62, PAM, infinite alleles, SLiM and general matrix mutation models. Varying rates along the genome, recurrent/back mutations, discrete or continuous genomic coordinates, overlaying multiple layers of mutations, exact times associated with mutations.</u>

Table 1: Major features of `msprime` 1.0 added since version 0.3.0 (Kelleher et al., 2016).

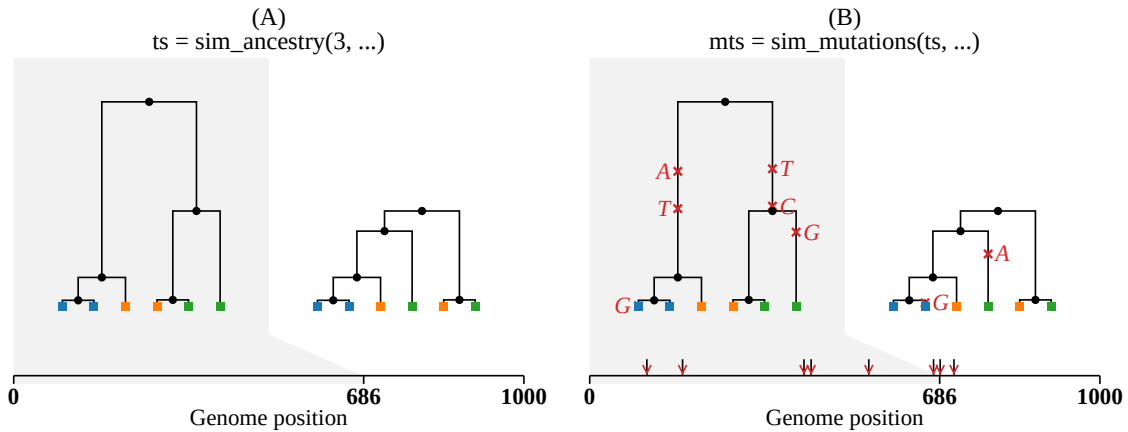


Figure 1: Visualisation of the separation between ancestry and mutation simulation. (A) The result of an invocation of `sim_ancestry(3, ...)` is two trees along a 1kb chunk of genome relating three diploid samples. Each diploid individual consists of two genomes (or nodes), indicated by colour. (B) This ancestry is provided as the input to `sim_mutations`, which adds mutations. Graphics produced using `tskit`'s `draw_svg` method.

line interfaces make it easy to run simple simulations, but as model complexity and the number of parameters increase, they become difficult to understand and error-prone (Ragsdale et al., 2020; ?) (Ragsdale et al., 2020; Gower et al., 2022). Specifying parameters through a text file alleviates this problem to a degree, but lacks flexibility, for example, when running simulations with parameters drawn from a distribution. In practice, for any reproducible simulation project users will write a script to generate the required command lines or input parameter files, invoke the simulation engine, and process the results in some way. This process is cumbersome and labour intensive, and a number of packages have been developed to allow simulations to be run directly in a high-level scripting language (Staab and Metzler, 2016; Parobek et al., 2017; Gladstein et al., 2018).

The more recent trend has been to move away from this file and command-line driven approach and to instead provide direct interfaces to the simulation engines via an Application Programming Interface (API) (e.g. Thornton, 2014; Kelleher et al., 2016; Becheler et al., 2019; Haller and Messer, 2019). The primary interface for `msprime` is through a thoroughly documented `and-stable` Python API, which has encouraged the development of an ecosystem of downstream tools (Terhorst et al., 2017; Chan et al., 2018; Spence and Song, 2019; Adrion et al., 2020a,b; Kamm et al., 2020; McKenzie and Eaton, 2020; Montinaro et al., 2020; Terasaki Hart et al., 2021; Rivera-Colón et al., 2021). As well as providing a stable and efficient platform for building downstream applications, `msprime`'s Python API makes it much easier to build reproducible simulation pipelines, as the entire workflow can be encapsulated in a single script, and package and version dependencies explicitly stated using the `pip` or `conda` package managers. For example, the errors made in the influential simulation analysis of Martin et al. (2017) were only detected because the pipeline could be easily run and reanalysed (Ragsdale et al., 2020; Martin et al., 2020).

A major change for the `msprime` 1.0 release is the introduction of a new set of APIs, designed in

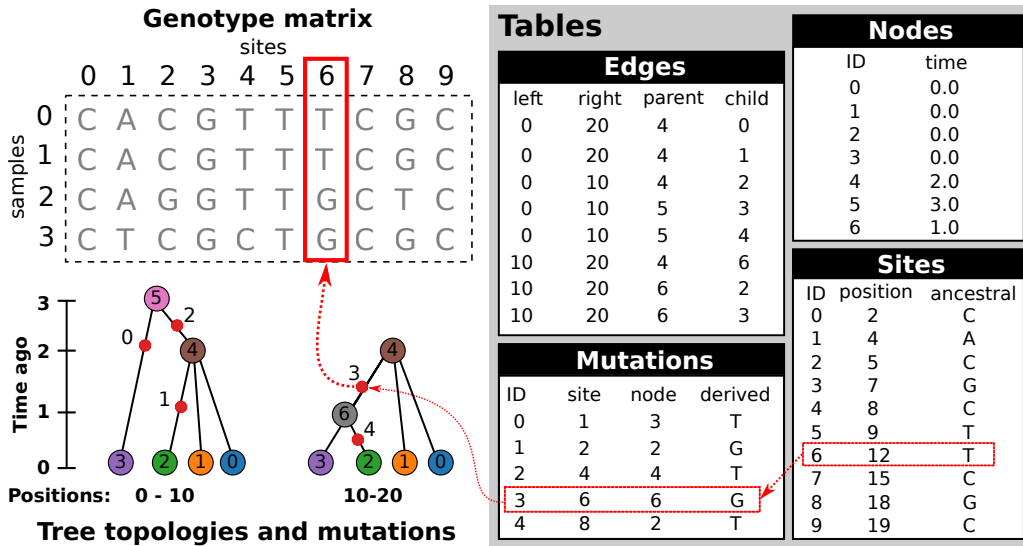


Figure 2: An example tree sequence describing genealogies and sequence variation for four samples at ten sites on a chromosome of twenty bases long. Information is stored in a set of tables (the tables shown here include only essential columns, and much more information can be associated with the various entities). The node table stores information about sampled and ancestral genomes. The edge table describes how these genomes are related along a chromosome, and defines the genealogical tree at each position. The site and mutation tables together describe sequence variation among the samples. [The genotype matrix and tree topologies shown on the left are derived from these tables.](#)

part to avoid sources of error (see the Demography section) but also to provide more appropriate defaults while keeping compatibility with existing code. In the new APIs, ancestry and mutation simulation are fully separated (see Fig. 1), with the `sim_ancestry` and `sim_mutations` functions replacing the legacy `simulate` function. Among other changes, the new APIs default to discrete genome coordinates and finite sites mutations, making the default settings more realistic and resolving a major source of confusion and error. The previous APIs are fully supported and tested, and will be maintained for the foreseeable future. The `msp` program ([a command line interface to the library](#)) has been extended to include new commands for simulating ancestry and mutations separately. A particularly useful feature is the ability to specify demographic models in Demes format (?) (Gower et al., 2022) from the command line, making simulation of complex demographies straightforward. We also provide an `ms` compatible command line interface to support existing workflows.

## Tree sequences

One of the key reasons for `msprime`'s substantial performance advantage over other simulators (Kelleher et al., 2016) is its use of the “succinct tree sequence” data structure to represent simulation results. The succinct tree sequence (usually abbreviated to “tree sequence”) was introduced by

Kelleher et al. (2016) to concisely encode genetic ancestry and sequence variation and was originally implemented as part of `msprime`. We subsequently extracted the core tree sequence functionality from `msprime` to create the `tskit` library, which provides a large suite of tools for processing genetic ancestry and variation data via APIs in the Python and C languages [\(?\) \(Tskit developers, 2022\)](#). The availability of `tskit` as a liberally licensed (MIT) open source toolkit has enabled several other projects (e.g. Kelleher et al., 2019; Haller and Messer, 2019; Wohns et al., 2021; Terasaki Hart et al., 2021) to take advantage of the same efficient data structures used in `msprime`, and we hope that many more will follow. While a full discussion of tree sequences and the capabilities of `tskit` is beyond the scope of this article, we summarise some aspects that are important for simulation.

Let us define a genome as the complete set of genetic material that a child inherits from one parent. Thus, a diploid individual has two (monoploid) genomes, one inherited from each parent. Since each diploid individual lies at the end of two distinct lineages of descent, they will be represented by *two* places (nodes) in any genealogical tree. In the tree sequence encoding a *node* therefore corresponds to a single genome, which is associated with its creation time (and other optional information), and recorded in a simple tabular format (Fig. 2). Genetic inheritance between genomes (nodes) is defined by edges. An *edge* consists of a parent node, a child node and the left and right coordinates of the contiguous chromosomal segment over which the child genome inherited genetic material from the parent genome. Parent and child nodes may correspond to ancestor and descendant genomes separated by many generations. Critically, edges can span multiple trees along the genome (usually referred to as “marginal” trees), and identical node IDs across different trees corresponds to the same ancestral genome. For example, in Fig. 2 the branch from node 0 to 4 is present in both marginal trees, and represented by a single edge (the first row in the edge table). This simple device, of explicitly associating tree nodes with specific ancestral genomes and recording the contiguous segments over which parent-child relationships exist, generalises the original “coalescence records” concept (Kelleher et al., 2016), and is the key to the efficiency of tree sequences (Kelleher et al., 2018, 2019; Ralph et al., 2020). [See Note that this formulation is fully compatible with the concept of an Ancestral Recombination Graph \(ARG\) and any ARG topology can be fully and efficiently encoded in the node and edge tables illustrated in Fig. 2; see the Ancestral Recombination Graphs section below for a discussion of this closely-related concept more details.](#)

The final output of most population genetic simulations is some representation of sequence variation among the specified samples. For coalescent simulations, we usually have three steps: (1) simulate the genetic ancestry, and optionally output the resulting marginal trees; (2) simulate sequence evolution conditioned on this ancestry by generating mutations (see the Simulating mutations section); and (3) output the resulting nucleotide sequences by percolating the effects of the mutations through the trees. Information about the mutations themselves—e.g., where they have occurred on the trees—is usually not retained or made available for subsequent analysis. In `msprime`, however, we skip step (3), instead using `tskit`’s combined data model of ancestry and mutations to represent the simulated sequences. As illustrated in Fig. 2, mutations are a fully integrated part of `tskit`’s tree sequence data model, and genetic variation is encoded by recording sites at which mutations have occurred, and where each mutation at those sites has occurred on the marginal tree. Crucially, the genome sequences themselves are never stored, or indeed directly represented in memory (although `tskit` can output the variant matrix in various formats, if required). It may at first seem inconvenient to have only this indirect representation of the genome sequences, but it is extremely powerful. Firstly, the storage space required for simulations is dramatically reduced. For a simulation of  $n$  samples with  $m$  variant sites, we would require  $O(nm)$  space to

store the sequence data as a variant matrix. However, if this simulation was of a recombining genome with  $t$  trees, then the `tskit` tree sequence encoding requires  $O(n + t + m)$  space, assuming we have  $O(1)$  mutations at each site (Kelleher et al., 2016). For large sample sizes, this difference is profound, making it conceivable, for example, to store the genetic ancestry and variation data for the entire human population on a laptop (Kelleher et al., 2019). As well as the huge difference in storage efficiency, it is often far more efficient to compute statistics of the sequence data from the trees and mutations than it is to work with the sequences themselves. For example, computing Tajima’s  $D$  from simulated data stored in the `tskit` format is several orders of magnitude faster than efficient variant matrix libraries for large sample sizes (Ralph et al., 2020).

The vast genomic datasets produced during the SARS-CoV-2 pandemic have highlighted the advantages of storing genetic variation data using the underlying trees. Turakhia et al. (2021) propose the Mutation Annotated Tree (MAT) format (consisting of a Newick tree and associated mutations in a binary format) and the `matUtils` program as an efficient way to store and process large viral datasets (McBroome et al., 2021), achieving excellent compression and processing performance. Similarly, `phastsim` (De Maio et al., 2021) was developed to simulate sequence evolution on such large SARS-CoV-2 phylogenies, and also outputs a Newick tree annotated with mutations (not in MAT format) to avoid the bottleneck of generating and storing the simulated sequences. While these methods illustrate the advantages of the general approach of storing ancestry and mutations rather than sequences, they do not generalise beyond their immediate settings, and no software library support is available.

The software ecosystem built around `tskit` is stable, mature and rapidly growing. Simulators such as  `fwdpy11`  (Thornton, 2014), `SLiM` (Haller and Messer, 2019), `stdpopsim` (Adrion et al., 2020a), `Geonomics` (Terasaki Hart et al., 2021) and `GSpace` (Virgoulay et al., 2021), and inference methods such as `tsinfer` (Kelleher et al., 2019), `tsdate` (Wohns et al., 2021) and `Relate` (Speidel et al., 2019) use either the Python or C APIs to support outputting results in tree sequence format. Tree sequences are stored in an efficient binary file format, and are fully portable across operating systems and processor architectures. The `tskit` library ensures interoperability between programs by having strict definitions of how the information in each of the tables is interpreted, and stringent checks for the internal consistency of the data model.

## Data analysis

The standard way of representing simulation data is to render the results in a text format, which must subsequently be parsed and processed as part of some analysis pipeline. For example, `ms` outputs a set of sequences and can also optionally output the marginal trees along the genome in Newick format, and variants of this approach are used by many simulators. Text files have many advantages, but are slow to process at scale. The ability to efficiently process simulation results is particularly important in simulation-based inference methods such as Approximate Bayesian Computation (ABC) (Beaumont et al., 2002; Csilléry et al., 2010; Wegmann et al., 2010) and machine learning based approaches (Sheehan and Song, 2016; Chan et al., 2018; Schrider and Kern, 2018; Flagel et al., 2019; Sanchez et al., 2020). Clearly, simulation efficiency is crucial since the size and number of simulations that can be performed determines the depth to which one can sample from the model and parameter space. Equally important, however, is the efficiency with which the simulation results can be transformed into the specific input required by the inference method. In the case of ABC, this is usually a set of summary statistics of the sequence data, and methods avoid the bottleneck of parsing text-based file formats to compute these statistics by either



developing their own simulators (e.g. Cornuet et al., 2008; Lopes et al., 2009) or creating forked versions (i.e., modified copies) of existing simulators (e.g. Thornton and Andolfatto, 2006; Hickerson et al., 2007; Pavlidis et al., 2010; Huang et al., 2011; Quinto-Cortés et al., 2018), tightly integrated with the inference method. Modern approaches to ABC such as ABC-RF (Raynal et al., 2019; Pudlo et al., 2016) and ABC-NN (Csilléry et al., 2012; Blum and François, 2010) use large numbers of weakly informative statistics, making the need to efficiently compute statistics from simulation results all the more acute. By using the stable APIs and efficient data interchange mechanisms provided by `tskit`, the results of an `msprime` simulation can be immediately processed, without format conversion overhead. The `tskit` library has a rich suite of population genetic statistics and other utilities, and is in many cases orders of magnitude faster than matrix-based methods for large sample sizes (Ralph et al., 2020). Thus, the combination of `msprime` and `tskit` substantially increases the overall efficiency of many simulation analysis pipelines.

Classical text based output formats like `ms` are inefficient to process, but also lack a great deal of important information about the simulated process. The tree-by-tree topology information output by simulators in Newick format lacks any concept of node identity, and means that we cannot reliably infer information about ancestors from the output. Because Newick stores branch lengths rather than node times, numerical precision issues also arise for large trees (McGill et al., 2013). Numerous forks of simulators have been created to access information not provided in the output. For example, `ms` has been forked to output information about migrating segments (Rosenzweig et al., 2016), ancestral lineages (Chen and Chen, 2013), and `ms`'s fork `msHOT` (Hellenthal and Stephens, 2007) has in turn been forked to output information on local ancestry (Racimo et al., 2017). All of this information is either directly available by default in `msprime`, or can be optionally stored via options such as `record_migrations` or `record_full_arg` (see the Ancestral Recombination Graphs section) and can be efficiently and conveniently processed via `tskit` APIs.

## Simulating mutations

Because coalescent simulations are usually concerned with neutral evolution (see the Selective sweeps section, however) the problem of generating synthetic genetic variation can be decomposed into two independent steps: firstly, simulating genetic ancestry (the trees), then subsequently simulating variation by superimposing mutation processes on those trees (see Fig. 1). A number of programs exist to place mutations on trees: for instance, the classical `Seq-Gen` program (Rambaut and Grassly, 1997) supports a range of different models of sequence evolution, and various extensions to the basic models have been proposed (e.g. Cartwright, 2005; Fletcher and Yang, 2009). Partly for efficiency and partly in the interest of simplicity for users (i.e., to avoid intermediate text format conversions), population genetic simulators have tended to include their own implementations of mutation simulation, with most supporting the infinite sites model (e.g. Hudson, 2002) but with several supporting a wide range of different models of sequence evolution (e.g. Mailund et al., 2005; Excoffier and Foll, 2011; Virgoulay et al., 2021). Thus, despite the logical separation between the tasks of simulating ancestry and neutral sequence evolution, the two have been conflated in practice.

Part of the reason for this poor record of software reuse and modularity is the lack of standardised file formats, and in particular, the absence of common library infrastructure to abstract the details of interchanging simulation data. Although `msprime` also supports simulating both ancestry and mutations, the two aspects are functionally independent within the software; both ancestry and mutation simulators are present in `msprime` for reasons of convenience and history, and could be split

into separate packages. The efficient C and Python interfaces for `tskit` make it straightforward to add further information to an existing file, and because of its efficient data interchange mechanisms, there is no performance penalty for ~~additional operations~~ operations being performed in a different software package. Thanks to this interoperability, `msprime`'s mutation generator can work with *any* `tskit` tree sequence, be it simulated using `SLiM` (Haller and Messer, 2019) or `fdpypy11` (Thornton, 2014), or estimated from real data (Kelleher et al., 2019; Speidel et al., 2019; Wohns et al., 2021). It is a modular component intended to fit into a larger software ecosystem, and is in no way dependent on `msprime`'s ancestry simulator.

~~As well as providing a new API that emphasises the logical split between ancestry and mutation simulation, we~~ We have greatly extended the sophistication of `msprime`'s mutation generation engine for version 1.0, achieving near feature-parity with `Seq-Gen`. We support a large number of mutation models, including the JC69 (Jukes et al., 1969), F84 (Felsenstein and Churchill, 1996), and GTR (Tavaré et al., 1986) nucleotide models and the BLOSUM62 (Henikoff and Henikoff, 1992) and PAM (Dayhoff et al., 1978) amino acid models. Other models, such as the Kimura two and three parameter models (Kimura, 1980, 1981), can be defined easily and efficiently in user code by specifying a transition matrix between any number of alleles ~~(which can be arbitrary unicode strings)~~. Mutation rates can vary along the genome, and multiple mutation models can be imposed on a tree sequence by overlaying mutations in multiple passes. We have extensively validated the results of mutation simulations against both theoretical expectations and output from `Seq-Gen` (Rambaut and Grassly, 1997) and `Pyvolve` (Spielman and Wilke, 2015).

Simulating mutations in `msprime` is efficient. Fig. 3 shows the time required to generate mutations (using the default JC69 model) on simulated tree sequences for a variety of mutation rates as we vary the number of samples (Fig. 3A) and the sequence length (Fig. 3B). For example, the longest running simulation in Fig. 3B required less than 2 seconds to generate an average of 1.5 million mutations over 137,081 trees in a tree sequence with 508,125 edges. This efficiency for large numbers of trees is possible because the tree sequence encoding allows us to generate mutations on an edge-by-edge basis (see Fig. 2 and the Mutation generation appendix), rather than tree-by-tree and branch-by-branch as would otherwise be required. ~~In the above example from Fig. 3B, if we generated mutations tree-by-tree, we would have to iterate over 273,887,838 branches (since there are 137,081 trees and 1,998 branches in each tree) rather than 508,125 edges, resulting in ~500 times more work. Even if we have a tree sequence consisting of~~ Simulating mutations on a single tree ~~(negating the advantage of working edge-by-edge), `msprime`'s mutation generator is still very efficient. For~~ is also very efficient; for example, we simulated mutations under the BLOSUM62 amino acid model for a tree with  $10^6$  leaves over  $10^4$  sites (resulting in  $\sim 260,000$  mutations) in about 0.8 seconds, including the time required for file input and output. We do not attempt a systematic benchmarking of `msprime`'s mutation generation code against other methods, because at this scale it is difficult to disentangle the effects of inefficient input and output formats from the mutation generation algorithms. Given ~~these~~ the above timings, it seems unlikely that generating mutations with `msprime` would be a bottleneck in any realistic analysis.

There are many ways in which the mutation generation code in `msprime` could be extended. For example, we intend to add support for microsatellites (Mailund et al., 2005), codon models (Arenas and Posada, 2007) and indels (Cartwright, 2005; Fletcher and Yang, 2009), although changes may be required to `tskit`'s data model which is currently based on the assumption of independent sites.

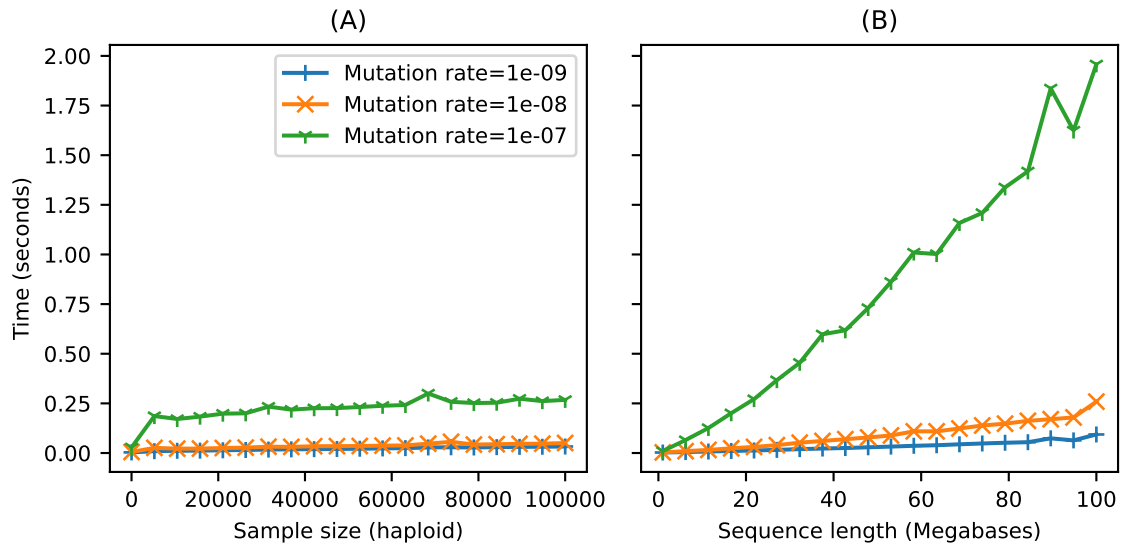


Figure 3: Time required to run `sim_mutations` on tree sequences generated by `sim_ancestry` (with a population size of  $10^4$  and recombination rate of  $10^{-8}$ ) for varying (haploid) sample size and sequence length. We ran 10 replicate mutation simulations each for three different mutation rates, and report the average CPU time required (Intel Core i7-9700). (A) Holding sequence length fixed at 10 megabases and varying the number of samples (tree tips) from 10 to 100,000. (B) Holding number of samples fixed at 1000, and varying the sequence length from 1 to 100 megabases.

## Recombination

Crossover recombination is implemented in `msprime` using Hudson’s algorithm, which works backwards in time, generating common ancestor and recombination events and tracking their effects on segments of ancestral material inherited from the sample (Hudson, 1983a, 1990; Kelleher et al., 2016). Common ancestor events merge the ancestral material of two lineages, and result in coalescences in the marginal trees when ancestral segments overlap. Recombination events split the ancestral material for some lineage at a breakpoint, creating two independent lineages. Using the appropriate data structures (Kelleher et al., 2016), this process is much more efficient to simulate than the equivalent left-to-right approach (Wiuf and Hein, 1999b,a). In `msprime` 1.0, recombination rates can vary along a chromosome, allowing us to simulate recombination hotspots and patterns of recombination from empirical maps. The implementation of recombination in `msprime` is extensively validated against analytical results (Hudson, 1983a; Kaplan and Hudson, 1985) and simulations by `ms`, `msHOT` and `SLiM`.

The Sequentially Markovian Coalescent (SMC) is an approximation of the coalescent with recombination (McVean and Cardin, 2005; Marjoram and Wall, 2006), and was primarily motivated by the need to simulate longer genomes than was possible using tools like `ms`. The SMC is a good approximation to the coalescent with recombination when we have fewer than five sampled genomes (Hobolth and Jensen, 2014; Wilton et al., 2015), but the effects of the approximation are less well understood for larger sample sizes, and several approaches have been proposed that allow simulations to more closely approximate the coalescent with recombination (Chen et al., 2009; Wang et al., 2014; Staab et al., 2015). The SMC and SMC’ models are supported in `msprime` 1.0. However, they are currently implemented using a naive rejection sampling approach, and are somewhat slower to simulate than the exact coalescent with recombination. These models are therefore currently only appropriate for studying the SMC approximations themselves, although we intend to implement them more efficiently in future versions.

**Running time for `msprime` for (A) small and (B) larger simulations on an Intel i7-6600U CPU. Each point is the run time of one simulation, for various values of effective population size ( $N_e$ ), chromosome length in Morgans ( $L$ ), and number of samples ( $n$ ). Run time scales quadratically with the product of  $N_e$  and  $L$ , shown on the horizontal axis. For example, (A) shows that 1,000 samples of 1 Morgan-length chromosomes from a population of  $N_e = 2,000$  diploids would take about 2 seconds, and (equivalently) that the same number of 0.01 Morgan segments with  $N_e = 200,000$  would take the same time. Since recombination rate in these simulations was  $10^{-8}$ ,  $L$  is the number of base pairs divided by  $10^8$ . The black lines are quadratic fits separately in each panel and sample size. Vertical grey lines show the approximate values of  $N_e L$  for chromosome 1 in three species, using values from the `stdpopsim` catalogue (Adrion et al., 2020a).**

In human-like parameter regimes and for large sample sizes, `msprime`’s implementation of the exact coalescent with recombination comprehensively outperforms all other simulators, including those based on SMC approximations (Kelleher et al., 2016). However, it is important to note that although the implementation of Hudson’s algorithm is very efficient, it is still quadratic in the population scaled recombination rate  $\rho = 4N_e L$ , where  $L$  is the length of the genome in units of recombination distance. This is because Hudson’s algorithm tracks recombinations not only in segments ancestral to the sample, but also between ancestral segments. As mentioned above, common ancestor events in which the ancestral material of two lineages is merged only result in coalescences in the marginal trees if their ancestral segments overlap. If there is no overlap, the merged segments represent an ancestral chromosome that is a genetic ancestor of the two lineages, but not the most recent common genetic ancestor at any location along the genome. When this

happens, the merged lineage carries “trapped” genetic material that is not ancestral to any samples, but where recombinations can still occur (Wiuf and Hein, 1999b). The SMC approximations work by disallowing common ancestor events that generate trapped material, greatly simplifying the process. However, this also removes subtle long-range correlations in the trees since there are many ways in which ancestry segments can merge without overlapping.

For large  $\rho$ , recombination events in trapped ancestral material will dominate, and so we can use this as a proxy for the overall number of events in Hudson’s algorithm. Hein et al. (2004, Eq. 5.10) gave

$$\rho(\rho + 1) \left( \sum_{i=1}^{n-1} \frac{1}{i} \right)^2 \quad (1)$$

as an upper bound on the number of recombination events within trapped ancestral material in Hudson’s algorithm for  $n$  samples. Fig. 8 shows the observed run time for simulations with a variety of population sizes, chromosome length and sample sizes, and demonstrates that Eq. correctly predicts. As discussed in the Time complexity of Hudson’s algorithm appendix, the quadratic dependence of simulation running time on  $\rho$ , as previously conjectured (Kelleher et al., 2016, Fig. 2). We also see that the dependence on  $n$  is quite weak, since increasing sample size 100-fold only increases run time by a factor of 2 or so. However, the  $\log^2 n$  factor implied by Eq. implied by (1) (the sum is a harmonic number and can be approximated by  $\log n$ ) is not well supported by observed run times (or numbers of events) except possibly at very large values of  $\rho$ . It therefore appears that a different dependence on  $n$  is required to accurately predict simulation time for a given  $\rho$  and  $n$ .

Fig. 8 is a useful yardstick, allowing us to predict how long simulations should take for a wide range of species. Taking a typical chromosome to be 1 Morgan in length, these plots show, roughly, that simulating chromosome-length samples from a population of thousands of individuals takes seconds, while samples from a population of tens of thousands take minutes. Simulating whole chromosomes for many species is very fast, with 1000 samples of chromosome 1 for *Arabidopsis thaliana* taking less than a second, and a few minutes for dogs and humans. However, the dependence on  $\rho$  is quadratic, and if  $\rho$  is sufficiently large simulations may not be feasible. For example, the *Drosophila melanogaster* chromosome 2L is about 23.5Mb long with an average recombination rate of around  $2.4 \times 10^{-8}$ , so  $L \approx 0.57$ , and with  $N_e = 1.7 \times 10^6$  (Li and Stephan, 2006),  $N_e L \approx 10^6$ , so extrapolating the curve in Fig. 8B predicts that simulation would require around 177 hours for 1000 samples. For such large values of  $\rho$  we recommend users consider approximate simulations. Since `msprime` does not currently have efficient implementations of approximate coalescent with recombination models, in these cases we recommend using SMC-based methods such as `scrm`, particularly if sample sizes are small. In practice, to predict the running time of a given simulation in `msprime`, we recommend that users measure run time in a series of simulations with short genome lengths and the desired sample size, and then predict run time by fitting a quadratic curve to genome length as in Fig. 8. It is important to note that the quadratic curves in the two panels of Fig. 8 are different, and predicting the run times of days-long simulations using the timing of seconds-long runs is unlikely to be very accurate.

What about simulations with changing population size? To understand how run time depends on demography it helps to consider why run time is quadratic in  $\rho$ . At any point in time, `msprime` must keep track of some number of lineages, each of which contains some number of chunks of genetic material. Common ancestor events reduce the number of lineages, and recombination events increase their number. However, with long genomes, only a small fraction of the common ancestor

events will involve overlapping segments of ancestry and lead to coalescence in the marginal trees. Such disjoint segments are often far apart (on average, about distance  $L/2$ ), and so recombine apart again immediately; it is these large numbers of rapid and inconsequential events that lead to the quadratic run time. The maximum number of lineages occurs when the increase and decrease in numbers of lineages due to common ancestor and recombination events balance out. To get an idea of run time we can estimate when this balance occurs. Suppose that the maximum number of lineages is  $M$ ; at this time the rate of common ancestor events is  $M(M-1)/(4N_e)$  and the total rate of recombination is  $M\ell$ , where  $\ell$  is the mean length of genome carried by each lineage (including “trapped” non-ancestral material). At the maximum, coalescence and recombination rates are equal, so a typical segment of ancestry will spend roughly half its time in a lineage with at least one other such segment—and, since such lineages carry at least two segments, at most one-third of the lineages carry long trapped segments of ancestry. Since the maximum number of lineages is reached very quickly (Nelson et al., 2020), this implies that  $\ell \approx L/6$ . Setting the rates of recombination and common ancestor events to be equal and solving for  $M$ , we find that  $M$  is roughly equal to  $LN_e$ . The number of lineages then decreases gradually from this maximum on the coalescent time scale, and therefore over roughly  $2N_e$  generations. Since the total rate of events when the maximum number of lineages is present is roughly  $L^2N_e/6$ , then the total number of events is proportional to  $(LN_e)^2$ —i.e., proportional to  $\rho^2$ .

What does this tell us about run time for simulating time-varying population sizes? The argument above implies that the work is spread out relatively evenly on the coalescent time scale. Suppose that population size today is  $N_1$ , while  $T$  generations ago it was  $N_2$ . Does the run time depend more on  $4N_1L$  or  $4N_2L$ ? The answer depends on how  $T$  compares to  $N_1$ : if  $T/N_1$  is large, then run time will be similar to a population of size  $N_1$ ; while if  $T/N_1$  is small, it will be similar to a population of size  $N_2$ . For instance, in many agricultural species  $N_1 \propto 100$ , while  $N_2 \propto 10^5$ , and the run time will depend critically on  $T$ —in other words, simulation will be quick in a species with a strong domestication bottleneck, and slow otherwise.

observations, and provides a useful means of predicting how long a particular simulation might require.

## Gene conversion

Gene conversion is a form of recombination that results in the transfer of a short segment of genetic material, for example between homologous chromosomes (Chen et al., 2007). Since gene conversion impacts much shorter segments than crossover recombination (typically below 1kb) it affects patterns of linkage disequilibrium differently (Korunes and Noor, 2017). Wiuf and Hein (2000) modelled gene conversion in the coalescent via a rate at which gene conversion events are initiated along the genome and a geometrically distributed tract length. In terms of the ancestral process, gene conversion differs from crossover recombination (as described in the previous section) in that it extracts a short tract of ancestry into an independent lineage, rather than splitting ancestry to the left and right of a given breakpoint. We have implemented this model of gene conversion in `msprime` 1.0, and validated the output against `ms` and analytical results (Wiuf and Hein, 2000).

Gene conversion is particularly useful to model homologous recombination in bacterial evolution, and so we compare the performance of `msprime` with gene conversion to two specialised bacterial simulators, `SimBac` (Brown et al., 2016) and `fastSimBac` (De Maio and Wilson, 2017). Figure 4A shows that `msprime` is far more efficient than both `SimBac` and the SMC-based approximation `fastSimBac`. Figure 4B shows that `msprime` requires somewhat more memory than `fastSimBac`, (as

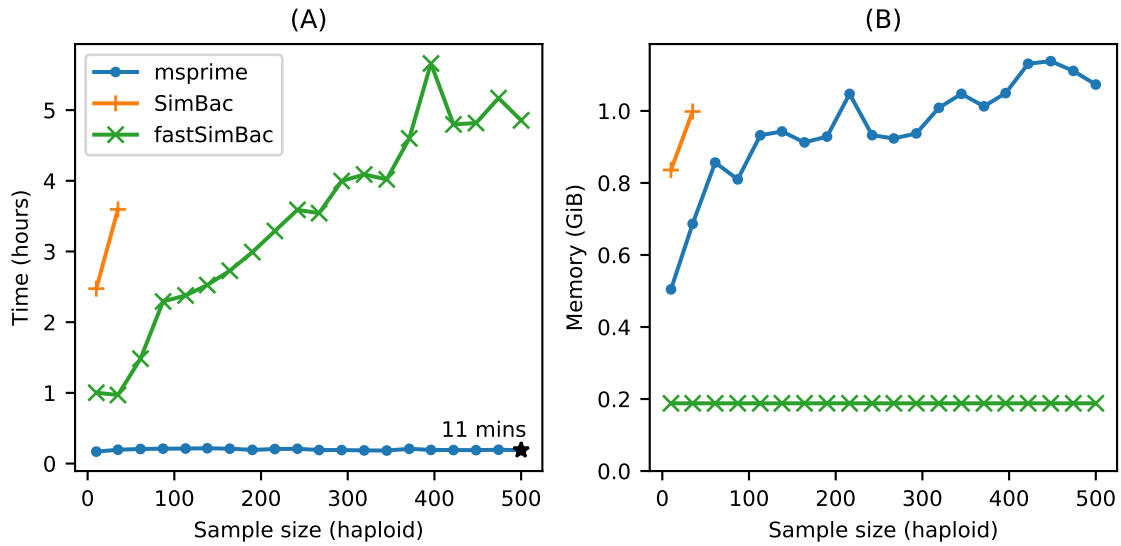


Figure 4: Comparison of simulation performance using `msprime` (`sim_ancestry`), `SimBac`, and `fastSimBac` for varying (haploid) sample sizes, with parameters roughly equivalent to and the current estimate estimates for *E. coli* parameters (Lapierre et al., 2016): a 4.54.6Mb genome with sealed,  $N_e = 1.8 \times 10^8$ , gene conversion rate of  $0.015 - 8.9 \times 10^{-11}$  per base and a mean tract length of 500-542. We report (A) the total CPU time and (B) maximum memory usage averaged over 5 replicates (Intel Xeon E5-2680 CPU). We did not run `SimBac` beyond first two data points because of the very long running times.

expected since `fastSimBac` uses a left-to-right SMC approximation) but is still reasonably modest at around 1GiB for a simulation of 500 whole *E. coli* genomes. However, `msprime` is currently lacking many of the specialised features required to model bacteria, and so an important avenue for future work is to add features such as circular genomes and bacterial gene transfer (Baumdicker and Pfaffelhuber, 2014).

~~In terms of predicting the run time for a simulation including gene conversion, we recommend following the same approach as discussed in the previous section: run a number of simulations for short genome lengths, fit a quadratic to the observed CPU times, and use this to predict run time for larger simulations. Depending on the relative contributions of gene conversion and crossover recombination, this may be an over-estimate since gene conversion events tend to generate less trapped ancestral material than crossovers. Thus, simulations using mammalian-like gene conversion parameters may run faster than simulations in which an equivalent amount of crossover recombination is imposed. Since each gene conversion creates two breakpoints and a crossover creates only one, we expect the output tree sequence for a given rate of gene conversion to be roughly twice the size of the output from a simulation with the same rate of crossovers.~~

## Demography

One of the key applications of population genetic simulations is to generate data for complex demographies. Beyond idealised cases such as stepping-stone or island models, or specialised cases such as isolation-with-migration models, analytical results are rarely possible. Simulation is therefore integral to the development and evaluation of methods for demographic inference. The demography model in `msprime` is directly derived from the approach used in `ms`, and supports an arbitrary number of randomly mating populations exchanging migrants at specified rates. A range of demographic events are supported, which allow for varying population sizes and growth rates, changing migration rates over time, as well as population splits, admixtures and pulse migrations. ~~The location of sampled lineages can be tracked through time in as much detail as required: each tree node is automatically associated with the population in which it arose, the location of lineages can be recorded at any given time via census events, or every lineage migration can be recorded. Large demographic models can be simulated efficiently in version `msprime` 1.0, since we only consider populations that contain lineages and have non-zero migration rates when generating migration event waiting times. This is a considerable improvement over version 0.x, which scaled quadratically with the number of populations.~~

A major change for `msprime` 1.0 is the introduction of the new Demography API, designed to address a design flaw in the `msprime` 0.x interface which led to a number of avoidable errors in downstream simulations (Ragsdale et al., 2020). ~~Briefly, the 0.x API required three separate parameters be provided to the `simulate` function to describe a demographic model, making it easy to accidentally omit information. The 1.0 API resolves this issue by creating a new `Demography` class, which encapsulates all information about the demographic model, and fully decouples the definition from other simulation details. An instance of this class is then provided as a parameter to the new `sim_ancestry` function, substantially reducing the potential for error. Another improvement over the 0.x APIs is the introduction of explicit population split and admixture events, and a population state machine that ensures that lineages cannot migrate into (or be sampled from) inactive populations. This demography model is compatible with the Demes standard (?), and the `Demography` class supports importing and exporting Demes models. Models previously constructed using the 0.x API can be seamlessly imported into the `Demography` class, and we also support~~



importing demographic models from Newick species trees and the output of programs like \*BEAST (Held and Drummond

The `DemographyDebugger` provides detailed information about demographic models as well as a new API is more user-friendly, providing the ability, for example, to refer to populations by name rather than their integer identifiers. We also provide numerical methods to make predictions about these models. For example, we can compute the coalescence rates for two or more lineages drawn from populations at specified times in the past, which can be inverted to obtain the “inverse instantaneous coalescence rate” of Chikhi et al. (2018). Many popular approaches in population genetics use the distribution of coalescence rates between pairs of lineages in one or more populations to infer effective population sizes over time (Li and Durbin, 2011; Sheehan et al., 2013; Schiffels and Durbin, 2014) or split times and subsequent migration rates between populations (Wang et al., 2020). These numerical methods provide a valuable ground-truth when evaluating such inference methods, as illustrated by Adrion et al. (2020a).

## Instantaneous bottlenecks

A common approach to modelling the effect of demographic history on genealogies is to assume that effective population size ( $N_e$ ) changes in discrete steps which define a series of epochs (Griffiths et al., 1994; Marth et al., 2004; Keightley and Eyre-Walker, 2007; Li and Durbin, 2011). In this setting of piecewise constant  $N_e$ , capturing a population bottleneck requires three epochs:  $N_e$  is reduced by some fraction  $b$  at the start of the bottleneck,  $T_{start}$ , and recovers to its initial value at time  $T_{end}$  (Marth et al., 2004). If bottlenecks are short both on the timescale of coalescence and mutations, there may be little information about the duration of a bottleneck ( $T_{end} - T_{start}$ ) in sequence data. Thus a simpler, alternative model is to assume that bottlenecks are instantaneous ( $T_{end} - T_{start} \rightarrow 0$ ) and generate a sudden burst of coalescence events (a multiple merger event) in the genealogy. The strength of the bottleneck  $B$  can be thought of as an (imaginary) time period during which coalescence events are collapsed, i.e. there is no growth in genealogical branches during  $B$  and the probability that a single pair of lineages entering the bottleneck coalesce during the bottleneck is  $1 - e^{-B}$ . Although this simple two parameter model of bottlenecks is attractive and both analytic results and empirical inference (Griffiths et al., 1994; Birkner et al., 2009; Galtier et al., 2000; Bunnefeld et al., 2015) have been developed under this model, there has been no software available to simulate data under instantaneous bottleneck histories.

We have implemented instantaneous bottlenecks in `msprime` 1.0 using a variant of Hudson’s linear time single-locus coalescent algorithm (Hudson, 1990). Instantaneous bottlenecks are specified by adding events to the `Demography` class (see the section) and can be used in combination with any other demographic modelling features. We have, and validated the results of these simulations by comparing against analytical expectations for coalescence times and the site frequency spectrum (Bunnefeld et al., 2015).

## Multiple merger coalescents

Kingman’s coalescent assumes that only two ancestral lineages can merge at each merger event. Although this is generally a reasonable approximation, there are certain situations in which the underlying mathematical assumptions are violated. For example in certain highly fecund organisms (Hedgecock, 1994; Beckenbach, 1994; Hedgecock and Pudovkin, 2011; Árnason, 2004; Irwin et al., 2016), where individuals have the ability to produce numbers of offspring on the order of

the population size and therefore a few individuals may produce the bulk of the offspring in any given generation (Hedgecock, 1994). These population dynamics violate basic assumptions of the Kingman coalescent, and are better modelled by ‘multiple-merger’ coalescents (Donnelly and Kurtz, 1999; Pitman, 1999; Sagitov, 1999; Schweinsberg, 2000; Möhle and Sagitov, 2001), in which more than two lineages can merge in a given event. Multiple-merger coalescent processes have also been shown to be relevant for modelling the effects of selection on gene genealogies (Gillespie, 2000; Durrett and Schweinsberg, 2004; Desai et al., 2013; Neher and Hallatschek, 2013; Schweinsberg, 2017).

Although multiple merger coalescents have been of significant theoretical interest for around two decades, there has been little practical software available to simulate these models. Kelleher et al. (2013, 2014) developed packages to simulate a related spatial continuum model (Barton et al., 2010), Zhu et al. (2015) simulate genealogies within a species tree based on a multiple-merger model, and Becheler and Knowles (2020) provide a general method for simulating multiple merger processes as part of the Quetzal framework (Becheler et al., 2019). The `Beta-Xi-Sim` simulator (Koskela, 2018; Koskela and Wilke Berenguer, 2019) also includes a number of extensions to the Beta-coalescent. None of these methods work with large genomes, and very little work has been performed on simulating multiple merger processes with recombination.

We have added two multiple merger coalescent models in `msprime` 1.0, the Beta-coalescent (Schweinsberg, 2003) and “Dirac”-coalescent (Birkner et al., 2013a), allowing us to efficiently simulate such models with recombination for the first time. These simulation models have been extensively validated against analytical results from the site frequency spectrum (Birkner et al., 2013b; Blath et al., 2016; Hobolth et al., 2019) as well as more general properties of coalescent processes. See the [Appendix-Multiple merger coalescent model appendix](#) for more details and model derivations.

## Ancestral Recombination Graphs

The Ancestral Recombination Graph (ARG) was introduced by Griffiths (Griffiths, 1991; Griffiths and Marjoram, 1997) to represent the stochastic process of the coalescent with recombination as a graph. This formulation is complementary to Hudson’s earlier work (Hudson, 1983a), and substantially increased our theoretical understanding of recombination. In Griffiths’ ARG formulation, a realisation of the coalescent with recombination is a graph in which vertices represent common ancestor or recombination events, and edges represent lineages. There is the “big” ARG, in which we track lineages arising out of recombinations regardless of whether they carry ancestral material (Ethier and Griffiths, 1990), and the “little” ARG in which we only track genetic ancestors. Over time, usage of the term has shifted away from its original definition as a stochastic process, to being interpreted as a representation of a particular genetic ancestry as a graph, without necessarily following the specific details of the Griffiths formulation (e.g. Minichiello and Durbin, 2006; Mathieson and Scally, 2020). Under the latter interpretation, the tree sequence encoding of genetic ancestry (described above) clearly *is* an ARG: the nodes and edges define a graph in which edges are annotated with the set of disjoint genomic intervals through which ancestry flows.

For our purposes, an ARG is a realisation of the coalescent with recombination, in the Griffiths (little ARG) sense. As described in detail by Kelleher et al. (2016), Hudson’s algorithm works by dynamically traversing a little ARG. The graph is not explicitly represented in memory, but is partially present through the extant lineages and the ancestral material they carry over time. We do not output the graph directly, but rather store the information required to recover the genealogical history as nodes and edges in a tree sequence. This is far more efficient than outputting the

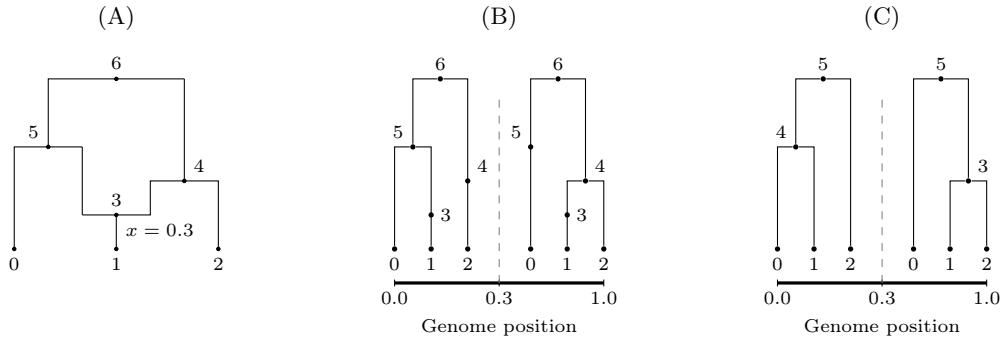


Figure 5: (A) A simple ARG in which a recombination occurs at position 0.3; (B) the equivalent topology depicted as a tree sequence, including the recombination node; (C) the same tree sequence topology “simplified” down to its minimal tree sequence representation. Note that the internal nodes have been renumbered in the simplified representation, so that, e.g., node 5 in (C) corresponds to node 6 in (A) and (B).

simulated ARG in its entirety. For a given scaled recombination rate  $\rho$  (setting aside the dependency on the sample size  $n$ ) we know from Eq. (1) that the number of nodes in an ARG is  $O(\rho^2)$ , whereas the size of the tree sequence encoding is  $O(\rho)$  (Kelleher et al., 2016). This difference between a quadratic and a linear dependency on  $\rho$  is profound, and shows why large simulations cannot output an ARG in practice.

Although by default `msprime` outputs tree sequences that contain full information about the genealogical trees, their correlation structure along the chromosome, and the ancestral genomes on which coalescences occurred, some information is lost in this mapping down from ARG space to the minimal tree sequence form. In particular, we lose information about ancestral genomes that were common ancestors but in which no coalescences occurred, and also information about the precise time and chromosomal location of recombination events. In most cases, such information is of little relevance as it is in principle unknowable, but there are occasions such as visualisation or computing likelihoods (see below) in which it is useful. We therefore provide the `record_full_arg` option in `msprime` to store a representation of the complete ARG traversed during simulation. This is done by storing extra nodes (marked with specific flags, so they can be easily identified later) and edges in the tree sequence (Fig. 5). One situation in which a record of the full ARG is necessary is when we wish to compute likelihoods during inference. The likelihood is a central quantity in evaluating the plausibility of a putative ancestry as an explanation of DNA sequence data, both directly through e.g. approaches based on maximum likelihood, and as an ingredient of methods such as Metropolis-Hastings (Kuhner et al., 2000; Nielsen, 2000; Wang and Rannala, 2008). We provide functions to compute the likelihood of ARG realisations and mutational patterns under the standard coalescent and infinite sites mutation model. ~~See the Appendix for details on these likelihood calculations~~ For details, see the appendix: Likelihood calculations.

## Selective sweeps

Another elaboration of the standard neutral coalescent with recombination is the addition of selective sweeps (Kaplan et al., 1989; Braverman et al., 1995; Kim and Stephan, 2002). Sweeps

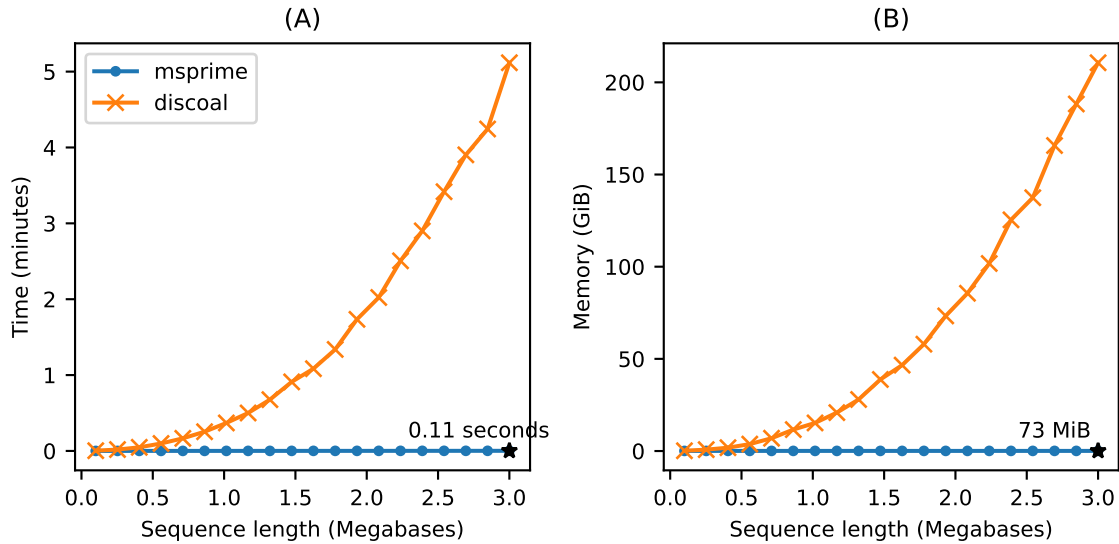


Figure 6: Comparison of selective sweep simulation performance in `msprime` (`sim_ancestry`) and `discoal` (Intel Xeon E5-2680 Gold 6148 CPU). We report the total average CPU time and maximum memory usage when simulating 100–3 replicates for 10–100 diploid samples in a model with a single selective sweep in its history, where the beneficial allele had a scaled selection coefficient of  $2Ns = 1000s = 0.05$ , a per-base recombination rate of  $10^{-9}$ – $10^{-8}$ , population size of  $N = 10^4$ , and sequence length varying from 1kb–200kb–100kb–3000kb.

are modelled by creating a structured population during the sojourn of the beneficial mutation through the population (i.e., the sweep phase) in which lineages may transit between favoured and unfavoured backgrounds through recombination. This approach allows for many selective sweep scenarios to be simulated efficiently, including recurrent, partial, and soft selective sweeps. However this efficiency comes at the cost of flexibility in comparison to forwards in time simulation. Several specialised simulators have been developed to simulate sweeps in the coalescent, including `SeISim` (Spencer and Coop, 2004), `mbs` (Teshima and Innan, 2009), `msms` (Ewing and Hermisson, 2010), `cosi2` (Shlyakhter et al., 2014) and `discoal` (Kern and Schrider, 2016).

Selective sweeps are implemented in the coalescent as a two step-process: first generating an allele frequency trajectory, and then simulating a structured coalescent process conditioned on that trajectory. Following `discoal`, we generate sweep trajectories in `msprime` using a jump process approximation to the conditional diffusion of an allele bound for fixation (Coop and Griffiths, 2004). The jump process moves back in time following the beneficial allele frequency,  $p$ , from some initial frequency (e.g.,  $p = 1$ ) back to the origination of the allele at  $p = 1/(2N)$ , tracking time in small increments  $\delta t$ . Then, given the frequency  $p$  at time  $t$ , the frequency  $p'$  at time  $t + \delta t$  is given by—

$$p' = \begin{cases} p + \mu(p)\delta t + \sqrt{p(1-p)}\delta t & \text{with probability } 1/2 \\ p + \mu(p)\delta t - \sqrt{p(1-p)}\delta t & \text{with probability } 1/2 \end{cases}$$

where

$$\mu(p) = \frac{\alpha p(1-p)}{\tanh(\alpha(1-p))}.$$

Here,  $\alpha = 2Ns$  and  $s$  is the fitness advantage in homozygotes. This model assumes genic selection (i.e., that the dominance coefficient  $h = 0.5$ ), but can be generalised straightforwardly to include arbitrary dominance. We can also define trajectories to model neutral alleles and soft selective sweeps, which we plan as future additions to `msprime`.

Then, given, as detailed in the [Selective sweeps model appendix](#), Given a randomly generated allele frequency trajectory under the above model, the simulation of a sweep works by assigning lineages to two different structured coalescent “labels”, based on whether they carry the beneficial allele. The allele frequency trajectory determines the relative sizes of the “populations” in these labels over time, and therefore the rates at which various events occur. Common ancestor events can then only merge lineages from *within* a label, but lineages can transfer from one label to the other (i.e., from the advantageous to disadvantageous backgrounds, and vice versa) as a result of recombination events. Once we have reached the end of the simulated trajectory the sweep is complete, and we remove the structured coalescent labels. Simulation may then resume under any other ancestry model.

Fig. 6 compares the performance of `msprime` and `discoal` under a simple sweep model, and shows that `msprime` has far better CPU time and memory performance. Since our implementation uses the abstract label system mentioned above, adding support for similar situations, such as inversions (Peischl et al., 2013), should be straightforward.

## Discrete time Wright-Fisher

The coalescent is an idealised model and makes many simplifying assumptions, but it is often surprisingly robust to violations of these assumptions (Wakeley et al., 2012). One situation in which the model does break down is the combination of large sample size and long recombining genomes, where the large number of recombination events in the recent past results in more than the biologically possible  $2^t$  ancestors in  $t$  diploid generations (Nelson et al., 2020). This pathological behaviour results in identity-by-descent, long-range linkage disequilibrium and ancestry patterns deviating from Wright-Fisher expectations, and the bias grows with larger sample sizes (Wakeley et al., 2012; Bhaskar et al., 2014; Nelson et al., 2020). Precisely this problem occurs when simulating modern human datasets, and we have implemented a Discrete Time Wright-Fisher (DTWF) model in `msprime` to address the issue. The DTWF simulates backwards in time generation-by-generation so that each gamete has a unique diploid parent, and multiple recombinations within a generation results in crossover events between the same two parental haploid copies. The method is described in [more](#) detail by Nelson et al. (2020).

Fig. 7 shows that `msprime` simulates the DTWF more quickly and requires substantially less memory than `ARGON` (Palamara, 2016), a specialised DTWF simulator. However, the generation-by-generation approach of the DTWF is less efficient than the coalescent with recombination when the number of lineages is significantly less than the population size (the regime where the coalescent is an accurate approximation), which usually happens in the quite recent past (Bhaskar et al., 2014). We therefore support changing the simulation model during a simulation so that we can run hybrid simulations, as proposed by Bhaskar et al. (2014). Any number of different simulation models can be combined, allowing for the flexible choice of simulation scenarios. As the `discrete`

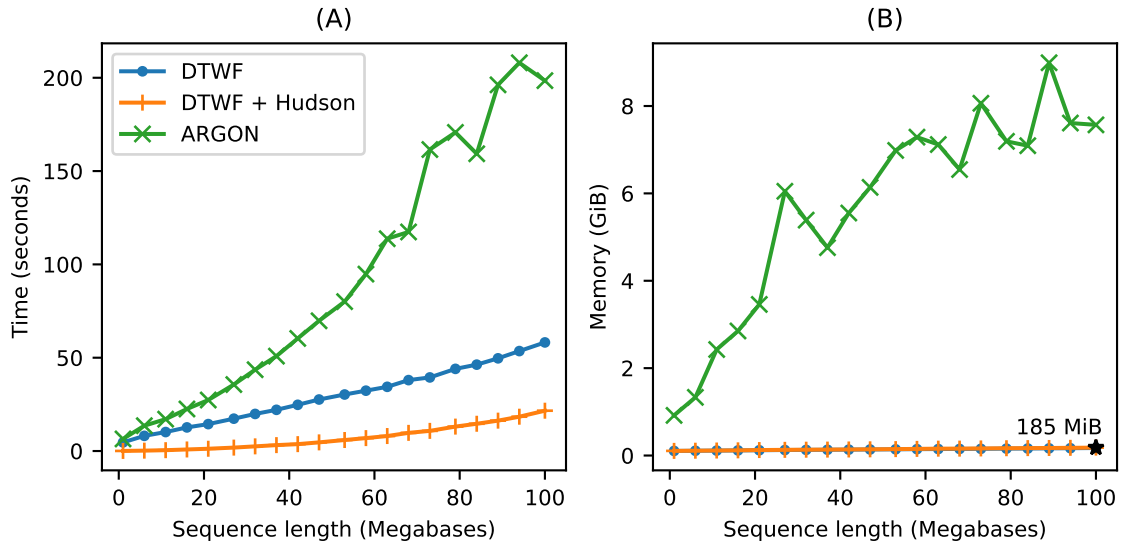


Figure 7: Comparison of Discrete Time Wright-Fisher (DTWF) simulation performance in `msprime` (`sim_ancestry`) and ARGON (Intel Xeon E5-2680 CPU). We simulated ancestry for a sample of 1000 haploids from ran simulations with a population size of  $10000 \cdot 10^4$  and recombination rate of  $10^{-8}$ , with 500 diploid samples and varying sequence length. We report the (A) total CPU time and (B) maximum memory usage for varying sequence lengths, and a per-base recombination rate of  $10^{-8}$ . Each point is the average over 5 replicate simulations. We show observations for ARGON, `msprime`'s DTWF implementation ("DTWF") and a hybrid simulation of 100 generations of the DTWF followed by the standard coalescent with recombination model ("DTWF + Hudson"). Memory usage for `msprime`'s DTWF and hybrid simulations are very similar. We ran ARGON with a mutation rate of 0 and with minimum output options, to ensure we are with a goal of measuring only ancestry simulation time. Memory usage for `msprime`'s DTWF and hybrid simulations are very similar.

~~time Wright-Fisher model~~ DTWF improves accuracy of genealogical patterns in the recent past, we can simulate the recent history using this model and then switch to the standard coalescent to more efficiently simulate the more ancient history.

## Integration with forward simulators

A unique feature of `msprime` is its ability to simulate genetic ancestries by extending an existing partial genetic ancestry. Given a tree sequence that is complete up until time  $t$  ago as input (where marginal trees may or may not have fully coalesced), `msprime` can efficiently obtain the segments of ancestral material present at this time, and then run the simulation backwards in time from there. This allows a simulated ancestry to be produced by any number of different processes across disjoint time slices. In practice this feature is used to “complete” forwards-time ancestry simulations (Kelleher et al., 2018) that may have not fully coalesced. This process (“recapitation”) can be orders of magnitude faster than the standard approach of neutral burn-in; see Haller et al. (2018) for more details and examples. This interoperability between simulators, where a partial ancestry simulation produced by `SLiM` (Haller and Messer, 2019) or  `fwdpy11`  (Thornton, 2014) can be picked up and completed by another simulator, with complete information retained—at scale—is unprecedented. There may be an opportunity for other forward genetic simulators (e.g. Gaynor et al., 2021) to leverage the tree sequence data format and associated tools.

## Development model

`Msprime` has a large number of features, encompassing the functionality of several more specialised simulators while maintaining excellent performance. It is developed by a geographically distributed team of volunteers under an open source community development model, with a strong emphasis on code quality, correctness, good documentation, and inclusive development. As in any large code base, unit tests play a key role in ensuring that new additions behave as expected and `msprime` has an extensive suite. These tests are run automatically on different operating systems on each pull request (where a contributor proposes a code change), using standard Continuous Integration (CI) methodology. Other CI services check for common errors, code formatting issues, and produce reports on the level of test coverage for the proposed change.

Unit tests are vital for ensuring software quality and correctness, but they are usually of little value in assessing the statistical properties of simulations. To validate the correctness of simulation output we maintain a suite of statistical tests (as of 1.0.0, 217 validation tests). These consist of running many replicate simulations to check the properties of the output against other simulators, and where possible against analytical results. For example, simulations of complex demography are validated against `ms`, selective sweeps against `discoal`, and Wright-Fisher simulations against forwards in time simulations in `SLiM`. This suite of tests is run before every release, to ensure that statistical errors have not been introduced.

More visibly to the end user, we also have a high standard for documentation, with precise, comprehensive, and cross-linked documentation that is automatically built from the code base and served through the website <https://tskit.dev>. With the goal of lowering the entry barrier to new users, we have invested significant effort in writing examples and introductions, and making common tasks discoverable. We also view contributions to documentation as equally important to the project as writing code or designing methods: what use would it be to write reliable, stable software if no-one used it?

An important goal of `msprime`'s development model is to maximise accessibility for prospective users and contributors, and to encourage diversity in our community. Gender and racial inequality caused by discrimination and marginalisation is a major problem across the sciences (Wellenreuther and Otto, 2016; Shannon et al., 2019) and in open source software development (Trinkenreich et al., 2021). Within our field, the contribution of women to early computational methods in population genetics was marginalised (Dung et al., 2019), and women continue to be under-represented in computational biology (Bonham and Stefan, 2017). The authorship of our paper reflects these trends, with a skew towards men and affiliations in the USA and Europe. We know the importance of creating and strengthening networks to develop and maintain a diverse community of contributors, and we are committed to fostering a supportive and collaborative environment that helps to address these inequalities in our field.

## Discussion

The 1.0 release of `msprime` marks a major increase in the breadth of available features and the potential biological realism of simulations. These abilities will allow researchers to perform more robust power analyses, more reliably test new methods, carry out more reliable inferences, and more thoroughly explore the properties of theoretical models. Despite this complexity and generality, `msprime`'s performance is state-of-the-art and all features are extensively tested and statistically validated. These advances have only been possible thanks to a distributed, collaborative model of software development, and the work of many people.

Even though simulation has long been a vital tool in population genetics, such collaborative software development has historically been uncommon. A huge proliferation of tools have been published (the references here are not exhaustive) and only a small minority of these are actively developed and maintained today. The ecosystem is highly fragmented, with numerous different ways of specifying parameters and representing results, and there are significant software quality issues at all stages. This is unsurprising, since the majority of simulation software development is performed by students, often without formal training in software development. The result resembles Haldane's sieve for new mutations: many new pieces of software stay permanently on a dusty shelf of supplementary materials, while some of those that prove particularly useful when new (like dominant alleles) are quickly adopted. Although this has produced many good tools and enabled decades of research, it also represents a missed opportunity to invest as a community in shared infrastructure and mentorship in good software development practice.

Scientific software is vital apparatus, and must be engineered to a high quality if we are to trust its results. There is a growing realisation across the sciences (e.g. Siepel, 2019; Harris et al., 2020; Gardner et al., 2021) that investing in shared community infrastructure produces better results than a proliferation of individually maintained tools, allowing scientists to focus on their specific questions rather than software engineering. `Msprime` 1.0 is the result of such a community process, with features added by motivated users, taking advantage of the established development practices and infrastructure. Software development in a welcoming community, with mentorship by experienced developers, is a useful experience for many users. The skills that contributors learn can lead to greatly increased productivity in subsequent work (e.g., through more reliable code and better debugging skills). We hope that users who find that features they require are missing will continue to contribute to `msprime`, leading to a community project that is both high quality and sustainable in the long term.

The succinct tree sequence data structure developed for `msprime` provides a view of not only



genetic variation, but also the genetic ancestry that produced that variation. Recent breakthroughs in methods to infer genetic ancestry in recombining organisms (Rasmussen et al., 2014; Kelleher et al., 2019; Speidel et al., 2019; Wohms et al., 2021; Schaefer et al., 2021; Speidel et al., 2021) have made it possible to estimate such ancestry from real data at scale for the first time (Harris, 2019; Tang, 2019). Given such inferred ancestry, many exciting applications become possible. For example, Osmond and Coop (2021) developed a method to estimate the location of genetic ancestors based on inferred trees, and other uses are sure to follow. Since the inferred genetic ancestry becomes the input for other downstream inferences, it is vitally important that these primary inferences are thoroughly validated, with the detailed properties of the inferred ancestries catalogued and understood. `Msprime` will continue to be an important tool for these inferences and validations, and in this context the ability to interoperate with other methods—particularly forwards simulators—through the succinct tree sequence data structure and `tskit` library will be essential.

## Availability

`Msprime` is freely available under the terms of the GNU General Public License v3.0, and can be installed from the Python Package Index (PyPI) or the `conda-forge` conda channel. Development is conducted openly on GitHub at <https://github.com/tskit-dev/msprime/>. The documentation for `msprime` is available at <https://tskit.dev/msprime/docs/>. The source code for all the evaluations and figures in this manuscript is available at <https://github.com/tskit-dev/msprime-1.0-paper/>.

## Acknowledgements

We [acknowledge the contributions of Ivan Krukov who we consider eligible for authorship, but were unable to contact for approval.](#) We would like to thank Iain Mathieson and Alywyn Scally for helpful comments on the manuscript. ADK was supported by NIH awards R01GM117241 and R01HG010774. BE was supported by DFG grant 273887127 through Priority Programme SPP 1819: Rapid Evolutionary Adaptation (grant STE 325/17-2) to Wolfgang Stephan; BE would also like to acknowledge funding through The Icelandic Research Centre (Rannís) through an Icelandic Research Fund Grant of Excellence nr. 185151-051 to Einar Árnason, Katrín Halldórsdóttir, Alison Etheridge, Wolfgang Stephan, and BE. FB is funded by the Deutsche Forschungsgemeinschaft EXC 2064/1 – Project number 390727645, and EXC 2124 – Project number 390838134. Graham Gower was supported by a Villum Fonden Young Investigator award to Fernando Racimo (project no. 00025300). Gregor Gorjanc is supported by the Chancellor’s Fellowship of the University of Edinburgh and the BBSRC grant to The Roslin Institute BBS/E/D/30002275. Jere Koskela is supported in part by EPSRC grant EP/R044732/1. Jerome Kelleher is supported by the Robertson Foundation. SG acknowledges funding from the Canada Research Chairs Program, from the Canadian Institutes of Health Research PJT 173300, and from the Canadian Foundation for Innovation.

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## Appendix 1261

### Mutation generation 1262

The algorithm that `msprime` uses to simulate mutations on a tree sequence proceeds in two steps: first, mutations are “placed” on the tree sequence (i.e., sampling their locations in time, along the genome, and on the marginal tree), and then the ancestral and derived alleles of each mutation are generated. All mutation models share the code to place mutations, but choose alleles in different ways. 1263  
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First, mutations are placed on the tree sequence under an inhomogeneous Poisson model by applying them independently to each edge. If an edge spans a region  $[a, b)$  of the genome and connected parent and child nodes with times  $s < t$ , and the mutation rate locally is  $\mu$ , then the number of mutations on the edge is Poisson with mean  $\mu(t - s)(b - a)$ , and each mutation is placed independently at a position chosen uniformly in  $[a, b)$  and a time uniformly in  $[s, t)$ . In a discrete genome, all positions are integers and so more than one mutation may occur at the same position 1268  
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on the same edge. Otherwise (i.e., for an infinite-sites model), positions are rejection sampled to obtain a unique floating-point number. If an edge spans a region of the genome with more than one mutation rate, this is done separately for each sub-region on which the mutation rate is constant. Since each edge is processed independently, the algorithm scales linearly with the number of edges in the tree sequence.

Next, alleles are chosen for each mutation. If the site was not previously mutated, then a new ancestral allele is chosen for the site, according to an input distribution of ancestral state allele probabilities. Then, each mutation on the tree is considered in turn, and a derived allele is randomly chosen based on the parental allele (which may be the ancestral allele or the derived allele of a previous mutation). Finally, information about the mutations are recorded in the site and mutation tables of the tree sequence.

A mutation model must, therefore, provide two things: a way of choosing an ancestral allele for each new variant site, and a way of choosing a derived allele given the parental allele at each mutation. Perhaps the simplest mutation model implemented in `msprime` is the `InfiniteAlleles` mutation model, which keeps an internal counter so that the requested alleles are assigned subsequent (and therefore unique) integers.

The distribution of ancestral alleles is used to choose the allele present at the root of the tree at each mutated site, i.e., the `root_distribution`. Mutation models with a finite possible set of alleles have a natural choice for this distribution—the *stationary distribution* of the mutation process. (All mutation models are Markovian, so this may be found as the top left eigenvector of the mutation matrix.) This is the default in most models, except, e.g., the `BinaryMutationModel`, whose alleles are 0 and 1 and always labels the ancestral allele “0”. However, mutational processes are not in general stationary, so we often allow a different root distribution to be specified.

Since the general algorithm above applies mutations at a single rate independent of ancestral state, a model in which different alleles mutate at different rates must necessarily produce some silent mutations, i.e., mutations in which the derived allele is equal to the parental allele. To illustrate this, consider a mutation model in which *A* or *T* mutates to a randomly chosen different nucleotide at rate  $\alpha$  and *C* or *G* mutates at rate  $\beta$ , with  $\beta < \alpha$ . To implement this, first place mutations at the largest total rate, which is  $\alpha$ . Then, at each site, choose an ancestral allele from the root distribution, and for each mutation, choose a derived allele as follows: if the parental allele is *A* or *T*, then choose a random derived allele different to the parental allele; if the parental allele is *C* or *G*, then choose the derived allele to be equal to the parent allele with probability  $\beta/(\alpha + \beta)$ , and randomly choose a different nucleotide otherwise. This produces the correct distribution by Poisson thinning: a Poisson process with rate  $\alpha$  in which each point is discarded independently with probability  $\beta/(\alpha + \beta)$  is equivalent to a Poisson process with rate  $\beta$ . All finite-state models (implemented under the generic `MatrixMutationModel` class) work in this way: mutations are placed at the maximum mutation rate, and then some silent mutations will result.

In previous versions of `msprime`, silent mutations were disallowed, and we could have removed them from the output entirely. However, we have chosen to leave them in, so that for instance simulating with the HKY mutation model will result in silent mutations if not all equilibrium frequencies are the same. The presence of silent mutations may at first be surprising but there is a good reason to leave them in: to allow layering of different mutation models. Suppose that we wanted to model the mutation process as a mixture of more than one model, e.g., Jukes-Cantor mutations at rate  $\mu_1$ , and HKY mutations occur at rate  $\mu_2$ . Layering multiple calls to `sim_mutations` is allowed, so we could first apply mutations with the JC69 model at rate  $\mu_1$  and then add more with the HKY model at rate  $\mu_2$ . However, there is a small statistical problem: suppose that after applying Jukes-Cantor

mutations we have an  $A \rightarrow C$  mutation, but then the HKY mutations inserts another mutation in the middle, resulting in  $A \rightarrow C \rightarrow C$ . If neither mutation model allows silent transitions, then this is clearly not correct, i.e., it is not equivalent to a model that simultaneously applies the two models. (The impact is small, however, as it only affects sites with more than one mutation.) The solution is to make the Jukes-Cantor model *state-independent* (also called “parent-independent”), by placing mutations at rate  $4/3\mu_1$  and then choosing the derived state for each mutation *independently* of the parent (so that 1/4 of mutations will be silent). If so—and, more generally, if the first mutational process put down is state-independent—then the result of sequentially applying the two mutation models is equivalent to the simultaneous model. To facilitate this, many mutation models have a `state_independent` option that increases the number of silent mutations and makes the model closer to state-independent.

Silent mutations are fully supported by `tskit`, which correctly accounts for their presence when computing statistics and performing other operations. For example, silent mutations have no effect on calculations of nucleotide site diversity.

## Time complexity of Hudson’s algorithm

As discussed in the Recombination section, the time complexity of Hudson’s algorithm is predicted to be quadratic in the population scaled recombination rate  $\rho = 4N_eL$  (where  $L$  is the length of the genome in units of recombination distance) by Eq. (1). Fig. 8 shows the running time for simulations with a variety of population sizes, chromosome length and sample sizes, and shows this quadratic prediction is well supported by observations (see also Kelleher et al., 2016, Fig. 2). We also see that the dependence on  $n$  is quite weak, since increasing sample size 100-fold only increases run time by a factor of 2 or so. However, the  $\log^2 n$  factor implied by Eq. (1) (the sum is a harmonic number and can be approximated by  $\log n$ ) is not well supported by observed run times (or numbers of events) except possibly at very large values of  $\rho$ . It therefore appears that a different dependence on  $n$  is required to accurately predict simulation time for a given  $\rho$  and  $n$ .

Fig. 8 is a useful yardstick, allowing us to predict how long simulations should take for a wide range of species. Taking a typical chromosome to be 1 Morgan in length, these plots show, roughly, that simulating chromosome-length samples from a population of thousands of individuals takes seconds, while samples from a population of tens of thousands take minutes. Simulating whole chromosomes for many species is very fast, with 1000 samples of chromosome 1 for *Arabidopsis thaliana* taking less than a second, and a few minutes for dogs and humans. However, the dependence on  $\rho$  is quadratic, and if  $\rho$  is sufficiently large simulations may not be feasible. For example, the *Drosophila melanogaster* chromosome 2L is about 23.5Mb long with an average recombination rate of around  $2.4 \times 10^{-8}$ , so  $L \approx 0.57$ , and with  $N_e = 1.7 \times 10^6$  (Li and Stephan, 2006),  $N_eL \approx 10^6$ , so extrapolating the curve in Fig. 8B predicts that simulation would require around 177 hours for 1000 samples. For such large values of  $\rho$  we recommend users consider approximate simulations. Since `msprime` does not currently have efficient implementations of approximate coalescent with recombination models, in these cases we recommend using SMC based methods such as `scrm`, particularly if sample sizes are small. In practice, to predict the running time of a given simulation in `msprime`, we recommend that users measure run time in a series of simulations with short genome lengths and the desired sample size, and then predict run time by fitting a quadratic curve to genome length as in Fig. 8. It is important to note that the quadratic curves in the two panels of Fig. 8 are different, and predicting the run times of days-long simulations using the timing of seconds-long runs is unlikely to be very accurate.



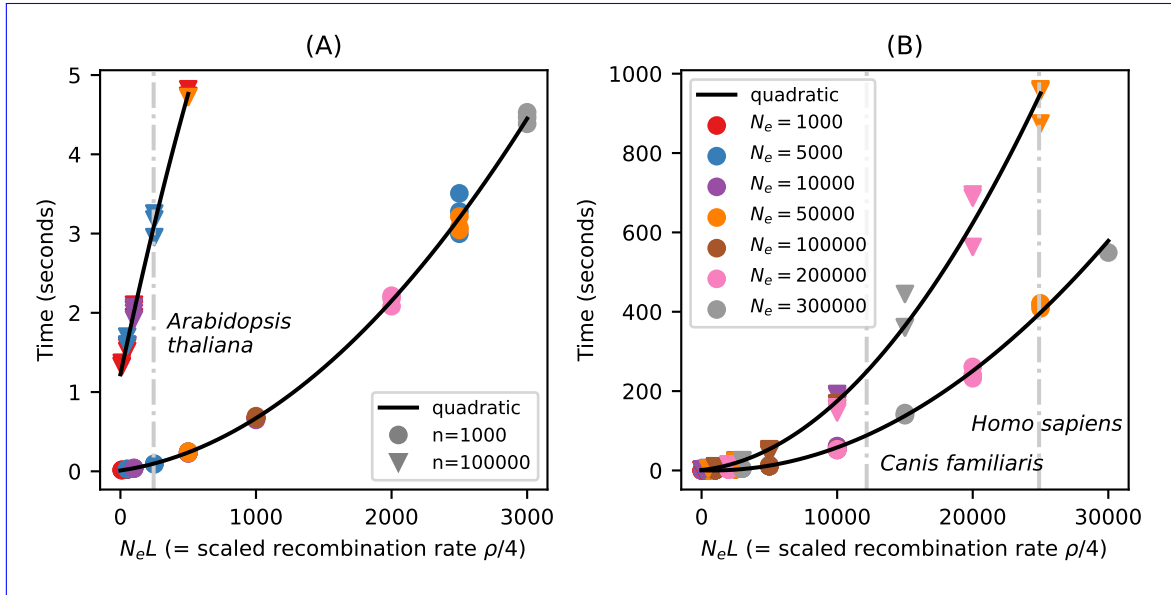


Figure 8: [Running time of `sim\_ancestry` for \(A\) small and \(B\) larger simulations on an Intel i7-6600U CPU.](#) Each point is the run time of one simulation, for various values of effective population size ( $N_e$ ), chromosome length in Morgans ( $L$ ), and number of diploid samples ( $n$ ). Run time scales quadratically with the product of  $N_e$  and  $L$ , shown on the horizontal axis. For example, (A) shows that 1,000 samples of 1 Morgan-length chromosomes from a population of  $N_e = 2,000$  diploids would take about 2 seconds, and (equivalently) that the same number of 0.01 Morgan segments with  $N_e = 200,000$  would take the same time. Since recombination rate in these simulations was  $10^{-8}$ ,  $L$  is the number of base pairs divided by  $10^8$ . The black lines are quadratic fits separately in each panel and sample size. Vertical grey lines show the approximate values of  $N_eL$  for chromosome 1 in three species, using values from the `stdpopsim` catalogue (Adrion et al., 2020a).

What about simulations with changing population size? To understand how run time depends on demography it helps to consider why run time is quadratic in  $\rho$ . At any point in time, `msprime` must keep track of some number of lineages, each of which contains some number of chunks of genetic material. Common ancestor events reduce the number of lineages, and recombination events increase their number. However, with long genomes, only a small fraction of the common ancestor events will involve overlapping segments of ancestry and lead to coalescence in the marginal trees. Such disjoint segments are often far apart (on average, about distance  $L/2$ ), and so recombine apart again immediately; it is these large numbers of rapid and inconsequential events that lead to the quadratic run time. The maximum number of lineages occurs when the increase and decrease in numbers of lineages due to common ancestor and recombination events balance out. To get an idea of run time we can estimate when this balance occurs. Suppose that the maximum number of lineages is  $M$ ; at this time the rate of common ancestor events is  $M(M-1)/(4N_e)$  and the total rate of recombination is  $M\ell$ , where  $\ell$  is the mean length of genome carried by each lineage (including “trapped” non-ancestral material). At the maximum, coalescence and recombination rates are equal, so a typical segment of ancestry will spend roughly half its time in a lineage with at least one other such segment—and, since such lineages carry at least two segments, at most one-third of the lineages carry long trapped segments of ancestry. Since the maximum number of lineages is reached very quickly (Nelson et al., 2020), this implies that  $\ell \approx L/6$ . Setting the rates of recombination and common ancestor events to be equal and solving for  $M$ , we find that  $M$  is roughly equal to  $LN_e$ . The number of lineages then decreases gradually from this maximum on the coalescent time scale, and therefore over roughly  $2N_e$  generations. Since the total rate of events when the maximum number of lineages is present is roughly  $L^2N_e/6$ , then the total number of events is proportional to  $(LN_e)^2$ —i.e., proportional to  $\rho^2$ .

What does this tell us about run time for simulating time-varying population sizes? Suppose that population size today is  $N_1$ , while  $T$  generations ago it was  $N_2$ . Does the run time depend more on  $4N_1L$  or  $4N_2L$ ? The answer depends on how  $T$  compares to  $N_1$ : if  $T/N_1$  is large, then run time will be similar to a population of size  $N_1$ ; while if  $T/N_1$  is small, it will be similar to a population of size  $N_2$ . For instance, in many agricultural species  $N_1 \propto 100$ , while  $N_2 \propto 10^5$ , and the run time will depend critically on  $T$ —in other words, simulation will be quick in a species with a strong domestication bottleneck, and slow otherwise.

## Selective sweeps model

Sweep trajectories are generated in `msprime` using a jump process approximation to the conditional diffusion of an allele bound for fixation (Coop and Griffiths, 2004). The jump process moves back in time following the beneficial allele frequency,  $p$ , from some initial frequency (e.g.,  $p = 1$ ) back to the origination of the allele at  $p = 1/(2N)$ , tracking time in small increments  $\delta t$ . Then, given the frequency  $p$  at time  $t$ , the frequency  $p'$  at time  $t + \delta t$  is given by

$$p' = \begin{cases} p + \mu(p)\delta t + \sqrt{p(1-p)\delta t} & \text{with probability } 1/2 \\ p + \mu(p)\delta t - \sqrt{p(1-p)\delta t} & \text{with probability } 1/2 \end{cases}$$

where

$$\mu(p) = \frac{\alpha p(1-p)}{\tanh(\alpha(1-p))}.$$

Here,  $\alpha = 2Ns$  and  $s$  is the fitness advantage in homozygotes. This model assumes genic selection (i.e., that the dominance coefficient  $h = 0.5$ ), but can be generalised straightforwardly to include arbitrary dominance. We can also define trajectories to model neutral alleles and soft selective sweeps, which we plan as future additions to `msprime`.

## Likelihood calculations

We provide two functions to facilitate likelihood-based inference. Both are implemented only for the simplest case of the standard ARG with a constant population size, and require tree sequences compatible with the `record_full_arg` option as their arguments.

The `msprime.log_arg_likelihood(ts, r, N)` function returns the natural logarithm of the sampling probability of the tree sequence `ts` under the ARG with per-link, per-generation recombination probability `r` and population size `N` (e.g. Kuhner et al., 2000, equation (1)). Specifically, the function returns the logarithm of

$$\left(\frac{1}{2N}\right)^{q_c} \left(\prod_{i:\mathcal{R}} r g_i\right) \exp\left(-\sum_{i=1}^q \left[\frac{1}{2N} \binom{k_i}{2} + r l_i\right] t_i\right),$$

where  $t_i$  is the number of generations between the  $(i-1)$ th and  $i$ th event,  $k_i$  is the number of extant ancestors in that interval,  $l_i$  is the number of links in that interval that would split ancestral material should they recombine,  $q$  is the total number of events in the tree sequence `ts`,  $q_c$  is the number of coalescences,  $\mathcal{R}$  is the set of indices of time intervals which end in a recombination, and  $g_i$  is the corresponding *gap*: the length of contiguous non-ancestral material around the link at which the recombination in question took place. The gap indicates the number of links (or length of genome in a continuous model) at which a recombination would result in exactly the observed pattern of ancestral material in the ARG. For a continuous model of the genome and a recombination in ancestral material, we set  $g_i = 1$  and interpret the result as a density.

The `msprime.unnormalised_log_mutation_likelihood(ts, m)` function returns the natural logarithm of the probability of the mutations recorded in the tree sequence `ts` given the corresponding ancestry, assuming the infinite sites model, up to a normalising constant which depends on the pattern of mutations, but not on the tree sequence or the per-site, per-generation mutation probability `m`. Specifically, the function returns the logarithm of

$$e^{-Tm/2} \frac{(Tm/2)^M}{M!} \prod_{\gamma \in \mathcal{M}} \frac{h_\gamma}{T},$$

where  $T$  and  $\mathcal{M}$  are the total branch length and set of mutations in `ts`, respectively, and for a mutation  $\gamma$ ,  $h_\gamma$  is the total branch length on which  $\gamma$  could have arisen while appearing on all of the leaves of `ts` it does, and on no others. Unary nodes on marginal trees arising from the `record_full_arg` option mean that, in general  $h_\gamma$  corresponds to the length of one or more edges.

## Multiple merger coalescent model

Multiple merger coalescents, in which a random number of ancestral lineages may merge into a common ancestor at a given time, are referred to as  $\Lambda$ -coalescents. The rate at which a given group of  $k$  out of a total of  $b$  lineages merges is

$$\lambda_{b,k} = \int_0^1 x^{k-2} (1-x)^{b-k} \Lambda(dx) + a \mathbb{1}_{\{k=2\}}, \quad 2 \leq k \leq b, \quad (2)$$

where  $\mathbb{1}_{\{A\}} := 1$  if  $A$  holds, and zero otherwise,  $a \geq 0$  is a constant, and  $\Lambda$  is a finite measure on the unit interval without an atom at zero (Donnelly and Kurtz, 1999; Pitman, 1999; Sagitov, 1999). There is also a larger class of simultaneous multiple merger coalescents involving simultaneous mergers of distinct groups of lineages into several common ancestors (Schweinsberg, 2000). These are commonly referred to as  $\Xi$ -coalescents, and often arise from population models incorporating diploidy or more general polyploidy (Birkner et al., 2013a; Blath et al., 2016). To describe a general  $\Xi$ -coalescent, let  $\Delta$  denote the infinite simplex

$$\Delta := \left\{ (x_1, x_2, \dots) : x_1 \geq x_2 \geq \dots \geq 0, \sum_{j=1}^{\infty} x_j \leq 1 \right\}.$$

The rate of mergers is determined by  $\Xi = \Xi_0 + a\delta_0$ , where  $a \geq 0$  is a constant,  $\delta_0$  is the Dirac delta measure, and  $\Xi_0$  is a finite measure on  $\Delta$  with no atom at  $(0, 0, \dots)$ . For an initial number of blocks  $b \geq 2$  and  $r \in \{1, 2, \dots, b-1\}$ , let  $k_1 \geq 2, \dots, k_r \geq 2$  be the sizes of  $r$  merger events and  $s = b - k_1 - \dots - k_r$  be the number of blocks not participating in any merger. The rate of each possible set of mergers with sizes  $(k_1, \dots, k_r)$  is

$$\begin{aligned} \lambda_{n; k_1, \dots, k_r; s} &= \int_{\Delta} \sum_{\ell=0}^s \sum_{\substack{i_1, \dots, i_{r+\ell}=1 \\ \text{all distinct}}}^{\infty} \binom{s}{\ell} x_{i_1}^{k_1} \dots x_{i_r}^{k_r} x_{i_{r+1}} \dots x_{i_{r+\ell}} \left(1 - \sum_{j=1}^{\infty} x_j\right)^{s-\ell} \frac{1}{\sum_{j=1}^{\infty} x_j^2} \Xi_0(dx) \\ &\quad + a \mathbb{1}_{\{r=1, k_1=2\}}, \end{aligned}$$

and the number of such  $(k_1, \dots, k_r)$  mergers is

$$\mathcal{N}(b; k_1, \dots, k_r) = \binom{b}{k_1 \dots k_r s} \frac{1}{\prod_{j=2}^b \ell_j!},$$

where  $\ell_j := \#\{i \in \{1, \dots, r\} : k_i = j\}$  is the number of mergers of size  $j \geq 2$  (Schweinsberg, 2000). 1421

Viewing coalescent processes strictly as mathematical objects, it is clear that the class of  $\Xi$ -coalescents contains  $\Lambda$ -coalescents as a specific example in which at most one group of lineages can merge at each time, and the class of  $\Lambda$ -coalescents contain the Kingman-coalescent as a special case. 1422  
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In `msprime` we have incorporated two examples of multiple-merger coalescents. One is a diploid extension (Birkner et al., 2013a) of the haploid Moran model adapted to sweepstakes reproduction considered by Eldon and Wakeley (2006). Let  $N$  denote the population size, and take  $\psi \in (0, 1]$  to be fixed. In every generation, with probability  $1 - \varepsilon_N$  a single individual (picked uniformly at random) perishes. With probability  $\varepsilon_N$ ,  $\lfloor \psi N \rfloor$  individuals picked uniformly without replacement perish instead. In either case, a surviving individual picked uniformly at random produces enough offspring to restore the population size back to  $N$ . Taking  $\varepsilon_N = 1/N^\gamma$  for some  $\gamma > 0$ , Eldon and Wakeley (2006) obtain  $\Lambda$ -coalescents for which the  $\Lambda$  measure in (2) is a point mass at  $\psi$ . The simplicity of this model does allow one to obtain some explicit mathematical results (see e.g. Der 1431  
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et al. (2012); Eldon and Freund (2018); Freund (2020); Matuszewski et al. (2018)), and the model has also been used to simulate gene genealogies within phylogenies (Zhu et al., 2015). As well as the haploid model of Eldon and Wakeley (2006), `msprime` provides the diploid version of Birkner et al. (2013a), in which individuals perish as above, but replacements are generated by sampling a single pair of diploid individuals as parents, with children sampling one chromosome from each parent. Hence, there are four parent chromosomes involved in each reproduction event, which can lead to up to four simultaneous mergers, giving rise to a  $\Xi$ -coalescent with merger rate

$$\lambda_{b;k_1,\dots,k_r;s}^{\text{Dirac}} = \frac{c\psi^2/4}{1+c\psi^2/4} \frac{4}{\psi^2} \sum_{\ell=0}^{s \wedge (4-r)} \binom{s}{\ell} (4)_{r+\ell} (1-\psi)^{s-\ell} \left(\frac{\psi}{4}\right)^{k_1+\dots+k_r+\ell} + \frac{\mathbb{1}_{\{r=1,k_1=2\}}}{1+c\psi^2/4}, \quad (3)$$

The interpretation of (3) is that ‘small’ reproduction events in which two lineages merge occur at rate  $1/(1+c\psi^2/4)$ , while large reproduction events with the potential to result in simultaneous multiple mergers occur at rate  $(c\psi^2/4)/(1+c\psi^2/4)$ .

The other multiple merger coalescent model incorporated in `msprime` is the haploid population model considered by Schweinsberg (2003), as well as its diploid extension (Birkner et al., 2018). In the haploid version, in each generation of fixed size  $N$ , individuals produce random numbers of juveniles  $(X_1, \dots, X_N)$  independently, each distributed according to a stable law satisfying

$$\lim_{k \rightarrow \infty} Ck^\alpha \mathbb{P}(X \geq k) = 1 \quad (4)$$

with index  $\alpha > 0$ , and where  $C > 0$  is a normalising constant. If the total number of juveniles  $S_N := X_1 + \dots + X_N$  produced in this way is at least  $N$ , then  $N$  juveniles are sampled uniformly at random without replacement to form the next generation. As long as  $\mathbb{E}[X_1] > 1$ , one can show that  $\{S_N < N\}$  has exponentially small probability in  $N$ , and does not affect the resulting coalescent as  $N \rightarrow \infty$  (Schweinsberg, 2003). If  $\alpha \geq 2$  the ancestral process converges to the Kingman-coalescent; if  $1 \leq \alpha < 2$  the ancestral process converges to a  $\Lambda$ -coalescent with  $\Lambda$  in (2) given by the Beta( $2 - \alpha, \alpha$ ) distribution, i.e.

$$\Lambda(dx) = \mathbb{1}_{\{0 < x \leq 1\}} \frac{1}{B(2 - \alpha, \alpha)} x^{1-\alpha} (1-x)^{\alpha-1} dx, \quad (5)$$

where  $B(a, b) = \Gamma(a)\Gamma(b)/\Gamma(a+b)$  for  $a, b > 0$  is the beta function (Schweinsberg, 2003). This model has been adapted to diploid populations by Birkner et al. (2018), and the resulting coalescent is  $\Xi$ -coalescent with merger rate

$$\lambda_{b;k_1,\dots,k_r;s}^{\text{Beta}} = \sum_{\ell=0}^{s \wedge (4-r)} \binom{s}{\ell} \frac{(4)_{r+\ell}}{4^{k+\ell}} \frac{B(k+\ell-\alpha, s-\ell+\alpha)}{B(2-\alpha, \alpha)}, \quad (6)$$

where  $k := k_1 + \dots + k_r$  (Blath et al., 2016; Birkner et al., 2018). The interpretation of (6) is that the random number of lineages participating in a potential merger is governed by the  $\Lambda$ -coalescent with rate (5), and all participating lineages are randomly allocated into one of four groups corresponding to the four parental chromosomes, giving rise to up to four simultaneous mergers.

The stable law (4) assumes that individuals can produce arbitrarily large numbers of juveniles. Since juveniles are at least fertilised eggs, it may be desirable to suppose that the number of juveniles surviving to reproductive maturity cannot be arbitrarily large. Hence we also consider an adaptation of the Schweinsberg (2003) model, where the random number of juveniles has a

deterministic upper bound  $\phi(N)$ , and the distribution of the number of juveniles produced by a given parent (or pair of parents in the diploid case) is 1472  
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$$\mathbb{P}(X = k) = \mathbb{1}_{\{1 \leq k \leq \phi(N)\}} \frac{\phi(N+1)^\alpha}{\phi(N+1)^\alpha - 1} \left( \frac{1}{k^\alpha} - \frac{1}{(k+1)^\alpha} \right). \quad (7)$$

See Eldon and Stephan (2018) for a related model. One can follow the calculations of Schweinsberg (2003) or Birkner et al. (2018) to show that if  $1 < \alpha < 2$  then, recalling that  $k = k_1 + \dots + k_r$ , the merger rate is 1474  
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$$\lambda_{b; k_1, \dots, k_r; s}^{\text{Beta}, M} = \sum_{\ell=0}^{s \wedge (4-r)} \binom{s}{\ell} \frac{(4)_{r+\ell}}{4^{k+\ell}} \frac{B(M; k + \ell - \alpha, s - \ell + \alpha)}{B(M; 2 - \alpha, \alpha)} \quad (8)$$

where  $B(z; a, b) := \int_0^z t^{a-1} (1-t)^{b-1} dt$  for  $a, b > 0$  and  $0 < z \leq 1$  is the incomplete beta function, and

$$M := \lim_{N \rightarrow \infty} \frac{\phi(N)/N}{\phi(N)/N + \mathbb{E}[X_1]} \in (0, 1]$$

(Chetwynd-Diggle et al., 2021). In other words, the measure  $\Lambda$  driving the multiple mergers is of the same form as in (5) with  $0 < x \leq M$  in the case  $1 < \alpha < 2$  and  $\lim_{N \rightarrow \infty} \phi(N)/N > 0$ . If  $\alpha \geq 2$  or  $\phi(N)/N \rightarrow 0$  then the ancestral process converges to the Kingman-coalescent (Chetwynd-Diggle et al., 2021). 1477  
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