# An Msprime Tutorial

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#### 1 Introduction

This tutorial introduces msprime [1]. It was written for a class that used msprime simulations to check for bias in legofit estimates. That effort involved the following steps:

- 1. Install msprime on your computer.
- 2. Modify the Python script described below, so that it simulates your own model of history.
- 3. Run the script to generate simulated data, and pipe the output into simpat, from the Legofit package, to generate site pattern frequencies in .opf format.
- 4. Use legofit to analyze the simulated data.
- 5. Make a graph that compares the true parameter values (the ones used in the simulation) with the estimates obtained by running legofit on the simulated data. This will show you whether the legofit estimates are biased.

## 2 Installing msprime

If you already have Python 3 on your computer, installation is very simple:

python3 -m pip install msprime

# 3 A Python script to run msprime

On the class website, you will find a Python script called msp.py. This script describes a simple model of history, involving 2 modern populations and 1 archaic. You will modify this code to describe your own model of history. Before explaining how it works, I will show you what it does.

By default, the script does not run a simulation. Instead, in invokes the msprime "demography debugger." To use it in this mode, type

```
python3 msp.py
```

and stare at the output. It's fairly self explanatory and you can google "msprime demography debugger" for details.

To simulate data, you must use the -r command-line option. Here's how to look at the first few lines of simulated data:

```
python3 msp.py -r | head
```

You should get something like

```
npops = 3
pop sampsize
x 1
y 1
n 1
0 1 1 0
0 0 0 1
0 0 0 1
0 1 1 0
0 1 1 0
```

This will be followed by an error message about a "Broken pipe," which is generated because the "head" program stopped reading after 10 lines, and so Python had no place to put its output. In this output, the first few lines are a header, which will be read by simpat. It gives the number of populations, their names, and sample sizes. After that (the lines beginning with "0") comes the data. Each line consists of four numbers: first the chromosome, and then the genotypes (0 or 1) of the haploid samples from "x," "y," and "z."

We now dive into the code, so that you will understand what you need to change to build your own model of history. Here is a listing of "msp.py:"

```
1
    #!/usr/bin/python3
    # Before running this: "python3 -m pip install msprime"
3
     import msprime as msp
     import os, sys
 6
    def usage():
         print("Usage: ./msp.py [options]")
9
        print(" where options may include:")
        print(" -r or --run: run simulation. Default: run DemographyDebugger")
         sys.exit(1)
12
    do_simulation = False
    for arg in sys.argv[1:]:
         if arg == "-r" or arg == "--run":
15
             do_simulation = True
         else:
18
             usage()
    # time parameters in generations
21
    Txyn = 25920
    Txy = 3788
    Ta = 1760
                         # age of Altai fossil
24
    Talpha = 1897
                         # time of Neanderthal admixture
    # population sizes
27
    Nxyn = 64964.1/2.0 # ancestral population
```

```
Nxy = 44869.2/2.0
    Nn = 9756.8/2.0
30
    Nx = 20000/2.0
                         # modern Africa
    Ny = 20000/2.0
                         # modern Europe
    # admixture
33
    mAlpha = 0.05
36
    nchromosomes = 10  # number of chromosomes: increase to 1000
                         # number of nucleotides per chromosome
    basepairs = 2e6
    u_per_site = 1.4e-8 \# mutation
39
    # Recombination rate. recomb is the probability of recombination
    # between sites at opposite ends of the simulated sequence.
     c = 1e-8 # rate per base pair per generation
42
    dem = msp.Demography()
45
    # Populations
    dem.add_population(
        name = "X",
48
         description = "African",
         initial\_size = Nx
51
    dem.add_population(
         name = "Y",
         description = "Eurasian",
54
         initial_size = Ny
57
    dem.add_population(
         name = "Y1",
         description = "Y before delta gene flow",
60
         initial_size = Ny
    dem.add_population(
        name = "N",
63
         description = "Neanderthal",
         initial_size = Nn
66
    dem.add_population(
        name = "XY",
         description = "early modern humans",
69
         initial_size = Nxy
    )
    dem.add_population(
72
         name = "XYN",
         description = "ancestral population",
75
         initial_size = Nxyn
```

```
)
     # N->Y gene flow
78
     dem.add_admixture(
         time = Talpha,
81
          derived = "Y",
          ancestral = ["Y1", "N"],
         proportions = [1-mAlpha, mAlpha]
84
     )
     # X-Y split
     dem.add_population_split(
87
          time = Txy,
          derived = ["X", "Y1"],
          ancestral = "XY"
90
     )
93
     # XY-N split
     dem.add_population_split(
          time = Txyn,
          derived = ["XY", "N"],
96
          ancestral = "XYN"
     )
99
     dem.sort_events()
     # One haploid sample from each of 3 populations: 2 modern (X,Y),
102
     # and 1 archaic (N).
     samples = [
105
         msp.SampleSet(num_samples=1, population="X", time=0, ploidy=1),
         msp.SampleSet(num_samples=1, population="Y", time=0, ploidy=1),
         msp.SampleSet(num_samples=1, population="N", time=Ta, ploidy=1),
108
     ]
     if do_simulation:
          # Seed for random number generator. Uses 4 bytes (32 bits) from
111
          # /dev/urandom.
          seed = int.from_bytes(os.urandom(4), "big")
114
          # Simulate gene genealogy. When sim_ancestry is called without
          # num_replicates, it returns a value of class
          # tskit.trees.TreeSequence. But when it's called with
117
          # num_replicates, it returns a "generator", which can be
          # used in a loop to deal with each replicate in turn.
          chr_generator = msp.sim_ancestry(samples = samples,
120
                                           demography = dem,
                                           sequence_length = basepairs,
123
                                           random_seed = seed,
```

```
recombination_rate = c,
                                            num_replicates = nchromosomes
          )
126
          # header
129
          lbl = ["x", "y", "n"]
         print("npops = %d" % len(lbl))
         print("%s %s" % ("pop", "sampsize"))
132
          for s in lbl:
              print("%s %d" % (s, 1))
          # Put mutations onto the gene genealogy
135
          for i, chr in enumerate(chr_generator):
              sim = msp.sim_mutations(chr, rate = u_per_site)
138
              for variant in sim.variants():
141
                  # Use only biallelic sites
                  if len(variant.alleles) != 2:
                      continue
144
                  print(i, end="")
                  for g in variant.genotypes:
                      print("", g, end="")
147
                  print()
      else:
          # Run demography debugger and quit
150
         print(dem.debug())
         print("Use \"./sim -r\" to run simulation")
```

In this program, line 4 imports msprime and abbreviates its name as "msp." Lines 7–11 define a function that prints a usage message message and then aborts. It is used if some problem is detected in parsing the command-line arguments. Line 13 sets do\_simulation equal to false. Unless this is changed by a command-line argument, the program will not execute a computer simulation. Lines 14–18 parse command-line arguments and set do\_simulation equal to true if the user specifies the -r argument. Lines 21–34 define population history parameters. These values should reflect the fitted parameters of your own model of history. Lines 36–42 define parameters relating to the simulation. In this code, I have set nchromosomes equal to 10. You will eventually want to increase it to 1000. The basepairs parameter is the length of a chromosome. I have set it to 2e6, because there isn't much linkage disequilibrium between sites that are farther apart than that. Using a large number of relatively short chromosomes makes the simulations faster.

Line 44 defines a variable called dem, which will hold our model of demographic history. It is an instance of the Demography class, which is defined by msprime. Lines 47–76 define the populations used in the model. Each population has a short name, a description, and an initial size. This size represents the diploid size of the population at the recent end of its temporal extent. My model of history does not include growth within populations, so the initial size of a population is its size throughout its history.

Lines 78–84 define an episode of admixture, giving the time at which it occurs, the name of

the derived population, those of the two ancestral populations, and the proportions of the derived population that came from each of the two parents. Msprime does not allow you to specify the same population as both derived and ancestral within the same episode of admixture. That is why I define Y1 to represent the pre-admixture portion of population Y.

Lines 86–98 define two population splits, one separating XY into X and Y, and the other separating XYN into XY and N. A population split is similar to admixture, except that there are two derived populations and only one derived one.

The add\_population\_split function is also needed in another context, which is not illustrated in our example. Suppose that two parent nodes, a and b, give rise to two daughter nodes, c and d, of which c is a mixture of a and b, but d derives only from b. To specify the relationship between b and d, you would use add\_population\_split like this:

```
dem.add_population_split(
    time = age_of_b,
    derived = ["d"],
    ancestral = "b"
)
```

Note that there is now only one derived population and that its name is enclosed not only in quotes but also in square brackets.

Continuing with the code above, line 100 sorts the events that have been added to dem. If you fail to do this, it's necessary to define the events in order of their time parameters.

Lines 102–108 define a list called samples, each element of which is an object of class SampleSet. The arguments are self explanatory. I set ploidy=1 for consistency with Legofit.

Lines 110–148 execute only if we are doing a simulation. The only line you will need to change is 129, which defines the labels of the populations. These should be consistent with the population parameters specified in lines 105–107. There is presumably a way to read these labels directly from the samples array, but I haven't yet figured out how to do that. This is why I define a separate 1b1 array on line 129.

Even though you won't need to change the other lines, let me explain what is going on in lines 111–148. The variable seed is set using random data provided by the operating system. This value is used to initialize the random number generator so that each run of the program will use a different sequence of pseudo-random numbers.

Lines 120-126 simulate the population history, using msprime's sim\_ancestry function. The type of the value returned by this function depends on whether the num\_replicates parameter is used. Without num\_replicates, sim\_ancestry returns an object of type TreeSequence, which is defined by msprime. With num\_replicates, it returns a Python "generator," which can be used to iterate over the TreeSequence objects of the replicates.

That is the purpose of the loop beginning at line 136. Each pass through that loop handles a different replicate, which I think of as a different chromosome. Within the loop, i is the number of the current chromosome, and chr is an object (defined by msprime) that describes the current chromosome. Line 137 creates sim, which holds all the data about genetic variation on the current chromosome. This variable can be thought of as a list of polymorphic loci, or "variants." We loop across these variants beginning at line 139, ignoring those that aren't biallelic (lines 142–143). Each variant includes a list of genotypes, which we loop over in lines 146–147.

### 4 Generate site pattern frequencies

The output of "msp.py" is designed for use with the "sitepat" program, which is part of Legofit. To run a simulation and tabulate site pattern frequencies, type

```
python3 msp.py -r | simpat
You should get something like this:
# simpat version 2.3.8-12-gf1f00c9
# Including singleton site patterns.
# Number of site patterns: 6
Doing single pass through data to tabulate patterns..
# Nucleotide sites: 66238
# Sites used: 66238
        SitePat
                             E[count]
                        13779.0000000
                        12838.0000000
              у
                        20232.0000000
              n
                        11146.0000000
            x:y
                         3535.0000000
            x:n
                         4679.0000000
            y:n
simpat is finished
```

As you can see, this run only generated 66,238 polymorphic sites. You'll need more sites than this in your analysis. To produce them, edit msp.py and increase the value of nchromosomes to 1000.

In the scaled-up analysis, you'll need an additional file called "sim.sh." This is a shell script that runs the simulation and puts the output into files whose names are determined by \$1, its first command-line argument. This file looks like this:

```
# msp.py is a Python script that executes msprime and generates output
# in the "sim" format, which is described in the document of simpat,
# within the Legofit package. simpat is a program that reads sim
# format and tabulates site patterns.
ofile=sim${1}.opf # standard output
efile=sim${1}.err # error output
python3 msp.py -r | simpat 1>${ofile} 2>${efile}
```

To generate 50 sets of simulated data, you will use a command I describe below, but don't do this on your laptop. The command launches 28 parallel processes, and your laptop probably dosn't have that many cores. Do it on the CHPC server, but not on one of the login nodes. (Users are not allowed to use these for substantial calculations.) Instead, start a job on one of my owner nodes by typing

```
salloc -t 2:00:00 -n 28 -N 1 -A rogersa-kp -p rogersa-kp
```

This starts an interactive session on an owner node. The command above will work on the kingspeak cluster. On notchpeak, change rogersa-kp to rogersa-np. Once you're in that interactive session, type:

```
seq 0 49 | xargs -n 1 -P 28 bash sim.sh
```

Here, "seq 0 49" prints the integers from 0 through 49. The "xargs" command reads this input, and launches processes that look like "bash sim.sh 0," "bash sim.sh 1," and so on up to "bash sim.sh 49." These jobs run in parallel, 28 at a time.

When these jobs finish, you will have files with names like sim0.opf, sim1.opf, and so on up to sim49.opf. You can use these in Legofit analyses similar to the ones you have already done.

### 5 Using Legofit to study simulated data

The analysis pipeline for simulated data hardly differs from the one you used earlier on a real data set and 50 bootstrap replicates. The only difference is that now there is no real data set, and you have simulation replicates instead of bootstrap replicates.

Here is the slurm script, al.slr, for stage 1 of the analysis:

```
#!/bin/bash
#SBATCH -J a1
#SBATCH --account=rogersa-np
#SBATCH --partition=rogersa-np
#SBATCH --time=0:10:00
#SBATCH --nodes 1
#SBATCH --ntasks 1
#SBATCH -o a1-%a.legofit # stdout
#SBATCH -e a1-%a.err # stderr
i=${SLURM_ARRAY_TASK_ID}
ifile='printf "../sim/sim%d.opf" $i'
                                         # input file
stateout='printf "a1-%d.state" $i'
time legofit -1 -d 0 --stateOut ${stateout} --tol 3e-6 -S 5000 a.lgo ${ifile}
On the server, you would launch this command like this:
sbatch --array=0-49 a1.slr
```

## 6 Graphing estimates from simulated data

Figure 1 is copied from Rogers [2]. It compares estimates from 50 simulated data sets with the true parameter values, as specified in the input to msprime. The horizontal spread of the blue values measures the uncertainty of each estimated parameter. The horizontal distance between the red crosses and the center of the blue distribution measures bias.

The code that produced this graph is on the class website in a file called b2.r. For your project, modify this code to reflect the parameter names and the true parameter values of your own project. You may want to remove the code that graphs  $m_{\alpha} + m_{\epsilon}$  and  $m_{\alpha} - m_{\epsilon}$ . These were relevant to a point I was making, but may not be of interest in your project.

Once you have edited b2.r as needed, you can run it from the command line by typing Rscript b2.r. This will produce a file called b2dot.pdf.

Here is a listing of b2.r:

```
# File b2.r: This script reads file "b2.flat", which contains
```

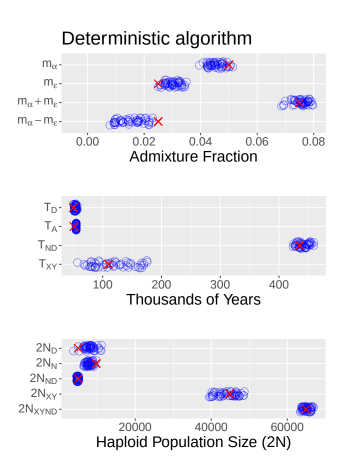


Figure 1: Parameter estimates from 50 simulated data sets. Blue circles are estimates and red crosses are the true parameter values. Symbols for simulation estimates have been jittered vertically.

```
# parameter estimates from several sets of simulated data. It then
    # makes a 1-dimensional scatter plot for each parameter, which
3
    # includes the estimate from each simulation replicate along with the
     # true parameter value, as specified in the input to msprime. The true
    # parameter values are hard coded into this file, rather than being
    # read as input.
    library(ggplot2)
    library(cowplot)
    library(tidyr)
12
    # Adjust text size. Default is 11
    mytheme = theme_get()
    mytheme$text$size = 18
15
    theme_set(mytheme)
18
    # Input file
    wide <- read.table("b2.flat", header=T)</pre>
21
    # parameter names
    parnames <- c("mAlpha", "mEpsilon", "Txy", "Tnd", "Ta", "Td",
                    "twoNxynd", "twoNxy", "twoNnd", "twoNn", "twoNd")
24
    # Subset data, keeping only the parameters we wish to plot.
    wide <- wide[,parnames]</pre>
27
    # Parameter labels, using R's "expression" syntax to make subscripts.
    lbl <- expression("m"[alpha], "m"[epsilon], "T"["XY"], "T"["ND"],</pre>
         "T"["A"], "T"["D"],
30
         "2N"["XYND"], "2N"["XY"], "2N"["ND"], "2N"["N"], "2N"["D"])
33
    # Make graph of migration parameters.
    mpar <- c("mAlpha", "mEpsilon")</pre>
    mdat <- subset(wide, select=mpar)</pre>
36
    ## Create two new parameters
    mdat$msum <- mdat$mAlpha + mdat$mEpsilon</pre>
39
    mdat$mdif <- mdat$mAlpha - mdat$mEpsilon</pre>
    ## Labels of migration parameters
42
    mlbl <- expression("m"[alpha], "m"[epsilon], "m"[alpha]+"m"[epsilon],</pre>
                         "m"[alpha]-"m"[epsilon])
    ## Convert "wide" data format to "long".
45
    mdat <- gather(mdat, par, value, mAlpha, mEpsilon, msum, mdif)</pre>
48
    ## Create column of true parameter values.
    mdat$tru <- rep(NA, nrow(mdat))</pre>
```

```
mdat$tru[mdat$par == "mAlpha"] <- 0.05</pre>
     mdat$tru[mdat$par == "mEpsilon"] <- 0.025</pre>
51
     mdat$tru[mdat$par == "msum"] <- 0.075</pre>
     mdat$tru[mdat$par == "mdif"] <- 0.025</pre>
     mpar <- c("mAlpha", "mEpsilon", "msum", "mdif")</pre>
54
     ## This controls the order in which parameters appear on the graph.
     mdat$par <- ordered(mdat$par, levels=rev(mpar))</pre>
57
     ## Plot migration rate
60
     mplt <- ggplot(mdat, aes(value, par)) +</pre>
         xlab("Admixture Fraction") +
         theme(aspect.ratio=0.2,
                axis.title.y = element_blank()) +
63
         scale_x_continuous(limits=c(-0.005,0.08)) +
         scale_y_discrete(labels=rev(mlbl)) +
66
         geom_jitter(height=0.2, width=0, shape=1, alpha=0.5, size=4, color="blue") +
         geom_point(mapping=aes(tru, par), shape=4, size=4, color="red") +
         ggtitle("Deterministic algorithm")
69
     # Make graph of time parameters
     tpar <- c("Txy", "Tnd", "Ta", "Td")</pre>
     tlbl <- expression("T"["XY"], "T"["ND"], "T"["A"], "T"["D"])</pre>
72
     tdat <- subset(wide, select=tpar)</pre>
     tdat <- gather(tdat, par,value)</pre>
     tdat$tru <- rep(NA, nrow(tdat))</pre>
75
     tdat$tru[tdat$par == "Txy"] <- 3788
     tdat$tru[tdat$par == "Tnd"] <- 15000
     tdat$tru[tdat$par == "Ta"] <- 1760
78
     tdat$tru[tdat$par == "Td"] <- 1734
     tdat$par <- ordered(tdat$par, levels=tpar)</pre>
81
     tplt <- ggplot(tdat, aes(29*value/1000, par)) +</pre>
         xlab("Thousands of Years") +
         theme(aspect.ratio=0.2,
84
                axis.title.y = element_blank()) +
         scale_y_discrete(labels=tlbl) +
87
         geom_jitter(height=0.2, width=0, shape=1, alpha=0.5, size=4, color="blue") +
         geom_point(mapping=aes(29*tru/1000, par), shape=4, size=4, color="red")
90
     # Make graph of population-size parameters
     npar <- c("twoNxynd", "twoNxy", "twoNnd", "twoNn", "twoNd")</pre>
     nlbl <- expression("2N"["XYND"], "2N"["XY"], "2N"["ND"], "2N"["N"],</pre>
         "2N"["D"])
93
     ndat <- subset(wide, select=npar)</pre>
     ndat <- gather(ndat, par, value)</pre>
96
     ndat$tru <- rep(NA, nrow(ndat))</pre>
     ndat$tru[ndat$par == "twoNxynd"] <- 64964.1</pre>
```

```
ndat$tru[ndat$par == "twoNxy"] <- 44869.2</pre>
     ndat$tru[ndat$par == "twoNnd"] <- 5000.0</pre>
99
     ndat$tru[ndat$par == "twoNn"] <- 9756.8</pre>
     ndat$tru[ndat$par == "twoNd"] <- 5000</pre>
     ndat$par <- ordered(ndat$par, levels=npar)</pre>
102
     nplt <- ggplot(ndat, aes(value, par)) +</pre>
          xlab("Haploid Population Size (2N)") +
105
          theme(aspect.ratio=0.3,
                axis.title.y = element_blank()) +
108
          scale_y_discrete(labels=nlbl) +
          geom_jitter(height=0.2, width=0, shape=1, alpha=0.5, size=4, color="blue") +
          geom_point(mapping=aes(tru, par), shape=4, size=4, color="red")
111
     # Combine the 3 graphs into a single graph
     plot_grid(mplt, tplt, nplt, align="hv", ncol=1)
114
     # Save graph in pdf format.
     ggsave("b2dot.pdf")
```

#### References

- [1] Jerome Kelleher, Alison M Etheridge, and Gilean McVean. "Efficient Coalescent Simulation and Genealogical Analysis for Large Sample Sizes". In: *PLoS Computational Biology* 12.5 (May 2016), pp. 1–22. DOI: 10.1371/journal.pcbi.1004842.
- [2] Alan R. Rogers. "An Efficient Algorithm for Estimating Population History from Genetic Data". In: *Peer Community Journal* 2 (2022), e32. DOI: 10.24072/pcjournal.132.