# Supplementary Material 1-A: Simulate SNV sequence data for pedigree founders

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This is the first in a series of RMarkdown documents describing how we simulated exome-sequencing data in pedigrees ascertained to have three or more relatives affected with lymphoid cancer. The following flow chart illustrates the overall workflow for this project.

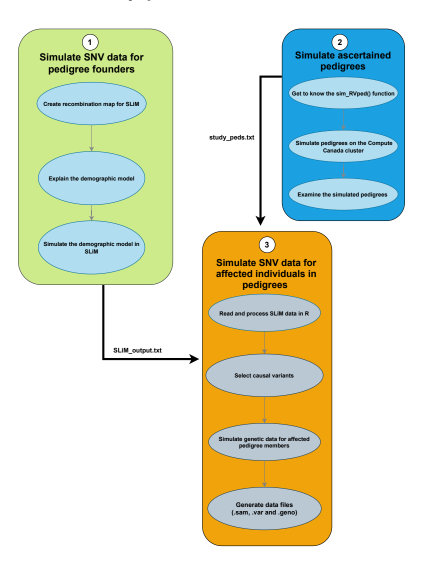


Figure 1: Flow chart showing the overall work flow for simulating the exome-sequencing data for ascertained pedigrees.

This document focuses on the part of the flowchart labelled as 1 (the green box). To start, we require single-nucleotide variant (SNV) sequences for pedigree founders. These founders are assumed to be sampled from an American Admixed population, which we simulate with the evolutionary simulation package SLiM (Haller et al. 2019). In particular, we simulate genome-wide sequences of exons only, to mimic exome sequencing.

The outline of this document as follows. Section 1 explains how we create the SLiM recombination map using

the create\_SlimMap() function in the SimRVSequence R package (Nieuwoudt, Brooks-Wilson, and Graham 2020). Section 2 explains the demographic model for the source population of the pedigree founders. Section 3 discusses how we set the parameters in our SLiM model to simulate the exon-only SNV sequences.

The final outcome of this RMarkdown document is the file SLiM\_output containing SNV exon sequences from a simulated American Admixed population. In the third RMarkdown document of this series, the population sequences file is sampled to get the founder sequences to drop down the ascertained pedigrees. The final gene-dropping step generates the exome-sequencing data in the family-based study of lymphoid cancer families.

## 1 Create recombination map for SLiM

To simulate the genome-wide exon-only sequences with SLiM, we need to supply a recombination map which reads the exon positions in chromosomes. We use the create\_SlimMap() function in the SimRVSequence (Nieuwoudt, Brooks-Wilson, and Graham 2020) R package, as shown in the next code chunk.

```
library(SimRVSequences)

# Load hg_exons data set in SimRVSequence package
data("hg_exons")

# Create recombination map for exon-only data using the hg_exons dataset
s_map <- create_slimMap(exon_df = hg_exons)
head(s_map)</pre>
```

```
##
     chrom segLength recRate mutRate
                                          exon simDist endPos
## 1
         1
                11873 0.00e+00
                                  0e+00 FALSE
                                                      1
                                                              1
## 2
         1
                  354 1.00e-08
                                  1e-08
                                          TRUE
                                                    354
                                                           355
## 3
         1
                  385 3.85e-06
                                  0e+00 FALSE
                                                      1
                                                           356
## 4
         1
                  109 1.00e-08
                                  1e-08
                                                    109
                                                           465
                                          TRUE
## 5
                  499 4.99e-06
                                  0e+00 FALSE
                                                      1
                                                           466
## 6
                 1609 1.00e-08
                                  1e-08
                                         TRUE
                                                   1609
                                                          2075
```

We use the hg\_exons dataset in the SimRVSequence package to specify the exon positions of each of the 22 human autosomes, based on the hg38 reference genome from the UCSC Genome Browser (Nieuwoudt, Brooks-Wilson, and Graham 2020). As shown above, the call to create\_SlimMap() returns a data frame with information about the genetic segments in each chromosome. As an example, the first row in the output above represents information about the genetic segment before the first exon on chromosome 1. The second row represents information about the first exon on chromosome 1. The exon contains 354 base pairs and the recombination and mutation rates in this exon are  $10^{-8}$  per site per generation. The other columns of the data frame are described in the SimRVSequences documentation. The recombination rate between adjacent exons is set to the number of base pairs in the intervening intronic segment (segLength) multiplied by  $10^{-8}$  per base pair per generation (recomb\_rate). Further, the gap between two unlinked chromosomes is set to be a single base pair and the recombination rate between them is set to be 0.5 per base pair per generation (Harris and Nielsen 2016). Since we are interested in exon-only data, the mutation rate outside exons is set to zero and mutation rates inside exons is set to  $10^{-8}$  per base pair per generation (Nieuwoudt, Brooks-Wilson, and Graham 2020).

We need three variables from s\_map() to create the recombination map for simulating exon-only data by SLiM: recRate, mutRate and endPos. We select these three variables and shift the endPos variable forward by one unit because SLiM reads arrays starting at position as 0 rather than 1. We save the resulting output as a text file (Slim\_Map\_chr.txt) to be used as a recombination map for SLiM.

```
# Restrict output to the variables required by SLiM
slimMap <- s_map[, c("recRate", "mutRate", "endPos")]</pre>
```

```
# Shift endPos up by one unit
slimMap$endPos <- slimMap$endPos - 1</pre>
# Print first four rows of slimMap
head(slimMap, n = 4)
##
      recRate mutRate endPos
## 1 0.00e+00
                 0e+00
                            0
## 2 1.00e-08
                 1e-08
                          354
## 3 3.85e-06
                 0e+00
                          355
## 4 1.00e-08
                 1e-08
                          464
# Write the results to a text file
write.table(slimMap, file ="Slim_Map_chr.txt")
```

The next section explains the demographic model we will use to simulate the population-level, exon-only SNV sequences. These sequences will be randomly sampled from the population to be assigned to the founders of our ascertained pedigrees in later steps of the workflow.

## 2 Explain the demographic model

Demographic models play a major role in understanding the genetic patterns in human populations. Throughout human evolution, different demographic events such as expansion, migration, splitting etc. have occurred, affecting genetic diversity (Ragsdale and Gravel 2019). The population-genetics literature has several established demographic models inferred from genetic data (Gutenkunst et al. 2009). Some of these models have been compiled in stdpopsim, a standard library of population-genetic simulation models (Adrion et al. 2020). At the time of writing, this library contains around nine demographic models. Among these, we select the American Admixture demographic model of Browning et al. (2018) because the family-based study motivating our work is in a North American population.

#### 2.1 American admixture demographic model

In the American-Admixture model (Browning et al. 2018), the pre-admixture model parameters are selected from the Out-of-Africa model of Gravel et al. (2011). The following figure illustrates the Out-of-Africa model.

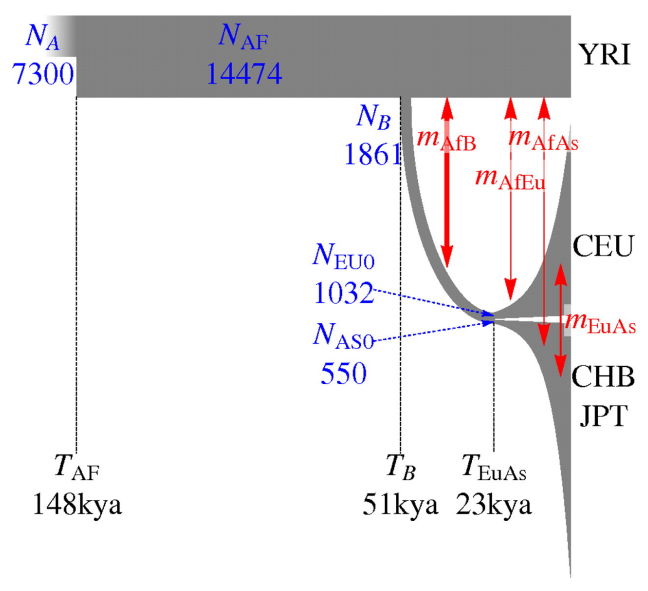


Figure 2: The inferred Out-of-Africa demographic model.

In the figure, the parameter estimates have been rounded and times are expressed in kilo-years before present (kya). The demographic model has three populations representing Africa, Europe and Asia. The initial effective population size of Africa was 7310 individuals which then increased to 14,474 individuals 5920 generations ago (148 kya, assuming a generation time of 25 years). About 2040 generations ago (51 kya), the out-of-Africa migration event occurred with a migrating effective population size of 1861 individuals. Then migration occurred between Africa and out-of-Africa populations with a rate of  $1.5 \times 10^{-4}$  per generation. About 920 generations ago (23 kya), the out-of-Africa population split into two populations, Europe and Asia, with effective sizes of 1032 and 554 individuals, respectively. These two populations then grew at rates of  $3.8 \times 10^{-3}$  per generation for Europe and  $4.8 \times 10^{-3}$  per generation for Asia. Further, between these three populations, (Africa , Europe and Asia) migrations occurred. The migration rates per generation were  $2.5 \times 10^{-5}$  between Africa and Europe,  $7.8 \times 10^{-6}$  between Africa and Asia, and  $3.11 \times 10^{-5}$  between Europe and Asia (Browning et al. 2018). Admixing started about 12 generations ago (0.3 kya) with the initial effective size of the admixed population being 30,000 individuals. The growth rate of the admixed population was 5% per generation with  $\frac{1}{6}$  of the admixed population originating from African ancestry,  $\frac{1}{3}$  from European ancestry and  $\frac{1}{2}$  from Asian ancestry (Browning et al. 2018).

As described in the next section, we use SLiM together with the inferred American-admixture demographic model to simulate population-level exon-only SNV sequences.

## 3 Simulate the demographic model in SLiM

The following SLiM script generates genome-wide exon-only SNV sequences for a population under the American-Admixture demographic model. The script is a .slim file embedded in an R code chunk (that is not run). The original SLiM\_American\_Admixture.slim file can be found on our GitHub page at https://github.com/SFUStatgen/SeqFamStudy/.

```
initialize() {
// Seed number which helps to reproduce the same result
setSeed(2181144364021)
// Read recombination map created by SimRVSequence R package
lines = readFile("~/Slim_Map_chr.txt");
Rrates = NULL;
Mrates = NULL;
ends = NULL;
for (line in lines)
components = strsplit(line);
ends = c(ends, asInteger(components[3]));
Rrates = c(Rrates, asFloat(components[1]));
Mrates = c(Mrates, asFloat(components[2]));
Exomelength = ends[size(ends)-1];
initializeRecombinationRate(Rrates, ends);
initializeMutationRate(Mrates, ends);
initializeSex("A"); // Specifies modeling of an autosome
initializeMutationType("m1", 0.5, "g", -0.043, 0.23); //non-synonymous
initializeMutationType("m2", 0.5, "f", 0.0); // synonymous
m1.mutationStackPolicy = "1";
m2.mutationStackPolicy = "1";
initializeGenomicElementType("g1", m1, 1); // positions 1 and 2
initializeGenomicElementType("g2", m2, 1); // positions 3
starts = repEach(seqLen(asInteger(round(Exomelength/3))) * 3, 2) +
  rep(c(0,2), asInteger(round(Exomelength/3)));
end pos = starts + rep(c(1,0), asInteger(round(Exomelength/3)));
types = rep(c(g1,g2), asInteger(round(length(starts)/2)));
initializeGenomicElement(types, starts, end_pos);
```

```
// Initialize the ancestral African population
1 { sim.addSubpop("p1", asInteger(round(7310.370867595234))); }
// End the burn-in period; expand the African population
73105 { p1.setSubpopulationSize(asInteger(round(14474.54608753566))); }
// Split Eurasians (p2) from Africans (p1) and set up migration
76968 {
sim.addSubpopSplit("p2", asInteger(round(1861.288190027689)), p1);
p1.setMigrationRates(c(p2), c(15.24422112e-5));
p2.setMigrationRates(c(p1), c(15.24422112e-5));
// Split p2 into European (p2) and East Asian (p3); resize; migration
78084 {
sim.addSubpopSplit("p3", asInteger(round(553.8181989)), p2);
p2.setSubpopulationSize(asInteger(round(1032.1046957333444)));
p1.setMigrationRates(c(p2, p3), c(2.54332678e-5, 0.7770583877e-5));
p2.setMigrationRates(c(p1, p3), c(2.54332678e-5, 3.115817913e-5));
p3.setMigrationRates(c(p1, p2), c(0.7770583877e-5, 3.115817913e-5));
// Set up exponential growth in Europe (p2) and East Asia (p3)
78084:79012{
t = sim.generation - 78084;
p2\_size = round(1032.1046957333444 * (1 + 0.003784324268)^t);
p3_size = round(553.8181989 * (1 + 0.004780219543)^t);
p2.setSubpopulationSize(asInteger(p2_size));
p3.setSubpopulationSize(asInteger(p3_size));
// Create the admix population
79012 early(){
sim.addSubpop("p4", 30000); //This new subpopulation is created with 30000 new empty individuals
p4.setMigrationRates(c(p1, p2, p3), c(0.1666667, 0.3333333, 0.5));
}
//After this early() event, SLiM will generate offspring, and the empty individuals in p4 will be
// discarded and replaced by migrant offspring from p1, p2 and p3 as requested.
79012 late(){
p4.setMigrationRates(c(p1, p2, p3), c(0, 0, 0));
// Set up exponential growth in admixture (p4)
79012:79024 {
t = sim.generation - 79012;
p4_new_size = round(30000 * (1 + 0.05)^t);
p4.setSubpopulationSize(asInteger(p4_new_size));
// Output and terminate
79024 late() {
p4.individuals.genomes.output(filePath = "~/Output_Full.txt");
```

}

Before the simulation starts, we need to initialize the mutation rate, recombination rate, genomic structure and so forth as the simulation parameters (Haller et al. 2019). We read the recombination map into SLiM using the readFile() function. Inside this function, we supply the path to our recombination map text file. Then we create three null vectors named Rrates, Mrates and ends to save the recombination rates, mutation rates and end positions of each exon in our recombination map, respectively.

Next, we use a for-loop to move along the genome, reading each line of the recombination map and: - save the recombination rate, mutation rate and end position of each genomic segment, - initialize the recombination rate for each genomic segment with the initializeRecombinationRate() function, by specifying the rate and the end position of the genomic segment, - initialize the mutation rate for each genomic segment with the initializeMutationRate() function, - specify that the genomic segment belongs to an autosomal chromosome with the initializeSex() function, - specify the mutation type for each genomic segment with the initializeMutationType() function (see below), - specify the mutation stacking policy for each genomic segment with the mutationStackPolicy command (see below), - specify the type for each genomic segment with the initializeGenomicElementType() function (see below).

In exons, the last base-pair position in a three base-pair codon (coding for an amino acid in a protein) is a synonymous site. Synonymous sites are viewed as selectively neutral in comparison to the first two base-pair positions in a codon, which are non-synonymous. Therefore, we simulate two types of mutations: synonymous and non-synonymous. The initializeMutationType("m1," 0.5, "g," -0.043, 0.23) callback in the for-loop explains all the parameters that are held by the "m1" mutation type. We use "m1" to represent the non-synonymous mutations. These non-synonymous mutations have a dominance coefficient of 0.5 and the selection coefficient is generated from a gamma distribution with mean -0.043 and shape parameter is 0.23 (Harris and Nielsen 2016). We initialize the synonymous mutations separately with another call to the initializeMutationType() function. In initializeMutationType("m2," 0.5, "f," 0.0) callback, the "m2" mutation type represents the synonymous mutations and they have a fixed selection coefficient denoted by "f." The selection coefficient of this type of mutation is always 0, as seen in the fourth argument of the function. The dominance coefficient in the second argument of the function is 0.5.

In SLiM (as in biology), the individuals rather than the mutations are under selection. Selection acts on the individual, through their fitness value. The fitness value of an individual is calculated from the fitness effects of all the mutations carried by that individual (based upon their selection coefficient, dominance coefficient, and heterozygous/homozygous state). All the fitness effects are multiplied together to produce the individual fitness. The individual fitness value then affects selection. Specifically, in the default Wright-Fisher (WF) model of SLiM (which we use), lower fitness means a lower probability of mating. As a result, deleterious mutations tend to decrease in frequency and beneficial mutations tend to increase in frequency.

SLiM allows for recurrent mutations at a given base position on a given sequence (Haller et al. 2019). By default, SLiM "stacks" any mutations that occur in the same location as pre-existing mutations on a given sequence. This default behaviour of "mutation stacking" ("s" for stacked), is changed to "l" (last) with the command m1.mutationStackPolicy = "l", so that new mutations occurring in the same location as pre-existing mutations on a given sequence replace the pre-existing mutations.

The next initialization task is to create the chromosome structure. In SLiM we can model different genomic structures in the chromosomes. We consider exons only, which have two genomic element types: one for non-synonymous sites (base positions 1 and 2 of a codon) and the other for synonymous sites (base position 3 of a codon). These genomic element types are called "g1" and "g2" and alternate as g1, g2, g1, g2, g1, etc. along the exome until the end position of a chromosome is reached. The first genomic-element type corresponds to non-synonymous sites, is initialized as "m1" and could have mutations with selection coefficients that come from the negative gamma distribution. The second genomic element type corresponds to the synonymous sites, is initialized as "m2" and could have neutral mutations. We use the initializeGenomicElementType() function to specify these two genomic elements and our exome structure. For example, initializeGenomicElementType("g1", m1, 1) specifies that genomic element type "g1" is defined as using mutation type "m1" for all of its mutations. The second genomic element type "g2" is defined

as using mutation type "m2" for all its mutations. Then we create the alternating start and end positions of the "g1" and "g2" genomic elements along the exome. Finally, we initialize the two genomic elements "g1" and "g2" with initializeGenomicElement(), supplying their starting and ending positions along the exome.

- After the initialize() callbacks end, we run the first generation of our simulation.
- We use the Out-of-Africa model described in the SLiM manual (Haller et al. 2019). Note that the SLiM manual uses the exact parameter estimates from the original model described in Gravel et al. (2011).
- In the first generation the first sub population is created with the inferred initial size and we label it as "p1."
- We use the function sim.addSubpop() to add "p1," the new sub population, by giving its initial size.
- Haller et al. (2019) start the model at 79024 generations back (approximately 1976 kya); this is generation 0 in the model.
- The model then takes 10\*African ancestral population size generations as the neutral burn-in time (Haller et al. 2019). An expansion of the African population occurs at 73105 generations.
- Therefore, at 73105 generations, we change the sub-population size of "p1" (Africa) from  $\sim$  7310 to  $\sim$  14474.
- After this event, at generation 76968, the African sub-population splits into the Eurasian ancestral sub-population.
- We create another sub-population, "p2," to represent the Eurasian ancestral sub-population and migration is started between these two populations (Haller et al. 2019).
- p1.setMigrationRates sets the migration rate from the African to the Eurasian ancestral subpopulation, while p2.setMigrationRates sets the migration rate from the Eurasian ancestral to the African sub-population.
- Then at 78084 generations, the "p2" Eurasian sub-population splits into European and Asian sub-populations. Therefore, we create a new sub-population, "p3," to represent the Asian sub-population. The Eurasian ancestral sub-population becomes the European sub-population.
- After that, we set the migration rates between three sub-populations accordingly.
- Then we set up exponential growth in European (p2) and Asian (p3) sub-populations starting from 78084 generations to 79012 generations.
- At 79012 generations, we create our admixed sub-population with an initial size 30000 and set the migration rates between the admixed and the other three sub-populations as given in Browning et al. (2018). After we create the admixed sub-population, we stop all migrations.
- Then between 79012 to 79024 generations we set exponential growth of this admixed sub-population with a rate of 5% (individuals per generation).
- Finally in generation 79024 we terminate our SLiM simulation and collect our output from the model.
- Since our main study is based on the Northern American population, we only consider the American admixed sub-population as the output.
- Therefore, we use the function p4.individuals.genomes.output() to get the all individual's genomes of the admixed sub-population.
- The following example represents the format of the output returned from the p4.individuals.genomes.output() function

 $\#OUT: 79024 \text{ GS } 107752 / \text{project} / 6007536 / \text{epasiedn} / \text{SLiM} / \text{American} Admixture} / \text{Output} _Full.txt Mutations:}$ 

7229 50171 m2 51287555 0 0.5 p1 5 60626 ...

```
\begin{array}{c} 13218\ 484904\ \mathrm{m2}\ 39812003\ 0\ 0.5\ \mathrm{p1}\ 45\ 9536\ \dots \\ 5202\ 762125\ \mathrm{m2}\ 36490340\ 0\ 0.5\ \mathrm{p1}\ 70\ 64099\ \dots \\ \\ \dots \\ \text{Genomes:} \\ p^*:0\ A\ 0\ 1\ 2\ 3\ 4\ 5\ 6\ 7\ 8\ 9\ 10\ 11\ \dots \\ p^*:1\ A\ 10605\ 1\ 2\ 3\ 10606\ 4\ 5\ 6\ 8\ 10607\ 10608\ 9\ \dots \\ p^*:2\ A\ 10605\ 1\ 2\ 4\ 15639\ 15640\ 6\ 10608\ 15641\ 15642\ 15643\ 15644\\ p^*:3\ A\ 0\ 1\ 2\ 19096\ 19097\ 4\ 6\ 19098\ 19099\ 9\ 10\ 19100\\ \dots \end{array}
```

- In the above example output, the first row starts with # OUT: and this is followed by the generation (79024) in which the output is generated. Then "GS" represents the "genomes SLiM format" and this is followed by the sample size in number of genomes (2\* number of individuals). Finally, the full path where we save the output is printed.
- Then the second line starts the mutation section.
- In the mutation section each row represents a mutation which is currently segregating in the population and the nine columns represent the mutation properties.
- The first column is the SLiM-generated identifier number which helps to identify the mutation easily. The second column is the mutation's identification number. The third field represents the type of the mutation. The fourth column is the base-pair position of the mutation on the chromosome. Fifth and sixth columns represent selection and dominance coefficients, respectively. The seventh column is the sub-population in which the mutation originated. The eighth column is the generation when the mutation arose. Finally, the ninth column represents the number of copies of the mutation in the sub-population.
- The last section in the output represents the genomes section.
- In the genome section a row corresponds to a genome in the sub-population and specifies all the mutations carried by the genome.
- Because we use p4.individuals.genomes.output(), we get all the genomes in p4 sub-population. The first line in the genomes section in the above example, "p\*: 0," means the 0th genome of the sub-population. Then "A" represents autosome, the type of the genome. This is followed by the SLiM-generated identification numbers of all the mutations carried by this genome. Recall that the SLiM-generated identification numbers are in the first column in the mutation section.

## 3.1 Simulation on Compute Canada Cluster

- This SLiM simulation is highly memory intensive and not suitable for most personal computers.
- We therefore use the Compute Canada cluster (http://www.computecanada.ca) as described next.
- First, we need to install the SLiM software. The way we install the software is exactly the same as how we install the software on our own computer. Use the SLiM manual guidelines for this task.
- After we install SLiM, we use a job scheduler in the cluster to run our jobs.
- In the Compute Canada Cluster we use the **Slurm Workload Manager** as the job scheduler.
- Slurm helps to allocate resources and time, and provides methods to execute our work.
- To run the SLiM script we use a Slurm script as follows.

```
#!/bin/bash

#SBATCH --account=def-jgraham

#SBATCH --ntasks=1

#SBATCH --time=7-05:05:00

#SBATCH --mem=64000M
```

```
module load StdEnv/2020 gcc/9.3.0 slim/3.4.0 slim SLiM_American_Admixture.slim
```

- Let's understand each line of the above batch script file.
- #SBATCH-account=def-jgraham specifies the account name. In this example, the account name is "def-jgraham."
- #SBATCH-ntasks=1 defines the number of processors. We request 1 processor to run the program.
- #SBATCH-time=7-05:05:00- specifies the time limit for the job. Usually, we allocate a time which is more than the expected time to run the program. This is because, if our simulation takes a longer time than the allocated time, the simulation will stop without executing when the given time limit is achieved.
- #SBATCH-mem=640000M specifies memory that we require to run our simulation. We request 64GB.
- Next, the executable commands are aligned in the script file. They are:
- module load StdEnv/2020 gcc/9.3.0 slim/3.4.0 loads the SLiM version which we installed.
- slim SLiM\_American\_Admixture.slim-calls for SLiM to run the SLiM script file: SLiM\_American\_Admixture.slim. This is the final command in the slurm script file.
- Then we use sbatch command to submit above slurm script file to run in the cluster. We type the sbatch command in our log in node as follows.

#### [epasiedn@gra-login2 American\_Admixture] \$ sbatch job\_serial.sh

- This simulation took approximately 3 days to complete.
- In our job script we allocate 64GB to run the simulation. Out of this 64GB, the job utilized 43.61GB.
- The output of this SLiM model is saved as SLiM\_output.txt and we use this file as one of the inputs for the third phase of our work flow.
- We will examine the SLiM\_output.txt output in our third R-markdown document.

#### 3.2 Summary Statistics

## Attaching package: 'Matrix'

• We obtain summary statistics of the American admixed population by using the SLiM output.

```
library(SimRVSequences)
library(tidyverse)
## -- Attaching packages -----
                                       ----- tidyverse 1.3.1 --
## v ggplot2 3.3.5
                     v purrr
                               0.3.4
## v tibble 3.1.4
                     v dplyr
                               1.0.7
## v tidyr
            1.1.3
                      v stringr 1.4.0
## v readr
            2.0.1
                     v forcats 0.5.1
## -- Conflicts -----
                                           ----- tidyverse conflicts() --
## x dplyr::filter() masks stats::filter()
## x dplyr::lag()
                   masks stats::lag()
library(Matrix)
```

```
## The following objects are masked from 'package:tidyr':
##
       expand, pack, unpack
##
library(data.table)
##
## Attaching package: 'data.table'
## The following objects are masked from 'package:dplyr':
##
       between, first, last
##
## The following object is masked from 'package:purrr':
##
##
       transpose
\# Read the SLiM output text file to R
exDat <- readLines("D:/SFU_Vault/SLiM_Output/Output_Full.txt")</pre>
```

- The SLiM\_output.txt file size is approximately 6 GB.
- We read the SLiM\_output.txt file to R and it takes approximately 1 minute to load to R on a Windows OS with an i7-8550U @ 1.8GHz,16GB of RAM.

#### ## [1] 53876

• The size of the simulated American admixed population is 53,876.

#### ## [1] 862243

- There are 862,243 number of mutations are currently segregating in the admixed population.
- Then we examine among these 862,243 mutations the percentage of allele frequency less than 1% in the population.

```
# Add 1 to temp ID so that we can easily associate mutations to columns.
# By default SLiM's first tempID is 0, not 1.
MutData$tempID <- MutData$tempID + 1
# First position in SLiM is 0, not 1
MutData$position <- MutData$position + 1

# Calculate the population derived allele frequency.
# Divide the allele count by the population size.
MutData$afreq <- MutData$count/(popCount)

# Get the percentage of SNVs whose allele frequency < 0.01
af_less <- which(MutData$afreq < 0.01)
af_less_per <- length(af_less)/ nrow(MutData)
af_less_per</pre>
```

#### ## [1] 0.9426565

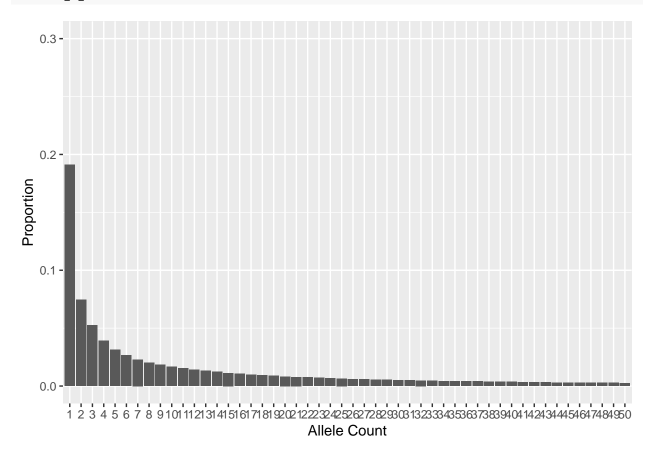
- Among these mutations, approximately 94% have frequencies less than 1%.
- We compare these results with the TopMed study (Taliun et al. 2021). The TopMed study has newly released empirical results for the American population and suggests that the majority of variants have alternate allele frequencies of less than 1%. According to Taliun et al. (2021), approximately 97% of variants in the TopMed Study are rare (i.e. have allele frequencies less than 1%).
- In our American admixed population also, we find that approximately 94% of the mutations are rare.
- We use the following code chuck to check the singleton percentage in the American admixed population.

```
# Use the prevalence (the number of times that the mutation occurs in any genome)
# column in MutData dataframe to calculate the singleton percentage
singelton <- MutData %>%
    count(count) %>%
    mutate(percentage = n/nrow(MutData))

colnames(singelton) <- c("number_of_allele", "count", "proportion")
head(singelton)</pre>
```

• Among 862,243 mutations, 19% of them are singletons. The following figure illustrates the allele frequency spectrum.

```
ylim(0, 0.3) +
scale_x_discrete(limits= as.character(1:50))
```



- In the TopMed study, about half of the variants are singletons (Taliun et al. 2021).
- The reason for the discrepancy between our results and TopMed is our lack of source populations. In our SLiM model we have only three source populations and we collect SNV data from the American admixed population only. But in the TopMed study, they consider the entire American population which consists of many more source populations.
- To check this, we combine data from all four populations in our SLiM model and check if the singleton percentage is increased. Due to the high computational cost we only simulate data from chromosome 8 and 9.
- Combining all four populations, around 26% of variants are singletons.
- The supplementary material 1-B discusses the commands to generate and summarize all the source populations and check the proportions of singletons after combining the four populations.

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