Supplementary Material 1-B: Combining Source Populations with the American-Admixed Population in SLiM

Nirodha Epasinghege Dona, Jinko Graham

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Supplementary Material 1-A discusses the SLiM simulation of the American-admixed population and the processing of its data. This document discusses how to obtain and process the data not just for the American-admixed population but also for all four populations in the simulation. We focus on chromosomes 8 and 9 only to reduce computational cost, and alter the original SLiM script as follows: 1. at the top of the script, we provide an abbreviated recombination map, Slim_Map.txt, containing only those lines of the original recombination map (in the data frame, s_map) pertaining to chromosomes 8 and 9, and 2. at the bottom of the script, we call sim.outputFull() rather than p4.individuals.genomes.output() to obtain the output. Otherwise, the SLiM script remains exactly the same as the original in Supplementary Material 1-A.

```
initialize() {
// Read abbreviated recombination map for chromosomes 8 and 9.
lines = readFile("~/Slim_Map.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 admixed population
79012{
p2_new_size = p2.individualCount;
p3_new_size = p3.individualCount;
defineConstant("pop_size", c(p2_new_size, p3_new_size));
sim.addSubpop("p4", 30000);
p4.setMigrationRates(c(p1, p2, p3), c(0.1666667, 0.3333333, 0.5));
79012 late(){
p4.setMigrationRates(c(p1, p2, p3), c(0, 0, 0));
```

```
// Setup exponential growth in Europe (p2) and East Asia (p3)
79012:79024 {
t = sim.generation - 79012;
p2_{new_size} = round(pop_size[0] * (1 + 0.003784324268)^t);
p3_{new_size} = round(pop_size[1] * (1 + 0.004780219543)^t);
p4_{new_size} = round(30000 * (1 + 0.05)^t);
p2.setSubpopulationSize(asInteger(p2_new_size));
p3.setSubpopulationSize(asInteger(p3_new_size));
p4.setSubpopulationSize(asInteger(p4_new_size));
// Output for all populations (not just p4) and terminate
79024 late() {
sim.outputFull("~/SLiM_output_chr8&9.txt");
}
We read SLiM_output_chr8&9.txt into R and obtain the number of individuals in each population and in
all populations combined.
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)
##
## Attaching package: '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
# Note: Change the path for the file as necessary.
exDat <- readLines("D:/SFU_Vault/SLiM_Output/SLiM_output_chr8&9.txt")
```

```
# Read the mutations and genomic sections in the output
MutHead <- which(exDat == "Mutations:")
GenHead <- which(exDat == "Genomes:")
PopHead <- which(exDat == "Populations:")
IndHead <- which(exDat == "Individuals:")

# Get the population count for each source population

popCount_1 <- as.numeric(unlist(strsplit(exDat[PopHead + 1], split = " "))[2])
popCount_2 <- as.numeric(unlist(strsplit(exDat[PopHead + 2], split = " "))[2])
popCount_3 <- as.numeric(unlist(strsplit(exDat[PopHead + 3], split = " "))[2])
popCount_4 <- as.numeric(unlist(strsplit(exDat[PopHead + 4], split = " "))[2])

# Get the total population count
popCount <- popCount_1 + popCount_2 + popCount_3 + popCount_4</pre>
```

The following table of population sizes summarizes the output of the above commands.

Table 1: Population sizes.

| Population | size |
|------------|---------|
| African | 14,475 |
| European | 35,815 |
| Asian | 48,765 |
| Admix | 53,876 |
| Total | 152,931 |

[1] 142549

On chromosomes 8 and 9, the number of mutations segregating in all four populations is 142,549.

```
# 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 derived-allele count by the population size.

MutData$afreq <- MutData$count/(popCount)

# Get the percentage of SNVs whose derived-allele frequency is < 0.01
```

```
af_less <- which(MutData$afreq < 0.01)
af_less_per <- length(af_less)/ nrow(MutData)
af_less_per</pre>
```

[1] 0.9616693

Among the 142,549 mutations on chromosomes 8 and 9, approximately 96% have frequencies less than 1%.

In Supplementary Material 1-A, we discuss how 26% of the variants in the combined populations were singletons. The following commands are used to calculate this percentage.

```
# Use the prevalence (the number of times that the mutation occurs in any genome)
# column in MutData dataframe to calculate the singleton percentage
singleton <- MutData %>% count(count) %>% mutate(percentage = n/nrow(MutData))
colnames(singleton) <- c("number_of_allele", "count", "proportion")
head(singleton)</pre>
```

The following figure illustrates the derived-allele frequency spectrum for chromosomes 8 and 9.

