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

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Commands for calculating summary statistics in the combined populations

- In Supplementary Material 1-A, we discuss the SLiM American admixed demographic model. However, in that we output only the American admixed population data because it is our main interest.
- In this document, we discuss the SLiM model which outputs the data for all four populations. We also show the commands to calculate summary statistics.
- The following is the SLiM script that we use to simulate all four populations in the American admixture demographic model. Note: We only simulate chromosome 8 and 9 exome data due to high computational cost.

```
initialize() {
// Read recombination map cretated by SimRVSequence R package
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 admix population
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 and terminate
79024 late() {
sim.outputFull("~/output_all_chr8&9.txt");
}
```

• We read the file output_all_chr8&9.txt into R and obtain the summary statistics that were described in the Supplementary Material 1-A document.

```
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 exDat <- readLines("D:/SFU_Vault/SLiM_Output/Output_all_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</pre>

• The following table illustrates the population size of each population.

popCount <- popCount_1 + popCount_2 + popCount_3 + popCount_4</pre>

Table 1: Population sizes.

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

[1] 142549

• There are 142,549 number of mutations are currently segregating.

```
# 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] 0.9616693

- Among all the mutations, 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
singelton <- MutData %>% count(count) %>% mutate(percentage = n/nrow(MutData))
colnames(singelton) <- c("number_of_allele", "count", "proportion")
head(singelton)</pre>
```

• The following figure illustrates the frequency spectrum.

