Relate practical: Using genealogies for population genetics

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In this practical, we will infer genealgies for the Simons Genome Diversity Project dataset, downloaded from https://reichdata.hms.harvard.edu/pub/datasets/sgdp/. This dataset comprises wholegenome sequencing data of 278 modern humans with sampling locations shown in Fig. 1.

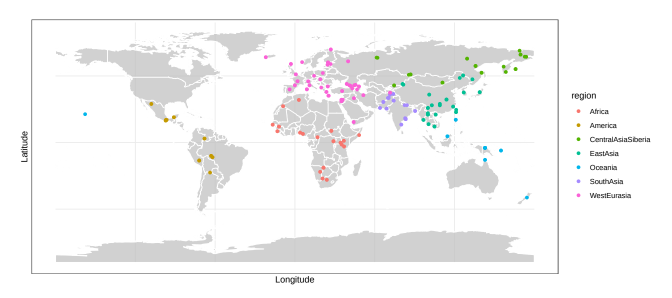


Figure 1: Sampling locations of the 278 modern humans in the Simons Genome Diversity Project. Samples are classified into seven regions shown by colours.

Note 1

The data was downloaded from:

- Phased genotypes: https://sharehost.hms.harvard.edu/genetics/reich_lab/sgdp/phased_data/PS2_multisample_public
- Genomic mask: https://reichdata.hms.harvard.edu/pub/datasets/sgdp/filters/all_samples/
- Human ancestral genome: ftp://ftp.1000genomes.ebi.ac.uk/vol1/ftp/phase1/analysis_results/supporting/ancestral_alignments/
- Recombination maps: https://mathgen.stats.ox.ac.uk/impute/1000GP_Phase3.html

We then used RelateFileFormats --mode ConvertFromVcf to convert to the haps/sample file format and the PrepareInputFiles.sh script to make sure ancestral alleles are denoted by 0 and to filter out regions according to the genomic mask.

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1 Relate

Relate estimates the joint genealogies of many thousands of modern individuals genome-wide. These genealogies describe how individuals are related through their most-recent common ancestors back in time and can be seen as the genetic analogue of a family tree for unrelated individuals.

The output of Relate is a sequence of binary trees, each describing the genealogical relationships locally in that part of the genome. Neighbouring genealogical trees differ because of recombination events that change the genetic relationships of individuals.

Note 2

A detailed documentation for Relate is available at https://myersgroup.github.io/relate.

Details about the method is published in:

- L. Speidel, M. Forest, S. Shi, S. R. Myers. A method for genome-wide genealogy estimation for thousands of samples. Nature Genetics **51**, 1321–1329 (2019).
- Leo Speidel, Lara Cassidy, Robert W. Davies, Garrett Hellenthal, Pontus Skoglund, Simon R. Myers. Inferring population histories for ancient genomes using genome-wide genealogies. Molecular Biology and Evolution 38, 3497–3511 (2021)

For this practical, we will use Relate binaries, bash scripts, and R.

The Relate binaries can be found in my home directory or downloaded from

https://myersgroup.github.io/relate. It will be convenient to define a variable storing the location of the Relate binaries, e.g.,

#adjust to where your relate binaries are stored
PATH_TO_RELATE="/home/wpsg/software/relate/"

1.1 Data requirements and file formats

Relate uses the **haps/sample** file format (output file format of SHAPEIT2) as input (see https://myersgroup.github.io/relate/input_data.html#FileFormat).

- You can convert from **vcf** to haps/sample and from **hap/legend/sample** to haps/sample using functions provided by Relate.

 $(see \ \mathtt{https://myersgroup.github.io/relate/input_data.html\#ConvertToHapsSample})$

Note 3: Data requirements

- Please separate your data by chromosome.
- Relate assumes whole-genome sequencing data as input.
- The data needs to be phased. This can be done, for instance, using SHAPEIT.
- Ancestral and derived states at single-nucleotide polymorphisms (SNPs) need to be known.
- A genomic mask should be specified that filters out "bad" regions

We provide a script to prepare your data under

\${PATH_TO_RELATE}/scripts/PrepareInputFiles/PrepareInputFiles.sh

(see https://myersgroup.github.io/relate/input_data.html#Prepare).

1.2 The arguments

Required arguments

mode	Mode in which to run Relate. Mode "All" will execute all stages of the algorithm
	and other modes can be used to execute individual stages (see here). This is
	useful, for instance, when parallelizing Relate.
-m,mutation_rate	Mutation rate per base per generation.
-N,effective_size	Haploid effective population size. (For diploid organisms, multiply the effective
	population size of individuals by 2)
haps	Filename of haps file.
sample	Filename of sample file.
map	Filename of genetic recombination map.
-o,output	Filename of output files without file extension.

Note 4: Optional arguments

There are other optional arguments for Relate, please consult the documentation for these (https://myersgroup.github.io/relate/getting_started.html#GettingStarted).

2 Running Relate on the Simons Genome Diversity Project dataset

Let's start with running Relate. Please copy the worksheet_data directory to your home directory

```
#assuming you are in your home directory
cp -r /home/wpsg/workshop_materials/a15_relate .
cd a15_relate
```

In this directory, you will find three subdirectories, named data (containing all input data), precomp_results (containing precomputed Relate output files), and Rscripts (containing R scripts for plotting results).

2.1 Running Relate

Note 5

Running Relate on all 278 samples and four cores took 52 minutes, so here we will run Relate only on the African samples of this dataset.

We run Relate on chromosome 15 for the 44 African samples of the Simons Genome Diversity Project.

```
#assuming you are under worksheet_data
mkdir results
cd results
lab="Africa"
chr=15
#This takes about 4 minutes
${PATH_TO_RELATE}/bin/Relate \
     --mode All \
     --haps "../data/data_subgroups/SGDP_input_${lab}_chr${chr}.haps.gz" \
     --sample "../data/data_subgroups/SGDP_input_${lab}_chr${chr}.sample.gz" \
     --map "../data/genetic_map_chr${chr}_combined_b37.txt" \
     --dist "../data/data_subgroups/SGDP_input_${lab}_chr${chr}.dist.gz" \
     -m 1.25e-8 \
     -N 30000 \
     -o "SGDP_${lab}_chr${chr}"
gzip SGDP_${lab}_chr15.*
```

The input haps/sample files store the genetic variation of our samples (similar to a vcf file). These are obtained from the PrepareInputFiles.sh script (see above). The genetic recombination map stores recombination rates along the genome. In addition, we specify a .dist file to adjust the distances (in units of BP) between SNPs removing regions of lower quality specified in the mask – this file is also outputted by the PrepareInputFiles.sh script and is necessary to adjust mutation rates (we can only observe mutations in regions that pass the filters of the mask).

Exercise 1

Let's plot a few trees. We use the TreeView.sh script provided with Relate: For this, we need an additional file storing assignment of individuals to populations. This file has four columns, named "sample", "population", "group", and "sex". For us, only the second column is of interest. The order in which individuals are listed has to be consistent with the order in the samples file.

Example:

```
ID POP GROUP SEX

S_Mozabite-1 Mozabite Africa NA

S_Mozabite-2 Mozabite Africa NA

S_Saharawi-1 Saharawi Africa NA

S_Saharawi-2 Saharawi Africa NA
```

We can now run the TreeView.sh script as follows (this will use R and requires ggplot2 and cowplot - these will be installed if missing):

```
${PATH_TO_RELATE}/scripts/TreeView/TreeView.sh \
     --haps "../data/data_subgroups/SGDP_input_Africa_chr${chr}.haps.gz" \
     --sample "../data/data_subgroups/SGDP_input_Africa_chr${chr}.sample.gz" \
     --anc "SGDP_Africa_chr15.anc.gz" \
     --mut "SGDP_Africa_chr15.mut.gz" \
     --poplabels "../data/poplabels/SGDP_Africa_lang.poplabels" \
     --bp_of_interest 200000000 \
     --years_per_gen 28 \
     -0 "SGDP_Africa_chr${chr}_BP20000000"
```

This will produce a pdf named SGDP_Africa_chr15_BP20000000.pdf. We can also plot trees for the precomputed trees of West Eurasian SGDP individuals:

Can you see anything unusual about this tree. You can also rerun the above replacing TreeView.sh by TreeViewMutation.sh; this will highlight branches carrying the mutation at chromosome 15, BP 48426484.

Feel free to plot a few other trees!

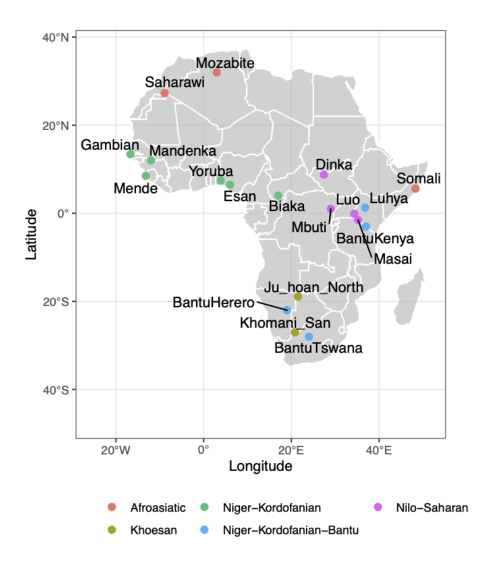


Figure 2: Map showing SGDP groups in Africa, coloured by language family.

3 Effective population sizes and split times

Note 6

We will use the genealogies for Africans obtained in Section 2.1 as an example here. You can replicate the analysis for other subgroups, or for all samples; each step will take longer for larger sample sizes (for all samples, computation time in Section 3.1 is around 15 min and for Section 3.2 around 70 min).

3.1 Estimating population sizes given a genealogy

We infer effective population sizes and split times from estimated genealogies of Africans obtained in Section 2.1. Extracting effective population sizes from genealogies is done as follows

```
#assuming you are in worksheet_data
mkdir popsizes

cd popsizes
lab="Africa"
chr=15

${PATH_TO_RELATE}/bin/RelateCoalescentRate \
    --mode EstimatePopulationSize \
    -i "../results/SGDP_${lab}_chr${chr}" \
    -o "SGDP_${lab}_chr${chr}" \
    --bins 3,7,0.25 \
    --poplabels "../data/poplabels/SGDP_${lab}.poplabels"
```

This will produce a file named SGDP_Africa_chr15.coal and SGDP_Africa_chr15.bin. The SGDP_Africa_chr15.coal stores cross-coalescence rates for all pairs of group labels in SGDP_Africa.poplabels of which there are 19. This might be too many to visualise easily, so we will aggregate some of these groups based on language family. The .bin file can be used in combination with different poplabels files to extract coalescence rates, e.g.

```
${PATH_TO_RELATE}/bin/RelateCoalescentRate \
    --mode FinalizePopulationSize \
    -o "SGDP_Africa_chr${chr}" \
    --poplabels "../data/poplabels/SGDP_Africa_lang.poplabels"
```

Exercise 2

Let's plot the estimated effective population sizes stored in SGDP_Africa_chr15.coal in R. You can use relater, which is an R package that can parse and manipulate Relate output files (preinstalled, available at https://github.com/leospeidel/relater):

```
library(relater)
library(ggplot2)
coal <- read.coal("SGDP_Africa_chr15.coal")
#diploid effective population size is the 0.5* inverse coalescence rate
coal$popsize <- 0.5/coal$haploid.coalescence.rate
#multiply epochs times by 28 to scale to years (assuming 28 years per generation)
coal$epoch.start <- 28 * coal$epoch.start

head(coal) #contains all vs all coalescence rates
#We subset to within group coalescence rates, but feel free to look at cross-population
    rates too
coal <- subset(coal, group1 == group2)
p <- ggplot(coal) +
    geom_step(aes(x = epoch.start, y = 0.5/haploid.coalescence.rate, color = group2)) +
    scale_x_continuous(trans = "log10", limit = c(1e2, 1e7)) +
    scale_y_continuous(trans = "log10", limit = c(1e3,1e7))</pre>
```

Can you describe how population sizes of groups speaking Afroasiatic languages differ from other groups?

3.2 Joint fitting of population size and branch lengths

So far, we used trees that assumed a pre-specified constant effective population size through time. Next, we use the EstimatePopulationSize.sh script to jointly fit effective population sizes and branch lengths.

```
chr=15
lab=Africa

# This will take around 4min

${PATH_TO_RELATE}/scripts/EstimatePopulationSize/EstimatePopulationSize.sh \
    -i "../results/SGDP_${lab}_chr${chr}" \
    -o "SGDP_${lab}_ne_chr${chr}" \
    --poplabels "../data/poplabels/SGDP_${lab}_lang.poplabels" \
    -m 1.25e-8 \
    --years_per_gen 28 \
    --bins 3,7,0.25 \
    --num_iter 2 \
    --threads 4
```

This will output SGDP_Africa_ne_chr15.anc.gz, SGDP_Africa_ne_chr15.mut.gz, and SGDP_Africa_ne_chr15.pairwise.coal. (The file SGDP_Africa_ne_chr15.coal stores coalescence rates grouping all samples into one group and can be ignored here.)

Note 7

To run this on more than one chromosome, the --chr argument can be useful (see documentation https://myersgroup.github.io/relate/modules.html#PopulationSizeScript.)

Exercise 3

- (i) Similarly to Exercise 2, let's plot the effective population sizes for SGDP_Africa_ne_chr15. pairwise.*. How is this plot different to the one generated in Exercise 2 (which assumed a constant population size through time)?

 (If you like, you can use the script under Rscripts/.)
- (ii) Now, let's get coalescence rates between all pairs of African individuals from SGDP_Africa_ne_chr15.pairwise using the poplabels file ../data/poplabels/SGDP_Africa_ind.poplabels. We can plot coalescence rates in a heatmap for different epochs to visualise how structure has evolved through time.

 (If you like, you can use the script under Rscripts/.)

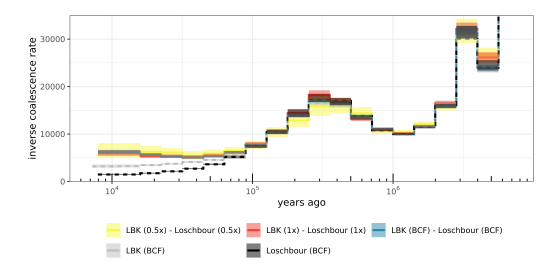


Figure 3: Illustration of what Colate can be used for. Here, we calculated coalescence rates between two ancient human individuals which were external to our inferred Relate genealogy. We downsampled both genomes, showing that the method works for low-coverage whole-genome sequencing data.

3.3 Colate: Inferring coalescence rates for low-coverage genomes

So far we have computed coalescence rates for samples built into the genealogies. However, using a tool called Colate (https://github.com/leospeidel/Colate), we can use the inferred genealogies to compute coalescence rates of external samples. This is particularly useful for samples that were sequenced to low coverage. Here, we use data of an early Neolithic farmer (Lazaridis et al., Nature 2014) and calculate coalescence rates against samples in our genealogy, but you could also calculate rates between two external samples.

Let us first define the following global variables in our bash terminal:

```
#assuming you are under worksheet_data
mkdir colate
cd colate
# path to reference genome, one fasta per chromosome
ref_genome="../data/GRCh37_chr15.fa.gz"
```

Genealogies date mutations segregating in the SGDP dataset and we use these in **Colate**. The **Relate** output files need to first be preprocessed using Colate --mode preprocess_mut; we have already done this and you can find the resulting file under precomp_results/Colate_ref_genealogy/. You can also download these from https://www.dropbox.com/s/pysvzyx68553wxx/SGDP_mutages.tar?dl=0.

We will first convert the BAM file of our ancient individual into Colate input format and will do the same with BCF files of SGDP subpopulations.

```
echo 15 > chr.txt #make a file containing chromosome names to be used
# path to mut file, which contains mutation dates.
# We use a precomputed file computed using all SGDP samples and subsetted to transversions
mut="../precomp_results/Colate_ref_genealogy/SGDP_half_ne_fixed_transv"
#compute Colate input files from BAM format
Colate --mode make_tmp \
    --mut ${mut} \
    --target_bam ../data/ancients/LBK_1x_chr15.bam \
    --ref_genome ${ref_genome} \
    --chr chr.txt \
    -o SGDP_LBK_1x
for pop in Africa WestEurasia SouthAsia EastAsia Oceania CentralAsiaSiberia America
do
  Colate --mode make_tmp \
    --mut ${mut} \
    --target_bcf ../data/data_subgroups/SGDP_input_${pop} \
    --ref_genome ${ref_genome} \
    --chr chr.txt \
    -o SGDP_${pop}
done
```

Now that we have the input files, we can compute coalescence rates between pairs of groups. In addition to these input files, we provide several parameters, including the age of samples (in this case our sample is 7244 years old), and the number of block-bootstrap iterations (here fixed to 1).

```
for pop in Africa WestEurasia SouthAsia EastAsia Oceania CentralAsiaSiberia America
do
    Colate --mode mut \
        --mut ${mut} \
        --target_tmp SGDP_LBK_1x.colate.in \
        --reference_tmp SGDP_${pop}.colate.in \
        --bins 3,7,0.2 \
        --chr chr.txt \
        --target_age 7244 \
        --reference_age 0 \
        --num_bootstraps 1 \
        -o SGDP_LBK_1x_${pop}}
done
```

Exercise 4

Plot the coalescence rates of LBK against each of the other modern groups in SGDP. Which modern group is closest to this individual?

(If you like, you can use the script under Rscripts/.)

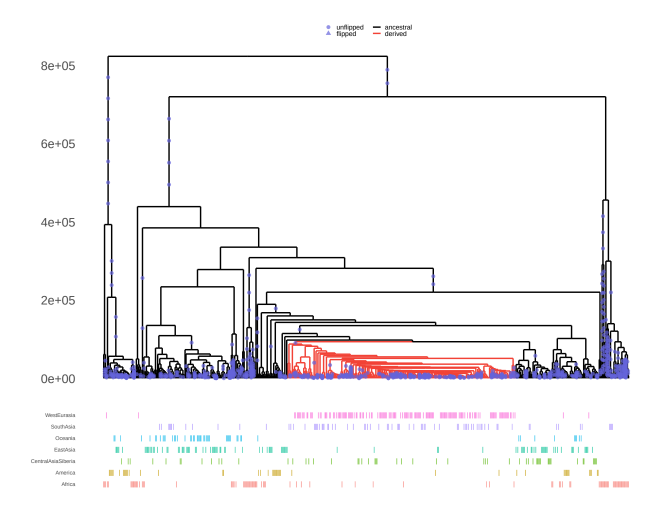


Figure 4: Marginal genealogical tree for rs1426654, a non-synonymous substitution associated with light skin pigmentation. The SNP is located in the SLC24A5 gene on chromosome 15 (BP=48,426,484). Derived allele carriers are shown in red.

4 Detecting evidence for positive selection

Positive natural selection on a derived allele is expected to lead to this allele spreading rapidly in a population, reflected in a burst of coalescence events. A well-known example is positive selection of a variant associated with lighter skin pigmentation in Europe and South Asia (Fig. 4). We have implemented a simple statistic that captures such events and measures the extend to which a mutation has out-competed other lineages.

To analyse selection, we will first apply functions provided with Relate to extract the relevant statistics, and then use the R package relater to analyse the output.

Note 8

We subset the sample into subgroups (e.g., WestEurasia) because the selection p-values assume a panmictic population and may get confounded by genetic structure.

In principle, the selection p-values are robust to misspecification of demographic histories, so we can use genealogies assuming a constant population size here, although we will use genealogies with jointly fitted population sizes and branch lengths below.

```
#assuming you are in worksheet_data
mkdir selection
cd selection
chr=15
lab="WestEurasia"
output="SGDP_${lab}_ne_chr${chr}"
# Copy anc/mut files for WestEurasia
cp ../precomp_results/WestEurasia/${output}* .
# Using these genealogies, calculate frequencies through time
# This will output a *.freq and *.lin file
${PATH_TO_RELATE}/bin/RelateSelection \
     --mode Frequency \
    -i ${output} \
     -o ${output}
# Next, calculate selection p-values.
# This will take the output of the previous step as input and output a *.sele file.
${PATH_TO_RELATE}/bin/RelateSelection \
     --mode Selection \
    -i ${output} \
     -o ${output}
# Also calculate a *.qual file storing statistics about the quality of trees
# (e.g., number of mutations mapping to the tree)
${PATH_TO_RELATE}/bin/RelateSelection \
     --mode Quality \
     -i ${output} \
     -o ${output}
```

Once we have generated these files, we use relater to analyse them in R.

```
library(relater)
output <- "./SGDP_WestEurasia_ne_chr15"

# parse files
mut <- read.mut(paste0(output, ".mut.gz"))
freq <- read.freq(paste0(output, ".freq"))
sele <- read.sele(paste0(output, ".sele"))
qual <- read.qual(paste0(output, ".qual"))

# combine these into a single data frame called allele_ages
allele_ages <- get.allele_ages(mut = mut, freq = freq, sele = sele)
# use the *qual file to filter out SNPs mapping to "bad" trees
allele_ages <- filter.allele_ages(allele_ages, qual)
allele_ages <- subset(allele_ages, !is.na(pvalue))

head(allele_ages[order(allele_ages$pvalue),])</pre>
```

Exercise 5

Let's analyse the allele_ages data frame. What is the top SNP with the lowest selection p-value?