EMBO Population Genomics Practical 1

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Introduction

For this practical, we will use admixtools v2, an R package that reimplements and expands the methods originally found in the classic ADMIXTOOLS program. The advantage of using admixtools v2 is that we can use R to issue commands and directly inspect and plot the outputs (a workflow in classic ADMIXTOOLS consisted a combination of sed/awk/shell scripting and manual editing of input files).

Combining and inspecting data to be passed to admixtools is also a challenge. These data often come in different formats, and might have different properties (e.g. modern vs ancient data). Data wrangling is often done in PLINK, but PLINK was not really designed for this type of work (it focuses on GWAS, not generic pop gen). In this practical, we will use tidypopgen, a new package developed to provide a clear grammar of population genetics, making data wrangling user friendly. tidypopgen has specific functions to prepare data for admixtools, and so the two blend together into a single R-centric workflow.

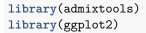
Using tidypopgen to wrangle data

tidypopgen follows the "tidyverse" logic and syntax. It is designed to make the processing and analysis of genetic data easy, reproducible and within a single programming language. The basic idea behind tidy data is that each observation should have its own row and each variable its own column, such that each value has its own cell. Applying this logic to population genetic data means that each individual should have its own row, with individual metadata (such as its population, sex, phenotype, etc) as the variables. Genotypes for each locus can also be thought of as variables, however, due to the large number of loci and the restricted values that each genotype can take, it would be very inefficient to store them as individual standard columns. They are stored in a file on disk, called a File Backed Matrix (FBM).

Hence, in tidypopgen, we represent data as a gen_tbl, a subclass of tibble which has two compulsory columns: id of the individual (as a character, which must be unique for each individual), and genotypes (stored in a compressed format as a File-Backed Matrix, with the vector in the tibble providing the appropriate link to those data).

library(tidypopgen)

```
## Loading required package: dplyr
##
## Attaching package: 'dplyr'
## The following objects are masked from 'package:stats':
##
## filter, lag
## The following objects are masked from 'package:base':
##
## intersect, setdiff, setequal, union
## Loading required package: tibble
```



Data types

This is additional reference information. It is not needed to run the practical, so you can skip it now, but it can be useful if you want to poke into the data files.

In this practical, we will use two types of data: PLINK binary BED files, and VCF (Variant Call Format) files. If you work with humans, you might also encounter PackedAncestry files. All these three formats can be read into a gen tibble in tidypopgen.

PLINK binary files are used to store genotype data in a compact and efficient binary format. They typically consist of three main files:

.bed file: Contains the binary genotype data. .bim file: Contains variant information. .fam file: Contains individual sample information.

The .bed file stores the genotype data in a compact binary format. It does not have a human-readable structure but is designed for efficient storage and access by PLINK and other compatible tools.

Genotype Encoding:

00: Homozygous for the reference allele (AA) 01: Missing genotype 10: Heterozygous (AB) 11: Homozygous for the alternate allele (BB)

The .bim file is a text file that contains variant information. Each row corresponds to a variant, and it has the following columns:

chrom: Chromosome number or ID variant ID: Unique identifier for the variant (e.g., rsID) genetic distance: Genetic distance (can be set to 0 if not available) base-pair position: Physical position of the variant on the chromosome allele 1: Reference allele (usually coded as the minor allele) allele 2: Alternate allele

The .fam file is a text file that contains information about each individual sample in the dataset. Each row corresponds to an individual and has the following columns:

Family ID: Identifier for the family (can be set to 0 if not applicable) Individual ID: Unique identifier for the individual Paternal ID: Identifier for the father (0 if not available) Maternal ID: Identifier for the mother (0 if not available) Sex: Sex of the individual (1 = male, 2 = female, 0 = unknown) Phenotype: Phenotype information (1 = unaffected, 2 = affected, -9 = missing)

VCF files are a standardized text file format widely used in bioinformatics for storing gene sequence variations. They consist of two main sections: the header and the data section. The header begins with ## and provides meta-information about the dataset, such as the version of the VCF file format, reference genome, and other key details. The header ends with a line starting with a single #, which specifies the column names for the data section.

The data section contains rows of variant calls, each representing a genetic variant. The columns, as defined in the header, typically include:

CHROM: Chromosome number or ID POS: Position of the variant on the chromosome ID: Identifier for the variant (e.g., dbSNP ID) REF: Reference allele ALT: Alternate allele(s) QUAL: Quality score of the variant call FILTER: Filter status indicating if the variant passed certain quality thresholds INFO: Additional information about the variant in a semi-colon-separated list of key-value pairs FORMAT: Format of the genotype fields in the subsequent columns sample1, sample2, ...: Genotype information for each sample, structured as per the FORMAT field

Loading the data

In this practical, we will use a panel of modern populations from the Human Origins genetics dataset, which is a collection of genetic data aimed at understanding human evolution, population structure, and migration patterns. The dataset contains genotypes from a wide array of modern human populations.

We will assume that you use a directory structure for your project where code sits in the code directory, and data sit in a data directory. So, from the code directory, we can look at the files in data with:

```
dir("./data")
```

```
## [1] "ancient_samples.vcf" "LBK_modern_pops.csv" "modern_samples.bed"
## [4] "modern_samples.bim" "modern_samples.fam"
```

We can see that there are three files with the prefix 'modern_samples', a .bed, a .bim and a .fam file. To convert them into a gen_tibble, we can simply use:

```
##
## gen_tibble saved to /home/andrea/git/f_stats_practical/data/modern_samples.gt
## using bigSNP file: /home/andrea/git/f_stats_practical/data/modern_samples.rds
## with backing file: /home/andrea/git/f_stats_practical/data/modern_samples.bk
## make sure that you do NOT delete those files!
## to reload the gen_tibble in another session, use:
## gt_load('/home/andrea/git/f_stats_practical/data/modern_samples.gt')
```

The message shows us where in our directories the gen_tibble(.gt) and its backing file (.bk) and R object file (.rds) are saved (these are used to store the genetic data for our gen_tibble). You only need to do this once. In the future, if you save your gen_tibble, you will be simply able to load it with gt_load().

Let's quickly inspect our data:

modern_gt

```
## # A gen_tibble: 588768 loci
## # A tibble:
                   413 x 3
##
      id
             population genotypes
##
      <chr> <chr>
                         <vctr_SNP>
##
   1 AD 006 AA
   2 AD_015 AA
                                  2
##
##
    3 AD 061 AA
                                  3
   4 AD_064 AA
                                  4
##
   5 AD 066 AA
                                  5
##
   6 AD 076 AA
                                  6
##
   7 AD 500 AA
                                  7
##
  8 AD_505 AA
                                  8
##
##
  9 AD_510 AA
                                  9
## 10 AD_511 AA
                                 10
## # i 403 more rows
```

We can see that we have >400 individuals and >500k markers. We can get a tally of how many individuals we have per population with:

```
modern_gt %>% group_by(population) %>% tally()
```

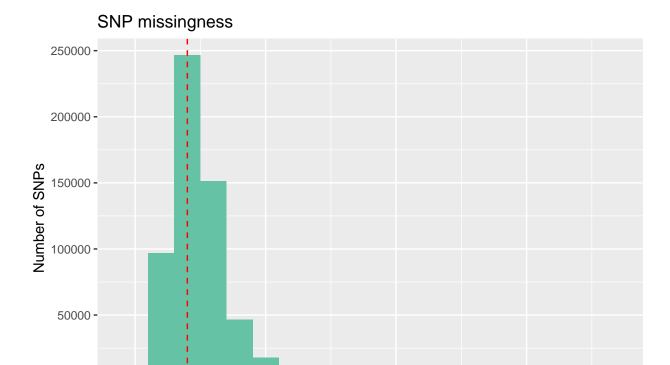
```
## # A tibble: 21 x 2
##
      population
      <chr>
                  <int>
##
##
    1 AA
                     12
##
    2 Basque
                     29
    3 Bedouin2
                     19
##
##
   4 Cypriot
                      8
                      7
##
   5 Dinka
##
   6 Druze
                     39
##
   7 French
                     25
   8 Han
##
                     33
## 9 Mayan
                     18
## 10 Mbuti
                     10
## # i 11 more rows
```

Note that we also have a chimp (pan_troglodytes), which we will use later on as an outgroup for certain analyses.

It is important to perform quality control (QC) on your data. Generally, we will be more stringent on the modern data, and allow more missingness on the ancient data. So, QC is often performed separately on different dataset depending on their nature. tidypopgen has two key functions to examine the quality of data, either across loci or across individuals. These functions are qc_report_loci and qc_report_indiv. We don't have space here to explore quality control in detail, but to exemplify, lets say we want to assess missingness across loci in our datasets. We can create a loci report and visualise our data. We will focus on missingness (the proportion of calls missing for each locus):

```
loci_report <- modern_gt %>%
  group_by(population) %>%
  qc_report_loci()

autoplot(loci_report, type = "missing")
```



There is a tail of a few loci with high missingness. Let's remove all sites that have missingness >= 4%. We can simply do that selecting the loci with missingness less than 0.04:

0.050

Proportion of missing data

0.075

0.025

0 -

0.000

```
modern_gt <- modern_gt %>%
select_loci_if(loci_missingness(genotypes)<0.04)</pre>
```

The select_loci_if() function can be used to filter data based on summary statistics such as missingness or minor allele frequency. In practice, this means that quality control of data can all be completed within R, without switching to command line software and rewriting files. QC is a very important process that we will not cover here, but make sure that you check the quality of your data BEFORE analysing them.

Then we can load our ancient data into a gen_tibble, this may take a moment, as we are reading from vcf format which is a large text file:

Ancient data can be low coverage, and often the are represented as pseudohaploids, i.e. sampling a random allele at each locus, and coding them as homozygotes:

```
ancient_gt <- gt_pseudohaploid(ancient_gt)</pre>
```

The ancient data contains ancient modern humans as well as Neanderthal and Denisovan samples:

```
ancient_gt$id
```

```
## [1] "AltaiNea" "Clovis" "Denisova" "GB20" "Otzi" "Kostenki" ## [7] "LBK" "Loschbour" "MA1" "UstIshim"
```

Having data in a tibble means that we can easily edit the metadata. Sample GB20 represents Mota, a \sim 4,000 year old individual from Ethiopia. Let's rename the ID of this sample to be more intuitive:

```
ancient_gt$id[ancient_gt$id == "GB20"] <- "Mota"</pre>
```

To use the data with admixtools, we need to assign a 'population' to all of our samples. For now, we can simply duplicate their individual id (so, each sample will be a population):

```
ancient_gt$population <- ancient_gt$id</pre>
```

Merging data

Once we are happy with the quality of our independent datasets, we can merge them to perform analyses.

Merging data from different sources is a common problem, especially in human population genetics where there is a wealth of SNP chips available. In tidypopgen, merging is enacted with an rbind operation between gen_tibbles.

If the datasets have the same loci, then the merge is trivial. If not, then it is necessary to subset to the same loci, and ensure that the data are coded with the same reference and alternate alleles (or swap them if needed).

Additionally, if data come from SNP chips, there is the added complication that the strand is not always consistent, so it might also be necessary to flip strand (in that case, ambiguous SNPs have to be filtered out). The rbind method for gen_tibbles has a number of parameters that allow us to control the behaviour of the merge.

To check our data compatibility before merging, we can run rbind_dry_run:

```
## harmonising loci between two datasets
## flip_strand = TRUE ; remove_ambiguous = TRUE
## ------
## dataset: reference
## number of SNPs: 581472 reduced to 581472
## ( 0 are ambiguous, of which 0 were removed)
## -------
## dataset: target
## number of SNPs: 588768 reduced to 581472
## ( 0 were flipped to match the reference set)
## ( 0 are ambiguous, of which 0 were removed)
```

The results show that merging these two datasets will cause a loss of around 7,000 SNPs from our 'target' dataset (in this case, the ancient samples as these are given as the second argument to rbind_dry_run, and are therefore the 'target').

Given the large overlap between our datasets, we can now merge using rbind (we need to give the name of the new backing file):

```
## harmonising loci between two datasets
## flip_strand = TRUE ; remove_ambiguous = TRUE
## -----
## dataset: reference
```

```
## number of SNPs: 581472 reduced to 581472
## ( 0 are ambiguous, of which 0 were removed)
## -------
## dataset: target
## number of SNPs: 588768 reduced to 581472
## ( 0 were flipped to match the reference set)
## ( 0 are ambiguous, of which 0 were removed)
##
## gen_tibble saved to /home/andrea/git/f_stats_practical/data/merged_samples.gt
## using bigSNP file: /home/andrea/git/f_stats_practical/data/merged_samples.rds
## with backing file: /home/andrea/git/f_stats_practical/data/merged_samples.bk
## make sure that you do NOT delete those files!
## to reload the gen_tibble in another session, use:
## gt_load('/home/andrea/git/f_stats_practical/data/merged_samples.gt')
```

f_2 statistics

Now that data are cleaned and merged, we can start our analysis. As all f statistics are combinations of pairs of f_2 statistics, we can begin by pre-computing f_2 's for all pairs of populations in our sample. First, we group our data by population, and then we can use $gt_extract_f2$ to save our f_2 pairs to disk into the outdir. This might take a minute or so.

```
## 202754 SNPs remain after filtering. 191339 are polymorphic.
```

```
\#\# i Allele frequency matrix for 202754 SNPs and 30 populations is 63 MB
```

i Computing pairwise f2 for all SNPs and population pairs requires 3796 MB RAM without splitting

i Computing without splitting since 3796 < 8000 (maxmem)...

Data written to ./data/f2_tidypopgen/

Now we can read read them in using admixtools:

```
f2_blocks = f2_from_precomp("./data/f2_tidypopgen")
```

```
## i Reading precomputed data for 30 populations...
```

```
## i Reading f2 data for pair 1 out of 465...i Reading f2 data for pair 2 out of 465...i Reading f2 dat
```

Ignore the warning about afprod = TRUE; it is a consequence of doing all possible f_2 in one go for multiple exercises. Usually, we would compute the f_2 for each project, but it would just take too long for this exercise.

Outgroup f_3

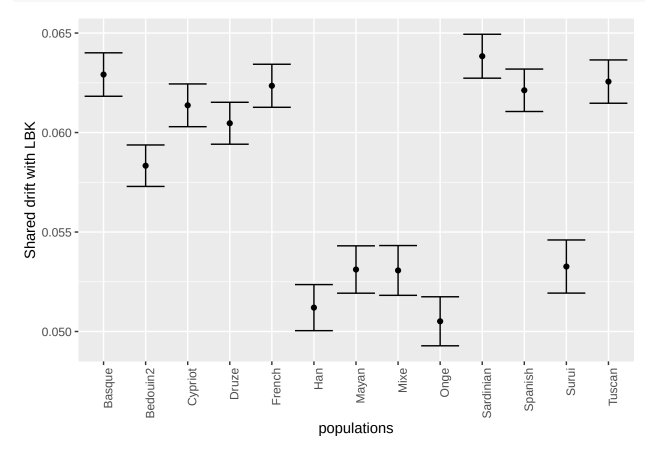
Imagine that we have a new mystery sample that we just sequenced, and we want to know what is the closest population in a panel of already characterised samples. For this exercise, we will use a Neolithic sample from the Linear Band Keramik culture, called LBK. We will use a set of contemporary populations from western Eurasia in our dataset:

```
lbk_modern_panel <- c("Basque", "Bedouin2", "Druze", "Cypriot", "Tuscan",</pre>
 "Sardinian", "French", "Spanish", "Onge", "Han", "Mayan", "Mixe", "Surui")
Now we compute the outgroup f_3, using Mbuti (a divergent African population) as an outgroup:
lbk_f3out <- f3(data = f2_blocks,</pre>
                pop1 = "Mbuti",
                pop2 = "LBK",
                pop3 = lbk_modern_panel)
1bk_f3out
## # A tibble: 13 x 7
##
      pop1 pop2
                  pop3
                                est
                                          se
                                                  Z
                                                        p
##
      <chr> <chr> <chr>
                              <dbl>
                                       <dbl> <dbl> <dbl>
##
   1 Mbuti LBK
                  Basque
                             0.0629 0.000546 115.
##
  2 Mbuti LBK
                  Bedouin2 0.0583 0.000521 112.
                                                        0
                  Cypriot
    3 Mbuti LBK
                             0.0614 0.000537 114.
                                                        0
                             0.0605 0.000527 115.
                                                        0
## 4 Mbuti LBK
                  Druze
## 5 Mbuti LBK
                  French
                             0.0624 0.000541 115.
## 6 Mbuti LBK
                             0.0512 0.000579
                                              88.4
                                                        0
                  Han
##
   7 Mbuti LBK
                  Mayan
                             0.0531 0.000595
                                              89.3
                                                        0
## 8 Mbuti LBK
                             0.0531 0.000626
                                                        0
                  Mixe
                                              84.8
## 9 Mbuti LBK
                             0.0505 0.000616
                                              82.0
                                                        0
                  Onge
## 10 Mbuti LBK
                  Sardinian 0.0638 0.000552 116.
                                                        0
## 11 Mbuti LBK
                  Spanish
                             0.0621 0.000533 117.
                                                        0
## 12 Mbuti LBK
                  Surui
                             0.0533 0.000668 79.8
                                                        0
## 13 Mbuti LBK
                  Tuscan
                             0.0626 0.000544 115.
                                                        0
Now, let's check which populations are closest to LBK (have the highest f_3)
lbk_f3out %>% arrange(desc(est))
## # A tibble: 13 x 7
##
      pop1 pop2 pop3
                                est
                                          se
                                                  z
                                                        p
##
      <chr> <chr> <chr>
                              <dbl>
                                       <dbl> <dbl> <dbl>
                  Sardinian 0.0638 0.000552 116.
##
   1 Mbuti LBK
                                                        0
  2 Mbuti LBK
                  Basque
                             0.0629 0.000546 115.
                                                        0
## 3 Mbuti LBK
                  Tuscan
                             0.0626 0.000544 115.
                                                        0
   4 Mbuti LBK
                  French
                             0.0624 0.000541 115.
##
                                                        0
## 5 Mbuti LBK
                  Spanish
                             0.0621 0.000533 117.
                                                        0
                             0.0614 0.000537 114.
## 6 Mbuti LBK
                  Cypriot
                                                        0
## 7 Mbuti LBK
                  Druze
                             0.0605 0.000527 115.
                                                        0
## 8 Mbuti LBK
                  Bedouin2 0.0583 0.000521 112.
                                                        0
## 9 Mbuti LBK
                             0.0533 0.000668
                                                        0
                  Surui
                                              79.8
## 10 Mbuti LBK
                  Mayan
                             0.0531 0.000595
                                              89.3
                                                        0
## 11 Mbuti LBK
                             0.0531 0.000626
                                                        0
                  Mixe
                                              84.8
## 12 Mbuti LBK
                             0.0512 0.000579
                                              88.4
                                                        0
                  Han
## 13 Mbuti LBK
                  Onge
                             0.0505 0.000616 82.0
QUESTION: Which populations share the most drift with LBK?
# The closest population is the Sardinian. Indeed they have been argued to be
# the modern population that is closest to Neolithic farmers based on ADMIXTURE
# clustering and PCA plots.
# Note that all these approaches, though, rely on
```

frequencies, so they are not completely independent lines of evidence.

You could produce a nice plot by using:

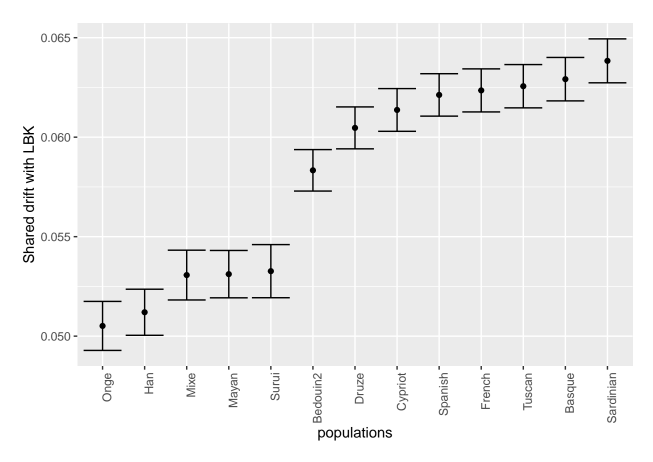
```
ggplot(lbk_f3out, aes(pop3, est)) +
  geom_point() +
  geom_errorbar(aes(ymin = est - 2 * se, ymax = est + 2 * se)) +
  labs(y = "Shared drift with LBK", x = "populations") +
  theme(axis.text.x = element_text(angle = 90, hjust = 1))
```



But it would be nicer if our populations were ordered by the level of shared drift. To do so, we need to reorder the levels of the population factor:

```
lbk_f3out$pop3<-factor(lbk_f3out$pop3, levels = lbk_f3out$pop3[order(lbk_f3out$est)])

ggplot(lbk_f3out, aes(pop3, est)) +
  geom_point() +
  geom_errorbar(aes(ymin = est - 2 * se, ymax = est + 2 * se)) +
  labs(y = "Shared drift with LBK", x = "populations") +
  theme(axis.text.x = element_text(angle = 90, hjust = 1))</pre>
```



QUESTION: Now we have another interesting sample from Central Asia that came into our lab, called 'MA1'. Investigate its relationship with contemporary populations in your sample.

```
##
  # A tibble: 13 x 7
##
      pop1
                   pop3
            pop2
                                 est
                                            se
                                                    z
                                                          p
##
                               <dbl>
                                         <dbl> <dbl>
                                                      <dbl>
      <chr> <chr>
                   <chr>
                              0.0618 0.000775
                                                79.7
                                                          0
##
    1 Mbuti MA1
                   Surui
                                                85.9
    2 Mbuti MA1
                              0.0617 0.000718
                                                          0
##
                   Mixe
    3 Mbuti MA1
                              0.0608 0.000685
                                                88.8
                                                          0
##
                   Mayan
    4 Mbuti MA1
                   French
                              0.0581 0.000591
                                                98.4
                                                          0
##
                                                          0
##
    5 Mbuti MA1
                   Basque
                              0.0579 0.000599
                                                96.7
                                                          0
##
    6 Mbuti MA1
                   Spanish
                              0.0569 0.000585
                                                97.3
    7 Mbuti MA1
                              0.0569 0.000600
                                                94.8
                                                          0
##
                   Tuscan
##
    8 Mbuti MA1
                   Sardinian 0.0561 0.000596
                                                94.1
                                                          0
##
    9 Mbuti MA1
                   Cypriot
                              0.0555 0.000596
                                                93.1
                                                          0
## 10 Mbuti MA1
                   Druze
                              0.0548 0.000569
                                                96.2
                                                          0
  11 Mbuti MA1
                   Han
                              0.0546 0.000617
                                                88.5
                                                          0
                              0.0539 0.000656
                                                82.3
                                                          0
  12 Mbuti MA1
                   Onge
                                                          0
## 13 Mbuti MA1
                   Bedouin2 0.0523 0.000557
                                                93.9
```

```
# So, this sample is actually closest to Native American populations. But note
# that our Asian representation in the panel is somewhat limited. However, MA1 is the
# Malt'a sample, which indeed has been argued to harbour ancestry that
# eventually made it to North America.
```

Admixture f_3

We now want to test whether a dataset of African Americans has detectable European ancestry (in other words, can they be modelled as a mixture of an African and a European population):

```
aa_f3admix <- f3(data = f2_blocks,</pre>
                  pop1 = "AA",
                  pop2 = "Yoruba",
                  pop3 = "French")
aa_f3admix
## # A tibble: 1 x 7
##
     pop1 pop2
                   pop3
                               est
                                          se
                                                 z
                                                        p
     <chr> <chr> <chr>
                             <dbl>
                                       <dbl> <dbl> <dbl>
           Yoruba French -0.00472 0.000120 -39.4
QUESTION: Do we have evidence for admixture?
```

Yes, we have a negative f3 value and z score that indicates a clear deviation # from 0 and suggests admixture.

Now, we want to test admixture in two East African target populations (Somali from Somalia, Dinka from Sudan). Source populations are Mota (\sim 4,000 year old individual from Ethiopia) and different modern and ancient (LBK) Eurasian populations:

```
eurasian_sources <- c("French", "Spanish", "Sardinian", "LBK")
```

We first test the Somali against all possible combinations of Mota and these Eurasian sources.

```
## # A tibble: 4 x 7
##
     pop1
            pop2
                      pop3
                                 est
                                           se
                                                             р
##
     <chr> <chr>
                      <chr>>
                               <dbl>
                                        <dbl> <dbl>
                      Mota -0.00456 0.000219 -20.8 2.27e- 96
## 1 Somali French
## 2 Somali LBK
                      Mota -0.00511 0.000353 -14.5 1.64e- 47
## 3 Somali Sardinian Mota -0.00495 0.000216 -22.9 4.24e-116
## 4 Somali Spanish
                      Mota -0.00450 0.000207 -21.7 7.86e-105
```

QUESTION: What can you conclude?

```
# We find significantly negative f3 statistics, which are evidence of admixture.
# The Somali population has a clear signal of admixture with Eurasian populations.
```

QUESTION: Now do the same for the Dinka. Do you get a similar result? If not, what can we conclude?

```
pop3 = "Mota")
dinka_f3admix
## # A tibble: 4 x 7
     pop1 pop2
                     pop3
                               est
                                          se
                                                           p
##
     <chr> <chr>
                     <chr>>
                              <dbl>
                                       <dbl> <dbl>
                                                       <dbl>
## 1 Dinka French
                     Mota
                           0.00606 0.000292
                                              20.8 1.09e- 95
## 2 Dinka LBK
                           0.00615 0.000400
                                             15.4 2.49e- 53
                     Mota
## 3 Dinka Sardinian Mota
                           0.00611 0.000294
                                              20.8 6.14e- 96
                           0.00613 0.000286
                                             21.4 5.63e-102
## 4 Dinka Spanish
                     Mota
# Here, we tested two East African populations (Somali and Dinka) considering
# several source populations from Europe and one ancient population from
# Ethiopia (Mota).
# For the Somali, we found significantly negative f3 statistics, which are
# evidence of admixture.
# For Dinka, we found significant f3 but they are POSITIVE. So, the significance
# is irrelevant, they key is that f3 is positive, so there is not evidence of
# admixture. But note that a positive f3 does not exclude
# admixture if the target population is heavily drifted.
```

f_4 or D statistics

A tibble: 1 x 8

pop1

Another way to look at admixture is to use the f_4 or D statistics. They do the same thing, and thus tend to give very similar results. D stats are easier to interpret in terms of magnitude (as they are standardised), but they can not be computed from f_2 . You can compute D stats if you start again from a file of genotypes (you would need to export a .bed file from the gen_tibble to be read directly in admixtools), but for this exercise we will use f_4 , as we already have all the f_2 .

For example, we might be interested in gene flow from a certain source, and we want to ask whether a population received it or not. The classical example is Neanderthal admixture into Eurasians. So, we want to ask whether Eurasian populations have more Neanderthal ancestry than Africans. We set up a test as:

```
neand_french_f4 <- f4(data = f2_blocks,</pre>
                       pop1 = "pan_troglodytes",
                       pop2 = "AltaiNea",
                      pop3 = "Mbuti",
                      pop4 = "French")
neand_french_f4
## # A tibble: 1 x 8
##
     pop1
                      pop2
                                pop3 pop4
                                                   est
                                                              se
                                                                      z
                                                                                  p
##
                       <chr>
                                <chr> <chr>
                                                 <dbl>
                                                           <dbl> <dbl>
                                                                             <dbl>
## 1 pan_troglodytes AltaiNea Mbuti French 0.00148 0.000328
                                                                 4.50 0.00000696
So, we can see that there is excess Neanderthal ancestry in the French. Now do the same for Han Chinese
(population 'Han')
neand_han_f4 <- f4(data = f2_blocks,</pre>
                    pop1 = "pan_troglodytes",
                    pop2 = "AltaiNea",
                    pop3 = "Mbuti",
                    pop4 = "Han")
neand_han_f4
```

est

z

p

se

pop3 pop4

pop2

QUESTION: Now we want to investigate the extend of Yamnaya ancestry that came into Europe during the Bronze age. If we assume that the LBK was the predominant genetic ancestry in Europe before the Yamnaya arrived, test:

- 1. that "French", "Basque" and "Spanish" all harbour Yamnaya ancestry
- 2. whether these populations differ in how much ancestry they share

Can you avoid writing down every comparison individually, and use vectors of populations to create some simple code to ask each question (look at what we did with the outgroup f_3).

```
target_eu <- c("French", "Basque", "Spanish")</pre>
# where did the Yamnaya get to?
f4(data = f2_blocks, pop1 = "Mbuti", pop2 = "Yamnaya",
  pop3 = "LBK", pop4 = target_eu)
## # A tibble: 3 x 8
##
    pop1 pop2
                   pop3 pop4
                                      est
                                                se
                                                       z
                                                                    p
##
     <chr> <chr>
                   <chr> <chr>
                                    <dbl>
                                             <dbl> <dbl>
                                                                <dbl>
## 1 Mbuti Yamnaya LBK
                        French 0.00202 0.000355
                                                   5.68 0.0000000136
## 2 Mbuti Yamnaya LBK
                        Basque 0.00182 0.000359
                                                    5.07 0.000000401
## 3 Mbuti Yamnaya LBK
                         Spanish 0.000757 0.000353 2.15 0.0317
# We see that the French and Basque have a clear Yamnaya signal, whilst we
# don't find one in the Spanish sample. Technically, the Spanish and LBK sample
# form a clade that is not broken by the Yamnaya, whilst this is not the case
# for French and Basque.
# do populations differ in their Yamnaya ancestry
f4(data = f2_blocks, pop1 = "Mbuti", pop2 = "Yamnaya",
  pop3 = target_eu, pop4 = target_eu)
## # A tibble: 9 x 8
##
    pop1 pop2
                   pop3
                           pop4
                                         est
                                                   se
     <chr> <chr>
                  <chr>
                           <chr>
                                       <dbl>
                                                <dbl>
                                                          <dbl>
                                                                    <dbl>
## 1 Mbuti Yamnaya French French
                                    2.32e-19 7.79e-17
                                                        0.00297 9.98e- 1
## 2 Mbuti Yamnaya French Basque -1.96e- 4 1.13e- 4 -1.73
                                                                8.29e- 2
## 3 Mbuti Yamnaya French Spanish -1.26e- 3 8.70e- 5 -14.5
                                                                1.21e-47
## 4 Mbuti Yamnaya Basque French
                                    1.96e- 4 1.13e- 4
                                                        1.73
                                                                8.29e- 2
## 5 Mbuti Yamnaya Basque Basque
                                  -2.07e-19 7.99e-17 -0.00259 9.98e- 1
## 6 Mbuti Yamnaya Basque Spanish -1.06e- 3 9.36e- 5 -11.4
                                                                6.13e-30
## 7 Mbuti Yamnaya Spanish French
                                    1.26e- 3 8.70e- 5 14.5
                                                                1.21e-47
## 8 Mbuti Yamnaya Spanish Basque
                                    1.06e- 3 9.36e- 5 11.4
                                                                6.13e-30
## 9 Mbuti Yamnaya Spanish Spanish -2.32e-19 7.85e-17 -0.00295 9.98e- 1
# Ignoring the comparisons of one population against itself, we see that
# French and Basque do not differ in their amount of Yamnaya ancestry, but they
# do differ from Spanish (in accordance to the test above).
```

f_4 ratio estimation

Let's go back to the question of European admixture into African Americans. We will use an f_4 ratio to estimate the proportion of European admixture. The order of populations is important. We have the sister taxon and the outgroup on the left, the target population in the centre, then the sister population, and the source of admixture to the right.

```
pops <- c("Han", "pan_troglodytes", "AA", "Yoruba", "French")</pre>
qpf4ratio(data = f2_blocks,
          pops = pops)
## # A tibble: 1 x 8
                                          pop5
##
     pop1 pop2
                            pop3 pop4
                                                 alpha
                                                 <dbl>
##
     <chr> <chr>
                            <chr> <chr>
                                          <chr>>
                                                          <dbl> <dbl>
           pan_troglodytes AA
                                   Yoruba French 0.158 0.00522
```

QUESTION: What is the proportion of European contribution into African Americans in the dataset?

```
# The alpha score of 0.158 and z-score of 30.36 indicates that 16% of African # American ancestry comes from a European source (and that amount is significant).
```

The order of our 'pops' is important. The population of interest, or target, is African American ('AA') and the population hypothesised to admix into the target are French. Here, we also use Yoruba as a proxy source population for African ancestry in African Americans (i.e. its ancestral source). You might want to draw a tree to make sure you are happy with who is going where.

- SISTER: refers to a sister population, one is related more closely to your hypothesised source of admixture than to the ancestral population of our target (in this case Han)
- OUTGROUP: refers to an outgroup to all other populations (in this case Pan troglodytes)
- TARGET: the admixed population under investigation (in this case, African American)
- ANCESTRAL: refers to the ancestral population that we know to be related to our admixed target (in this case, Yoruba)
- HYPOTHESISED SOURCE OF ADMIXTURE: refers to the population who is suggested to have admixed into the target (in this case, French)

QUESTION: Can you write some code to compute the f_4 ratio by hand (i.e. estimate the numerator and denominator f_4 , and compute the ratio by hand)?

```
#To illustrate the necessary order, the above f4 ratio can be written out as:
#numerator <- f4(f2_blocks, SISTER, OUTGROUP, TARGET, ANCESTRAL)$est
#denominator <- f4(f2_blocks, SISTER, OUTGROUP, HYPOTHESISED SOURCE OF ADMIXTURE, ANCESTRAL)$est
#AA_f4ratio_by_hand <- numerator/denominator

# F4 ratio
numerator <- f4(f2_blocks, "Han", "pan_troglodytes", "AA", "Yoruba")$est
denominator <- f4(f2_blocks, "Han", "pan_troglodytes", "French", "Yoruba")$est

AA_f4ratio_by_hand <- numerator/denominator

AA_f4ratio_by_hand</pre>
```

[1] 0.1580174

QUESTION: Repeat the analysis using other European populations (Sardinian, Spanish and Basque). Do the results change when using different European populations?

```
pops <- c("Han", "pan_troglodytes", "AA", "Yoruba", "Sardinian")</pre>
qpf4ratio(data = f2_blocks, pops = pops)
## # A tibble: 1 x 8
##
     pop1 pop2
                            pop3 pop4
                                         pop5
                                                    alpha
                                                               se
##
     <chr> <chr>
                            <chr> <chr> <chr>
                                                    <dbl>
                                                            <dbl> <dbl>
## 1 Han
           pan_troglodytes AA
                                  Yoruba Sardinian 0.162 0.00536 30.3
pops <- c("Han", "pan_troglodytes", "AA", "Yoruba", "Spanish")</pre>
qpf4ratio(data = f2_blocks, pops = pops)
## # A tibble: 1 x 8
##
     pop1 pop2
                           pop3 pop4
                                         pop5
                                                 alpha
                                                                    z
##
     <chr> <chr>
                            <chr> <chr> <chr>
                                                 <dbl>
                                                          <dbl> <dbl>
## 1 Han
           pan_troglodytes AA
                                  Yoruba Spanish 0.161 0.00531 30.3
pops <- c("Han", "pan_troglodytes", "AA", "Yoruba", "Basque")</pre>
qpf4ratio(data = f2_blocks, pops = pops)
## # A tibble: 1 x 8
    pop1 pop2
                           pop3 pop4
                                         pop5
                                                alpha
                            <chr> <chr> <chr> <chr> <dbl>
     <chr> <chr>
                                                         <dbl> <dbl>
                                  Yoruba Basque 0.159 0.00523 30.3
           pan_troglodytes AA
# Neither the ratio nor the z-score changes significantly with a different
# choice of European population. So, the result is robust with respect to
# our choice of admixture source.
```

QUESTION: A final little challenge. **qpf4ratio** can take a matrix of populations, where each line gives the 5 pops for an f_4 ratio. Can you write the code to run the 4 comparisons we just run as a single command?

```
## # A tibble: 4 x 8
##
    pop1 pop2
                         pop3 pop4
                                      pop5
                                                alpha
    <chr> <chr>
                         <chr> <chr> <chr>
                                                <dbl>
                                                       <dbl> <dbl>
                               Yoruba French
                                                0.158 0.00522 30.3
## 1 Han
          pan_troglodytes AA
## 2 Han pan_troglodytes AA
                               Yoruba Sardinian 0.162 0.00536
## 3 Han pan_troglodytes AA
                               Yoruba Spanish 0.161 0.00531 30.3
## 4 Han pan_troglodytes AA
                               Yoruba Basque
                                                0.159 0.00523 30.3
```

Combining outgroup f_3 with PCA

In case you have extra time, or simply do it at home.

We should always explore our data before running f statistics. In the practical above we skipped that step as it would have taken too long. Let's have a look at the outgroup f_3 analysis we did to place LBK, and complement it with a PCA

```
lbk_modern_panel <- c("Basque", "Bedouin2", "Druze", "Cypriot", "Tuscan",
    "Sardinian", "French", "Spanish", "Onge", "Han", "Mayan", "Mixe", "Surui")
modern_panel_gt <- merged_gt %>% filter (population %in% lbk_modern_panel)
# remove monomorphic sites (ungrouping the tibble, as we want to use global frequencies
# rather than population frequencies)
modern_panel_gt <- modern_panel_gt %>% ungroup() %>% select_loci_if(loci_maf(genotypes)>0)
```

PCA can not cope with missing data, so we will have to impute any missing data. Before doing so, we create a new backing file so that we can update it:

modern_panel_gt <- gt_update_backingfile(modern_panel_gt)</pre>

```
## Genetic distances are not sorted, setting them to zero
##
## gen_backing files updated, now
## using bigSNP file: /home/andrea/git/f_stats_practical/data/merged_samples_v2.rds
## with backing file: /home/andrea/git/f_stats_practical/data/merged_samples_v2.bk
## make sure that you do NOT delete those files!
# reset the ploidy for this tibble, as it is now all diploids
attr(modern_panel_gt$genotypes, "ploidy") <- 2
modern_panel_gt <- modern_panel_gt %>% gt_impute_simple(method="mode")

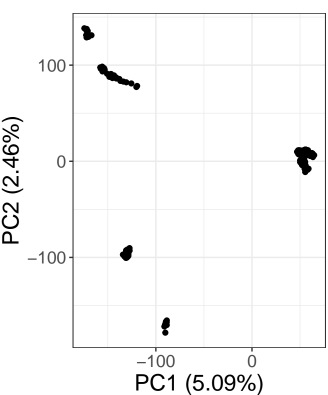
The final step is to filter for LD:
modern_panel_gt <- modern_panel_gt %>% select_loci_if(loci_ld_clump(genotypes))

And finally we fit the PCA:
modern_panel_gt <- modern_panel_gt %>% gt_pca_randomSVD()
```

We can now quickly plot the PCA:

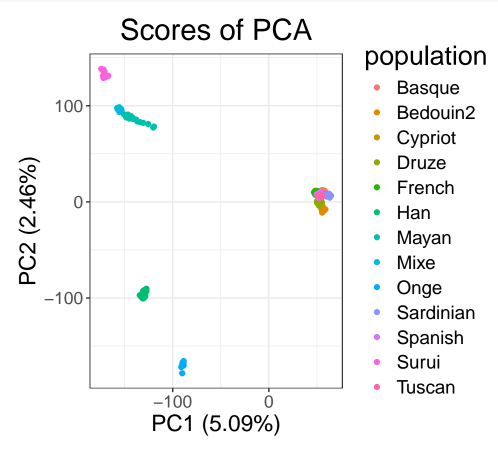
```
autoplot(modern_pca, type = "scores")
```

Scores of PCA



To colour our points, we can simply decorate the basic autoplot() output with ggplot2

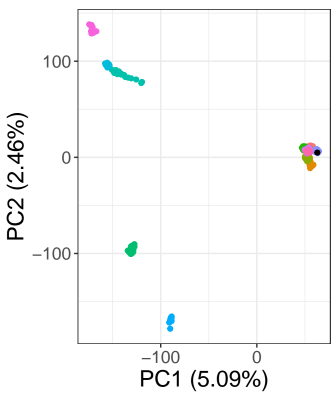
```
library(ggplot2)
autoplot(modern_pca, type = "scores") +
  aes(color = modern_panel_gt$population) +
  labs(color = "population")
```



We can now project the LBK onto the modern PCA:

geom_point(data=lbk_pca_scores, mapping=aes(x=.data\$.PC1, y=.data\$.PC2), col = "black")





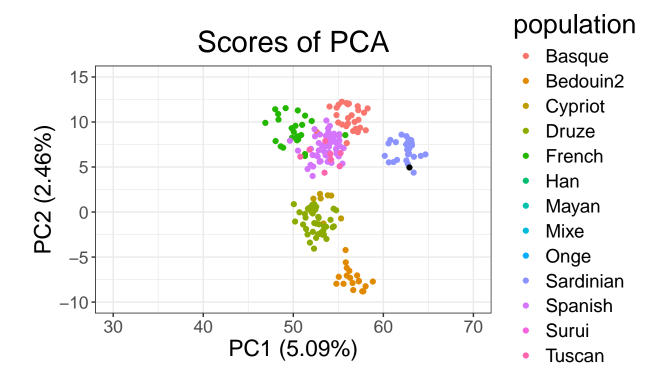
population

- Basque
- Bedouin2
- Cypriot
- Druze
- French
- Han
- Mayan
- Mixe
- Onge
- Sardinian
- Spanish
- Surui
- Tuscan

Let us zoom in onto Europe

```
autoplot(modern_pca, type = "scores") +
  aes(color = modern_panel_gt$population) +
  labs(color = "population") +
  geom_point(data=lbk_pca_scores, mapping=aes(x=.data$.PC1, y=.data$.PC2), col = "black") +
  lims(x=c(30, 70), y = c(-10, 15))
```

Warning: Removed 85 rows containing missing values or values outside the scale range
(`geom_point()`).



We can see that LBK falls squarely onto the Sardinians, confirming our f_3 analysis.

A few notes to make you think

In some tests, we combined modern and ancient populations. If you have a target of one type and sources of a different kind, then that strategy is generally fine. But if you mix things up, there is a risk of attraction between populations of the same type. For example, if you have a modern population and an ancient population and you test them against a modern population, then the modern populations will looks more similar to each other than they should. That artefact is mostly driven by damage in the ancient samples. A potential solution is to use transversions only, which are not affected by damage. We could have used tidypopgen at the beginning to filter out transitions.