

Analysis of Human Origins (plus Pacific) data

Alejandro Ochoa and John D. Storey

2019-05-29

Introduction

In this R markdown file we demonstrate the basic analysis of Human Origins dataset presented in our paper “New kinship and FST estimates reveal higher levels of differentiation in the global human population” by Ochoa and Storey. The analysis presented here relies exclusively on the data files provided in this repository (<https://github.com/StoreyLab/human-differentiation-manuscript>), which are the public subset of Human Origins and Pacific datasets.

This vignette takes about 8 minutes to compile on a single thread with a 3.6G processor and 31G of memory. The slowest steps are estimation of two kinship matrices (our new `popkin` method, and the existing Standard Kinship formula), and the two existing FST methods (Weir-Cockerham and Hudson K).

R package dependencies

To run this code, you will need the following packages:

```
library(BEDMatrix)    # to load genotype matrices
library(popkin)        # to estimate "population kinship" matrices
library(popkinsuppl)  # implements competing methods, available on OchoaLab GitHub
library(genio)         # to parse FAM file
library(readr)         # to read additional tables
library(tibble)        # to create tables
library(RColorBrewer)  # for colors
library(dplyr)         # for sanity checks
```

```
##
## 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
```

Estimate the kinship matrix

Load the genotype matrix (smartly, actually doesn't load into memory):

```
X <- BEDMatrix('data/human_origins_and_pacific_public', simple_names = TRUE)
```

```
## Extracting number of samples and rownames from human_origins_and_pacific_public.fam...
## Extracting number of variants and colnames from human_origins_and_pacific_public.bim...
```

This function from `genio` loads individual annotations (in plink FAM format), including subpopulations.

```
fam <- read_fam('data/human_origins_and_pacific_public')
```

```
## Reading: data/human_origins_and_pacific_public.fam
```

Now estimate the kinship matrix. We rely on the subpopulation labels to estimate the minimum kinship value to be treated as unrelated (zero). This step takes about 5 minutes.

```
kinship <- popkin(X, subpops = fam$fam)
```

Sub-subpopulation annotations

Before plotting, we need to reorder individuals so things look reasonable. To achieve this and other tasks, we load an additional table with subpopulation annotations.

```
info <- read_tsv(
  'data/human_origins_and_pacific_public_subpops.txt',
  col_types = 'ccddc'
)
# inspect
info
```

```
## # A tibble: 248 x 5
##   subsubpop      subpop      x      y country
##   <chr>         <chr>   <dbl> <dbl> <chr>
## 1 Ju_hoan_South SAfrica  20.7 -21.2 Botswana
## 2 Ju_hoan_North SAfrica  21.5 -18.9 Namibia
## 3 Taa_West      SAfrica  20.3 -23.6 Botswana
## 4 Taa_East      SAfrica  22.8 -24.2 Botswana
## 5 Taa_North     SAfrica  22.4 -23   Botswana
## 6 Naro         SAfrica  21.6 -22   Botswana
## 7 Gui          SAfrica  23.3 -21.5 Botswana
## 8 Hoan         SAfrica  23.4 -24   Botswana
## 9 Xuun         SAfrica  19.7 -18.7 Namibia
## 10 Gana        SAfrica  23.4 -21.7 Botswana
## # ... with 238 more rows
```

This table has one row per sub-subpopulation, for each of the 248 sub-subpopulations in the full dataset (including singleton and other sub-subpopulations removed in our publication). The table contains geographical coordinates (x and y) and country annotations, which we won't use here. These sub-subpopulations are grouped into 11 subpopulations (second column) that represent continental-level regions. We will use the `subsubpop` and `subpop` labels, and their order in this table, to sort individuals and create plots. The order and subpopulation assignments were defined manually, informed by geography and refined iteratively using the kinship plots to keep the most similar sub-subpopulations together.

Reorder individuals using sub-subpopulation annotations

Let's reorder individuals. First we ensure that every sub-subpopulation in the genotype table is present in the annotations table:

```
stopifnot(
  fam$fam %in% info$subsubpop
)
```

Now we reorder individuals so their sub-subpopulations appear in the desired order

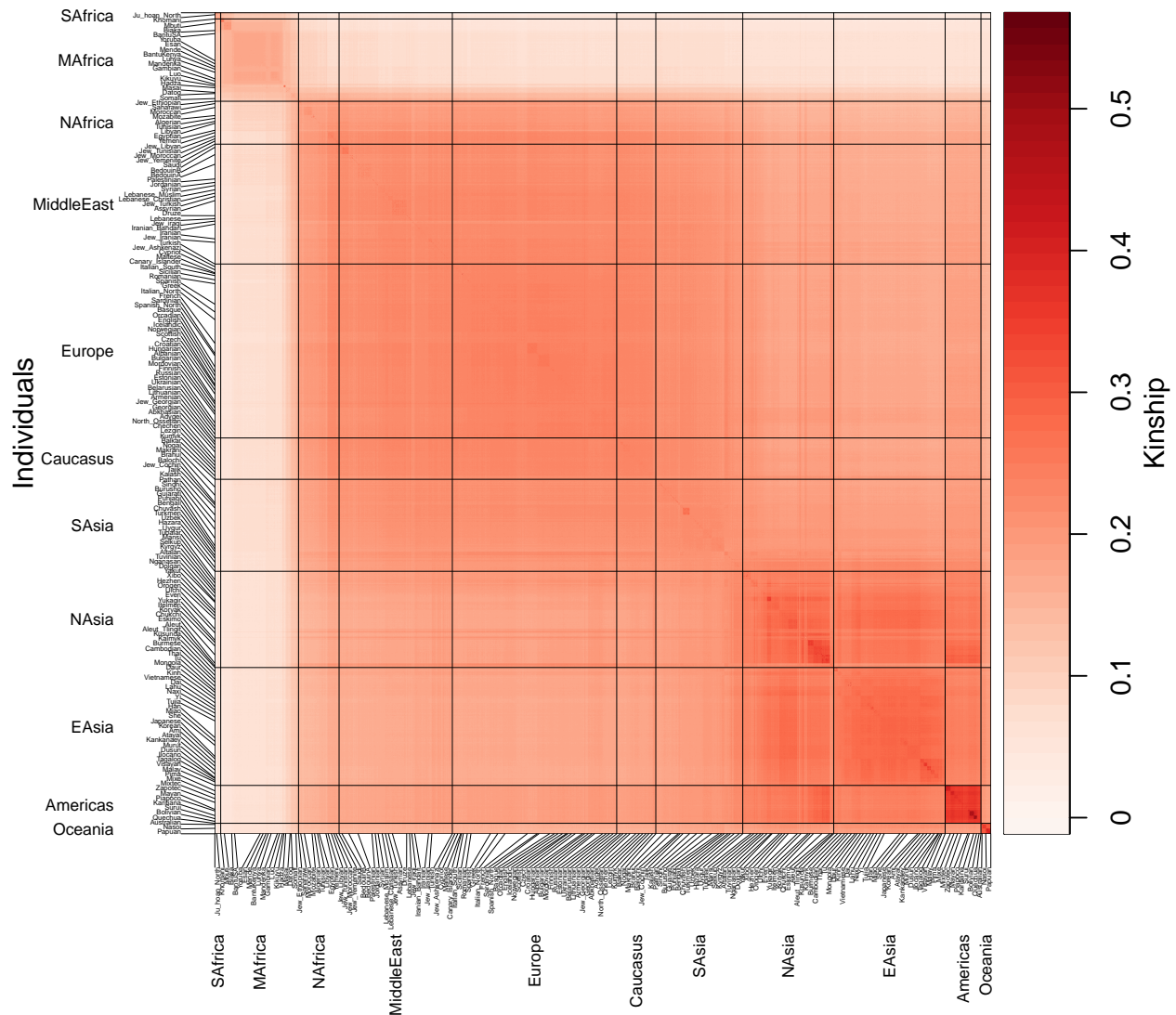
```
# the rank ties every individual in the same sub-subpopulation
fam_rank <- match(fam$fam, info$subsubpop)
# this order breaks ties by order of appearance
indexes <- order(fam_rank)
# before reordering, let's map subpopulations onto the FAM table
fam$subpop <- info$subpop[fam_rank]
# keep a copy of fam in the original order (used by FST estimators later)
# this copy does have the new 'subpop' column
fam_orig <- fam
# apply order to kinship matrix (both dimensions)
kinship <- kinship[indexes, indexes]
# and to individual annotations (rows only)
fam <- fam[indexes, ]
```

Kinship matrix plot

We are ready to visualize the estimated kinship matrix. One slight difference is that these values are not capped (in the publication they are capped to the top 1 percentile of the inbreeding value).

```
# line position for outer labels
line <- 3
# inner margins
par(mar = c(0, 0, 0, 0) + 0.2)
# outer margins
par(oma = c(line + 2.2, line + 3.2, 0, 3))

# main plot with all labeling bells and whistles
plot_popkin(
  # plot inbreeding along diagonal
  kinship = inbr_diag( kinship ),
  # leave a lot of space for labels
  ylab_line = line + 2,
  # two levels of labels
  labs = cbind(fam$fam, fam$subpop),
  labs_cex = c(0.25, 0.6),
  labs_las = 2,
  labs_line = c(1, line),
  labs_lwd = c(0.1, 0.5),
  labs_sep = c(FALSE, TRUE),
  labs_even = c(TRUE, FALSE)
)
```



Compared to the full dataset, this version limited to the public data has very few samples from **SAfrica** and **Oceania**.

Standard Kinship estimate

We use the `kinship_std` function from the `popkinsupl` package, which also works on `BEDMatrix` objects and controls memory just like `popkin`. The `mean_of_ratios = TRUE` option ensures that we use the most common formulation of this estimator.

```
kinship_standard <- kinship_std(X, mean_of_ratios = TRUE)
```

Next we reorder individuals to match the kinship matrix estimated earlier with `popkin` and ordered using an annotations table. Both kinship matrices have individual IDs stored in the row and column names, which we use for sorting.

```
# indexes to reorder individuals
indexes <- match(rownames(kinship), rownames(kinship_standard))
# apply reordering to both dimensions
kinship_standard <- kinship_standard[ indexes, indexes ]
```

```

# sanity check to make sure it was done correctly
stopifnot( rownames(kinship_standard) == rownames(kinship) )
# transform diagonal to have inbreeding values
# make a copy of the matrix, so we retain the original
kinship_standard_copy <- inbr_diag(kinship_standard)

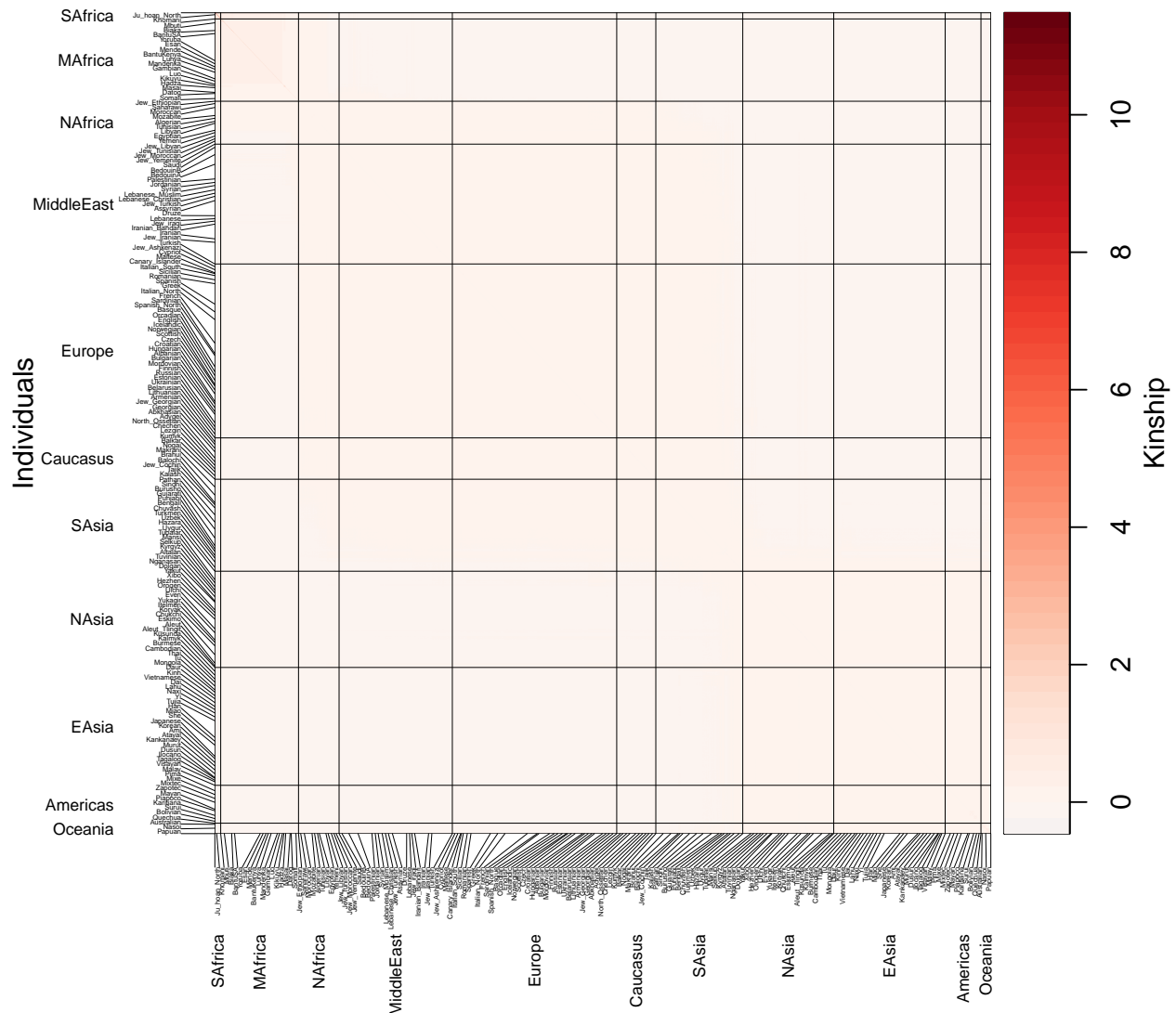
```

Now we plot this kinship matrix estimate.

```

line <- 3
par(mar = c(0, 0, 0, 0) + 0.2)
par(oma = c(line + 2.2, line + 3.2, 0, 3))
plot_popkin(
  kinship = kinship_standard_copy,
  ylab_line = line + 2,
  labs = cbind(fam$fam, fam$subpop),
  labs_cex = c(0.25, 0.6),
  labs_las = 2,
  labs_line = c(1, line),
  labs_lwd = c(0.1, 0.5),
  labs_sep = c(FALSE, TRUE),
  labs_even = c(TRUE, FALSE)
)

```



The biases along the diagonal are so extreme that they make everything else look too small. Note in the color key above that the most extreme values go higher than 11 (present in the top left corner of the kinship matrix heatmap)! Since the diagonal is the minority of the data, we cap using quantiles of the overall data.

```
# cap the estimate to this percentile to maximize dynamic range
alpha <- 0.01
# high cap
cap_hi <- quantile(kinship_standard_copy, probs = 1 - alpha)
# low cap
cap_lo <- quantile(kinship_standard_copy, probs = alpha)
# apply caps to the copy only
kinship_standard_copy[kinship_standard_copy > cap_hi] <- cap_hi
kinship_standard_copy[kinship_standard_copy < cap_lo] <- cap_lo
```

Now we plot the capped version:

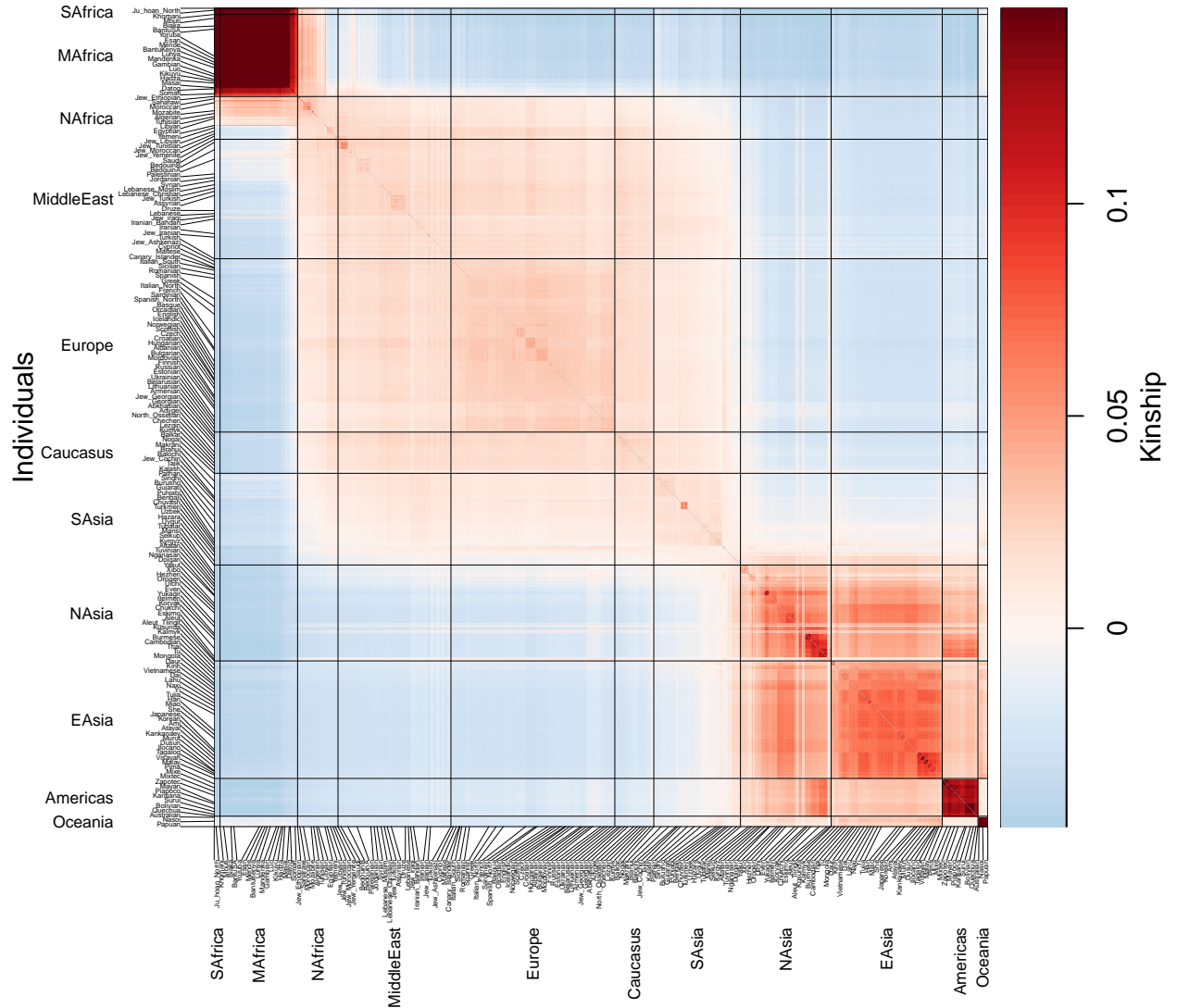
```
line <- 3
par(mar = c(0, 0, 0, 0) + 0.2)
par(oma = c(line + 2.2, line + 3.2, 0, 3))
plot_popkin(
  kinship = kinship_standard_copy,
```



```

ylab_line = line + 2,
labs = cbind(fam$fam, fam$subpop),
labs_cex = c(0.25, 0.6),
labs_las = 2,
labs_line = c(1, line),
labs_lwd = c(0.1, 0.5),
labs_sep = c(FALSE, TRUE),
labs_even = c(TRUE, FALSE)
)

```



The many negative values and distortions are clearly visible here. This estimate does not agree with the African Origins model of serial founder effects, which would assign Sub-Saharan Africans the lowest kinship, while this estimate assigns them the highest kinship values.

Weights to balance subpopulations and sub-subpopulations

This function calculates weights for individuals such that:

- every subpopulation has equal weight, and
- every sub-subpopulation has equal weight within its subpopulation.

```
# wrap code around function
get_weights_human_origins <- function(fam, info) {
  # this processing will be wrong if info contains more subpops than are present in fam,
  # so let's make them agree internally
  info <- info[ info$subsubpop %in% fam$fam, ]

  # get counts for sub-subpops and subpops
  # number of individuals in each sub-subpopulation
  subsubpop_to_counts <- table( fam$fam )
  # number of sub-subpopulations in each subpopulation
  subpop_to_counts <- table( info$subpop )
  # number of unique subpopulations
  K <- length( subpop_to_counts )

  # construct weights!
  weights <- 1 / (K * subsubpop_to_counts[fam$fam] * subpop_to_counts[fam$subpop] )
}

# apply function
weights <- get_weights_human_origins(fam, info)

# sanity checks

# do the weights sum to one? Yes!
sum( weights )
```

```
## [1] 1
```

```
# next steps are best performed on a tibble
weights_tibble <- tibble(
  weight = as.numeric( weights ),
  subpop = fam$subpop,
  subsubpop = fam$fam
)

# inspect
weights_tibble
```

```
## # A tibble: 2,124 x 3
##   weight subpop subsubpop
##   <dbl> <chr> <chr>
## 1 0.00909 SAfrica Ju_hoan_North
## 2 0.00909 SAfrica Ju_hoan_North
## 3 0.00909 SAfrica Ju_hoan_North
## 4 0.00909 SAfrica Ju_hoan_North
## 5 0.00909 SAfrica Ju_hoan_North
## 6 0.00413 SAfrica Khomani
## 7 0.00413 SAfrica Khomani
## 8 0.00413 SAfrica Khomani
## 9 0.00413 SAfrica Khomani
## 10 0.00413 SAfrica Khomani
## # ... with 2,114 more rows
```



```
# are subpopulations weighed equally? Also yes!
```

```
weights_tibble %>%  
  group_by( subpop ) %>%  
  summarize(sum = sum(weight))
```

```
## # A tibble: 11 x 2  
##   subpop      sum  
##   <chr>     <dbl>  
## 1 Americas 0.0909  
## 2 Caucasus 0.0909  
## 3 EAsia    0.0909  
## 4 Europe   0.0909  
## 5 MAfrica  0.0909  
## 6 MiddleEast 0.0909  
## 7 NAfrica  0.0909  
## 8 NAsia    0.0909  
## 9 Oceania  0.0909  
## 10 SAfrica 0.0909  
## 11 SAsia   0.0909
```

```
# lastly, the sub-subpopulations are weighed equally within their subpopulation.  
# We test that with 'Americas' only, for brevity:
```

```
weights_tibble %>%  
  filter( subpop == 'Americas' ) %>%  
  group_by( subsubpop ) %>%  
  summarize(sum = sum(weight))
```

```
## # A tibble: 10 x 2  
##   subsubpop      sum  
##   <chr>     <dbl>  
## 1 Bolivian 0.00909  
## 2 Karitiana 0.00909  
## 3 Mayan    0.00909  
## 4 Mixe     0.00909  
## 5 Mixtec   0.00909  
## 6 Piapoco  0.00909  
## 7 Pima     0.00909  
## 8 Quechua  0.00909  
## 9 Surui    0.00909  
## 10 Zapotec 0.00909
```

FST estimates

This is the FST estimate we obtain, which is made more robust by using the weights we previously calculated to balance subpopulations and sub-subpopulations:

```
Fst_popkin <- fst( kinship, weights )
```

```
# inspect  
Fst_popkin
```

```
## [1] 0.2622629
```

Nice! The value we estimated in the publication, using the full dataset, was 0.260.

Comparison to existing FST estimators

We use the following functions from the `popkinsuppl` package to estimate FST using the existing Weir-Cockerham estimator (Weir and Cockerham 1984) and “Hudson” estimator (Bhatia *et al.* 2013) generalized to more than two subpopulations in our work (Ochoa and Storey 2016). As subpopulation labels (required for these approaches), we use the 11 subpopulations (rather than the sub-subpopulations).

The genotype matrix `X` is in the original order for individuals, but the individual annotations table `fam` was reordered (along with the kinship matrix) using the annotations table `info`. Below we use `fam_orig` instead, which is the same as `fam` but in the original order.

```
# make sure that X and fam_orig agree
stopifnot(
  fam_orig$id == rownames(X)
)

# compute and store the values
Fst_wc <- fst_wc( X, labs = fam_orig$subpop )$fst
Fst_hudson_k <- fst_hudson_k( X, labs = fam_orig$subpop )$fst

# inspect them
Fst_wc

## [1] 0.07708545
Fst_hudson_k

## [1] 0.1108861
```

As reported in our publication, these values are unrealistically small because they treat the subpopulations as independent, which introduces downward biases in FST estimation. Our kinship estimates and other evidence conclusively show that these human subpopulations are not independent (otherwise the off-diagonal kinship values would all be zero).

Density of inbreeding coefficients

This code plots density curves of inbreeding coefficients for each subpopulation in a different color:

```
subpops <- unique( info$subpop )
# color palette for subpopulations
colors_subpops <- rev(brewer.pal(length(subpops), 'Spectral'))

# extract inbreeding vector
inbrs <- inbr( kinship )
# initialize y max, will grow below
y_max <- 0
# collect densities of interest.
# Complicated because samples are weighed to balance sub-subpopulations
# within each subpopulation
subpop_to_density <- list()
for (subpop in subpops) {
  # filter of data
  indexes <- fam$subpop == subpop
  # get subset weights too
  weights_subpop <- weights[indexes]
```

```

# renormalize when filters are done!
weights_subpop <- weights_subpop / sum(weights_subpop)
# estimate density using these weights
density_subpop <- density( inbrs[indexes], weights = weights_subpop )
# add to list of densities
subpop_to_density[[subpop]] <- density_subpop
# grow y max as needed
y_max <- max(y_max, density_subpop$y)
}

# shrink default margins
par( mar = c(4, 4, 0, 0) )

# initialize plotting region
plot(
  NA,
  xlim = range(0, inbrs),
  ylim = c(0, y_max),
  xlab = 'Inbreeding Coefficient',
  ylab = 'Density'
)

# add each subpopulation density
for (subpop in rev(subpops)) {
  # get color for this subpopulation
  color_subpop <- colors_subpops[match(subpop, subpops)]
  # plot density
  lines( subpop_to_density[[subpop]], col = color_subpop )
}

# add new Fst estimate (dashed line)
abline(v = Fst_popkin, lty = 2)

# legend
legend(
  'topright',
  subpops,
  title = 'Subpopulations',
  lty = 1,
  col = colors_subpops,
  cex = 0.7
)

```

