# A bioinformatics workflow for detecting signatures of selection in genomic data

Murray Cadzow $^1$ , James Boocock $^1$ , Hoang Tan Nguyen $^{1,2}$ , Phillip Wilcox $^{1,3}$ , Tony R $_1$  Merriman $^1$  and Michael A $_2$  Black $^1$ 

<sup>1</sup>Department of Biochemistry, University of Otago <sup>2</sup>Department of Mathematics and Statistics, University of Otago <sup>3</sup>Scion Research, Rotorua, New Zealand

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## Contents

1	Inti	roduction	4
2	Get	eting Started	4
	2.1	Prerequisites	4
	2.2	Python dependencies	5
	2.3	Downloading	5
	2.4	Installation	5
	2.5	Genetic Maps and Impute Haplotypes	5
	2.6	Ancestral Fasta Files	6
3	Tut	orial	6
	3.1	Selection Signatures at the Lactase Locus	6
		3.1.1 Getting the Data	6
	3.2	Setting up the Pipeline Run	7
	3.3	Population Files	7
	3.4	Run The Tutorial	7
	3.5	Data Visualisation	8
		3.5.1 Visualizing $F_{ST}$ values	8
		3.5.2 Fay and Wu's H	8
		3.5.3 iHS	13
		3.5.4 Tajima's D	16
		3.5.5. Reb	18

4	Out	out Files	19
	4.1	multi_population	19
		4.1.1 Fst	19
	4.2	selection_pipeline	19
		4.2.1 Fay and Wu's H	19
		4.2.2 iHS	19
		4.2.3 iHH	19
		4.2.4 Tajima's D	19
5	Con	mand line Arguments	20
	5.1		20
			20
		•	20
			20
			20
	5.2		21
			21
		5.2.2 Output Files	21
			21
			21
	5.3		22
		5.3.1 Input Files	23
		5.3.2 Output Files	23
		5.3.3 Other parameters	23
	5.4	Configuration File	23
		5.4.1 system	24
		5.4.2 environment	24
		5.4.3 selection_pipeline	24
		5.4.4 vcf_tools	24
		5.4.5 shapeit	24
		5.4.6 impute2	25
		5.4.7 plink	25
		5.4.8 Rscript	25
		5.4.9 haps_scripts	26
		5.4.10 ancestral_allele	26
		5.4.11 qctool	26
		5.4.12 multicore_ihh	26
6	Log	Files	27
	6.1		27
	6.2	<del>-</del>	27

7	Extra Features	27
	7.1 Galaxy Intergration	27
8	$\mathbf{F.A.Q}$	27

## 1 Introduction

This selection analysis workflow utilizes genotype data derived from next-generation sequencing (NGS) or highdenisity microarray (e.g., "SNP chip") experiments to identify the presence of signatures of selection. The tools used to detect selection are dependent on the selection signature being investigated (Sabeti et al., 2006). The pipeline presented here generates various output files containing within- and between-population selection signatures. The starting point for the analysis is a variant call format (VCF) file of the genotype data and populations of interest (Danecek et al., 2011). Both  $F_{ST}$  and Tajima's D can be calculated from standard genotype data (Weir and Cockerham, 1984; Tajima, 1989). To compute iHS, Rsb and Fay and Wu's H requires haplotype information, and thus the genotype data must be phased prior to calculation of these statistics (Voight et al., 2006; Gautier and Vitalis, 2012; Fay and Wu, 2000). For phasing, shapeit2 is used, and for imputation impute2 is used (Howie et al., 2009; Delaneau et al., 2013). Furthermore these statistics also require ancestral allele information (Flicek et al., 2012). The pipeline performs phasing if the VCF files do not contain phase information, and then performs ancestral allele annotation. [\*\*\* JAMES - IS THIS ONLY TRUE FOR HUMAN DATA? \*\*\*] Once complete, the rehh package for R provides a simple interface for implementing EHH-based analyses (Gautier and Vitalis, 2012). Here we have extended rehh to include penalties for gaps that match those used in the original iHS paper (Voight et al., 2006). rehh is used to calculate iHH, iHS, iES and Rsb. To calculate Fay and Wu's H, a C program, variscan, was utilised (Vilella et al., 2005). The pipeline is implemented in Python, and takes a VCF file as input. The output is a collection of files relating to selection signatures dected by the various software tools.

# 2 Getting Started

## 2.1 Prerequisites

The selection pipeline was developed on a 64-bit Ubuntu 13.04 system and has been tested on 64-bit Centos and Ubuntu 13.10 installations. The pipeline should work on any 64-bit linux derivative assuming some basic libraries and tools are installed on the system. 8GB of RAM should be sufficient for all computation steps (imputation is the most RAM-intensive component of the pipeline).

- Python > = 2.6
- Bourne-again Shell (Bash)
- Perl5
- R >= 3.0.0 (with the ability to install packages into a library within your home directory structure)
- GNU Autotools
- GCC
- Git

The software is installed with the same permissions as the user than runs the script: if the user is not root then a local (i.e., user-writeable) R library is required. The program also installs the scripts to the user's /.local/bin

directory. This directory should be added to the system PATH to give direct access to the programs from the command-line.

## 2.2 Python dependencies

The following Python packages are required for the pipeline:

- python-setuptools
- python-numpy
- python-scipy

if using python < 2.7 the package argparse will need to be installed.

It can be installed using the following command.

```
easy\_install-2.6 argparse
```

Most linux distributions provide these packages through the official package management repositories.

## 2.3 Downloading

The selection pipeline can be obtained at the url: https://github.com/smilefreak/selectionTools

To download run the following command in a terminal.

```
git clone https://github.com/smilefreak/selectionTools
```

## 2.4 Installation

To perform an automatic installation of the selection analysis pipeline, run the following command in the root of the directory in which the pipeline was installed (i.e., within the "selectionTools" directory).

```
./install.sh
```

The installation process creates a default configuration file located in the base directory of the pipeline. It also adds a program called selection\_pipeline to the system path. To test that the program is installed correctly, run the following command at a terminal prompt.

```
selection_pipeline -h
```

If the above command does not work, make sure that /.local/bin/ is included in the PATH environment variable.

## 2.5 Genetic Maps and Impute Haplotypes

To use the phasing and imputation features of the pipeline requires both genetic map files and haplotype files. For humans, files that conform to the format required for shapeit and impute2 can be found here. For impute2, one reference is available here. Download and extract the archive to referencefiles/impute\_ref and uncompress the contents. For shapeit2, a genetic map can be found here. Download and extract the archive to reference-files/shapeit ref.

To use other reference files with the selection pipeline requires setting the following options in the config file. The question mark character "?" in the config is substituted by the chromosome number: this is used for reference files that are split on chromosomes.

```
genetic_map_prefix=genetic_map_chr?_combined_b37.txt
...
impute_map_prefix=genetic_map_chr?_combined_b37.txt
impute_reference_prefix=ALL_1000G_phase1integrated_v3_chr?_impute
...
```

If you decide to store the reference files in another location, further options require alteration in the config file:

```
genetic_map_dir= ${HOME}/MerrimanSelectionPipeline/referencefiles/shapeit_ref
...
impute_map_dir= ${HOME}/MerrimanSelectionPipeline/referencefiles/impute_ref
impute_reference_dir= \${HOME}/MerrimanSelectionPipeline/referencefiles/impute_ref
...
```

#### 2.6 Ancestral Fasta Files

The generation of results for iHS requires assigning the ancestral allele. The selection pipeline uses the ancestral alleles from the 6-way EPO (Enredo-Pecan-Ortheus) alignment pipeline. The files can be downloaded from here. Be sure to extract the contents of the archive after download. The default directory to store the ancestral reference files is

```
referencefiles/ancestral_ref/
```

If you downloaded your reference to a different location you can alter the following setting in your config file.

```
ancestral_fasta_dir = # directory you downloaded alignment to #
...
```

## 3 Tutorial

## 3.1 Selection Signatures at the Lactase Locus

## 3.1.1 Getting the Data

In humans, lactase is encoded by the LCT gene, which is located on Chromosome 2 at the coordinates: 136,545,410-136,594,750. For this example we will use a 10 megabase region containing the LCT, and genotype data from the CEU and YRI populations from the 1000 Genomes Project. In order to demonstrate the functionality of the pipeline we will use the chromosome 2 region 130,000,000-140,000,000. The lactase gene is an example of

strong selection in the last 5,000-10,000 years in human populations, specifically those of European ancestory (Bersaglieri *et al.*, 2004). The tutorial commands will be given as examples using phasing and imputation or taking advantage of the fact that the 1000 genomes VCF files actually contain phasing information. If you choose to run the tutorial without using phasing and imputation you will only need to download the Ancestral allele information. To download the example dataset enter the following command:

```
git clone https://github.com/smilefreak/SelectionPipelineTestData
```

Navigate to the created SelectionPipelineTestData folder and extract selection pipeline tutorial.tar.gz.

```
cd SelectionPipelineTestData
tar xzf selection_pipeline_tutorial.tar.gz
```

## 3.2 Setting up the Pipeline Run

## 3.3 Population Files

Population files are required for any cross population comparisions. The commands below will initiate the data generation step. Population files are line separated files, where the first line contains the population name, and every successive line contains an individual ID from that population.

```
<POPULATION_IDENTIFIER>
<INDIVIDUAL ID 1>
<INDIVIDUAL ID 2>
......
<INDIVIDUAL ID N>
```

### 3.4 Run The Tutorial

The default configuration file is located in the base directory of the selection pipeline. To run the pipeline with phasing and imputation, execute the command below in the folder in which the example data were extracted, and change the –config-file parameter to match the location where the pipeline is installed.

```
multipop_selection_pipeline -p CEU_ids.txt -p YRI_ids.txt \
   -i CEU_YRI_lactase.vcf --config-file defaults.cfg \
   -a "--imputation"
```

As the 1000 genomes datasets are actually phased VCF to run the pipeline without phasing run the following command. The –phased-vcf option significantly speeds up the selection pipeline.

```
multipop_selection_pipeline -p CEU_id.txt -p YRI_ids.txt \
-i CEU_YRI_lactase.vcf --config-file defaults.cfg \
--a "--phased-vcf"
```

The generated folders and current folder have all the data required to perform further selection analysis. Within each population folder four output files are generated. These contain Tajima's D, iHH, an updated VCF, and Fay and Wu's H statistic. The files are located in the results folder inside each population subfolder.  $F_{ST}$  is

calculated between each population and results are located in the fst folder.  $F_{ST}$  results are calculated using the Weir and Cockerham estimator.

#### 3.5 Data Visualisation

The purpose of the analysis pipeline is to generate standard signatures of selection from a VCF formatted input file. In order to assist with exploring/interpretting the results, visualization of the pipeline output can be extrenely useful. The next section describes some basic approaches to plotting these data using the R programming language. All of the following commands are run in an R session with the working directory set as the base directory from which the tutorial is being run. In each of the plots that follow, the vertical blue lines indicate the position of the lactase gene (LCT).

## 3.5.1 Visualizing $F_{ST}$ values

Read in the  $F_{ST}$  data:

```
CEUYRIfst=read.table("fst/2CEUYRI.fst", header=TRUE)
```

Plot the  $F_{ST}$  results across the entire 10 megabase region for the tutorial. Mean  $F_{ST}$  is being plotted in each case.

Plot mean Weir FST values.

```
weirFST = read.table('fst/2CEUYRI.weir.fst',header=T)
plot(weirFST[,6]~ weirFST[,2], pch=16, cex=.4,
type="p",ylab=expression(F[ST]),xlab="Chromosome position (bp)")
abline(h=mean(weirFST[,6]) + 3 * sd (weirFST[,6]),col='red')
rect(136545410,-1900,136594750,100,border="Blue")
```

Plot mean HapMap FST values.

```
hapmapFST = read.table('fst/2CEUYRI.hapmap.fst',header=T)
plot(hapmapFST[,6]~ hapmapFST[,2], pch=16, cex=.4,
type="p",ylab=expression(F[ST]),xlab="Chromosome position (bp)")
abline(h=mean(hapmapFST[,6]) + 3 * sd (hapmapFST[,6]),col='red')
rect(136545410,-1900,136594750,100,border="Blue")
```

Figures (1 and 2) show the weir and hapmap FST values for the tutorial region.

## 3.5.2 Fay and Wu's H

Plot the Fay and Wu's H values for the CEU population:

```
CEUFay=read.table('CEU/results/CEU2.faw',comment.char="#")

#Plot Fay and Wu's H

plot(CEUFay[,15] ~ CEUFay[,1],xlim=c(136545410-1e6,136594750+1e6),

pch='.',cex=2,ylim=c(-50,0),

xlab='Chromosome position (bp)',ylab="H Statistic")

rect(136545410,-1000,136594750,100,border="Blue")
```

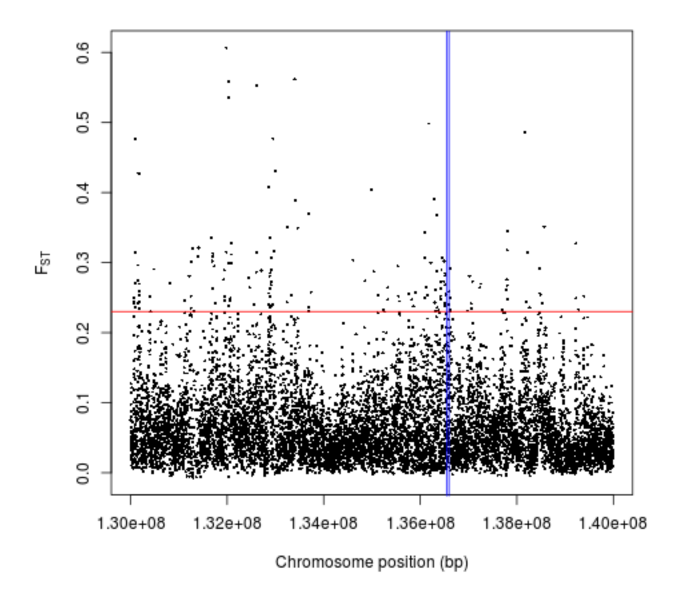


Figure 1: Values of Weir  ${\cal F}_{ST}$  mean FST between the CEU and YRI populations

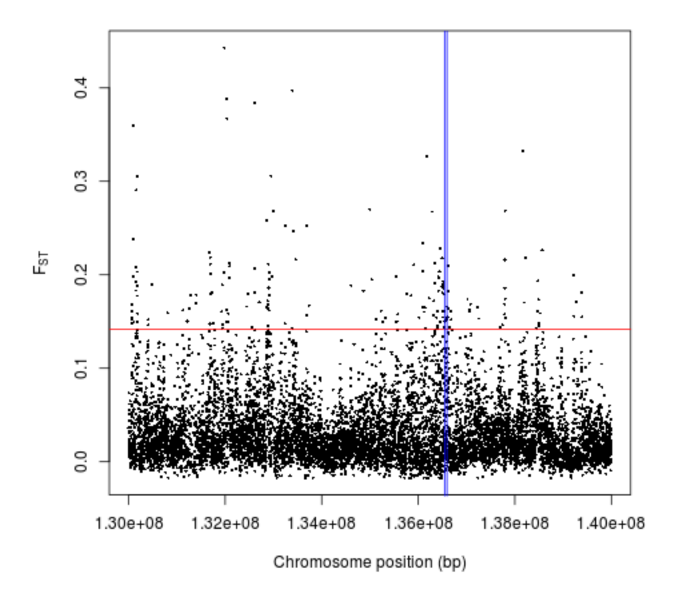


Figure 2: Values of HapMap  ${\cal F}_{ST}$  mean FST between the CEU and YRI populations

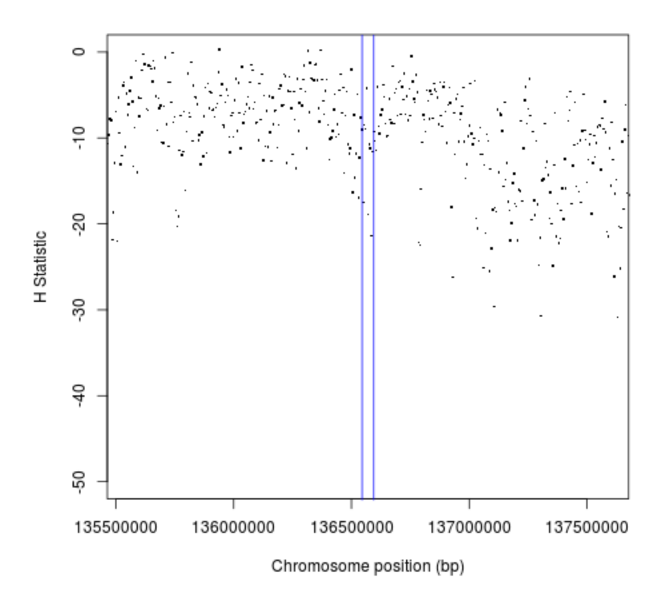


Figure 3: Fay and Wu's H statistic in the CEU population, across a two megabase region around the LCT gene.

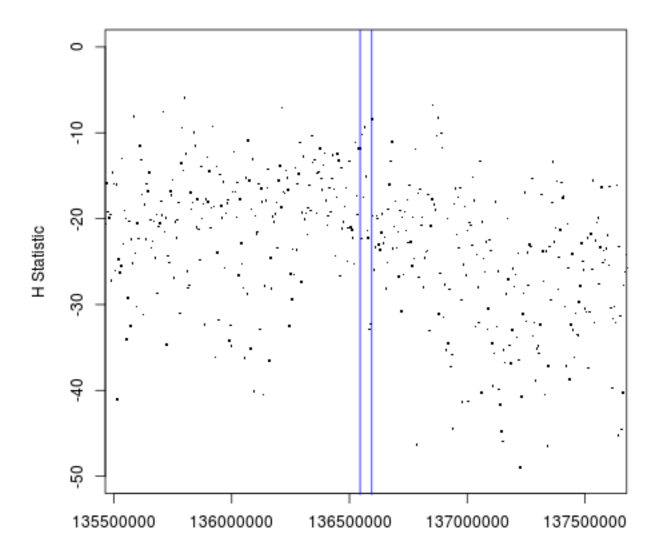


Figure 4: Fay and Wu's H statistic in the YRI population, across a two megabase region around the LCT gene.

Figure 3 shows the Fay and Wu's H statistic one megabase downstream and upstream of the lactase gene. Plot the Fay and Wu's H values for the YRI population:

```
YRIFay=read.table('YRI/results/YRI2.faw',comment.char="#")

#Plot Fay and Wu's H

plot(YRIFay[,15] ~ YRIFay[,1],xlim=c(136545410-1e6,136594750+1e6),pch='.',cex=2,ylim=c(-50,0),
xlab='',ylab="H Statistic")

rect(136545410,-1900,136594750,100,border="Blue")
```

Figure 4 shows Fay and Wu's H statistic one megabase downstream and upstream of the lactase gene.

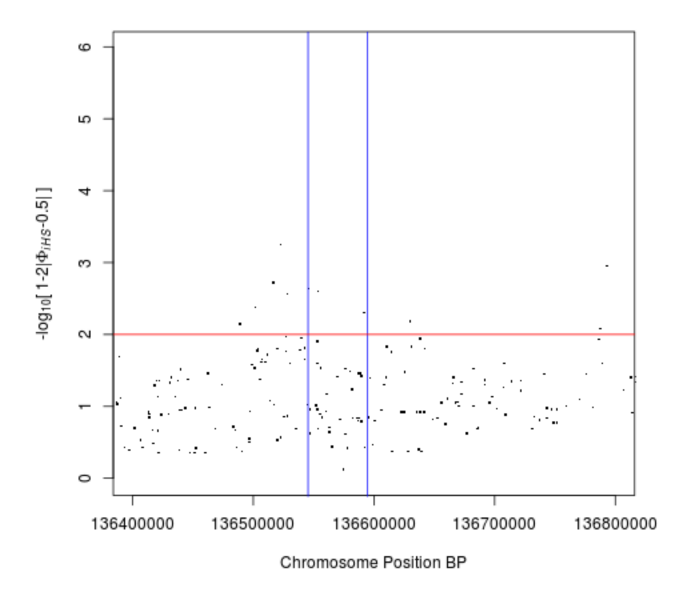


Figure 5: iHS statistic in the CEU population, across a 400 kilobase region around the LCT gene.

#### 3.5.3 iHS

Plot the iHS values around the lactase gene for the CEU population:

```
CEUihs = read.table('CEU/results/CEUchr2.ihs')

#plot IHS pvalues

plot(CEUihs[,4] ~ CEUihs[,2],xlim=c(1.364e8,1.368e8),pch='.',cex=2,

ylab=expression("-" * log[10] * "[" ~ "1-2|" * Phi[scriptstyle(italic(iHS))] * "-0.5|" ~ "]"),

xlab="Chromosome Position BP")

rect(136545410,-10,136594750,10,border="Blue")

abline(h=2,col="red")
```

Figure 5 shows iHS pvalues around the lactase gene in the CEU population.

Plot the iHS values around the lactase gene for the YRI population:

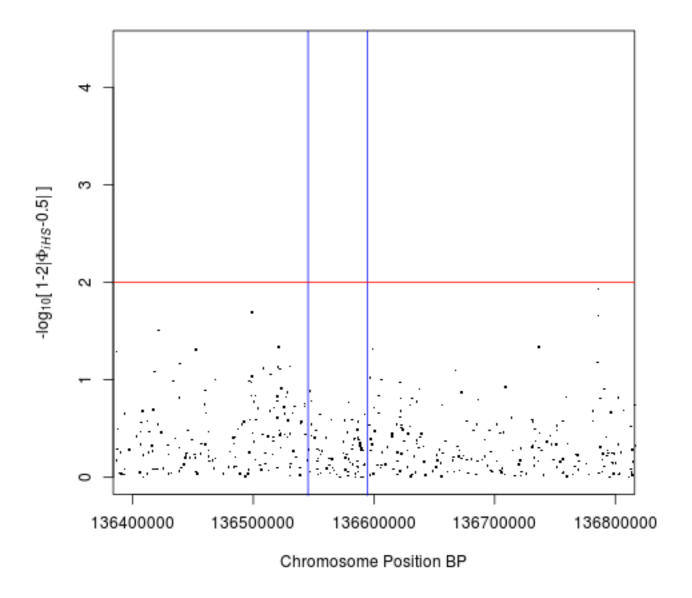


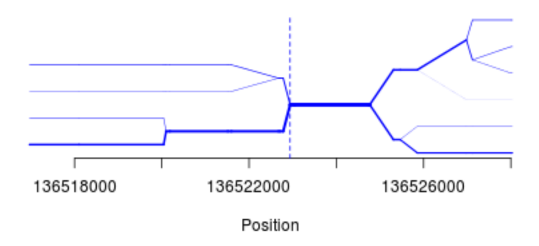
Figure 6: iHS statistic in the YRI population, across a 400 kilobase region around the LCT gene.

```
YRIihs = read.table('YRI/results/YRIchr2.ihs')
#plot IHS pvalues
plot(YRIihs[,4] ~ YRIihs[,2],xlim=c(1.364e8,1.368e8),pch='.',cex=2,
ylab=expression("-" * log[10] * "[" ~ "1-2|" * Phi[scriptstyle(italic(iHS))] * "-0.5|" ~ "]"),
xlab="Chromosome Position BP")
rect(136545410,-10,136594750,10,border="Blue")
abline(h=2,col="red")
```

Figure 6 shows iHS pvalues around the lactase.

To visualize individual SNPs, a haplotype bifurication diagram can be used (Gautier and Vitalis, 2012). The SNP rs28453840 displayed a strong signal of selection using iHS in the CEU population. The following commands construct a bifurcation diagram for both alleles at this SNP using rehh.

# rs28453840 (Ancestral Allele)



# rs28453840 (Derived Allele)

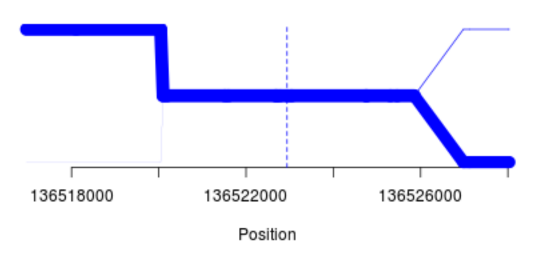


Figure 7: Bifurcation Diagram for rs28453840

```
# Plot bifurcation diagram.
load('CEU/results/CEUchr2.RData')
par(mfrow=c(2,1))
bifurcation.diagram(haplohh=d,mrk_foc=13750,all_foc=1)
bifurcation.diagram(haplohh=d,mrk_foc=13750,all_foc=2)
```

Figure 7 shows the bifurcation plot for rs28453840 a snp that had a iHS score greater that 3 at position 136522941 on chromosome 2 just upstream of lactase.

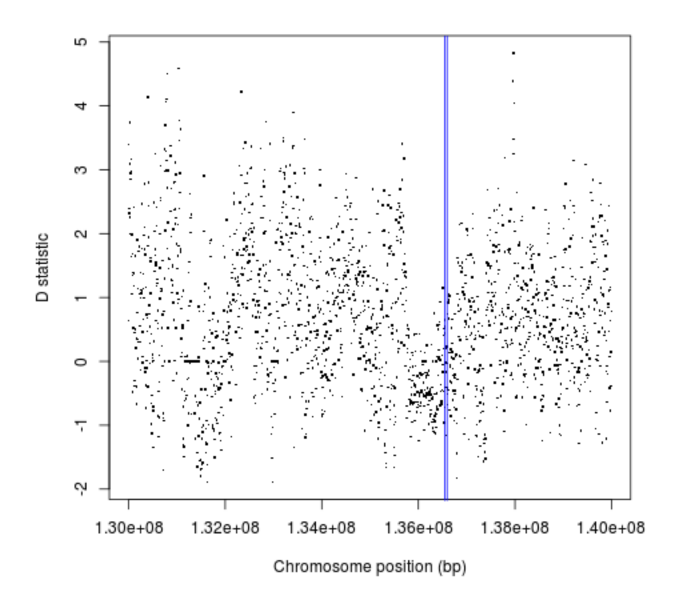


Figure 8: Tajima's D statistic in the CEU population, across the full 10 megabase pipeline region.

## 3.5.4 Tajima's D

```
tajimaD=read.table(file="CEU/results/CEU2.taj_d", header=TRUE)

plot(tajimaD[,4] ~ tajimaD[,2],pch='.',cex=2,xlab="Chromosome position (bp)", ylab="D statistic")

rect(136545410,-10,136594750,10,border="Blue")

Figure 8 show the Tajima's D statistic across the full tutorial region.

tajimaD=read.table(file="YRI/results/YRI2.taj_d", header=TRUE)

plot(tajimaD[,4] ~ tajimaD[,2],pch='.',cex=2,xlab="Chromosome position (bp)", ylab="D statistic")
```

Figure 9 show the Tajima's D statistic across the full tutorial region.

rect(136545410,-10,136594750,10,border="Blue")

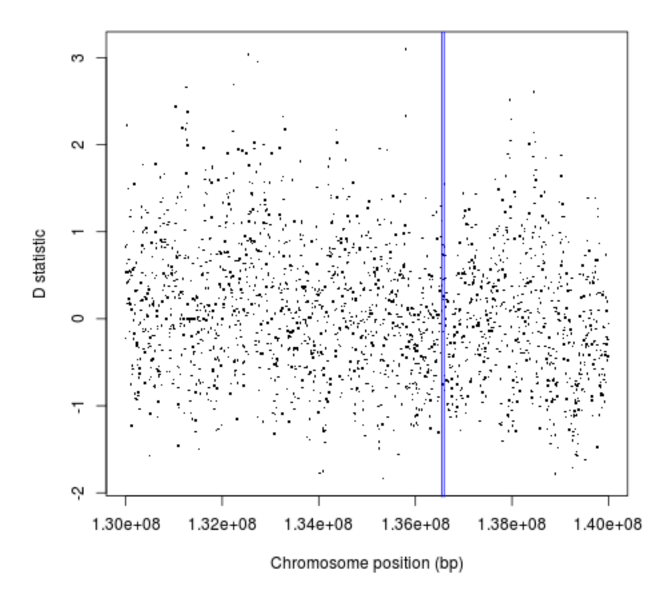


Figure 9: Tajima's D statistic in the YRI population for the tutorial region

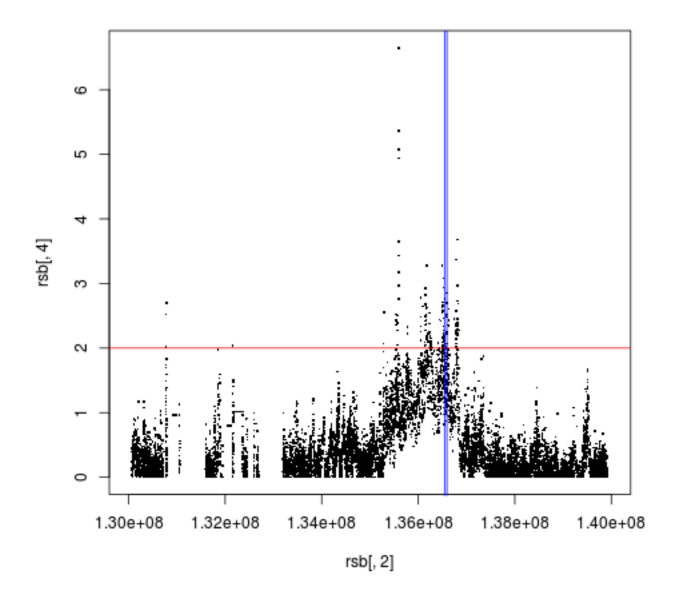


Figure 10: RSB statistic between the CEU and YRI population for the tutorial region

## 3.5.5 Rsb

Rsb makes the differences between the two populations very clear. The following code shows how to plot the Rsb bilateral P-values for the 10 megabase tutorial region.

Figure 10 shows the Rsb statistic for the entire region for the tutorial, it is visually striking the difference at the LCT locus.

# 4 Output Files

The output files are preserved in the same state as the original output from the program used to generate the data.

## 4.1 multi population

#### 4.1.1 Fst

Located in the fst folder. Tab-delimited data file containing 1 header line followed data on each subsequent line

CHROM	BIN_START	BIN_END N_VARIA	NTS	WEIGHTED_FST	MEAN_FST
2	13000001	130010000	94	0.133102	0.0680276

## 4.2 selection pipeline

All the outputs for each population are contained in the results folder. If you ran the tool using multi\_population the outputs are located in <pop name>/results.

#### 4.2.1 Fay and Wu's H

Space-delimited data file containing header line that start with a hash character (#). Contains lots of columns. If you are only interested in Fay and Wu's H, column 1 provides the position and column 15 provides the H statistic.

```
# RefStart Refend ... FayWu_H
130000040 130005039 ... -22.2438460
```

#### 4.2.2 iHS

Space-delimited data file containing one header line followed by data on each subsequent line.

```
"CHR" "POSITION" "iHS" "Pvalue"
"rs4662641" 2 130000272 0.0644902912148128 0.0229261107107533
```

#### 4.2.3 iHH

Space-delimited data file containing one header line followed by data on each subsequent line.

```
"CHR" "POSITION" "FREQ_a" "IHHa" "IHHd" "IES"
"rs1251176" 2 130000040 0.9823 11558.89 83915.49 11571.13
```

#### 4.2.4 Tajima's D

Space-delimited data file containing one header line followed by data on each subsequent line.

CHROM	BIN_START	N_SNPS	TajimaD
2	130000000	22	0.775224

# 5 Command line Arguments

The selection pipeline contains three programs: selection\_pipeline, aa\_annotate and multi\_population. The selection pipeline does all the intra-population statistics calculations. The multi\_population program calculates all the inter-population statistics and calls the selection pipeline. The aa\_annotate program annotates a haplotype file or a phased vcf file with the ancestral allele from the 6-way EPO alignment, for other species or alternative ancestral annotation, the feauture will be added in the future.

## 5.1 Multipopulation

#### 5.1.1 Input Files

• -i <vcf input file> VCF file containing all the populations you want to analyse from one chromosome or a part of a chromosome only.

#### 5.1.2 Output Files

• FST

Fst results are stored in the fst folder with the chromosome number followed by the two populations. e.g 2CEUYRI.fst

• Selection Pipeline Results

All single population pipeline results are stored in the subdirectory of the population in a folder named results. These contain the iHH, Tajima's D and a population VCF file.

#### 5.1.3 Other parameters (Compulsory)

• -l <log file>

Name for the log file. Moved into the logs folder at the end of program run.

• -c <Chromosome>

Integer for the chromosome being used.

• -a <Arguments to the selection pipeline>

Quoted string containing any extra arguments to the selection pipeline program. e.g "-imputation"

• -config-file < path to config file>

Path to the selection pipeline config file an example config file is located in the base directory of the extracted package.

## 5.1.4 Other parameters (Optional)

• -no-clean-up

Do not clean up intermediate data files.

• -fst-window-size <FST window size>

Fst calculation sliding window size in kilobases (default = 1kb).

-fst-window-step <FST window step>
 Fst calculation windoow jump in kilobases, if window size is equal to the jump size non-overlapping windows are used (default = 1kb).

• -cores

Number of cores avaliable for the pipeline overrides setting in the config file.

## 5.2 Selection Pipeline

#### 5.2.1 Input Files

 $\bullet$  -i <VCF input file>

Single population single chromosome VCF input file. VCf should be unzipped

## 5.2.2 Output Files

The Results directory contains all the output files.

• .ihh file

The outputted iHH data for each SNP

• .taj d file

Tajima's D output

• .vcf file

Single population VCF updated by the pipeline, can contain.

## 5.2.3 Other parameters(Compulsory)

• -config-file < Config File path>

Path to the selection pipeline config file an example config file is located in the base directory of the extracted package.

#### 5.2.4 Other parameters(Optional)

• -l <log file>

Name for the log file. Moved into the logs folder at the end of program run.

• -maf <minimum MAF>

Minor allele frequency filter threshold any SNPs below this threshold will be discarded from the analysis (default = 0.01).

• -hwe <hardy-weinberg minimum p-value>

A hardy weinberg test is performed on every snp any snps failing the test will be discarded (default = 0.001).

• -remove-missing <Inclusion threshold for missing genotypes>

Inclusion criteria for SNPs with missing data. SNPs with less than this value will be removed from analysis (default = 0.99).

• -fay-Window-Width <window width>

Sliding window width for Fay and Wu's H calculation in kb (default = 5kb).

• -fay-Window-Jump < Window Jump>

Window jump for Fay and Wu's H calculation, if equal to fay-Window-Width non-overlapping windows will be used (default = 5kb).

• -TajimaD <tajimas D bin size>

Tajima's D statistic bin size in kb (default = 5kb).

• -no-clean-up

Do not clean up intermediate data files

 $\bullet$  -ehh-window-size

Window size for multicore rehh calculations in megabases (Mb) (default = 5Mb).

• -ehh-overlap

Window overlap for multicore rehh calculations in megabases (Mb) (defaut = 2Mb).

• -daf <Minimum derived allele frequency>

Derived allele frequencies below this minimum will be discarded (default = 0.0).

• -big-gap

Gap size in kb for not calculating iHH if the gap is too large. If set to zero the big-gap rule is not applied (default = 0kb).

 $\bullet$  -small-gap

Gap size in kb for applying a penalty to the area calculated by iHH. If set to zero the small-gap rule is not applied (default = 0kb).

 $\bullet$  -small-gap-penalty

Penalty multiplier for intergration step in iHH calculation.  $multiplier/gap\_size*area$  is the formula we use. Setting the multiplier to the same value as the small gap threshold is recommended (default = 0kb).

• -cores

Number of cores avaliable for the pipeline overrides setting in the config file.

## 5.3 Ancestral Annotation

The program ancestral\_annotation is installed in the program path. The program annotates haps and .vcf files with ancestral allele annotation from the 6-way IPO alignment or the human reference genome.

#### 5.3.1 Input Files

- -i or -haps <HAPS File>Haplotype File (.haps)
- -v <Phased VCF file>

Phased VCF file (.vcf), phased VCF genotypes denoted by a bar (  $\mid$  ) for each sample.

• -a or -aa <Ancestral allele fasta>

Ancestral allele annotation file. Currently only works on a the full 1000 Genomes Project GRCh3764 reference file or the single chromosome fasta files from the 6-way EPO alignment.

#### 5.3.2 Output Files

• -o or -output <Output file name>

Output file name optional argument by default output is sent to the stdout stream.

• -s or -sample-file <Sample file output>

Sample file output name ( currently only works with phased vcf option)

#### 5.3.3 Other parameters

• -header-regex

Compulsory argument: it is a regex (regular expression) with a question mark denoting substitution for the chromosome number. The regex should match the header in the fasta file and when the question mark is replaced selectively return only the chromosome of interest. Optional for single chromosome fasta files.

 $\bullet$  -single-chromosome

Single chromosome pipeline run option.

• -c <chromosome number>

The number or symbol of the chromosome being used.

ullet -f or -format <format>

The 6-way EPO alignment denotes ancestral alleles with both high and low confidence. To use only ancestral alleles with high confidence use –format upper. To use both high and low confident alleles use –format lower. By default the program will use only highly confident alleles. The highly confident alleles are in uppercase.

By default it uses all the valid bases in the file that are one of either A, T, C, G, a, t, c or g

## 5.4 Configuration File

The selection pipeline requires a configuration file. By default the program looks in the current working directory for a file named defaults.cfg but you can point the program to any file using command line argument –config-file <config file location>. There are two main programs in the selection pipeline namely selection pipeline and

multi\_population. These programs share a config file but certain configuration parameters can be ommitted when using the selection\_pipeline program exclusively. A clean install of the pipeline generates an example configuration file containing default arguments for all the compulsory parameters. The default config file contains an example of the format.

## **5.4.1** system

• cores avaliable

Certain programs in the pipeline can take advantage of multicore computers. This option instructs the pipeline about the maximum number of concurrent processes it is allowed to use.

#### 5.4.2 environment

## • LD LIBRARY PATH

Set the library path when running the pipeline, this enables the pipeline to use the shared libraries that are used for some programs in the pipeline (alter this option with caution!)

#### • PERL5LIB

Sets the PERL5LIB environment variable, this enables the pipeline to use the perl libraries required by VCFTOOLS (alter this option with caution!)

## 5.4.3 selection pipeline

• selection pipeline executable

Points to the location of the selection pipeline executable.

## 5.4.4 vcf tools

• vcf tools executable

Points to the vcftools executable, by default it points to the vcftools executable installed with the pipeline.

 $\bullet \ vcf\_subset\_executable \\$ 

Points to the vcf-subset executable, by default pointing to the vcf-subset installed with the pipeline.

• vcf merge executable

Points to the vcf-merge executable, by default pointing to the vcf-subset installed with the pipeline.

• extra args

A quoted string containing extra arguments to send to the vcf\_tools executable.

#### 5.4.5 shapeit

• shapeit executable

Location of the shapeit executable.

• genetic map dir

Directory containing the genetic map for shapeit.

• genetic map prefix

The full file for the genetic map files with a "?" character representing the changing chromosome number.

• extra\_args extra arguments to send to shapeit (Warning: Certain options could potentially break the pipeline - use with caution).

## 5.4.6 impute2

• impute executable

Location of the impute2 executable

• impute map dir

Directory containing the genetic map for impute2

• impute reference dir

Directory containing the reference panel ( .legend and .hap) files for impute2.

• chromosome split size

Window size for imputation calculation.

• impute map prefix

The full file name for the genetic map files with a "?" character representing the changing chromosome number

• impute reference prefix

The full file name for the reference panels minus the extension with a "?" character representing the changing chromosome number.

• extra args

extra arguments to send to impute (Warning: Certain options could potentially break to pipeline use with caution).

#### 5.4.7 plink

• plink\_executable

Location of the plink executable

#### 5.4.8 Rscript

• rscript executable

Location of the rscript executable (Program usually in path so just Rscript is the default).

 $\bullet \ \ indel\_filter$ 

Location of the rscript indel filter (hap indel and maf filter.R)i

## 5.4.9 haps scripts

- haps\_to\_hapmap\_script
   Location of the haps to hapmap script
- haps\_filter\_scriptLocation of the haps\_filter script

## 5.4.10 ancestral allele

• split by chromosome

Option determines whether the ancestral fasta file is split by chromosome or not. If the ancestral fasta is split by chromosome the ancestral\_fasta\_dir and ancestral\_prefix arguments are required. If the ancestral fasta is just one flat file the ancestral\_fasta\_file options is required which is just merely the location of the ancestral fasta option.

• ancestral fasta header regex

This options specifies a regular expression that will extract each chromosome from the fasta file. The '?' denotes the chromosome number and will be replaced in the regular expression for the chromosome number of interest. Even when using one fasta per chromosome this is required as the python library used to extract the sequence uses the fasta header for extraction. Only required if your fasta file is not split by chromosome.

• ancestral\_fasta\_file

Ancestral fasta file used if the split by chromosome is set to false.

• ancestral\_allele\_script

Location of the ancestral annotation script (ancestral annotation)

ancestral\_fasta\_dir
 Directory containing the ancestral reference files

ancestral\_prefix
 Full file name for ancestral fasta files containing a "?" character

#### 5.4.11 qctool

qctool\_executable
 Location of the qctool executable.

#### 5.4.12 multicore ihh

multicore\_ihh
 Location of the multicore iHH.R script

# 6 Log Files

The log files contain all the information for a pipeline run including any errors if for some reason some part of the pipeline does not complete.

## 6.1 multi population

The location of the log file for *multi\_population* defaults is located in the log directory. It contains all the logging information for the between population selection signature calculations.

## 6.2 selection pipeline

The location of the log file for *selection\_pipeline* is located in the log directory. The log file contains all the logging information for the within population selection signature calculations.

## 7 Extra Features

## 7.1 Galaxy Intergration

The galaxy folder contains the scripts required to add the selection pipeline to a local galaxy installation. The pipeline is also avaliable on the galaxy toolshed at galaxy url. To do intergrate the pipeline into galaxy.

[\*\*\* JAMES - is this stuff available for Galaxy? \*\*\*]

# 8 F.A.Q

1. How do I run *multi\_population* with a phased VCF?

In the -a argument for multi\_population merely add -phased-vcf between the quotes. This will ensure phasing and imputation will be skipped when *selection\_pipeline* is called.

2. My populations are in seperate VCF-Files: how do I run multi population?

To run the pipeline you will need to merge the VCF-files into one large multipopulation VCF file and generate the appropriate population files.

To merge your vcfs you can use the vcf-merge program. For this to work correctly outside the selection pipeline you will need to add the following to your .bashrc file or equivalent dot-file for another shell.

```
export PERL5LIB=\${PERL5LIB}:<path to selection pipeline>/lib/perl5
```

The command to run vcf-merge is as follows.

```
vcf-merge <vcf1.vcf> <vcf2.vcf> ..... > big_vcf.vcf
```

3. My VCF file is not split by chromosome how do I get my VCF into a single chromosome?

The vcftools program can be used to extract each chromosome from your full vcf file. If you do not have the vcftools program installed the bin/ directory contains exactly what you need. For example, for human 1000 genomes data to extract chromosome 2 from your VCF file use the following command.

```
vcf-tools --vcf big_vcf.vcf --chr 2 --out chr2 --recode
```

The command will generate a vcf file named chr2.recode.vcf containing only data from chromosome 2.

## 4. Rehh alterations

The rehh package source included with the pipeline has been altered to match the output filters used in Voight's paper. If the EHH > 0.05 reaches the end of a chromosome or the start of a gap > big\_gap, then no value is returded for the core snp. The small\_gap specifies the gap distance to reduce the distance spanned by the gap by a multiplicative factor specified by small\_gap\_multiplier. The formula for the penalty is  $\frac{small\_gap\_multiplier}{gap\_size}$ . To match the parameters used by Voight, a value of 200,000 should be used for big\_gap\_threshold, 20,000 for small\_gap\_threshold and 20,000 for small\_gap\_multiplier (Voight et al., 2006).

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