Supporting Information How to choose sets of ancestry informative markers: A supervised feature selection approach

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Abstract

We describe some details for the analysis carried out in [6]. In particular, we give details to the R-scripts and simulations used in this study.

1 Prerequisites

For data analysis as well as for our simulation studies, we rely on scripts in the statistical language R [7].

- 1. The simulation studies are performed using msprime as implemented by [3] (and most easily installed via pip3 install msprime), which is a fast coalescent simulator. In particular, structured populations (with varying population sizes etc) can be simulated. We are using the python-interface of msprime. See https://msprime.readthedocs.io/en/stable/ for more information.
- 2. For multicore-computing, we require the R-package parallel.
- 3. Since data from the 1000 genomes project [1], which is analysed here, and the data from coalescent simulations come both in vcf-format (which stands for *variant-call-format*), we require the R-package vcfR; see [4].
- 4. For some steps in the analysis, we use vcftools [1] and bcftools [5].

In addition, data from the 1000 genomes project (phase 3) was downloaded, as well as information on the sampling locations. For this, we used

wget ftp://ftp.1000genomes.ebi.ac.uk/vol1/ftp/release/20130502/ALL.*
wget ftp://ftp.1000genomes.ebi.ac.uk/vol1/ftp/release/20130502/
integrated_call_samples_v3.20130502.ALL.panel. The latter file was renamed
1000G_SampleListWithLocations.txt, and the first row (the header) was removed.

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2 File structure

We are using the following filestructure:

• code: Contains the functions tools.r as described in Section 5, as well as commands for using the coalescent simulations, as described in Section 4;

- data/1000G: Contains the data of the 1000 genomes project (see the wget-commands from above);
- data/sim: Contains the simulations; subfolders are ooa for the Out-Of-Africa model of human evolution, and si for the symmetric island model.
- data/AIMsets: Contains the AIMsets and further analysis we find using our method;
- doc: Contains the documentation.

We note that several files pipeline*.r exist with all commands for obtaining interesting results, as described below. In particular, the master directory contains a file pipeline_example.r, which serves as a starting point to check the installation and for a first impression. We briefly describe it next.

3 A small example from pipeline_example.r

Using the example datafile data/sim/ooa/ooa_chromosome_1_example.vcf.gz, together with the information contained in ooa_SampleListWithLocations.txt on sampling locations, all steps as described in Section 6 are carried out. The main output is in the following files:

- AIMs: A list of AIMs is produced here. Note that the identifiers of SNPs in the simulated dataset which is used here, are of the form chr_pos.
- AIMs.vcf.gz: Here, the data is stored for all AIMs in AIMs and all 2400 individuals from the dataset (800 diploids from the simulated version of Africa, Europe and Asia each).

addition, the numbers of segregating stores sites put files. Since there is only one input file in this example, sinooa_SampleListWithLocations.txt gle number is stored. ated from the datafile and contains the names of individuals and their locations. training_ooa_SampleListWithLocations.txt and sampling test_ooa_SampleListWithLocations.txt are the subsets of training and test data. predictions.csv store the posterior probabilities for each individual to come from any of the continents. classifications.tab stores for each individual which deme has the maximal posteriori probability. sampleAIMs.rds is an R-file, which stores an intermediate step. Some more files are generated within the folder tmp.

4 Coalescent simulations

Once msprime is installed, the python interface can be used. In code/simulate4biogeo.py, we collected the commands needed to run msprime. A file pipeline_simulated_ooa_data.r used for analyzing the simulations is described in Section 7.

We ran two sets of simulations: (i) a symmetric island model; (ii) the model for human population history from [2], briefly called the out-of-Africa model (ooa) in the sequel. Both cases come with the parameters:

- num_chromosomes: This is the number of chromosomes which are simulated. Here, chromosomes are treated as (stochastically) independent copies of the same coalescent process.
- random_seed: In order to have reproducible results, a seed can (but does not have to) be set.
- ploidy: Since the output of msprime usually only treats haploid samples, they have to be combined in order to form diploids. So, one might want to set ploidy = 2 for humans.

In order to have the same pipeline for simulated and real data, we use the function write_vcfgz in order to obtain the simulations and real data in the same format.

4.1 Symmetric island model

Consider a population structure given by 5 continents or islands with constant bidirectional and equal migration rate m between all pairs of continents. For each migration rate m = 1, 2, ..., 8 (which stands for the expected number of haploid individuals migrating between any (fixed) pair of continents. We performed 10 independent simulations, where each dataset consists of 800 individuals from each of the five continents. Simulations are generated with the command:

python3 code/simulate4biogeo.py si *m* random_seed num_chromosomes

4.2 Out of africa model

The human demographic history as inferred by Gutenkunst et al. [2] was used to mimic the human population structure. Adopting the model for human genetic history from [2] (see their Table 1 and Figure 2), we have populations 1, 2 and 3, modeling Africa/YRI, Europe/CEU and Asia/CHB, respectively. We simulated 10 independent datasets under this model with a constant mutation rate of 0.125 and a recombination rate of 1.25. For each simulation run 1600 (haploid) individuals for each of the three continents (Africa, Europe and Asia) were generated. Simulations are generated with the command:

python3 code/simulate4biogeo.py ooa seed 20

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5 R-functions

Here is a short description of used variables, all of which are used in the functions described below:

- filenameInds: A tab-delimited file, read by read.table, which is required to have the following columns 1st: identifier for an individual; 3rd: population of the individual; the file must not have a header and then the same number of lines as filenameData; There are two more variables, filenameIndsTraining and filenameIndsTest, which have the same structure, and contain training and test sets;
- inds: A matrix with two columns; the first column contains the individual identifiers, the second contains the sampling location;
- samplesInDeme: A vector of integers containing the number of individuals for all demes;
- nss: A vector of integers containing the number of segregating sites for all files;
- snplist: A file containing SNP-identifiers. Usually used for extracting these SNPs using vcftools;
- sample: A matrix of nss columns (the SNPs) and sum(samplesInDeme) rows (the individuals). Entries are either 0/1 (if diploid = FALSE) or 0/1/2 (if diploid = TRUE);
- samLoc: A vector of length nrows(inds), indicating which sample comes from which deme;
- demes = unique(samLoc) are all sampling locations;
- freqs (Usually obtained through getfreqs(sample, samLoc, diploid)): A matrix with length(demes) rows and nss columns, containing the frequencies of the derived allele in the sample;
- error.type: The type of error by which the classification method is evaluated; either misclassification or logloss;
- filenameData: The name of the file of the data, which must be a ...vcf.gz file. Note that filenameData must not contain this suffix. This file must only contain bi-allelic markers and no missing data;
- filenameDataVCFGZ: Same as filenameData, but including the suffix .vcf.gz;
- method: The method used for classification; either naiveBayes (most often) or informativeness; other methods, such as logistic regression, are not implemented;

Here is a list of functions contained in tools.r:

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 appTestError(sample.test, freqs.training, samplesInDeme.training, samLoc.test, method = "naiveBayes", error.type="misclassification", diploid = FALSE)

For error.type, which is either misclassification of logloss, we give the error when classifying sample.test, and freqs.training is used to obtain the predictions;

- cut.VCF(filenamesDataVCFGZ, stepsize, removeUnnamed = TRUE)
 For handling of large files and limited memory, it might be beneficial to cut it in smaller pieces, containing only a subset of SNPs. Here, stepsize determines the number of SNPs in the smaller files. By default, SNPs without an identifier are excluded. The resulting files are tmp/chr1_segment1.recode.vcf.gz etc.;
- extract.and.merge(filenamesDataVCFGZ, snplist, outfile)
 Here, filenamesDataVCFGZ is a vector of filenames and snplist is a file containing a list of SNPs (e.g. possible AIMs). The result is a file outfile of vcf.gz-format, which stores exactly the sample information for SNPs in snplist;
- find.AIMs.fromRDS(filenamesRDS, sampleAIMs = NULL, inds, numAIM, method = "naiveBayes", error.type="misclassification", diploid = FALSE, outfile = "AIMs") filenamesRDS is a vector of rds-files (i.e. R-files containing a single R-object) containing the samples, i.e. all files in filenamesRDS contain the same individuals, but different SNPs. numAIM is the number of AIMs which should be found. It is possible to start already with a few AIMs, stored in sampleAIMs. During runtime, when a new AIM is found, it is appended to the file AIMs, and sampleAIMs.rds stores the data at the AIM sets;
- find.nextAIM.fromRDS(filenameRDS, sampleAIMs, inds, method, error.type, diploid)
 This function is called repeatedly from find.AIMs.fromRDS in order to find a single new AIM. This is called stepwise forwards search of AIMs;
- find.SNPs.inVCFGZ(filenamesDataVCFGZ, snplist)
 Given the SNPs in the file snplist, determine which of them are present in the set of files filenamesDataVCFGZ;
- getData(filenameDataVCFGZ, skip=0, windowSize=-1, inds, snps = TRUE, diploid = FALSE)

 Using the function read.vcfR from the package vcfR, the data is read. Using skip and windowSize, only a subset of SNPs from filenameDataVCFGZ can be obtained; the value is a list containing sample and samLoc;
- getfreqs(sample, samLoc, diploid = FALSE)
 From sample and samLoc, we know which frequencies are observed in which demes.
 The result is given here, a matrix with length(demes) rows and nss columns;

- getfreqs.from.vcf(filenameData, filenameInds, snplist=FALSE)
 For large files, getfreqs from above might have memory issues. This funtion uses vcftools with its freqs2-option in order to write .frq-files for frequencies in all demes. These are subsequenty extracted and reported in the file paste(filenameData, "_freqs");
- getIdentifiers(filenameDataVCFGZ) Returns the SNP-identifiers used in file filenameDataVCFGZ;
- getNss(filenameDataVCFGZ) Returns the number of segregating sites in the file filenameDataVCFGZ;
- getTestandtrainingset(filenameInds, filenameIndsTraining, filenameIndsTest, size)
 Using a random number generator, the individuals are split into a training and a test set. filenameIndsTraining and filenameIndsTest are the filenames where the individuals for these two sets are stored; size is a vector containing the sizes of the testset in all populations;
- informativeness.global(freqs)
 A vector of length nss. This function computes the informativeness for all SNPs in freqs.
 This function needsxlogx2(x) and xlogx(x);
- logloss.error(prediction, true.value)
 The logarithmic loss;
- log.prediction.naiveBayes(sample.test, freqs.training, samplesInDeme.training, diploid) and prediction.naiveBayes(sample.test, freqs.training, samplesInDeme.training, diploid)

 Based on freqs.training, probabilities are computed which give the chances that some individual from sample.test comes from one of the demes. Since this is based on the naive Bayes approach, we also need number how many samples were taken from each deme (samplesInDeme.training); We use here prediction.naiveBayes = exp(log.prediction.naiveBayes) for numerical stability;
- misclassification.error(prediction, true.value)
 The misclassification error;
- setcores(cores=1):
 Determines the number of cores used for classification (in step6 below); this requires the R-package parallel;

6 Analyzing data from the 1000 genomes project

We now describe the file pipeline_1000G_AIMs_noAMR.r in detail. All other files called pipeline*.r work similarly. After importing necessary packages, we set the number of cores

using setcores to a maximum number on our machine. These will be used in parallel to find AIMs in step6 below. Then, some parameters need to be set:

The variable testindsperdeme = 100 indicates that 100 samples deme/population/continent are taken. The rest of the data is used as training set. Since we ignore all samples in population AMR, we only have demes=4. The informativenessBound = 0.9 is used as a presection step for the AIMs. This number means that only SNPs which have an informativeness in the top 10% of all observed informativenesses qualify to be chosen as AIM. Then, stepsize = 10000 is used when chopping the data to smaller files. We found this number to work well with 64GB of RAM on a machine with 32 cores. Then, numAIM = 30 will eventually produce a set of 30 AIMs. Finally, we use the standard setting, as also described in the main text: method = "naiveBayes", error.type="logloss" (which is used to find the AIMs, although we finally report the misclassification error) and diploid = TRUE. The user might want to set a seed using something like set.seed(31) in order to have reproducable results. (This is only used for creating the training and test set of individuals. The filenames filenameInds, filenameIndsTraining, filenameIndsTest and the vector filenameData is created for future reference.

Then, the following steps are carried out (which can be in- or excluded depending on where the user wants to make progress):

- prepare: Based on the file data/1000G/1000G_SampleListWithLocations.txt, we exclude AMRs and write the resulting file in the current directory. Test and training set are also written using getTestandtrainingset.
- step0: Using vcftools, we reduce dataset to bi-allelic SNPs, and remove indels. The resulting files are called tmp/chr1_biallelic.recode.vcf.gz etc.
- step1: Using the function getfreqs.from.vcf, we obtain frequencies for all SNPs in all populations for the training set.
- step2: Filter the most informative SNPs (i.e. SNPs with informativeness above the informativenessBound quantile) and write a new vcf.gz-File, denoted tmp/chr1_informative.recode.vcf.gz etc. In these files, we have all SNPs which are eligible to become AIMs.
- step3: For better handling, we cut the files in smaller pieces with stepsize SNPs each using the function cut.VCF. The resulting files are tmp/chr1_segment1.recode.vcf.gz etc.
- step4: Since they are needed in the remaining steps, we read off the number of informative sites for all chromosomes are write them into the file nss.
- step5: We use all files from step3, read in the data in the format we need, and store the result in an R-file (i.e. ending with .rds). The advantage is that these files are very quick to read in R.
- step6: The main (i.e. most time-consuming) step in the analysis is where we find the AIMs. Each of the rds-files is read, the best SNP is found in each file (i.e. the SNP which,

if included as AIM, has the smallest logloss-error), and we finally take the SNP with the smallest such error. We look stepwise for numAIM AIMs. However, each time a new AIM is found, it is added to the file AIMs, and an rds-file containing all AIMs ist stored in sampleAIMs.rds.

- step7: Finally, we have all AIMs and create (using extract.and.merge) a vcf.gz-file for all SNPs from step6. The resulting file is called AIMs.vcf.gz.
- step8: From AIMs.vcf.gz, we obtain for all individuals the posteriori probabilities to come from one of the four continents. The results are stored in predictions.csv. In addition, classifications.tab stores the continents with the maximal posteriori probability.

7 Analyzing simulated data from the Out-Of-Africa model

Since simulated data also comes in the vcf.gz-format, we can use the same pipeline as in Section 6. The main difference is that the data is already bi-allelic, but we have to simulate the data before we can begin:

- simulate: A multicore call is used for simulating the model as described in Section 4.2. The data is stored in data/sim/ooa/tmp/.
- prepare: Here, we know that the samples come in the order of the coalescent simulations. This means that in the data file, there are 800 diploids from AFR, EUR and ASI. The identifiers of the individuals are stored in ooa_SampleListWithLocations.txt. Test and training-set are obtained as in the analysis of the data from the 1000 gebomes.
- step1 step8 are carried out as in Section 6.

References

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