
TRIBES

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TRIBES is a user-friendly platform for relatedness detection in genomic data. **TRIBES** is the first tool which is both accurate (up to 7th degree) and combines essential data processing steps in a single platform.

Accurately classifying the degree of relatedness between pairs of individuals has multiple important applications, including disease gene discovery, removal of confounding relatives in genome wide association studies (GWAS) and family planning. Currently no tools are available which are accurate beyond 3rd degree and combine the necessary data processing steps for accuracy and ease of use. To address this we have developed '**TRIBES**', a user-friendly platform which leverages the GERMLINE algorithm to accurately identify distant relatives. **TRIBES** enables user-guided data pruning, phasing of genomes, IBD segment recovery, masking of artefactual IBD segments and finally relationship estimation. To facilitate ease-of-use we employ 'Snakemake', a workflow tool which enables flexibility and reproducibility.

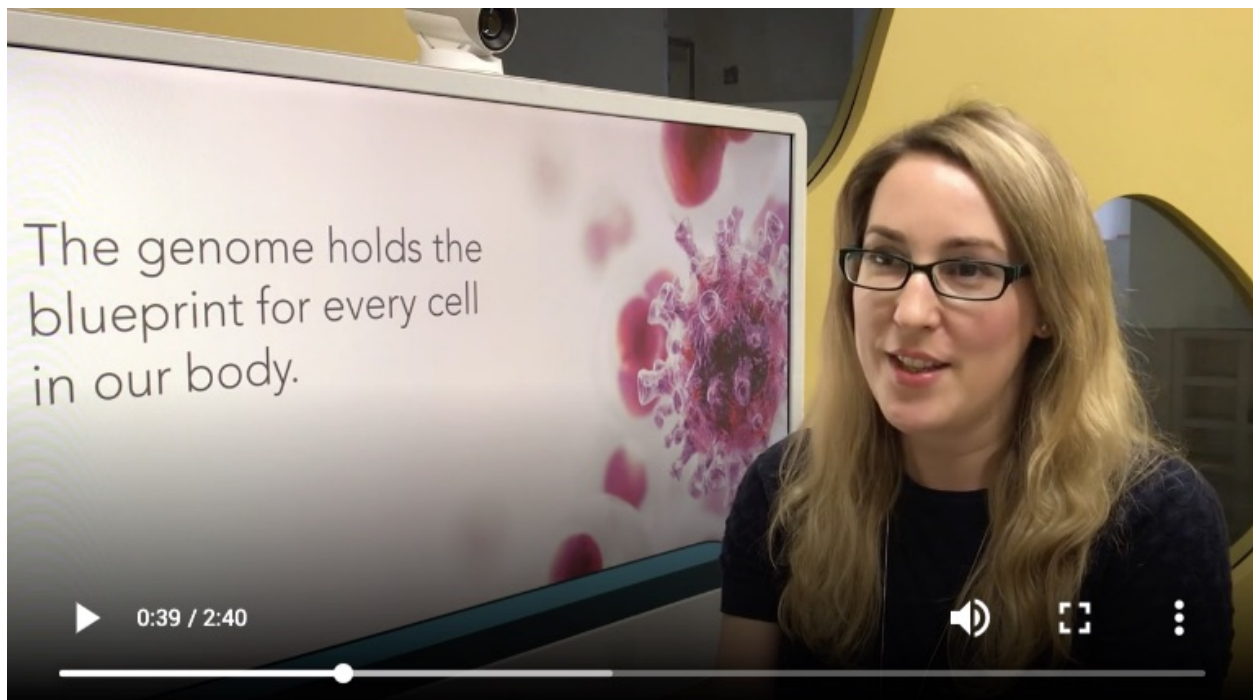
We demonstrate the accuracy of **TRIBES** in our publications [here](<https://www.biorxiv.org/content/10.1101/686253v1>) and [here](<https://www.biorxiv.org/content/10.1101/685925v2>)

Briefly, input data to **TRIBES** is quality control filtered, joint sample VCF. *TRIBES* then follows these steps:

1. The VCF is filtered using quality metrics contained within the VCF file.
2. The resultant VCF is then phased using BEAGLE.
3. IBD Segments are then estimated using GERMLINE.
4. Artefactual IBD is masked using a reference file by adjusting segment endpoints.
5. Adjusted IBD Segments are then summed to estimate relationships.
6. **TRIBES** returns result files, including .csv of estimated relationships.

The full **TRIBES** pipeline is described in detail in [Supplementary Material](<https://www.biorxiv.org/content/10.1101/686253v1.supplementary-material>).

Watch a short video introducing *TRIBES* and its applications



1.1 Getting Started

This section describes the analysis of an example dataset. We advise that you run **TRIBES** on the example data first, to confirm you have installed **TRIBES** correctly. To run **TRIBES** on your own datasets, refer to instructions from [Installation](#) onwards.

TRIBES requires a 64-bit version of Linux, MacOS or Windows 10, and about 10GB of free disk space for software, reference and example data. Install **TRIBES** using one of the methods described in the [Installation](#) section.

Alternatively you can run **TRIBES** from a pre-packaged docker image using `docker` or `singularity` (see: [Containers](#) section).

After installation return to [Testing installation on example dataset](#)

1.1.1 Testing installation on example dataset

To demonstrate how **TRIBES** works we will use an example dataset (TFCEu) with reference data from 1000 Genomes ‘EUR’ superpopulation (REF-G1K_EUR).

Create and navigate to a directory for reference and sample data, e.g `$HOME/tribes-data`

```
mkdir -p $HOME/tribes-data
cd $HOME/tribes-data
```

Download and uncompress reference data (1.2 GB)

```
wget https://d3o0p4nu4e38rq.cloudfront.net/downloads/reference/1.0/REF_G1K-EUR_0.001.
→tar.gz
tar -xzf REF_G1K-EUR_0.001.tar.gz
rm REF_G1K-EUR_0.001.tar.gz (optionally)
```

The reference data is subset of 1000 genomes dataset with unrelated ‘EUR’ individuals and it’s used in various stages of preprocessing (e.g. LD pruning, phasing and filtering on MAF).

Download and uncompress example data (390 MB)

```
wget https://d3o0p4nu4e38rq.cloudfront.net/downloads/examples/0.2/TFceu.tar.gz
tar -xzf TFceu.tar.gz
rm TFceu.tar.gz (optionally)
```

The sample data is a synthetic pedigree created from unrelated 1000 Genomes ‘CEU’ individuals. For more info on the dataset see the [Installation](#) section. Inside the TFceu directory you will find the following files:

- TF-CEU-15-2.vcf.gz - the source multisample VCF files
- TF-CEU-15-2.true.rel - the true pairwise relations
- glk_ceu_family_15_2.ped - pedigree
- config.yaml - the configuration file describing the steps taken in **TRIBES** pipeline.

The config.yaml provides configuration for the pipeline defining the location and name of reference data and the true relations file, as well as the name of the input VCF file and the preprocessing steps required prior to IBD/relatedness estimation, e.g.:

```
rel_sample: "TF-CEU-15-2_BiSnp_MAF@0.01_LD"
```

identifies TF-CEU-15.vcf.gz as the input file and applies 3 pre-processing steps: filtering on biallelic SNPs and a minor allele frequency (MAF) of 0.01 plus LD pruning. All steps that can be used in *TRIBES* pipeline are described below in *Preparing a custom pipeline*

Note: Please note that the IBD estimation requires a phased VCF file. If the input file is not phased, pre-processing must include phasing (usually last the last step, after filtering), e.g. TF-CEU-15-2_BiSnp_MAF@0.01_LD_PH (where ‘PH’ in the file name indicates to phase without reference) or TF-CEU-15-2_BiSnp_MAF@0.01_LD_RPH (with ‘RPH’ in the filename indicates to phase with reference). This is not required in this example because the input VCF is phased.

To run from a local installation do to your **TRIBES** installation directory and run **TRIBES** with:

```
./tribes -d $HOME/tribes-data/TFceu -j <no_cpu_cores> estimate_degree_vs_true
```

To run using docker:

```
docker run -it -rm -v "$HOME/tribes-data:$HOME/tribes-data" docker.io/piotrszul/
↳ tribes -d $HOME/tribes-data/TFceu -j <no_cpu_cores> estimate_degree_vs_true
```

To run using singularity:

```
singularity run -e docker://docker.io/piotrszul/tribes -d $HOME/tribes-data/TFceu -j
↳ <no_cpu_cores> estimate_degree_vs_true
```

Where no_cpu_cores is the number of CPU cores to use. estimate_degree_vs_true calls **TRIBES** to perform all relatedness estimation steps described in IBD/Relatedness steps: under *Preparing a custom pipeline*.

It takes about 20 minutes to to run the entire pipeline using 4 cores.

1.1.2 TRIBES output for example dataset

Upon the successful completion, you can find the final and intermediate stages of the pipeline in \$HOME/tribes-data/TFceu/ (~2.3GB). In particular:

- TF-CEU-15-2_BiSnp_MAF@0.01_LD_PH_GRM-allchr_FPI_IBD.csv - includes the pairwise estimate of the degree of relatedness (EstDegree)

- TF-CEU-15-2_BiSnp_MAF@0.01_LD_PH_GRM-allchr_FPI_IBD_RVT.html - notebook which compares estimated degrees vs the reported (true) ones.

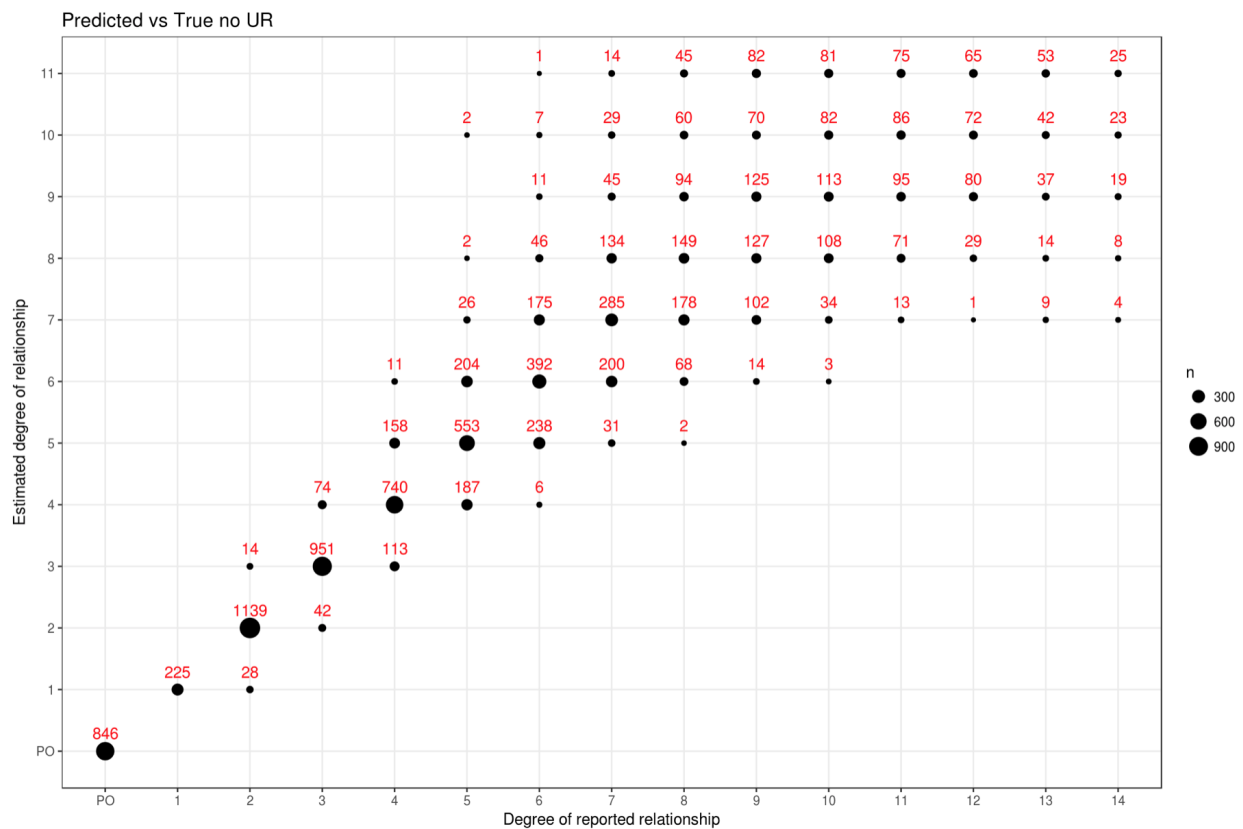
The estimated relatedness is in CSV format with the following columns and data:

```
Id1, Id2, IBD0.cM, IBD1.cM, IBD2.cM, EstDegree
NA07347, NA11919, 0.999073851764529, NA, NA, 11
NA12058, NA12829, 0.999107459568523, NA, NA, 11
```

To see the comparison results you can open the report in your preferred browser (e.g. firefox):

```
firefox $HOME/tribes-data/TFceu/TF-CEU-15-2_BiSnp_MAF@0.01_LD_PH_GRM-allchr_FPI_IBD_
↪RVT.html
```

The comparison is presented in the form of a dot chart like this:



1.2 Installation

1.2.1 Installation for workstation use

TRIBES requires a 64-bit version of Linux, MacOS or Windows 10.

Windows Subsystem for Linux (WSL)

To run TRIBES on Windows 10, first install [Ubuntu](#) from the Microsoft Store.

Then open the Ubuntu app from the Start menu.

Miniconda

TRIBES has a list of dependencies required to be installed prior to running the analysis pipeline. For this, we use *Miniconda*.

Install miniconda (3.7 or 2.7) from <https://docs.conda.io/en/latest/miniconda.html>:

```
wget https://repo.anaconda.com/miniconda/Miniconda3-latest-Linux-x86_64.sh
sh Miniconda3-latest-Linux-x86_64.sh
```

Download *TRIBES*

Download the latest release of *TRIBES* from <https://github.com/aehtc/TRIBES/releases> and extract it to your selected directory.

Alternatively you can clone the most recent (unstable) version from github:

```
git clone https://github.com/aehtc/TRIBES.git
```

Installing *TRIBES*

Go to the *TRIBES* installation directory.

Install dependencies (requires about 500 MB for software packages):

```
./setup/install-with-conda.sh
```

This will create a conda environment named `tribes` and install all required dependencies, as well as create the appropriate *TRIBES* configuration file at `~/.tribesrc`

To check the installation run:

```
./tribes
```

This should display among others usage info.

After you install *TRIBES*, return to [Testing installation on example dataset](#) to test the installation on example data.

1.2.2 Manual installation

TRIBES is implemented as a *snakemake* pipeline and relies a number of bioinformatics tools for processing, such as *bcftools*, *bgzip*, *tabix*, *vcftools*, *germline*, *beagle* as well as a number of *python* and *R* packages.

The complete list of dependencies and their required (minimal) versions can be inferred from the conda environment file at: [setup/environment.yaml](#)

They can be installed using the OS specific way (e.g. using *apt* or *yum* on Linux or *brew* on MacOS)

In addition *TRIBES* requires `tribes.tools` *R* packages which can be installed from sources with:

```
Rscript --vanilla -e "install.packages('R/tribes.tools', repos=NULL)"
```

1.2.3 Installation on HPC Cluster

snakemake and thus *TRIBES* can run on HPC clusters (for example with `slurm`).

An example setup for CSIRO HPC cluster is described in [README-CSIRO.md](#) and can be used as a guide to configure *TRIBES* on other clusters.

For more information on running `snakemake` on HPC clusters please check the `snakemake` documentation <https://snakemake.readthedocs.io/en/stable/>

1.3 Usage

Read the sections below to run *TRIBES* on your own data, with a custom pipeline

1.3.1 Input data

TRIBES requires the following input files:

- `filename.vcf.gz` - multi-sample VCF file containing sample genotypes
- `filename.true.rel` - true pairwise relations (optional, only if a user has known relations and wants to calculate accuracy of estimated relationships)
- `config.yaml` - pipeline configuration file defining the location and name of reference data, the true relations file, the input filename and the preprocessing steps required before IBD/relatedness estimation

Refer to files inside example dataset `TFCEu/` directory for correct format for these input files.

1.3.2 Preparing a custom pipeline

A key strength of *TRIBES* is that is a flexible pipeline, utilizing `snakemake`, to enable the user to specify which pre-processing steps they want to include.

The following steps can be used in the pipeline.

Preprocessing:

- `NM`: retain only loci with non-missing genotypes
- `BiSnp`: retain only bi-allelic SNPs
- `BiSnpNM`: combines `BiSnp` and `NM` in a single step
- `MAF@<maf-threshold>`: filters for `MAF >= maf-threshold`, e.g. `MAF@0.01`. `MAF` is determined from the reference data `AF` annotation which is also added to the output in `REF_AF` annotation.
- `LD`: prune on LD with the reference defined in `G1K_SNP_EUR` (`bcftools +prune -l 0.95 -w 1kb`)
- `QC`: filter on quality (with `bcftools`: `INFO/MQ>59 & INFO/MQRankSum>-2 & AVG (FORMAT/DP)>20 & AVG (FORMAT/DP)<100 & INFO/QD>15 & INFO/BaseQRankSum>-2 & INFO/SOR<1`)
- `PH`: phase (using `beagle`) without reference
- `RPH`: phase (using `beagle`) with reference defined in `ref_sample` config parameter

IBD/Relatedness steps:

- GRM: detect pairwise IBD segments using `germline`
- FPI: filter out IBD segments using a mask defined in the reference data.
- IBD: estimate pairwise degree of relatedness based on IBD0
- RVT: compare the estimated degree to the known degree, reflecting accuracy

1.3.3 Examples

Example 1

For example, a user may wish to identify relationships using an unphased input VCF. They wish to filter on allele frequency of $MAF = 0.01$ and then phase the data using reference file and estimate relatedness. They would then need to edit the `config.yaml` file from the example data `TFCEu` directory to reflect their input VCF filename and processing steps. Their input VCF file should be in the same `TFCEu` directory, for the `config.yaml` file to work.

Their `config.yaml` file would look like this:

- `rel_sample`: `filename_BiSnpNM_MAF@0.01_RPH` [where `filename` refers to the input VCF filename]
- `ref_dir`: `../REF_G1K-EUR_0.001` [where `ref_dir` is the location of the reference directory, which hosts the cohort used for filtering on MAF and LD, phasing and masking steps]

The user would then run *TRIBES* from the installation directory as in the *Getting started* section

```
./tribes -d $HOME/tribes-data/TFCEu -j <no_cpu_cores> estimate_degree
```

where `estimate_degree` is an alias which calls *TRIBES* to perform the GRM, FPI and IBD steps described under ‘IBD/Relatedness steps’ in *Preparing a custom pipeline*

Example 2

Alternatively, a user may want to identify novel relationship, as well as confirm known relationships. They wish to pre-process the VCF to filter on $MAF = 0.01$ and quality metrics, then phase the data using reference, estimate relationships and compare estimated with known relationships.

Their `config.yaml` file would look like this:

- `rel_sample`: `filename_BiSnpNM_MAF@0.01_QC_RPH`
- `ref_dir`: `../REF_G1K-EUR_0.001`
- `rel_true`: `filename.true.rel` [a reference file containing known relationships, required if step RVT is used in the pipeline]

The user would then run *TRIBES* from the installation directory as in the *Getting started* section

```
./tribes -d $HOME/tribes-data/TFCEu -j <no_cpu_cores> estimate_degree_vs_true
```

If users provide a `rel_true`: file in the `config.yaml` file, they can call `estimate_degree_vs_true` which is an alias that calls *TRIBES* to perform the GRM, FPI, IBD and RVT steps described under ‘IBD/Relatedness steps’ in *Preparing a custom pipeline*

1.4 Datasets

1.4.1 1000 Genomes EUR (REF_G1K-EUR_0.001)

Location: https://d3o0p4nu4e38rq.cloudfront.net/downloads/reference/1.0/REF_G1K-EUR_0.001.tar.gz

This is a reference dataset used for MAF filtering, LD pruning and phasing. It's based on the data from release 3 of [1000 Genomes Project](#). It includes all biallelic SNPs with $MAF > 0.001$ for unrelated individuals from 'EUR' superpopulation.

- `VCF` : all sample genotypes (separate file per chromosome)
- `sample.txt` : list of included EUR samples
- `ersa-mask.tsv`: list of regions with excessive IBD (generated with [ersa](#) for this sample)
- `plink.chrALL.GRCh37.map.gz`: genetic map (included for convenience)

1.4.2 TrueFamily CEU (TFCEu)

Location: <https://d3o0p4nu4e38rq.cloudfront.net/downloads/examples/0.2/TFCEu.tar.gz>

This is synthetic dataset with simulated genotypes based on unrelated individuals from CEU population of [1000 Genomes Project](#). The pedigree is defined in `g1k_ceu_family_15_2.ped` and includes 15 generations.

- `TF-CEU-15-2.vcf.gz` : VCF file for the simulated genotypes
- `g1k_ceu_family_15_2.ped`: pedigree
- `TF-CEU-15-2.true.rel` : true relations

1.5 Containers

TRIBES docker image includes the pipeline and all the dependences and it's publicly available from <https://hub.docker.com/r/piotrszul/tribes> as `docker.io/piotrszul/tribes` and the most recent version can be pulled with:

```
docker pull docker.io/piotrszul/tribes
```

To use a specific version e.g.: `0.2.0` please use `docker.io/piotrszul/tribes:0.2.0` as the docker image name.

It's an executable image with `snakemake` as an entry point.

When running using docker it's necessary to mount the reference data and pipeline data volumes (or local filesystem) so that the container have access to both, e.g:

```
docker run -it --rm -v <path-to-ref-data>:<path-to-ref-data> -v <path-to-data>:<path-to-data> docker.io/piotrszul/tribes -d <path-to-data> <other_options> ...
```

When running with `singularity` this may not be needed if the volumes with data and reference data are mounted as per configuration. One important consideration though is to use `-e` flag as some host environment variables (e.g. related to python) cause issues while running in the container:

```
singularity run -e docker://docker.io/piotrszul/tribes -d <path-to-data> <other_options> ...
```


CHAPTER 2

Contact

Please report any issues or ideas at: <https://github.com/aeherc/TRIBES/issues>

Or contact the *TRIBES* team at: TBP

CHAPTER 3

Indices and tables

- `genindex`
- `modindex`
- `search`