The STRAF Book

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Preface

What is this book?

This is the online version of **The STRAF Book**, which is currently under active development. It is dedicated to the STRAF software, a web application for the analysis of genetic data in forensic practice.

Forensic and population genetics, lost sisters

Genetics has many faces, and population and forensic genetics are two of them. If we were to briefly summarise their respective scopes, we could say that the former aims at understanding genetic differences within and between populations and the latter is the application of those findings to legal matters.

Forensic genetics and population genetics have always been tightly linked disciplines. This is likely because quite a number of questions they address are similar. Even though problems in forensics and population genetics seem different, they often correspond to the same question, simply phrased differently.

In population genetics, a common goal is to characterise the genetic diversity of a set of populations, by looking at how related individuals are within and between populations. DNA profiling used in criminal investigations aims at matching different DNA samples. To be able to evaluate the probative strength of such a match, forensic geneticists need to know how much the members of the relevant population are ge-

netically related to each other. The same applies to the calculation of probabilities for different hypotheses of kinship, e.g. in paternity tests. Both fields aim at **understanding** and **quantifying** the **relatedness** of individuals based on their DNA.

Software and metrics developed in the population genetics for the study of the evolution of species are now used routinely in forensic genetics practice. But forensics is not just *applied population genetics*. The legal implications and unique situations encountered in the forensics world also led to the development of relevant statistical tools and metrics with a more specific purpose.

And then there was STRAF

STRAF was born from the encounter of two scientists: a forensic geneticist and a population geneticist, in 2017, in the beautiful city of Bern (Switzerland). Martin came to visit a population genetics lab, where Alexandre was pursuing his Ph.D. thesis at that time. This encounter led to a fruitful collaboration when they realised that some tools used in population genetics should be made more accessible to the forensics community.

This encounter led to a fruitful collaboration when they realised that some tools used in population genetics could be leveraged by the forensics community. The most striking example is the computation of forensics parameters, that describe for example how good are our loci at discriminating samples. These parameters were typically computed using a spreadsheet that had been created by one of the suppliers of assays used to genotype samples. It is the mythical PowerStats v1.2 spreadsheet, allowing to compute forensic statistics and allele frequencies in Microsoft Excel. It has been since then removed from the Internet, and forensic geneticists started sharing this spreadsheet among each other, circulating almost secretly, "under the cloak" as French speakers would say.

As similar operations were done in routine in population genetics, we already had some scripts for the analysis of STR data. Then, after we applied them to an existing dataset, we decided to put everything into a web application so that the forensics community could benefit from it.

A few weeks later, STRAF was born [Gouy and Zieger, 2017], and after four year,

STRAF had become a widely used tool by the forensics community, but not only. It has been used as a support for teaching population genetics, and has been used in evolutionary biology studies. The positive reception of the software in the community motivated its development over the years until the release of STRAF 2.0 in 2021.

What will you learn?

By reading this book, our hope is that you will:

- Get a brief overview of common concepts in forensic and population genetics
- Learn how to use the STRAF software for STR data analysis through practical applications
- Be able to interpret common metrics and analyses used in forensics practice

Outline

The book is organised as follows:

- In Chapter 1, we'll start by an introduction to essential forensic and population genetics concepts.
- In Chapter 2, we will focus on data, from its generation to its preparation for downstream analysis in STRAF.
- In Chapter 3, we will review forensic parameters that can be computed in STRAF, and discuss their interpretation.
- In Chapter 4, we will review essential population genetics concepts and describe population genetics indices that can be computed in STRAF.
- In Chapter 5, we will focus on multivariate statistics and how they can provide insights into population structure, with a particular focus on Principal Component Analysis (PCA) and Multidimensional Scaling (MDS), two widely used approaches in genetics.

In Chapter 6, we will explain how to compare samples of interest to reference
populations by loading reference allele frequencies into the software and
performing a Multidimensional Scaling analysis.

 In Chapter 7, we gather recommendations around potential next analysis steps by presenting STRAF's file conversion capabilities and useful methods implemented in other software.

Chapter 1

Introduction

In this chapter, we will briefly introduce some essential concepts in genetics.

DNA and genetic variation

Each of our cells contains 23 pairs of **chromosomes**, composed of a long **DNA** (deoxyribonucleic acid) molecule. Under this somewhat barbaric name is hiding a simple concept. This molecule is the carrier of the information the body need for its function and development. This information is encoded by a chain of **nucleotides** of four types that can be referred to using the letters A (Adenine), T (Thymine), C (Cytosine) and G (Guanine).

Developments in biotechnologies enabled the characterisation of the DNA of individuals. These techniques also led to the discovery that this DNA varies between individuals. This **genetic variation**, also called **polymorphism**, can be used to characterise individuals and populations based on their DNA.

Markers of polymorphism

DNA variation can take different forms: it can for example be a **Single Nucleotide Polymorphism** (**SNP**), when a mutation occurs and changes a nucleotide at a given position in the genome. In that case, we would observe different nucleotides in a population at a single position.

There can also be **insertions** and **deletions** (sometimes referred to as **InDels**), of one or multiple nucleotides.

Finally, other markers of genetic variation are **Copy Number Variants** (**CNVs**), when a sequence is repeated a certain number of times. They can contain more or less repetitive units. These units can contain more or less nucleotides. **Short Tandem Repeats** (**STRs**) are a type of genetic polymorphism consisting in short sequences from 2 to 7 base pairs that are repeated several times. The **number of repeats varies** among individuals, therefore characterizing the length of those repeats regions can be useful to identify individuals. STRs are the most common markers used in forensic genetics.

Polymorphism and forensics

DNA profiling and typing

As DNA varies between individuals, DNA typing became a central element of the forensic scientist toolkit. For example, typical questions forensic genetics aims at answering include:

- What is the probability that a DNA profile at a crime scene does not only match
 the person of interest, but also a person picked randomly from the relevant
 population?
- Based on the detected genetic variants: Is it more likely to detect these combinations if the two persons of interest are brother and sister or if they are unrelated?

The role of population genetics

To answer these questions, it is crucial to first get a good characterisation of **genetic variant** (or **allele**) frequencies in populations of interest, at different **loci** across the genome. Indeed, these frequencies can vary widely among populations.

In this context, STRAF has been designed to facilitate the analysis of **population** data in forensic genetics.

Chapter 2

Importing data

In this chapter, we will explain how to prepare the input file containing genotypic data, and how to upload it into STRAF.

2.1 STR data and STRAF input format

STR data consists in various **observations**: the **genotypes** (number of STR repeats) of **each individual**, at **each locus**.

In genetics, we potentially observe two values per individual and per locus, if the markers are diploid, that is two copies are present per sample.

STRAF's input file is a text file containing the genotypes of each sample:

- The first column, named ind, needs to contain the sample ID
- The second column, named pop, contains the population ID (this column must exist even if a single population is studied)
- The next columns correspond to genotypes: for haploid samples, one column
 per locus must be reported; for diploid data, two columns per locus (with the

same name)

- Genotypes must be encoded as numbers (STRAF accepts point alleles, e.g. 9.3 in TH01)
- Missing data (e.g. null alleles) must be indicated with a "0".

For diploid data, the table should look like this:

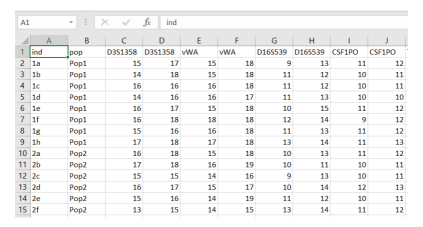
ind	pop	Locus1	Locus1	Locus2	Locus2
Α	Bern	12	14	17	17
В	Bern	14	14	13	15.2
С	Lausanne	12	16	15.2	17

For haploid data, the table would look like this:

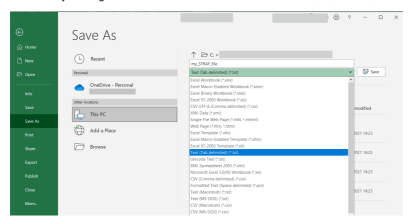
ind	рор	Locus1	Locus2
Α	Bern	12	17
В	Bern	14	13
С	Lausanne	12	15.2

2.2 Generating the input data from Excel

It only takes a few steps to generate an input file in a format that is suitable for use in STRAF. From Excel, for example, we can start from a spreadsheet looking like this:



Then, one simply needs to save this table as a tab-delimited text file. This can be achieved by clicking on Save As > Text (Tab-delimited) (*.txt)

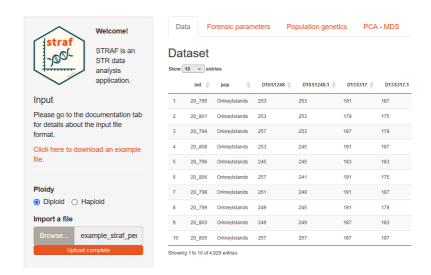


2.3 Uploading the data to STRAF

Once the input file has been prepared, it is possible to upload it into STRAF. Alternatively, you can **download an example file** by clicking on the link in the sidebar.

Then, you need to select the **ploidy** of your data (for example, *Diploid* for autosomal markers, and *Haploid* for Y-chromosome markers). After that, you can upload the file by clicking on *Browse* and selecting the file on your computer.

Once the file is uploaded, a preview will be displayed on the right and all the analyses will be available. If not, it is likely that an error has occurred. If the error message is not explicit, you can refer to the **Input file checklist** below that gathers common issues with the input file.



2.4 Common issues

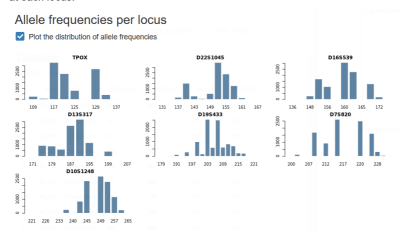
Even though you've been very careful in the generation of STRAF's input file, it is possible that you still run into an error after uploading the file to STRAF. In case STRAF cannot read your input file, we've put together a checklist to identify common issues with the input file.

Input file checklist

- Check input parameters in the sidebar: do they actually correspond to the input data?
- Check locus names: are they all different for haploid data? Do both columns for a single locus for diploid data have the exact same name?
- · Check that all missing data have been encoded with a "0"
- Try to remove any special characters from sample and locus names
- · Check for the presence of empty spaces at the end of each line
- · Check if alleles are exclusively encoded with numbers
- · Check if values are separated by tabs and not spaces
- · Check if the first two columns are names "ind" and "pop"

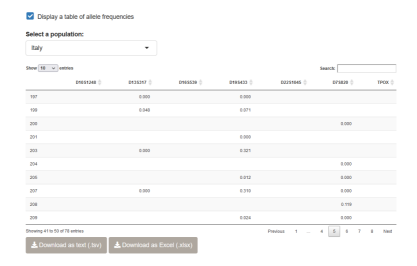
2.5 Having a first look at the data

Below the dataset preview, you will be able to generate a plot of allele frequencies at each locus.



You can also generate an allele frequency table, which is standard practice when

reporting new population data in a forensics journal. You can either download the allele frequencies in a text format (TSV), or as an Excel (XLSX) file. Note that you have the ability to select a specific population using the drop-down menu above the table.



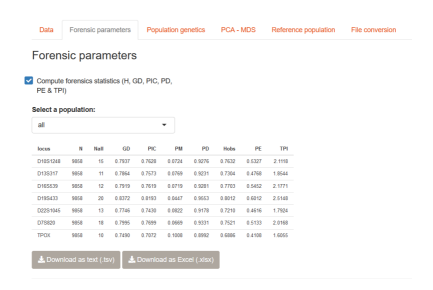
Chapter 3

Forensic parameters

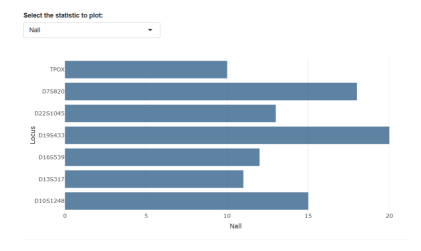
In this chapter, we'll show how to compute forensic parameters using STRAF, and provide details on how they are computed and should be interpreted. We'll introduce a few equations, but please do not be afraid! The goal of this chapter is to translate each of them into plain English.

3.1 How to compute forensic parameters in STRAF

Once your data has been uploaded, you can go to the **Forensic parameters** tab and check the *Compute forensics statistics* box. The computation will be performed and a table containing the values per locus will be displayed. The computation is done per population and overall, a drop-down menu is present to select the population.



Below the forensic parameters table, You can select the metric you would like to represent using the drop-down menu.



3.2 Details on the forensic parameters

3.2.1 Random match probability (PM)

The **Random match probability**, or probability of matching (PM), is defined as the probability of observing a random match between two unrelated individuals.

Formula

$$PM = \sum_{i} (G_i)^2,$$

where G_i is the frequency of the genotype i at a given locus in the population.

Interpretation

Computing PM means calculating, for a given locus, the frequency of each genotypes. Then we take the square of each frequency, i.e. we multiply it by itself. Finally, we sum the values of each genotype.

The intuition behind it is that if we observe a random match in a population when looking at a single locus, it means that our two samples have the same genotype at that locus. In terms of probabilities, sampling a specific genotype in the population has a probability equal to its frequency. And sampling the same genotype a second time (i.e., observing a match), is the probability of sampling this genotype multiplied by itself.

As an example, say the genotype "12-14" has a frequency of 5% in the population, the probability of having a random match between two individuals having the same genotype is 0.05×0.05 .

To get an overall probability of matching, we sum this over all possible genotypes in our population.

3.3 Power of Discrimination (PD)

The power of discrimination (PD) is defined as the probability of discriminating between two unrelated individuals.

Formula

$$PD = 1 - PM$$

Interpretation

PD is simply 1 - PM. Instead of looking at the probability of matching, we are interested in the probability of "not matching", i.e. the probability of discrimination.

3.4 Gene diversity

Gene diversity (GD, sometimes simply D), also called **expected heterozygosity** $(H_{\rm exp})$, is computed using the following estimator:

Formula

$$H_{\mathrm{exp}} = GD = \frac{n}{n-1} \left(1 - \sum_{i=1}^n (p_i)^2\right),$$

where n is the number of gene copies sampled (i.e., the sample size for haploid samples, and twice the sample size for diploid samples) and p_i is the frequency of the i^{th} allele in the population.

Interpretation

It is the probability that an individual will be heterozygous at a given locus.

As an example, a value of GD of 0.6 means that there is a 60% chance of being heterozygote at this locus.

It depends directly on the genetic diversity at this locus, which itself depends on allele frequencies in your population.

Say we have two alleles in a given population, genetic diversity will be higher in a population with allele frequencies 0.5 and 0.5 than in a population where frequencies are 0.1 and 0.9 (as less heterozygotes can be made with rare alleles). This rationale can be extended to any number of alleles.

3.5 Polymorphism Information Content (PIC)

Formula

The Polymorphism Information Content (PIC) is computed as follow:

$$PIC = 1 - \sum_{i=1}^{n} p_i^2 - \sum_{i=1}^{n-1} \sum_{j=i+1}^{n} 2p_i^2 p_j^2,$$

where \boldsymbol{p}_i and \boldsymbol{p}_j are allele frequencies.

Interpretation

The PIC can be interpreted as:

- · the probability that the maternal and paternal alleles of a child are deducible
- or, the probability of being able to deduce which allele a parent has transmitted to the child

3.6 Power of Exclusion (PE)

Formula

The **power of exclusion** (PE) is defined as:

$$PE = h^2 \left(1 - 2hH^2 \right),$$

where h is the proportion of heterozygous individuals and H the proportion of homozygous individuals in the population sample.

Interpretation

The power of exclusion depends on the observed proportions of heterozygous and homozygous individuals in a population. These proportions, multiplied as in the equation above, give the **mean exclusion probability for a paternity test**. Intuitively, the more diverse the population is, the greater this value is, as diversity is related to heterozygosity (see above).

3.7 Typical Paternity Index (TPI)

Finally, the typical paternity index (TPI) reflects the "mean PI for random non-excluded men" in trio cases (father-mother-child) for a given locus.

Formula

Let H be the frequency of homozygotes, then

$$TPI = \frac{1}{2H}$$

Interpretation

Unlike the other parameters, the typical paternity index values are not in the 0-1 range and cannot be interpreted as a probability. It is a ratio of 1 divided by twice the frequency of homozygotes in the population. This is an **odds ratio**, measuring how many times more likely that a possible father is the actual father than a randomly selected man in the population, on average (*typical* case).

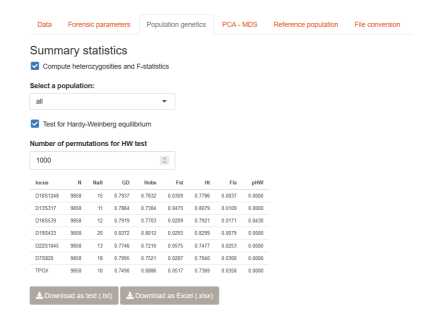
Chapter 4

Population genetics indices

In this chapter, we will see how to compute some population genetics indices in STRAF

4.1 Computing population genetics parameters in STRAF

Once you have uploaded your genotypes in STRAF, you can go to the **Population genetics** tab to compute relevant population genetics indices. It is also possible to perform a Hardy-Weinberg equilibrium test by checking the relevant box.



4.2 Details on population genetics indices

4.2.1 Hardy-Weinberg equilibrium

A population is considered at Hardy-Weinberg equilibrium (HWE) when the observed genotypic frequencies are in agreement with the expectations in an "ideal" population, which assumes for example random mating in the population. This is important as the assumptions of the Hardy-Weinberg model allow to derive quantities such as forensic parameters and population genetic indices. Therefore, if some assumptions of the model are violated, conclusions drawn from metrics computed assuming HWE could be challenged.

If a locus presents a significant deviation from HWE, it means that a process is influencing the distribution of allele and genotype frequencies in the population. It could for example be due to **inbreeding**, **hidden population structure**, or **natural selec-**

tion.

STRAF reports the **p-value** of a test for HWE. A low p-value indicates a significant deviation from HWE.

4.2.2 Heterozygosities

In STRAF, several measures of **heterozygosity** are computed. They capture different aspects of genetic diversity.

- The expected heterozygosity (or Gene diversity, ${\cal H}_{exp}$ or GD) has been defined in the previous chapter.
- The observed heterozygosity (H_{obs}) is the proportion of heterozygous genotypes at this locus in the population.
- The **total heterozygosity** (H_T) is the heterozygosity expected if all the individuals in all the subpopulations were behaving as a population at HWE.

4.2.3 F-statistics

STRAF reports two F-statistics.

- The $F_{\rm IS}$ is a measure of genetic relatedness within a population. It is sometimes called the **inbreeding coefficient**. High values indicate a high degree of inbreeding.
- The $F_{\rm ST}$ is a measure of genetic differentiation between populations. It takes values between 0 (no differentiation) and 1 (full differentiation).

One concept, multiple estimators.

Several **estimators** of $F_{\rm ST}$ exist (for example, Weir and Cockerham's, Nei's, Hudson's $F_{\rm ST}$). It's like if each population geneticist had decided to develop their own estimator!

Why is that? In statistics, what we call an **estimator** is a metric aiming at estimating a given quantity based on **observed data**. It is important to keep in mind that these estimators rely on a specific **model**, with underlying assumptions. It explains why some estimators are more or less reliable depending on the case and observed data, and each of them has been developed for a different situation. In the case of $F_{\rm ST}$ for example, different estimators assume different demographic models.

4.3 Linkage disequilibrium (LD)

4.3.1 What is linkage disequilibrium?

Linkage disequilibrium is an important quantity to be measured in genetics. It is defined as the **nonindependence of genotypes at distinct loci**. It means that it is more likely to observe the co-occurrence of some genotypes at different loci. This can be influenced by population history.

However, most of the times, LD is explained by the physical proximity between loci. If two loci are next to each other on the genome, recombination events between them will be rare and genotypes won't be shuffled. Genotypes at these loci will be correlated and linkage disequilibrium will be high. On the other hand, two loci found on two different chromosomes are not expected to show any LD signals as genotypes will be systematically shuffled at each generation.

4.3.2 How to compute LD in STRAF?

It is possible to test for the presence of LD in the dataset you uploaded using STRAF. After checking the *Display pairwise LD p-values matrix*, LD tests between each pair of loci will be performed and p-values will be reported.

31

Linkage disequilibrium

✓ Display pairwise LD p-values matrix

	D10S1248	D13S317	D16S539	D19S433	D22S1045	D7S820	TPOX
TPOX	0.0000	0.0000	0.0001	0.0000	0.0000	0.0000	
D7S820	0.4549	0.0000	0.0506	0.0000	0.0000		
D22S1045	0.0000	0.0000	0.0000	0.0000			
D19S433	0.0000	0.0000	0.0000				
D16S539	0.2147	0.0000					
D13S317	0.0000						
D10S1248							



Important note: Other population genetics software, **Genepop** and **Arlequin**, implement more reliable versions of the LD test that should be preferred. They are currently not implemented in STRAF because of performance limitations. If you need to perform such a test, the **File conversion** utilities (cf. Chapter 6) should facilitate the workflow

Chapter 5

Multivariate statistics

In this chapter, we will describe the two multivariate statistics methods implemented in STRAF, **Principal Component Analysis (PCA)** and **Multidimensional Scaling (MDS)**.

5.1 Principal Component Analysis (PCA)

PCA is a method of **dimensionality reduction**. What is does it that it tries to **capture most of the variation** present in the data and **project** it onto a small number of new axes called **principal components** (PCs).

This is a useful method to capture variation from a large number of variables and allows to discover hidden patterns by increasing interpretability.

In our case, if we consider that **each allele at each locus** is a variable, and that our individual observations are the presence / absence of each allele for each sample, we end up with a highly dimensional dataset (we have as many variables as we have alleles!). It gets even worse if you analyse genome sequences, where you can have millions of variables in your dataset. This is definitely not an interpretable dataset if you are not able to easily extract relevant information.

PCA allows to bring most of the variation existing between our samples onto a few axes capturing most of the variation (PCs).

In STRAF, you can perform a PCA by going into the **PCA - MDS** tab and checking the following *Run and plot a PCA* checkbox.

Principal Component Analysis (PCA)

Run and plot a PCA (Principal Component Analysis) PCA axis 1 2 3 PCA projection PCA projection PCA projection PCA projection PCA projection PCA projection

It will trigger the PCA computation and return a graph. You are also able to download the coordinates of the samples on different PCs (also called eigenvectors).

PC1 (2.39 %)

Interpreting PCA results

PCA has become a popular tool in population genetics, as the relationships between individuals on the PCA projection tend to reflect their genetic **relatedness**. The closer individuals are on the PCA projection, the more genetically related they tend to be.

It can also be used as a **quality control** tool. For example, if a sample is very far from all the others, including the ones that are part of the same population, it is possible that there is an issue with the data. One would need to inspect the raw data and check if any strange pattern can be found. It is important to be aware of the influence of non-demographic processes on the PCA projection. For example, **imbalanced sample sizes** between populations can drive some patterns. When populations are sampled unevenly, the projection will be **distorted** and distances observed on the projection can be driven by these differences and not by their evolutionary history.

Hence, as multiple processes influence the results, PCA should remain an **exploratory approach** and further analyses should be conducted before drawing any major conclusions on the relationships between populations and individuals.

5.2 Multidimensional Scaling (MDS)

An **MDS** is conceptually similar to a PCA. One of the main differences is that it requires input data to be in a different format. As PCA uses raw genotypes and can accommodate data at the individual level, **pairwise distances** between data points are used for an MDS.

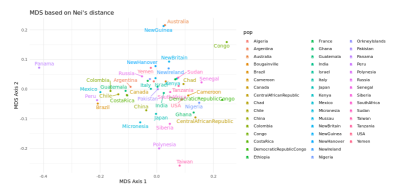
In forensics, it is typically used to compare **populations** and not **individuals**, even though it would be theoretically possible.

As **pairwise distances between populations** are used, and MDS can be based on any genetic distance: one could for example compute pairwise **FSTs**, **Nei's genetic distance**, or any other metric.

Based on these distances, the MDS will **project** the populations onto a **lower-dimensional space**. This projection facilitates the interpretation of relationships between populations.

Multidimensional Scaling (MDS) based on Nei's distance

Compute Nei's genetic distance between populations and run MDS



Interpreting MDS results

Just like with a PCA, the **closer** populations are on the MDS projection, the more **genetically related** they tend to be, based on the markers that have been used to compute their pairwise genetic distances.

Again, like for the PCA, the MDS should remain an **exploratory approach** and further analyses should be conducted before drawing any major conclusions on the relationships between populations and individuals.

Reference populations analysis

In this chapter, we will see how to compare the uploaded populations to reference populations (based on allele frequencies).

6.1 MDS on reference frequencies

STRAF implements an MDS computation based on allele frequency data. By default, allele frequencies from the STRiDER database [Bodner et al., 2016] are used (loci with less than 10 populations have been excluded). If you are not familiar with the MDS method and interpretation, you can find details in **Chapter 4**.

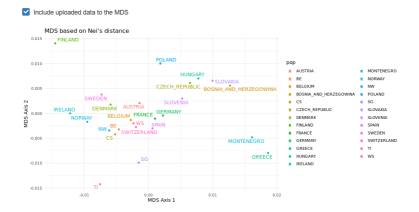
You can see the output of the MDS in the **Reference population** tab. It includes the MDS projection in two dimensions for 19 European populations, as well as the population tree.

Custom allele frequency database Import allele frequencies (if empty, the STRidER database will be used.) No file selected Include uploaded data to the MDS MDS based on Nei's distance pop AUSTRIA 8 BELGIUM BOSNIA AND HERZEGOWINA CZECH_REPUBLIC n DENMARK 8 FINLAND GREECE • FRANCE GERMANY MDS Axis 2 GREECE SWITZERLAND AUSTRIA SPAIN CZECH_REPUBLIC HUNGARY GERMANY IRELAND MONTENEGRO NORWAY IRELAND HUNGARY POLAND BOSNIA_AND_HERZEGOWINA FRANCE 8 SLOVENIA 8 SPAIN SWEDEN 8 SWITZERLAND -0.01 0.00 MDS Axis 1 0.01 -SWITZERLAND -GERMANY -BELGIUM -AUSTRIA -SPAIN -FRANCE -NORWAY -IRELAND -SWEDEN -DENMARK -POLAND -HUNGARY -BOSNIA AND HERZEGOWINA -FINLAND -SLOVENIA -CZECH REPUBLIC -SLOVAKIA -MONTENEGRO -GREECE

The drop-down menu below the plots allows one to select or unselect populations to be used in the MDS analysis.



Provided that some loci are in common between the reference allele frequencies and the genotypes you uploaded, it is possible to add your own populations to the existing MDS by checking the *Include uploaded data to the MDS* box.



You will then see your populations added to the MDS plots.

6.2 Preparing a custom allele frequency database

If you want to use other reference databases than STRIDER, for example if you're working with non-European samples or with different loci, it is possible to upload a custom allele frequency database in STRAF.

The data must be formatted as follows:

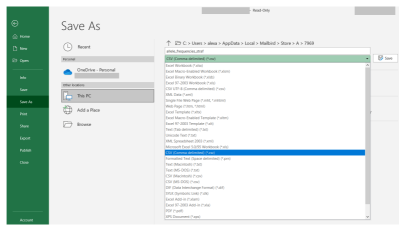
D1S1656			
Allele	Switzerland	Germany	France
9	0.12	0.09	0
10	0.40	0.35	0.28
10.2	0.31	0.41	0.5
11	0.17	0.15	0.22
D2S1338			
Allele	Switzerland	Germany	France
19	0.40	0.38	0.42
20	0.29	0.26	0.28
21	0.31	0.36	0.3

Starting from an Excel file for example, we can start with a spreadsheet that looks like this:

4	Α	В	С	D	Е
1	D8S1179				
2	Allele	Pop1	Pop2	Pop3	Pop4
3	8	0.013	0.014	0.004	0.0146
4	9	0.0095	0.017	0.005	0.0049
5	10	0.0969	0.017	0.061	0.0777
6	11	0.0674	0.062	0.081	0.1165
7	12	0.1129	0.167	0.121	0.1359
8	13	0.2907	0.315	0.292	0.2136
9	14	0.2123	0.245	0.26	0.1408
10	15	0.1394	0.092	0.144	0.2184
11	16	0.048	0.022	0.028	0.0583
12	17	0.0095	0.001	0.003	0.0194
13	18	0	0	0.001	0
14	19	0	0	0	0
15	D21S11				
16	Allele	Pop1	Pop2	Pop3	Pop4
17	12	0	0	0.129	0
18	13	0	0	0.024	0
19	19	0.001	0	0	0
20	24.2	0	0	0	0
21	24.3	0	0	0	0
22	25	0.002	0.001	0.001	0
23	25.2	0	0	0	0
24	25.3	0	0	0	0
25	26	0.0025	0.003	0.014	0
26	27	0.011	0.03	0.034	0.0248

Then, one simply needs to save this table as a CSV (Comma-Separated Values)

file. This can be achieved by clicking on Save As > CSV (Comma-delimited) (*.csv).



Then, you can upload your data and use it as described for the STRIDER database.

File conversion

As STRAF is a web application and can be used simultaneously by multiple users, computing resources are limited. Therefore, the most computationally intensive analyses are not available in STRAF. In order to ease the path to other software, file conversion utilities have been implemented. It is possible to convert the input file to the **Genepop**, **Arlequin** and **Familias** formats. They are all available in the **File conversion** tab of the application.

7.1 How to convert a file in STRAF

Once you have imported your file, it is straightforward to convert it to another format. You can go to the **File conversion** tab and click on one of the following buttons to download your genotypes in another format.



7.2 Genepop and Arlequin formats

Genepop [Rousset, 2008] and **Arlequin** [Excoffier and Lischer, 2010] softwares implement several population genetics methods, including ones that are part of standard forensics practice:

- · linkage disequilibrium computation
- · Hardy-Weinberg tests

STRAF currently implements these computations, however the ones implemented in Genepop are overall more reliable as they can rely on more permutations. They are overall preferable to the HW and LD tests implemented in STRAF.

7.3 Familias

Here a file containing allele frequencies is created. This file can be used in Familias [Kling et al., 2014] to provide allele frequency reference data for relationship testing.

Analysing SNP data

8.1 STRs vs. SNPs

STRs have been and remain the dominant marker used in forensic genetics.

This is mainly due to the fact that STRs have a high mutation rate, therefore are more diverse in human populations. This explains their high **power of discrimination**. Furthermore, they can also be used in deciphering mixture components, a very common case in forensics. Finally, they can be combined in multiplex assays, which is convenient when lowamounts of biological material can be recovered.

Even though STRs will still remain the standard markers in forensic genetics for many years, the application of sequencing data, especially the analyses of SNPs became more and more popular in recent years.

Indeed, it nowadays easier and cheaper to generate whole-genome sequences than before. One could wonder why STRs are still so popular in forensic practice, and have not been replaced by SNPs that are easily generated from Next-Generation Sequencing (NGS) data.

The advantages of SNPs are a higher multiplex capacity (with STRs and CE we can

analyse not much more than 25 markers in parallel) and therefore a higher discriminative power and shorter amplicon size (interesting for degraded samples).

8.2 How to analyse SNP data in STRAF?

It is in theory possible to use any marker type in STRAF, as long as the input format corresponds to what is expected from the software, that is:

- · 2 columns per locus for diploid data, 1 for haploid;
- alleles are encoded by a number: for example, a diallelic SNP could be encoded as 100 for the reference allele, and 101 for the alternative allele.

The main challenge remains encoding these values and converting from the initial data format. As an example, a genotype table is easier to convert than raw sequences or bioinformatics-specific formats such as the Variant Calling Format (VCF).

Conclusion

You have now reached the end of the STRAF Book. This book has been written to give a broad overview of the concepts and tools you can find in the application. We hope it has been useful to you.

If you want to know more about STRAF, you will find more information about the software in the **Appendix** section.

You can now go back to straf.fr and use the application for your forensics project! Any feedback is always welcome and you can contact us anytime. STRAF is continuously improving thanks to your precious feedback!

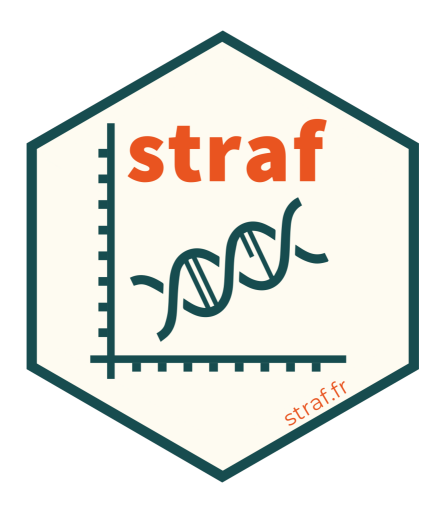


Figure 9.1: The STRAF logo.

References

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Appendix A

Technical details

A.1 Basic information

- STRAF is hosted on an Amazon Web Services (AWS) server located in Ireland.
- STRAF has **2 CPUs** and **4 gigabytes of memory** available on the server. It is enough for the vast majority of cases, but might not be enough if your dataset is too large (too many samples, or too many loci).
- The web application is based on the R-Shiny framework. R is an open source programming language particularly popular in data science. Shiny is an open source R package that provides a web framework for building web applications using R.

A.2 Understanding STRAF's performance

When you open STRAF, a new session. When you close your browser or the tab, the session is closed and your data and computations are deleted.



Figure A.1: The R logo.

STRAF can accept an unlimited number of user connections at the same time. However, the computational resources are shared between users. In other words, multiple users can connect at the same time, but computations will be ran one after the other. If you wait longer than usual for a given task, it is likely that someone else is using the server. Please keep this in mind and try to avoid unnecessary computations!

STRAF has limited resources, as described above, and this can be limiting if your dataset is too large because you have too many samples or loci. In that case, you could consider trying to run STRAF locally as outlined in **Chapter ??**. You could even host it on your own institution's server, in that case feel free to contact us as we would be happy to help in the process.

A.3 External dependencies

STRAF wouldn't exist without the amazing community of R developers who contributed great resources and packages.

The application uses several genetics packages:

- genepop
- · edegenet
- · hierfstat
- · pegas

More generic data science packages are also used:

- Shiny-related packages: shiny, colourpicker, shinycssloaders, shinyWidgets
- The "tidyverse": dplyr, tidyr, ggplot2, magrittr, openxlsx, reshape2, ggrepe1

Appendix B

STRAF's commitments

B.1 STRAF will always remain free and open source

STRAF is a **free** and **open-source** software. As free software advocates, we believe anyone should be able to **run** the software, **study** the software, **modify** the software, and to **share** copies of the software. That is why STRAF is licensed under the permissive **GNU General Public License version 3**. Here is the full text of the GPL-3 license

For the most curious of you, you can view all the source code of STRAF on GitHub.

B.2 STRAF is carbon-neutral

Hosting a web application onto a server requires energy, and energy is becoming more and more precious. For that reason, STRAF is hosted on a server that is on a carbon-neutral data center and that will be fully powered by renewable energies by 2025.

This book and STRAF's code source are hosted onto GitHub servers, which are carbon-neutral as well since 2019.

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