Bioinformatics and Statistical Genetics	Descriptive analysis of genetic markers
Iván Galván-Femenía-Jan Graffelman	November 2, 2016

,	7.	(1p) Compute the minor allele frequencies for all markers, and make a histogram of it.
8	3.	(1p) What percentage of the markers have a maf below 0.05? And below 0.01?
(9.	(2p) Compute for each marker its expected heterozygosity , where the expected heterozygosity for a bi-allelic markers is defined as $1 - \sum_{i=1}^{k} p_i^2$, where p_i is the frequency of the <i>i</i> th allele. Compute the average expected heterozygosity over all markers. Make a histogram of the expected heterozygosity.
2		STR dataset
-	1.	The file FrenchStrs.dat contains genotype information (STRs) of individuals from a French population. The first column of the data set contains an identifier the individual. STR data starts at the second column. Load this data into the R environment.
	2.	(1p) How many individuals and how many STRs contains the database?
;	3.	(1p) The value −9 indicates a missing value. Replace all missing values by NA. What percentage of the total amount of datavalues is missing?
2	1.	(2p) Write a function that determines the number of alleles for a STR. Determine the number of alleles for each STR in the database. Compute basic descriptive statistics of the number of alleles (mean, standard deviation, median, minimum, maximum).
į	5.	(2p) Make a boxplot and a histogram of the number of alleles per STR. What is the most common number of alleles for an STR?
(3.	(2p) Compute the expected heterozygosity for each STR. Make a histogram of the expected heterozygosity over all STRS. Compute the average expected heterozygosity over all STRs.

7.	(2p) Compare the results you obtained for the SNP database with those you obtained for the STR
	database. What differences do you observe between these two types of genetic markers?