Practical 03 SG: LD and Haplotype estimation

Kathryn Weissman & Irene Fernández

2022-11-24

LD (15p.)

- 1. The file FOXP2.zip contains genetic information of individuals of a Japanese population of unrelated individuals. The genotype information concerns SNPs of the Forkhead box protein P2 (FOXP2) gene region, located the long arm of chromosome number 7. This gene plays an important role in the development of speech and language. The FOXP2.zip file contains:
- FOXP2.dat: a text file with the genotype data which can be read in with R.
- FOXP2.fam: a PLINK file with data on the individuals (family id, individual id, ids of parents, sex and phenotype).
- FOXP2.bed: a PLINK file with binary genotype data.
- FOXP2.bim: a PLINK file with data on the genetic variants (chromosome, SNP identifier, basepair position along the chromosome and alleles).
- 2. (1p) Load the **FOXP2.dat** file into the R environment. How many individuals and how many SNPs are there in the database? What percentage of the data is missing?
- There are 104 individuals in the database.
- There are 544 SNPs in the database.
- 0% of the data is missing.

```
file <- "~/Downloads/FOXP2/FOXP2.dat"
data <- fread(file, header = TRUE)
data[1:10, 1:10]</pre>
```

##		id	rs34684677	rs1839115	rs4727804	rs4727805	rs200888633	rs12534908
##	1:	NA18939	T/G	C/T	G/A	T/G	T/G	G/A
##	2:	NA18940	G/G	T/T	A/A	G/G	T/G	A/A
##	3:	NA18941	G/G	T/T	A/A	G/G	T/G	A/A
##	4:	NA18942	G/G	T/T	A/A	G/G	T/T	A/A
##	5:	NA18943	G/G	T/T	A/A	G/G	T/T	A/A
##	6:	NA18944	T/T	C/C	G/G	T/G	G/G	G/G
##	7:	NA18945	G/G	T/T	A/A	G/G	G/G	A/A
##	8:	NA18946	T/G	C/T	G/A	G/G	G/G	G/A
##	9:	NA18947	T/G	C/T	G/A	G/G	T/G	G/A
##	10:	NA18948	G/G	T/T	A/A	G/G	G/G	A/A
##		rs125330)49 rs778613	356 rs6945	561			
##	1:	(C/T T	T/T (C/T			
##	2:	7	T/T T	T/T 1	Γ/Τ			

```
3:
                T/T
                              T/T
                                          T/T
##
                                          T/T
##
    4:
                T/T
                              T/T
##
    5:
                T/T
                              T/T
                                          T/T
    6:
                C/C
                              T/T
                                          C/C
##
##
    7:
                T/T
                              T/T
                                          T/T
    8:
                C/T
                              T/T
                                          C/T
##
##
    9:
                C/T
                              T/T
                                          C/T
## 10:
                T/T
                              A/T
                                          T/T
```

```
n <- nrow(data); n # number of samples
```

[1] 104

```
p <- ncol(data); p # number of variables</pre>
```

[1] 544

```
perc.mis <- 100*sum(is.na(data))/(n*p); perc.mis # percentage of missing data overall
```

[1] 0

- 3. (1p) Determine the genotype counts for each SNP, and depict all SNPs simultaeneously in a ternary plot, and comment on your result. For how many variants do you reject Hardy-Weinberg equilibrium using an ordinary chi-square test without continuity correction? (hint: you can read the .bim in R in order to determine the alleles of each SNP, and use function MakeCounts from the HardyWeinberg package to create a matrix of genotype counts).
- 4. (1p) Using the function LD from the genetics package, compute the LD statistic D for the SNPs rs34684677 and rs2894715 of the database. Is there significant association between the alleles of these two SNPs?
- 5. (2p) Also compute the LD statistic D for the SNPs rs34684677 and rs998302 of the database. Is there significant association between these two SNPs? Is there any reason why rs998302 could have stronger or weaker correlation than rs2894715?
- 6. (2p) Given your previous estimate of D for SNPs rs34684677 and rs2894715, infer the haplotype frequencies. Which haplotype is the most common?
- 7. (2p) Compute the LD statistics R^2 for all the marker pairs in this data base, using the LD function of the packages genetics. Be prepared that this make take a few minutes. Also compute an alternative estimate of R^2 obtained by using the PLINK program. For this purpose you should:
- Download and install PLINK 1.90 from https://www.cog-genomics.org/plink2/
- Take care to store the files FOXP2.bim, FOXP2.fam and FOXP2.bed in a directory where PLINK can find them.
- Compute LD estimates with PLINK using plink --bfile FOXP2 --r2 --matrix --out FOXP2 This creates a file with extension FOXP2.ld that contains a matrix with all R^2 statistics. Read this file into the R environment. Make a scatter plot for R's LD estimates against PLINK's LD estimates. Are they identical or do they at least correlate? What's the difference between these two estimators? Which estimator would your prefer and why?
- 8. (2p) Compute a distance matrix with the distance in base pairs between all possible pairs of SNPs, using the basepair position of each SNP given in the .bim file. Make a plot of R's R^2 statistics against the distance (expressed as the number of basepairs) between the markers. Comment on your results.

- 9. (2p) Make an LD heatmap of the markers in this database, using the R^2 statistic with the LD function. Make another heatmap obtained by filtering out all variants with a MAF below 0.35, and redoing the computations to obtain the R^2 statistics in R. Can you explain any differences observed between the two heatmaps?
- 10. (1p) Can you distinguish blocks of correlated markers in the area of the FOXP2 gene? How many blocks do you think that at least seem to exist?
- 11. (1p) Simulate independent SNPs under the assumption of Hardy-Weinberg equilibrium, using R's sample instruction sample(c("AA", "AB", "BB"),n,replace=TRUE,prob=c(p*p,2*p*q,q*q))). Simulate as many SNPs as you have in your database, and take care to match each SNP in your database with a simulated SNP that has the same sample size and allele frequency. Make an LD heatmap of the simulated SNPs, using R^2 as your statistic. Compare the results with the LD heatmap of the FOXP2 region. What do you observe? State your conclusions.

Haplotype estimation (10p.)

- 1. Apolipoprotein E (APOE) is a protein involved in Alzheimer's disease. The corresponding gene APOE has been mapped to chromosome 19. The file APOE.dat contains genotype information of unrelated individuals for a set of SNPs in this gene. Load this data into the R environment. APOE.zip contains the corresponding .bim, .fam and .bed files. You can use the .bim file to obtain information about the alleles of each polymorphism.
- 2. (1p) How many individuals and how many SNPs are there in the database? What percentage of the data is missing?
- 3. (1p) Assuming all SNPs are bi-allelic, how many haplotypes can theoretically be found for this data set?
- 4. (2p) Estimate haplotype frequencies using the haplo.stats package (set the minimum posterior probability to 0.001). How many haplotypes do you find? List the estimated probabilities in decreasing order. Which haplotype number is the most common?
- 5. (2p) Is the haplotypic constitution of any of the individuals in the database ambiguous or uncertain? For how many? What is the most likely haplotypic constitution of individual NA20763? (identify the constitution by the corresponding haplotype numbers).
- 6. (1p) Suppose we would delete polymorphism rs374311741 from the database prior to haplotype estimation. Would this affect the results obtained? Justify your answer.
- 7. (1p) Remove all genetic variants that have a minor allele frequency below 0.10 from the database, and re-run haplo.em. How does this affect the number of haplotypes?
- 8. (2p) We could consider the newly created haplotypes in our last run of haplo.em as the alleles of a new superlocus. Which is, under the assumption of Hardy-Weinberg equilibrium, the most likely genotype at this new locus? What is the probability of this genotype? Which genotype is the second most likely, and what is its probability?