

# GenABEL tutorial

Yurii Aulchenko

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## 1 Overview

**GenABEL** is an R library developed to facilitate Genome-Wide Association (GWA) analysis of binary and quantitative traits. **GenABEL** is implemented as an R library. R is a free, open source language and environment for general-purpose

statistical analysis (available at <http://www.r-project.org/>). It implements powerful data management and analysis tools. Though it is not strictly necessary to learn everything about R to run GenABEL, it is highly recommended as this knowledge will improve flexibility and quality of your analysis.

This tutorial was originally written to serve as a set of exercises for the "Advances in population-based studies of complex genetic disorders" (GE03) course of the Netherlands Institute of Health Sciences (Nihes).

If you read this tutorial not as a part of the GE03 course, and you are eager to start with your GWA analysis without reading all the not-so-strictly-necessary stuff, start directly from the section 5 ("Genome-wide association analysis").

Otherwise, you can start with R basics and simple association analyses using few SNPs in section 2, "Introduction to R". In the next section, 4 ("Introduction to GenABEL") you will learn how to work with the `gwaa.data-class`, which is used to store GWA data in GenABEL and will perform some simple large-scale analyses.

In the next section, 5 ("Genome-wide association analysis"), you will do quality control of genetic data and do association analysis under realistic conditions. This section is the core of this tutorial.

The section 6 ("GWA in presence of genetic stratification") is dedicated to analysis in presence of populational stratification and analysis of family-based data.

Genetic data imputations are covered in section 8, "Genetic data imputations".

The last section, 10 ("Analysis of selected region"), is dedicated to analysis of haplotype association and analysis of SNP interactions.

Information on importing the data from different formats to GenABEL is given in appendix A. Answers to exercises are provided in appendix B.

Experienced R users start directly with the section (4, "Introduction to GenABEL").

## 2 Introduction to R

In this section we will consider basic R data types and operations, tools for analysis of qualitative and quantitative traits. Needless to say, that only basic R functionality may be covered within such short section. If you decide to improve your knowledge of R, we recommend you excellent manuals, such as 'An introduction to R', 'Simple R', 'Practical Regression and Anova using R', and others, available free of charge from the R project web-site (<http://www.r-project.org>).

In the first parts of this section, we will cover basic types of R objects and will work with them. Next, we will analyse qualitative and quantitative traits, using only basic R functionality.

### 2.1 Basic R data types and operations

On the contrast to many other statistical analysis package, analysis in R is not based on graphic user interface, but is rather command-based. When you first start R, a command prompt appears. To get help and overview of R, type `help.start()` in the command line and press `enter`. This will start your internet browser and open R documentation.

Let us first exploit R as a calculator. You can directly operate with numbers in R. Try adding three to two:

```
> 2 + 3
```

```
[1] 5
```

Square roots, base-10 logarithm, and exponentiation can be done straightforwardly with

```
> sqrt(5)
```

```
[1] 2.236068
```

```
> log10(2.24)
```

```
[1] 0.350248
```

```
> exp(0.35)
```

```
[1] 1.419068
```

The arithmetic operations and functions can be nested:

```
> exp(log10(sqrt(2 + 3)))
```

```
[1] 1.418337
```

R functions include standard ones, such as logarithms, power, but also a wide range of statistical function, for example, distribution and probability density of many distributions.

For any function with name say 'fun', help may be obtained by typing '`help(fun)`' on the command line. If you do not know the exact name for the function you look for, try '`help.search("query")`', where `query` is the keyword.

**Exercise 1** Try to find out what are the functions to do

1. Wilcoxon test
2. Fisher exact test
3. T-test

R help pages have standard layout, documenting usage of the function, explaining function arguments, providing details of implementation, explaining the value returned by the function, and giving references and examples of function usage.

**Exercise 2** Explore the help page for the Wilcoxon test and answer the questions:

1. When exact Wilcoxon test is computed by default?
2. If the default conditions for the exact test are not satisfied, what approximation is used?

Most of the documented functions will have examples of their usage at the end of the 'help' page, and these examples can be evaluated in R. Try '`example(wilcox.test)`'.

One of important R operations is *assignment*, which is done with '`<-`' operator. For example, we if want to assign value '2' to variable 'a', and value '3' to the variable 'b':

```
> a <- 2  
> b <- 3
```

Typing the variable name in R command line will return its' value, e.g.

```
> b
```

```
[1] 3
```

Naturally, evaluation of

```
> exp(log10(sqrt(a + b)))  
[1] 1.418337
```

gives the expected result we have obtained above using numerical arguments.

Just introduced variables 'a' and 'b' contain single values. More practically important are the variables containg *vectors*. Let us create an example vector and experiment with it:

```
> v <- c(1, 3, 5, 7, 11)
```

Now, let us try different operations with this vector:

```
> v + 1  
[1] 2 4 6 8 12  
  
> 1/v  
[1] 1.0000000 0.3333333 0.2000000 0.1428571 0.0909091  
  
> log(v)  
[1] 0.000000 1.098612 1.609438 1.945910 2.397895
```

It is easy to see that these operations give a vector with elements being the result of application of the same operation to different initial elements.

What happens if two vectors are supplied as function arguments? Let us define new vector

```
> ov <- c(1, 2, 3, 4, 5)
```

and add it and previous vector up:

```
> v + ov
```

```
[1] 2 5 8 11 16
```

multiply them

```
> v * ov
```

```
[1] 1 6 15 28 55
```

or compute v to the power of ov:

```
> v^ov
```

```
[1] 1 9 125 2401 161051
```

Here, the operation was done over each of the two corresponding elements of the vectors, resulting, again, in the vector of the same length.

Other functions may evaluate a vector as a whole and return a single value as output. For example, to obtain a sum of vector's elements, use

```
> sum(v)
```

```
[1] 27
```

Other examples of such functions involve `length`, returning number of elements of a vector, `mean`, returning the mean, `var`, returning the variance, etc. For example:

```
> length(v)
```

```
[1] 5
```

```
> mean(v)
```

```
[1] 5.4
```

```
> var(v)
```

```
[1] 14.8
```

One of the most important data operations in R is *sub-setting*. This refers to operations which help you derive a subset of the data. Let us create a short vector and play a bit with sub-setting. This vector will contain 5 simple character strings:

```

> a <- c("I am element 1", "I am element 2", "I am element 3",
+      "I am element 4", "I am element 5")
> a

[1] "I am element 1" "I am element 2" "I am element 3" "I am element 4"
[5] "I am element 5"

```

To find out what is the value of the  $i$ -th element of this vector, you can sub-set it by `a[i]`. For example the 3rd elements is:

```

> a[3]

[1] "I am element 3"

```

You can also select bigger sub-sets, e.g. all elements from 2 to 4:

```

> a[c(2:4)]

[1] "I am element 2" "I am element 3" "I am element 4"

```

Here, operation `c(2:4)` is equivalent to 'sequence from 2 to 4', that is `c(2,3,4)`.

We can easily get disjoint elements; e.g. if you want to retrieve elements 1, 3, and 5, you can do

```

> dje <- c(1, 3, 5)
> dje

[1] 1 3 5

> a[dje]

[1] "I am element 1" "I am element 3" "I am element 5"

```

One of very attractive features of R data objects is possibility to derive sub-sets based on some condition. Let us consider two vectors, `tmpheight`, containing the height of some subjects, and `tmpids`, containing their IDs:

```

> tmpheight <- c(150, 175, 182, 173, 192, 168)
> tmpheight

[1] 150 175 182 173 192 168

> tmpids <- c("fem1", "fem2", "man1", "fem3", "man2", "man3")
> tmpids

[1] "fem1" "fem2" "man1" "fem3" "man2" "man3"

```

Imagine you want to derive the IDs of people with height over 170 cm. The way to do it is to combine several steps. First, you can run the logical function `>170` on the height data:

```

> vec <- (tmpheight > 170)
> vec

[1] FALSE TRUE TRUE TRUE TRUE FALSE

```

This returns logical vector whose elements are true, when particular element of `tmpht` satisfies the condition `>170`. Such logical vector, in turn, may be applied to sub-set any other vector of the same length<sup>1</sup>, including itself. If you want to see what are the heights in people taller than 170 cm, you can use

```
> tmpht[vec]
```

```
[1] 175 182 173 192
```

or you can get IDs of these people with

```
> tmpids[vec]
```

```
[1] "fem2" "man1" "fem3" "man2"
```

You can combine more than one logical condition to derive sub-sets. For example, to see what are the IDs of people taller than 170 but shorter than 190 cm, you can use

```
> vec <- (tmpht > 170 & tmpht < 190)
> vec
```

```
[1] FALSE TRUE TRUE TRUE FALSE FALSE
```

```
> tmpids[vec]
```

```
[1] "fem2" "man1" "fem3"
```

Other, and easier<sup>2</sup> way to do the same is to use `which()` function. This function reports which elements pass logical condition. To obtain above results you can run:

```
> vec <- which(tmpht > 170 & tmpht < 190)
> vec
```

```
[1] 2 3 4
```

```
> tmpids[vec]
```

```
[1] "fem2" "man1" "fem3"
```

You can remove `tmpht` and `tmpids` variable so they will not interfere with our future analysis:

```
> ls()
```

```
[1] "a"      "b"      "dje"     "ov"      "tmpht"   "tmpids"  "v"      "vec"
```

```
> rm(tmpht, tmpids)
```

and check if the removal was successful:

---

<sup>1</sup>Actually, you can apply it to a longer vector too, and then the logical vector will be "expanded" to total length by repeating the original vector head-to-tail. However, we will not use this in our exercises.

<sup>2</sup>Because it treats NAs for you

```
> ls()
[1] "a"    "b"    "dje"  "ov"   "v"    "vec"
```

Sub-setting for 2D objects (matrices) is done in similar manner. Let us construct a simple matrix and do several sub-setting operations on it:

```
> a <- matrix(c(11, 12, 13, 21, 22, 23, 31, 32, 33), nrow = 3,
+               ncol = 3)
> a
[,1] [,2] [,3]
[1,]   11   21   31
[2,]   12   22   32
[3,]   13   23   33
```

To obtain the element in the 2nd row and 2nd column

```
> a[2, 2]
[1] 22
```

...the second row and third column:

```
> a[2, 3]
[1] 32
```

To obtain the 2x2 set of elements contained in upper left corner, you can do

```
> a[1:2, 1:2]
```

```
[,1] [,2]
[1,]   11   21
[2,]   12   22
```

Or you can even get the variables, which reside in corners:

```
> a[c(1, 3), c(1, 3)]
[,1] [,2]
[1,]   11   31
[2,]   13   33
```

If one of the dimensions is not specified, complete list is returned for this dimension. For example, here we retrieve the first row

```
> a[1, ]
[1] 11 21 31
...and third column
> a[, 3]
[1] 31 32 33
```

...or columns 1 and 3:

```
> a[, c(1, 3)]  
 [,1] [,2]  
[1,] 11 31  
[2,] 12 32  
[3,] 13 33
```

As well as with vectors, you can sub-set matrices using logical conditions or indexes. GWA genetic data are stored in matrices, and you can sub-set them using the methods described above.

For example, if we want to see what elements of a are greater than 21, we can run

```
> a > 21  
 [,1] [,2] [,3]  
[1,] FALSE FALSE TRUE  
[2,] FALSE TRUE TRUE  
[3,] FALSE TRUE TRUE
```

or obtain these elements by

```
> a[a > 21]  
[1] 22 23 31 32 33
```

At this point, you can exit R.

### Summary:

- You can get a top-level access to R documentation by `help.start()`. To search help for some keyword `key`, try `help.search(key)`. To get description of some function `fun`, try `help(fun)`.
- You can use R as a powerful calculator.
- It is possible to get sub-sets of vectors and matrices by specifying index value or a logical condition (of the same length as the vector / matrix) between square brackets (`[, ]`)
- When you obtain an element of a matrix with `[i, j]`, `i` is the row and `j` is the column of the matrix.
- Function `which(A)` returns index of the elements of A which are "true".

**Exercise 3** In this exercise, you will explore few vectors representing different data on study subjects described in `srdta` example data set supplied together with `GenABEL`. First, you need to load `GenABEL` by typing

```
> library(GenABEL)
```

and load the data by

```
> data(srdta)
```

The vector containing study subjects sex can be accessed through `srdta@gtdata@male`; this vector's value is one when the corresponding person is male and zero otherwise. The vector containing SNP names can be accessed via `srdta@gtdata@snpnames`, chromosome ID – through `srdta@gtdata@chromosome` and map – through `srdta@gtdata@map`. Explore these vectors and answer the questions.

1. What is the ID and sex of the first person in the data set?
2. Of the 22nd person?
3. How many males are observed among first hundred subjects?
4. How many FEMALES are among 4th hundred?
5. What is the male proportion in first 1000 people?
6. What is the FEMALE proportion in second 1000 (1001:2000) people?
7. What is name, chromosome and map position of 33rd marker?
8. What is distance between markers 25 and 26?

## 2.2 Data frames

Start R with double-click on the file named `assocbase.RData`. You can see the names of the loaded objects by using the command `ls()`:

```
> ls()  
[1] "assoc"
```

You can see that there is a single object the class of the object can be interrogated by using `class` function:

```
> class(assoc)  
[1] "data.frame"
```

Thus, the file you have loaded contains one *data frame*. A data frame is an R term for a data table. In such tables, it is usually assumed that rows correspond to subjects (observations) and columns correspond to variables.

We will investigate the data presented in the `assoc` data frame. The nice feature of data frames is that columns carry names for the variables, and the data stored there can be retrieved by referencing these names.<sup>3</sup> To see what are the variable names, use the command `names()`:

```
> names(assoc)
```

```
[1] "subj" "sex"   "aff"    "qt"     "snp4"  "snp5"  "snp6"
```

The 7 variables correspond to the personal ID, sex, affection status, quantitative trait *qt* and several SNPs.

A variable from a data frame `frame`, which has some name `name` can be accessed through `frame$name`. This will return a conventional vector. For example to see the affection status (`aff`) in the data frame `assoc`, use

> assoc\$aff

**Exercise 4** Investigate types of the variables presented in data frame `assoc`. For each variable, write down the class.

We can easily check how many people are described in the data set by checking the length of any variable (all variables will be the same length in a data frame):

```
> length(assoc$sex)
```

[1] 250

or through asking to report the dimensionality of the data frame:

> `dim(assoc)`

[1] 250 7

Here, the first number corresponds to the number of rows (subjects) and the second to the number of columns (variables).

Let us go into more details and check what are the ID, sex and affection status for the person 75:

---

<sup>3</sup>This may also be true for matrices; more fundamental difference is though that a matrix always contains variables of the same type, e.g. character or integer, while a data frame may contain variables of different types.

```
> assoc$subj[75]
```

```
[1] 1409
```

```
> assoc$sex[75]
```

```
[1] 1
```

```
> assoc$aff[75]
```

```
[1] 0
```

Data frame may be thought of as a matrix, or as a collection of vectors. All sub-setting operations discussed before for vectors/matrices are applicable to a data frame. Therefore an alternative way to see all the data for person 75 is:

```
> assoc[75, ]
```

subj	sex	aff	qt	snp4	snp5	snp6
1409	1409	1	0	1.014664	A/B	B/A

In the same manner, you can get data for e.g. subjects 5 to 15 (a new data frame containing data only on first 10 people) by

```
> assoc[5:15, ]
```

subj	sex	aff	qt	snp4	snp5	snp6
1533	1533	0	0	0.1009220	A/B	B/A
2466	2466	1	0	-0.1724321	A/B	A/A
2425	2425	0	0	-0.3378473	B/B	A/A
1068	1068	0	0	-1.7112925	A/A	B/B
198	198	1	0	-0.4815822	A/B	B/A
1496	1496	1	0	1.2281232	A/A	B/B
909	909	0	0	0.5993945	A/B	B/A
1213	1213	0	0	1.9792190	A/A	B/B
181	181	1	0	1.5435921	A/A	B/B
1783	1783	0	0	-1.6242738	A/B	B/A
1914	1914	0	0	-0.5160331	A/A	B/B

As well as with vectors, it is possible to sub-set elements of a data frame based on (a combination of) logical conditions. For example, if we want to check what are the IDs of the people with qt over 1.4, we can run

```
> vec <- which(assoc$qt > 1.4)
> vec
```

```
[1] 12 13 33 41 54 68 72 76 89 106 118 142 156 161 175 181 193 219 241
```

```
> assoc$subj[vec]
```

```
[1] 1213 181 1737 1319 516 1355 186 1426 1284 822 2129 212 1443 704 1648
[16] 1628 562 858 698
```

Or, if we are interested to see subject-number and SNP information for these people, we can see that with

```

> assoc[vec, c(1, 5, 6, 7)]

  subj .snp4.snp5.snp6
1213 1213   A/A   B/B   B/B
181   181   A/A   B/B   B/B
1737 1737   A/A   B/B   B/B
1319 1319   A/A   B/A   B/A
516   516   A/B   B/A   B/A
1355 1355   A/A   B/B   B/B
186   186   A/A   B/A   B/A
1426 1426   A/B   B/A   B/A
1284 1284   A/A   B/B   B/B
822   822   A/B   B/A   B/A
2129 2129   A/B   B/A   B/A
212   212   A/B   B/A   B/A
1443 1443   A/A   B/B   B/B
704   704   A/B   B/A   B/A
1648 1648   A/B   B/A   B/A
1628 1628   A/B   B/A   B/A
562   562   A/A   B/B   B/B
858   858   A/B   B/A   B/A
698   698   B/B   A/A   A/A

```

here, we select people identified by `vec` in the first dimension (subjects), and by `c(1,5,6,7)` we select first, fifth, sixth and seventh column (variable).

Next, we can check how many of the subjects are males by

```
> sum(assoc$sex)
```

```
[1] 129
```

and what is male sex proportion:

```
> sum(assoc$sex)/length(assoc$sex)
```

```
[1] 0.516
```

or

```
> mean(assoc$sex)
```

```
[1] 0.516
```

or

```
> sum(assoc$sex == 1)/length(assoc$sex)
```

```
[1] 0.516
```

You can also explore and modify the raw data contained in a data frame by using `fix()` command (e.g. `fix(assoc)`). However, normally this is not necessary.

The function `table(x)` produces a frequency table for the variable `x`. Thus, we can use

```
> table(assoc$sex)
```

0	1
121	129

which, again, tells us that there are 129 males and 121 females in this data set.

Tables of other qualitative variables, such as affection and SNPs, can be generated in the same manner.

**Exercise 5** *Explore qualitative variables presented in assoc*

1. *How many affected and unaffected are present in the data set?*
2. *What is the proportion of affected?*
3. *What is the distribution of snp4?*

A more convenient way to access data presented in a data frame is through "attaching" it to the R search path by

```
> attach(assoc)
```

After that, the variables can be accessed directly, e.g.

```
> table(sex)
```

sex	
0	1
121	129

As with conventional vectors, conditions may be used to subset variables in a data frame. If for example, you want to check distribution of number of affected in men and women separately, you can do that by

```
> table(aff[sex == 1])
```

0	1
98	31

```
> table(aff[sex == 0])
```

0	1
96	25

On R command line pressing the "up-arrow" button makes the last typed command re-appear (pressing it one more time will bring you to the one before the last, so on). This is very handy when you have to repeat the same analysis of different variables

**Summary:**

- The list of available objects can be viewed with `ls()`; a class of some object `obj` can be interrogated with `class(obj)`.
- Simple summary statistics for numeric variables can be generated by using `summary` function
- Histogram for some variable `var` can be generated by `hist(var)`
- A variable with name `name` from a data frame `frame`, can be accessed through `frame$name`.
- You can attach the data frame to the search path by `attach(frame)`. Then the variables contained in this data frame may be accessed directly. To detach the data frame (because, e.g., you are now interested in other data frame), use `detach(frame)`.

**Exercise 6** An `srdta` data supplied with *GenABEL* contains a data frame with phenotypes. This data frame may be accessed through `srdta@phdata`. Explore this data frame and answer the questions

1. What is the value of the 4th variable for subject number 75?
2. What is the value of variable 1 for person 75? Check what is the value of this variable for the first ten people. Can you guess what variable 1 is?
3. What is sum of the variable 2? Can you guess what it is?

**Exercise 7** Explore `phdata` slot of `srdta`

1. How many people has age over 65 years?
2. What is the sex distribution in the people over 65 years old?

## 2.3 Exploratory analysis of qualitative and quantitative traits

Contingency tables of more than one variable may be generated in R using the `table` command we have been using before to explore frequency distributions. For example, if you want cross-tabulate sex and affection status in the data frame `assoc`, you can use

```
> table(sex, aff)

  aff
sex 0 1
  0 96 25
  1 98 31
```

Here, the first variable (sex) is presented in rows and the second (affection status) in columns.

As is usual case with R, the output may be saved as a new object (of class 'table', which is a variety of a matrix):

```
> a <- table(sex, aff)
> a

  aff
sex 0 1
  0 96 25
  1 98 31
```

and this object may be analysed further.

For example, we can easily get the number of affected male with

```
> a[2, 2]
[1] 31
```

Alternatively, we can analyse the resulting contingency table **a** with more complex functions. If we want to see proportions in this table, we can use

```
> prop.table(a)

  aff
sex      0      1
  0 0.384 0.100
  1 0.392 0.124
```

Needless to say, this is equivalent to

```
> prop.table(table(assoc$sex, assoc$aff))

  0      1
  0 0.384 0.100
  1 0.392 0.124
```

In the above table, we see what proportion of people belong to four different classes (affected male, affected female, unaffected male and unaffected female). We may be though interested in the proportion of males in affected and unaffected. This may be achieved by

```
> prop.table(a, 2)

  aff
sex      0      1
  0 0.4948454 0.4464286
  1 0.5051546 0.5535714
```

saying us that 55.4% of affected are male.

Alternatively, we may be interested in proportion of affected among males/females. To answer this question, run

```
> prop.table(a, 1)

    aff
sex      0      1
0 0.7933884 0.2066116
1 0.7596899 0.2403101
```

saying us that 55.4% of male are affected.

Other useful contingency table analysis function is `fisher.test`, which implements the Fisher Exact Test of independence:

```
> fisher.test(a)

Fisher's Exact Test for Count Data

data: a
p-value = 0.547
alternative hypothesis: true odds ratio is not equal to 1
95 percent confidence interval:
0.6409648 2.3156591
sample estimates:
odds ratio
1.213747
```

Exploration of genetic data within base R, though possible, may be a bit of a pain. For example, we can easily generate contingency table of SNP5 vs affected status:

```
> a <- table(aff, snp5)
> a

snp5
aff A/A B/A B/B
0 31 88 71
1 9 26 17
```

We can also look up what is the proportion of affected among different genotypic groups

```
> prop.table(a, 2)

snp5
aff      A/A      B/A      B/B
0 0.7750000 0.7719298 0.8068182
1 0.2250000 0.2280702 0.1931818
```

showing that proportion of cases is similar in 'A/A' and 'A/B' genotypic groups and somewhat decreased in 'B/B'. It is easy to test if this affection is statistically independent of genotype by

```

> chisq.test(a)

Pearson's Chi-squared test

data: a
X-squared = 0.3874, df = 2, p-value = 0.8239

```

which gives (insignificant) genotypic association test on two degrees of freedom.

However, testing Hardy-Weinberg equilibrium, testing allelic effects, and even computation of allelic frequency is not so straightforward. Such specific genetic tests are implemented in special R libraries, such as `genetics` and `GenABEL` and will be covered in later sections of this document.

At this moment we will switch to exploratory analysis of quantitative traits. We will make use of the `srdta` data supplied with `GenABEL`. As you can remember from an exercise, the library is loaded with `library(GenABEL)` and the data are loaded with `data(srdta)`: Then the phenotypic data frame may be accessed through `srdta@phdata`.

**Exercise 8** Explore `srdta@phdata`. How many observations and variables are presented in the data frame? What are the classes of these variables?

As it was mentioned before, the function `summary()` generates a summary statistics for an object. For example, to see summary for trait `qt1`, we can use

```

> summary(srdta@phdata$qt1)

  Min. 1st Qu. Median     Mean 3rd Qu.    Max.    NA's
-4.6000 -0.9500 -0.3100 -0.2981  0.3800  3.2000  3.0000

```

`summary` is quite useful function which may operate in different ways for objects of different classes. Try `summary(srdta@phdata)`.

With R, it is also easy to explore the data graphically. For example, the histogram for `qt1` may be generated by

```

> hist(srdta@phdata$qt1)

(resulting histogram is shown at figure 1)

In similar manner, scatter-plots may be generated. To see relation between
qt1 and qt3, you can run

> plot(srdta@phdata$qt1, srdta@phdata$qt3)

```

(resulting plot is shown at figure 2)

The mean, median, minimum and maximum of the distribution of the trait may be found out using functions `mean`, `median`, `min` and `max`, respectively. The variance and standard deviation can be computed with `var` and `sd`.

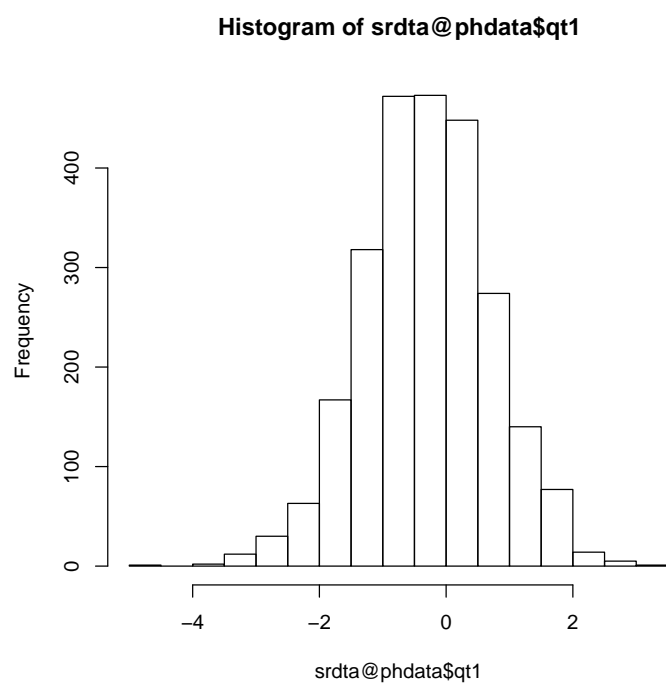


Figure 1: Histogram of qt1

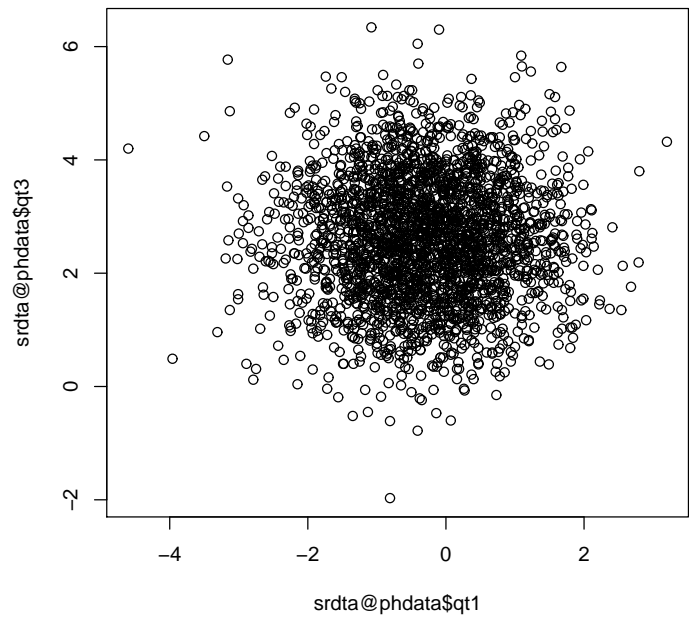


Figure 2: Scatter-plot of qt1 against qt3

To compute correlation between two variables (or all variables in a matrix/data frame), use `cor`.

In `GenABEL`, there is a special function designed to facilitate phenotypic quality control. This function takes names of variables and a data frame as an input, and returns summary statistics, list of outliers (using False Discovery Rate) and graphs.

For example, to do QC of sex, age and qt3, try

```
> check.trait(c("sex", "age", "qt3"), srdta@phdata)

-----
Trait sex has 2500 measurements
Missing: 0 ( 0 %)
Mean = 0.51 ; s.d. = 0.5
NO outliers discovered for trait sex
-----

Trait age has 2500 measurements
Missing: 0 ( 0 %)
Mean = 50.0378 ; s.d. = 7.060125
NO outliers discovered for trait age
-----

Trait qt3 has 2489 measurements
Missing: 11 ( 0.44 %)
Mean = 2.60859 ; s.d. = 1.101154
NO outliers discovered for trait qt3
```

The corresponding graph is depicted at figure ??.

Before you start with the exercise: if a function returns unexpected results, and you are confident that syntax was right, checking help page is always a good idea!

#### Exercise 9 Explore variables in `phdata` slot of `srdta`

1. What is the mean, median, minimum and maximum age in the sample?
2. Compare the distribution of qt3 in people younger and older than 65 years. Use function `sd(A)` to get standard deviation of A.
3. Produce distributions of different traits. Do you see something special?
4. What is correlation between qt3 and age?

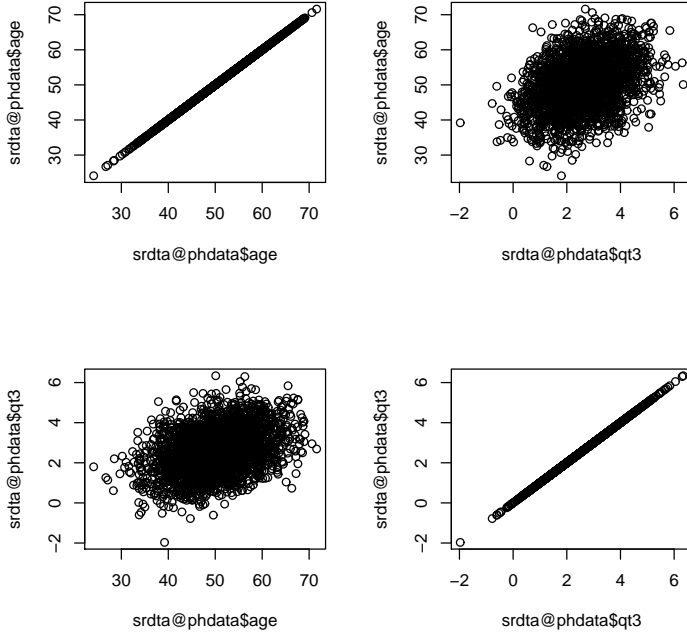


Figure 3: Quality control graph for `sex`, `age`, `qt3`

## 2.4 Regression analysis

While contingency tables, bi-plots and correlation are powerful tools to analyse relations between pairs of variable, a more general framework allowing investigation of relation of an outcome to multiple predictors is regression. In R, function `lm` implements linear regression modelling, and function `glm` implements generalised linear regression. In this section, we will use these two functions to analyse quantitative and binary outcomes.

You can do linear regression to check if trait `qt2` has relation with sex and age by

```
> a <- lm(srdta@phdata$qt2 ~ srdta@phdata$age + srdta@phdata$sex)
```

The results of analysis are stored in object 'a', which has class 'lm' and contains many sub-objects:

```
> class(a)
[1] "lm"
> names(a)
[1] "coefficients"      "residuals"        "effects"          "rank"
[5] "fitted.values"     "assign"           "qr"              "df.residual"
[9] "xlevels"            "call"             "terms"           "model"
```

At this moment you do not need to understand all these sub-objects; the meaningful summary of analysis is produced with

```
> summary(a)

Call:
lm(formula = srdta@phdata$qt2 ~ srdta@phdata$age + srdta@phdata$sex)

Residuals:
    Min      1Q   Median      3Q      Max 
-5.6498 -1.7953 -1.0328 -0.3148 883.0808 

Coefficients:
            Estimate Std. Error t value Pr(>|t|)    
(Intercept) -1.55892   4.41667  -0.353   0.724    
srdta@phdata$age  0.14022   0.08668   1.618   0.106    
srdta@phdata$sex  1.30377   1.22393   1.065   0.287    

Residual standard error: 30.59 on 2497 degrees of freedom
Multiple R-squared: 0.001518,          Adjusted R-squared: 0.0007181 
F-statistic: 1.898 on 2 and 2497 DF, p-value: 0.1501
```

You can see that `qt2` is not associated with age or sex.

As before, to make easy access to your data (basically, to avoid typing `srdta@phdata` before every trait name, you may attach the data to the search path:

```
> attach(srdta@phdata)
```

Then, the above expression to run linear regression analysis simplifies to:

```
> summary(lm(qt2 ~ age + sex))

Call:
lm(formula = qt2 ~ age + sex)

Residuals:
    Min      1Q   Median      3Q      Max 
-5.6498 -1.7953 -1.0328 -0.3148 883.0808 

Coefficients:
            Estimate Std. Error t value Pr(>|t|)    
(Intercept) -1.55892   4.41667  -0.353   0.724    
age          0.14022   0.08668   1.618   0.106    
sex          1.30377   1.22393   1.065   0.287    

Residual standard error: 30.59 on 2497 degrees of freedom
Multiple R-squared: 0.001518,          Adjusted R-squared: 0.0007181 
F-statistic: 1.898 on 2 and 2497 DF, p-value: 0.1501
```

with the same results.

Analysis of binary outcomes may be performed using `glm` function, using *binomial family* for the error distribution and the link function. For example, to figure out if your binary trait (`bt`) is associated with sex and age, you need to tell that this is binary trait:

```
> a <- glm(bt ~ age + sex, family = "binomial")
> summary(a)

Call:
glm(formula = bt ~ age + sex, family = "binomial")

Deviance Residuals:
    Min      1Q  Median      3Q     Max 
-1.992 -1.091 -0.444  1.094  1.917 

Coefficients:
            Estimate Std. Error z value Pr(>|z|)    
(Intercept) -4.639958   0.330519 -14.038 < 2e-16 ***
age          0.088860   0.006463  13.749 < 2e-16 ***
sex          0.379593   0.084138   4.512 6.44e-06 ***
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

(Dispersion parameter for binomial family taken to be 1)

Null deviance: 3450.5 on 2488 degrees of freedom
Residual deviance: 3216.5 on 2486 degrees of freedom
(11 observations deleted due to missingness)
AIC: 3222.5
```

Number of Fisher Scoring iterations: 4

There is strong association between `bt` and sex and age. If you want to characterise the strength of association to a binary trait with Odds Ratios, take the exponents of the regression coefficient. For example, the odds ratio associated with male is

```
> exp(0.3796)
[1] 1.461700
```

### 3 Introduction to association analysis in R

When analyzing several (dozens of) SNPs, facilities of base R are sufficient and efficient for data storage and analysis. Few specific test, such as these of Hardy-Weinberg Equilibrium (HWE) and Linkage Disequilibrium (LD), are implemented in different libraries, e.g. `genetics` and `GenABEL`.

In this section, we will describe library `genetics` and will make use of it to guide you through simple genetic analysis exercise using a small example data set. In the last part, you will investigate a bigger data set as based on the knowledge obtained in the first part, and will answer the questions.

### 3.1 Exploring genetic data with library genetics

Library `genetics` was written by Gregory R. Warnes to facilitate analysis of genetic data in R. This library

- Implements genetic analysis tests, such as test for Hardy-Weinberg equilibrium and Linkage disequilibrium.
- Implements new data classes, such as `genotype`, `haplotype` and `LD.data.frame`.
- Modifies default R functions, such as `summary` and `plot` to correctly analyse and present these new classes.
- Facilitates export of the data from R to the formats supported by such genetic analysis packages as GenePop and QTDT.

Start R by double-click on the file `ge03d1p1.RData`. Load library `genetics`, which we will need for testing HWE and computations of LD by

```
> library(genetics)
```

The file you have loaded contains single data frame `assocg`. Let us briefly explore it:

```
> class(assocg)
[1] "data.frame"
> names(assocg)
[1] "subj" "sex"   "aff"   "qt"    "snp4"  "snp5"  "snp6"
> dim(assocg)
[1] 250   7
```

You can see that `assocg` looks remarkably similar to the previously explored data frame `assoc` (section 2.2, page 11). Indeed, they are almost equivalent. Let us present the data for the subjects 5 to 15 and compare this output to that presented on page 13:

```
> assocg[5:15, ]
  subj sex aff      qt.snp4.snp5.snp6
  1533 1533 0 0.1009220 A/B B/A B/A
  2466 2466 1 0 -0.1724321 A/B A/A A/A
  2425 2425 0 0 -0.3378473 B/B A/A A/A
  1068 1068 0 0 -1.7112925 A/A B/B <NA>
  198 198 1 0 -0.4815822 A/B B/A B/A
  1496 1496 1 0 1.2281232 A/A B/B B/B
  909 909 0 0 0.5993945 A/B B/A B/A
  1213 1213 0 0 1.9792190 A/A B/B B/B
  181 181 1 0 1.5435921 A/A B/B B/B
  1783 1783 0 0 -1.6242738 A/B B/A B/A
  1914 1914 0 0 -0.5160331 A/A B/B B/B
```

The data are identical. However, the SNP data presented in the new data frame have special class `genotype`, as implemented in library `genetics`:

```
> class(assocg$snp4)
[1] "genotype" "factor"
```

Previously, the SNP genotypes were coded as characters. This new way of presentation allows library `genetics` to recognise the SNP data as genetic and analyse them accordingly.

Let us attach the `assocg` data frame and explore what data analysis advantages are achieved by application of library `genetics`.

```
> attach(assocg)
```

As we noted in section 2.2, testing Hardy-Weinberg equilibrium, testing allelic effects, and even computation of allelic frequency is not so straightforward in base R. These tests, are, however, easy with library `genetics`. To see the allelic frequencies and other summary statistics for a SNP, you can use

```
> summary(snp4)

Number of samples typed: 243 (97.2%)
```

```
Allele Frequency: (2 alleles)
  Count Proportion
A     323      0.66
B     163      0.34
NA    14       NA
```

```
Genotype Frequency:
  Count Proportion
B/B    29      0.12
A/B   105      0.43
A/A   109      0.45
NA     7       NA
```

```
Heterozygosity (Hu) = 0.4467269
Poly. Inf. Content   = 0.3464355
```

To check these characteristics in controls and cases separately, you can use

```
> summary(snp4[aff == 0])
```

```
Number of samples typed: 190 (97.9%)
```

```
Allele Frequency: (2 alleles)
  Count Proportion
A     255      0.67
B     125      0.33
NA    8        NA
```

```

Genotype Frequency:
  Count Proportion
B/B     22      0.12
A/B     81      0.43
A/A     87      0.46
NA      4       NA

Heterozygosity (Hu) = 0.4426469
Poly. Inf. Content  = 0.3440288

> summary(snp4[aff == 1])

Number of samples typed: 53 (94.6%)

Allele Frequency: (2 alleles)
  Count Proportion
A      68      0.64
B      38      0.36
NA     6       NA

Genotype Frequency:
  Count Proportion
B/B     7       0.13
A/B    24      0.45
A/A    22      0.42
NA     3       NA

Heterozygosity (Hu) = 0.4643306
Poly. Inf. Content  = 0.3541731

```

Let us check if HWE holds for the SNPs described in this data frame. We can do exact test for HWE by

```

> HWE.exact(snp4)

Exact Test for Hardy-Weinberg Equilibrium

data: snp4
N11 = 109, N12 = 105, N22 = 29, N1 = 323, N2 = 163, p-value = 0.666

```

If you want to check HWE using controls only, you can do it by

```

> HWE.exact(snp4[aff == 0])

Exact Test for Hardy-Weinberg Equilibrium

data: snp4[aff == 0]
N11 = 87, N12 = 81, N22 = 22, N1 = 255, N2 = 125, p-value = 0.6244

```

Let us check if there is LD between snp4 and snp5:

```

> LD(snp4, snp5)

Pairwise LD
-----
          D      D'      Corr
Estimates: 0.2009042 0.9997352 0.8683117

          X^2 P-value   N
LD Test: 354.3636      0 235

```

The output shows results of the test for significance of LD, and estimates of the magnitude of LD ( $D'$  and correlation,  $r$ ). To obtain  $r^2$ , you can either square the correlation manually

```
> 0.8683117 * 0.8683117
```

```
[1] 0.7539652
```

or simply ask `LD()` to report it by

```
> LD(snp4, snp5)$"R^2"
```

```
[1] 0.7539652
```

The latter command is possible because the `LD()` function actually computes more things than it reports. This is quite common for R functions. You can apply `names()` function to the analysis objects to see (at least part of) what was actually computed. Try

```
> ld45 <- LD(snp4, snp5)
```

and check what are the sub-objects contained in this analysis object

```
> names(ld45)
```

```
[1] "call"      "D"        "D'"       "r"        "R^2"      "n"        "X^2"
[8] "P-value"
```

Any of these variables can be accessed through `object$var` syntax, e.g. to check  $D'$  we can use

```
> ld45$"D"
```

```
[1] 0.9997352
```

To check LD for more than two SNPs, we can compute an LD analysis object by

```
> ldall <- LD(data.frame(snp4, snp5, snp6))
```

and later check

```
> ldall$"P-value"
```

	snp4	snp5	snp6
snp4	NA	0	0
snp5	NA	NA	0
snp6	NA	NA	NA

to see significance,

```
> ldall$"D"
```

	snp4	snp5	snp6
snp4	NA	0.9997352	0.8039577
snp5	NA	NA	0.9997231
snp6	NA	NA	NA

for  $D'$  and

```
> ldall$"R^2"
```

	snp4	snp5	snp6
snp4	NA	0.7539652	0.5886602
snp5	NA	NA	0.8278328
snp6	NA	NA	NA

for  $r^2$ .

You can also present e.g.  $r^2$  matrix as a plot by

```
> image(ldall$"R^2")
```

A more neat way to present it requires specification of the set of threshold (break points) and colors to be used (you do not need to try this example if you do not want):

```
> image(ldall$"R^2", breaks = c(0.5, 0.6, 0.7, 0.8, 0.9, 1), col = heat.colors(5))
```

Resulting plot is shown at figure 4.

For any R command, you can get help by typing `help(command)`. Try `help(image)` if you are interested to understand what are "breaks" and "col"; or try `help(heat.colors)` to figure this color schema out.

Similar to our HWE checks, we may want to compute (and compare) LD in cases and controls separately:

```
> ldcases <- LD(data.frame(snp4, snp5, snp6)[aff == 1, ])
> ldcases$"R^2"

snp4      snp5      snp6
snp4    NA 0.7615923 0.6891558
snp5    NA        NA 0.8943495
snp6    NA        NA        NA

> ldcontr <- LD(data.frame(snp4, snp5, snp6)[aff == 0, ])
> ldcontr$"R^2"

snp4      snp5      snp6
snp4    NA 0.7512458 0.5616395
snp5    NA        NA 0.8075894
snp6    NA        NA        NA
```

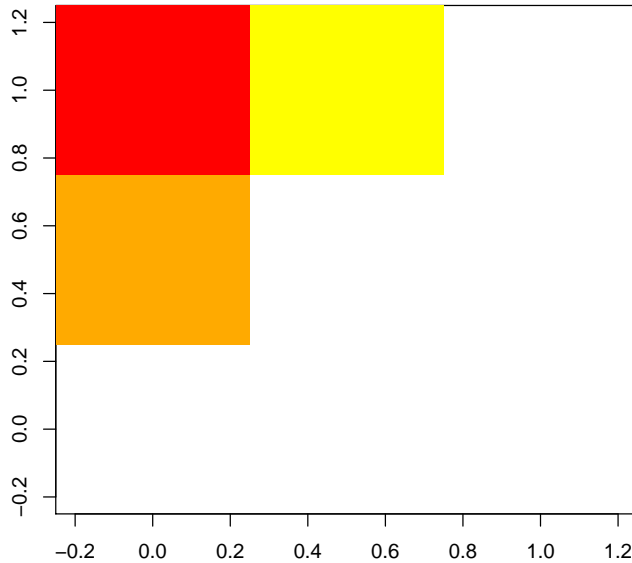


Figure 4:  $r^2$  plot for snp4, snp5 and snp6

and even present it results for cases and controls on the same graph (you do not need to produce this graph, which is presented at the figure 5):

```
> image(ldcases$"R^2", breaks = c(0.5, 0.6, 0.7, 0.8, 0.9, 1),
+       col = heat.colors(5))
> image(t(ldcontr$"R^2"), breaks = c(0.5, 0.6, 0.7, 0.8, 0.9, 1),
+       col = heat.colors(5), add = T)
```

### 3.2 Example association analysis

Now, after we have described genetic and phenotypic data separately, we are ready to test association between these two. In previous sections, we showed that association between a binary trait and genotype may be analysed using contingency tables (functions `table`, `prop.table`, `fisher.test`, etc.). The association between a quantitative trait and genotype may be done using correlations, T-test, etc.

However, a more flexible analysis is possible when using regression modelling. First, we will investigate relation between the quantitative trait `qt` and the SNPs by using linear regression

```
> mg <- lm(qt ~ snp4)
```

The `lm` command fits linear regression model to the data and returns an analysis object. The summary of analysis may be generated with

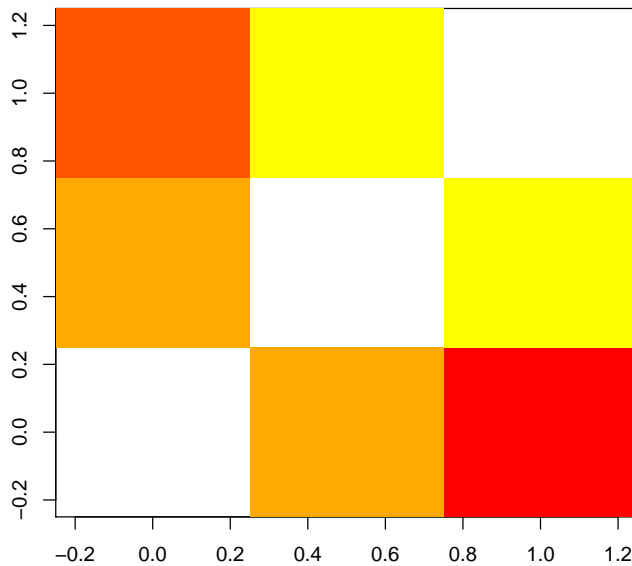


Figure 5:  $r^2$  plot for snp4, snp5 and snp6. Above diagonal: LD in cases; below: controls

```
> summary(mg)

Call:
lm(formula = qt ~ snp4)

Residuals:
    Min      1Q   Median      3Q     Max 
-2.63700 -0.62291 -0.01225  0.58922  3.05561 

Coefficients:
            Estimate Std. Error t value Pr(>|t|)    
(Intercept) -0.081114  0.092517  -0.877   0.382    
snp4A/B     -0.108366  0.132079  -0.820   0.413    
snp4B/B     -0.006041  0.201820  -0.030   0.976    

Residual standard error: 0.9659 on 240 degrees of freedom
(7 observations deleted due to missingness)
Multiple R-squared:  0.003049,    Adjusted R-squared: -0.005259 
F-statistic: 0.367 on 2 and 240 DF,  p-value: 0.6932
```

From the summary output, it is clear that the model assumes arbitrary (estimated) effects of the genotypes AA, AB and BB. Neither effect of AB nor BB is significant in this case. The global test on two degrees of freedom (bottom of

the output) is also not significant.

If you want to include some covariate into your model, e.g. sex, you can easily do that by adding the term to the formula:

```
> summary(lm(qt ~ sex +.snp4))

Call:
lm(formula = qt ~ sex + .snp4)

Residuals:
    Min      1Q   Median      3Q     Max 
-2.664422 -0.624169 -0.008752  0.597045  3.080857 

Coefficients:
            Estimate Std. Error t value Pr(>|t|)    
(Intercept) -0.110298   0.115260  -0.957   0.340    
sex          0.053018   0.124493   0.426   0.671    
.snp4A/B     -0.104429   0.132628  -0.787   0.432    
.snp4B/B     -0.002452   0.202340  -0.012   0.990    

Residual standard error: 0.9676 on 239 degrees of freedom
(7 observations deleted due to missingness)
Multiple R-squared:  0.003805,    Adjusted R-squared: -0.0087 
F-statistic: 0.3043 on 3 and 239 DF,  p-value: 0.8223
```

You can also allow for interaction by using the "\*" operator

```
> summary(lm(qt ~ sex * .snp4))

Call:
lm(formula = qt ~ sex * .snp4)

Residuals:
    Min      1Q   Median      3Q     Max 
-2.570485 -0.645961 -0.002641  0.610938  3.019696 

Coefficients:
            Estimate Std. Error t value Pr(>|t|)    
(Intercept) -0.20579   0.13834  -1.487   0.138    
sex          0.22649   0.18647   1.215   0.226    
.snp4A/B     0.05222   0.19024   0.274   0.784    
.snp4B/B     0.18071   0.28576   0.632   0.528    
sex:.snp4A/B -0.30191   0.26566  -1.136   0.257    
sex:.snp4B/B -0.35508   0.40531  -0.876   0.382    

Residual standard error: 0.9684 on 237 degrees of freedom
(7 observations deleted due to missingness)
Multiple R-squared:  0.01041,    Adjusted R-squared: -0.01047 
F-statistic: 0.4984 on 5 and 237 DF,  p-value: 0.7773
```

Note that both main effects of sex and .snp4, and also effects of interaction are estimated in this model.

Of interest in genetic studies may be three other models: additive, dominant and recessive.

The additive model assumes that the difference between mean trait's values between 'AA' and 'BB' is twice the difference between 'AA' and 'BB', that is the mean value of the trait for heterozygous genotypes is right in between the two homozygotes. To test additive model, we first need to recode the predictor (genotype) as a numeric factor to be used as covariate. This can be easily done with function `as.numeric`:

```
> add4 <- as.numeric(snp4) - 1
```

We can check if recoding was done correctly by producing the table

```
> table(snp4, add4)
```

add4			
snp4	0	1	2
A/A	109	0	0
A/B	0	105	0
B/B	0	0	29

Now to test the additive model run

```
> summary(lm(qt ~ add4))
```

Call:

```
lm(formula = qt ~ add4)
```

Residuals:

Min	1Q	Median	3Q	Max
-2.54813	-0.62104	-0.02754	0.60584	3.00652

Coefficients:

	Estimate	Std. Error	t value	Pr(> t )
(Intercept)	-0.10476	0.08710	-1.203	0.230
add4	-0.03563	0.09133	-0.390	0.697

Residual standard error: 0.9651 on 241 degrees of freedom  
(7 observations deleted due to missingness)

Multiple R-squared: 0.0006313, Adjusted R-squared: -0.003516  
F-statistic: 0.1522 on 1 and 241 DF, p-value: 0.6968

The model assuming dominant action of the 'A' allele means that the means of genotypes 'AA' and 'AB' are the same. This is equivalent to the model of recessive action of 'B' allele. To code SNP4 according to this model, we can use function `replace`:

```
> dom4 <- add4
> dom4[dom4 == 2] <- 1
> table(snp4, dom4)
```

dom4		
snp4	0	1

```

A/A 109   0
A/B   0 105
B/B   0   29

```

To test association with a binary outcome, we will use function `glm` with `binomial` family:

```

> summary(glm(aff ~ snp4, family = "binomial"))

Call:
glm(formula = aff ~ snp4, family = "binomial")

Deviance Residuals:
    Min      1Q  Median      3Q     Max 
-0.7433 -0.7204 -0.6715 -0.6715  1.7890 

Coefficients:
            Estimate Std. Error z value Pr(>|z|)    
(Intercept) -1.3749    0.2386 -5.761 8.35e-09 ***  
snp4A/B      0.1585    0.3331  0.476   0.634    
snp4B/B      0.2297    0.4952  0.464   0.643    
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

(Dispersion parameter for binomial family taken to be 1)

Null deviance: 254.91 on 242 degrees of freedom
Residual deviance: 254.58 on 240 degrees of freedom
(7 observations deleted due to missingness)
AIC: 260.58

Number of Fisher Scoring iterations: 4

To make a test of global significance of the SNP effect, you can use
> anova(glm(aff ~ snp4, family = "binomial"), test = "Chisq")
Analysis of Deviance Table

Model: binomial, link: logit

Response: aff

Terms added sequentially (first to last)


```

	Df	Deviance	Resid. Df	Resid. Dev	P(> Chi )
NULL		242	254.908		
snp4	2	0.329	240	254.579	0.848

In the manner similar to that described for quantitative traits, additive and dominance/recessive models can be tested by proper coding of the genotypic variable, e.g. to test the additive model, use

```

> summary(glm(aff ~ as.numeric(snp4), family = "binomial"))

Call:
glm(formula = aff ~ as.numeric(snp4), family = "binomial")

Deviance Residuals:
    Min      1Q  Median      3Q     Max 
-0.7548 -0.7139 -0.6747 -0.6747  1.7842 

Coefficients:
            Estimate Std. Error z value Pr(>|z|)    
(Intercept) -1.4913    0.4164 -3.581 0.000342 ***  
as.numeric(snp4) 0.1272    0.2268  0.561 0.574994    
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

(Dispersion parameter for binomial family taken to be 1)

Null deviance: 254.91 on 242 degrees of freedom
Residual deviance: 254.60 on 241 degrees of freedom
(7 observations deleted due to missingness)
AIC: 258.60

Number of Fisher Scoring iterations: 4

```

Now you have learned all commands necessary to answer the questions of the next section.

Exit R by typing `q()` command (do not save image) and proceed to the self exercise.

### 3.3 Exercise

Start R by double-click over the file `ge03d1p2.RData`. Explore the data frame present and answer the questions.

**Exercise 10** *How many SNPs are described in this data frame?*

**Exercise 11** *What is the frequency (proportion) of `snp1` allele A? What is its frequency in these affected (`aff==1`)?*

**Exercise 12** *How many cases and controls are present?*

**Exercise 13** *If all subjects are used to test HWE, are there any SNPs out of HWE at nominal  $P \leq 0.05$ ? Which ones?*

**Exercise 14** *If only controls are used to test the SNPs which are out of HWE in total sample, are these still out of HWE?*

**Exercise 15** *Which SNP pairs are in strong LD ( $r^2 \geq 0.8$ )?*

**Exercise 16** *For SNPs in strong LD, what is  $r^2$  for separate samples of cases and controls?*

**Exercise 17** Is there significant association between affection status and sex? What is P-value for association?

**Exercise 18** Is association between the disease and *qt* significant?

**Exercise 19** Which SNPs are associated with the quantitative trait *qt* at nominal  $P \leq 0.05$ ? Use 2 d.f. test.

**Exercise 20** Test each SNP for association with the affection status, using 2 d.f. test. Which SNPs are significantly associated at nominal  $P \leq 0.05$ ? How can you describe the model of action of the significant SNPs?

**Exercise 21** For the SNPs selected in previous question, test association using additive model. Which SNPs are still associated?

**Exercise 22** If you adjust the analysis under additive model (question 21) for significant covariates which you discovered in questions 17 and 18, are these findings still significant?

**Exercise 23** Test association between *aff* and *snp5* and *snp10*, allowing for the SNPs interaction effect. Use arbitrary (not an additive) model. Do you observe significant interaction? How can you describe the model of concert action of *snp5* and *snp10*?

## 4 Introduction to GenABEL

In this section, you will become familiar with the **GenABEL** library, designed for GWA analysis. Compared to **genetics** package, it provides specific facilities for storage and manipulation of large amounts of data, very fast tests for GWA analysis, and special functions to analyse and graphically present the results of GWA analysis (thus "analysis of analysis").

Start R and load **GenABEL** library using command

```
> library(GenABEL)
```

After that, load the data with the command

```
> data(srdata)
```

### 4.1 General description of gwaa.data-class

The object you have loaded, **srdata**, belongs to the **gwaa.data** class. This is a special class developed to facilitate GWA analysis.

In GWA analysis, different types of data are used. These include the phenotypic and genotypic data on the study participants and chromosome and location of every SNP. For every SNP, it is desirable to know the details of coding (what are alleles? – A, T, G, C? – and what is the strand – '+' or '–', 'top' or 'bot'? – this coding is for).

One could attempt to store all phenotypes and genotypes together in a single table, using, e.g. one row per study subject; then the columns will correspond to study phenotypes and SNPs. For a typical GWA data set, this would lead

to a table of few thousands rows and few hundreds of thousands of columns. Such a format is generated when one downloads HapMap data for a region. To store GWA data in such tables internally, within R, proves to be inefficient. In **GenABEL**, special data class, **gwaa.data-class** is used to store GWA data. The structure of this data class is shown at the figure 6.

An object of some class has "slots" which may contain actual data or objects of other classes. The information stored at a particular **slot** of an **object** can be accessed by command **object@slot**.

At the first level, a **gwaa.data-class** object has slot **phdata**, which contains all phenotypic information in a data frame (**data.frame-class** object). The rows of this data frame correspond to study subjects, and the columns correspond to the variables. There are two default variables, which are always present in **phdata**. The first of these is "id", which contains study subject identification code. This identification code can be arbitrary character, but every person must be coded with an unique ID. The second default variable is "sex", where males are coded with ones ("1") and females are coded with zero ("0"). It is important to understand that this data frame is not supposed to be directly modified by the user. In particular, it is extremely important to remember that one should not directly add subjects to the table, change the values of "id" and "sex", and change the order of subjects in **phdata** unless this one is really understands the way **GenABEL** works. One also should not run such data manipulation functions as **merge**, **cbind** and **rbind** – exactly because they may change the number of subjects or interfere with the order. On the other hand, it is OK to add more variables to the data frame through direct computations, for example, if one wishes to add computed body mass index, it is OK to run the command like

```
obj@phdata$bmi <- obj@phdata$weight/((obj@phdata$height)^2)
```

To add many variables to **phdata**, special **GenABEL** function **add.phdata** should be used.

The other slot of an object of **gwaa.data-class** is slot **gtodata**, which contains all GWA genetic information in an object of class **snp.data** class (figure 6). This class, in turn, has slots **nids**, containing the number of study subjects, **idnames**, containing all ID names of these subjects, **nsnps**, containing the number of SNPs typed, **snpnames**, containing the SNP names, **chromosome**, containing the name of the chromosome the SNPs belong to and slot **map** with map position of SNPs, and slot **male**, containing the sex code for the subjects (1=males, 0=females). The latter is identical to the "sex" variable contained in the **phdata**, but is duplicated here because many operations with purely genetic data, in particular these concerning analysis of sex chromosomes, depend on the sex. The strand information is presented in the slot **strand**. **GenABEL** codes strand as "+" (forward), "-" (reverse) or "u" (unknown). Of course, if you prefer top/bottom coding, this information may be stored in the same form – you will just need to remember that "+" corresponds to e.g. "top", and "-" to "bottom" strand. The allelic coding is presented in slot **coding**. Coding for every allele is presented with a pair of characters, for example "AG". Thus, for such polymorphism, you may expect "AA", "AG" and "GG" genotypes to be found in population. The order (that is "AG" vs "GA") is important – the first allele reported is the one which will be used as a reference in association analysis, and thus the effects are reported for the second allele. To avoid memory overheads, the strand and coding information is internally stored as **snp.strand-class** and **snp.coding-class**. Information can be converted to human-readable format

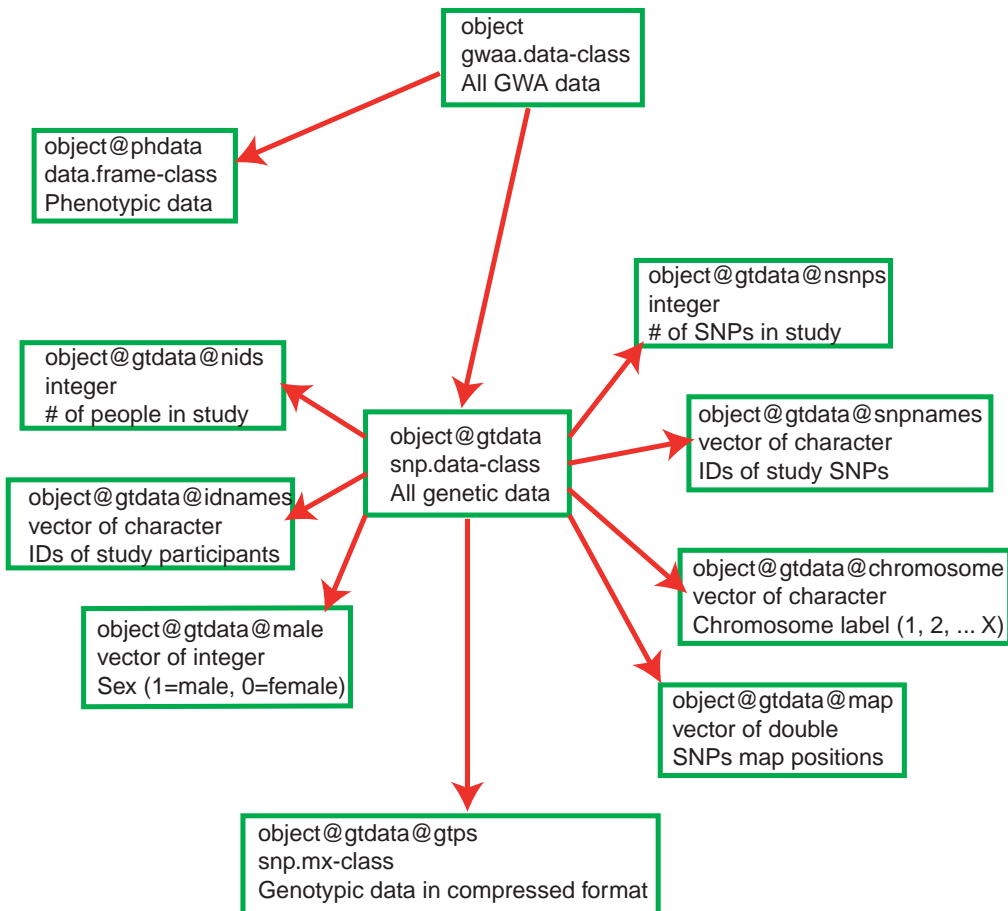


Figure 6: Structure of `gwaa.data-class`. In every box, first line contains the object and slot names, second line describes the class of this object, and third line describes what information is contained. NEEDS TO BE UPDATED WITH `object@gtdat@coding`; `snp.coding-class`; `snp allele coding` and `object@gtdat@strand`; `snp.strand-class`; `snp allele strand`

using `as.character` function.

If, for example, you would like to know, how many SNPs were included in the study (slot `nsnps` of the slot `gtdata` of `srdta`), you need to run command

```
> srdta@gtdata@nsnps
```

```
[1] 833
```

Thus, 833 SNPs were typed in the study. You can access information stored in any slot in this manner.

You may want to read the general GenABEL man page using `help(GenABEL)`. To see help on `gwaa.data-class`, you can use `help("gwaa.data-class")` (mind the quotation marks!).

#### Summary:

- An object of some class has "slots" which may contain actual data or objects of other classes. The information stored at a particular slot of an object can be accessed by command `object@slot`.
- GenABEL uses special data class, `gwaa.data-class`, to store GWA data.

#### Exercise 24 Explore `srdta`.

1. How many people are included in the study?
2. How many of these are males?
3. How many are females?
4. What is male proportion?

#### Exercise 25 Explore slot containing map (`map`) and slot containing SNP names (`snpnames`) of the `gtdata` slot of `srdta`.

1. What are names of markers located after 2,490,000 b.p.?
2. Between 1,100,000 and 1,105,000 b.p.?

## 4.2 Sub-setting and coercing `gwaa.data`

It is possible to sub-set the object, which stores the GWA data in the manner similar to the described above. Very primitively, you may think of an object of class `gwaa.data` as a matrix whose rows correspond to study subjects and columns correspond to SNPs studied (though the actual object is a way more complicated). For example, if we would like to investigate what is the content of `srdta` for the first 5 people and 3 SNPs, we can run

```
> ssubs <- srdta[1:5, 1:3]
> class(ssubs)

[1] "gwaa.data"
attr(,"package")
[1] "GenABEL"

> ssubs

  id sex age  qt1   qt2   qt3 bt
1 p1  1 43.4 -0.58  4.46 1.43  0
2 p2  1 48.2  0.80  6.32 3.90  1
3 p3  0 37.9 -0.52  3.26 5.05  1
4 p4  1 53.8 -1.55 888.00 3.76  1
5 p5  1 47.5  0.25  5.70 2.89  1

@nids = 5
@nsnps = 3
@nbytes = 2
@idnames = p1 p2 p3 p4 p5
@snpnames = rs10 rs18 rs29
@chromosome = 1 1 1
@coding = 08 0b 0c
@strand = 01 01 02
@map = 2500 3500 5750
@male = 1 1 0 1 1
@gtps =
40 40 40
40 40 00
```

As you can see, by sub-setting we obtained a smaller object of `gwaa.data-class`, with all its slots. Most of the information is straightforward and does not need further explanation.

There are though three slots which are not human-readable, i.e. `@coding`, `@strand`, and `@gtps`. These are coded using new classes based in R `raw` data type; these can be converted in human-readable format using a variety of '`as.*`' functions. For example, to see human-readable information on coding and strand, let us try

```
> as.character(ssubs@gtdata@coding)

rs10 rs18 rs29
"TG" "GA" "GT"
```

This tells what alleles are observed in the three SNPs, and which alleles will be used as a reference in association analysis.

To see the strand, use

```
> as.character(ssubs@gtdata@strand)

rs10 rs18 rs29
"+" "+" "-"
```

The slot `gtp`s contains the SNP data, and is not readable, because the information is compressed (each element contains data on up to four genotypes). To get human-readable information, an object of class `snp.data-class` (e.g. `srdta@gtdata`) can be coerced to a character using the same `as.character()` function:

```
> as.character(ssubs@gtdata)

rs10 rs18 rs29
p1 "T/T" "G/G" "G/G"
p2 "T/T" "G/G" NA
p3 "T/T" "G/G" NA
p4 "T/T" "G/G" NA
p5 "T/T" "G/A" "G/G"
```

For `@gtp`s conversion to other formats are possible as well. Other useful coercion is to "numeric":

```
> as.numeric(ssubs@gtdata)

rs10 rs18 rs29
p1    0    0    0
p2    0    0    NA
p3    0    0    NA
p4    0    0    NA
p5    0    1    0
```

You can see that conversion to numeric data type uses information on both underlying genotypes and coding. When coding is "GA", as is for the `rs18` (the second SNP), homozygotes for the first allele, as specified by coding ("G") are converted to zeros ("0"), heterozygotes are converted to ones ("1"), and homozygotes for the second allele ("A") are converted to twos ("2"). Clearly, when numerically converted data are used for association analysis, the effects will be estimated for the second allele, while first will be used as a reference.

Genotypic data converted to standard R "numeric" format can be used in any further analysis. Homozygotes of one type are coded as "0", heterozygotes are coded as "1" and other type of homozygotes is coded as "2". You can think of this as the number of allele of "B" type.

Several useful genetic analysis libraries were developed for R. These include `genetics` (analysis of linkage disequilibrium and many other useful functions) and `haplo.stats` (analysis of association between traits and haplotypes). These use their own genetic data formats.

One can translate `GenABEL` genetic data to the format used by "genetics" library by `as.genotype()`:

```
> as.genotype(ssubs@gtdata)

      rs10 rs18 rs29
p1   T/T  G/G  G/G
p2   T/T  G/G <NA>
p3   T/T  G/G <NA>
p4   T/T  G/G <NA>
p5   T/T  G/A  G/G
```

To translate GenABEL data to the format used by "haplo.stats" you can use function `as.hsgeno()`

```
> as.hsgeno(ssubs@gtdata)

      rs10.a1 rs10.a2 rs18.a1 rs18.a2 rs29.a1 rs29.a2
p1       1       1       1       1       1       1
p2       1       1       1       1     NA     NA
p3       1       1       1       1     NA     NA
p4       1       1       1       1     NA     NA
p5       1       1       1       2       1       1
```

Actually, most users will not need the latter function, as GenABEL provides a functional interface to "haplo.stats" (such GenABEL functions as `scan.haplo()` and `scan.haplo.2D()`).

It is possible to select sub-sets of `gwaa.data-class` based not only on index (e.g. first 10 people and SNP number 33), but also based on names.

For example, if we would like to retrieve phenotypic data on people with IDs "p141", "p147" and "p2000", we can use

```
> srdta[c("p141", "p147", "p2000"), ]@phdata

      id sex age qt1 qt2 qt3 bt
141  p141  0 47.2  0.51 5.23 2.17  0
147  p147  0 43.2  0.14 4.47 1.73  0
2000 p2000 0 43.1 -1.53 2.78 2.70  1
```

here, the first part of expression sub-sets `srdta` on selected IDs, and the second tells which part of the retrieved sub-set we want to see. You can try `srdta[c("p141", "p147", "p2000"), ]`, but be prepared to see long output, as all information will be reported.

In similar manner, we can also select on SNP name. For example, if we are interested to see information on SNPs "rs10" and "rs29" for above people, we can run

```
> srdta[c("p141", "p147", "p2000"), c("rs10", "rs29")]

      id sex age qt1 qt2 qt3 bt
141  p141  0 47.2  0.51 5.23 2.17  0
147  p147  0 43.2  0.14 4.47 1.73  0
2000 p2000 0 43.1 -1.53 2.78 2.70  1
@nids = 3
@nsnps = 2
```

```

@nbytes = 1
@idnames = p141 p147 p2000
@snpnames = rs10 rs29
@chromosome = 1 1
@coding = 08 0c
@strand = 01 02
@map = 2500 5750
@male = 0 0 0
@gtps =
40 40

```

To see the actual genotypes for the above three people and two SNPs, use

```
> as.character(srdata[c("p141", "p147", "p2000"), c("rs10", "rs29")])
```

	rs10	rs29
p141	"T/T"	"G/G"
p147	"T/T"	"G/G"
p2000	"T/G"	"G/T"

or

```
> as.numeric(srdata[c("p141", "p147", "p2000"), c("rs10", "rs29")])
```

	rs10	rs29
p141	0	0
p147	0	0
p2000	1	1

**Exercise 26** Explore genotypes for SNP "rs114".

1. What is the coding and which allele is the reference one?
2. What is the frequency of **non-reference** ("effective") allele in total sample?
3. What is the frequency of effective allele in male?
4. What is the frequency of effective allele in female?
5. What is the frequency of the **reference** allele in total sample, males and females?

**Summary:**

- It is possible to obtain subsets of objects of **gwaa.data-class** and **snp.data-class** using standard 2D sub-setting model [i,j], where i corresponds to study subjects and j corresponds to SNPs.

- It is possible to provide ID and SNP names instead of indexes when subsetting an object of class `gwaa.data-class`.
- Function `as.numeric()` converts genotypic data from `snp.data-class` to regular integer numbers, which can be used in analysis with R.
- Function `as.character()` converts genotypic data from `snp.data-class` to character format.
- Function `as.genotype()` converts genotypic data from `snp.data-class` to the format used by library `genetics`.
- Function `as.hsgeno()` converts genotypic data from `snp.data-class` to the format used by library `haplo.stats`.

### 4.3 Exploring genetic data

Implementation of function `summary()` to `snp.data` class is very useful in genetic data exploration and quality control (QC). Let us try application of this function to the `ssubs`:

```
> a <- summary(ssubs)
> a

$phdata
      id          sex         age        qt1
Length:5      Min.   :0.0   Min.   :37.90  Min.   :-1.55
Class :character  1st Qu.:1.0   1st Qu.:43.40  1st Qu.:-0.58
Mode  :character  Median :1.0   Median :47.50  Median :-0.52
                  Mean   :0.8   Mean   :46.16  Mean   :-0.32
                  3rd Qu.:1.0   3rd Qu.:48.20  3rd Qu.: 0.25
                  Max.   :1.0   Max.   :53.80  Max.   : 0.80
      qt2          qt3          bt
Min.   : 3.26  Min.   :1.430  Min.   :0.0
1st Qu.: 4.46  1st Qu.:2.890  1st Qu.:1.0
Median : 5.70  Median :3.760  Median :1.0
Mean   :181.55  Mean   :3.406  Mean   :0.8
3rd Qu.: 6.32  3rd Qu.:3.900  3rd Qu.:1.0
Max.   :888.00  Max.   :5.050  Max.   :1.0

$gtdata
      NoMeasured CallRate Q.2 P.11 P.12 P.22 Pexact      Fmax      Plrt
rs10          5     1.0 0.0    5    0    0      1  0.0000000 1.0000000
rs18          5     1.0 0.1    4    1    0      1 -0.1111111 0.7386227
rs29          2     0.4 0.0    2    0    0      1  0.0000000 1.0000000
      Chromosome
rs10          1
rs18          1
rs29          1
```

In the first section, the summary is generated for phenotypic data. In the second section, summary is generated for genotypic data. In this section, **NoMeasured** refers to the number of genotypes scores, **CallRate** to the proportion of these, **Q.2** is the frequency of the 'B' allele. The counts in three genotypic classes are provided next. **Pexact** refers to exact P-value for the test of Hardy-Weinberg equilibrium.

As you've seen above, an object of the class **gwaa.data-class** or **snp.data-class** is sub-settable in standard manner: [i,j], where i is an index of a study subject and j is an index of a SNP. Importantly, i could be a list of indexes:

```
> vec <- which(srdta@phdata$age >= 65)
> vec

[1] 64 122 186 206 207 286 385 386 492 514 525 536 545 565 613
[16] 632 649 673 701 779 799 981 1008 1131 1186 1223 1281 1383 1471 1489
[31] 1501 1565 1584 1673 1679 1782 1821 1832 1866 1891 1953 2081 2085 2140 2224
[46] 2268 2291 2384 2420 2453

> summary(srdta@gtdata[vec, 1:3])

  NoMeasured CallRate      Q.2 P.11 P.12 P.22     Pexact      Fmax
rs10          48    0.96 0.1354167   36    11     1 1.0000000  0.02131603
rs18          47    0.94 0.2765957   25    18     4 0.7245853  0.04298643
rs29          45    0.90 0.1555556   32    12     1 1.0000000 -0.01503759

  Plrt Chromosome
rs10 0.8843626      1
rs18 0.7697067      1
rs29 0.9188943      1
```

This shows summary of first three genotypes for people with age greater than or equal to 65 y.o. The same result may be achieved by sub-setting using a vector of logical values:

```
> vec <- (srdta@phdata$age >= 65)
> table(vec)

vec
FALSE  TRUE
2450    50

> summary(srdta@gtdata[vec, 1:3])

  NoMeasured CallRate      Q.2 P.11 P.12 P.22     Pexact      Fmax
rs10          48    0.96 0.1354167   36    11     1 1.0000000  0.02131603
rs18          47    0.94 0.2765957   25    18     4 0.7245853  0.04298643
rs29          45    0.90 0.1555556   32    12     1 1.0000000 -0.01503759

  Plrt Chromosome
rs10 0.8843626      1
rs18 0.7697067      1
rs29 0.9188943      1
```

or a list with IDs of study subjects:

```

> vec1 <- srdta@gtdata@idnames[vec]
> vec1

[1] "p64"    "p122"   "p186"   "p206"   "p207"   "p286"   "p385"   "p386"   "p492"
[10] "p514"   "p525"   "p536"   "p545"   "p565"   "p613"   "p632"   "p649"   "p673"
[19] "p701"   "p779"   "p799"   "p981"   "p1008"  "p1131"  "p1186"  "p1223"  "p1281"
[28] "p1383"  "p1471"  "p1489"  "p1501"  "p1565"  "p1584"  "p1673"  "p1679"  "p1782"
[37] "p1821"  "p1832"  "p1866"  "p1891"  "p1953"  "p2081"  "p2085"  "p2140"  "p2224"
[46] "p2268"  "p2291"  "p2384"  "p2420"  "p2453"

> summary(srdta@gtdata[vec1, 1:3])

      NoMeasured CallRate      Q.2 P.11 P.12 P.22     Pexact      Fmax
rs10          48    0.96 0.1354167    36    11     1 1.0000000  0.02131603
rs18          47    0.94 0.2765957    25    18     4 0.7245853  0.04298643
rs29          45    0.90 0.1555556    32    12     1 1.0000000 -0.01503759

      Plrt Chromosome
rs10 0.8843626      1
rs18 0.7697067      1
rs29 0.9188943      1

```

Let us explore the object returned by `summary` function when applied to `snp.data` class in more details:

```

> a <- summary(srdta@gtdata[vec1, 1:3])
> class(a)

[1] "data.frame"

```

Thus, the object returned is a `data.frame`. Therefore it should have dimensions and names:

```

> dim(a)

[1] 3 10

> names(a)

[1] "NoMeasured" "CallRate"    "Q.2"        "P.11"        "P.12"
[6] "P.22"       "Pexact"      "Fmax"       "Plrt"       "Chromosome"

```

Indeed, we derived 8 characteristics ("NoMeasured", "CallRate", "Q.2", "P.11", "P.12", "P.22", "Pexact", "Chromosome") for the first 3 SNPs.

**Exercise 27** *Test if Hardy-Weinberg equilibrium holds for the first 10 SNPs*

1. Total sample
2. In cases (`bt` is 1)
3. In controls (`bt` is 0)

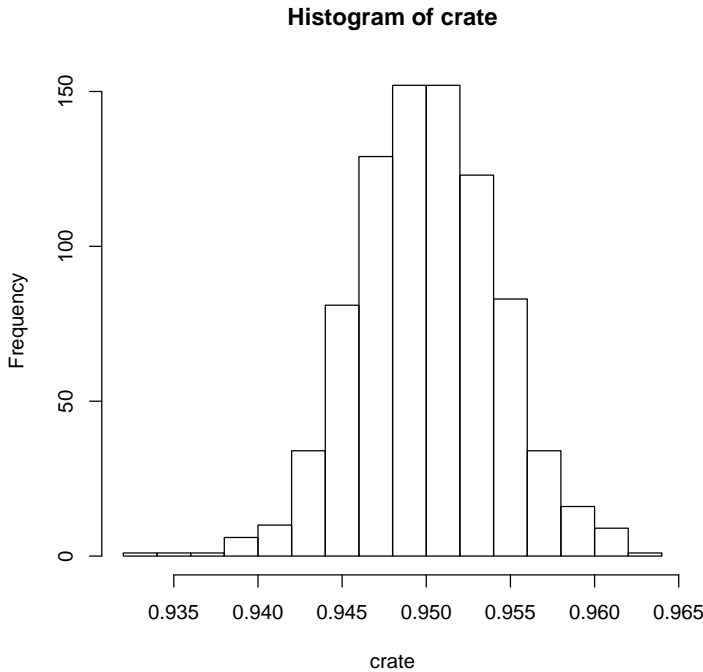


Figure 7: Histogram of the call rate

Let us analyse the distribution of call rate in the whole study. For this, we first need to obtain the vector of call rates:

```
> sumgt <- summary(srdata@gtdata)
> crate <- sumgt[, "CallRate"]
```

This vector may be presented by a histogram

```
> hist(crate)
```

which shows that most SNPs have call rate between 93 and 97% (figure 7).

As next step, you would like to produce a summary table, showing how many markers had call rate lower than, say, 93%, between 93 and 95%, between 95 and 99% and more than 99%. You can use `catable()` command for that:

```
> catable(crate, c(0.93, 0.95, 0.99))

X<=0.93 0.93<X<=0.95 0.95<X<=0.99 X>0.99
No      0        415.000     418.000      0
Prop     0        0.498       0.502       0
```

Similar procedure may be applied to see deviation from HWE:

```
> hwp <- sumgt[, "Pexact"]
> catable(hwp, c((0.05/srdata@nsnps), 0.01, 0.05, 0.1))
```

```

X<=6.00240096038415e-05 6.00240096038415e-05<X<=0.01 0.01<X<=0.05
No 2.000 7.000 23.000
Prop 0.002 0.008 0.028
0.05<X<=0.1 X>0.1
No 31.000 770.000
Prop 0.037 0.924

```

The first cut-off category will detect SNPs which are deviating from HWE at the Bonferroni-corrected P-level.

However, for these data it will make more sense to table cumulative distribution:

```

> catable(hwp, c((0.05/srdta@gtdata@nsnps), 0.01, 0.05, 0.1), cum = T)

X<=6.00240096038415e-05 X<=0.01 X<=0.05 X<=0.1 all X
No 2.000 9.000 32.000 63.000 833
Prop 0.002 0.011 0.038 0.076 1

```

If you would like to investigate the minor allele frequency (MAF) distribution, the same logic would apply. First, derive MAF with

```

> afr <- sumgt[, "Q.2"]
> maf <- pmin(afr, (1 - afr))

```

Next, generate histograms for frequency and MAF:

```

> par(mfcol = c(2, 1))
> hist(afr)
> hist(maf)

```

(shown at the figure 8) and then generate table describing frequency distribution:

```

> catable(afr, c(0.01, 0.05, 0.1, 0.2, 0.5, 0.8, 0.9, 0.95, 0.99))

X<=0.01 0.01<X<=0.05 0.05<X<=0.1 0.1<X<=0.2 0.2<X<=0.5 0.5<X<=0.8
No 22.000 53.000 99.000 132.000 313.000 187.000
Prop 0.026 0.064 0.119 0.158 0.376 0.224
0.8<X<=0.9 0.9<X<=0.95 0.95<X<=0.99 X>0.99
No 18.000 8.00 1.000 0
Prop 0.022 0.01 0.001 0

> catable(maf, c(0, 0.01, 0.05, 0.1, 0.2), cum = T)

X<=0 X<=0.01 X<=0.05 X<=0.1 X<=0.2 all X
No 0 22.000 76.000 183.00 333.0 833
Prop 0 0.026 0.091 0.22 0.4 1

```

Note that we used "0" as the first category – this will give you the number of monomorphic SNPs which we recommend to exclude from analysis.

Other function, `perid.summary`, produces summary SNP statistics per person. Let us try producing this summary for the first 10 people:

```

> perid.summary(srdta[1:10, ])

```

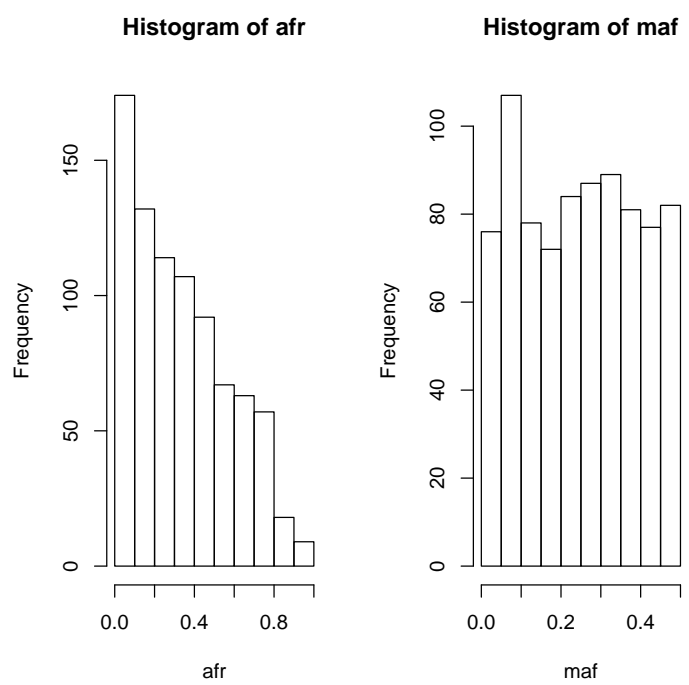


Figure 8: Histogram of the call rate

	NoMeasured	CallPP	Het
p1	790	0.9483794	0.2012658
p2	792	0.9507803	0.2525253
p3	783	0.9399760	0.3793103
p4	789	0.9471789	0.3929024
p5	790	0.9483794	0.3341772
p6	787	0.9447779	0.2337992
p7	794	0.9531813	0.3690176
p8	793	0.9519808	0.2976040
p9	788	0.9459784	0.3324873
p10	797	0.9567827	0.3412798

This table lists the number of genotypes scored for the person, call rate, and heterozygosity. The outliers who have increased average heterozygosity may be suggestive of contaminated DNA samples.

Let us analyse the distribution of heterozygosity:

```
> het <- perid.summary(srdata)$Het
> mean(het)

[1] 0.3309457

> catable(het, c(0.1, 0.25, 0.3, 0.35, 0.5))

X<=0.1 0.1<X<=0.25 0.25<X<=0.3 0.3<X<=0.35 0.35<X<=0.5 X>0.5
No    7.000      73.000     339.000    1281.000      800.00      0
Prop   0.003      0.029      0.136      0.512      0.32        0

> plot(het)
```

The resulting histogram is presented in figure 9. It is easy to see that few people have very low heterozygosity, but there are no outliers with extremely high values.

In this section, we covered low-level functions `summary` and `perid.summary`. Base on these, an upper-level genetic data quality control function, `check.marker`, is based. That function will be covered in the next section.

### Summary:

- When `summary()` function is applied to an object of `snp.data-class`, it return summary statistics for SNPs, including exact test for Hardy-Weinberg equilibrium.
- When `perid.summary()` function is applied to an object of `snp.data-class`, it return per-person summary statistics, including the call rate within this person and its' heterozygosity.

**Histogram of het**

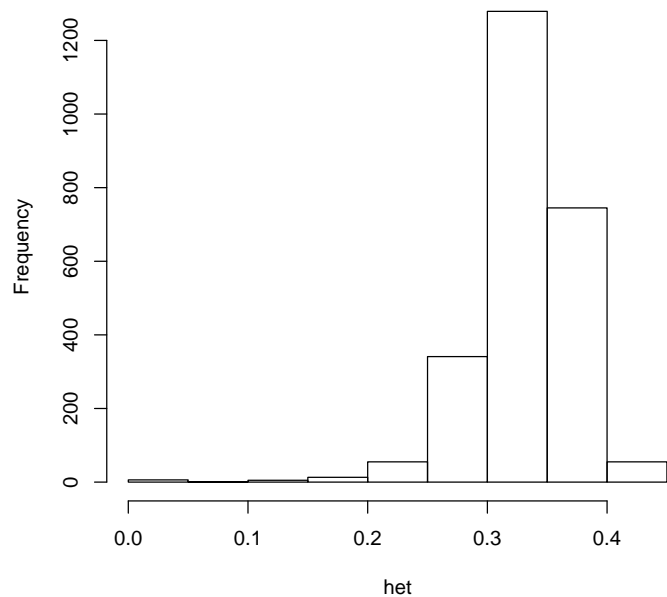


Figure 9: Histogram of heterozygosity

**Exercise 28** Characterise the distribution of call rates within study subjects and produce a histogram. How many people have call rate below 93%?

## 5 Genome-wide association analysis

In the first parts of this section you will be guided through a GWA analysis of a small data set. In the last part you will investigate a larger data set by yourself, do a verification study and will answer the questions. All data sets used assume a study in a relatively homogeneous population. Try to finish the first part in the morning and the second part in the afternoon.

Though only few thousands of markers located at four small chromosomes are used in the scan, we still going to call it Genome-Wide (GW), as the amount of data we will use is approaches the amount to be expected in a real experiment. However, because the regions are small, and the LD between SNPs is high, some specific features (e.g. relatively high residual inflation, which occurs because large proportion of SNPs are in LD with the really associated ones) are specific features of this data set, which are not observed in true GWA studies.

Start R and load GenABEL library by typing

```
> library(GenABEL)
```

and load the data which we will use in this section by

```
> data(ge03d2ex)
```

Investigate the objects loaded by command

```
> ls()
```

```
[1] "ge03d2ex"
```

The ge03d2ex is an object of the class gwaa.data:

```
> class(ge03d2ex)
```

```
[1] "gwaa.data"
attr(,"package")
[1] "GenABEL"
```

To check what are the names of variables in the phenotypic data frame, use

```
> names(ge03d2ex@phdata)
```

```
[1] "id"      "sex"     "age"     "dm2"     "height"   "weight"   "diet"     "bmi"
```

We can attach this data frame to the R search path by

```
> attach(ge03d2ex@phdata)
```

## 5.1 Data descriptives and first round of GWA analysis

Let us investigate what are the traits presented in the data frame loaded and what are the characteristics of the distribution by using specific `GenABEL` function `descriptive.trait`:

```
> descriptives.trait(ge03d2ex)
```

	No	Mean	SD
id	136	NA	NA
sex	136	0.529	0.501
age	136	49.069	12.926
dm2	136	0.632	0.484
height	135	169.440	9.814
weight	135	87.397	25.510
diet	136	0.059	0.236
bmi	135	30.301	8.082

You can see that phenotypic frame contains the data on 136 people; the data on sex, age, height, weight, diet and body mass index (BMI) are available. Our trait of interest is `dm2` (type 2 diabetes). Note that every single piece of information in this data set is simulated; however, we tried to keep our simulations in a way we think the control of T2D may work.

You can produce a summary for cases and controls separately and compare distributions of the traits by

```
> descriptives.trait(ge03d2ex, by = dm2)
```

	No(by.var=1)	Mean	SD	No(by.var=0)	Mean	SD	Ptt	Pkw
id	86	NA	NA	50	NA	NA	NA	NA
sex	86	0.593	0.494	50	0.420	0.499	0.053	0.052
age	86	50.250	12.206	50	47.038	13.971	0.179	0.205
dm2	86	NA	NA	50	NA	NA	NA	NA
height	86	170.448	10.362	49	167.671	8.586	0.097	0.141
weight	86	93.587	27.337	49	76.534	17.441	0.000	0.000
diet	86	0.058	0.235	50	0.060	0.240	0.965	0.965
bmi	86	32.008	8.441	49	27.304	6.463	0.000	0.001
Pexact								
id	NA							
sex	0.074							
age	NA							
dm2	NA							
height	NA							
weight	NA							
diet	1.000							
bmi	NA							

Here, the `by` argument specifies the grouping variable. You can see that cases and controls are different in weight, which is expected, as T2D is associated with obesity.

Similarly, you can produce grand GW descriptives of the marker data by using

```

> descriptives.marker(ge03d2ex)

$`Minor allele frequency distribution`
  X<=0.01 0.01<X<=0.05 0.05<X<=0.1 0.1<X<=0.2      X>0.2
No    146.000      684.000      711.000      904.000  1555.000
Prop   0.036        0.171        0.178        0.226      0.389

$`Cumulative distr. of number of SNPs out of HWE, at different alpha`
  X<=1e-04 X<=0.001 X<=0.01 X<=0.05 all X
No     46.000     71.000   125.000   275.000    4000
Prop   0.011     0.018     0.031     0.069      1

$`Distribution of porportion of successful genotypes (per person)`
  X<=0.9 0.9<X<=0.95 0.95<X<=0.98 0.98<X<=0.99 X>0.99
No     1.000       0           0       135.000      0
Prop   0.007       0           0       0.993      0

$`Distribution of porportion of successful genotypes (per SNP)`
  X<=0.9 0.9<X<=0.95 0.95<X<=0.98 0.98<X<=0.99 X>0.99
No     37.000      6.000     996.000   1177.000  1784.000
Prop   0.009      0.002      0.249      0.294     0.446

$`Mean heterozygosity for a SNP`
[1] 0.2582298

$`Standard deviation of the mean heterozygosity for a SNP`
[1] 0.1592255

$`Mean heterozygosity for a person`
[1] 0.2476507

$`Standard deviation of mean heterozygosity for a person`
[1] 0.04291038

```

It is of note that we can see inflation of the proportion of the tests for HWE at particular threshold, as compared to the expected. This may indicate poor genotyping quality and/or genetic stratification.

We can test the GW marker characteristics in controls by

```

> descriptives.marker(ge03d2ex, ids = (dm2 == 0))

$`Minor allele frequency distribution`
  X<=0.01 0.01<X<=0.05 0.05<X<=0.1 0.1<X<=0.2      X>0.2
No    233.000      676.000      671.000      898.000  1522.000
Prop   0.058        0.169        0.168        0.225      0.381

$`Cumulative distr. of number of SNPs out of HWE, at different alpha`
  X<=1e-04 X<=0.001 X<=0.01 X<=0.05 all X
No         0     3.000    14.000    98.000    4000
Prop       0     0.001    0.003    0.025      1

```

```
$`Distribution of porportion of successful genotypes (per person)`
X<=0.9 0.9<X<=0.95 0.95<X<=0.98 0.98<X<=0.99 X>0.99
No      0          0          0         50          0
Prop    0          0          0         1          0

$`Distribution of porportion of successful genotypes (per SNP)`
X<=0.9 0.9<X<=0.95 0.95<X<=0.98 0.98<X<=0.99 X>0.99
No    37.000     49.000    1523.000      0 2391.000
Prop   0.009      0.012      0.381      0      0.598

$`Mean heterozygosity for a SNP`
[1] 0.2555009

$`Standard deviation of the mean heterozygosity for a SNP`
[1] 0.1618707

$`Mean heterozygosity for a person`
[1] 0.2525720

$`Standard deviation of mean heterozygosity for a person`
[1] 0.04714886
```

Apparently, HWE distribution holds better in controls than in the total sample.

Let us check whether there are indications that deviation from HWE is due to cases. At this stage we are only interested in HWE distribution table, and therefore will ask to report only table two:

```
> descriptives.marker(ge03d2ex, ids = (dm2 == 1))[2]

$`Cumulative distr. of number of SNPs out of HWE, at different alpha`
X<=1e-04 X<=0.001 X<=0.01 X<=0.05 all X
No      45.000     79.00 136.000 268.000 4000
Prop    0.011      0.02   0.034   0.067    1
```

It seems that indeed excessive number of markers are out of HWE in cases. If no laboratory procedure (e.g. DNA extraction, genotyping, calling) were done for cases and controls separately, this may indicate possible heterogeneity specific for cases.

It may be interesting to plot a  $\chi^2 - \chi^2$  plot contrasting observed and expected distributions for the test for HWE in cases. First, we need to compute summary SNP statistics by

```
> s <- summary(ge03d2ex@gtda[(dm2 == 1), ])
```

Note the you have produced the summary for the gtda slot of ge03d2ex; this is the slot which actually contain all genetic data in special compressed format.

You can see first 10 elements of this very long table by

```
> s[1:10, ]
```

NoMeasured	CallRate	Q.2	P.11	P.12	P.22	Pexact
rs7435137	84 0.9767442	0.52380952	17	46	21	0.510978370
rs7725697	85 0.9883721	0.01176471	83	2	0	1.0000000000
rs664063	86 1.0000000	0.08720930	71	15	0	1.0000000000
rs4670072	60 0.6976744	0.11666667	53	0	7	0.001701645
rs546570	84 0.9767442	0.89880952	1	15	68	1.0000000000
rs7908680	83 0.9651163	0.03012048	78	5	0	1.0000000000
rs166732	83 0.9651163	0.04216867	76	7	0	1.0000000000
rs4257079	86 1.0000000	0.07558140	73	13	0	1.0000000000
rs5150804	84 0.9767442	0.39880952	31	39	14	0.820496827
rs3508821	83 0.9651163	0.20481928	52	28	3	1.0000000000
Fmax	Plrt	Chromosome				
rs7435137	-0.09772727	0.3699602726		1		
rs7725697	-0.01190476	0.8773691192		3		
rs664063	-0.09554140	0.2308999066		2		
rs4670072	1.000000000	0.0002510899		X		
rs546570	0.01830931	0.8693645189		2		
rs7908680	-0.03105590	0.6935168932		1		
rs166732	-0.04402516	0.5787401988		1		
rs4257079	-0.08176101	0.3022863648		1		
rs5150804	0.03177183	0.7710793180		2		
rs3508821	-0.03565062	0.7419587406		2		

Note that the column before the last provides P-exact we need. We can extract these to a separate vector by

```
> pexcas <- s[, "Pexact"]
```

and produce the  $\chi^2 - \chi^2$  plot and estimate inflation factor by command `estlambda()`, which operates with a vector of P-values or  $\chi^2$ s:

```
> estlambda(pexcas)
```

```
$estimate
```

```
[1] 1.068184
```

```
$se
```

```
[1] 0.02614764
```

By default, this function also produces a  $\chi^2 - \chi^2$  plot, at which you can see some extreme deviation of observed from expected. The resulting plot (figure 10) shows extreme deviation for high values of the test. Looking at the  $\lambda$  estimate, we indeed see inflation of the test statistics.

You can repeat this test for the controls, if time permits.

The 'se' produced by `estlambda` *can not* be used to test if inflation is significant and make conclusions about presence of stratification.

Let us first try do GWA scan using raw (before quality control) data. We will use the score test, as implemented in the `qtscore()` function of `GenABEL` for testing:

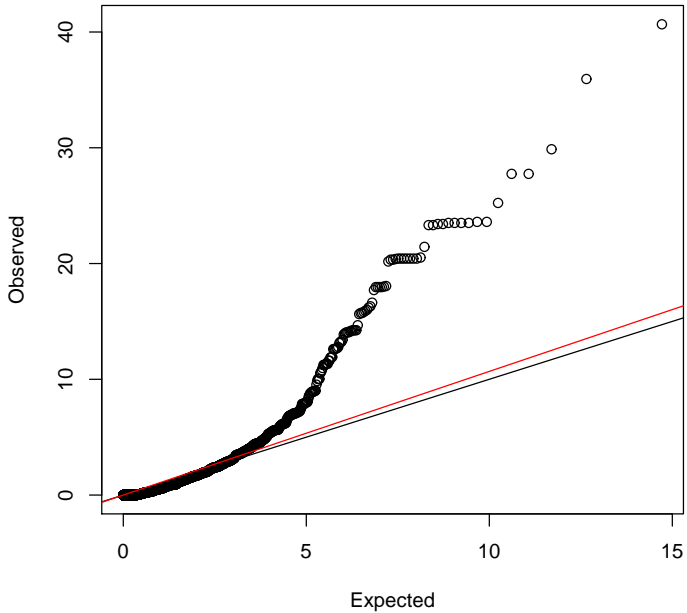


Figure 10:  $\chi^2 - \chi^2$  plot for the exact test for HWE. Black line of slope 1: expected under no inflation; Red line: fitted slope.

```
> an0 <- qtscore(dm2, ge03d2ex, trait = "binomial")
```

The first argument used describes the model; here it is rather simple — the affection status, `dm2`, is supposed to depend on SNP genotype only.

You can see what objects are returned by this function by using

```
> names(an0)

[1] "chi2.1df"    "chi2.2df"    "P1df"        "P2df"        "Pc1df"
[6] "Pc2df"       "lambda"      "effB"        "effAB"       "effBB"
[11] "N"           "snppnames"   "idnames"     "map"         "chromosome"
[16] "formula"     "family"
```

Here, `P1df`, `P2df` and `Pc1df` are most interesting; the first two are vectors of 1 and 2 d.f. P-values obtained in the GWA analysis, the last one is 1 d.f. P-value corrected for inflation factor  $\lambda$  (which is presented in `lambda` object).

Let us see if there is evidence for the inflation of the test statistics

```
> an0$lambda
```

```
$estimate
[1] 1.047846
```

```
$se
```

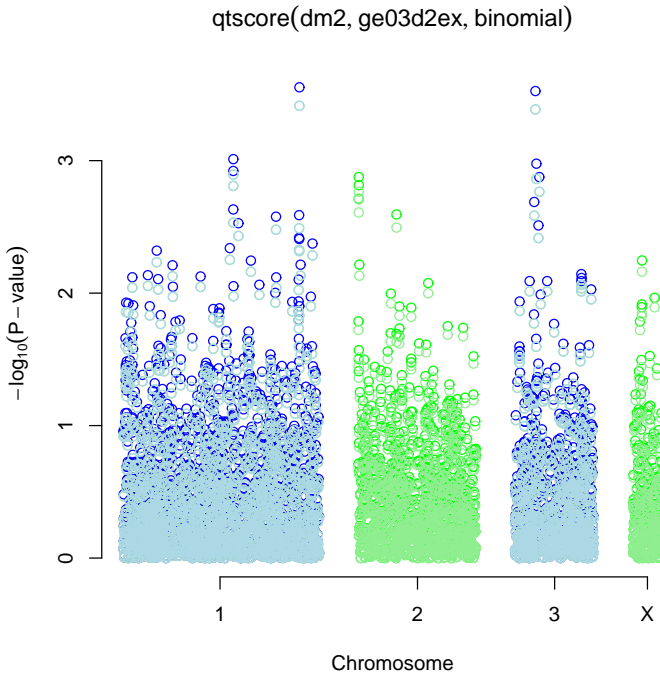


Figure 11:  $-\log_{10}(P\text{-value})$  from the genome scan before QC procedure. Raw analysis: black; corrected analysis: green

```
[1] 0.0009700066
```

```
$iz0
[1] 1.002987
```

```
$iz2
[1] 1
```

The estimate of  $\lambda$  is 1.05, suggesting inflation of the test and some degree of stratification.

We can plot the results of analysis by

```
> plot(an0)
```

The resulting plot is presented in the figure 11. By default,  $-\log_{10}(P\text{-value})$  on 1 d.f. are presented; see help to figure out how this behaviour can be changed.

We can also add the corrected P-values to the plot with

```
> add.plot(an0, df = "Pc1df", col = c("lightblue", "lightgreen"))
```

You can also generate a descriptive table for the "top" (as ranked by P-value) results by

```
> descriptives.scan(an0)
```

	Chromosome	Position	N	effB	P1df	Pc1df
rs1719133	1	4495479	136	0.33729339	0.0002795623	0.0003857992
rs2975760	3	10518480	134	3.80380024	0.0002983731	0.0004106050
rs7418878	1	2808520	136	3.08123060	0.0009743183	0.0012744665
rs5308595	3	10543128	133	3.98254950	0.0010544366	0.0013746240
rs4804634	1	2807417	132	0.43411456	0.0011970132	0.0015520826
rs3224311	2	6009769	135	3.15831710	0.0013290907	0.0017156757
rs26325	3	10617781	135	0.09742793	0.0013313876	0.0017185144
rs8835506	2	6010852	132	3.17720829	0.0015321522	0.0019658784
rs3925525	2	6008501	135	2.98416931	0.0019400358	0.0024644132
rs2521089	3	10487652	135	2.50239493	0.0020524092	0.0026009460
	effAB	effBB	P2df			
rs1719133	0.4004237	0.0000000	0.0006333052			
rs2975760	3.4545455	10.0000000	0.0011434877			
rs7418878	3.6051282	4.8717949	0.0022642036			
rs5308595	3.3171429	Inf	0.0045930101			
rs4804634	0.5240642	0.1739130	0.0036964462			
rs3224311	3.4151786	4.2500000	0.0029405999			
rs26325	0.1097724	NA	0.0013313876			
rs8835506	3.4903846	4.1250000	0.0031618340			
rs3925525	3.2380952	4.1212121	0.0045554384			
rs2521089	2.5717703	4.7727273	0.0069661425			

Here you see top 10 results, sorted by P-value with 1 d.f. If you want to sort by the corrected P-value, you can use `descriptives.scan(an0,sort="Pc1df")`; to see more than 10 (e.g. 25) top results, use `descriptives.scan(an0,top=25)`. You can combine all these options. Large part of results reports NA as effect estimates and 9.99 as *P-value* for 2 d.f. test – for these markers only two out of three possible genotypes were observed, and consequently 2 d.f. test could not be performed.

Now let us apply `emp.qtscore()` function, which computes empirical GW (or experiment-wise) significance

```
> an0.e <- emp.qtscore(dm2, ge03d2ex)

100%

> descriptives.scan(an0.e, sort = "Pc1df")



|           | Chromosome | Position | N   | effB        | P1df | Pc1df | effAB      | effBB      |
|-----------|------------|----------|-----|-------------|------|-------|------------|------------|
| rs1719133 | 1          | 4495479  | 136 | -0.26520640 | 0.40 | 0.53  | -0.2080882 | -0.7375000 |
| rs2975760 | 3          | 10518480 | 134 | 0.23406549  | 0.44 | 0.56  | 0.2755102  | 0.4090909  |
| rs5308595 | 3          | 10543128 | 133 | 0.24455163  | 0.88 | 0.91  | 0.2564832  | 0.4623656  |
| rs7418878 | 1          | 2808520  | 136 | 0.20890976  | 0.88 | 0.91  | 0.2807405  | 0.3268398  |
| rs4804634 | 1          | 2807417  | 132 | -0.20504486 | 0.91 | 0.93  | -0.1193830 | -0.3845238 |
| rs3224311 | 2          | 6009769  | 135 | 0.21336328  | 0.91 | 0.95  | 0.2778634  | 0.3151515  |
| rs26325   | 3          | 10617781 | 135 | -0.48753666 | 0.91 | 0.95  | -0.4875367 | NA         |
| rs8835506 | 2          | 6010852  | 132 | 0.21120000  | 0.93 | 0.97  | 0.2796221  | 0.3076923  |
| rs3925525 | 2          | 6008501  | 135 | 0.20570949  | 0.97 | 0.99  | 0.2660834  | 0.3074627  |
| rs7435137 | 1          | 4259040  | 134 | 0.07345101  | 1.00 | 1.00  | 0.1166373  | 0.1461694  |
|           | P2df       |          |     |             |      |       |            |            |


```

```

rs1719133 0.74
rs2975760 0.85
rs5308595 1.00
rs7418878 0.98
rs4804634 1.00
rs3224311 1.00
rs26325   1.00
rs8835506 1.00
rs3925525 1.00
rs7435137 1.00

```

None of the SNPs hits GW significance. If, actually, any did pass the threshold, we could not trust the results, because the distribution of the HWE test and presence of inflation factor for the association test statistics suggest that the data may contain multiple errors (indeed they do). Therefore before association analysis we need to do rigorous Quality Control (QC).

Note that at certain SNP, corrected *P-values* become equal to 1 – at this point order is arbitrary because sorting could not be done.

#### Summary:

- The **descriptives** family of functions was developed to facilitate production of tables which can be directly used in a manuscript — it is possible to save the output as a file, which can be open by Excel or Word. See e.g. `help(descriptives.trait)` for details.
- The inflation of test statistics compared to null (1 d.f.) may be estimated with `estlambda` function.

## 5.2 Genetic data QC

The major genetic data QC function of GenABEL is `check.marker()`. We will try to run it; the output is rather self-explaining. As it was detailed at the lecture, in the first round of the QC we do not want to check for HWE. This can be achieved by setting HWE P-value selection threshold to zero (`p.level=0`):

```

> qc1 <- check.marker(ge03d2ex, p.level = 0)

Excluding people/markers with extremely low call rate...
4000 markers and 136 people in total
0 people excluded because of call rate < 0.1
6 markers excluded because of call rate < 0.1
Passed: 3994 markers and 136 people

Running sex chromosome checks...
197 heterozygous X-linked male genotypes found
1 X-linked markers are likely to be autosomal (odds > 1000 )
2 male are likely to be female (odds > 1000 )
0 female are likely to be male (odds > 1000 )

```

If these people/markers are removed, 0 heterozygous male genotypes are left  
Passed: 3993 markers and 134 people

no X/Y/mtDNA-errors to fix

RUN 1

3993 markers and 134 people in total  
304 (7.613323%) markers excluded as having low (<1.865672%) minor allele frequency  
36 (0.9015778%) markers excluded because of low (<95%) call rate  
0 (0%) markers excluded because they are out of HWE (P <0)  
1 (0.7462687%) people excluded because of low (<95%) call rate  
Mean autosomal HET is 0.2747262 (s.e. 0.03721277)  
3 (2.238806%) people excluded because too high autosomal heterozygosity (FDR <1%)  
Excluded people had HET >= 0.4856887  
Mean IBS is 0.7724447 (s.e. 0.02098092), as based on 2000 autosomal markers  
1 (0.7462687%) people excluded because of too high IBS (>=0.95)  
In total, 3653 (91.4851%) markers passed all criteria  
In total, 129 (96.26866%) people passed all criteria

RUN 2

3653 markers and 129 people in total  
72 (1.970983%) markers excluded as having low (<1.937984%) minor allele frequency  
0 (0%) markers excluded because of low (<95%) call rate  
0 (0%) markers excluded because they are out of HWE (P <0)  
0 (0%) people excluded because of low (<95%) call rate  
Mean autosomal HET is 0.2744972 (s.e. 0.01706096)  
0 people excluded because too high autosomal heterozygosity (FDR <1%)  
Mean IBS is 0.7709758 (s.e. 0.01861282), as based on 2000 autosomal markers  
0 (0%) people excluded because of too high IBS (>=0.95)  
In total, 3581 (98.02902%) markers passed all criteria  
In total, 129 (100%) people passed all criteria

RUN 3

3581 markers and 129 people in total  
0 (0%) markers excluded as having low (<1.937984%) minor allele frequency  
0 (0%) markers excluded because of low (<95%) call rate  
0 (0%) markers excluded because they are out of HWE (P <0)  
0 (0%) people excluded because of low (<95%) call rate  
Mean autosomal HET is 0.2744972 (s.e. 0.01706096)  
0 people excluded because too high autosomal heterozygosity (FDR <1%)  
Mean IBS is 0.7694881 (s.e. 0.01830812), as based on 2000 autosomal markers  
0 (0%) people excluded because of too high IBS (>=0.95)  
In total, 3581 (100%) markers passed all criteria  
In total, 129 (100%) people passed all criteria

The computation of all pairwise proportion of alleles identical-by-state (IBS) by ibs() function, which is also called by check.markers() may take quite some

time, which is proportional to the square of the number of subjects. This is not a problem with the small number of people we use for this example or when modern computers are used. However, the computers in the Nihes computer room are very old. Therefore be prepared to wait for long time when you will do a self-exercise with 1,000 people.

From the output you can see that QC starts with checking the data for SNPs and people with extremely low call rate. Six markers are excluded from further analysis due to very low call rate. Next, X-chromosomal errors are identified. The function finds out that all errors (heterozygous male X-genotypes) are due to two people with wrong sex assigned and one marker, which looks like an autosomal one. This actually could be a marker from pseudoautosomal region, which should have been arranged as a separate "autosome".

Then, the procedure finds the markers with low call rate ( $\leq 0.95$ ) across people, markers with low MAF (by default, low MAF is defined as less than few copies of the rare allele, see help for details); people with low call rate ( $\leq 0.95$ ) across SNPs, people with extreme heterozygosity (at FDR 0.01) and these who have GW IBS  $\geq 0.95$ . These default parameters may be changed if you wish (consult help).

Because some of the people fail to pass the tests, the data set is not guaranteed to be really "clean" after single iteration, e.g. some marker may not pass the call threshold after we exclude few informative (but apparently wrong) people. Therefore the QC is repeated iteratively until no further errors are found.

You can generate short summary of QC by marker and by person through

```
> summary(qc1)

$`Per-SNP fails statistics`
      NoCall NoMAF NoHWE Redundant Xsnpfail
NoCall      42     0     0       0       0
NoMAF      NA   376     0       0       0
NoHWE      NA     NA     0       0       0
Redundant    NA     NA     NA      0       0
Xsnpfail    NA     NA     NA      NA      1

$`Per-person fails statistics`
      IDnoCall HetFail IBSFail isfemale ismale isXXY
IDnoCall      1      0      0       0       0       0
HetFail      NA      3      0       0       0       0
IBSFail      NA     NA      1       0       0       0
isfemale     NA     NA     NA      2       0       0
ismale       NA     NA     NA      NA      0       0
isXXY        NA     NA     NA      NA      NA      0
```

Note that the original data, `ge03d2ex`, are not modified during the procedure; rather, `check.markers()` generate a list of markers and people which pass or do not pass certain QC criteria. The objects returned by `check.markers()` are:

```
> names(qc1)
```

```
[1] "nofreq"   "nocall"    "nohwe"      "Xmrkfail"  "hetfail"    "idnocall"
[7] "ibsfail"  "isfemale"   "ismale"     "snpok"      "idok"       "call"
```

The element `idok` provides the list of people who passed all QC criteria, and `snpok` provides the list of SNPs which passed all criteria. You can easily generate a new data set, which will consist only of these people and markers by

```
> data1 <- ge03d2ex[qc1$idok, qc1$snpok]
```

If there are any residual sporadic X-errors (male heterozygosity), these can be fixed (set to NA) by

```
> data1 <- Xfix(data1)
```

```
no X/Y/mtDNA-errors to fix
```

Applying this function does not make any difference for the example data set, but you will need to use it for the bigger data set.

At this point, we are ready to work with the new, cleaned, data set `data1`. However, if we try

```
> length(dm2)
```

```
[1] 136
```

we can see that the original phenotypic data are attached to the search path (there are only 129 people left in the 'clean' data set). Therefore we need to detach the data by

```
> detach(ge03d2ex@phdata)
```

and attach new data by

```
> attach(data1@phdata)
```

At this stage, let us check if the first round of QC solves the problem of inflated test for HWE, which may be the case if this inflation is due to genotypic errors we managed to eliminate:

```
> descriptives.marker(data1)[2]

$`Cumulative distr. of number of SNPs out of HWE, at different alpha`
  X<=1e-04 X<=0.001 X<=0.01 X<=0.05 all X
  No      43.000   65.000 121.000 240.000  3581
  Prop     0.012    0.018   0.034   0.067     1

> descriptives.marker(data1[dm2 == 1])[2]

$`Cumulative distr. of number of SNPs out of HWE, at different alpha`
  X<=1e-04 X<=0.001 X<=0.01 X<=0.05 all X
  No      46.000   70.00 127.000 228.000  3581
  Prop     0.013    0.02   0.035   0.064     1

> estlambda(summary(data1@gtdata[dm2 == 1, ])[, "Pexact"])
```

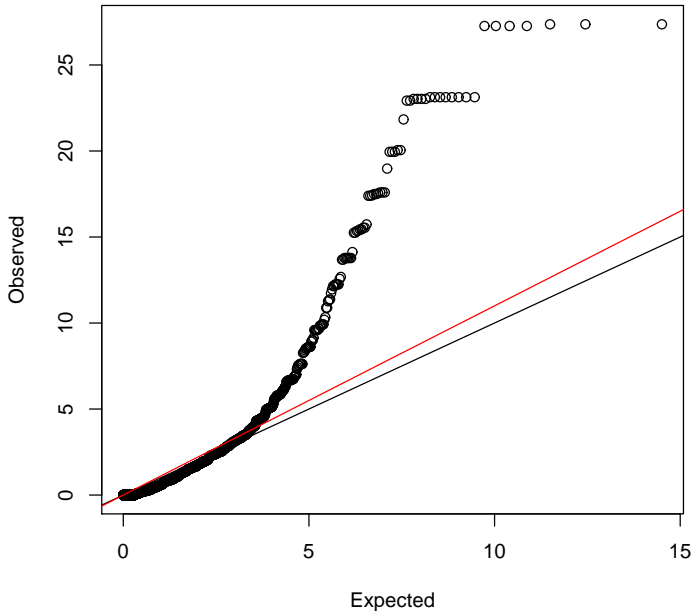


Figure 12:  $\chi^2 - \chi^2$  plot for the exact test for HWE. Black line of slope 1: expected under no inflation; Red line: fitted slope.

```
$estimate
[1] 1.099913

$se
[1] 0.02770376
```

Apparently, the distribution (figure 12) looks better (note the scale difference between the graphs), but the test statistics is still inflated.

### 5.3 Finding genetic sub-structure

Now, we are ready for the second round of QC, detection of genetic outliers which may contaminate our data. We will detect genetic outliers using a technique, which resembles the one suggested by Price et al.

As a first step, we will compute a matrix of genomic kinship between all pairs of people, using only autosomal markers by

```
> data1.gkin <- ibs(data1[, data1@gtdata@chromosome != "X"], weight = "freq")
```

You can see the 5x5 upper left sub-matrix by

```
> data1.gkin[1:5, 1:5]
```

	id199	id300	id403	id415	id666
id199	0.496562976	3262.0000000	3261.0000000	3249.0000000	3265.0000000
id300	-0.012267995	0.49486164	3268.0000000	3257.0000000	3271.0000000
id403	-0.012464681	-0.01262302	0.51836209	3255.0000000	3270.0000000
id415	-0.002399026	0.01363777	-0.02528089	0.56733776	3259.0000000
id666	-0.019116287	-0.02110468	0.02096914	-0.02025885	0.5684808

This step may take few minutes on large data sets or when using old computers!

The numbers below the diagonal show genomic kinship (IBD), the numbers on the diagonal correspond to 0.5 plus the genomic homozigosity, and the numbers above the diagonal tell how many SNPs were typed successfully for both subjects (thus the IBD estimate is derived using this number of SNPs).

Second, we transform this matrix to a distance matrix using standard R command

```
> data1.dist <- as.dist(0.5 - data1.gkin)
```

Finally, we perform Classical Multidimensional Scaling by

```
> data1.mds <- cmdscale(data1.dist)
```

by default, the first two principal components are computed and returned.

This may take few minutes on large data sets or when using old computers!

We can present the results graphically by

```
> plot(data1.mds)
```

The resulting plot is presented in figure 13. Each point on the plot corresponds to a person, and the 2D distances between points were fitted to be as close as possible to these presented in the original IBS matrix. You can see that study subjects clearly cluster in two groups.

You can identify the points belonging to clusters by

```
> km <- kmeans(data1.mds, centers = 2, nstart = 1000)
> cl1 <- names(which(km$cluster == 1))
> cl2 <- names(which(km$cluster == 2))
> cl1
[1] "id199"  "id300"  "id403"  "id415"  "id666"  "id689"  "id765"  "id830"
[9] "id908"  "id980"  "id994"  "id1193" "id1423" "id1505" "id1737" "id1827"
[17] "id1841" "id2068" "id2094" "id2151" "id2317" "id2618" "id2842" "id2894"
[25] "id2985" "id3354" "id3368" "id3641" "id3831" "id3983" "id4097" "id4328"
[33] "id4380" "id4395" "id4512" "id4552" "id4710" "id4717" "id4883" "id4904"
[41] "id4934" "id4961" "id5014" "id5078" "id5274" "id5275" "id5454" "id5853"
```

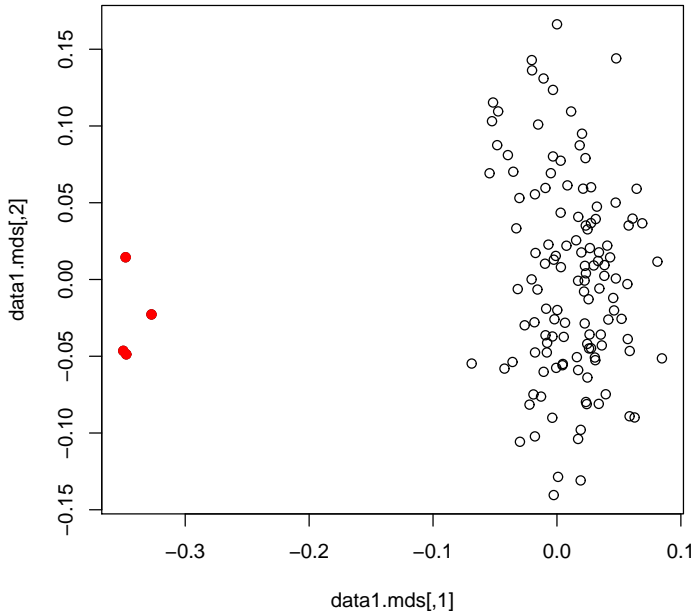


Figure 13: Mapping samples on the space of the first two Principle Components resulting from analysis of genomic kinship. Red dots identify genetic outliers

```
[49] "id5926" "id5969" "id6237" "id6278" "id6352" "id6501" "id6554" "id6663"
[57] "id6723" "id7499" "id7514" "id7541" "id7598" "id7623" "id7949" "id8059"
[65] "id8128" "id8281" "id8370" "id8400" "id8433" "id8772" "id8880" "id8890"
[73] "id8957" "id8996" "id9082" "id9901" "id9930" "id1857" "id2528" "id4862"
[81] "id9184" "id5677" "id6407" "id5472" "id2135" "id8545" "id4333" "id1670"
[89] "id1536" "id6917" "id6424" "id3917" "id9628" "id9635" "id4729" "id5190"
[97] "id6399" "id6062" "id620" "id1116" "id6486" "id41" "id677" "id4947"
[105] "id9749" "id6428" "id7488" "id5949" "id2924" "id5783" "id4096" "id903"
[113] "id9049" "id185" "id1002" "id362" "id9014" "id5044" "id2749" "id5437"
[121] "id2286" "id4743" "id4185" "id8330" "id6934"

> c12
[1] "id2097" "id6954" "id2136" "id858"
```

Four outliers are presented in the smaller cluster.

Now you will need to use the BIGGER cluster for to select study subjects.  
Whether this will be c11 or c12 in you case, is totally random.

We can form a data set which is free from outliers by using only people from the bigger cluster:

```
> data2 <- data1[c12, ]
```

After we dropped the outliers, we need to repeat QC using `check.markers()`. At this stage, we want to allow for HWE checks (we will use only controls and exclude markers with  $FDR \leq 0.2$ ):

```
> qc2 <- check.marker(data2, hweids = (data2@phdata$dm2 == 0),
+      fdr = 0.2)

Excluding people/markers with extremely low call rate...
3581 markers and 125 people in total
0 people excluded because of call rate < 0.1
0 markers excluded because of call rate < 0.1
Passed: 3581 markers and 125 people

Running sex chromosome checks...
0 heterozygous X-linked male genotypes found
0 X-linked markers are likely to be autosomal (odds > 1000 )
0 male are likely to be female (odds > 1000 )
0 female are likely to be male (odds > 1000 )
If these people/markers are removed, 0 heterozygous male genotypes are left
Passed: 3581 markers and 125 people

no X/Y/mtDNA-errors to fix
```

#### RUN 1

```
3581 markers and 125 people in total
40 (1.117006%) markers excluded as having low (<2%) minor allele frequency
0 (0%) markers excluded because of low (<95%) call rate
0 (0%) markers excluded because they are out of HWE (FDR <0.2)
0 (0%) people excluded because of low (<95%) call rate
Mean autosomal HET is 0.2776868 (s.e. 0.01655360)
0 people excluded because too high autosomal heterozygosity (FDR <1%)
Mean IBS is 0.7719071 (s.e. 0.01239049), as based on 2000 autosomal markers
0 (0%) people excluded because of too high IBS (>=0.95)
In total, 3541 (98.883%) markers passed all criteria
In total, 125 (100%) people passed all criteria
```

#### RUN 2

```
3541 markers and 125 people in total
0 (0%) markers excluded as having low (<2%) minor allele frequency
0 (0%) markers excluded because of low (<95%) call rate
0 (0%) markers excluded because they are out of HWE (FDR <0.2)
0 (0%) people excluded because of low (<95%) call rate
Mean autosomal HET is 0.2776868 (s.e. 0.01655360)
0 people excluded because too high autosomal heterozygosity (FDR <1%)
Mean IBS is 0.769734 (s.e. 0.01269325), as based on 2000 autosomal markers
0 (0%) people excluded because of too high IBS (>=0.95)
In total, 3541 (100%) markers passed all criteria
In total, 125 (100%) people passed all criteria
```

```

> summary(qc2)

$`Per-SNP fails statistics`
      NoCall  NoMAF  NoHWE Redundant Xsnpfail
NoCall       0       0       0       0       0
NoMAF      NA     40       0       0       0
NoHWE      NA      NA       0       0       0
Redundant   NA      NA      NA       0       0
Xsnpfail    NA      NA      NA      NA       0

$`Per-person fails statistics`
      IDnoCall HetFail IBSFail isfemale ismale isXXY
IDnoCall       0       0       0       0       0       0
HetFail      NA       0       0       0       0       0
IBSFail      NA      NA       0       0       0       0
isfemale     NA      NA      NA       0       0       0
ismale       NA      NA      NA      NA       0       0
isXXY        NA      NA      NA      NA      NA       0

```

If the procedure did not run, check previous Note.

Indeed, in the updated data set few markers do not pass our QC criteria and we need to drop a few markers. This is done by

```
> data2 <- data2[qc2$idok, qc2$snpok]
```

This is going to be our final analysis data set, therefore let us attach the phenotypic data to the search path, then we do not need to type `data2@phdata$...` to access `dm2` status or other variables:

```
> detach(data1@phdata)
> attach(data2@phdata)
```

## 5.4 GWA association analysis

Let us start again with descriptives of the phenotypic and marker data

```
> descriptives.trait(data2, by = dm2)
```

	No(by.var=1)	Mean	SD	No(by.var=0)	Mean	SD	Ptt	Pkw
id	77	NA	NA	48	NA	NA	NA	NA
sex	77	0.597	0.494	48	0.438	0.501	0.084	0.082
age	77	50.593	12.465	48	46.378	13.865	0.089	0.097
dm2	77	NA	NA	48	NA	NA	NA	NA
height	77	170.423	10.646	47	167.988	8.610	0.166	0.223
weight	77	94.160	26.963	47	77.273	17.427	0.000	0.000
diet	77	0.065	0.248	48	0.062	0.245	0.957	0.957
bmi	77	32.235	8.335	47	27.485	6.539	0.001	0.001
Pexact								

```

id      NA
sex     0.098
age     NA
dm2     NA
height   NA
weight   NA
diet    1.000
bmi     NA

```

You can see that relation to weight is maintained in this smaller, but hopefully cleaner, data set; moreover, relation to age becomes boundary significant.

If you check descriptives of markers (only HWE part shown)

```

> descriptives.marker(data2)[2]

$`Cumulative distr. of number of SNPs out of HWE, at different alpha`
X<=1e-04 X<=0.001 X<=0.01 X<=0.05 all X
No          1    2.000  22.000 108.00 3541
Prop        0    0.001   0.006   0.03    1

```

you can see that the problems with HWE are apparently fixed; we may guess that these were caused by the Wahlund's effect.

Run the score test on the cleaned data by

```
> data2.qt <- qtscore(dm2, data2)
```

and check lambda

```
> data2.qt$lambda
```

```
$estimate
[1] 1.052329
```

```
$se
[1] 0.001500786
```

```
$iz0
[1] 1.018069
```

```
$iz2
[1] 1
```

there is still some inflation, which is explained by the fact that we investigate only few short chromosomes with high LD and few causative variants.

Produce the association analysis plot by

```
> plot(data2.qt, df = "Pc1df")
```

(figure 14).

Produce the scan summary by

```
> descriptives.scan(data2.qt, sort = "Pc1df")
```

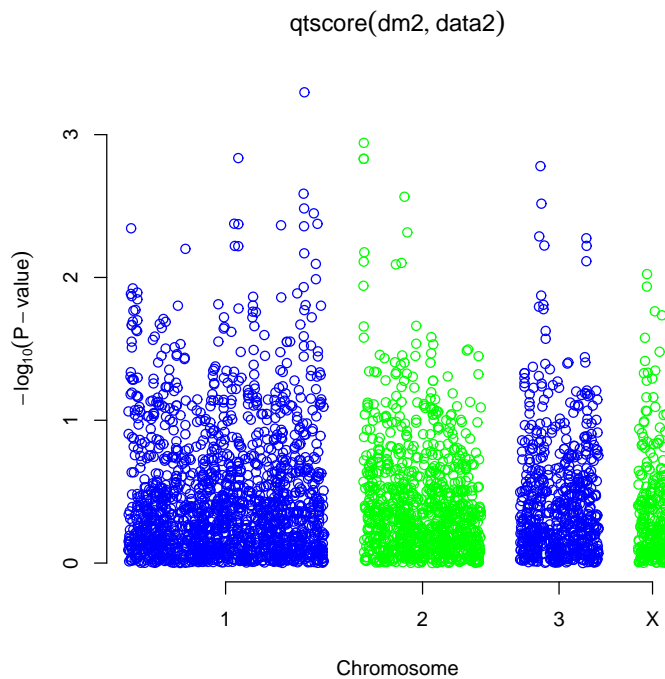


Figure 14:  $-\log_{10}(\text{Corrected } P\text{-value})$  from the genome scan after the QC procedure

	Chromosome	Position	N	effB	P1df	Pc1df
rs1719133	1	4495479	125	-0.2709715	0.0003599745	0.0005052222
rs8835506	2	6010852	122	0.2346446	0.0008465818	0.0011416640
rs4804634	1	2807417	122	-0.2161383	0.0010949169	0.0014590088
rs3925525	2	6008501	125	0.2280847	0.0011081904	0.0014758702
rs3224311	2	6009769	125	0.2280847	0.0011081904	0.0014758702
rs2975760	3	10518480	124	0.2244898	0.0012545339	0.0016611805
rs4534929	1	4474374	124	-0.1864322	0.0019996882	0.0025913669
rs6079246	2	7048058	124	-0.4730491	0.0021060217	0.0027226464
rs5308595	3	10543128	123	0.2731788	0.0023674963	0.0030442101
rs1013473	1	4487262	125	0.1864494	0.0025659783	0.0032872160
				effAB	effBB	P2df
rs1719133		-0.2153891		-0.7260274	0.0009290908	
rs8835506		0.3201507		0.3201507	0.0012972329	
rs4804634		-0.0971660		-0.3947368	0.0026493765	
rs3925525		0.3046200		0.3187614	0.0020102814	
rs3224311		0.3046200		0.3187614	0.0020102814	
rs2975760		0.2570423		0.3959311	0.0047945922	
rs4534929		-0.1487634		-0.3968254	0.0074296050	
rs6079246		-0.4730491		NA	0.0021060217	
rs5308595		0.2844101		0.4719101	0.0095505795	
rs1013473		0.2714833		0.3782654	0.0067821374	

Comparison with the top 10 from the scan before QC shows that results changed substantially with only few markers overlapping.

You can see similar results when accessing empirical GW significance:

```
> data2.qte <- emp.qtscore(dm2, data2)

100%

> descriptives.scan(data2.qte, sort = "Pc1df")

      Chromosome Position   N      effB P1df  Pc1df      effAB      effBB
rs1719133       1 4495479 125 -0.27097152 0.44  0.55 -0.21538910 -0.7260274
rs8835506       2 6010852 122  0.23464458 0.76  0.85  0.32015066  0.3201507
rs4804634       1 2807417 122 -0.21613833 0.82  0.94 -0.09716599 -0.3947368
rs3925525       2 6008501 125  0.22808468 0.82  0.94  0.30461997  0.3187614
rs3224311       2 6009769 125  0.22808468 0.82  0.94  0.30461997  0.3187614
rs2975760       3 10518480 124  0.22448980 0.90  0.95  0.25704225  0.3959311
rs7435137       1 4259040 124  0.08916655 1.00  1.00  0.11073664  0.1761787
rs664063        2 7288020 125 -0.13014211 1.00  1.00 -0.09180036 -0.6372549
rs546570        2 6120257 124 -0.15740741 1.00  1.00  0.75000000  0.5925926
rs7908680       1 2311762 122  0.19316239 1.00  1.00  0.19316239      NA

      P2df
rs1719133  0.64
rs8835506  0.74
rs4804634  1.00
rs3925525  0.91
rs3224311  0.91
rs2975760  1.00
rs7435137  1.00
rs664063   1.00
rs546570   1.00
rs7908680  1.00
```

Again, none of the SNPs hits GW 5% significance. Still, you can see that after QC top markers achieve somewhat "better" significance.

In the last part, we will do several adjusted and stratified analyses. Only empirical P-values will be estimated to make the story shorter. To adjust for sex and age, we can

```
> data2.qtae <- emp.qtscore(dm2 ~ sex + age, data2)

100%

> descriptives.scan(data2.qtae)

      Chromosome Position   N      effB P1df  Pc1df      effAB      effBB
rs1719133       1 4495479 125 -0.25890157 0.59  0.79 -0.22773582 -0.6210215
rs4804634       1 2807417 122 -0.20651613 0.94  0.98 -0.09071397 -0.3764928
rs3925525       2 6008501 125  0.21871930 0.94  0.98  0.30498571  0.2825598
rs8835506       2 6010852 122  0.22215111 0.94  0.96  0.31550163  0.2814815
rs3224311       2 6009769 125  0.21871930 0.94  0.98  0.30498571  0.2825598
rs6079246       2 7048058 124 -0.46876373 0.96  0.99 -0.46876373      NA
```

rs2939190	1	71843	125	-0.21514616	0.99	1.00	-0.27260946	-0.3173458
rs7504607	1	2704056	124	-0.39615225	0.99	1.00	-0.39615225	NA
rs7435137	1	4259040	124	0.06938366	1.00	1.00	0.10716530	0.1349937
rs664063	2	7288020	125	-0.11264399	1.00	1.00	-0.07332386	-0.6118779
			P2df					
rs1719133	0.93							
rs4804634	0.99							
rs3925525	0.92							
rs8835506	0.84							
rs3224311	0.92							
rs6079246	1.00							
rs2939190	1.00							
rs7504607	1.00							
rs7435137	1.00							
rs664063	1.00							

You can see that there is little difference between adjusted and unadjusted analysis, but this is not always the case; adjustment may make your study much more powerful when covariates explain a large proportion of environmental trait variation.

Finally, let us do stratified (by BMI) analysis. We will contract obese ( $BMI \geq 30$ ) cases to all controls.

```
> data2.qtse <- emp.qtscore(dm2 ~ sex + age, data2, ids = ((bmi >
+      30 & dm2 == 1) | dm2 == 0))

100%

> descriptives.scan(data2.qtse, sort = "Pc1df")

      Chromosome Position N          effB P1df  Pc1df      effAB
rs1891586        1 2297398 89 -0.2525525541 0.80  0.87 -0.1844921696
rs794264        1 2534738 88  0.2285092556 0.90  0.96  0.3184520260
rs9178808        1 2536431 89 -0.2265653369 0.92  0.99 -0.1652286996
rs5032886        1 2537020 89 -0.2265653369 0.92  0.99 -0.1652286996
rs7435137        1 4259040 89  0.0810467038 1.00  1.00  0.1298496587
rs664063        2 7288020 89 -0.0649051305 1.00  1.00 -0.0159823043
rs546570        2 6120257 89 -0.1239122686 1.00  1.00  0.1099895418
rs7908680        1 2311762 87  0.2309587412 1.00  1.00  0.2309587412
rs166732        1 4716343 88  0.0078773947 1.00  1.00  0.0078773947
rs4257079        1 3455895 89  0.0000729184 1.00  1.00  0.0000729184

      effBB P2df
rs1891586 -0.47441121  1
rs794264   0.49475804  1
rs9178808 -0.49835345  1
rs5032886 -0.49835345  1
rs7435137  0.15517100  1
rs664063  -0.51438083  1
rs546570  -0.01392273  1
rs7908680    NA       1
rs166732    NA       1
rs4257079    NA       1
```

Again, noting interesting at GW significance level. If we would have had found something, naturally, we would not known if we mapped a T2D or obesity gene (or a gene for obesity in presence of T2D, or the one for T2D in presence of obesity).

At this point, you acquired the knowledge necessary for the self-exercise. Please close R by `q()` command and proceed to the next section.

## 5.5 GWA exercise

During the exercise, you will work with a larger data set (approximately 1,000 people and 7,000+ SNPs). You are to do complete three-round QC; perform GWA analysis with `dm2` as the outcome of interest and identify 10 SNPs which you would like to take to the stage 2 (replication) scan. You will do replication analysis using a confirmatory data set. If you did everything right, the SNPs which you identified as significant or replicated will be located in know T2D genes.

Please keep in mind that the data are simulated, and do not take your findings too seriously!

Start R by going to "Start -> Programs -> R -> R-2.4.1". Load `GenABEL` library by

```
> library(GenABEL)
```

The two data sets we will use in this exercise are part of the `GenABEL` distribution. The first one ("discovery" set) can be loaded by

```
> data(ge03d2)
```

Please move along the lines detailed in the guided exercise and try to answer following questions:

**Exercise 29** *How many cases and controls are presented in the original data set?*

**Exercise 30** *How many markers are presented in the original data set?*

**Exercise 31** *Is there evidence for inflation of the HWE test staistics?*

**Exercise 32** *Perform GWA analysis of the raw data, using asymptotic test and plot the results. Try to think how you can produce  $\chi^2 - \chi^2$  plot for the P-values on 1 d.f.. What is the estimate of  $\lambda$  for the 1 d.f. test?*

**Exercise 33** *Analyse empirical GW significance. How many SNPs pass genome-wide significance threshold, after correction for the inflation factor? Write down the names of these SNPs for further comparison.*

Perform complete three steps of the genetic data QC.

**Exercise 34** *How many male turned apparently female?*

**Exercise 35** *How many sporadic X errors do you still observe even when the female male and non-X X-markers are removed? (do not forget to `Xfix()` these!)*

**Exercise 36** How many "twin" DNAs did you discover?

**Exercise 37** How many genetic outliers did you discover?

After you have finished QC, answer the questions:

**Exercise 38** How many cases and controls are presented in the data after QC?

**Exercise 39** How many markers are presented in the data after QC?

**Exercise 40** Is there evidence for inflation of the HWE test staistics?

**Exercise 41** Perform GWA analysis of the cleaned data, using assymptotic test and plot the results. What is the estimate of  $\lambda$  for the 1 d.f. test?

**Exercise 42** Analyse empirical GW significance. How many SNPs pass genome-wide significance threshold, after correction for the inflation factor? Do these SNPs overlap much with the ones ranked at the top before the QC? If not, what could be the reason?

If time permits, do analysis with adjustment for covariates and stratified analysis.

Select 10 SNPs which you would like to follow-up. Say, you've selected rs1646456, rs7950586, rs4785242, rs4435802, rs2847446, rs946364, rs299251, rs2456488, rs1292700, and rs8183220.

Make a vector of these SNPs with

```
> vec12 <- c("rs1646456", "rs7950586", "rs4785242", "rs4435802",
+           "rs2847446", "rs946364", "rs299251", "rs2456488", "rs1292700",
+           "rs8183220")
```

Load the stage 2 (replicaton) data set by

```
> data(ge03d2c)
```

and select the subset of SNPs you need by

```
> confdat <- ge03d2c[, vec12]
```

Analyse the `confdat` for association with `dm2`.

**Exercise 43** Given the two-stage design, and applying the puristic criteria specified in the lecture, for how many SNPs you can claim a significant finding?

**Exercise 44** Using the same criteria, for how many SNPs you can claim a replicated finding?

You can check if any of the SNPs you have identified as significant or replicated are the ones which were simulated to be associated with `dm2` by using the command

```
> show.ncbi(c("snpname1", "snpname2", "snpname3"))
```

where `snpnameX` stands for the name of your identified SNP. The "true" SNPs can be found on NCBI and are located in known T2D genes (just because we used these names to name the "significant" ones).

If time permits, characterise the mode of inheritance of the significant SNPs. You can convert data from `GenABEL` format to the format used by `dgc.genetics` and `genetics` libraries by using `as.genotype()` function. Consult help for details. Please do not attempt to convert more than few dozens SNPs: the format of `genetics` is not compressed, which means conversion may take long and your low-memory computer may even crash if you attempt to convert the whole data set.

If time permits, try to do first round of QC allowing for HWE checks (assume FDR of 0.1 for total sample). In this case, can you still detect stratification in the "cleaned" data?

## 6 GWA in presence of genetic stratification

Both ethnic admixture and presence of close relationships represents examples of confounding in association analysis. However, the methods to correct for stratification as resulting from mixture of subjects coming from different genetic populations, and methods to correct for family relations may be slightly different, and will be described separately in the next two sections.

## 6.1 Analysis with ethnic admixture

In previous section we detected genetic stratification by analysis of genomic kinship matrix and excluded genetic outliers from our further analysis. When there are only a few such outliers, exclusion them from analysis is a good option. However, in large studies cases and controls are usually selected across a number of locations and genetic populations, and stratification is expected by design. In such case, analysis of association should account for this stratification.

One of the ways to do that is to perform *structured association* analysis. In such analysis, effect size estimate and its' variance is estimated within each strata separately, and then these estimates are pooled to generate global statistics. The strata can be known from design (e.g. place of birth or ethnicity of parents) or estimated from GWA data.

Let us do structured association analysis using the `data1` data derived in previous section. First, we need to define the variable which will tell what population the study subjects belong to. In previous section, we stored the names of 'outlier' subjects in variable `c11`:

> c11

```
[1] "id2097" "id6954" "id2136" "id858"
```

We can use function `%in%` to find out what names of subjects are in `c11`:

```
> pop <- as.numeric(data1@phdata$id %in% cl1)
> pop
```

Now, structured association may be done with `qtscore` function by specifying `strata`:

```
> data1.sa <- qtscore(dm2, data = data1, strata = pop)
```

We can compare results of analysis excluding outliers (black dots) and structured association analysis (green) by

```
> plot(data2.qt, cex = 0.5, pch = 19, ylim = c(1, 5))
> add.plot(data1.sa, col = c("lightgreen", "lightblue"), cex = 1.2)
```

The resulting plot is presented at figure 15.

In this case, there is very little difference, because all people belonging to the smaller sub-population are cases.

Other way to adjust for genetic (sub)structure is to apply the method of Price et al., which make use of principal components of the genomic kinship matrix to adjust both phenotypes and genotypes for possible stratification. In GenABEL, such analysis is done using `egsScore` function:

```
> data1.eg <- egsscore(dm2, data = data1, kin = data1.gkin)
```

The analysis plot may be added to the previous one by

```
> add.plot(data1.eg, col = c("red"), cex = 1.3)
```

The resulting plot is presented at figure 15.

Again, the difference between three analysis methods is marginal because there are no highly differentiated SNPs in the data set, and one sub-population is presented by cases only. Still, the signals at chromosome one and three slightly improved, while these at two and X went down.

**Exercise 45** Load and analyse the data set presented in file `stratified.RData`. GWA data presented in this file concern a study containing data from several populations. All these populations originate from the same base population some generations ago. Some of these populations maintained large size and some were small. There was little (2.5%) migration between populations.

Two traits (`quat` and `bint`) are available for analysis. Investigate relations between phenotypes and covariates. Perform association analysis using structured association and `egscore`. Answer the questions

1. How many SNPs and IDs are presented in the data set?
  2. How many SNPs and IDs pass the quality control?
  3. Is there evidence for stratification coming from the distribution of GW test for HWE (what is lambda?)
  4. Is there evidence that the test statistics for trait *quat* is inflated (what is lambda?)

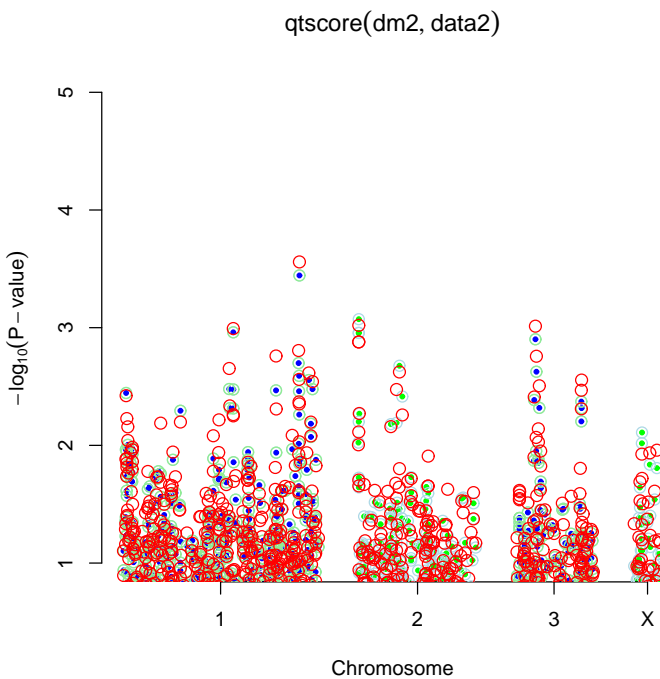


Figure 15: Comparison of structured association analysis (green), method of Price et al. (red) and analysis excluding genetic outliers (solid black).

- 5. Is there evidence that the test statistics for trait `bint` is inflated (what is lambda?)
- 6. How many genetically distinct populations are present in the data set? How many people belong to each population?
- 7. Please make some inferences on the characteristics / history of these populations.
- 8. What is the strongest SNP associated with trait `quat`? What model (method and covariates used) gives best results? Is the finding GW-significant?
- 9. What is the strongest SNP associated with trait `bint`? What model (method and covariates used) gives best results? Is the finding GW-significant?

## 6.2 Analysis of family data

In this section we will consider analysis of quantitative traits in a family-based cohort, where participants were not selected for the value of the trait under analysis. Such data may be generated in any study selecting participants based

on kinship (e.g. collections of sibships, nuclear or extended families); also any study in a genetically isolated population is likely to end up with a large proportion of relatively closely related individuals, even if ascertainment was random with the respect to kinship.

In pedigree-based association analysis the pedigree works as a confounder – exactly in the same manner as ethnic origin may work in a population-based study. Any genetic polymorphism is inherited through genealogy, and therefore genotypes are more similar between close relatives. In the same manner, any other heritable trait will be also more similar between relatives, and therefore certain degree of association is expected between *any* genetic marker and *any* heritable trait in a family-based sample. If additive 1 d.f. test for association is considered, the effect of confounding by pedigree can be shown to inflate the resulting null distribution of presumably  $\chi^2_1$  test statistics by a certain constant  $\lambda$ .

As you remember, this is exactly what happens when simple test for association is applied to a population-based data with ethnic admixture. In a population-based study with strong admixture (both in terms of the proportion and ethnic "distance"), some genomic regions may have been differentially selected in different populations. In such situation, use of genomic control does not prevent false-positive association between a trait and these regions, and other methods, such as EIGENSTRAT or Structured Association, are to be used.

For pedigree-based data coming from (relatively) genetically homogeneous population it can be shown that  $\lambda$  is a function of trait's heritability and pedigree structure, expressed as kinship matrix. Thus, genomic control is a simple and valid method to study association in genetically homogeneous families. However, this method reduces (or summarises if you prefer) all the abundant information about heritability and relationship into a single parameter  $\lambda$ , therefore it is not the most powerful method.

In quantitative genetics, a mixed polygenic model of inheritance may be considered as "industrial standard" – this model has sound theoretical bases and is proven by time to describe well inheritance of complex quantitative traits. This model describes the vector of observed quantitative traits as

$$Y = \mu + G + e \quad (1)$$

where  $\mu$  is the intercept,  $G$  is contribution from polygene, and  $e$  is random residual.

It is assumed that for each individual its "personal" random residual  $e_i$  is distributed as Normal with mean zero and variance  $\sigma_e^2$ . As these residuals are independent between pedigree members, the joint distribution of residuals in the pedigree can be modelled using multivariate normal distribution with variance-covariance matrix proportional to the identity matrix  $I$  (this is a matrix with diagonal elements equal to 1, and off-diagonal elements equal to zero):  $e \sim MVN(0, I\sigma_e^2)$ .

The polygenic component  $G$  describes the contribution from multiple independently segregating genes all having a small additive effect onto the trait (infinitesimal model). For a person for whom parents are not known, it is assumed that  $G_i$  is distributed as Normal with mean zero and variance  $\sigma_G^2$ . Assuming model of infinitely large number of genes, it can be shown that given polygenic values for parents, the distribution of polygene in offspring follows Normal distribution with mean  $(G_m + G_f)/2$  and variance  $\sigma_G^2/2$ , where  $G_m$  is maternal

and  $G_f$  is paternal polygenic values. From this, it can be shown that jointly the distribution of polygenic component in a pedigree can be described as multivariate normal with variance-covariance matrix proportional to the relationship matrix  $\Phi$ :  $G \sim MVN(0, \Phi\sigma_G^2)$ .

Thus the log-likelihood for this model can be written as a function of three parameters:

$$L(\mu, \sigma_G^2, \sigma_e^2) = -\frac{1}{2} \cdot \log_e |(\Phi \cdot \sigma_G^2 + I \cdot \sigma_e^2)| \\ + (Y - \mu)^T \cdot (\Phi \cdot \sigma_G^2 + I \cdot \sigma_e^2)^{-1} \cdot (Y - \mu) \quad (2)$$

where  $\mu$  is intercept,  $\sigma_G^2$  is the proportion of variance explained by the polygenic component, and  $\sigma_e^2$  is the residual variance.

Covariates such as sex, age, or a genetic marker studied for association can be easily included into the model:

$$Y = \mu + \sum_j \beta_j \cdot C_j + G + e$$

Here,  $C_j$  is the vector containing  $j$ -th covariate and  $\beta_j$  is the coefficient of regression of  $Y$  onto that covariate.

This mixed model leads to likelihood

$$L(\mu, \sigma_G^2, \sigma_e^2, \beta_1, \beta_2, \dots) = -\frac{1}{2} \cdot \log_e |(\Phi \cdot \sigma_G^2 + I \cdot \sigma_e^2)| \\ + \left( Y - (\mu + \sum_j \beta_j \cdot C_j) \right)^T \cdot (\Phi \cdot \sigma_G^2 + I \cdot \sigma_e^2)^{-1} \\ \cdot \left( Y - (\mu + \sum_j \beta_j \cdot C_j) \right) \quad (3)$$

This general formulation can be easily adopted to test genetic association; for example, an effect of a SNP can be incorporated into regression model

$$Y = \mu + \beta_g \cdot g + G + e$$

where  $g$  is the vector containing genotypic values. In this mode, you can specify a variety of 1 d.f. models by different coding of the vector  $g$ . For example, if you consider an "AG" polymorphism and want to estimate and test additive effect of the allele "G", you should code "AA" as 0 (zero), "AG" as 1 and "GG" as 2. Under this coding, the  $\beta_g$  will estimate additive contribution from the "G" allele. If you are willing to consider dominant model for G, you should code "AA" and "AG" as 0 and "GG" as 1. Recessive and over-dominant models can be specified in a similar manner. If, however, you want to estimate general 2 d.f. model, the specification should be different:

$$Y = \mu + \beta_a \cdot g + \beta_d \cdot I_{g=2} + G + e$$

where  $g$  is coded as 0, 1 or 2, exactly the same as in the additive model, and  $I_{g=2}$  is the binary indicator which takes value of one when  $g$  is equal to 2 and zero otherwise. In this model,  $\beta_a$  will estimate the additive and  $\beta_d$  – the dominance

effect. There may be other, alternative coding(s) allowing for essentially the same model, for example

$$Y = \mu + \beta_1 \cdot I_{g=1} + \beta_2 \cdot I_{g=2} + G + e$$

would estimate trait's deviation in these with  $g = 1$  ( $\beta_1$ ) and these with  $g = 2$  ( $\beta_2$ ) from the reference ( $g = 0$ ).

The classical way to estimate mixed polygenic model and test for significance is Maximum Likelihood (ML) or Restricted ML (REML) using equation (3). However, when large pedigrees are analysed, ML/REML solution may take prohibitively long time, i.e. from minutes to hours for single SNP analysis, making study of hundreds of thousand of SNPs impossible. Therefore fast approximate tests were developed for the purposes of GWA association analysis in samples of relatives.

Here we will cover two of fast approximations available, FAmily-based Score Test for Association (FASTA, Chen & Abecasis, 2007) and Genome-wide Rapid Analysis using Mixed Models And Score test (GRAMMAS, Amin et al, 2007). Both tests are based on the classical polygenic mixed model and are performed in two steps.

First, polygenic model as specified by equation (1) and likelihood (2) is estimated using available data.

Secondly, the maximum likelihood estimates (MLEs) of the intercept,  $\hat{\mu}$ , proportion of variance explained by the polygenic component,  $\hat{\sigma}_G^2$ , and residual variance,  $\hat{\sigma}_e^2$ , are used to compute the FASTA test statistics

$$T_F^2 = \frac{((g - E[g])^T \cdot (\Phi \cdot \hat{\sigma}_G^2 + I \cdot \hat{\sigma}_e^2)^{-1} \cdot (Y - \hat{\mu}))^2}{(g - E[g])^T \cdot (\Phi \cdot \hat{\sigma}_G^2 + I \cdot \hat{\sigma}_e^2)^{-1} \cdot (g - E[g])}$$

It can be shown that  $T_F^2$  follows  $\chi_1^2$  when pedigree structure is 100% complete and 100% correct. As this is never actually the case, application of GC to correct for residual inflation is recommended.

FASTA test results in unbiased estimates of the SNP effect and correct  $P - values$ . Please keep in mind that this is correct – as for any score test – only when alternative is reasonably close to the null, i.e. when the SNP explains small proportion of trait's variance. Disadvantages of this test are that it can be relatively slow when thousands of study subjects are analysed, and that permutation procedures can not be applied to estimate genome-wide significance, because the data structure is not exchangeable.

Other test, GRAMMAS, also exploits MLEs from the polygenic model (1). However, these are used to first compute the vector of environmental residuals  $\hat{e}$ , using standard equation

$$\hat{e} = \hat{\sigma}_e^2 \cdot (\Phi \cdot \hat{\sigma}_G^2 + I \cdot \hat{\sigma}_e^2)^{-1} \cdot (Y - \hat{\mu})$$

These residuals, in turn, are used to run simple score test:

$$T_G^2 = \frac{((g - E[g])^T \cdot \hat{e})^2}{(g - E[g])^T \cdot (g - E[g])}$$

This test is conservative, but GC can be used to correct for the deflation of the test statistics.

The fact that environmental residuals  $\hat{e}$  are not dependent on pedigree structure leads to a nice property of the GRAMMAS test: the data structure becomes exchangeable and permutations may be used to estimate genome-wide significance. When used in combination with GC,  $P$  – values derived from GRAMMAS test are correct; however, there is a downward bias in estimates of SNP effects.

When using FASTA or GRAMMAS test, it is recommended to estimate genomic kinship matrix from available genome-wide data, and use it in analysis instead of pedigree kinship. This solution firstly does not rely on the completeness and quality of pedigree; secondly, genomic kinship is more likely to give a better estimate of a *true* covariance between individual genomes, while pedigree kinship provides one with expectation. Therefore use of genomic kinship is expected to lead to better estimates of polygenic model, and thus better power to detect association in GWA analysis. This being said, we generally advocate use of genomic, and not pedigree kinship. Of course, you can only implement this solution when you have GWA data; in a candidate gene study you will have to rely on the pedigree structure to estimate kinship matrix.

### 6.3 Example GWA analysis using family-based data

In this section, we will explore small data set (150 people, 5827 SNPs). Let us load and explore it:

```
> load("RData/erfsmal.RData")
> ls()

[1] "erfs"   "pkins"

> class(erfs)

[1] "gwaa.data"
attr(,"package")
[1] "GenABEL"

> class(pkinds)

[1] "matrix"
```

You can see that there are two objects, `erfs` and `pkinds`, presented in the data. The class of the first object is standard GenABEL's `gwaa.data-class`; this is the object containing GWA data. The other object contains kinship matrix, as estimated from pedigree data.

You can check the number of people and SNPs in the data set with

```
> erfs@gtdata@nids

[1] 150

> erfs@gtdata@nsnps

[1] 5827
```

As usual, it is advisable to check the distribution of SNPs by chromosome:

```

> table(erfs@gtdata@chromosome)

  1   10   11   12   13   14   15   16   17   18   19   2   20   21   22   23   3   4   5   6
484  251  224  285  188  210  206  189  159  170  149  481  130  100  132  26  403  300  320  397
    7     8     9     X
286  253  206  278

```

(here, 23 stays for pseudo-autosomal region of the X chromosome); you can see that markers are evenly spread over the chromosomes.

Summary marker statistics can be generated by

```

> descriptives.marker(erfs@gtdata)

$`Minor allele frequency distribution`
  X<=0.01 0.01<X<=0.05 0.05<X<=0.1 0.1<X<=0.2      X>0.2
No      17.000        26.000        75.000      437.000  5272.000
Prop     0.003        0.004        0.013        0.075      0.905

$`Cumulative distr. of number of SNPs out of HWE, at different alpha`
  X<=1e-04 X<=0.001 X<=0.01 X<=0.05 all X
No          2    14.000    83.000   319.000    5827
Prop        0    0.002    0.014    0.055       1

$`Distribution of porportion of successful genotypes (per person)`
  X<=0.9 0.9<X<=0.95 0.95<X<=0.98 0.98<X<=0.99 X>0.99
No          0     1.000     2.000      12.00    135.0
Prop        0     0.007     0.013      0.08      0.9

$`Distribution of porportion of successful genotypes (per SNP)`
  X<=0.9 0.9<X<=0.95 0.95<X<=0.98 0.98<X<=0.99 X>0.99
No      77.000        33.000        214.000      208.000  5295.000
Prop     0.013        0.006        0.037        0.036      0.909

$`Mean heterozygosity for a SNP`
[1] 0.4402752

$`Standard deviation of the mean heterozygosity for a SNP`
[1] 0.08287253

$`Mean heterozygosity for a person`
[1] 0.4354001

$`Standard deviation of mean heterozygosity for a person`
[1] 0.01305448

```

You can see that the quality of genotypic data is quite reasonable: call rate is generally high, both per-person and per SNP, and there is little deviation from Hardy-Weinberg equilibrium.

Let us explore pedigree kinship matrix. First, let us just look how this matrix looks like by displaying few elements from the upper-left corner:

```
> pkins[1:5, 1:5]
```

```

      id1      id2      id3      id4      id5
id1 5.00000e-01 8.56146e-05 1.01984e-04 2.33397e-04 8.56146e-05
id2 8.56146e-05 5.00000e-01 3.96513e-03 2.56896e-05 2.51269e-01
id3 1.01984e-04 3.96513e-03 5.00000e-01 1.21593e-05 3.96513e-03
id4 2.33397e-04 2.56896e-05 1.21593e-05 5.00000e-01 2.56896e-05
id5 8.56146e-05 2.51269e-01 3.96513e-03 2.56896e-05 5.00000e-01

```

By definition, pedigree kinship should take values between 0 and 0.5 (plus some small amount from inbreeding); kinship between (non-inbred) sibs or an offspring and the parent is 1/4. You can see that in the upper-left corner there is one inbred sib-pair (or parent-offspring pair; "id2" and "id5"). You can also see that this matrix is symmetric around the diagonal.

Let us summarise the distribution of kinship coefficients; in doing this we want to generate the summary for every off-diagonal element only once. Function `lower.tri` can be used to get the "lower triangle" sub-matrix elements:

```

> summary(pkins[lower.tri(pkins)])

```

Min.	1st Qu.	Median	Mean	3rd Qu.	Max.
0.0000000	0.0004999	0.0028500	0.0062810	0.0053780	0.2633000

As you can see, average relationship corresponds to that expected between second cousins ( $1/64 = 0.015625$ ) and third cousins ( $1/256 = 0.00390625$ ).

We can also draw a histogram of the distribution of the kinship coefficients (shown at figure 16A):

```
> hist(pkins[lower.tri(pkins)])
```

and see that most elations are indeed remote.

Let us estimate genomic kinship matrix using autosomal data with the command `ibs`, and look up the elements in the upper-left corner:

```

> gkins <- ibs(erfs[, autosomal(erfs)], weight = "freq")
> gkins[1:5, 1:5]

      id1      id2      id3      id4      id5
id1 0.531729506 5.439000e+03 5.446000e+03 5441.00000000 5440.0000000
id2 -0.012569446 4.921817e-01 5.524000e+03 5524.00000000 5521.0000000
id3 0.001516184 -8.551323e-03 5.144766e-01 5529.00000000 5528.0000000
id4 0.010459422 -1.471218e-02 -3.467894e-03 0.52042474 5523.0000000
id5 -0.007957596 2.561484e-01 -8.925127e-03 -0.02205665 0.4943998

```

Here, the estimated kinship is shown below the diagonal, and the number of informative SNP pairs used for estimation is shown above the diagonal.

You can see that "genomic kinship" coefficients may take values lower than zero, which is consequence of the fact that in effect "genomic kinship" is simply covariance between the vectors of individual genotypes. This quantity, though it provides an unbiased estimate of kinship, can be lower than zero.

```

> summary(gkins[lower.tri(gkins)])

```

Min.	1st Qu.	Median	Mean	3rd Qu.	Max.
-0.038980	-0.011930	-0.005826	-0.003362	0.000481	0.268000

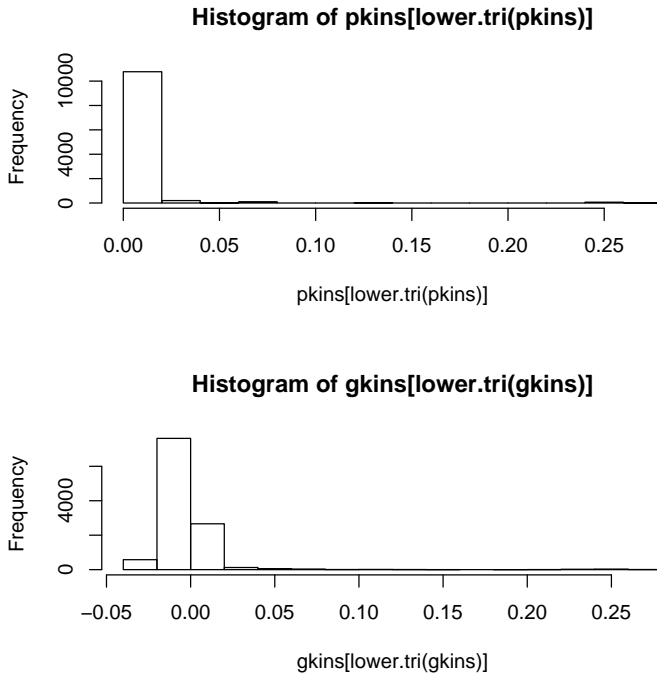


Figure 16: Distribution of the pedigree (upper histogram) and genomic (lower histogram) kinship coefficients for `erfs` data set.

here, the average is quite close to that obtained with pedigree kinship.

We can also draw a histogram of the distribution of "genomic kinship" coefficients (shown at figure 16B):

```
> hist(gkins[lower.tri(gkins)])
```

and can easily graphically present relations between genomic and pedigree kinship with

```
> plot(pkins[lower.tri(pkins)], gkins[lower.tri(gkins)])
```

(shown at figure 17), and estimate correlation between the two with

```
> cor(pkins[lower.tri(pkins)], gkins[lower.tri(gkins)])
```

```
[1] 0.91615
```

From the graph, you can clearly see that, though there is a very strong correlation between genomic and pedigree kinships, these are not identical.

In real data, you may find that there are some points where pedigree data clearly suggest relation different from that suggested by genomic data. Which one to believe? Generally, pedigrees are more prone to errors than genotypic data. In the data containing close relatives it is better to rely on "genomic kinship".

Let us first analyse the data using plain GC method:

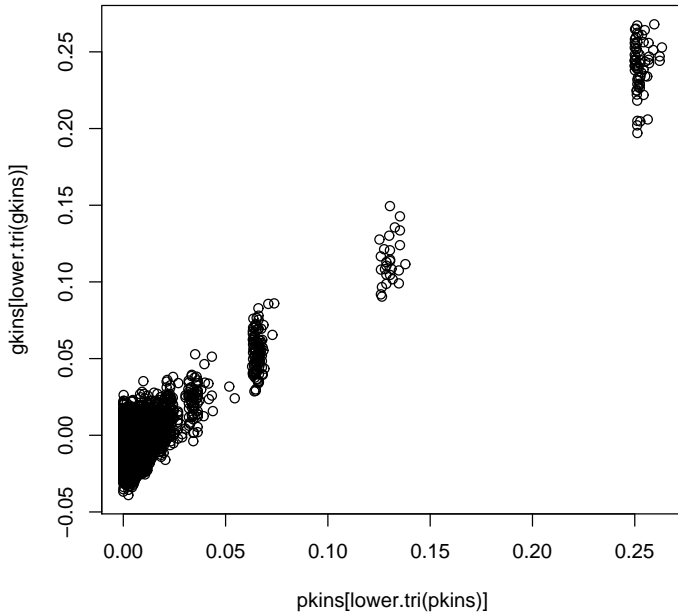


Figure 17: Scatter-plot relating pedigree and genomic kinships for `erfs` data set.

```
> qts <- qtscore(qtbas, data = erfs)
```

You can check the estimate of the inflation factor  $\lambda$  with

```
> qts$lam$est
```

```
[1] 1.158683
```

This is relatively high value, suggesting presence of close relatives in data and high heritability of the trait.

The top 10 hits from GWA analysis can be displayed with

```
> descriptives.scan(qts, sort = "Pc1df")
```

	Chromosome	Position	N	effB	P1df	Pc1df
rs1781670	13	2694164735	150	0.4554848	0.0001804816	0.0005032759
rs1054889	2	324154337	150	0.4760959	0.0002711808	0.0007173608
rs1861659	16	3171591122	150	0.4844231	0.0002811280	0.0007402184
rs1860991	5	1213061568	150	0.4616985	0.0002834991	0.0007456515
rs1043883	8	1745120471	128	0.4338657	0.0005729702	0.0013762794
rs1872087	17	3329938025	150	-0.4111272	0.0006091602	0.0014517276
rs1982904	4	961162979	150	0.5369605	0.0006904958	0.0016192705
rs2141693	12	2590653716	150	1.2178255	0.0007070577	0.0016530692
rs7141672	14	2851979738	150	0.4595401	0.0007473279	0.0017348360

```

rs1075456      15 3036078968 150 -0.3943230 0.0008094764 0.0018599411
                effAB     effBB      P2df
rs1781670 0.55710736 0.9015131 0.0007574863
rs1054889 0.32660599 1.0001217 0.0008416365
rs1861659 0.30184684 1.0290361 0.0006959544
rs1860991 0.49412227 0.9080701 0.0013452354
rs1043883 0.23834763 0.9427557 0.0015609770
rs1872087 -0.50149169 -0.8092418 0.0024403173
rs1982904 0.60470761 0.8678924 0.0026214066
rs2141693 1.21782551      NA 0.0007070577
rs7141672 0.37923110 1.0172357 0.0028443093
rs1075456 -0.01486038 -0.8540990 0.0002978615

```

here, nominal  $P$ -values after genomic control are given in column named "Pc1df".

We can estimate genome-wide empirical significance by using the same function with `times` argument, which tells the number of permutations:

```

> qts.e <- qtscore(qtbas, data = erfs, times = 200)
100%
> descriptives.scan(qts.e, sort = "Pc1df")

      Chromosome   Position    N      effB    P1df  Pc1df      effAB
rs1781670        13 2694164735 150 0.45548481 0.545 0.860 0.5571074
rs1054889         2 324154337 150 0.47609594 0.680 0.935 0.3266060
rs1860991         5 1213061568 150 0.46169852 0.685 0.935 0.4941223
rs1861659        16 3171591122 150 0.48442305 0.685 0.935 0.3018468
rs1043883         8 1745120471 128 0.43386568 0.895 0.995 0.2383476
rs1872087        17 3329938025 150 -0.41112719 0.905 0.995 -0.5014917
rs2017143          1 2252811 150 0.06610235 1.000 1.000 0.3084208
rs2840531          1 2318570 150 -0.31150314 1.000 1.000 -0.3273450
rs2477703          1 2458154 150 -0.16765976 1.000 1.000 -0.2043520
rs734999           1 2545378 150 -0.04010750 1.000 1.000 0.2438654
                effBB      P2df
rs1781670 0.90151313 0.970
rs1054889 1.00012170 0.975
rs1860991 0.90807011 1.000
rs1861659 1.02903607 0.935
rs1043883 0.94275573 1.000
rs1872087 -0.80924177 1.000
rs2017143 0.05342849 1.000
rs2840531 -0.47957726 1.000
rs2477703 -0.29647369 1.000
rs734999 -0.09548236 1.000

```

As you can see, in this analysis nothing comes even close to genome-wide significance, as indicated by genome-wide corrected  $P$ -values (column "Pc1df") all  $>> 0.05$ .

Let us try to estimate polygenic model with

```
> h2 <- polygenic(qtbas, kin = gkins, data = erfs)
```

The results of estimation are contained in "h2an" element of the resulting analysis object:

```
> h2$h2an

$minimum
[1] 139.9891

$estimate
[1] 0.03828507 0.78448170 1.21485857

$gradient
[1] -4.0470110 -0.0636133 -3.0506775

$code
[1] 2

$iterations
[1] 4
```

In the "estimate" list, the MLEs shown correspond to intercept  $\hat{\mu}$ , heritability  $\hat{h}^2 = \hat{\sigma}_G^2 / (\hat{\sigma}_G^2 + \hat{\sigma}_e^2)$ , and total variance  $\hat{\sigma}_T^2 = \hat{\sigma}_G^2 + \hat{\sigma}_e^2$ . You can see that heritability of the trait is indeed high – almost 80%.

Under these conditions (high heritability, presence of close relatives) we may expect that FASTA and GRAMMAS analysis exploiting heritability model and relationship matrix in exact manner may have better power compared to simple GC.

Let us run FASTA test using estimated polygenic model, as specified by `h2` object:

```
> mms <- mmscore(h2, data = erfs)
```

There is little residual inflation left when we use "genomic kinship" matrix:

```
> mms$lam$est
```

```
[1] 1.024464
```

And the significance of "top" hit becomes an order of magnitude better compared to plain GC:

```
> descriptives.scan(mms, sort = "Pc1df")
```

	Chromosome	Position	N	effB	P1df	Pc1df	effAB
rs1075456	15	3036078968	150	-0.3251590	0.0000119801	0.0000152282	NA
rs1054889	2	324154337	150	0.2561733	0.0002076476	0.0002473709	NA
rs1781670	13	2694164735	150	0.2443012	0.0002398577	0.0002848148	NA
rs1264007	X	4256199578	150	-0.2720459	0.0006816474	0.0007906349	NA
rs774033	12	2531521679	150	-0.2094525	0.0009456325	0.0010888321	NA
rs7141672	14	2851979738	150	0.3351830	0.0010869290	0.0012476437	NA
rs537848	15	3020539910	150	0.2127141	0.0012151234	0.0013913324	NA

rs760109	X	4353106270	150	0.2686623	0.0013788938	0.0015744148	NA
rs1860991	5	1213061568	150	0.2517586	0.0014521085	0.0016561046	NA
rs1262129	14	2878536332	150	-0.1843508	0.0017034834	0.0019359148	NA
	effBB	P2df					
rs1075456	NA	9.99					
rs1054889	NA	9.99					
rs1781670	NA	9.99					
rs1264007	NA	9.99					
rs774033	NA	9.99					
rs7141672	NA	9.99					
rs537848	NA	9.99					
rs760109	NA	9.99					
rs1860991	NA	9.99					
rs1262129	NA	9.99					

If you compare these results to that obtained with simple GC, you can also see that the ranks of top hits have changed quite a bit; unbiased estimated of genetic effects were obtained.

However, we can not estimate genome-wide significance with FASTA, because the data structure is not exchangeable.

Using GRAMMAS method, you can estimate nominal  $P - values$  by

```
> grs <- qtscore(h2$pgres, data = erfs, clam = FALSE)
> grs$lam$est
[1] 0.7987019
```

In the above analysis, note that the estimated "inflation" factor  $\lambda$  is less than one, i.e. now it is the GRAMMAS *deflation* factor. In order to obtain non-conservative test statistics, we had to say to qtscore that deflation is OK (parameter `clam=FALSE`).

We can see "top" nominal corrected  $P - values$  with

```
> descriptives.scan(grs, sort = "Pc1df")

```

Chromosome	Position	N	effB	P1df	Pc1df	
rs1075456	15	3036078968	150	-0.10704009	0.0001344706	0.0000193502
rs1054889	2	324154337	150	0.10245323	0.0009999196	0.0002314709
rs1781670	13	2694164735	150	0.09165369	0.0015542793	0.0003990239
rs1264007	X	4256199578	150	-0.07764670	0.0023993775	0.0006817145
rs774033	12	2531521679	150	-0.09282392	0.0026100856	0.0007562579
rs7141672	14	2851979738	150	0.09341129	0.0040026811	0.0012807435
rs537848	15	3020539910	150	0.08489591	0.0042136070	0.0013643467
rs1860991	5	1213061568	150	0.08655964	0.0042663779	0.0013854152
rs760109	X	4353106270	150	0.07432363	0.0045810469	0.0015122637
rs2639197	15	3068765468	150	-0.09085234	0.0052630062	0.0017939216
	effAB	effBB	P2df			
rs1075456	-0.04882437	-0.2241217	0.0002410666			
rs1054889	0.04392601	0.2236716	0.0013187233			
rs1781670	0.08025377	0.1843682	0.0064411019			
rs1264007	-0.09842481	-0.1512012	0.0099102406			

```

rs774033 -0.09430729 -0.1856043 0.0107644457
rs7141672 0.08525171 0.1967954 0.0153935223
rs537848 0.08848437 0.1695158 0.0165978319
rs1860991 0.10807561 0.1629485 0.0140947764
rs760109 0.06139187 0.1491903 0.0179490427
rs2639197 -0.09197031 -0.1804026 0.0203630654

```

By comparing this output to that from FASTA test, you can see that  $P - values$  are quite close, but the effects are underestimated with GRAMMAS, as expected.

However, the strengths of GRAMMAS test is not only its speed, but also possibility to estimate genome-wide significance. This can be done by

```

> grs.e <- qtscore(h2$pgres, data = erfs, times = 200, clam = FALSE)
100%

> descriptives.scan(grs.e, sort = "Pc1df")

  Chromosome   Position    N      effB  P1df  Pc1df      effAB
rs1075456       15 3036078968 150 -0.10704009 0.405 0.055 -0.04882437
rs1054889        2 324154337 150 0.10245323 0.980 0.530 0.04392601
rs1781670       13 2694164735 150 0.09165369 0.995 0.750 0.08025377
rs1264007        X 4256199578 150 -0.07764670 1.000 0.915 -0.09842481
rs774033         12 2531521679 150 -0.09282392 1.000 0.945 -0.09430729
rs1860991         5 1213061568 150 0.08655964 1.000 0.985 0.10807561
rs7141672        14 2851979738 150 0.09341129 1.000 0.985 0.08525171
rs537848          15 3020539910 150 0.08489591 1.000 0.985 0.08848437
rs760109          X 4353106270 150 0.07432363 1.000 0.990 0.06139187
rs2017143         1 2252811 150 0.02373127 1.000 1.000 0.05393011

      effBB  P2df
rs1075456 -0.2241217 0.595
rs1054889  0.2236716 0.995
rs1781670  0.1843682 1.000
rs1264007 -0.1512012 1.000
rs774033  -0.1856043 1.000
rs1860991  0.1629485 1.000
rs7141672  0.1967954 1.000
rs537848  0.1695158 1.000
rs760109  0.1491903 1.000
rs2017143  0.0376451 1.000

```

As you can see, now the "top" hit starts approaching genome-wide significance (genome-wide  $P - value \sim 10\%$ ), showing the power of kinship-based methods under high heritability model.

Finally, let us plot  $-\log_{10}$  nominal  $P - values$  from different methods across the genome. Let black dots correspond to GC, green to GRAMMAS and red to FASTA (figure 18):

```

> plot(mms, df = "Pc1df")
> add.plot(grs, df = "Pc1df", col = c("lightgreen", "lightblue"),
+           cex = 1.2)
> add.plot(qts)

```

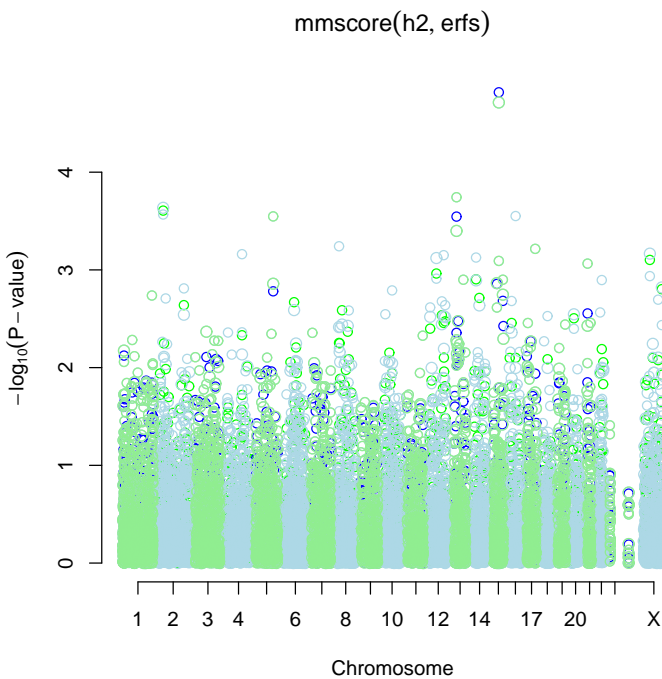


Figure 18: Comparison of FASTA (red), GRAMMAS (green), and plain GC (black).

You can see that there is a great degree of correlation between the FASTA and GRAMMAS  $P$  – values, while plain GC really stands apart.

#### 6.4 Exercise: analysis of family data

**Exercise 46** Repeat heritability estimation, FASTA and GRAMMAS analysis of previous section using pedigree kinship (`pkins` object). Discuss the results.

In the next section, you will explore a small (695 people) subset of people from ERF, a family-based study with participants coming from a genetically isolated population and sampled based on kinship (all living descendants of 22 couples living in the area in mid-XIX<sup>th</sup> century). The study participants were genotyped using Illumina 6K "linkage" array. QC was already performed. Your trait of interest is "qtbas".

Explore the data set and answer the questions:

**Exercise 47** Describe the trait "qt". Can you detect significant outliers at visual inspection? Is trait distributed normally? What are significant covariates?

**Exercise 48** Explore relations between genomic and pedigree kinship (these are provided in data as `gkin` and `pkin` data objects, respectively). What are your conclusions? Which matrix would you use later on?

**Exercise 49** What is the heritability of the trait (take care: polygenic analysis may rung for a long while)? Based on heritability analysis, how would you rank different methods of GWA analysis for this trait (and why)?

**Exercise 50** Do GWA analysis using simple score test with genomic control. Estimate genome-wide significance. What are your conclusions?

Run GWA analysis using the "best" method and model as you have decided in previous exercises. Estimate genome-wide significance. What are your conclusions? Did they change compared to simple analysis?

**Exercise 51** Repeat the last "best" analysis using pedigree kinship. How your results change?

**Exercise 52** If you have any time left – repeat analysis using "qt" trait. This one is much more fun, but also more laborous to analyse.

## 7 Exploring and using public databases

See help for `show.ncbi`

More functions coming soon.

## 8 Genetic data imputations

With very dense SNP panels available nowadays, linkage disequilibrium between neighbouring SNP is high. This allows us, based on the context, predict genotypes with high degree of accuracy. The genetic data imputations refer to such prediction of genotypes of missing SNPs,

The uses of genetic data imputations are basically two-fold. The first situation concerns imputations of partly missing genotypes of SNPs, typed, in a study, based on the context provided by SNPs typed in the same study. This allows decreasing the proportion of missing SNPs, potentially improving the power of the study. The second situation concerns prediction of genotypes at SNPs, which were not typed in the study, based on the context provided by the SNPs typed and using the data coming from another study with more dense genotyping, which may be potentially done using other population. Though imputations may be done using the same method under the two scenarios, and difference between them is quantitative rather than qualitative, for practical reasons it is useful to distinguish these two situations.

GenABEL is well connected with MACH imputations software, developed by Y. Li and G. Abecasis. Please read abou MACH at <http://www.sph.umich.edu/csg/abecasis/MACH/>.

### 8.1 Imputing partly missing genotypes

Assume you have `gwaa.data` object "data" and you want to impute missing genotypes for chromosome 1. For this, you will first need to export chromosome one data in MACH format. The MACH data format bears remarkable similarity to the format used by MERLIN, an excellent linkage analysis program. This is not surprising: MERLIN was developed by G. Abecasis. Therefore we can export the data for MACH using our function `export.merlin`:

```
> export.merlin(data[,data@gtdata@chromosome=="1"],pedf="chr1.ped",
+                           dataf="chr1.dat",mapf="chr1.map")
```

where the first argument supplies the data to be exported and others specify the names for the output files containing MACH genetic data (pedigree-file), and descriptives (data-file). It will also save map information in file 'chr1.map'; this information is not necessarily used by MACH, but is required by MERLIN, and also required by GenABEL .

Given the data are exported, we can use MACH for imputations of missing genotypes. Read MACH documentation for details. Basically, under UNIX, you need to run command

```
bash> mach1 -p chr1.ped -d chr1.dat --states 200 --rounds 20
--mle --mldetails --prefix chr1.out
```

This should produce files, of which we will use `chr1.out.mlgeno`, containing the imputed genotypes and `chr1.out.mlinfo`, containing information about imputations process, including quality score.

Now we need to read the imputed data back into R environment. First, we need convert MACH output to GenABEL format:

```
> convert.snp.mach(pedf="chr1.out.mlgeno",mapf="chr1.map",
+                   info="chr1.out.mlinfo",outf="chr1.raw",quality=0.9)
```

This will convert MACH output to file 'chr1.raw', containing genotypic data in GenABEL format. All SNPs with quality < 0.9 will be dropped.

Conversion of data from MACH to GenABEL format may be a lengthy procedure. You can improve the speed much by replacing all "/" to space (" ") in 'chr1.out.mlgeno'.

As soon as conversion is done, you can import the data by

```
> chr1.imp <- load.gwaa.data(phe="pheno.dat",gen="chr1.raw")
```

This assumes that your phenotypes are in file "pheno.dat", probably easiest way to generate it is by

```
> save.gwaa.data(data,phen="pheno.dat",geno="tmp.raw")
```

Later, you can even replace your original data with imputed:

```
> data@gtdata@gtps[,data@gtdata@chromosome=="1"] <- chr1.imp@gtdata@gtps
```

## 8.2 Inferences from other data sets

If you imported your data to GenABEL before version 1.2-6, you will need to re-load your data to GenABEL , to provide the information about actual coding and strand (easily available for Affymetrix). Please check documentation for convert.snp.text, convert.snp.illumina, convert.snp.ped. For the latter two take care that strand="file".

Assume that your data are in `gwaa.data` object `data`; you want to impute SNPs in the region between SNP 100 and 500.

Go to HapMap web-site and download phased haplotype data for the region between SNP 100 and 500. We have a script which generates MACH input files from HapMap dump-files (`dump2mach.pl`, available upon request). Note that in HapMap phased haplotype data, SNP coding is given for "+" strand.

Assume that haplotype and SNP file names are `reg1.haplo`, `reg1.snps` (read MACH documentation for details). Check that all of your SNPs (from 100 to 500) are in HapMap data; else generate the vector of indexes/names of the ones which are in HapMap (only these will pass through MACH). Assuming all 500 are in HapMap, export your regional data in MACH format by:

```
> export.merlin(data[,c(100:500)],pedf="reg1.ped",
+                 dataf="reg1.dat",mapf="reg1.map",strand="+")
```

To do imputations, in bash type

```
bash> mach1 -p reg1.ped -d reg1.dat -h reg1.haplo -s reg1.snps
      -greedy --rounds 20 --mle --mldetails --prefix reg1.out
```

This should produce files, of which we will use `reg1.out.mlgeno`, containing the imputed genotypes, and `reg1.out.mlinfo`, containing information about imputations process, including quality score.

Back in R, convert the data to GenABEL format by:

```
> convert.snp.mach(pedf="reg1.out.mlgeno",mapf="reg1.large.map",
+                     info="reg1.out.mlinfo",outf="reg1.raw",quality=0.9)
```

Here, `reg1.large.map` provides map for all HapMap SNPs in the region (this one is easily made from the dump-file, or with `dump2mach.pl`) The above command will convert MACH output to GenABEL format, dropping all SNPs with quality < 0.9.

Conversion of data from MACH to GenABEL format may be a lengthy procedure. You can improve the speed much by replacing all "/" to space (" ") in 'chr1.out.mlgeno'.

Now, you can import the data by

```
> reg1.imp <- load.gwaa.data(phe="pheno.dat",gen="reg1.raw")
```

## 9 Meta-analysis of GWA scans

### 9.1 Standard meta-analysis methods

Imagine you are interested in the effect of a certain polymorphism onto a particular disease. After scanning literature, you find some studies that implicate certain allele as significantly increasing the risk of the disease, but you will typically find also that other studies were inconclusive (no significant association), and that even some of the studies implicated the same allele as "protective".

Your gut feeling may be that the allele is indeed the risk one, because you feel that the studies contradicting to this hypothesis were based on small number of subjects; however, how do you quantify this feeling? In this situation you need to perform meta-analysis of available data to come up with the joint effect size estimate and P-value, as based on all available data.

Let us first consider a situation when you are interested in the effect of the allele on a quantitative phenotype, expressed as a coefficient of regression of the trait onto the number of this allele in the genotype. Under a favourable scenario, from every individual study you would know the estimate of this regression coefficient, and the standard error of the estimate (or, equivalently, the *P-value* or the test statistics value for association).

One of approaches frequently used in meta-analysis of the data coming from a number of independent studies is the inverse variance method. In essence, this method is equivalent to combining likelihoods coming from separate studies, using quadratic approximation. Denote coefficients of regression estimated in  $N$  studies as  $\beta_i$ , and associated squared standard errors of the estimates as  $s_i^2$  where  $i \in 1, 2, \dots, N$ . Note that the regression coefficient should be reported on the same scale, e.g. centimeters, meters, or using observations reported on the standard normal scale. Define weights for individual studies as

$$w_i = \frac{1}{s_i^2}$$

Then the pooled estimate of the regression coefficient is

$$\beta = \frac{\sum_{i=1}^N w_i \beta_i}{\sum_{i=1}^N w_i}$$

As you can see, the weights have straightforward interpretation: the bigger the weight of the study (meaning the small is the standard error in the study), the larger is the contribution from this study onto the pooled estimate.

The standard error of the pooled estimate is computed as

$$s^2 = \frac{1}{\sum_{i=1}^N w_i}$$

and the  $\chi^2$ -test for association is computed in standard manner as

$$T^2 = \frac{\beta^2}{s^2} = \frac{\left(\sum_{i=1}^N w_i \beta_i\right)^2}{\sum_{i=1}^N w_i}$$

or, alternatively, the Z-test is

$$Z = \frac{\beta}{s} = \frac{\sum_{i=1}^N w_i \beta_i}{\sqrt{\sum_{i=1}^N w_i}}$$

Let us try to do meta-analysis using the inverse variance pooling method. Imagine we have information from four different studies reporting effect and the standard error of the same allele:

Let us try to access the joint significance of the association using these data. First, let us define a vector of regression coefficients and squared standard errors:

Table 1: Estimated regression coefficients from four studies

Study	$n$	$\beta$	$s_\beta$	$\chi^2$
1	225	0.16	0.07	5.224
2	560	0.091	0.042	4.694
3	437	0.072	0.048	2.25
4	89	-0.03	0.12	0.062
Total	1311	?	?	?

```
> beta <- c(0.16, 0.091, 0.072, -0.03)
> s <- c(0.07, 0.042, 0.048, 0.12)
> s2 <- s * s
> s2

[1] 0.004900 0.001764 0.002304 0.014400
```

Compute the weight for individual studies as

```
> w <- 1/s2
> w

[1] 204.08163 566.89342 434.02778 69.44444
```

Estimate pooled regression coefficient as

```
> pbeta <- sum(w * beta)/sum(w)
> pbeta
```

```
[1] 0.08898527
```

and pooled squared standard error as

```
> ps2 <- 1/sum(w)
> ps2

[1] 0.0007846539
```

To access significance of association in meta-analysis, let us compute  $\chi^2$  test statistics and the  $P$  – value with

```
> pchi2 <- pbeta * pbeta/ps2
> pchi2
```

```
[1] 10.09155
```

```
> ppvalue <- 1 - pchisq(pchi2, 1)
> ppvalue
```

```
[1] 0.001489504
```

We conclude that there is a significant association in meta-analysis.

There is an important effect which should be considered when doing meta-analysis of published data. Given numerous polymorphisms available in human

genome, a particular polymorphism usually becomes a focus of interest only when it was shown to be significantly associated in some study which reports it. Put it other way around: only when a significant association was detected and reported, more studies are likely to be performed on the same polymorphism. This first report, however, is not guaranteed to demonstrate a true association: it may well report a false-positive, or, even in presence of association, it is likely to over-estimate the effect of the polymorphism. Thus there is a positive bias in literature reports, and this bias is particularly strong for the first report, a phenomenon frequently referenced to as "champion's" or "winner's curse".

The observations we have just considered are quite typical in that the first study, where the association was originally discovered, reports the biggest effect and most significant effect, while the follow-up studies suggest smaller effect.

Therefore, when you meta-analyse data from publications it is always good idea to exclude the first report (in case it is positive – and it is always positive!) and check if significant association is still observed. Let us try to do that:

```
> beta <- beta[2:4]
> s2 <- s2[2:4]
> w <- w[2:4]
> pbeta <- sum(w * beta)/sum(w)
> pbeta

[1] 0.07544522

> ps2 <- 1/sum(w)
> ps2

[1] 0.0009342602

> pchi2 <- pbeta * pbeta/ps2
> pchi2

[1] 6.092501

> ppvalue <- 1 - pchisq(pchi2, 1)
> ppvalue

[1] 0.01357568
```

Indeed, when the first "champion" report is excluded, the overall evidence is decreased and results become less significant, though still pointing to the same direction.

When binary traits are studied, and results are reported as Odds Ratios with  $P-values$ , it is also possible to apply inverse variance method. For this, you need to transform your Odds Ratios using natural logarithm, and, on this scale, estimate the standard error. Generic inverse variance pooling may be applied to the data transformed this way; the final results are back-transformed onto Odds Ratio scale using exponentiation.

Let us consider a simple example. Let Odds Ratios and  $\chi^2$  test statistics values coming from four studies of a binary phenotype are as following:  $\theta_1 = 1.5$  ( $\chi^2 = 5.1$ ),  $\theta_2 = 1.3$  ( $\chi^2 = 2.2$ ),  $\theta_3 = 0.9$  ( $\chi^2 = 0.5$ ),  $\theta_4 = 1.2$  ( $\chi^2 = 3.1$ ).

Let us first transform the Odds Ratio to the logarithmic scale with

```

> or <- c(1.5, 1.3, 0.9, 1.2)
> lnor <- log(or)
> lnor

[1] 0.4054651 0.2623643 -0.1053605 0.1823216

```

To compute standard errors from known  $\chi^2$  values, one can use simple relation

$$\chi^2 = \frac{\beta^2}{s^2}$$

and thus

$$s^2 = \frac{\beta^2}{\chi^2}$$

Thus to compute the square standard errors corresponding to the log-Odds Ratio, we can use

```

> chi2or <- c(5.1, 2.2, 0.5, 3.1)
> s2lnor <- lnor * lnor/chi2or
> s2lnor

[1] 0.03223568 0.03128864 0.02220168 0.01072295

```

Now we can combine log-Odds Ratios and corresponding standard errors using inverse variance method:

```

> w <- 1/s2lnor
> plnor <- sum(w * lnor)/sum(w)
> plnor

[1] 0.1650462

> ps2 <- 1/sum(w)
> ps2

[1] 0.004968165

> pchi2 <- plnor * plnor/ps2
> pchi2

[1] 5.482958

> ppval <- 1 - pchisq(pchi2, 1)
> ppval

[1] 0.01920274

```

And the corresponding estimate of pooled Odds Ratio is

```

> exp(plnor)

[1] 1.179448

```

and 95% confidence interval is

```

> exp(plnor - 1.96 * sqrt(ps2))
[1] 1.02726
> exp(plnor + 1.96 * sqrt(ps2))
[1] 1.354181

```

Some times, effects are reported on different scales, and/or there may be suspect that these effects are not translatable across studies because of the differences in experimental design or for some other reasons. In this case, it may be better to poll the data without use of the effect estimate in exact manner, based only on the sign of association and its significance. This can be done by pooling Z-score values. Z-score refers to the test statistics, which has standard normal distribution under the null and can be derived e.g. by dividing estimate of the regression coefficient onto its standard error:

$$Z_i = \frac{\beta_i}{s_i}$$

The Z-score pooling methods can be derived from the inverse variance pooling by exploiting the fact that generally standard error of the estimate is proportional to  $1/\sqrt{n}$ , where  $n$  is the number of observations used for estimation. Therefore individual scores are assigned weights which are proportional to the square root of number of independent observations used in individual study,  $w_i = \sqrt{n_i}$ . The pooled Z-score statistics is computed as

$$Z = \frac{\sum_{i=0}^N w_i Z_i}{\sqrt{\sum_{i=0}^N w_i^2}}$$

We can now repeat the analysis of our first data set using Z-score pooling method. First, our data from table 1 are

```

> n <- c(225, 560, 437, 89)
> beta <- c(0.16, 0.091, 0.072, -0.03)
> s <- c(0.07, 0.042, 0.048, 0.12)

```

The Z-scores and weights are are:

```

> z <- beta/s
> z
[1] 2.285714 2.166667 1.500000 -0.250000

> w <- sqrt(n)
> w
[1] 15.000000 23.664319 20.904545 9.433981

```

The pooled estimate of Z-score is

```

> pz <- sum(w * z)/sqrt(sum(w^2))
> pz

```

```
[1] 3.163875
```

and corresponding  $P - value$  is

```
> 1 - pchisq(pz * pz, 1)
```

```
[1] 0.001556839
```

which is almost the same  $P - value$  we have obtained previously using the inverse variance method. Note, however, that now we do not know the "pooled" estimate of the regression coefficient.

Other important aspects of meta-analysis, such as heterogeneity, and a wide range of methods different from the inverse variance and Z-score based methods are not covered here, and we refer the reader to more epidemiologically-oriented literature for a better review.

## 9.2 Exercise: meta-analysis of literature data

In this exercise, you will perform meta-analysis of results collected from literature. These results resemble those obtained for association analysis between Pro12Ala polymorphism of the PPAR- $\gamma$  gene and type 2 diabetes. The data collected from literature are presented in the table 2.

Table 2: Summary of six studies of association between T2D and Pro12Ala polymorphism of the PPAR- $\gamma$  gene.  $n$ : number of subjects; effective allele: the allele for which the OR was estimated.

Study	Effective allele	$n$	$OR_E$	$P - value$
1	Ala	221	0.67	0.013
2	Pro	306	0.93	0.60
3	Pro	71	1.08	0.84
4	Ala	164	0.83	0.40
5	Pro	242	1.22	0.25
6	Pro	471	1.23	0.07

As you can see, only the original study report significant association, while other four are insignificant and one point in opposite direction.

Answer the following questions:

**Exercise 53** Perform meta-analysis of the data presented in table 2. Which allele is the risk one? Is this risk significant? What is pooled Odds Ratio and 95% confidence interval? Do analysis using at least two methods. Which method is better (best) in this situation? Why?

**Exercise 54** Perform meta-analysys excluding the original report (study 1). Is there still significant association between Pro12Ala and diabetes?

## 9.3 Reporting GWA results for future meta-analysis

In this section, we will discuss specifics of GWA analysis when meta-analysis is aimed at later stage. In order to perform meta-analysis at later stage, using either inverse variance or Z-score based method, you generally need to report

only effect estimates, standard errors of The estimates (or, equivalently,  $P$  – values or test statistics values), and number of observations used for estimation.

It is also clear that it is crucial to know for which allele the effect is reported, and this is the point where meta-analysis of genetic data may be very confusing. Generally, one may think that reporting what couple of nucleotide bases correspond to the polymorphism under the study and defining what allele was used as reference in the regression model may be enough. This, however, is not true for certain class of polymorphisms and may be a source of great confusion.

Consider a DNA molecule; as you know it is made of two complementary strands (forward or "+" and reverse or "-"). As you may guess, depending on the strand, what is an "A/G" polymorphism when reported on "+" strand becomes "T/C" polymorphism on the "-" strand (using complementarity rule  $A \leftrightarrow T$  and  $G \leftrightarrow C$ ). This is not a big problem for most of the polymorphism classes, because if say you know that for a first study  $\beta_1$  is reported for the "G" allele of the "A/G" polymorphism and in the second study  $\beta_2$  is the estimate of the effect "C" allele of the same polymorphism, but coded as "T/C" (thus other strand), you can easily spot that and say the "C" is the same as "G" in this situation, and pool the two betas straightforwardly.

However, for two types of polymorphisms, "A/T" and "G/C", where you can not get away without knowing what the strand was: what is reported as the effect of "T" in "A/T" polymorphism in study one; and seemingly corresponding effect of "T" in "A/T" polymorphism in study two may be apparently reports for two opposite alleles, if strands used for reporting in two studies were different.

The story may become even more complex, because the forward/reverse orientation depends on the genomic build<sup>4</sup>.

Thus if you want to pool your results with the results of others, there are quite a few SNP characteristics which are absolutely crucial to report, namely, the nucleotide bases describing the polymorphism, with indication which one was used as reference, and which one as "effective", strand, genomic build, and only than the effect estimate, standard error of the effect, and number of observations used to do estimation.

Other characteristics which are also recommended for reporting because they describe the quality characteristics of the sample and/or provide redundant information, which is good for double checks. Such characteristics include: frequency of the effective or reference allele, call rate, P-value for Hardy-Weinberg equilibrium and may be some parameter describing what is the direction of deviation from HWE (e.g.  $F_{max}$ ). When reporting results for imputed SNPs, more quality control characteristics should be included, suh as average maximal posterior probability,  $R^2$ , etc.

Let us start with arranging two data sets which could then be used for meta-analysis. Basically, we will use cleaned data from the GWA exercise you did in section 5 (Genome-wide association analysis, page 53), and split that is two parts.

Let us first load the data and re-name the data object:

```
> data(ge03d2ex.clean)
> data2 <- ge03d2ex.clean
> rm(ge03d2.clean)
```

---

<sup>4</sup>There is alternative, top/bottom, strand designation, which does not depend on genomic build. However, it is not always used.

and then split it in two parts:

```
> data2@gtdata@nids  
[1] 125  
  
> mdata1 <- data2[1:40, ]  
> mdata2 <- data2[41:data2@gtdata@nids, ]
```

We will analyse body mass index. If you pooling results of analysis of studies which are designed in approximately the same manner, you may think of reporting the effect estimates on the same scale and use of the inverse variance method for meta-analysis.

However, in meta-analysis of multiple data sets different individual studies are likely to assess different population, will use different designs, measure different covariates, and so on. Therefore you should think of some standardisation of the outcome variable (or apply Z-score method).

Therefore for the purpose of future meta-analysis, it becomes conventional to analyse pre-adjusted data which are scaled to Standard Normal (mean of zero and variance of unity). Note that this argument applies only to meta-analysis – you may and should report effects on the original scale (e.g. in centimeters and grams) in analysis of individual studies, in order to have better interpretability.

Moreover, in meta-analysis you heavily rely on the large numbers approximation when estimating *P-values*; while for individual study you can always apply empirical, e.g. permutation-based, procedures to derive the correct *P-value* whatever is the distribution of the trait, in meta-analysis the Normality assumption becomes crucial, and you do not want few outliers spoiling your results by screwing up *P-values*. Therefore some transformation improving normality is desirable. Note that transformation to Standard Normal does not improve the fit to normality; to do that other transformation should be applied. Probably the most famous transformations are log- and square root ones, then one may think of Box-Cox transformation. At the same time there is a transformation, called a Rank Transformation to Normality which guarantees perfect fit to Normal in absence of heavy ties<sup>5</sup>. We advocate the use of Rank Transformation to Normal for meta-analysis purposes.

GenABEL implements the `ztransform` function for the purposes of Z-transformation. This function takes two (actually three – see help for details) arguments: formula (or just the variable name) and data. `ztransform` function will perform (generalised) linear regression using the specified formula, and will transform the residuals from analysis onto Z-scale by subtracting the mean and division by the standard deviation.

Let us consider what this function does practically. Let us first transform BMI from the first set without using covariates:

```
> zbmio <- ztransform(bmi, mdata1)
```

The histogram of the transformed variable and scatter-plot of raw against transformed BMI is given at figure 19, column 1. Note that the fit to Normality is not improved by this transformation; with the original BMI, the Shapiro test for deviation from normality gives

---

<sup>5</sup>Ties are generated by the subjects with exactly the same trait values

```

> shapiro.test(mpta1@phdata$bmi)

Shapiro-Wilk normality test

data: mpta1@phdata$bmi
W = 0.9328, p-value = 0.01990

```

with identical results from the transformed variable:

```

> shapiro.test(zbmi0)

Shapiro-Wilk normality test

data: zbmi0
W = 0.9328, p-value = 0.01990

```

This is quite natural: as you can note from scatter-plot in column 1 of figure 19, only the centering and the spread of the scales are different for X (original BMI) and Y ( $x_0$ ), otherwise there is an exact linear correspondence between the two.

We can also do transformation using sex and age-adjusted residuals with

```
> zbm1 <- ztransform(bmi ~ sex + age, mpta1)
```

The scatter-plot of raw against transformed BMI is given at figure 19, column 2. Note that this transformation may slightly change the fit to Normal, which happens because we factor out the effects of sex and age:

```

> shapiro.test(zbm1)

Shapiro-Wilk normality test

data: zbm1
W = 0.9263, p-value = 0.01224

```

From the scatter-plot in column 2 of figure 19, it is quite clear what happens: the residuals from linear regression are not corresponding to the original BMI in exact linear manner.

A similar function, which performs rank-transformation to normality, is named **rnttransform**. For example if we want to adjust BMI for sex and age and rank-transform the residuals to Normal, we can use

```
> rnbm1 <- rnttransform(bmi ~ sex + age, mpta1)
```

This transformation, however, indeed improves the fit to Normal:

```

> shapiro.test(rnbm1)

Shapiro-Wilk normality test

data: rnbm1
W = 0.999, p-value = 1

```

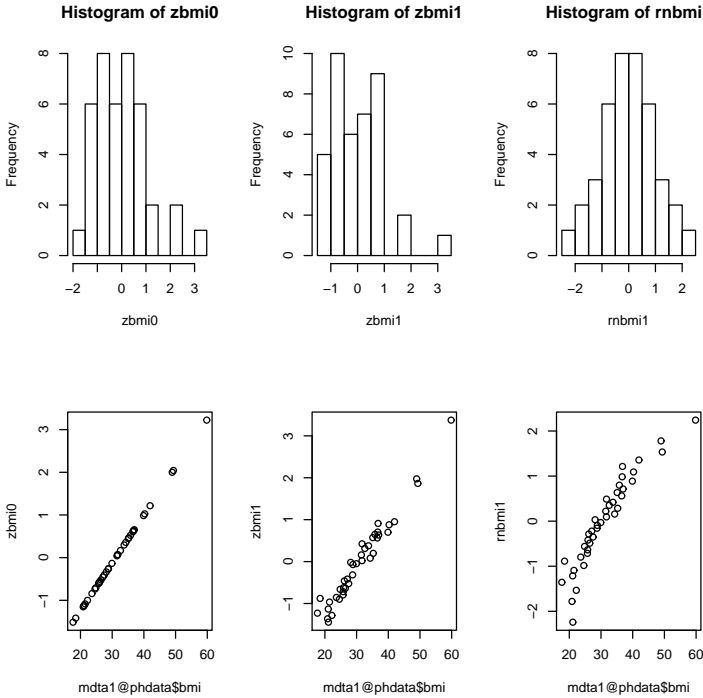


Figure 19: Histogram of transformed BMI and scatter-plots of the raw BMI against transformed BMI. Column 1: Z-transformation without covariates. Column 2: Z-transformation with adjustment for age and sex. Column 3: Rank-transformation to normality, after adjustment for sex and age.

In essence, the  $P - value$  of 1 means perfect fit to Normal – and this is what should have occurred when this transformation is used on the data without ties. Perfectly Normal distribution of the transformed trait may be enjoyed at the histogram presented at column 3 of figure 19.

Let us analyse Rank-Normal-transformed, sex and age-adjusted BMI in the two data sets, using `qtscore` function. Analysis of the first study is done with

```
> qts1 <- qtscore(rnbmi1, mdta1)
```

and analysis of the second study is done with

```
> zbm12 <- ztransform(bmi ~ sex + age, mdta2)
> qts2 <- qtscore(zbm12, mdta2)
```

The analysis looks very simple – is not it? However, the real difficulty did not start yet: now we need to extract coding, reference allele, strand, etc. – otherwise we can not do right meta-analysis later on!

Let us assume that you want to summarise the GW results from additive 1 d.f. test using following variables: (1) SNP name (2) chromosome (3) position (4) number of people with data available for this SNP test (5) effect of the allele (6) standard error of the effect (7) P-value for the test (8) corrected P-value (we will use Genomic Control) (9) coding, with reference allele coming first (10)

strand (11) frequency of the reference allele and, finally, (11) *P-value* from the exact test for HWE (that would be good for QC checks later on).

In the above list of output parameters, two are not directly available – frequency of the reference allele, which can be computed with

```
> refallfreq <- 1 - summary(mdata1@gtdata)$Q.2
```

and the standard error of the effect estimate. Though **GenABEL** does not report the s.e., it can be computed from the  $\chi^2$  value of the test. By definition,

$$\left( \frac{\beta}{SE_\beta} \right)^2 = \chi_1^2$$

and thus given value of the  $\chi^2$  test and knowing the effect, we can compute standard error with

$$SE_\beta = \frac{\beta}{\sqrt{\chi^2}}$$

Thus, s.e. is

```
> seeff <- qts1$effB/sqrt(qts1$chi2.1df)
```

At this moment we can arrange the required data frame:

```
> mdf1 <- data.frame(name = qts1$snpnames, chrom = qts1$chromosome,
+   pos = qts1$map, n = qts1$N, beta = qts1$effB, sebeta = seeff,
+   p = qts1$P1df, pgc = qts1$Pc1df, coding = as.character(mdata1@gtdata@coding),
+   strand = as.character(mdata1@gtdata@strand), reffreq = refallfreq)
```

Let us inspect the first 10 raws of the resulting output:

```
> mdf1[1:5, ]
```

		name	chrom	pos	n	beta	sebeta	p	pgc	
rs7435137	rs7435137			1	4259040	39	0.05838361	0.2141932	0.7851803	0.7851803
rs664063	rs664063			2	7288020	40	0.01829962	0.4705679	0.9689794	0.9689794
rs546570	rs546570			2	6120257	40	0.27341721	0.3890643	0.4822079	0.4822079
rs7908680	rs7908680			1	2311762	40	0.67913083	0.5908536	0.2503885	0.2503885
rs166732	rs166732			1	4716343	39	0.35418295	0.7213534	0.6234280	0.6234280
					coding	strand	reffreq			
rs7435137		CT		-	0.5128205					
rs664063		GC		-	0.9375000					
rs546570		TA		+	0.1000000					
rs7908680		CA		-	0.9625000					
rs166732		TG		-	0.9743590					

However, it is not recommended that you perform above-described reporting actions unless you develop your own format. In case if you plan to use **MetABEL** for meta-analysis, you best use **formetascore** function, which basically performs operations similar to described, and reports results in format compatible with **MetABEL**.

Thus, if you plan to use **MetABEL** for meta-analysis, required tables can be generated with single command:

```
> mdf1 <- formetascore(bmi ~ sex + age, mpta1, transform = rntransform)
```

You can see that results are the same as previously:

```
> mdf1[1:5, ]
```

		name	chromosome	position	strand	allele1	allele2	effallele
rs7435137	rs7435137		1	4259040	-	C	T	T
rs664063	rs664063		2	7288020	-	G	C	C
rs546570	rs546570		2	6120257	+	T	A	A
rs7908680	rs7908680		1	2311762	-	C	A	A
rs166732	rs166732		1	4716343	-	T	G	G
		effallelefreq	n	beta	sebeta	p	pgc	pexhwe
rs7435137		0.48717949	39	0.05838361	0.2141932	0.7851803	0.7851803	0.7484124
rs664063		0.06250000	40	0.01829962	0.4705679	0.9689794	0.9689794	1.0000000
rs546570		0.90000000	40	0.27341721	0.3890643	0.4822079	0.4822079	1.0000000
rs7908680		0.03750000	40	0.67913083	0.5908536	0.2503885	0.2503885	1.0000000
rs166732		0.02564103	39	0.35418295	0.7213534	0.6234280	0.6234280	1.0000000
		call						
rs7435137		0.975						
rs664063		1.000						
rs546570		1.000						
rs7908680		1.000						
rs166732		0.975						

To write all the data to a file, we can use standard R `write.csv` function:

```
> write.csv(mdf1, file = "RData/part1.rnbmisexage.csv", row.names = F)
```

Similar analysis is applied to the second data set:

```
> mdf2 <- formetascore(bmi ~ sex + age, mpta2, transform = rntransform)
```

We can inspect the first five lines of the output with

```
> mdf2[1:5, ]
```

		name	chromosome	position	strand	allele1	allele2	effallele
rs7435137	rs7435137		1	4259040	-	C	T	T
rs664063	rs664063		2	7288020	-	G	C	C
rs546570	rs546570		2	6120257	+	T	A	A
rs7908680	rs7908680		1	2311762	-	C	A	A
rs166732	rs166732		1	4716343	-	T	G	G
		effallelefreq	n	beta	sebeta	p	pgc	pexhwe
rs7435137		0.47619048	84	0.1701252	0.1658456	0.3049836	0.3239662	0.2729091
rs664063		0.11309524	84	0.2235208	0.2428365	0.3573334	0.3761317	1.0000000
rs546570		0.95180723	83	-0.2957072	0.3683184	0.4220571	0.4401322	1.0000000
rs7908680		0.01234568	81	-0.2475810	0.7150117	0.7291462	0.7391794	1.0000000
rs166732		0.02976190	84	-0.3871952	0.4576614	0.3975362	0.4159422	1.0000000
		call						
rs7435137		1.0000000						
rs664063		1.0000000						
rs546570		0.9880952						
rs7908680		0.9642857						
rs166732		1.0000000						

Let us write the data to a file:

```
> write.csv(mdf2, file = "RData/part2.rnrbmisexage.csv", row.names = F)
```

Finally let us analyse and save results for another data set, ge03d2c:

```
> data(ge03d2c)
> mdf3 <- formetascore(bmi ~ sex + age, ge03d2c, transform = rntransform)
> write.csv(mdf3, file = "RData/part3.rnrbmisexage.csv", row.names = F)
```

## 9.4 Meta-analysis with MetABEL

Now we are ready to meta-analyse GWA data coming from three studies. For this we will need to use **MetABEL** package, implementing simple meta-analysis functions for GWA data. Start with loading the package:

```
> library(MetABEL)
```

We will first meta-analyse the three studies using the data frames generated in previous section, **mdf1**, **mdf2** and **mdf3**. For this we will use the core function of **MetABEL**, **metagwa.tables**. This function takes four arguments: two data frames with results from individual studies, and two arguments supplying the study names. Pooling of multiple studies is possible by sequential application of this function.

Let us pool two first data frames:

```
> pooled <- metagwa.tables(mdf1, mdf2, name.x = "Part1", name.y = "Part2")
analysing ...
Lambda Part1 = 0.9418417
Lambda Part2 = 1.087699
Corrected Lambda Part1 = 0.9418417
Corrected Lambda Part2 = 1
Lambda POOLED data = 0.9980875
... DONE
```

The pooled data frame contains results of meta-analysis and essential details of the original studies:

```
> pooled[1:5, ]
```

	name	strand	allele1	allele2	effallele	chromosome	position	n	npops
1	rs100616	-	G	C	C	1	1911712	123	2
2	rs1006497	+	T	G	G	1	2658810	123	2
3	rs1011580	+	A	G	G	3	10048771	122	2
4	rs1011953	+	A	G	G	2	6464510	124	2
5	rs1013473	+	A	T	T	1	4487262	124	2

	beta	sebeta	effallelefreq	call	pexhwe	obetaPart1
1	0.11129808	0.1841270	0.1260163	0.9919667	4.7183287	0.08819818
2	-0.07517940	0.1931907	0.1463415	0.9920732	0.1698876	-0.25425751
3	-0.06827273	0.1393526	0.5040984	0.9839091	3.8552114	-0.06755996
4	0.07605436	0.1366477	0.3306452	1.0000000	0.1037354	0.18982068
5	0.30190019	0.1317848	0.5161290	1.0000000	0.0000000	0.31522388

```

    obetaPart2  osePart1  osePart2      chi2      p
1 0.118757384 0.3726827 0.2117793 0.3653763 0.54553545
2 0.003368901 0.3498751 0.2317182 0.1514346 0.69716811
3 -0.068805418 0.2130800 0.1842065 0.2400295 0.62418479
4 0.007546243 0.2228918 0.1729650 0.3097732 0.57781939
5 0.295031422 0.2259542 0.1622361 5.2480290 0.02197165

```

If one needs to pool more studies, this data frame should be used as the first argument of the `metagwa.tables`, and `name.x` argument should take special value "POOLED":

```

> pooled <- metagwa.tables(pooled, mdf3, name.x = "POOLED", name.y = "mdf3")

NA for betas in both populaions
18 SNPs removed
analysing ...
Lambda mdf3 = 1.125465
Corrected Lambda mdf3 = 1
Lambda POOLED data = 1.357255
... DONE

> pooled[1:5, ]

  name strand allele1 allele2 effallele chromosome position   n npops
1 rs1000475    +      T      C      C       X 13721802  91     1
2 rs1000909    -      A      G      G       2  8531681 190     1
3 rs1006092    -      T      G      G       X 13527448 190     1
4 rs100616     -      G      C      C       1  1911712 311     3
5 rs1006497    +      T      G      G       1 2658810 316     3

  beta    sebeta effallelefreq      call    pexhwe  obetaPart1
1 -0.05521349 0.11172204    0.5824176 0.4690722 5.9671619      NA
2 -0.08504871 0.13203441    0.8157895 0.9793814 2.8521518      NA
3 -0.03712065 0.08513945    0.5078947 0.9793814 6.8378385      NA
4  0.11689015 0.12052467    0.1237942 0.9781269 6.6661003 0.08819818
5 -0.02259297 0.11384385    0.1613924 0.9937663 0.2330977 -0.25425751

  obetaPart2    obetamdf3  osePart1  osePart2  osemfdf3      chi2      p
1        NA -0.055213494      NA      NA 0.11172204 0.24423775 0.6211620
2        NA -0.085048708      NA      NA 0.13203441 0.41491697 0.5194843
3        NA -0.037120652      NA      NA 0.08513945 0.19009450 0.6628380
4 0.118757384 0.121082405 0.3726827 0.2117793 0.15942451 0.94059776 0.3321242
5 0.003368901 0.005382408 0.3498751 0.2317182 0.14090855 0.03938469 0.8426885

```

This procedure may become quite laborious if multiple studies are to be pooled. In this case, it is possible to run meta-analysis using data provided in files, by applying function `metagwa.files`. As the first argument, this function takes the path to the directory where the files with results of individual studies are stored. It is assumed that the file names are made of two parts: population/study name and an extension. Thus the second argument of the `metagwa.files` function is the vector with names of studies, and the third one provides extension. Other arguments, "maf", "call" and "phwe" provide the threshold for QC filtering of SNPs in individual studies.

The function does not return any value, but rather creates a new file named POOLEDextens, where "extens" is the argument supplied to the function, in the source directory. To run analysis on the three files in the directory "RData" we can use

```
> metagwa.files(dir = "RData", pops = c("part1", "part2", "part3"),
+     extens = ".rnbmisexage.csv", maf = 1e-04, call = 0.93, phwe = 1e-08)

Population part1 , reading RData/part1.rnbmisexage.csv done
Dimensions after filters are 3678 15
population part2, reading RData/part2.rnbmisexage.csv done
Dimensions after filters are 3823 15
AT coding without +/-TOP/BOT strand info in part1
649 SNPs nullified
GC coding without +/-TOP/BOT strand info in part1
599 SNPs nullified
AT coding without +/-TOP/BOT strand info in part2
682 SNPs nullified
GC coding without +/-TOP/BOT strand info in part2
616 SNPs nullified
NA for betas in both populaions
1316 SNPs removed
analysing ...
Lambda part1 = 0.9170982
Lambda part2 = 1.081238
Corrected Lambda part1 = 0.9170982
Corrected Lambda part2 = 1
Lambda POOLED data = 0.9485962
... DONE
Dimensions after pooling are 2552 20
population part3, reading RData/part3.rnbmisexage.csv done
Dimensions after filters are 7493 15
AT coding without +/-TOP/BOT strand info in part3
1277 SNPs nullified
GC coding without +/-TOP/BOT strand info in part3
1210 SNPs nullified
NA for betas in both populaions
2487 SNPs removed
analysing ...
Lambda part3 = 1.128227
Corrected Lambda part3 = 1
Lambda POOLED data = 1.337598
... DONE
Dimensions after pooling are 5006 22
$analysed.pops
[1] "part1" "part2" "part3"

> poolf <- read.csv("RData/POOLED.rnbmisexage.csv", strings = F)
> poolf[1:5, ]

      name strand allele1 allele2 effallele chromosome position    n npops
1 rs1000909        0         A         G         G        2 8531681 190       1
```

```

2 rs1006092      0      T      G      G      X 13527448 190      1
3 rs1006497      0      T      G      G      1 2658810 316      3
4 rs1010481      0      A      C      C      2 8409087 190      1
5 rs1011580      0      A      G      G      3 10048771 313      3
          beta     sebeta effallelefreq      call      pexhwe obetapart1
1 -0.08504871 0.13219628    0.8157895 0.9793814 2.8521518      NA
2 -0.03712065 0.08524383    0.5078947 0.9793814 6.8378385      NA
3 -0.02260022 0.11385269    0.1613924 0.9937663 0.2330977 -0.25425751
4 -0.03241475 0.12378575    0.2657895 0.9793814 0.0000000      NA
5 -0.09718027 0.08691427    0.5223642 0.9842917 4.6164715 -0.06755996
          obetapart2 obetapart3 osepart1 osepart2 osepart3      chi2      p
1           NA -0.085048708      NA      NA 0.13219628 0.41390146 0.5199959
2           NA -0.037120652      NA      NA 0.08524383 0.18962924 0.6632254
3  0.003368901 0.005382408 0.3498751 0.2310290 0.14108131 0.03940385 0.8426507
4           NA -0.032414750      NA      NA 0.12378575 0.06857156 0.7934281
5 -0.068805418 -0.115686832 0.2130800 0.1836586 0.11131267 1.25018424 0.2635173

```

## 10 Analysis of selected region

### 10.1 Exploring linkage disequilibrium

See help for `r2fast`.

### 10.2 Haplotype analysis

Use

```
> gtfor1d <- as.hsgeno(srdta[, 1:5])
```

to convert part of your SNPs to `haplo.stats` format.

You can also use interface function to do sliding widow analysis

```
> h2 <- scan.haplo("qt1~CRSNP", srdta, snps = c(1:5))
```

### 10.3 Analysis of interactions

See help for `scan.haplo.2D` and `scan.glm.2D`

## A Importing data to GenABEL

As described in section 4.1, **GenABEL** gwaa.data-class consist of phenotypic data frame and an object of.snp.data-class, which contains all genetic data. To import data to **GenABEL**, you need to prepare two files: one containing the phenotypic, and the other containing genotypic data.

The phenotype file relates study subject IDs with values of covariates and outcomes. In the phenotypic data file, the first line gives a description (variable name) of the data contained in a particular column; the names should better be unique, otherwise R will change them.

The first column of the phenotype file **must** contain the subjects' unique ID, named "id". The IDs listed here, and in the genotypic data file, must be the same. It is recommended that the id names are given in quotation marks (see example below), which will save you a possible troubles with e.g. leading zeros.

There also should also be a column named "sex" and giving sex information (0 = female, 1 = male). Other columns in the file should contain phenotypic information.

Missing values should be coded with "NA"; binary traits should have values 0 or 1.

All subjects present in the genotypic files **must** be listed in the phenotypic file as well, because sex information provided by the phenotypic file is an essential part of the genotypic QC procedure.

An example of few first lines of a phenotype file is as follows:

id	sex	age	bt1	qt	qt1
"cd289982"	0	30.33	NA	NA	3.93
"cd325285"	0	36.514	1	0.49	3.61
"cd357273"	1	37.811	0	1.65	5.30
"cd872422"	1	20.393	0	1.95	4.07
"cd1005389"	1	28.21	1	0.35	3.90

This file tells us that, for example, person 325286 is female (0 in second column), and she has "1" (usually this means a "case") value for the trait "bt1", so on. Person 289982 has measurements only for sex, age and qt1, while other measurements are missing (NA, Not Available).

If you need to add phenotypes to a **gwaa.data-class** object already created, you can use function **add.phdata**. This function allows you to add variables contained in some data frame to the existing **data@phdata** object. The data frame to be added should contain "id" variable, identical to that existing in the object, and **should not** contain any other variables with names identical to these already existing.

The second file you need should contains genotypic data. As described in section 4.1 (General description of **gwaa.data-class**, page 37), **GenABEL** **snp.data-class** contains different types of information. For every SNP, information on map position, chromosome, and strand should be provided. For every person, every SNP genotype should be provided. **GenABEL** provides a number of function to convert these data from different formats to the internal **GenABEL** raw format. We will first consider our preferred format, which we informally call "Illumina"-like.

## A.1 Converting from preferred format

We will consider use of `convert.snp.illumina` procedure; details of other procedures are given later. Note that what we call "illumina" format is not really a proprietary format from that company, it is just one of the possible text output format from the Illumina BeadStudio; similar formats are accepted/generated by HapMap and Affymetrix.

The file of the "Illumina" format contains SNPs in rows and IDs in columns. The first line is a "header", containing column names. The first three columns should contain information on SNP name, chromosome, and position. There is an optional (though highly recommended!) fourth column, containing strand information (acceptable codes: "+", "-", "u", the last stands for "unknown"). After that column, each of the residual ones corresponds to an individual, with ID as the column name. Genotypes should be presented by two consecutive characters (no separator).

An example of few first lines of the "illumina" genotypic file is as follows:

name	chr	pos	strand	"cd289982"	"cd325285"	"cd357273"	"cd872422"	"cd1005389"
rs1001	1	1235	+	AA	AG	AG	AA	GG
rs6679	9	2344	+	GT	GG	GG	TG	GG
rs2401	22	3455	+	AA	CC	CC	CC	AC
rs123	X	32535	-	TT	GT	TT	TT	TT
rs6679	XY	2344	-	GT	GG	GG	TG	GG
rs876	Y	23556	+	OO	OO	TT	GG	TT
mitoA1	mt	24245	-	AA	CC	OO	OO	OO

It is clear that is not quite conventional Illumina file – because in BeadStudio the alleles are reported using the "top" strand; rather, this is an Affymetrix or HapMap-type of a file. Anyways, this file contains all required genotypic information, and this file format is the preferred one for import. Assume that the file with the genotypic data is called "gen0.illu", and is stored in the directory "RData". You can convert the data to GenABEL raw format by

```
> convert.snp.illumina(inf = "RData/gen0.illu", out = "RData/gen0i.raw",
+   strand = "file")
```

Reading genotypes from file 'RData/gen0.illu' ...  
Writing to file 'RData/gen0i.raw' ...  
... done.

Here is the content of the converted file "gen0i.raw" – internal raw data representation:

```
#GenABEL raw data version 0.1
"cd289982" "cd325285" "cd357273" "cd872422" "cd1005389"
rs1001 rs6679 rs2401 rs123 rs6679 rs876 mitoA1
1 9 22 X XY Y mt
1235 2344 3455 32535 2344 23556 24245
04 0c 0f 08 0c 08 0f
01 01 01 02 02 01 02
69 c0
96 40
```

```
d5 80
65 40
96 40
07 40
d0 00
```

Note the option `strand="file"` – it is telling `GenABEL` that strand information is provided in the file.

At this moment, you can load the data into `GenABEL`. Assume that the phenotypic file described above is called "phe0.dat" and the converted genotypic file in the raw `GenABEL` format is called "gen0i.raw". You can load the data using the command

```
> df <- load.gwaa.data(phe = "RData/phe0.dat", gen = "RData/gen0i.raw",
+ force = T)

ids loaded...
marker names loaded...
chromosome data loaded...
map data loaded...
allele coding data loaded...
strand data loaded...
genotype data loaded...
snp.data object created...
assignment of gwaa.data object FORCED; X-errors were not checked!
```

The option "force=TRUE" tells that `GenABEL` should load the data even if it finds sex errors.

You can inspect the loaded data; let us first look into phenotypic data by

```
> df@phdata
```

		id	sex	age	bt1	qt	qt1
cd289982	cd289982	cd289982	0	30.330	NA	NA	3.93
cd325285	cd325285	cd325285	0	36.514	1	0.49	3.61
cd357273	cd357273	cd357273	1	37.811	0	1.65	5.30
cd872422	cd872422	cd872422	1	20.393	0	1.95	4.07
cd1005389	cd1005389	cd1005389	1	28.210	1	0.35	3.90

... and than check that the genotypes have imported right:

```
> g0 <- as.character(df@gtdata)
> g0

rs1001 rs6679 rs2401 mitoA1 rs123 rs6679 rs876
cd289982 "A/A" "G/T" "A/A" "A/A" "T/T" "G/T" NA
cd325285 "A/G" "G/G" "C/C" "C/C" "T/G" "G/G" NA
cd357273 "A/G" "G/G" "C/C" NA "T/T" "G/G" "T/T"
cd872422 "A/A" "G/T" "C/C" NA "T/T" "G/T" "G/G"
cd1005389 "G/G" "G/G" "C/A" NA "T/T" "G/G" "T/T"

> as.character(df@gtdata@strand)
```

```

rs1001 rs6679 rs2401 mitoA1  rs123 rs6679  rs876
 "+"    "+"    "+"    "-"    "--"   "--"   "+"
> as.character(df@gtdata@coding)

rs1001 rs6679 rs2401 mitoA1  rs123 rs6679  rs876
 "AG"   "GT"   "CA"   "CA"   "TG"   "GT"   "TG"

```

In a real Illumina file, a coding on the TOP strand is supplied. Then, the file will normally look like

name	chr	pos	"cd289982"	"cd325285"	"cd357273"	"cd872422"	"cd1005389"
rs1001	1	1235	AA	AG	AG	AA	GG
rs6679	9	2344	GT	GG	GG	TG	GG
rs2401	22	3455	AA	CC	CC	CC	AC
rs123	X	32535	TT	GT	TT	TT	TT
rs6679	XY	2344	GT	GG	GG	TG	GG
rs876	Y	23556	00	00	TT	GG	TT
mitoA1	mt	24245	AA	CC	00	00	00

and the conversion command will be

```

> convert.snp.illumina(inf = "RData/gen0.illuwos", out = "RData/gen0iwos.raw",
+   strand = "+")
Reading genotypes from file 'RData/gen0.illuwos' ...
Writing to file 'RData/gen0iwos.raw' ...
... done.

```

In this particular data set, after conversion, the "+" strand will actually mean not "forward", but TOP – something you should remember for this particular data. The resulting file will look like this:

You can load the data with

```

> df <- load.gwaa.data(phe = "RData/phe0.dat", gen = "RData/gen0iwos.raw",
+   force = T)
ids loaded...
marker names loaded...
chromosome data loaded...
map data loaded...
allele coding data loaded...
strand data loaded...
genotype data loaded...
snp.data object created...
assignment of gwaa.data object FORCED; X-errors were not checked!

```

Obviously, the "strand" is always "+" (here it means TOP):

```

> g1 <- as.character(df@gtdata)
> g1

```

```

rs1001 rs6679 rs2401 mitoA1 rs123 rs6679 rs876
cd289982 "A/A" "G/T" "A/A" "A/A" "T/T" "G/T" NA
cd325285 "A/G" "G/G" "C/C" "C/C" "T/G" "G/G" NA
cd357273 "A/G" "G/G" "C/C" NA "T/T" "G/G" "T/T"
cd872422 "A/A" "G/T" "C/C" NA "T/T" "G/T" "G/G"
cd1005389 "G/G" "G/G" "C/A" NA "T/T" "G/G" "T/T"

> as.character(df@gtdata@strand)

rs1001 rs6679 rs2401 mitoA1 rs123 rs6679 rs876
 "+" "+" "+" "+" "+" "+" "+"

> as.character(df@gtdata@coding)

rs1001 rs6679 rs2401 mitoA1 rs123 rs6679 rs876
 "AG" "GT" "CA" "CA" "TG" "GT" "TG"

```

We can see that the genotypes are identical to ones we imported previously, as should be the case:

```

> g0 == g1

rs1001 rs6679 rs2401 mitoA1 rs123 rs6679 rs876
cd289982 TRUE TRUE TRUE TRUE TRUE TRUE NA
cd325285 TRUE TRUE TRUE TRUE TRUE TRUE NA
cd357273 TRUE TRUE TRUE NA TRUE TRUE TRUE
cd872422 TRUE TRUE TRUE NA TRUE TRUE TRUE
cd1005389 TRUE TRUE TRUE NA TRUE TRUE TRUE

```

## A.2 Converting PLINK tped files

PLINK TPED (transposed-pedigree) format bears notable similarity to what we call "Illumina" format, with few exceptions: (1) there is no header line giving field names (and therefore IDs are stored in a separate file) (2) the first column gives chromosome, second – SNP name, third *genetic* map (usually kept as zeroes), the fourth – physical position, and, starting with the fifth column, genotypic data are listed, (3) finally, within a genotype, alleles are separated with a space. In TPED format, the data we already worked with would look like

```

1  rs1001  0   1235  A A  A G  A G  A A  G G
9  rs6679  0   2344  G T  G G  G G  T G  G G
22 rs2401  0   3455  A A  C C  C C  C C  A C
X   rs123  0   32535 T T  G T  T T  T T  T T
XY  rs6679  0   2344  G T  G G  G G  T G  G G
Y   rs876  0   23556 O O  O O  T T  G G  T T
mt  mitoA1 0   24245 A A  C C  O O  O O  O O

```

Obviously, a separate file is needed to keep correspondence between genotypes and IDs. This file emulated standard pedigree file without a header line. The file, conventionally called a TFAM-file, should contain six columns, corresponding to pedigree ID, ID, father, mother, sex, and affection. Only the

second column is used by **GenABEL** – please make sure you use unique IDs. Consequently, it does not matter what pedigree ID, father/mother, sex, or affection status you assign in the file – the real information is coming from the phenotypic data file. The TFAM file for our data will look like this:

```
1 cd289982 0 0 1 0
1 cd325285 0 0 1 0
1 cd357273 0 0 1 0
1 cd872422 0 0 1 0
1 cd1005389 0 0 1 0
```

You can convert the data from PLINK TPED format to the **GenABEL** format using command `convert.snp.tped`:

```
> convert.snp.tped(tped = "RData/gen0.tped", tfam = "RData/gen0.tfam",
+       out = "RData/gen0tped.raw", strand = "+")

Reading individual ids from file 'RData/gen0.tfam' ...
... done. Read 5 individual ids from file 'RData/gen0.tfam'
Reading genotypes from file 'RData/gen0.tped' ...
...done. Read 7 SNPs from file 'RData/gen0.tped'
Writing to file 'RData/gen0tped.raw' ...
... done.
```

and load the data with

```
> df <- load.gwaa.data(phe = "RData/phe0.dat", gen = "RData/gen0tped.raw",
+   force = T)

ids loaded...
marker names loaded...
chromosome data loaded...
map data loaded...
allele coding data loaded...
strand data loaded...
genotype data loaded...
snp.data object created...
assignment of gwaa.data object FORCED; X-errors were not checked!
```

Obviously, the "strand" is always "+" (meaning TOP):

```
> g1 <- as.character(df@gtdata)
> g1

rs1001 rs6679 rs2401 mitoA1 rs123 rs6679 rs876
cd289982 "A/A" "G/T" "A/A" "A/A" "T/T" "G/T" NA
cd325285 "A/G" "G/G" "C/C" "C/C" "T/G" "G/G" NA
cd357273 "A/G" "G/G" "C/C" NA "T/T" "G/G" "T/T"
cd872422 "A/A" "G/T" "C/C" NA "T/T" "G/T" "G/G"
cd1005389 "G/G" "G/G" "C/A" NA "T/T" "G/G" "T/T"

> as.character(df@gtdata@strand)
```

```
rs1001 rs6679 rs2401 mitoA1 rs123 rs6679 rs876
 "+"   "+"   "+"   "+"   "+"   "+"   "+"
```

```
> as.character(df@gtdata@coding)
```

```
rs1001 rs6679 rs2401 mitoA1 rs123 rs6679 rs876
 "AG"  "GT"  "CA"  "CA"  "TG"  "GT"  "TG"
```

We can see that the genotypes are identical to ones we imported previously, as should be the case:

```
> g0 == g1
```

```
rs1001 rs6679 rs2401 mitoA1 rs123 rs6679 rs876
cd289982 TRUE  TRUE  TRUE  TRUE  TRUE  TRUE  NA
cd325285 TRUE  TRUE  TRUE  TRUE  TRUE  TRUE  NA
cd357273 TRUE  TRUE  TRUE  NA  TRUE  TRUE  TRUE
cd872422 TRUE  TRUE  TRUE  NA  TRUE  TRUE  TRUE
cd1005389 TRUE  TRUE  TRUE  NA  TRUE  TRUE  TRUE
```

### A.3 Converting linkage-like files

Linkage-like files, also known as pre-makeped files, or pedigree files, represent a historic format which dates back to the time when only few markers could be typed – thus the number of subjects was usually greater than the number of markers. In that situation, it was natural and obvious to keep IDs in rows and markers in columns. In the first six columns, standard linkage-like file would contain pedigree ID, ID, father's ID, mother's ID, sex (coded as 1 = male and 2 = female), and affection status (0 = unknown, 1 = unaffected, 2 = affected). In the following columns, genotypic information is provided. Alleles of the same genotype could be separated by a space, or by a slash ("/"). Thus the data we are working with could be presented as

```
1 cd289982 0 0 1 0 A A G T A A T T G T 0 0 A A
1 cd325285 0 0 1 0 A G G G C C G T G G 0 0 C C
1 cd357273 0 0 1 0 A G G G C C T T G G T T 0 0
1 cd872422 0 0 1 0 A A T G C C T T T G G G G 0 0
1 cd1005389 0 0 1 0 G G G G A C T T G G T T 0 0
```

As you can see, this file misses header line, and information what are the SNP names, position, etc. should be provided in a separate MAP-file. GenABEL accepts map in Merlin format, and an extended format. A map in Merlin format consist of header line, giving column names, and three columns with chromosome, name and position information, for example:

chr	name	pos
1	rs1001	1235
9	rs6679	2344
22	rs2401	3455
X	rs123	32535
XY	rs6679	2344
Y	rs876	23556
mt	mitoA1	24245

The data can be converted to the internal GenABEL format with

```
> convert.snp.ped(ped = "RData/gen0.ped", map = "RData/map0.dat",
+      out = "RData/gen0pedwos.raw", strand = "+")  
  
Reading map from file 'RData/map0.dat' ...
... done. Read positions of 7 markers from file 'RData/map0.dat'
Reading genotypes from file 'RData/gen0.ped' ...
...done. Read information for 5 people from file 'RData/gen0.ped'
Analysing marker information ...
Writing to file 'RData/gen0pedwos.raw' ...
... done.
```

and loaded with

```
> df <- load.gwaa.data(phe = "RData/phe0.dat", gen = "RData/gen0pedwos.raw",
+      force = T)  
  
ids loaded...
marker names loaded...
chromosome data loaded...
map data loaded...
allele coding data loaded...
strand data loaded...
genotype data loaded...
snp.data object created...
assignment of gwaa.data object FORCED; X-errors were not checked!
```

We can inspect the genotypic data and check that conversion results are identical to previous runs with

```
> g1 <- as.character(df@gtdata)
> g1  
  
rs1001 rs6679 rs2401 mitoA1 rs123 rs6679 rs876
cd289982 "A/A" "G/T" "A/A" "A/A" "T/T" "G/T" NA
cd325285 "A/G" "G/G" "C/C" "C/C" "T/G" "G/G" NA
cd357273 "A/G" "G/G" "C/C" NA "T/T" "G/G" "T/T"
cd872422 "A/A" "G/T" "C/C" NA "T/T" "G/T" "G/G"
cd1005389 "G/G" "G/G" "C/A" NA "T/T" "G/G" "T/T"  
  
> as.character(df@gtdata@strand)  
  
rs1001 rs6679 rs2401 mitoA1 rs123 rs6679 rs876
 "+"     "+"     "+"     "+"     "+"     "+"  
  
> as.character(df@gtdata@coding)  
  
rs1001 rs6679 rs2401 mitoA1 rs123 rs6679 rs876
 "AG"    "GT"   "CA"   "CA"   "TG"   "GT"   "TG"
```

```
> g0 == g1
```

	rs1001	rs6679	rs2401	mitoA1	rs123	rs6679	rs876
cd289982	TRUE	TRUE	TRUE	TRUE	TRUE	TRUE	NA
cd325285	TRUE	TRUE	TRUE	TRUE	TRUE	TRUE	NA
cd357273	TRUE	TRUE	TRUE	NA	TRUE	TRUE	TRUE
cd872422	TRUE	TRUE	TRUE	NA	TRUE	TRUE	TRUE
cd1005389	TRUE	TRUE	TRUE	NA	TRUE	TRUE	TRUE

If you are willing to import strand information, you can make use of the *extended* map format. In this format the strand information is added to the map-file:

chr	name	pos	strand	coding
1	rs1001	1235	+	AG
9	rs6679	2344	+	TG
22	rs2401	3455	+	AC
X	rs123	32535	-	GT
XY	rs6679	2344	-	GT
Y	rs876	23556	+	GT
mt	mitoA1	24245	-	AC

The data can be converted to the internal GenABEL format with

```
> convert.snp.ped(ped = "RData/gen0.ped", map = "RData/emap0.dat",
+      out = "RData/gen0ped.raw", strand = "file")

Reading map from file 'RData/emap0.dat' ...
... done. Read positions of 7 markers from file 'RData/emap0.dat'
Reading genotypes from file 'RData/gen0.ped' ...
...done. Read information for 5 people from file 'RData/gen0.ped'
Analysing marker information ...
Writing to file 'RData/gen0ped.raw' ...
... done.
```

Note that option `strand==file` was used to specify that the extended map format should be used. The data can be loaded with

```
> df <- load.gwaa.data(phe = "RData/phe0.dat", gen = "RData/gen0ped.raw",
+   force = T)

ids loaded...
marker names loaded...
chromosome data loaded...
map data loaded...
allele coding data loaded...
strand data loaded...
genotype data loaded...
snp.data object created...
assignment of gwaa.data object FORCED; X-errors were not checked!
```

We can inspect the genotypic data and check that conversion results are identical to previous runs with

```
> g1 <- as.character(df@gtdata)
> g1
```

```

rs1001 rs6679 rs2401 mitoA1 rs123 rs6679 rs876
cd289982 "A/A" "G/T" "A/A" "A/A" "T/T" "G/T" NA
cd325285 "A/G" "G/G" "C/C" "C/C" "T/G" "G/G" NA
cd357273 "A/G" "G/G" "C/C" NA "T/T" "G/G" "T/T"
cd872422 "A/A" "G/T" "C/C" NA "T/T" "G/T" "G/G"
cd1005389 "G/G" "G/G" "C/A" NA "T/T" "G/G" "T/T"

> as.character(df@gtdata@strand)

rs1001 rs6679 rs2401 mitoA1 rs123 rs6679 rs876
 "+" "+" "+" "-" "-" "-" "+"

> as.character(df@gtdata@coding)

rs1001 rs6679 rs2401 mitoA1 rs123 rs6679 rs876
 "AG" "GT" "CA" "CA" "TG" "GT" "TG"

> g0 == g1

rs1001 rs6679 rs2401 mitoA1 rs123 rs6679 rs876
cd289982 TRUE TRUE TRUE TRUE TRUE TRUE NA
cd325285 TRUE TRUE TRUE TRUE TRUE TRUE NA
cd357273 TRUE TRUE TRUE NA TRUE TRUE TRUE
cd872422 TRUE TRUE TRUE NA TRUE TRUE TRUE
cd1005389 TRUE TRUE TRUE NA TRUE TRUE TRUE

```

#### A.4 Converting from MACH format

The data from MACH format can be converted by using `convert.snp.mach`. This function actually calls `convert.snp.ped` in specific format. MACH software is widely used for SNP imputations. For our needs we consider two files produced by MACH: pedigree file with (the imputed) genotypic data, and info-file, containing information about quality of imputations for particular SNP.

SEE HELP FOR `convert.snp.mach` for further details.

#### A.5 Converting from text format

## B Answers to exercises

### B.1 Exercise 3:

For the first person id is "p1" and sex code is 1 (1=male, 2=female)

```
> srdta@gtdata@idnames[1]  
[1] "p1"  
  
> srdta@gtdata@male[1]  
  
p1  
1
```

For the 22nd person id is "p22" and sex code is 1:

```
> srdta@gtdata@idnames[22]  
[1] "p22"  
  
> srdta@gtdata@male[22]  
  
p22  
1
```

Among first 100 subjects, there are 53 males:

```
> sum(srdta@gtdata@male[1:100])  
[1] 53
```

Among 4th hundred subjects there are 45 females:

```
> 100 - sum(srdta@gtdata@male[301:400])  
[1] 45
```

Male proportion among first 1000 people is

```
> mean(srdta@gtdata@male[1:1000])  
[1] 0.508
```

Female proportion among second 1000 people is

```
> 1 - mean(srdta@gtdata@male[1001:2000])  
[1] 0.476
```

Name, chromosome and map position of the 33rd marker are:

```
> srdta@gtdata@snpnames[33]  
[1] "rs422"  
  
> srdta@gtdata@chromosome[33]
```

```

rs422
 1
Levels: 1

> srdta@gtdata@map[33]

rs422
105500

```

The map positions for and distance between markers 25 and 26 are:

```

> pos25 <- srdta@gtdata@map[25]
> pos25

```

```

rs365
91250

```

```

> pos26 <- srdta@gtdata@map[26]
> pos26

```

```

rs372
92750

```

```

> pos26 - pos25

```

```

rs372
1500

```

## B.2 Exercise 6:

Value of the 4th variable of person 75:

```

> srdta@phdata[75, 4]

```

```

[1] -0.04

```

Value for the variable 1 is

```

> srdta@phdata[75, 1]

```

```

[1] "p75"

```

Also, if we check first 10 elements we see

```

> srdta@phdata[1:10, 1]

```

```

[1] "p1"  "p2"  "p3"  "p4"  "p5"  "p6"  "p7"  "p8"  "p9"  "p10"

```

This is personal ID.

The sum for variable 2 is

```

> sum(srdta@phdata[, 2])

```

```

[1] 1275

```

This is sex variable.

### B.3 Exercise 7:

To obtain the number of people with age >65 y.o., you can use any of the following

```
> sum(srdta@phdata$age > 65)
[1] 48

> vec <- which(srdta@phdata$age > 65)
> length(vec)

[1] 48
```

To get sex of these people use any of:

```
> sx65 <- srdta@phdata$sex[srdta@phdata$age > 65]
> sx65

[1] 1 1 0 0 0 0 1 1 1 1 0 0 1 1 0 1 1 1 0 0 0 0 1 1 1 0 1 1 1 1 0 1 1 1 0 0 0 0
[39] 1 0 1 0 0 0 0 1 1 1

> sx65 <- srdta@phdata$sex[vec]
> sx65

[1] 1 1 0 0 0 0 1 1 1 1 0 0 1 1 0 1 1 1 0 0 0 0 1 1 1 0 1 1 1 1 0 1 1 1 0 0 0 0
[39] 1 0 1 0 0 0 0 1 1 1
```

Thus, number of males is:

```
> sum(sx65)
```

```
[1] 26
```

To conclude, the proportion of male is 0.541666666666667

Distribution of qt3 in people younger and older than 65 are:

```
> summary(srdta@phdata$qt3[vec])

   Min. 1st Qu. Median     Mean 3rd Qu.    Max.
0.730   2.690   3.480   3.499   4.265   5.840

> sd(srdta@phdata$qt3[vec], na.rm = TRUE)

[1] 1.128701

> young <- which(srdta@phdata$age < 65)
> summary(srdta@phdata$qt3[young])

   Min. 1st Qu. Median     Mean 3rd Qu.    Max.    NA's
-1.97   1.83   2.58   2.59   3.35   6.34   11.00

> sd(srdta@phdata$qt3[young], na.rm = TRUE)

[1] 1.093374
```

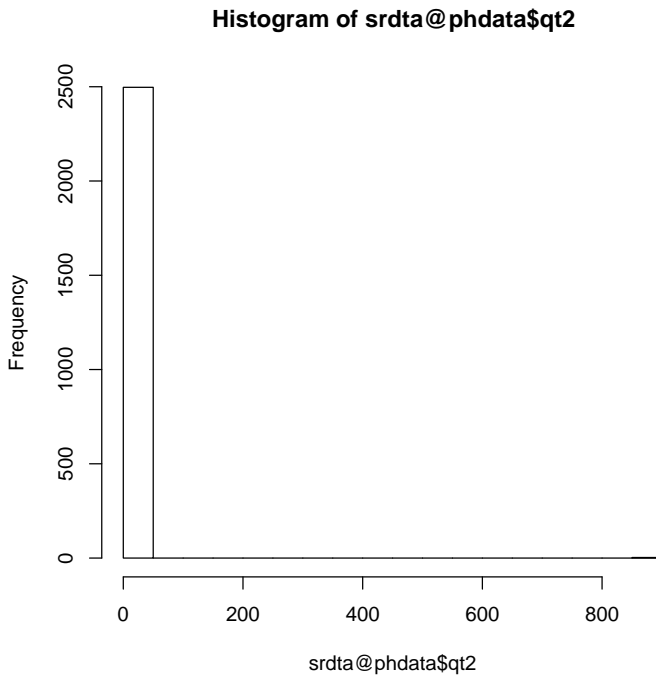


Figure 20: Histogram of qt2

#### B.4 Exercise 9:

```
> summary(srdta@phdata$age)
   Min. 1st Qu. Median   Mean 3rd Qu.   Max.
24.10  45.10  50.00  50.04  54.80  71.60
```

The histogram for qt2 looks strange: it seems there are few very strong outliers (figure 20) You can also see that with `summary`:

```
> summary(srdta@phdata$qt2)
   Min. 1st Qu. Median   Mean 3rd Qu.   Max.
0.000  4.220  5.045  6.122  5.910 888.000
```

#### B.5 Exercise 10:

How many SNPs are described in this data frame?

```
> attach(popdat)
The following object(s) are masked from data2@phdata :
sex
> names(popdat)
```

```
[1] "subj"   "sex"    "aff"     "qt"      "snp1"   "snp2"   "snp3"   "snp4"   "snp5"  
[10] "snp6"   "snp7"   "snp8"   "snp9"   "snp10"
```

The answer is 10 snps

### B.6 Exercise 11:

What is the frequency (proportion) of snp1 allele A? What is its frequency in these affected (aff==1)?

```
> summary(snp1)
```

Number of samples typed: 2374 (95%)

Allele Frequency: (2 alleles)

	Count	Proportion
A	3462	0.73
B	1286	0.27
NA	252	NA

Genotype Frequency:

	Count	Proportion
B/B	199	0.08
A/B	888	0.37
A/A	1287	0.54
NA	126	NA

Heterozygosity (Hu) = 0.3950646

Poly. Inf. Content = 0.3169762

The frequency of A in all subjects is 0.73.

```
> summary(snp1[aff == 1])
```

Number of samples typed: 519 (94.5%)

Allele Frequency: (2 alleles)

	Count	Proportion
A	729	0.7
B	309	0.3
NA	60	NA

Genotype Frequency:

	Count	Proportion
B/B	48	0.09
A/B	213	0.41
A/A	258	0.50
NA	30	NA

Heterozygosity (Hu) = 0.4185428

Poly. Inf. Content = 0.3307192

The frequency of A in affected subjects is 0.7.

### B.7 Exercise 12:

How many cases and controls are present?

```
> table(aff)
```

aff	
0	1
1951	549

There are 549 cases and 1951 controls.

### B.8 Exercise 13:

If all subjects are used to test HWE, are there any SNPs out of HWE at nominal  $P \leq 0.05$ ? Which ones?

```
> HWE.exact(snp1)
```

```
Exact Test for Hardy-Weinberg Equilibrium
```

```
data: snp1
N11 = 1287, N12 = 888, N22 = 199, N1 = 3462, N2 = 1286, p-value =
0.01083
```

```
...
```

```
> HWE.exact(snp10)
```

```
Exact Test for Hardy-Weinberg Equilibrium
```

```
data: snp10
N11 = 1792, N12 = 552, N22 = 40, N1 = 4136, N2 = 632, p-value = 0.7897
```

Only SNP 1 is out of HWE in the total sample.

### B.9 Exercise 14:

If only controls are used to test the SNPs which are out of HWE in total sample, are these still out of HWE?

```
> HWE.exact(snp1[aff == 0])
```

```
Exact Test for Hardy-Weinberg Equilibrium
```

```
data: snp1[aff == 0]
N11 = 1029, N12 = 675, N22 = 151, N1 = 2733, N2 = 977, p-value =
0.008393
```

Yes, SNP 1 is out of HWE also in controls.

## B.10 Exercise 15:

Which SNP pairs are in strong LD ( $r^2 \geq 0.8$ )?

```
> LD(popdat[, 5:14])$"R^2"
```

	snp1	snp2	snp3	snp4	snp5	snp6	snp7	snp8	snp9	snp10
snp1	NA	0.016	0.235	0.206	0.258	0.227	0.152	0.117	0.090	0.000
snp2	NA	NA	0.004	0.004	0.005	0.004	0.000	0.000	0.000	0.000
snp3	NA	NA	NA	0.602	0.457	0.346	0.641	0.031	0.042	0.001
snp4	NA	NA	NA	NA	0.803	0.650	0.729	0.027	0.037	0.002
snp5	NA	NA	NA	NA	NA	0.874	0.586	0.034	0.046	0.002
snp6	NA	NA	NA	NA	NA	NA	0.670	0.030	0.040	0.002
snp7	NA	NA	NA	NA	NA	NA	NA	0.020	0.027	0.003
snp8	NA	NA	NA	NA	NA	NA	NA	NA	0.002	0.000
snp9	NA	NA	NA	NA	NA	NA	NA	NA	NA	0.001
snp10	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA

SNP pairs 4-5 and 5-6 have  $r^2 \geq 0.8$ .

## B.11 Exercise 16:

For SNPs in strong LD, what is  $r^2$  for separate samples of cases and controls?

For controls,

```
> LD(data.frame(snp4, snp5, snp6)[aff == 0, ])$"R^2"
```

	snp4	snp5	snp6
snp4	NA	0.806591	0.6419715
snp5	NA	NA	0.8661005
snp6	NA	NA	NA

For cases,

```
> LD(data.frame(snp4, snp5, snp6)[aff == 1, ])$"R^2"
```

	snp4	snp5	snp6
snp4	NA	0.7951475	0.6773275
snp5	NA	NA	0.9083237
snp6	NA	NA	NA

Note that the fact that LD is higher in cases may mean nothing because the estimates of LD are biased upwards with smaller sample sizes. For example in a small sample (5 people) of controls we expect even higher LD because of strong upward bias:

```
> LD(popdat[which(aff == 0)[1:5], 8:10])$"R^2"
```

	snp4	snp5	snp6
snp4	NA	0.9995876	0.9995876
snp5	NA	NA	0.9995876
snp6	NA	NA	NA

More elaborate methods, such as that by Zaykin, are required to contrast LD between sample of unequal size.

### B.12 Exercise 17:

Is there significant association between affection status and sex? What is  $P$ -value for association?

```
> glm(aff ~ sex, family = binomial())  
  
Call: glm(formula = aff ~ sex, family = binomial())  
  
Coefficients:  
(Intercept)          sex  
-1.3197        0.1006  
  
Degrees of Freedom: 2499 Total (i.e. Null); 2498 Residual  
Null Deviance:      2632  
Residual Deviance: 2631           AIC: 2635
```

There is significant ( $P = 0.03$ ) association.

### B.13 Exercise 18:

Is association between the disease and qt significant?

```
> glm(aff ~ qt, family = binomial())  
  
Call: glm(formula = aff ~ qt, family = binomial())  
  
Coefficients:  
(Intercept)          qt  
-1.26769       -0.02514  
  
Degrees of Freedom: 2499 Total (i.e. Null); 2498 Residual  
Null Deviance:      2632  
Residual Deviance: 2632           AIC: 2636
```

There is no significant ( $P = 0.6$ ) association.

### B.14 Exercise 19:

Which SNPs are associated with the quantitative trait qt at nominal  $P \leq 0.05$ ? Use 2 d.f. test.

```
> summary(lm(qt ~ snp1))  
  
Call:  
lm(formula = qt ~ snp1)  
  
Residuals:  
    Min     1Q   Median     3Q    Max  
-3.52609 -0.66427 -0.01110  0.67648  3.54622  
  
Coefficients:  
            Estimate Std. Error t value Pr(>|t|)
```

```

(Intercept) -0.02846    0.02758   -1.032    0.3022
snp1A/B      0.08200    0.04316   1.900    0.0575 .
snp1B/B      0.18644    0.07536   2.474    0.0134 *
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Residual standard error: 0.9893 on 2371 degrees of freedom
(126 observations deleted due to missingness)
Multiple R-squared: 0.00335,          Adjusted R-squared: 0.002509
F-statistic: 3.985 on 2 and 2371 DF,  p-value: 0.01873

...
> summary(lm(qt ~ snp10))

Call:
lm(formula = qt ~ snp10)

Residuals:
    Min      1Q      Median      3Q      Max 
-3.586464 -0.677484  0.001935  0.673270  3.412527 

Coefficients:
            Estimate Std. Error t value Pr(>|t|)    
(Intercept) 0.01915   0.02344   0.817   0.414    
snp10A/B   0.01277   0.04829   0.264   0.792    
snp10B/B   0.17178   0.15860   1.083   0.279    

Residual standard error: 0.9921 on 2381 degrees of freedom
(116 observations deleted due to missingness)
Multiple R-squared: 0.0005072,          Adjusted R-squared: -0.0003324
F-statistic: 0.6041 on 2 and 2381 DF,  p-value: 0.5467

```

SNPs 1, 4, 5 an 9 are significantly associated at nominal  $P \leq 0.05$ .

### B.15 Exercise 20:

Test each SNP for association with the affection status, using 2 d.f. test. Which SNPs are significantly associated at nominal  $P \leq 0.05$ ? How can you describe the model of action of the significant SNPs?

```

> x <- glm(aff ~ snp5, family = binomial())
> x

Call: glm(formula = aff ~ snp5, family = binomial())

Coefficients:
(Intercept)     snp5A/A       snp5B/B      
-1.4868        0.2112       0.3387      

Degrees of Freedom: 2382 Total (i.e. Null); 2380 Residual
(117 observations deleted due to missingness)

```

```

Null Deviance: 2440
Residual Deviance: 2431 AIC: 2437

> anova(x, test = "Chisq")

Analysis of Deviance Table

Model: binomial, link: logit

Response: aff

Terms added sequentially (first to last)

          Df Deviance Resid. Df Resid. Dev P(>|Chi|)
NULL           2382    2440.40
snp5      2     9.24      2380    2431.16      0.01

...
> x <- glm(aff ~ snp10, family = binomial())
> x

Call: glm(formula = aff ~ snp10, family = binomial())

Coefficients:
(Intercept)    snp10A/B    snp10B/B
-1.3703        0.2909     -0.1803

Degrees of Freedom: 2383 Total (i.e. Null); 2381 Residual
(116 observations deleted due to missingness)
Null Deviance: 2475
Residual Deviance: 2468 AIC: 2474

> anova(x, test = "Chisq")

Analysis of Deviance Table

Model: binomial, link: logit

Response: aff

Terms added sequentially (first to last)

          Df Deviance Resid. Df Resid. Dev P(>|Chi|)
NULL           2383    2475.13
snp10      2     6.73      2381    2468.39      0.03

```

The SNPs 5 and 10 are significantly associated at  $P \leq 0.05$ . The model of action of SNP5 can be described as recessive (while the risk for AA and AB is not significantly different, there is 1.4 times increased risk for these homozygous for

BB). The SNP 10 demonstrates somewhat weird action with the risk increased in heterozygous AB individuals. However, the confidence interval for BB is large and therefore we can not claim that BB is not increasing the risk.

### B.16 Exercise 21:

For the SNPs selected in previous question, test association using additive model. Which SNPs are still associated?

```
> glm(aff ~ as.numeric(snp5), family = binomial())
Call: glm(formula = aff ~ as.numeric(snp5), family = binomial())

Coefficients:
(Intercept) as.numeric(snp5)
-1.5695      0.1094

Degrees of Freedom: 2382 Total (i.e. Null); 2381 Residual
(117 observations deleted due to missingness)
Null Deviance: 2440
Residual Deviance: 2438      AIC: 2442

> glm(aff ~ as.numeric(snp10), family = binomial())
Call: glm(formula = aff ~ as.numeric(snp10), family = binomial())

Coefficients:
(Intercept) as.numeric(snp10)
-1.5539      0.1976

Degrees of Freedom: 2383 Total (i.e. Null); 2382 Residual
(116 observations deleted due to missingness)
Null Deviance: 2475
Residual Deviance: 2471      AIC: 2475
```

Only SNP 10 is significantly associated under the additive model.

### B.17 Exercise 22:

If you adjust the analysis under additive model (question 21) for significant covariates which you discovered in questions 17 and 18, are these findings still significant?

```
> glm(aff ~ sex +.snp10, family = binomial())
Call: glm(formula = aff ~ sex + .snp10, family = binomial())

Coefficients:
(Intercept)       sex     .snp10A/B     .snp10B/B
-1.41894      0.09513     0.29230     -0.18471

Degrees of Freedom: 2383 Total (i.e. Null); 2380 Residual
```

```

(116 observations deleted due to missingness)
Null Deviance: 2475
Residual Deviance: 2467 AIC: 2475

```

Yes, SNP 10 becomes even a bit more significantly associated after adjusting for sex.

### B.18 Exercise 23:

Test association between `aff` and `snp5` and `snp10`, allowing for the SNPs interaction effect. Use arbitrary (not an additive) model. Do you observe significant interaction? How can you describe the model of concert action of `snp5` and `snp10`?

```

> glm(aff ~ snp5 * snp10, family = binomial())
Call: glm(formula = aff ~ snp5 * snp10, family = binomial())

Coefficients:
(Intercept)      snp5A/A      snp5B/B      snp10A/B
-1.50840       -0.41802       0.33441      -0.01403
snp10B/B  snp5A/A:snp10A/B  snp5B/B:snp10A/B  snp5A/A:snp10B/B
-0.14983        1.48369       0.12989       0.82348
snp5B/B:snp10B/B
-0.28562

Degrees of Freedom: 2268 Total (i.e. Null); 2260 Residual
(231 observations deleted due to missingness)
Null Deviance: 2282
Residual Deviance: 2243 AIC: 2261

```

It appears that SNP10 genotype is only relevant in those who are homozygous for the low-risk A allele at the SNP5; in such cases SNP 10 allele B is risk increasing. In these homozygous for SNP 5 A, we observe highly significant increase in risk for heterozygotes for SNP10 and increased (though not significantly) risk for SNP 10 BB.

100

### B.19 Exercise 24:

```

> data(srdta)

Number of people:
> srdta@gtdata@nids
[1] 2500

Number of males:
> sum(srdta@gtdata@male)
[1] 1275

```

Number of females:

```
> srdta@gtdata@nids - sum(srdta@gtdata@male)  
[1] 1225
```

... or you could get the same answer like this<sup>6</sup>:

```
> sum(srdta@gtdata@male == 0)  
[1] 1225
```

The proportion of males can be computed using above results

```
> sum(srdta@gtdata@male)/srdfa@gtdata@nids  
[1] 0.51
```

or by using `mean()` function:

```
> mean(srdta@gtdata@male)  
[1] 0.51
```

## B.20 Exercise 25:

The names of markers located after 2,490,000 b.p. are

```
> vec <- (srdfa@gtdata@map > 2490000)  
> srdfa@gtdata@snpnames[vec]  
  
[1] "rs9273" "rs9277" "rs9279" "rs9283"
```

Between 1,100,000 and 1,105,000 b.p.:

```
> vec <- (srdfa@gtdata@map > 1100000 & srdfa@gtdata@map < 1105000)  
> srdfa@gtdata@snpnames[vec]  
  
[1] "rs4180" "rs4186" "rs4187"
```

## B.21 Exercise 26:

To learn what allele of "rs114" is the reference you need to run

```
> as.character(srdfa@gtdata@coding["rs114"])  
  
<NA>  
"AT"
```

Here, the first ("A") allele is the reference and thus the second ("T") is the effective one. Remember that when using `as.numeric` function to convert the genotypes to human-readable and R-operable format, the homozygotes for reference will be coded as "0", heterozygotes as "1" and the non-reference ("effective") homozygotes will be coded as "2":

---

<sup>6</sup>This is something covered later in the section ?? ("??")

```

> table(as.character(srdta@gtdata[, "rs114"]), as.numeric(srdta@gtdata[,
+      "rs114"]))

      0     1     2
A/A 1868    0    0
A/T      0 491    0
T/T      0    0   34

```

To compute frequency of the effective allele of SNP "rs114" in total sample, you can go two ways. First, we can try to take a sum of all rs114 genotypes and divide it by twice the number of people:

```

> a <- as.numeric(srdta@gtdata[, "rs114"])
> sum(a)

[1] NA

```

This, however, returns NA, because some of the genotypes are missing. We can deal with this problem by running `sum()` with the option `na.rm=TRUE`:

```

> sum(a, na.rm = T)

[1] 559

```

However, now we do not know what was the number of people for whom the genotype was measured!

An easier way would be to compute mean value of rs114 with the `mean( ... ,na.rm=TRUE)` function and divide it by 2:

```

> mean(a, na.rm = T)/2

[1] 0.116799

```

To compute frequency of the effective allele of "rs114" in males, you can use

```

> amale <- as.numeric(srdta@gtdata[srdta@phdata$sex == 1, "rs114"])
> mean(amale, na.rm = T)/2

[1] 0.1164216

```

To compute frequency of the effective allele in females, you can use

```

> afemale <- as.numeric(srdta@gtdata[srdta@phdata$sex == 0, "rs114"])
> mean(afemale, na.rm = T)/2

[1] 0.1171942

```

Actually, the problem that we do not know how many people are measured, can be easily dealt with. This can be done by using `is.na(A)` function which returns true when some element of A is not measured. Thus, the number of people with measured genotype for "rs114" is

```

> a <- as.numeric(srdta@gtdata[, "rs114"])
> sum(!is.na(a))

```

```
[1] 2393
```

And the allele frequency estimate is

```
> sum(a, na.rm = T)/(2 * sum(!is.na(a)))
```

```
[1] 0.116799
```

exactly the same as above.

The frequencies of the reference allele are computed very simply as one minus the frequency of the effective allele.

## B.22 Exercise 27:

To test for HWE in first 10 SNPs in total sample

```
> summary(srdta@gtdta[, 1:10])
```

	NoMeasured	CallRate	Q.2	P.11	P.12	P.22	Pexact	Fmax
rs10	2384	0.9536	0.13255034	1792	552	40	7.897327e-01	-0.00680004
rs18	2385	0.9540	0.28029350	1232	969	184	7.608230e-01	-0.007017332
rs29	2374	0.9496	0.13774221	1763	568	43	7.955141e-01	-0.007241148
rs65	2378	0.9512	0.71972246	182	969	1227	6.475412e-01	-0.010016746
rs73	2385	0.9540	0.01341719	2331	44	10	1.792470e-12	0.303150234
rs114	2393	0.9572	0.11679900	1868	491	34	7.663683e-01	0.005487764
rs128	2391	0.9564	0.02488499	2281	101	9	9.408599e-06	0.129600629
rs130	2379	0.9516	0.69377890	222	1013	1144	9.615127e-01	-0.002140946
rs143	2377	0.9508	0.47728229	655	1175	547	6.512540e-01	0.009313705
rs150	2369	0.9476	0.65998312	267	1077	1025	5.518478e-01	-0.012948436

	Plrt	Chromosome
rs10	7.355343e-01	1
rs18	7.315304e-01	1
rs29	7.227853e-01	1
rs65	6.246577e-01	1
rs73	1.281239e-12	1
rs114	7.894076e-01	1
rs128	1.000431e-05	1
rs130	9.168114e-01	1
rs143	6.497695e-01	1
rs150	5.281254e-01	1

To test it in cases

```
> summary(srdta@gtdta[srdta@phdata$bt == 1, 1:10])
```

	NoMeasured	CallRate	Q.2	P.11	P.12	P.22	Pexact	Fmax
rs10	1197	0.9622186	0.13700919	888	290	19	4.635677e-01	-0.024514202
rs18	1189	0.9557878	0.28511354	605	490	94	7.759191e-01	-0.010949158
rs29	1176	0.9453376	0.14285714	859	298	19	2.832575e-01	-0.034722222
rs65	1185	0.9525723	0.72700422	83	481	621	4.647357e-01	-0.022595469
rs73	1187	0.9541801	0.01053075	1167	15	5	3.988770e-08	0.393614304
rs114	1190	0.9565916	0.12184874	918	254	18	8.924018e-01	0.002606831

```

rs128      1183 0.9509646 0.02409129 1129   51    3 2.747904e-02  0.083175674
rs130      1188 0.9549839 0.68392256 117    517   554 8.407527e-01 -0.006569292
rs143      1192 0.9581994 0.48489933 320    588   284 6.848365e-01  0.012522119
rs150      1182 0.9501608 0.66624365 127    535   520 5.568363e-01 -0.017756050
          Plrt Chromosome
rs10      3.871421e-01     1
rs18      7.052930e-01     1
rs29      2.214580e-01     1
rs65      4.348023e-01     1
rs73      2.423624e-08     1
rs114     9.285104e-01     1
rs128     3.157174e-02     1
rs130     8.207476e-01     1
rs143     6.654994e-01     1
rs150     5.409408e-01     1

in controls

> summary(srdta@gtdata[srdta@phdata$bt == 0, 1:10])

      NoMeasured CallRate       Q.2 P.11 P.12 P.22      Pexact      Fmax
rs10      1177 0.9453815 0.12744265 897   260   20 7.933317e-01  0.006751055
rs18      1185 0.9518072 0.27426160 623   474   88 9.418133e-01 -0.004812165
rs29      1188 0.9542169 0.13215488 897   268   23 5.288436e-01  0.016525913
rs65      1183 0.9502008 0.71344041 98    482   603 8.871139e-01  0.003540522
rs73      1188 0.9542169 0.01641414 1154   29    5 6.941219e-06  0.244001185
rs114     1192 0.9574297 0.11157718 941   236   15 8.846527e-01  0.001356081
rs128     1197 0.9614458 0.02589808 1141   50    6 7.745807e-05  0.172107564
rs130     1181 0.9485944 0.70491109 104   489   588 8.887439e-01  0.004728114
rs143     1174 0.9429719 0.46805792 334   581   259 8.604122e-01  0.006165442
rs150     1176 0.9445783 0.65306122 139   538   499 7.968462e-01 -0.009574142
          Plrt Chromosome
rs10      8.178295e-01     1
rs18      8.683219e-01     1
rs29      5.737373e-01     1
rs65      9.031273e-01     1
rs73      5.537568e-06     1
rs114     9.627084e-01     1
rs128     7.552399e-05     1
rs130     8.710047e-01     1
rs143     8.326938e-01     1
rs150     7.424986e-01     1

```

### B.23 Exercise 53:

Perform meta-analysis of the data presented in table 2. Which allele is the risk one? Is this risk significant? What is pooled Odds Ratio and 95% confidence interval? Do analysis using at least two methods. Which method is better (best) in this situation? Why?

We first need to unify Odds Ratios by using the same effective allele. Let that be the "risk" allele, as may be guessed from a glance to the data, namely "Pro".

When the effects are reported for the other, "Ala" allele, the corresponding ORs for the "Pro" allele can be found using simple relation  $OR_{Pro} = 1/OR_{Ala}$ .

Thus, the vector of Odds Ratios for "Pro" allele is

```
> or.pro <- c(1/0.67, 0.93, 1.08, 1/0.83, 1.22, 1.23)
> or.pro
[1] 1.492537 0.930000 1.080000 1.204819 1.220000 1.230000
```

The corresponding  $P - values$  are

```
> p <- c(0.013, 0.6, 0.84, 0.4, 0.25, 0.07)
```

Let us find log-ORs

```
> logor.pro <- log(or.pro)
> logor.pro
[1] 0.40047757 -0.07257069 0.07696104 0.18632958 0.19885086 0.20701417
```

Corresponding squared standard errors are

```
> s2 <- logor.pro * logor.pro/qchisq(1 - p, 1)
> s2
[1] 0.02599764 0.01915121 0.14531060 0.04901514 0.02988102 0.01305349
```

and weights are

```
> w <- 1/s2
> w
[1] 38.46503 52.21601 6.88181 20.40186 33.46606 76.60788
```

Thus the pooled estimate of log-OR is

```
> p.logor.pro <- sum(w * logor.pro)/sum(w)
> p.logor.pro
[1] 0.1686548
```

and the standard error is

```
> p.s <- 1/sqrt(sum(w))
> p.s
[1] 0.066221
```

Thus the pooled estimate of Odds Ratio from association between type 2 diabetes and "Ala" allele is

```
> exp(p.logor.pro)
[1] 1.183711
```

and the 95% confidence interval is

```

> exp(p.logor.pro - 1.96 * p.s)
[1] 1.039627
> exp(p.logor.pro + 1.96 * p.s)
[1] 1.347765

```

The  $\chi^2$  test for association and corresponding  $P - value$  are

```

> p.chi2 <- (p.logor.pro/p.s)^2
> p.chi2
[1] 6.486429

> p.pval <- 1 - pchisq(p.chi2, 1)
> p.pval
[1] 0.01087011

```

Z-score pooling though may be more appropriate method for such differentially designed studies (e.g. control groups are very different). To get Z-score pooling working, we need first find Z-scores from P-values

```

> p <- c(0.013, 0.6, 0.84, 0.4, 0.25, 0.07)
> z <- sqrt(qchisq(1 - p, 1))
> z
[1] 2.4837693 0.5244005 0.2018935 0.8416212 1.1503494 1.8119107

```

and assign the right sign (let "+" is for the risk effect of "Pro").

```

> effsig <- c(1, -1, 1, 1, 1, 1)
> z <- z * effsig
> z
[1] 2.4837693 -0.5244005 0.2018935 0.8416212 1.1503494 1.8119107

```

Now, we need to assign weights to the studies as

```

> n <- c(221, 306, 71, 164, 242, 471)
> w <- sqrt(n)

```

and the pooled estimate of Z and corresponding  $P - value$  are

```

> zpoo <- sum(w * z)/sqrt(sum(w^2))
> zpoo
[1] 2.537333

> 1 - pchisq(zpoo * zpoo, 1)
[1] 0.01117008

```

As you can see the results are almost identical to the previous obtained with inverse variance pooling.

## B.24 Exercise 54:

Perform meta-analys excluding the original report (study 1). Is there still significant association between Pro12Ala and diabetes?

The answer to this exercise can be obtained in exactly the same manner, as for the previous one, limiting our consideration to the last five studies.

Thus, the vector of Odds Ratios for "Pro" allele is

```
> or.pro <- c(0.93, 1.08, 1/0.83, 1.22, 1.23)
> or.pro
[1] 0.930000 1.080000 1.204819 1.220000 1.230000
```

The corresponding  $P$ -values are

```
> p <- c(0.6, 0.84, 0.4, 0.25, 0.07)
```

Let us find log-ORs

```
> logor.pro <- log(or.pro)
> logor.pro
[1] -0.07257069 0.07696104 0.18632958 0.19885086 0.20701417
```

Corresponding squared standard errors are

```
> s2 <- logor.pro * logor.pro/qchisq(1 - p, 1)
> s2
[1] 0.01915121 0.14531060 0.04901514 0.02988102 0.01305349
```

and weights are

```
> w <- 1/s2
> w
[1] 52.21601 6.88181 20.40186 33.46606 76.60788
```

Thus the pooled estimate of log-OR is

```
> p.logor.pro <- sum(w * logor.pro)/sum(w)
> p.logor.pro
[1] 0.1216172
```

and the standard error is

```
> p.s <- 1/sqrt(sum(w))
> p.s
[1] 0.07262917
```

Thus the pooled estimate of Odds Ratio from association between type 2 diabetes and "Ala" allele is

```
> exp(p.logor.pro)
```

```
[1] 1.129322
```

and the 95% confidence interval is

```
> exp(p.logor.pro - 1.96 * p.s)
[1] 0.9794776
> exp(p.logor.pro + 1.96 * p.s)
[1] 1.302090
```

The  $\chi^2$  test for association and corresponding  $P-value$  are

```
> p.chi2 <- (p.logor.pro/p.s)^2
> p.chi2
[1] 2.803937
> p.pval <- 1 - pchisq(p.chi2, 1)
> p.pval
[1] 0.09403318
```

Z-score pooling though may be more appropriate method for such differentially designed studies (e.g. control groups are very different). To get Z-score pooling working, we need first find Z-scores from P-values

```
> p <- c(0.6, 0.84, 0.4, 0.25, 0.07)
> z <- sqrt(qchisq(1 - p, 1))
> z
[1] 0.5244005 0.2018935 0.8416212 1.1503494 1.8119107
```

and assign the right sign (let "+" is for the risk effect of "Pro").

```
> effsig <- c(-1, 1, 1, 1, 1)
> z <- z * effsig
> z
[1] -0.5244005 0.2018935 0.8416212 1.1503494 1.8119107
```

Now, we need to assign weights to the studies as

```
> n <- c(306, 71, 164, 242, 471)
> w <- sqrt(n)
```

ane the pooled estimate of Z and corresponding  $P-value$  are

```
> zpoo <- sum(w * z)/sqrt(sum(w^2))
> zpoo
[1] 1.709151
> 1 - pchisq(zpoo * zpoo, 1)
[1] 0.08742297
```

As you can see the results are almost identical to the previous obtained with inverse variance pooling.

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