# Livestock Breeding and Genomics - Exercise 3

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## Problem 1: Breeding Value

We are considering a quantitative trait that depends on a given bi-allelic locus G. The frequency of the favorable allele corresponds to 0.08. Suppose that genotype frequencies follow the Hardy-Weinberg equilibrium. The difference between the homozygous genotypes corresponds to 15. The heterozygous genotype has a value of -1.5.

- a) Compute the breeding values and the dominance deviations for the three genotypes.
- b) Because of selecting the positive allele the frequency has increased to 0.096. How does this increased allele frequency change the breeding values?

**Hint**: Have a look at the summary table of all values in the course notes.

### Problem 2: Allele Substitution

What is the meaning of the term **allele substituion** an how big is it in 1a) and 1b)?

## Problem 3: Reading Data into R

You can download a file in csv-format from the course website. The URL is https://charlotte-ngs.github.io/LBGFS2018/ex/w04/iris.csv. Read the data from that csv-file into R using the function read.csv2(). Test the consequences of specifying the option stringsAsFactors=FALSE.

#### Hints:

- 1. You can first download the csv-file to your local computer and then read the data, or you can directly indicate the URL when reading the data. You get more information with the command ?read.csv2 at the R-console.
- 2. Assign the result of read.csv2() to a variable
- 3. Use the function str() on the result of read.csv2() to see the difference between the two results of reading the data.
- 4. Use the description at https://bookdown.org/rdpeng/rprogdatascience/getting-data-in-and-out-of-r. html as a reference to read data into R. There is also a video on the same subject under https://youtu.be/Z\_dc\_FADyi4.

# Additional Problem: Create a plot in R

Plot the values in the columns Sepal.Length and Petal.Length of the Iris data set. The plot should look like the following figure.