# Livestock Breeding and Genomics - Solution 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/LBGFS2019/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 <code>?read.csv2</code> 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.