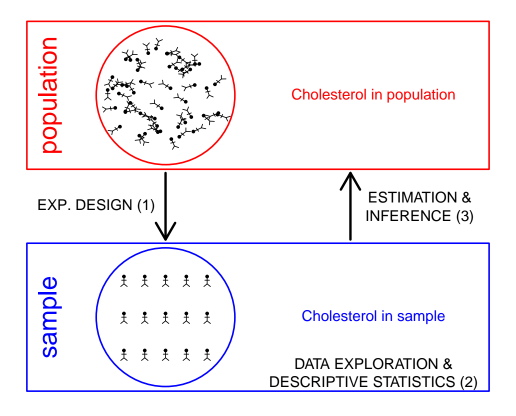
# 4. Data Exploration

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# 1 Why data exploration and descriptive statistics?

- 1. Reporting on results: not useful to only report a table with the results for each subject.
- 2. Letting the data speak by summarizing and visualizing it.
- 3. Getting insight in the data.
- 4. Discover errors, anomalies or even fraud.
- 5. Check assumptions that are required for the downstream statistical inference, e.g. are the data Normally distributed.

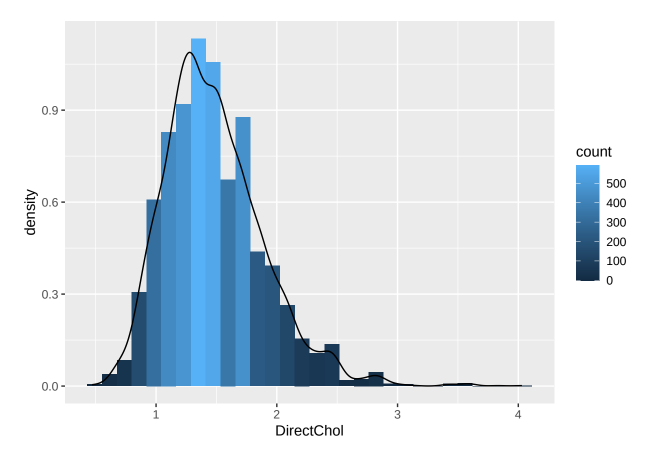
# 2 Univariate exploration of quantitative variables

### 2.1 Histogram

A histogram is very useful to provide an estimate of the distribution of the data without making distributional assumptions.

For a histogram you typically require to have enough observations. A sample size of 30 observations is a bare minimum to construct a histogram.

```
NHANES %>%
  filter(Gender == "female") %>%
  ggplot(aes(x = DirectChol)) +
  geom_histogram(aes(y = ..density.., fill = ..count..), bins = 30) +
  geom_density(aes(y = ..density..))
```



1. Select females and pipe results to ggplot.

NHANES %>% filter(Gender=="female")

2. Select data to plot.

ggplot(aes(x=DirectChol)) +

- 3. Equal bins for interpretation, the number of bins can be selected with the bins argument to the geom\_hist.
- 4. Relative frequenties to enable visual comparison between histograms.

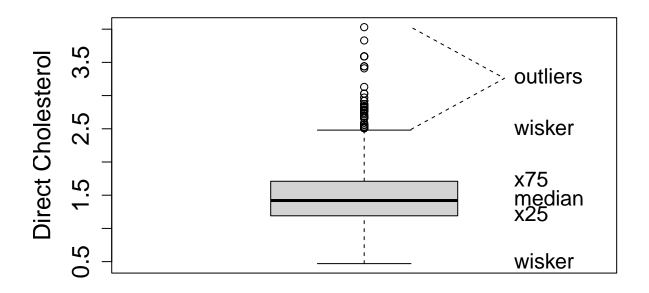
geom\_histogram(aes(y=..density.., fill=..count..)) +

5. If we have enough observations we can use a kernel density estimator of f(x).

geom\_density(aes(y=..density..))

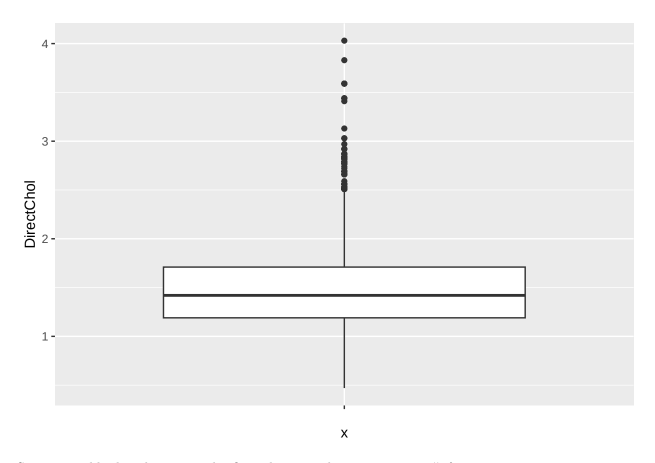
## 2.2 Boxplot

• A quantile,  $x_{a\%}$ , is the value of the random variable that correspond to a certain probability  $F(x_{a\%}) = P[X \le x_{a\%} = a\%]$ .



With ggplot we always have to define an x variable if we make a boxplot. If we use a string then all data is considered to originate from one category and one boxplot is constructed.

```
NHANES %>%
filter(Gender == "female") %>%
ggplot(aes(x = "", y = DirectChol)) +
geom_boxplot()
```



So we can add a boxplot to a ggplot figure by using the geom\_boxplot() function.

If the dataset is small to moderate in size we can also add the raw data to the plot with the <code>geom\_point()</code> function and the <code>position="jitter"</code> argument. Note, that we then also set the <code>outlier.shape</code> argument in the <code>geom\_boxplot</code> function on NA so that the outliers are not plotted twice.

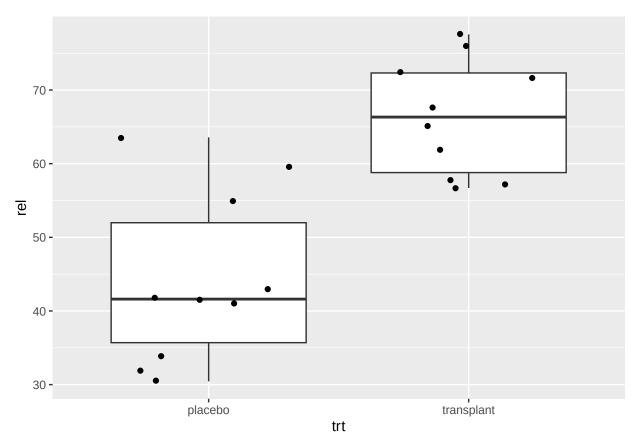
Here, we will plot again the relative abundances of **Staphylococcus** from the armpit transplant experiment.

```
ap <- read_csv("https://raw.githubusercontent.com/GTPB/PSLS20/master/data/armpit.csv")
ap</pre>
```

```
# A tibble: 20 \times 2
   trt
                 rel
   <chr>
               <dbl>
 1 placebo
               55.0
 2 placebo
               31.8
 3 placebo
                41.1
               59.5
 4 placebo
5 placebo
               63.6
6 placebo
               41.5
7 placebo
               30.4
8 placebo
               43.0
9 placebo
               41.7
10 placebo
                33.9
11 transplant
               57.2
12 transplant
               72.5
13 transplant
               61.9
14 transplant
               56.7
```

```
15 transplant 76
16 transplant 71.7
17 transplant 57.8
18 transplant 65.1
19 transplant 67.5
20 transplant 77.6

ap %>%
    ggplot(aes(x = trt, y = rel)) +
    geom_boxplot(outlier.shape = NA) +
    geom_point(position = "jitter")
```



When we specify a factor variable for x, we get a boxplot for each treatment group.

## 2.3 Descriptive statistics

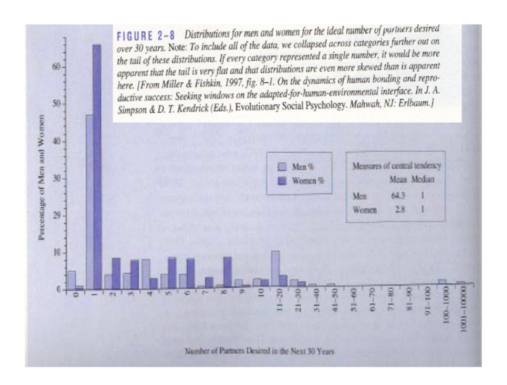
#### 2.3.1 Central location: Mean or Median?

## 2.3.1.1 Mean

• In a period of 30 years, males hope to have on average 64.3 partners and females 2.8 (Miller and Fishkin, 1997)

#### 2.3.1.2 Median

• The median of the number of partners males and females want to have is both 1 (Miller and Fishkin, 1997)



## 2.3.1.3 What happens?

• Mean is very sensitive towards outliers!

Figuur 2: Partners

#### 2.3.2 Geometric mean

$$\sqrt[n]{\prod_{i=1}^n x_i} = \exp\left\{\frac{1}{n}\sum_{i=1}^n \log(x_i)\right\}$$

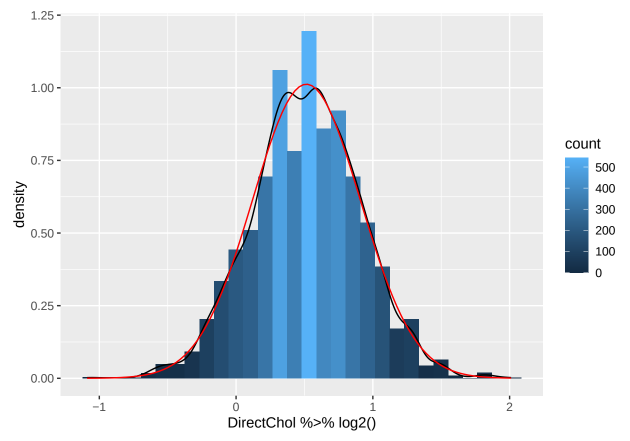
- Geometric mean is closer to the median then the mean
- log-transformation removes skewness
- Often a more useful measure for the central location than median:
- 1. Uses all observations: is more precise
- 2. It is the ordinary mean on log-transformed data  $\rightarrow$  classical statistical methods can be directly applied, e.g. hypothesis tests and confidence intervals (see chapter 5)
- 3. Useful for many biological characteristics e.g. concentrations that cannot be negative.
- 4. Differences on a log scale have the interpretation of a log fold change:

$$\log(B) - \log(A) = \log(\frac{B}{A}) = \log(FC_{\rm B~vs~A})$$

In Genomics often the  $log_2$  transformation is used. A difference of 1 corresponds to a FC = 2.

```
logSummary <-
NHANES %>%
filter(Gender == "female") %>%
summarize(logMean = mean(DirectChol %>% log2(), na.rm = TRUE), sd = sd(DirectChol %>% log2(), na.rm =
mutate(geoMean = 2^logMean)

NHANES %>%
filter(Gender == "female") %>%
ggplot(aes(x = DirectChol %>% log2())) +
geom_histogram(aes(y = ..density.., fill = ..count..), bins = 30) +
geom_density(aes(y = ..density..)) +
stat_function(fun = dnorm, color = "red", args = list(mean = logSummary$logMean, sd = logSummary$sd))
```



#### logSummary

- Indeed the mean is pulled to larger values by the skewed data.
- The geometric mean is closer to the median.
- The cholesterol data are much more symmetric upon log transformation and the approximation by a Normal distribution is good.

## 2.4 Descriptive Statistics for Variability

The variability around the central value is crucial:

- 1. Biologists are often interested in how animals or plants are spread in the study region.
- 2. Compare groups: the group effect is more clear when the response has less variability. Quantifying variability is crucial to distinguish between systematic and random patterns.
- The response varies between and within individuals and is the reason why we need statistics.
- Crucial to describe both the central location and the variability.
- Which part of the variability can we explain (e.g. with characteristics treatment, age, etc.) and which part is unexplained?

#### 2.4.1 Sample variance and sample standard deviation

• Sample variance:

$$s_X^2 = \sum_{i=1}^n \frac{(X - \bar{X})^2}{n - 1}$$

- Interpretation is often difficult because it is in another unit than the measurements.
- Standard deviation:

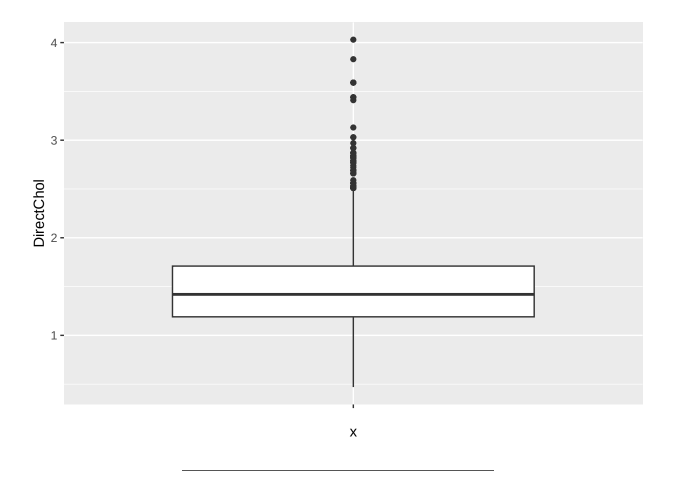
$$s_X = \sqrt{s_x^2}$$

- Very useful for Normal distributed observations:
  - 68% of the observations falls in the interval  $\bar{x} s_x$  en  $\bar{x} + s_x$
  - -95% of the observations falls in the interval  $\bar{x}-2s_x$  en  $\bar{x}+2s_x$ .
- These intervals are referred to as 68% en 95% reference-intervals.
- If the data are not Normally distributed, reference intervals are not valid.

#### 2.4.2 Interquartile range

For skewed data the standard deviation is not useful

- It is very sensitive to outliers
- Inter Quartile Range: Distance between first and third quartile
- Width of the boxplot!



# 3 Normal approximation

- $\bullet\,$  Biological and chemical data are often Normally distributed upon transformation.
- If this is the case we can get a lot of insight in the data using just two descriptive statistics: mean  $\mu$  and standard deviation  $\sigma$ .

### 3.1 Evaluation with QQ-plots

If your analysis builds upon the assumption that the data are Normally distributed, it has to be verified.

We use QQ-plots or quantile-quantile plots.

- Observed quantiles from the observations in the sample are plotted against quantiles from the Normal distribution.
- If the data are Normally distributed both quantiles have to be in line.
- Dots in the plot are expected on a straight line.
- Systematic deviations of the straight line indicate that the data are not Normally distributed.
- Note, that we will always observe some random deviations from the straight line in the plot because of random biological variability, which is not indicative for deviations from Normality.
- So it is important to train yourself to learn to distinguish systematic from random deviations.

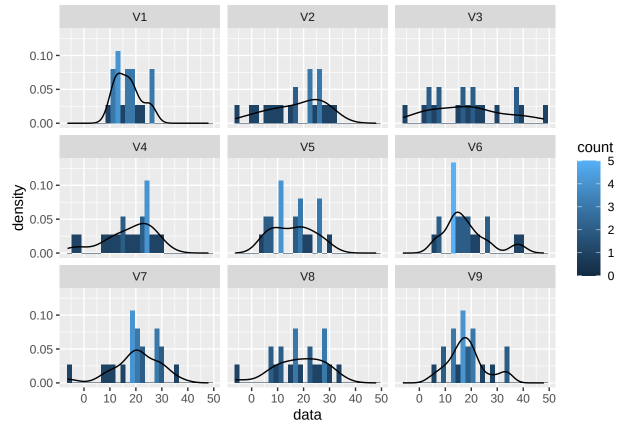
#### 3.1.1 Normal data

- We will first simulate data from the Normal distribution to show how the plots look like for data that is meeting the assumptions.
- We will simulate data from 9 samples with a mean of 18 and standard deviation of 9.

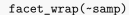
```
n <- 20
mu <- 18
sigma <- 9
nSamp <- 9

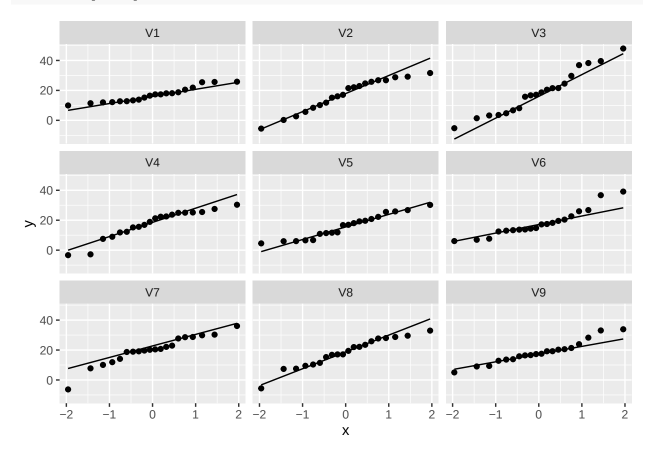
normSim <- matrix(rnorm(n * nSamp, mean = mu, sd = sigma), nrow = n) %>% as.data.frame()

normSim %>%
    gather(samp, data) %>%
    ggplot(aes(x = data)) +
    geom_histogram(aes(y = ..density.., fill = ..count..), bins = 30) +
    geom_density(aes(y = ..density..)) +
    facet_wrap(~samp)
```



```
normSim %>%
  gather(samp, data) %>%
  ggplot(aes(sample = data)) +
  geom_qq() +
  geom_qq_line() +
```



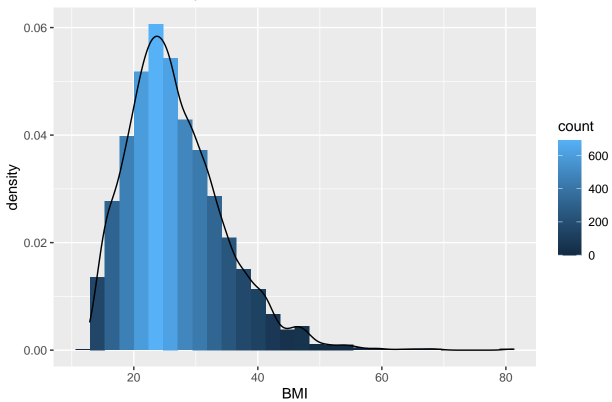


So even for Normal data we observe some deviations due to sampling variability!

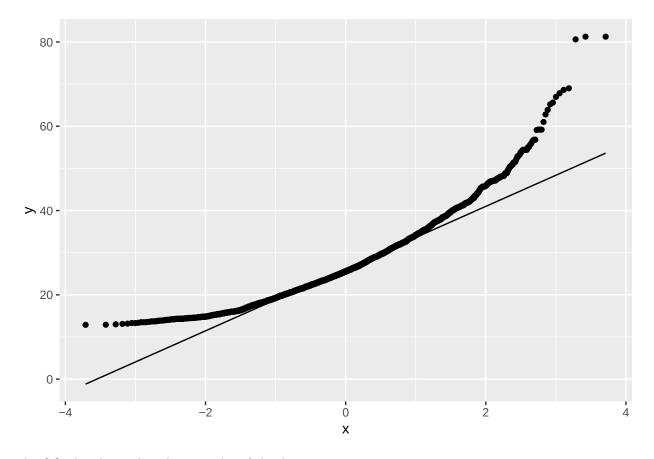
#### 3.1.2 Real data

```
NHANES %>%
  filter(Gender == "female" & !is.na(BMI)) %>%
  ggplot(aes(x = BMI)) +
  geom_histogram(aes(y = ..density.., fill = ..count..)) +
  xlab("BMI") +
  ggtitle("All females in study") +
  geom_density(aes(y = ..density..))
```





```
NHANES %>%
  filter(Gender == "female" & !is.na(BMI)) %>%
  ggplot(aes(sample = BMI)) +
  geom_qq() +
  geom_qq_line()
```



The QQ-plot shows that the quantiles of the data

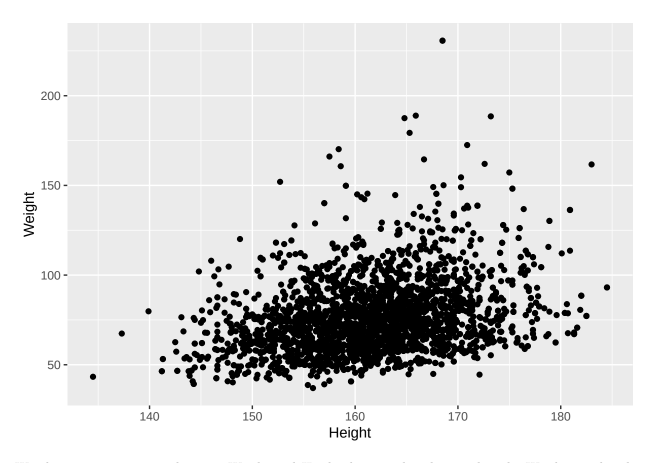
- are larger (above the line) than these from the Normal in the left tail: compression of the lower tail,
- are larger (above the line) than these from the Normal in the right tail: long tail to the right.

We can clearly see that the data are right-skewed.

# 4 Two continuous variables: Correlation

- NHANES study
- $\bullet\,$  Height and Weight Example for females

```
NHANES %>%
filter(Age > 25 & Gender == "female") %>%
ggplot(aes(x = Height, y = Weight)) +
geom_point()
```

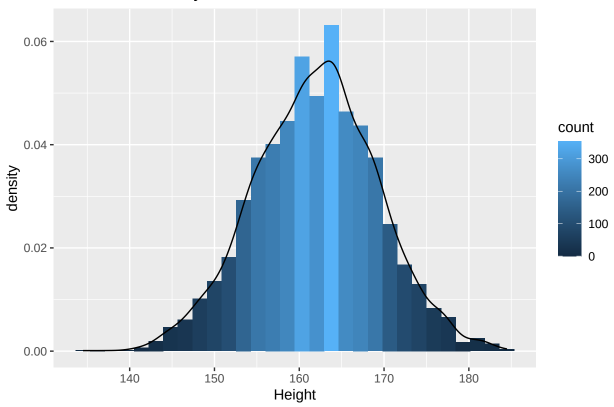


We observe an association between Weight and Height, but we also observe that the Weights tend to be right-skewed.

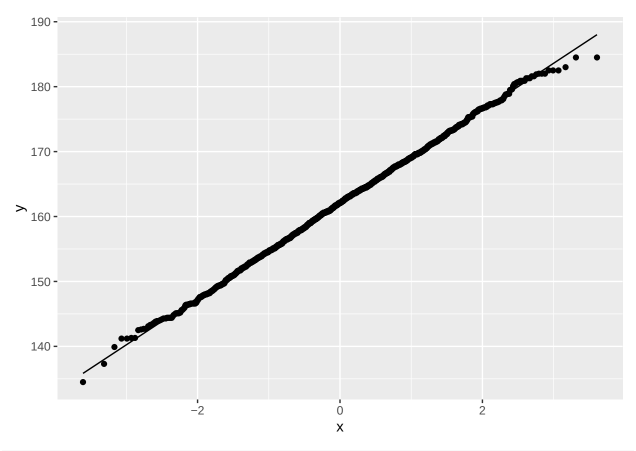
Lets look at the univariate distributions first.

```
NHANES %>%
  filter(Age > 25 & Gender == "female") %>%
  ggplot(aes(x = Height)) +
  geom_histogram(aes(y = ..density.., fill = ..count..)) +
  xlab("Height") +
  ggtitle("All females in study") +
  geom_density(aes(y = ..density..))
```

# All females in study

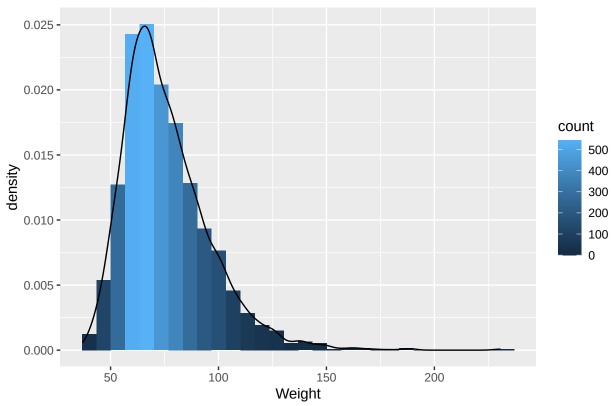


```
NHANES %>%
  filter(Age > 25 & Gender == "female") %>%
  ggplot(aes(sample = Height)) +
  geom_qq() +
  geom_qq_line()
```

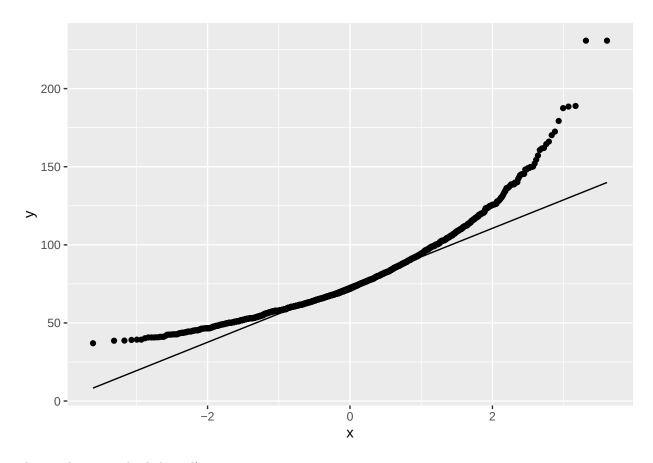


```
NHANES %>%
  filter(Age > 25 & Gender == "female") %>%
  ggplot(aes(x = Weight)) +
  geom_histogram(aes(y = ..density.., fill = ..count..)) +
  xlab("Weight") +
  ggtitle("All females in study") +
  geom_density(aes(y = ..density..))
```





```
NHANES %>%
filter(Age > 25 & Gender == "female") %>%
ggplot(aes(sample = Weight)) +
geom_qq() +
geom_qq_line()
```

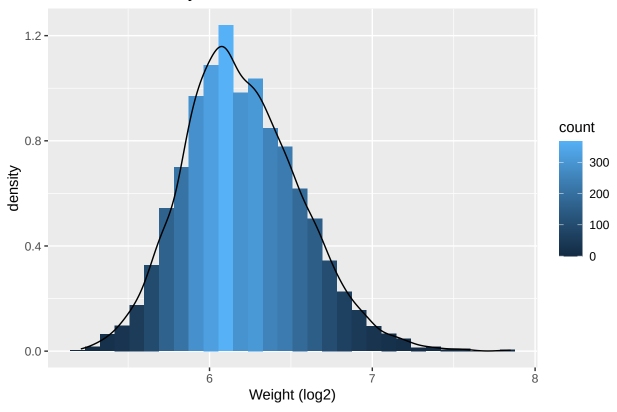


The weights are indeed skewed!

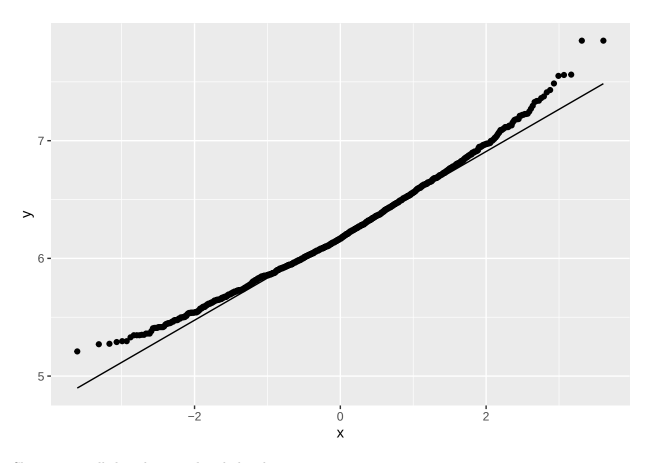
Upon log transformaton the Weights are less skewed, but still not Normally distributed.

```
NHANES %>%
  filter(Age > 25 & Gender == "female") %>%
  ggplot(aes(x = Weight %>% log2())) +
  geom_histogram(aes(y = ..density.., fill = ..count..)) +
  xlab("Weight (log2)") +
  ggtitle("All females in study") +
  geom_density(aes(y = ..density..))
```

# All females in study

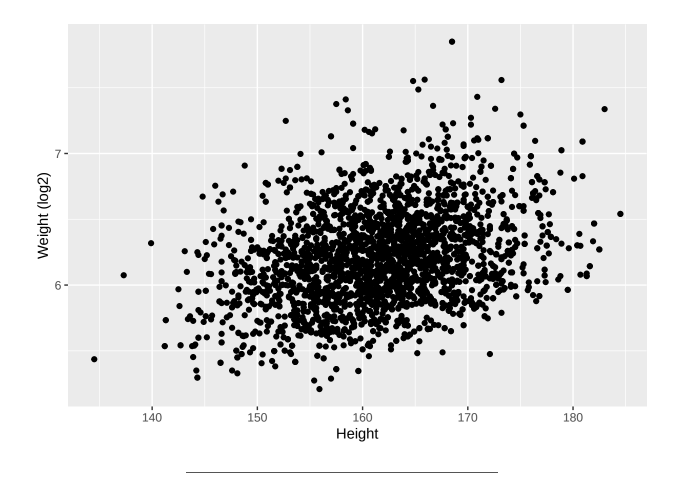


```
NHANES %>%
  filter(Age > 25 & Gender == "female") %>%
  ggplot(aes(sample = Weight %>% log2())) +
  geom_qq() +
  geom_qq_line()
```



Skewness is still there but is reduced already.

```
NHANES %>%
  filter(Age > 25 & Gender == "female") %>%
  ggplot(aes(x = Height, y = Weight %>% log2())) +
  ylab("Weight (log2)") +
  geom_point()
```



#### 4.1 Covariance and Correlation

- Let X and Y be to continuous random variables, and for each subject i we observe  $(X_i, Y_i)$ .
- Covariance: how deviate X\_i and Y\_i around their means?

$$Covar(X, Y) = E[(X - E[X])(Y - E[Y])]$$

• Correlation: standardise the covariance according to the variability in each variable:

$$\mathrm{Cor}(X,Y) = \frac{E[(X - E[X])(Y - E[Y])]}{\sqrt{E[(X - E[X])^2}\sqrt{E[(Y - E[Y])^2}}$$

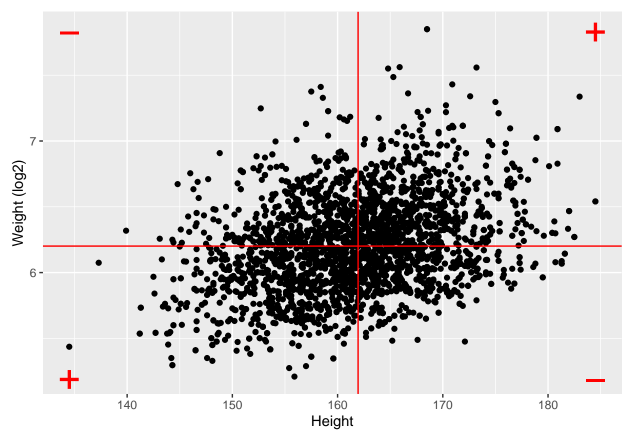
### 4.2 Pearson Correlation

• Association between two continuous covariate:

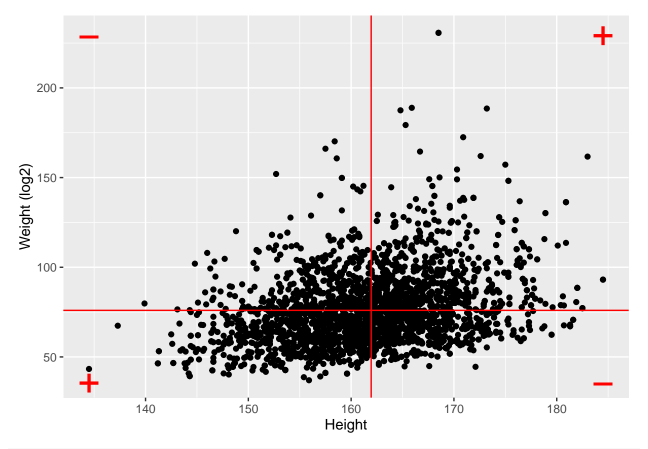
$$\operatorname{Cor}(X,Y) = \frac{\sum_{i=1}^n (X_i - \bar{X})(Y_i - \bar{Y})}{(n-1)s_X s_Y}$$

- Positive correlation:  $x \nearrow \Rightarrow y \nearrow$
- Negative correlation:  $x \nearrow \Rightarrow y \searrow$
- Correlation always between -1 en 1

```
means <- NHANES %>%
  filter(Age > 25 & Gender == "female") %>%
  select(Weight, Height) %>%
  mutate(log2Weight = Weight %>% log2()) %>%
  apply(., 2, mean, na.rm = TRUE)
ranges <- NHANES %>%
  filter(Age > 25 & Gender == "female") %>%
  select(Weight, Height) %>%
  mutate(log2Weight = Weight %>% log2()) %>%
  apply(., 2, range, na.rm = TRUE)
NHANES %>%
  filter(Age > 25 & Gender == "female") %>%
  ggplot(aes(x = Height, y = Weight %>% log2())) +
  ylab("Weight (log2)") +
  geom_point() +
  geom_hline(yintercept = means["log2Weight"], color = "red") +
  geom_vline(xintercept = means["Height"], color = "red") +
  annotate(
    geom = "text",
    x = c(ranges[1, "Height"], ranges[1, "Height"], ranges[2, "Height"], ranges[2, "Height"]),
    y = c(ranges[1, "log2Weight"], ranges[2, "log2Weight"], ranges[1, "log2Weight"], ranges[2, "log2Wei
    label = c("+", "-", "-", "+"), color = "red", size = 10
```



```
NHANES %>%
  filter(Age > 25 & Gender == "female") %>%
  ggplot(aes(x = Height, y = Weight)) +
  ylab("Weight (log2)") +
  geom_point() +
  geom_hline(yintercept = means["Weight"], color = "red") +
  geom_vline(xintercept = means["Height"], color = "red") +
  annotate(
    geom = "text",
    x = c(ranges[1, "Height"], ranges[1, "Height"], ranges[2, "Height"]),
    y = c(ranges[1, "Weight"], ranges[2, "Weight"], ranges[1, "Weight"], ranges[2, "Weight"]),
    label = c("+", "-", "-", "+"), color = "red", size = 10
)
```



```
NHANES %>%
  filter(Age > 25 & Gender == "female") %>%
  select(Weight, Height) %>%
  mutate(log2Weight = Weight %>% log2()) %>%
  na.exclude() %>%
  cor()
```

```
      Weight
      Height
      log2Weight

      Weight
      1.0000000
      0.3029834
      0.9812646

      Height
      0.3029834
      1.0000000
      0.3240824

      log2Weight
      0.9812646
      0.3240824
      1.0000000
```

• Note, that the correlation is lower when the data are not transformed.

- The Pearson correlation is sensitive to outliers!
- Do not use Pearson correlation for skewed distributions or for data with outliers.

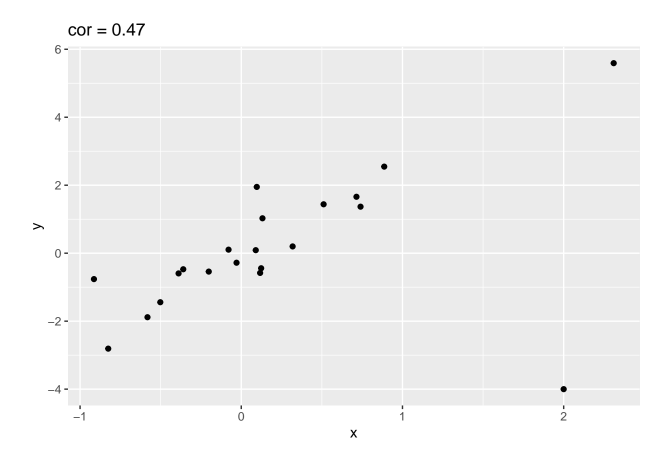
#### 4.2.1 Impact of outliers

Illustration with simulated data that has one outlier.

```
set.seed(100)
x <- rnorm(20)
simData <- data.frame(x = x, y = x * 2 + rnorm(length(x)))
simData %% ggplot(aes(x = x, y = y)) +
   geom_point() +
   ggtitle(paste("cor =", cor(simData[, 1], simData[, 2]) %>% round(., 2)))
```

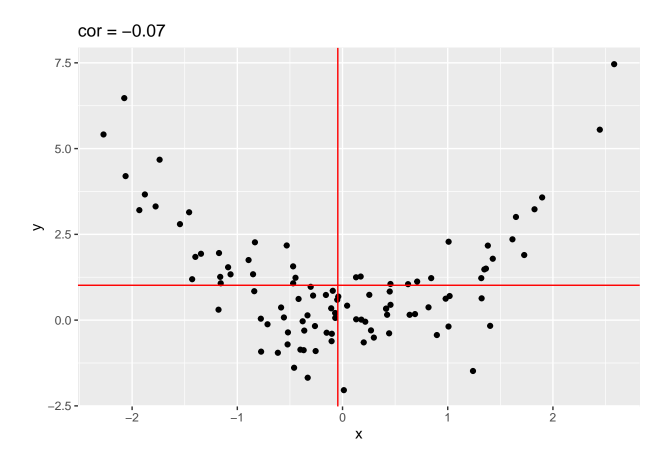
```
cor = 0.93
```

```
outlier <- rbind(simData, c(2, -4))
outlier %>% ggplot(aes(x = x, y = y)) +
  geom_point() +
  ggtitle(paste("cor =", cor(outlier[, 1], outlier[, 2]) %>% round(., 2)))
```



## 4.2.2 Only linear association

```
x <- rnorm(100)
quadratic <- data.frame(x = x, y = x^2 + rnorm(length(x)))
quadratic %>% ggplot(aes(x = x, y = y)) +
    geom_point() +
    ggtitle(paste("cor =", cor(quadratic[, 1], quadratic[, 2]) %>% round(., 2))) +
    geom_hline(yintercept = mean(quadratic[, 2]), col = "red") +
    geom_vline(xintercept = mean(quadratic[, 1]), col = "red")
```

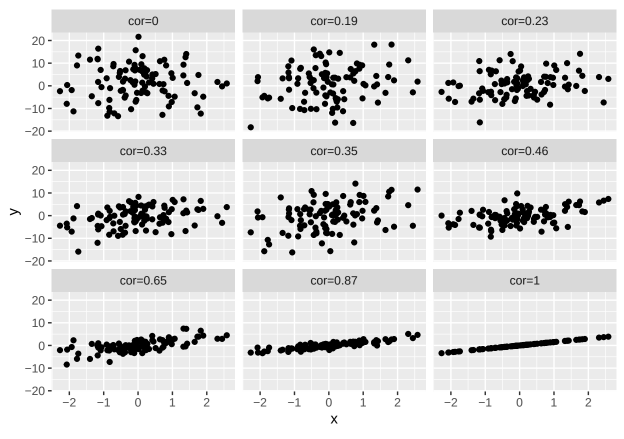


# 4.3 Different magnitudes of correlation

```
set.seed(100)
x <- rnorm(100)
simData2 <- cbind(x, 1.5 * x, sapply(1:7, function(sd, x) 1.5 * x + rnorm(length(x), sd = sd), x = x), sep = "=")

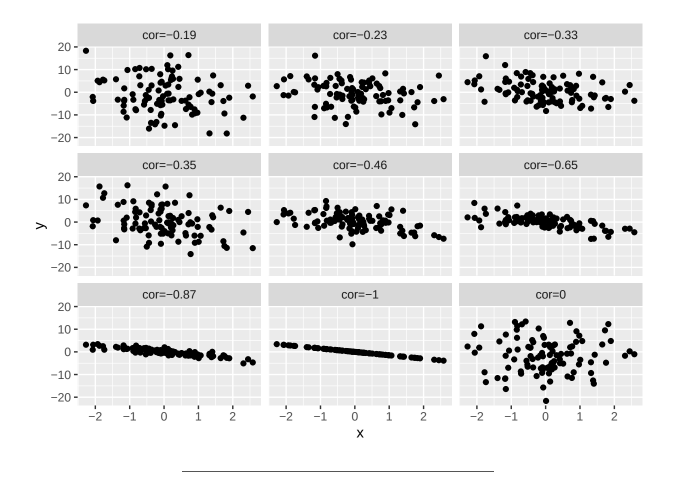
colnames(simData2)[-1] <- paste("cor", round(cor(simData2)[1, -1], 2), sep = "=")

simData2 %>%
    as.data.frame() %>%
    gather(cor, y, -x) %>%
    ggplot(aes(x = x, y = y)) +
    geom_point() +
    facet_wrap(~cor)
```



```
simData3 <- simData2
simData3[, -1] <- -simData2[, -1]
colnames(simData3)[-1] <- paste("cor", round(cor(simData3)[1, -1], 2), sep = "=")

simData3 %>%
   as.data.frame() %>%
   gather(cor, y, -x) %>%
   ggplot(aes(x = x, y = y)) +
   geom_point() +
   facet_wrap(~cor)
```



## 4.4 Spearman correlation

The Spearman correlation is the Pearson correlation after transforming the data to ranks.

• Pearson correlation

cor(outlier)

• Spearman correlation

```
cor(outlier, method = "spearman")
```

```
x y
x 1.0000000 0.6571429
y 0.6571429 1.0000000
```

- Spearman correlation is less sensitive to outliers.
- Pearson correlation on ranks

### y 0.6571429 1.0000000

• NHANES example

```
NHANES %>%
  filter(Age > 25 & Gender == "female") %>%
  select(Weight, Height) %>%
  mutate(log2Weight = Weight %>% log2()) %>%
  na.exclude() %>%
  cor(method = "spearman")
```

 Weight
 Height
 log2Weight

 Weight
 1.0000000
 0.3039481
 1.000000

 Height
 0.3039481
 1.0000000
 0.3039481

 log2Weight
 1.0000000
 0.3039481
 1.0000000