# Statistical tests

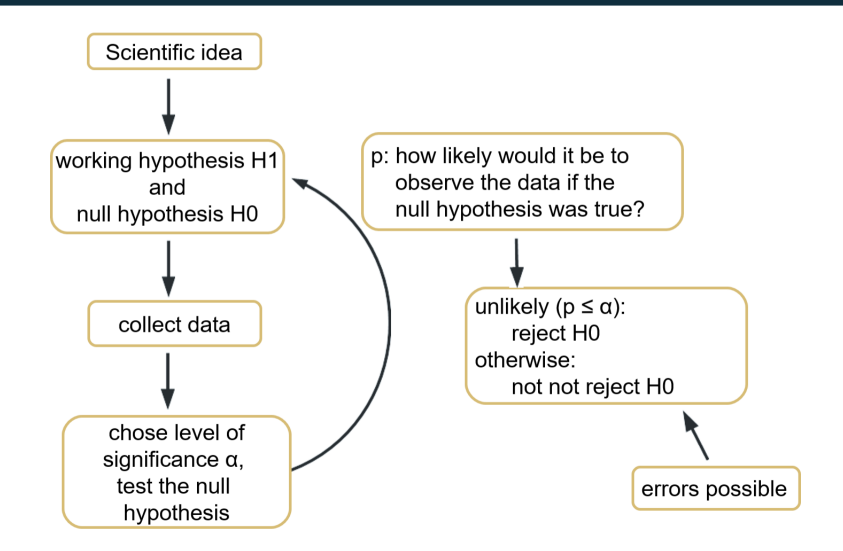
Introduction to R - Day 3

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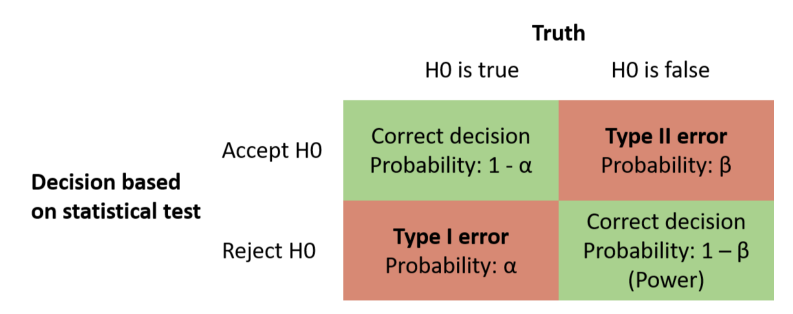
Freie Universität Berlin - Theoretical Ecology

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# Typical approach in statistics



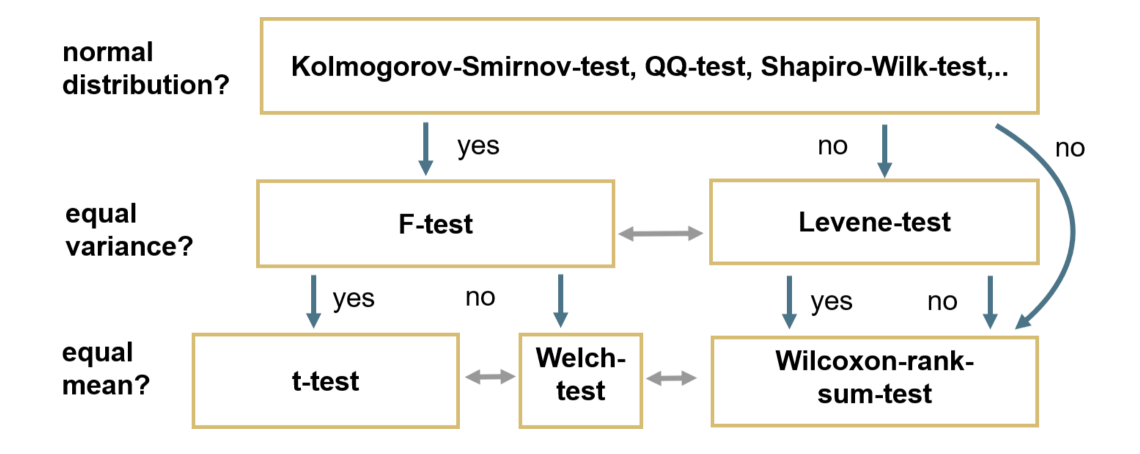
# Types of errors



Keep error probability as low as possible

- Decrease type I errors: choose smaller  $\alpha \rightarrow$  but increases type II errors
- Decrease type II errors: more data or different test
- → There are trade-offs and errors can't be completely avoided, especially with multiple tests applied in a row

### Overview of tests



# Tests for normal distribution

### Test for normal distribution

There are various tests and the outcome might differ!

#### Kolmogorov-Smirnov-Test (KS-Test)

- how much does the cumulative probability of observed data differs from normal distribution
- very general test (also works for other distributions)
- ullet low power: data not normally distributed but  $H_0$  not rejected

#### Shapiro-Wilk-Test

- how much does variance of observed data differ from normal distribution.
- specific test only for normal distribution
- high power, also for few data points

#### Visual tests: QQ-Plot

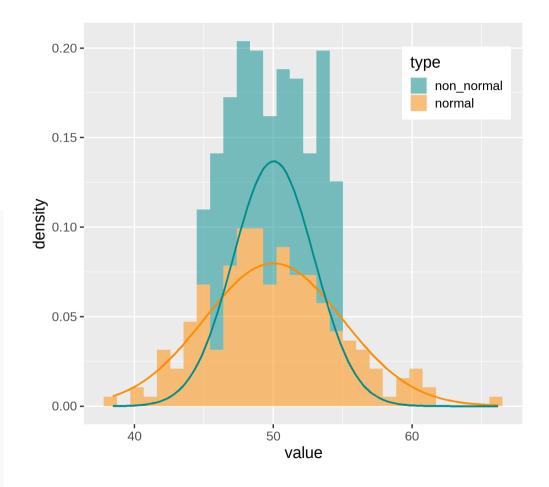
- quantiles of observed data plotted against quantiles of normal distribution
- scientist has to decide if normal or not

### The data

#### Create a tibble with two variables

- normal: 200 normally distributed values with mean 50 and standard deviation 5
- non\_normal: 200 uniformly distributed values between 45 and 55

```
set.seed(123)
mydata <- tibble(
  normal = rnorm(
    n = 200,
    mean = 50,
    sd = 5
),
  non_normal = runif(
    n = 200,
    min = 45,
    max = 55
)
)</pre>
```



### Short deviation: The \$ operator

With the \$ operator, you can select columns from a data frame / tibble.

The \$ operator will return the column as a vector

```
mydata$normal
## [1] 47.19762 48.84911 57.79354 50.35254 50.64644 58.57532 52.30458 43.67469
```

In comparison to dplyr::select() which will return the colum as a tibble

```
select(mydata, normal)
## # A tibble: 200 x 1
## normal
```

```
## normal
## <dbl>
## 1 47.2
## 2 48.8
## 3 57.8
## # ... with 197 more rows
```

This is important because most tests work with vectors as input.

# Kolmogorov-Smirnov-Test

 $H_0$ : Data does not differ from a normal distribution

```
ks.test(mydata$normal,
   "pnorm",
   mean = mean(mydata$normal),
   sd = sd(mydata$normal)
)

##

## One-sample Kolmogorov-Smirnov test
##

## data: mydata$normal
## D = 0.054249, p-value = 0.5983
## alternative hypothesis: two-sided
```

- D: test statistic, largest difference between cumulative distributions
- ullet p-value: probability to observe data if  $H_0$  was true
- two-sided test: we don't know in which direction our data deviates from normal distribution

Result: The data does not deviate significantly from a normal distribution (KS-Test, D = 0.054, p > 0.05)

# Kolmogorov-Smirnov-Test

 $H_0$ : Data does not differ from a normal distribution

```
ks.test(mydata$non_normal,
    "pnorm",
    mean = mean(mydata$non_normal),
    sd = sd(mydata$non_normal)
)
##
## One-sample Kolmogorov-Smirnov test
##
## data: mydata$non_normal
## D = 0.077198, p-value = 0.1843
## alternative hypothesis: two-sided
```

Result: The data does not deviate significantly from a normal distribution (KS-Test, D = 0.077, p > 0.05)

But: We know that our data is not normally distributed. Nevertheless, we would not reject  $H_0 \Rightarrow$  we would make a type II error here

# Shapiro-Wilk-Test

 $H_0$ : Data does not differ from a normal distribution

```
shapiro.test(mydata$normal)
##
## Shapiro-Wilk normality test
##
## data: mydata$normal
## W = 0.99076, p-value = 0.2298
```

- W: test statistic
- ullet p-value: probability to observe the data if  $H_0$  was true

The data does not deviate significantly from a normal distribution (Shapiro-Wilk-Test, W = 0.991, p > 0.05)

# Shapiro-Wilk-Test

 $H_0$ : Data does not differ from a normal distribution

```
shapiro.test(mydata$non_normal)
##
## Shapiro-Wilk normality test
##
## data: mydata$non_normal
## W = 0.95114, p-value = 2.435e-06
```

The data deviates significantly from a normal distribution (Shapiro-Wilk-Test, W = 0.95, p < 0.05)

### Visual test with QQ-Plot

Points should match the straight line. Small deviations are okay.

```
ggplot(mydata, aes(sample = normal)) +
  stat_qq() +
  stat_qq_line()
```

```
ggplot(mydata, aes(sample = non_normal)) +
  stat_qq() +
  stat_qq_line()
```

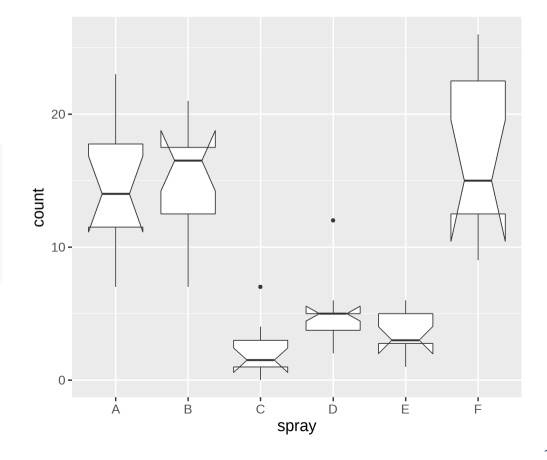
# Tests for equal variance

### The data

Counts of insects in agricultural units treated with different insecticides.

#### Compare treatments A, B and E:

 Create subsets before: count variable for each treatment as a vector



### Test for equal variance

First, test for normal distribution!

#### F-Test

- normal distribution of groups
- calculates ratio of variances (if equal, ratio = 1)
- p: How likely is ratio if variances were equal?
- only for normal distribution

#### Levene test

- non-normal distribution of groups
- compare difference between data sets with difference within data sets
- works with all distributions

# Test for equal variances

If we want to compare variances between treatments A, B and E, we first test for normal distribution

```
shapiro.test(TreatA)
     Shapiro-Wilk normality test
## data: TreatA
\#\#\ W = 0.95757, p-value = 0.7487
shapiro.test(TreatB)
##
      Shapiro-Wilk normality test
## data: TreatB
## W = 0.95031, p-value = 0.6415
shapiro.test(TreatE)
       Shapiro-Wilk normality test
## data: TreatE
\#\#\ W = 0.92128, p-value = 0.2967
```

Result: All 3 treatments are normally distributed.

#### F-Test

#### $H_0$ : Variances do not differ between groups

```
var.test(TreatA, TreatB)
##
## F test to compare two variances
##
## data: TreatA and TreatB
## F = 1.2209, num df = 11, denom df = 11, p-value = 0.7464
## alternative hypothesis: true ratio of variances is not equal to 1
## 95 percent confidence interval:
## 0.3514784 4.2411442
## sample estimates:
## ratio of variances
## 1.22093
```

- F: test statistics, ratio of variances (if F = 1, variances are equal)
- df: degrees of freedom of both groups
- ullet p-value: how likely is it to observe the data if  $H_0$  was true?

Result: The variances of sprays A and B do not differ significantly (F-Test,  $F_{11,11}$  = 1.22, p > 0.05)

#### F-Test

#### $H_0$ : Variances do not differ between groups

```
var.test(TreatA, TreatE)
##
## F test to compare two variances
##
## data: TreatA and TreatE
## F = 7.4242, num df = 11, denom df = 11, p-value = 0.002435
## alternative hypothesis: true ratio of variances is not equal to 1
## 95 percent confidence interval:
## 2.137273 25.789584
## sample estimates:
## ratio of variances
## 7.424242
```

Result: The variances of sprays A and E differ significantly (F-Test,  $F_{11,11}$  = 7.42, p < 0.05)

# Test for equal means

# Test for equal means

First test for normality and equal variances in the groups!

#### t-test

- normal distribution AND equal variance
- compares if mean values are within range or standard error of each other
- p: how likely is the difference if the means were equal

#### Welch-Test

- normal distribution but unequal variance
- corrected t-test

#### Wilcoxon rank sum test

- non-normal distribution and unequal variance
- compares rank sums of the data
- non-parametric



### t-test

 $H_0$ : The samples do not differ in their mean

Treatment A and B: normally distributed and equal variance

```
t.test(TreatA, TreatB, var.equal = TRUE)
##

## Two Sample t-test
##

## data: TreatA and TreatB
## t = -0.45352, df = 22, p-value = 0.6546
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -4.643994 2.977327
## sample estimates:
## mean of x mean of y
## 14.50000 15.33333
```

- t: test statistics (t = 0 means equal means)
- df: degrees of freedom of t-statistics
- ullet p-value: how likely is it to observe the data if  $H_0$  was true?

**Result:** The means of spray A and B do not differ significantly (t = -0.45, df = 22, p > 0.05)

### Welch-Test

 $H_0$ : The samples do not differ in their mean

Treatment A and E: normally distributed and non-equal variance

```
t.test(TreatA, TreatE, var.equal = FALSE)
##
## Welch Two Sample t-test
##
## data: TreatA and TreatE
## t = 7.5798, df = 13.91, p-value = 2.655e-06
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## 7.885546 14.114454
## sample estimates:
## mean of x mean of y
## 14.5 3.5
```

**Result:** The means of spray A and E do differ significantly (t = 7.58, df = 13.9, p < 0.05)

### Wilcoxon-rank-sum Test

 $H_0$ : The samples do not differ in their mean

We don't need the Wilcoxon test to compare treatment A and B, but for the sake of an example:

```
wilcox.test(TreatA, TreatB)
##
## Wilcoxon rank sum test with continuity correction
##
## data: TreatA and TreatB
## W = 62, p-value = 0.5812
## alternative hypothesis: true location shift is not equal to 0
```

**Result:** The means of spray A and E do not differ significantly (W = 62, p > 0.05)

### Paired values

Are there pairs of data points?

**Example:** samples of invertebrates across various rivers before and after sewage plants.

- for each plant, there is a pair of data points (before and after the plant)
- Question: Is the change (before-after) significant

Use paired = TRUE in the test.

```
t.test(treatA, treatB, var.equal = TRUE, paired = TRUE)
t.test(treatA, treatB, var.equal = FALSE, paired = TRUE)
wilcoxon.test(treatA, treatB, paired = TRUE)
```

Careful: your treatment vector both have to have the same order

# Now you

Task 1: Statistical tests

Find the task description here