Course 02402 Introduction to Statistics Lecture 5:

One-sample hypothesis test and model control

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Motivating example - sleeping medicine

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- One-sample t-test and p-value
- Oritical value and relation to confidence interval
- 4 Hypothesis test in general
 - The alternative hypothesis
 - The general method
 - Errors in hypothesis testing
- 6 Checking the normality assumption
 - The Normal QQ plot
 - Transformation towards normality

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Motivating example - sleeping medicine

Difference of sleeping medicines?

In a study the aim is to compare two kinds of sleeping medicine A and B. 10 test persons tried both kinds of medicine and the following 10 DIFFERENCES between the two medicine types were measured: (For person 1, sleep medicine B was 1.2 sleep hour better than medicine A, etc.):

	person	x = Beffect - Aeffect
Sample, $n = 10$:	1	1.2
1 7	2	2.4
	3	1.3
	4	1.3
	5	0.9
	6	1.0
	7	1.8
	8	0.8
	9	4.6
	10	1.4

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Example - sleeping medicine

The hypothesis of no difference:

$$H_0: \mu = 0$$

Sample mean and standard deviation:

$$\bar{x} = 1.670 = \hat{\mu}$$
$$s = 1.13 = \hat{\sigma}$$

Is data in acoordance with the null hypothesis H_0 ?

Data:
$$\bar{x} = 1.67, H_0: \mu = 0$$

NEW:p-value:

$$p - \text{value} = 0.00117$$

(Computed under the scenario, that H_0 is true)

NEW: Conclusion:

As the data is unlike far away from H_0 , we **reject** H_0 - we have found a **significant effect** of sleep medicine B as compared to A.

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One-sample t-test and p-value

Method 3.22: One-sample *t*-test and *p*-value

How to compute the *p*-value?

For a (quantitative) one sample situation, the (non-directional) p-value is given by:

$$p$$
 – value = $2 \cdot P(T > |t_{obs}|)$

where T follows a t-distribution with (n-1) degrees of freedom. The observed value of the test statistics to be computed is

$$t_{\rm obs} = \frac{\bar{x} - \mu_0}{s / \sqrt{n}}$$

where μ_0 is the value of μ under the null hypothesis:

$$H_0: \mu = \mu_0$$

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One-sample t-test and p-value

The definition and interpretation of the p-value (COMPLETELY general)

The p-value expresses the *evidence* against the null hypothesis – Table 3.1:

p < 0.001	Very strong evidence against H_0
$0.001 \le p < 0.01$	Strong evidence against H_0
$0.01 \le p < 0.05$	Some evidence against H_0
$0.05 \le p < 0.1$	Weak evidence against H_0
$p \ge 0.1$	Little or no evidence against H_0

Definition 3.21 of the *p*-value:

The p-value is the probability of obtaining a test statistic that is at least as extreme as the test statistic that was actually observed. This probability is calculated under the assumption that the null hypothesis is true.

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Example - sleeping medicine

The hypothesis of no difference:

$$H_0: \mu = 0$$

Compute the *p*-value:

Compute the test-statistic:

$$t_{\text{obs}} = \frac{1.67 - 0}{1.13 / \sqrt{10}} = 4.67$$

$$2P(T > 4.67) = 0.00117$$

2 * (1-pt(4.67, 9))

Interpretation of the *p*-value in light of Table 3.1:

There is strong evidence agains the null hypothesis.

One-sample t-test and p-value

Example - sleeping medicine - in R - with inbuilt function

```
t.test(x)
    One Sample t-test
##
## data: x
## t = 4.7, df = 9, p-value = 0.001
## alternative hypothesis: true mean is not equal to 0
## 95 percent confidence interval:
## 0.8613 2.4787
## sample estimates:
## mean of x
        1.67
```

Example - sleeping medicine - in R - manually

```
## Enter data:
x \leftarrow c(1.2, 2.4, 1.3, 1.3, 0.9, 1.0, 1.8, 0.8, 4.6, 1.4)
n <- length(x)
## Compute the tobs - the observed test statistic:
tobs \leftarrow (mean(x) - 0) / (sd(x) / sqrt(n))
## Compute the p-value as a tail-probability
## in the t-distribution:
pvalue \leftarrow 2 * (1-pt(abs(tobs), df=n-1))
pvalue
## [1] 0.001166
```

One-sample t-test and p-value

The definition of hypothesis test and significance (generally)

Definition 3.23. Hypothesis test:

We say that we carry out a hypothesis test when we decide against a null hypothesis or not using the data.

A null hypothesis is *rejected* if the *p*-value, calculated after the data has been observed, is less than some α , that is if the p-value $< \alpha$, where α is some pre-specifed (so-called) significance level. And if not, then the null hypothesis is said to be accepted.

Definition 3.28. Statistical significance:

An effect is said to be (statistically) significant if the p-value is less than the significance level α .

(OFTEN we use $\alpha = 0.05$)

Example - sleeping medicine

With $\alpha = 0.05$ we can conclude:

Since the p-value is less than α so we **reject** the null hypothesis.

And hence:

We have found a **significant effect** af medicine B as compared to A. (And hence that B works better than A)

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Critical value and relation to confidence interva

Critical value

Definition 3.30 - the critical values of the *t*-test:

The $(1-\alpha)100\%$ critical values for the (non-directional) one-sample t-test are the $(\alpha/2)100\%$ and $(1-\alpha/2)100\%$ quantiles of the *t*-distribution with n-1 degrees of freedom:

$$t_{\alpha/2}$$
 and $t_{1-\alpha/2}$

Metode 3.31: One-sample *t*-test by critical value:

A null hypothesis is *rejected* if the observed test-statistic is more extreme than the critical values:

If
$$|t_{\text{obs}}| > t_{1-\alpha/2}$$
 then reject

otherwise accept.

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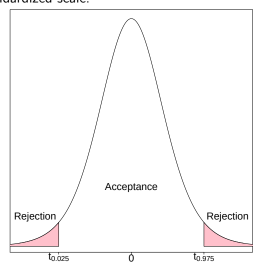
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Critical value and relation to confidence interval

Critical value and hypothesis test

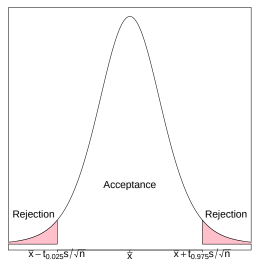
The acceptance region are the values for μ not too far away from the data - here on the standardized scale:



Critical value and relation to confidence interval

Critical value and hypothesis test

The acceptance region are the values for μ not too far away from the data - now on the original scale:



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Critical value and relation to confidence interval

Proof:

Remark 3.33

A μ_0 inside the confidence interval will fullfill that

$$|\bar{x} - \mu_0| < t_{1-\alpha/2} \cdot \frac{s}{\sqrt{n}}$$

which is equivalent to

$$\frac{|\bar{x} - \mu_0|}{\frac{s}{\sqrt{n}}} < t_{1-\alpha/2}$$

and again to

$$|t_{\rm obs}| < t_{1-\alpha/2}$$

which then exactly states that μ_0 is accepted, since the $t_{\rm obs}$ is within the critical values.

Critical value, confidence interval and hypothesis test

Theorem 3.32: Critical value method = Confidence interval method

We consider a $(1-\alpha) \cdot 100\%$ confidence interval for μ :

$$\bar{x} \pm t_{1-\alpha/2} \cdot \frac{s}{\sqrt{n}}$$

The confidence interval corresponds to the acceptance region for H_0 when testing the (non-directional) hypothesis

$$H_0: \mu = \mu_0$$

(New) interpretation of the confidence interval:

The confidence interval covers those values of the parameter that we believe in given the data.

Those values that we accept by the corresponding hypothesis test.

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Hypothesis test in general

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The alternative hypothesis

So far - implied: (= non-directional)

The alternative to H_0 : $\mu = \mu_0$ is : H_1 : $\mu \neq \mu_0$

BUT there are other possible settings, e.g. one-sided (=directional), "less":

The alternative to H_0 : $\mu = \mu_0$ is : H_1 : $\mu < \mu_0$

But we stick to the "non-directional" in this course

Hypothesis test in general The general method

The one-sample t-test again

Method 3.35 The level α test is:

- Compute t_{obs} as before
- **Q** Compute the evidence against the *null hypothesis* H_0 : $\mu = \mu_0$ vs. the alternative hypothesis $H_1: \mu \neq \mu_0$ by the

$$p$$
-value = $2 \cdot P(T > |t_{obs}|)$

where the *t*-distribution with n-1 degrees of freedom is used.

• If p-value $< \alpha$: We reject H_0 , otherwise we accept H_0 .

OR:

The rejection/acceptance conclusion could alternatively, but equivalently, be made based on the critical value(s) $\pm t_{1-\alpha/2}$:

If $|t_{\text{obs}}| > t_{1-\alpha/2}$ we reject H_0 , otherwise we accept H_0 .

Steps by hypothesis tests - an overview

Generelly a hypothesis test consists of the following steps:

- Formulate the hypotheses and choose the level of significance α (choose the "risk-level")
- Calculate, using the data, the value of the test statistic
- Calculate the p-value using the test statistic and the relevant sampling distribution, and compare the p-value and the significance level α and make a conclusion

OR:

Alternatively, make a conclusion based on the relevant critical value(s)

Hypothesis test in general Errors in hypothesis testing

Errors in hypothesis testing

Two kind of errors can occur (but only one at a time!)

Type I: Rejection of H_0 when H_0 is true

Type II: Non-rejection (acceptance) of H_0 when H_1 is true

The risks of the two types or errors:

$$P(\mathsf{Type}\;\mathsf{I}\;\mathsf{error}) = \alpha$$

$$P(\mathsf{Type}\;\mathsf{II}\;\mathsf{error}) = \beta$$

Hypothesis test in general Errors in hypothesis testing

Court of law analogy

A man is standing in a court of law:

A man is standing in a court of law accused of criminal activity. The null- and the the alternative hypotheses are:

The man is not guilty

The man is guilty

That you cannot be proved guilty is not the same as being proved innocent

Or differently put:

Accepting a null hypothesis is NOT a statistical proof of the null hypothesis being true!

Checking the normality assumptio

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Errors in hypothesis testing

Theorem 3.38: Significance level = The risk of a Type I error

The significance level α in hypothesis testing is the overall Type I risk:

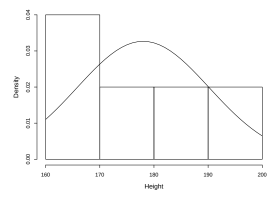
$$P(\mathsf{Type} \; \mathsf{I} \; \mathsf{error}) = P(\mathsf{Rejection} \; \mathsf{of} \; H_0 \; \mathsf{when} \; H_0 \; \mathsf{is} \; \mathsf{true}) = \alpha$$

Two possible truths vs. two possible conclusions:

	Reject H_0	Fail to reject H_0
H_0 is true	Type I error $(lpha)$	Correct acceptance of H_0
H_0 is false	Correct rejection of H_0 (Power)	Type II error $(oldsymbol{eta})$

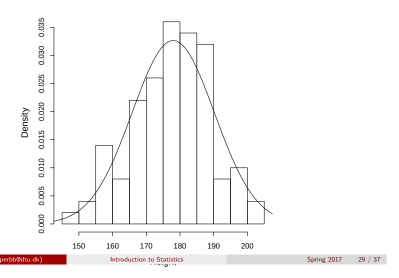
Example - student heights - are they normally distributed?

```
x \leftarrow c(168, 161, 167, 179, 184, 166, 198, 187, 191, 179)
hist(x, xlab="Height", main="", freq = FALSE)
lines(seq(160, 200, 1), dnorm(seq(160, 200, 1), mean(x), sd(x)))
```



Example - 100 observations from a normal distribution:

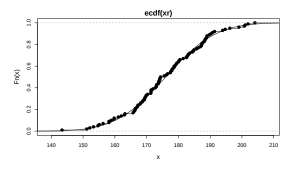
```
xr \leftarrow rnorm(100, mean(x), sd(x))
hist(xr, xlab="Height", main="", freq = FALSE)
lines(seq(130, 230, 1), dnorm(seq(130, 230, 1), mean(x), sd(x)))
```



Checking the normality assumption The Normal QQ plot

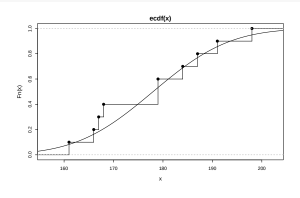
Example - 100 observations from a normal distribution, ecdf:

```
xr \leftarrow rnorm(100, mean(x), sd(x))
plot(ecdf(xr), verticals = TRUE)
xp \leftarrow seq(0.9*min(xr), 1.1*max(xr), length.out = 100)
lines(xp, pnorm(xp, mean(xr), sd(xr)))
```



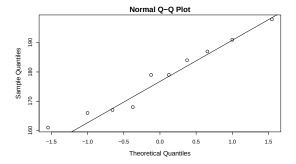
Example - student heights - ecdf

```
plot(ecdf(x), verticals = TRUE)
xp \leftarrow seq(0.9*min(x), 1.1*max(x), length.out = 100)
lines(xp, pnorm(xp, mean(x), sd(x)))
```



Example - student heights - Normal Q-Q plot

```
qqnorm(x)
qqline(x)
```

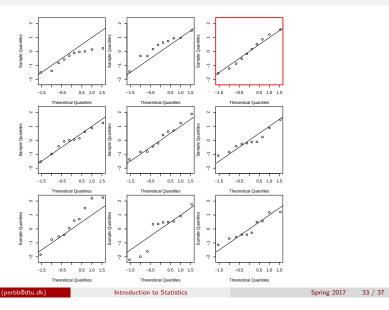


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Checking the normality assumption The Normal QQ plot

Checking the normality assumption The Normal QQ plot

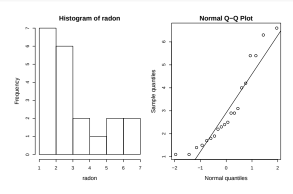
Example - student heights - Normal Q-Q plot - compare with other simulated normally distributed data



Checking the normality assumption Transformation towards normality

Example - Radon data

```
## READING IN THE DATA
radon<-c(2.4, 4.2, 1.8, 2.5, 5.4, 2.2, 4.0, 1.1, 1.5, 5.4, 6.3,
        1.9, 1.7, 1.1, 6.6, 3.1, 2.3, 1.4, 2.9, 2.9)
##A HISTOGRAM AND A QQ-PLOT
par(mfrow=c(1,2))
hist(radon)
qqnorm(radon,ylab = 'Sample quantiles',xlab = "Normal quantiles")
qqline(radon)
```



Normal Q-Q plot

Metode 3.41- The formal definition

The ordered observations $x_{(1)}, \dots, x_{(n)}$ are plotted versus a set of expected normal quantiles z_{p_1}, \ldots, z_{p_n} . Different definitions of p_1, \ldots, p_n exist:

• In R, when n > 10:

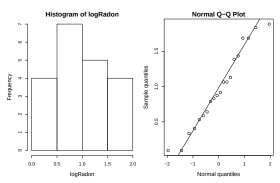
$$p_i = \frac{i - 0.5}{n + 1}, i = 1, \dots, n$$

• In R, when $n \le 10$:

$$p_i = \frac{i-3/8}{n+1/4}, i = 1, \dots, n$$

Example - Radon data - log-transformed are closer to a normal distribution

```
##TRANSFORM USING NATURAL LOGARITHM
logRadon <-log(radon)
hist(logRadon)
qqnorm(logRadon,ylab = 'Sample quantiles',xlab = "Normal quantiles")
qqline(logRadon)
```



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