# MA22004 – Statistics and Probability II

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# **Course Documents**

# Welcome

Welcome to MA22004 at the University of Dundee.

This course covers the basics of statistical inference. The first part of this document contains information of a practical nature regarding the mechanics of the course. The second part of this document contains the content.

These are some trying times.	
Please try to stay healthy.	

These notes are available at dundeemath.github.io/MA22004/ and also as a PDF (visit the page and click on the PDF icon to download).

## **Course Guide**

#### **Organisation**

This module runs for 11 teaching weeks and is worth 20 SCQF credits (equivalently, 10 ECTS points). All organisation and teaching will be carried out by:



Dr Eric Hall ehall001@dundee.ac.uk Mathematics Division Room TBA, Fulton Building 01382 TBA

This course uses Blackboard Ultra (look for course MA22004\_SEM0000\_2021) for communicating all announcements/deadlines and also for running online meetings. This course also uses Gradescope for submission of some of the continuous assessment items and Perusall for collaborative engagement with reading materials.

If you have a problem regarding the course, then you should make an appointment to see Dr Hall. You may also bring matters of concern about the course to the attention of the Mathematics Division Staff/Student Committee, which meets once each semester. A volunteer from Level 2 Mathematics will act as class representative to sit on the Staff/Student Committee (see Ultra for contact details).

#### **Timetable**



Due to COVID19, these plans may be subject to change.

The delivery of this module consists of a blend of synchronous and asynchronous content delivered both in-person and online. On an average week, there will be six planned teaching and learning activities.

The anticipated student effort is 200 hours over the length of the module. You are expected to be "present" for all synchronous timetabled activities except for the online office hours, which are optional. You may engage with the asynchronous material at your own pace, keeping in mind to meet any deadlines for engagement and/or attainment that will be posted to Ultra and discussed.

Activity	Timetabled	Group	Hours	Delivery
Reading	asynchronous	individually & in groups	6	online
Investigation	asynchronous	individually	1	online
Seminar	synchronous	whole class	1	online
Computer Lab	asynchronous	individually	6	online
Workshop Preparation	asynchronous	individually or in groups	2	online
Workshop	synchronous	in groups	1	face-to-face

#### **Pre-requisites**

To take this course, you must have passed module MA12003 or equivalent.

#### **Syllabus**

#### **Sampling Distributions**

Mean and standard deviation of samples, sampling from a single population, sampling from two populations, shape of sampling distributions. Normal distribution,  $\chi^2$ -square distribution, F-distribution.

#### **Hypothesis tests**

Null and Alternate hypotheses, inferences, confidence intervals, estimating means, proportions and standard deviations.

#### **Linear Regression**

Least squares, assessing usefulness of a model, using a model.

#### **Industrial Quality Control**

Control Charts, acceptance sampling.

#### R software package

Appropriate use of computational software to carry out statistical and probabilistic calculations.

#### **Recommended Books**

In addition to the course notes, here are some textbooks you may wish to consult.

- Probability and Statistics for Engineering and the Sciences, [Devore, 2016, §6-10, 12, 14, 16]
- Probability and Statistics, [DeGroot and Schervish, 2001, §7-10]
- Mathematical statistics and data analysis, [Rice, 2006, §6-12]
- All of Statistics, [Wasserman, 2004, Concise general reference]



You do not need to purchase these books.

#### Assessment

The module will be *continuously* assessed using coursework and examinations. Deadlines, as well as test dates, will be posted on Ultra and announcements made in the class hours. The module assessment weighting is as follows.

Assessment	Weight
Assignments	20%
Midterm Exam 1	20%
Midterm Exam 2	20%
Final Exam	40%

#### Coursework

Assessed coursework includes:

- six hand-in laboratory reports (there will be seven labs, the lowest grade will be dropped) and
- weekly engagement with the reading material using Perusall.

There will also be alternative means of demonstrating your mastery of course material through:

- one (group) lab presentation and
- short seminar quizzes (announced in advance).

#### **Examinations**

The **Midterm Exams** will be computer-assessed and will be one (1) hour in scope. These will likely be in weeks 4 and 8.

The **Final Exam** will be a two (2) hour hand-written exam that will be submitted using Gradescope. This process will be thoroughly discussed and trialled with a dummy exam in advance of the real submission. The Final Exam will be in week 11 (i.e., during the last week of the term).

To pass this module, you must:

- obtain an overall grade of at least D3 in the overall assessment and
- obtain a grade of at least M1 for the exam and
- obtain a grade of at least M1 for the coursework.

For those who fail the module, there may be an opportunity to take a two-hour resit examination paper at the next available exam diet.



Resit marks are based on the resit exam only.

Unless you have mitigating circumstances, if you fail to achieve a module grade of CF or above at first attempt, then you may not be permitted to resit the exam. Also, unless you have mitigating circumstances, any pass after a resit will be capped at a grade of D3 regardless of the weighted average mark obtained.

#### **Your Commitment**

You should attend all synchronous timetabled sessions except on medical grounds or with the special permission of Dr Hall. If you are unable to attend the degree examination or complete elements of the coursework on time, then you should inform Dr Hall and submit a medical certificate. Medical certificates should be submitted to your School Office as soon as possible after the absence. You must also submit a Mitigating Circumstances form to explain which aspects of assessment have been affected by your

absence. A Medical Certificate will only be taken into account if accompanied by a completed Mitigating Circumstances form that refers to the medical certificate.

#### **Approved Calculators**

The Casio FX83 and the Casio FX85 are the only calculators approved for use in assessments in the School of Engineering, Physics and Mathematics.

#### **Study Support**

If you are having difficulty with the course, you are encouraged to seek help at an early stage by making an appointment with Dr Hall. You may also obtain additional help from the Maths Base (see Ultra for details).

#### **Disability**

The University of Dundee is committed to making reasonable, effective and appropriate accommodations to meet the needs of students with disabilities and to create an inclusive and barrier-free campus. If you require accommodation for a documented disability, then you are advised to register with Disability Services. Please communicate any needs you may have directly with Dr Hall and as soon as possible to ensure timely management of any accommodations.

#### **Academic Honesty**

Honesty in scholarship and research is integral to the integrity of the academic enterprise of any higher education institution. Therefore, all students at the University of Dundee must practice academic honesty. Academic dishonesty includes cheating, fabrication, plagiarism, and facilitating dishonesty. Cases of academic dishonesty will be subject to appropriate sanctions and ignorance of such standards is not sufficient evidence of lack of intent. Please see the *Code of Practice on Academic Misconduct by Students* for more information about what constitutes academic dishonesty.

#### **End of Module Questionaire**

You will have the opportunity to complete a confidential questionnaire regarding the content and presentation of the module periodically. These questionaires form an important element in the University's Academic Standards procedures. Thank you in advance for your cooperation.

# Lab Guide

You will learn about the statistical programming language R and the software RStudio by working through 7 interactive lab tutorials and by completing lab reports.

These tutorials and all associated materials (templates, data sets, etc) are available as an R package at:  $\frac{1}{2} \left( \frac{1}{2} \right) = \frac{1}{2} \left( \frac{1}{2} \right) \left( \frac{1}{2} \right)$ 

https://github.com/dundeemath/MA22004labs.

# **Deadlines**

Below we have collected all the deadlines and important dates for the module. The three class tests (T1, T2, and T2) are on Wednesdays (W) from 13:00-15:00 and 14:00-17:00, respectively; we will practice (P) at uploading a test on Gradescope on Friday of week 1. Please keep in mind that you must engage with the weekly readings on Perusall **before** the seminar meeting on Wednesday afternoon. The lab reports (L1, L2, L3, L4, L5, L6, L7) are due on Thursday evenings at 17:00.

Further details about the practice upload and the class tests will be circulated nearer in time to the relevant deadline.

Week	Reading	Class Tests	Labs	Feedback (Planned)
1	Guide, §1	P (F15)	Install R, RStudio	
2	<b>§</b> 2		L1	P
3	§3.1		L2	L1
4	§3.2, 3.3	T1 (W13-15)		L2
5	§4.1, 4.2		L3	T1
6	§4.3, 4.4		L4	L3
7	§5		L5	L4
8	<b>§</b> 6	T2 (W13-15)		L5
9	§7		L6	T2
10	<b>§</b> 8		L7	L6
11	Review	T3 (W14-17)		L7
12				Т3

# **Course Notes**

# **Preliminaries**

#### **Notation**

Uppercase roman letters, e.g., X, will typically denote random variables (rvs); lower case letters, e.g., x, will denote a particular value (observation) of a rv. Rvs have probability distributions. Distributions are typically characterized by *parameters* which are fixed real numbers. Parameters describe population characteristics that are often unknown and must be estimated from data. Statistical inference is a tool that will help us to do this.



Statistical models comprise both rvs and parameters. Be careful not to confuse them!

#### **Abbreviations**

Abbreviation	Expanded
pdf	probability density function
cdf	cumulative distribution function
rv	random variable
iid	independent and identically distributed
obs	observations
CI	confidence interval
df	degrees of freedom

# Topic 1

# **Sampling distributions**

A **statistic** is a quantity that can be calculated from sample data. Prior to obtaining data, a statistic is an unknown quantity and is therefore a rv. We refer to the probability distribution for a statistic as a **sampling distribution** to emphasize how the distribution will vary across all possible sample data.

#### 1.1 Normal distribution

Normal distributions play an important role in probability and statistics as they describes many natural phenomenon. For instance, the Central Limit Theorem tells us that sums of rvs are approximately normal in distribution.

**Definition 1.1.** A continuous rv X has a **normal distribution** with parameters  $\mu$  and  $\sigma^2$ , where  $-\infty < \mu < \infty$  and  $\sigma > 0$ , if X has pdf

$$f(x; \mu, \sigma) = \frac{1}{\sqrt{2\pi}\sigma} e^{-(x-\mu)^2/(2\sigma^2)}, \quad -\infty < x < \infty.$$

We write  $X \sim N(\mu, \sigma^2)$ .

For  $X \sim N(\mu, \sigma^2)$ , it can be shown that  $\mathbf{E}(X) = \mu$  and  $\mathrm{Var}(X) = \sigma^2$ , that is,  $\mu$  is the *mean* and  $\sigma^2$  is the *variance* of X. The pdf takes the form of a bell-shaped curve that is symmetric about  $\mu$ . The value  $\sigma$  (*standard deviation*) is the distance from  $\mu$  to the inflection points of the curve. Thus, the position (location) and spread of the distribution depends on  $\mu$  and  $\sigma$ .

**Definition 1.2.** We say that X has a **standard normal distribution** if  $\mu = 0$  and  $\sigma = 1$  and we will usually denote standard Normal rvs by Z (why? tradition!).

#### 1.1.1 Some useful facts about normal variates

Here are some useful facts about how to manipulate Normal rvs.

- 1. If  $X \sim N(\mu, \sigma^2)$ , then  $Z = (X \mu)/\sigma \sim N(0, 1)$ .
- 2. If  $Z \sim N(0, 1)$ , then  $X = \mu + \sigma Z \sim N(\mu, \sigma^2)$ .
- 3. If  $X_i \sim N(\mu_i, \sigma_i^2)$  for i = 1, ..., n are independent rvs, then

$$\sum_{i=1}^{n} X_i \sim N\left(\sum_{i=1}^{n} \mu_i, \sum_{i=1}^{n} \sigma_i^2\right).$$

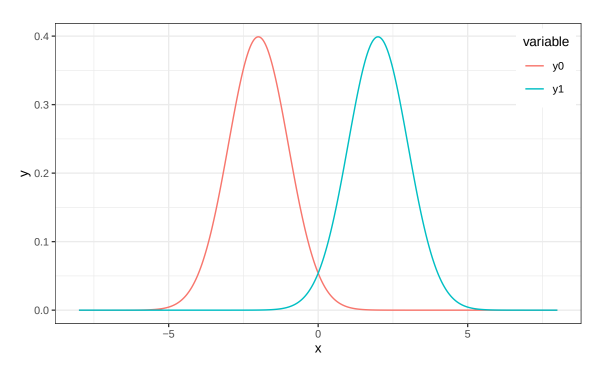


Figure 1.1: The pdfs of two normal rvs with different means and the same standard deviations.

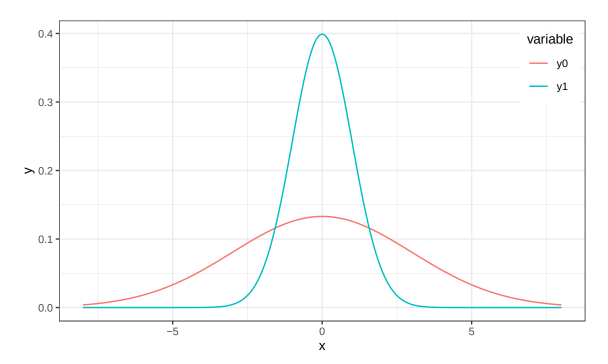


Figure 1.2: The pdfs of two normal rvs with the same means and different standard deviations.

In particular, we note that for differences of independent rvs  $X_1 \sim N(\mu_1, \sigma_1^2)$  and  $X_2 \sim N(\mu_2, \sigma_2^2)$  then the variances also add:

$$X_1 - X_2 \sim N(\mu_1 + \mu_2, \sigma_1^2 + \sigma_2^2)$$
.

Probabilities  $P(a \le X \le b)$  are found by converting the problem in  $X \sim N(\mu, \sigma^2)$  to the *standard normal* distribution  $Z \sim N(0, 1)$  whose probability values  $\Phi(z) = P(Z \le z)$  can then be looked up in a table. From (1.) above,

$$\begin{split} P(a < X < b) &= P\left(\frac{a - \mu}{\sigma} < Z < \frac{b - \mu}{\sigma}\right) \\ &= \Phi\left(\frac{b - \mu}{\sigma}\right) - \Phi\left(\frac{a - \mu}{\sigma}\right) \,. \end{split}$$

This process is often referred to as *standardizing* (the normal rv).

**Example 1.1.** Let  $X \sim N(5,9)$  and find  $P(X \ge 5.5)$ .

$$P(X \ge 5.5) = P\left(Z \ge \frac{5.5 - 5}{3}\right)$$

$$= P(Z \ge 0.1667)$$

$$= 1 - P(Z \le 0.1667)$$

$$= 1 - \Phi(0.1667)$$

$$= 1 - 0.5662$$

$$= 0.4338,$$

where we look up the value of  $\Phi(z) = P(Z \le z)$  in a table of standard normal curve areas.

The probability corresponds to the shaded area under the normal density  $\varphi(x) = \Phi'(x)$  corresponding to  $x \ge 5.5$ .

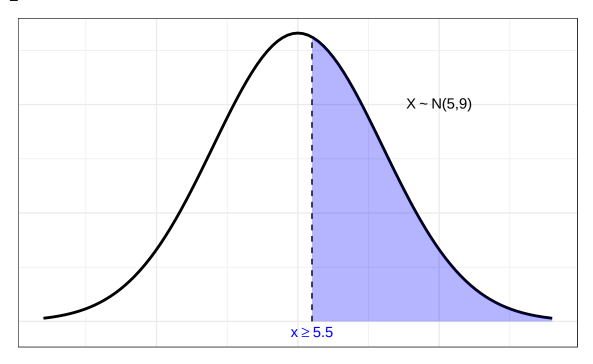


Figure 1.3: The normal density from the exercise with the (one-sided) interval shaded in blue.

Alternatively, we can use the r code: pnorm(5.5, mean = 5, sd = 3, lower.tail = FALSE).  $\Diamond$ 

**Example 1.2.** Let  $X \sim N(5, 9)$  and find  $P(4 \le X \le 5.25)$ .

$$P(4 \le X \le 5.25) = P\left(\frac{4-5}{3} \le Z \le \frac{5.25-5}{3}\right)$$

$$= P(-0.3333 \le Z \le 0.0833)$$

$$= \Phi(0.0833) - \Phi(-0.3333)$$

$$= 0.5332 - 0.3694$$

$$= 0.1638.$$

where we look up the value of  $\Phi(z) = P(Z \le z)$  in a table of standard normal curve areas.

The probability corresponds to the shaded area under the normal density  $\varphi(x) = \Phi'(x)$  corresponding to  $4 \le x \le 5.25$ .

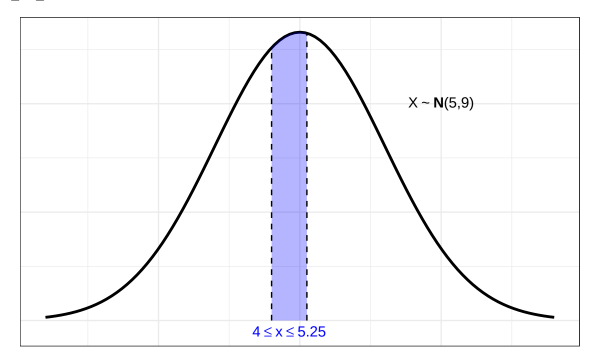


Figure 1.4: The normal density from the exercise with the interval shaded in blue.

Alternatively, we can use the r code: pnorm(5.25, mean = 5, sd = 3) - pnorm(4, mean = 5, sd = 3).  $\Diamond$ 

#### **1.1.2** Empirical rule (68 – 95 – 99.7 rule)

For samples from a normal distribution, the percentage of values that lie within one, two, and three standard deviations of the mean are 68.27%, 95.45%, and 99.73%, respectively. That is, for  $X \sim N(\mu, \sigma^2)$ ,

$$P(\mu - 1\sigma \le X \le \mu + 1\sigma) \approx 0.6827,$$
 
$$P(\mu - 2\sigma \le X \le \mu + 2\sigma) \approx 0.9545,$$
 
$$P(\mu - 3\sigma \le X \le \mu + 3\sigma) \approx 0.9973.$$

For a normal population, this means that nearly all the values lie within "three-sigmas" of the mean.

#### 1.2 t distribution

Student's t distribution gets its peculiar name as it was first published under the pseudonym "Student". This bit of obfuscation was to protect the identity of his employer, and thereby vital trade secrets, in a highly competitive and lucrative industry.

**Definition 1.3.** A continuous rv X has a t **distribution** with parameter  $\nu > 0$ , if X has pdf

$$f(x; v) = \frac{\Gamma\left(\frac{v+1}{2}\right)}{\sqrt{v\pi}\Gamma\left(\frac{v}{2}\right)} \left(1 + \frac{x^2}{v}\right)^{-\frac{v+1}{2}}, \quad -\infty < x < \infty.$$

We write  $X \sim \mathsf{t}(v)$ .

The density for t(v) for several values of v are plotted below.

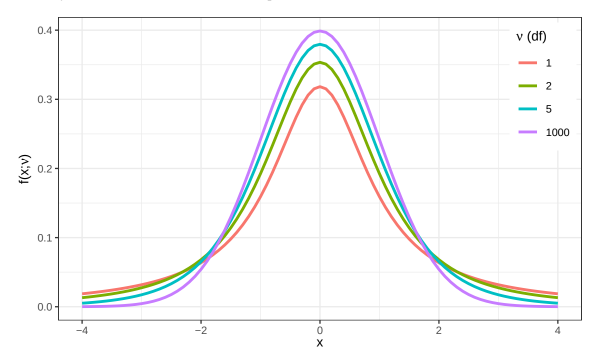


Figure 1.5: The density for t(v) for several values of v (df).

#### 1.2.1 Properties of t distributions

- 1. The density for t(v) is a bell-shaped curve centered at 0.
- 2. The density for t(v) is more spread out than the standard normal density (i.e., it has "fatter tails" than the normal).
- 3. As  $v \to \infty$ , the spread of the corresponding t(v) density converges to the standard normal density (i.e., the spread of the t(v) density decreases relative to the standard normal).

If  $X \sim \mathsf{t}(v)$ , then  $\mathbf{E}[X] = 0$  for v > 1 (otherwise the mean is undefined).

<sup>&</sup>lt;sup>1</sup>https://mathshistory.st-andrews.ac.uk/Biographies/Gosset/

<sup>&</sup>lt;sup>2</sup>https://www.wikiwand.com/en/Guinness Brewery

#### 1.3 $\chi^2$ distribution

The  $\chi^2$  distribution arises as the distribution of a sum of the squares of  $\nu$  independent standard normal rvs.

**Definition 1.4.** A continuous rv X has a  $\chi^2$  distribution with parameter  $\nu \in \mathbb{N}_>$ , if X has pdf

$$f(x; v) = \frac{1}{2^{\nu/2} \Gamma(\nu/2)} x^{(\nu/2)-1} e^{-x/2},$$

with support  $x \in (0, \infty)$  if v = 1, otherwise  $x \in [0, \infty)$ . We write  $X \sim \chi^2(v)$ .

The pdf f(x; v) of the  $\chi^2(v)$  distribution depends on a positive integer v referred to as the df. The density for several values of v are plotted below.

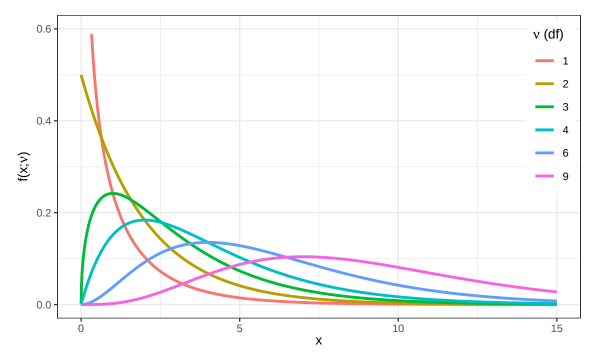


Figure 1.6: The density for  $\chi^2(\nu)$  for several values of  $\nu$  (df).

The density f(x; v) is positively skewed, i.e., the right tail is longer and hence the mass is concentrated to the left of the figure. The distribution becomes more symmetric as v increases. We denote critical values of the  $\chi^2(v)$  distribution by  $\chi^2_{\alpha,v}$ .



Unlike the normal and t distributions, the  $\chi^2$  distribution is not symmetric. This means that the critical values e.g.  $\chi^2_{.99,\nu}$  and  $\chi^2_{0.01,\nu}$  are **not** equal. Hence, it will be necessary to look up both values for CIs based on  $\chi^2$  critical values.

If  $X \sim \chi^2(\nu)$ , then  $\mathbf{E}[X] = \nu$  and  $Var[X] = 2\nu$ .

#### 1.4 F distribution

The F distribution arises as a test statistic when comparing population variances and in ANOVA.

**Definition 1.5.** A continuous rv X has an F **distribution** with df parameters  $v_1$  and  $v_2$ , if X has pdf

$$f(x;v_1,v_2) = \frac{\Gamma\left(\frac{v_1+v_2}{2}\right)v_1^{v_1/2}v_2^{v_2/2}}{\Gamma\left(\frac{v_1}{2}\right)\Gamma\left(\frac{v_2}{2}\right)} \frac{x^{v_1/2-1}}{(v_2+v_1x)^{(v_1+v_2)/2}} \,.$$

The pdf  $f(x; v_1, v_2)$  of the  $F(v_1, v_2)$  distribution depends on two positive integers  $v_1$  and  $v_2$  referred to, respectively, as the numerator and denominator df. The density is plotted below for several combinations of  $(v_1, v_2)$ .

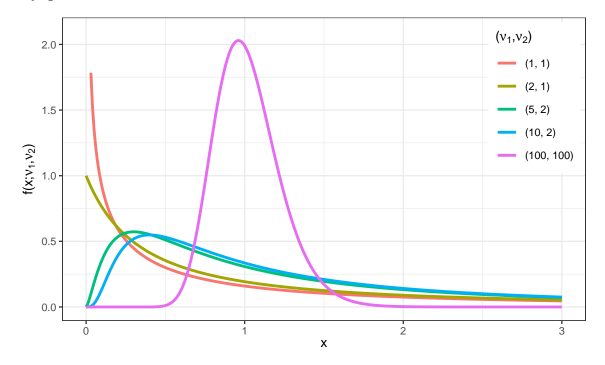


Figure 1.7: The density for  $F(v_1, v_2)$  for several combinations of  $(v_1, v_2)$ .

Where do the terms numerator and denominator df come from? The F distribution is related to ratios of  $\chi^2$  rvs.

**Theorem 1.1.** If  $X_1 \sim \chi^2(v_1)$  and  $X_2 \sim \chi^2(v_2)$  are independent rvs, then the rv

$$F = \frac{X_1/\nu_1}{X_2/\nu_2} \sim \mathsf{F}(\nu_1, \nu_2),$$

that comprises the ratio of two  $\chi^2$  rvs divided by their respective df has an  $F(v_1, v_2)$  distribution.

# **Topic 2**

## **Basics of statistical inference**

We discuss point estimation, confidence intervals, and hypothesis testing in Sections 2.1, 2.2, and 2.3, respectively. These three tools will form the basis for making inferences about a population.

#### 2.1 Point estimation

Statistical inference seeks to draw conclusions about the characteristics of a population from data. For example, suppose we are botanists interested in taxonomic classification of iris flowers. Let  $\mu$  denote the true average petal length (in cm) of the *Iris setosa*<sup>1</sup> (AKA the bristle-pointed iris). The parameter  $\mu$  is a characteristic of the whole population of the *setosa* species. Before we collect data, the petal lengths of m independent *setosa* flowers are denoted by rvs  $X_1, X_2, \ldots, X_m$ . Any function of the  $X_i$ 's, such as the sample mean,

$$\overline{X} = \frac{1}{m} \sum_{i=1}^{m} X_i, \qquad (2.1)$$

or the sample variance,

$$S^{2} = \frac{1}{m-1} \sum_{i=1}^{m} (X_{i} - \overline{X})^{2}, \qquad (2.2)$$

is also a rv.

Suppose we actually find and measure the petal length of 50 independent *setosa* flowers resulting in observations  $x_1, x_2, \ldots, x_{50}$ ; the distribution (counts) of 50 such petal length measurements are displayed in Figure 2.1. The sample mean  $\bar{x}$  for petal length can then be used to draw a conclusion about the (true) value of the population mean  $\mu$ . Based on the data in Figure 2.1 and using (2.1), the value of the sample mean is  $\bar{x} = 1.462$ . The value  $\bar{x}$  provides a "best guess" or point estimate for the true value of  $\mu$  based on the m = 50 samples.



The botonist Edgar Anderson's **Iris Data** contains 50 obs. of four features (sepal length [cm], sepal width [cm], petal length [cm], and petal width [cm]) for each of three plant species (*setosa*, *virginica*, *versicolor*) for 150 obs. total. This data set can be accessed in r by loading library(datasets) and then calling data(iris).

<sup>&</sup>lt;sup>1</sup>https://www.wikiwand.com/en/Iris\_setosa

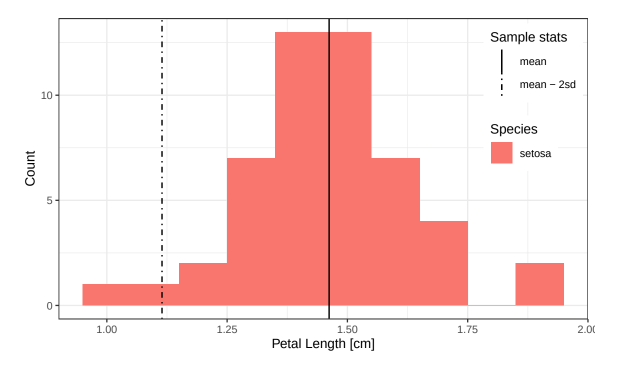


Figure 2.1: The distribution (counts) of m = 50 setosa petal length measurments.

**Definition 2.1.** A **point estimate** of a parameter  $\theta$  (recall: a fixed, unknown quantity) is a single number that we regard as a sensible value for  $\theta$ . Consider iid  $X_1, X_2, \dots, X_m \sim F(\theta)$ . A **point estimator**  $\hat{\theta}_m$  of  $\theta$  is obtained by selecting a suitable statistic g,

$$\hat{\theta}_n = g(X_1, \dots, X_m).$$

A point estimate  $\hat{\theta}_m$  can then be computed from the estimator using sample data.



The symbol  $\hat{\theta}_m$  (or simply  $\hat{\theta}$  when the sample size m is clear from context) is typically used to denote both the estimator and the point estimate resulting from a given sample. Note that writing, e.g.,  $\hat{\theta} = 42$  does not indicate how the point estimate was obtained. Therefore, it is essential to report both the estimator and the resulting point estimate.

Definition 2.1 does not say how to select an appropriate statistic. For the *setosa* example, the sample mean  $\overline{X}$  is suggested as a good estimator of the population mean  $\mu$ . That is,  $\widehat{\mu} = \overline{X}$  or "the point estimator of  $\mu$  is the sample mean  $\overline{X}$ ". Here, while  $\mu$  and  $\sigma^2$  are fixed quantities representing characteristics of the population,  $\overline{X}$  and  $S^2$  are rvs with sampling distributions. If the population is *normally distributed* or if the *sample is large* then the sampling distribution for  $\overline{X}$  has a known form:  $\overline{X}$  is normal with mean  $\mu_{\overline{X}} = \mu$  and variance  $\sigma^2_{\overline{X}} = \sigma^2/m$ , i.e.,

$$\overline{X} \sim N(\mu, \sigma^2/m)$$
,

where m is the sample size and  $\mu$  and  $\sigma$  are the (typically unknown) population parameters.

**Example 2.1.** Let us consider the heights (measured in inches) of 31 black cherry trees (sorted, for your enjoyment) in Table 2.1.

Table 2.1: Observations of m = 31 felled black cherry trees.

Height [in]

63, 64, 65, 66, 69, 70, 71, 72, 72, 74, 74, 75, 75, 75, 76, 76, 77, 78, 79, 80, 80, 80, 80, 80, 81, 81, 82, 83, 85, 86, 87



The **Cherry Tree Data** contains 31 obs. of three features (diameter, height, and volume) and can be accessed in r by loading library(datasets) and then calling data(trees).

The quantile-quantile plot in Figure 2.2, that compares the quantiles of this data to the quantiles of a normal distribution, is fairly straight.<sup>2</sup> Therefore, we assume that the distribution of black cherry tree heights is (at least approximately) normal with a mean value  $\mu$ ; i.e., that the population of heights is distributed  $N(\mu, \sigma^2)$  where  $\mu$  is a parameter to be estimated and  $\sigma^2$  is unknown. The observations  $X_1, \ldots, X_{31}$  are then assumed to be a random sample from this normal distribution, i.e., iid

$$X_1,\ldots,X_{31} \sim \mathsf{N}(\mu,\sigma^2)$$
.

Consider the following three different estimators and the resulting point estimates for  $\mu$  based on the 31 samples in Table 2.1.

- a. Estimator (sample mean)  $\overline{X}$  as in (2.1) and estimate  $\overline{x} = \sum x_i/n = 2356/31 = 76$ .
- b. Estimator (average of extreme heights)  $\widetilde{X} = [\min(X_i) + \max(X_i)]/2$  and estimate  $\widetilde{x} = (63 + 87)/2 = 75$ .
- c. Estimator (10% trimmed mean i.e., in this instance exclude the smallest and largest three values)  $\overline{X}_{tr(10)}$  and estimate  $\overline{x}_{tr(10)} = (2356 63 64 65 87 86 85)/25 = 76.24$ .

Each estimator above uses a different notion of center for the sample data, i.e., a different statistic. An interesting question to think about is: which estimator will tend to produce estimates closest to the true parameter value? Will the estimators work universally well for all distributions?  $\Diamond$ 

In addition to reporting a point estimate together with its estimator, some indication of its precision should be given. One measure of the precision of an estimate is its standard error.

**Definition 2.2.** The **standard error** of an estimator  $\hat{\theta}$  is the standard deviation

$$\sigma_{\hat{\theta}} = \sqrt{\operatorname{Var}(\hat{\theta})}$$
.

Often, the standard error depends on unknown parameters and must also be estimated. The **estimated standard error** is denoted by  $\hat{\sigma}_{\hat{\theta}}$  or simply  $s_{\hat{\theta}}$ .

#### 2.2 Confidence intervals

An alternative to reporting a point estimate for a parameter is to report an interval estimate suggesting an entire range of plausible values for the parameter of interest. A confidence interval is an interval estimate

<sup>&</sup>lt;sup>2</sup>How do we tell whether a population is normal? Constructing a normal quantile-quantile plot is one way of assessing whether a normality assumption is reasonable; such a plot compares the quantiles of the sample data  $x_i$  against the (theoretical) standard normal quantiles (see Figure 2.2). If the sample data is consistent with a sample from a normal distribution, then the points will lie on a straight line (more or less). The QQ plot Figure 2.2 comparing quantiles of cherry tree heights from Table 2.1 to normal quantiles.

<sup>&</sup>lt;sup>3</sup>The standard error is sometimes denoted se = se( $\hat{\theta}$ ) and the estimated standard error by se.

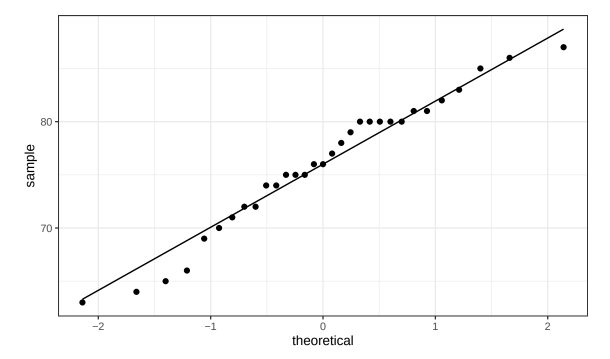


Figure 2.2: Normal quantile-quantile plot for the **Cherry Tree Data**.

that makes a probability statement about the degree of reliability, or the confidence level, of the interval. The first step in computing a confidence interval is to select the confidence level  $\alpha$ . A popular choice is a 95% confidence interval which corresponds to level  $\alpha = 0.05$ .

**Definition 2.3.** A  $100(1-\alpha)\%$  **confidence interval** for a parameter  $\theta$  is a *random* interval  $C_m = (L_m, U_m)$ , where  $L_m = \ell(X_1, \dots, X_m)$  and  $U_m = u(X_1, \dots, X_m)$  are functions of the data, such that

$$P_{\theta}(L_m < \theta < U_m) = 1 - \alpha, \qquad (2.3)$$

for all  $\theta \in \Theta$ .

My favorite interpretation of a confidence interval is due to [Wasserman, 2004, p 92]:

On day 1, you collect data and construct a 95 percent confidence interval for a parameter  $\theta_1$ . On day 2, you collect new data and construct a 95 percent confidence interval for an unrelated parameter  $\theta_2$ . On day 3, you collect new data and construct a 95 percent confidence interval for an unrelated parameter  $\theta_3$ . You continue this way constructing confidence intervals for a sequence of unrelated parameters  $\theta_1$ ,  $\theta_2$ , ... Then 95 percent of your intervals will trap the true parameter value. There is no need to introduce the idea of repeating the same experiment over and over.

This interpretation makes clear that a confidence interval is not a probability statement about the parameter  $\theta$ . In Definition 2.3, note that  $\theta$  is fixed ( $\theta$  is not a rv) and the interval  $C_m$  is random. After data has been collected and a point estimator has been calculated, the resulting CIs either contain the true parameter value or they do not, as illustrated in Figure 2.3.

#### 2.3 Hypothesis testing

Sections 2.1 and 2.2 reviewed how to estimate a parameter by a single number (point estimate) or range of plausible values (confidence interval), respectively. Next we discuss methods for determining which

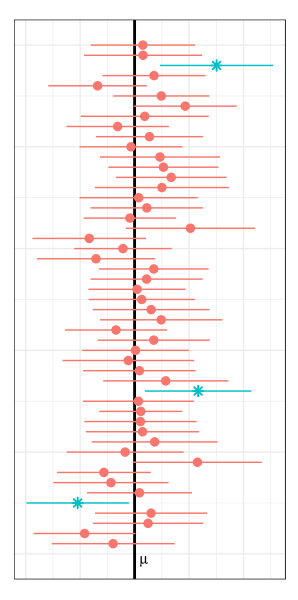


Figure 2.3: Fifty 95% CIs for a population mean  $\mu$ . After a sample is taken, the computed interval estimate either contains  $\mu$  or it does not (asterisk identify intervals that do not include  $\mu$ ). When drawing such a large number of 95% CIs, we would anticipate that approximately 5% (ca. 2 or 3) would fail to cover the true parameter  $\mu$ .

of two contradictory claims, or hypotheses, about a parameter is correct.

**Definition 2.4.** The **null hypothesis**, denoted by  $H_0$ , is a claim that we intially assume to be true by dafault. The **alternative hypothesis**, denoted by  $H_a$ , is an assertion that is contradictory to  $H_0$ .

Typically, we shall consider hypothesis concerning a parameter  $\theta \in \Theta$  taking values in a parameter space. The statistical hypothesis are contradictory in that  $H_0$  and  $H_a$  divide  $\Theta$  into two disjoint sets. For example, for a statistical hypothesis regarding the *equality* of a parameter  $\theta$  with a fixed quantity  $\theta_0$ , the null and alternative hypotheses will usually take one of the following forms in Table 2.2.

Table 2.2: Typical null hypothesis and corresponding alternative hypothesis.

Null hypothesis	Alternative hypothesis	Test form
$H_0$ : $\theta = \theta_0$	$H_a:\theta\neq\theta_0$	two-sided test
$H_0: \theta \leq \theta_0$	$H_a: \theta > \theta_0$	one-sided test
$H_0: \theta \ge \theta_0$	$H_a: \theta < \theta_0$	one-sided test

These pairs of hypothesis are associated with either a one-sided or two-sided test; what this means will become clear in the sequel. The value  $\theta_0$ , called the **null value**, separates the alternative from the null.

**Definition 2.5.** A **hypothesis test** asks if the available data provides sufficient evidence to reject  $H_0$ . If the observations disagree with  $H_0$ , then we reject the null hypothesis. If the sample evidence does not strongly contradict  $H_0$ , then we continue to believe  $H_0$ . The two possible conclustions of a hypothesis test are: reject  $H_0$  or fail to reject  $H_0$ .<sup>4</sup>

A procedure for carrying out a hypothesis test is based on specifying two additional items: a test statistic and a corresponding rejection region. A **test statistic** T is a function of the sample data (like an estimator). The decision to reject or fail to reject  $H_0$  will involve computing the test statistic. The **rejection region** R is the collection of values of the test statistic for which  $H_0$  is to be rejected in favor of the alternative, e.g.,

$$R = \{x : T(x) > c\},$$

where c is referred to as a **critical value**. If a given sample falls in the rejection region, then we reject  $H_0$ . That is, if  $X \in R$  (e.g., the calculated test statistic exceeds some critical value), then we reject  $H_0$ . The alternative is that  $X \notin R$  and in this case we fail to reject the null.

When carrying out a hypothesis test, two types of errors can be made. The basis for choosing a rejection region involves considering these errors.

**Definition 2.6.** A **type I** error occurs if  $H_0$  is rejected when  $H_0$  is actually true. A **type II** error is made if we fail to reject  $H_0$  when  $H_0$  is actually false.

If the maximal type I error of a test is fixed at an acceptably small value, then the type II error decreases as the sample size increases. In particular, a conclusion is reached in a hypothesis test by selecting a **significance level**  $\alpha$  for the test linked to the maximal type I error rate. Typically,  $\alpha = 0.10, 0.05, 0.01$ , or 0.001 is selected for the significance level.

**Definition 2.7.** A *P*-value is the probability, calculated assuming  $H_0$  is true, of obtaining a value of the test statistic at least as contradictory to  $H_0$  as the value calculated from the sample data.

Smaller P-values indicate stronger evidence against  $H_0$  in favor of  $H_a$ . If  $P \leq \alpha$  then we reject  $H_0$  at significance level  $\alpha$ . If  $P \ge \alpha$  we fail to reject  $H_0$  at significance level  $\alpha$ .



The P-value is a probability calculated assuming that  $H_0$  is true. However, the P-value is not the probability that:

1.  $H_0$  is TRUE,

2.  $H_0$  is FALSE, or

3. a wrong conclusion is reached.

#### **Proposition 2.1.** The hypothesis test procedure that

$$\begin{cases} \textit{rejects } H_0 & \textit{if } P \leq \alpha \\ \textit{fails to reject } H_0 & \textit{otherwise} \end{cases}$$

has  $P(type\ I\ error) = \alpha$ .

**Example 2.2.** Churchill claims that he will receive half the votes for the House of Commons seat for the constituency of Dundee.<sup>5</sup> If we do not believe Churchill's claim and we are doubtful of his popularity, then we would seek to test an alternative hypothesis. How should we write down our research hypotheses?

If we let p be the fraction of the population voting for Churchill, then we have the null hypothesis,

$$H_0: p = 0.5$$
,

and the alternative hypothesis (we believe Churchill is less popular than he claims),

$$H_a: p < 0.5$$
.

Support for the alternative hypothesis is obtain by showing lack of support for it's converse hypothesis (the null hypothesis).  $\Diamond$ 

**Example 2.3.** Suppose that m = 15 voters are selected from Dundee and X, the number favoring Churchill, is recorded. Based on observing X, we construct a rejection region  $R = \{x : x \le k\}$ . If k is small compared to m, then the rejection region would provide pretty strong evidence to reject  $H_0$ . How should one choose the rejection region?

Assume now that m = 15 voters are polled and that we select k = 2 to have a rejection region  $R = \{x \le 2\}$ . For this choice of k, the rejection region R provides strong support to reject  $H_0$ . If we assume the null hypothesis is true, then we would expect that approximately half of the 15 voters (ca. 7) would plump for Churchill. Observing x = 0, x = 1 or x = 2 (the values that would place us in the rejection region) would provide strong evidence against  $H_0$ .

We can calculate the probability of a type I error. From the definition of type I error,

$$\alpha = P(\text{type I error})$$

$$= P(\text{rejecting } H_0 \text{ when } H_0 \text{ is true})$$

$$= P(X \in R \text{ when } H_0 \text{ is true})$$

$$= P(X \le 2 \text{ when } p = 0.5).$$

<sup>&</sup>lt;sup>5</sup>Sir Winston Churchill was Member of Parliament for Dundee from 1908–1922 [https://www.wikiwand.com/en/Winsto n\_Churchill].

Since  $X \sim \text{Binom}(15, 0.50)$ , we calculate that  $\alpha = 0.00369$ . Thus, for this particular choice of rejection region R, the risk of concluding that Churchill will lose if in fact he is the winner is very small.

For this rejection region, how good is the test at protecting us from type II errors, i.e., concluding that Churchill is the winner if in fact he will lose? Suppose that Churchill receive 25 of the votes (p = 0.25). The probability of a type II error  $\beta$ ,

```
\beta = P(\text{type II error})
  = P(\text{fail to reject } H_0 \text{ when } H_0 \text{ false})
  = P(X \notin R \text{ when } H_0 \text{ false})
  = P(X > 2 \text{ when } p = 0.3).
```

For  $X \sim \text{Binom}(15, 0.25)$ , we calculate  $\beta = 0.764$ . If we use  $R = \{x \le 2\}$  then our test will lead us to conclude that Churchill is the winner with probability 0.764 even if p is as low as 0.25!

If we repeat these calculations for  $R^* = \{x \le 5\}$ , we find  $\alpha = 0.151$  versus  $\beta = 0.148$ , even if p is as low as 0.25, which is a much better balance between type I and type II errors.  $\Diamond$ 



To summarize, the elements of a statistical test are:

- 1. Null hypothesis  $(H_0)$
- Null hypothesis (H<sub>a</sub>)
   Alternative hypothesis (H<sub>a</sub>)
   Test statistic
   Rejection region
   Significance level (α)

 $<sup>^{6}</sup>X$  is a binomial random variable because it can be modeled as m independent Bernoulli trails each with probability p of success (i.e. votes Churchill) as long as the sample size m is much smaller than the population of Dundee. If we had the means to canvas nearly the whole population, what goes wrong conceptually?

# **Topic 3**

# Inferences based on a single sample

In a few situations we can derive the sampling distribution for the statistic of interest and use this as the basis for constructing confidence intervals and hypothesis tests. Presently we estimate population means  $\mu$  in Section 3.1, population proportions p in Section 3.2, and population variances  $\sigma^2$  in Section 3.3 in some special cases.

#### 3.1 Estimating means

If the parameter of interest is the population mean  $\theta = \mu$ , then what can be said about the distribution of the sample mean estimator  $\hat{\theta} = \overline{X}$  in (2.1)? We will consider three cases,

- 1. normal population with known  $\sigma^2$ ,
- 2. any population with unknown  $\sigma^2$ , when the sample size m is large,
- 3. normal population with unknown  $\sigma^2$ , when the sample size m is small,

where the form of the confidence interval and hypothesis test statistic for  $\mu$  can be derived using the approximate normality of the sample mean.

In general, the confidence intervals for the mean based on normality theory will have the form:

point estimate 
$$\mu \pm$$
 (critical value of reference dist.) · (precision of point estimate), (3.1)

where the reference distribution will be the standard normal (for 1. and 2.) and the Student's t distribution (for 3.). The critical value corresponds to the value under the reference distribution that yields the two-sided (symmetric) tail areas summing to  $1 - \alpha$ .

#### 3.1.1 Mean of a normal population with known variance

When sampling from a normal population with known mean and variance, the estimator for the sample mean is also normal with mean  $\mu$  and variance  $\sigma^2/m$  where m is the sample size. Standardizing,

$$\frac{\overline{X} - \mu}{\sigma / \sqrt{m}} \sim N(0, 1) \tag{3.2}$$

we see that

$$P\left(-z_{\alpha/2} < \frac{\overline{X} - \mu}{\sigma/\sqrt{m}} < z_{\alpha/2}\right) = 1 - \alpha.$$

Based on knowing the sampling distribution of the estimator, we state the following CI.

**Definition 3.1.** A  $100(1-\alpha)\%$  **confidence interval** for the mean  $\mu$  of a normal population when the value of  $\sigma^2$  is known is given by

$$\left(\overline{x} - z_{\alpha/2} \cdot \frac{\sigma}{\sqrt{m}}, \overline{x} + z_{\alpha/2} \cdot \frac{\sigma}{\sqrt{m}}\right),\tag{3.3}$$

or  $\overline{x} \pm z_{\alpha/2} \cdot \sigma / \sqrt{m}$ , where m is the sample size.

The CI for the mean (3.3) can be expressed (cf. (3.1)) as

point estimate  $\mu \pm (z \text{ critical value}) \cdot (\text{standard error of mean})$ .

The z critical value is related to the tail areas under the standard normal curve; we need to find the z-score having a cumulative probability equal to  $1 - \alpha$  according to Definition 2.3.

**Example 3.1.** Consider 400 samples from a normal population with a known standard deviation  $\sigma = 17000$  with mean  $\overline{x} = 20992$  (as depicted in 3.1). How do we construct a 95% confidence interval for  $\mu$ ?

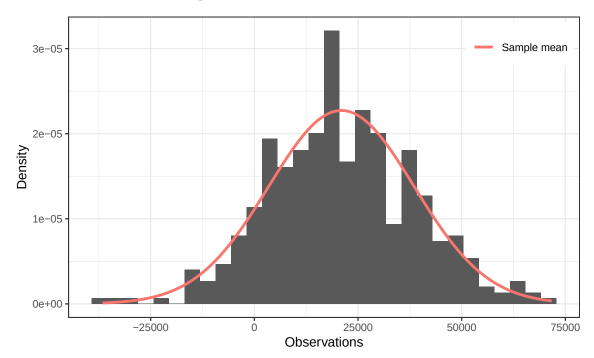


Figure 3.1: 400 samples from a normal population with known variance  $\sigma = 17000$  together with the corresponding (normal) sampling distribution for the observed mean.

For  $\alpha = 0.05$ , the critical value  $z_{0.025} = 1.96$ ; this value can be found by looking in a table of critical z values or using the r code qnorm(1-.05/2). From Definition 3.1,

$$\left(\overline{x} - z_{\alpha/2} \frac{\sigma}{\sqrt{m}}, \overline{x} + z_{\alpha/2} \frac{\sigma}{\sqrt{m}}\right) = \left(20992 - 1.96 \frac{17000}{\sqrt{400}}, 20992 + 1.96 \frac{17000}{\sqrt{400}}\right)$$
$$= (19326, 22658).$$

The data above was generated with a true population parameter  $\mu=21500$ , and, incidentally, the CI actually contains the parameter value.  $\Diamond$ 

As noted in (3.1) and (3.3), the width of a CI is related to the estimator's precision. The confidence level (or reliability) is inversely related to this precision. When the population is normal and the variance is known,

then an appealing strategy is to determine the sample size necessary to achieve a desired confidence level and precision. A general formula for the sample size  $m^*$  necessary to achieve an interval width w is obtained at confidence level  $\alpha$  is obtained by equating w to  $2z_{\alpha/2} \cdot \sigma/\sqrt{m^*}$  and then solving for  $m^*$ .

**Proposition 3.1.** The sample size n required to achieve a CI for  $\mu$  with width w at level  $\alpha$  is given by,

$$m^* = \left(2z_{\alpha/2} \cdot \frac{\sigma}{w}\right)^2.$$

From Proposition 3.1, we see that the smaller the desired w then the larger  $m^*$  must be (and subsequently, the more effort that must be allocated to data collection).

**Example 3.2.** In Example 3.1 we identified a 95% confidence interval for a normal population with known variance. The range (width) of that interval was 22658 - 19326 = 3332. By how much would m need to increase to halve the interval width?

Using Proposition 3.1,

$$m = \left(2 \cdot 1.96 \cdot \frac{17000}{1666}\right)^2 = (40)^2 = 1600.$$

Thus, we find that for the same level  $\alpha = 0.05$ , we would need to quadruple our original sample size to halve the interval. It is expensive to remove uncertainty!  $\Diamond$ 

Suppose now that we would like to consider a hypothesis test for the population mean, such as  $H_0$ :  $\mu = \mu_0$ . Starting from (3.2) and assuming that the null hypothesis is true, we find

$$Z = \frac{\overline{X} - \mu_0}{\sigma / \sqrt{m}} \,.$$

The statistic Z measures the distance (measured in units of  $\operatorname{sd}[\overline{X}]$ ) between  $\overline{X}$  and its expected value under the null hypothesis. We will use the statistic Z to determine if there is substantial evidence against  $H_0$  i.e. if the distance is too far in a direction consistent with  $H_a$ .

**Proposition 3.2.** Assume that we sample  $X_1, \ldots, X_m$  from a normal population with mean  $\mu$  and known variance  $\sigma^2$ 

Consider  $H_0$ :  $\mu = \mu_0$ . The test statistic is

$$Z = \frac{\overline{X} - \mu_0}{\sigma / \sqrt{m}} \,. \tag{3.4}$$

For a hypothesis test at level  $\alpha$ , we use the following procedure:

If  $H_a: \mu > \mu_0$ , then  $P = 1 - \Phi(z)$ , i.e., upper-tail  $R = \{z > z_a\}$ .

If  $H_a$ :  $\mu < \mu_0$ , then  $P = \Phi(z)$ , i.e., lower-tail  $R = \{z < -z_a\}$ .

If 
$$H_a: \mu \neq \mu_0$$
, then  $P = 2(1 - \Phi(|z|))$ , i.e., two-tailed  $R = \{|z| > z_{\alpha/2}\}$ .

We recall that  $\Phi(z)$  is the area in the lower-tail of the standard normal density, i.e., to the *left* of the calculated value of z. Thus  $1 - \Phi(z)$  is the area in the upper-tail and  $2(1 - \Phi(|z|))$  is twice the area capture in the upper-tail by |z| i.e. the sum of the area in the tails corresponding to  $\pm z$ . If  $P < \alpha$  then we reject  $H_0$  at level  $\alpha$  as the data provides sufficient evidence at the  $\alpha$  level against the null hypothesis.

**Example 3.3.** Let's return to the data in Example 3.1, where we sample from a normal population with a known standard deviation  $\sigma = 17000$ . Suppose that someone cliams the true mean is  $\mu_0 = 20000$ . Does our sample mean  $\overline{x} = 20992$  based on m = 400 samples provide evidence to contradict this claim at the  $\alpha = 0.05$  level?

The first thing to record is our parameter of interest:  $\mu$ , the true population mean. The null hypothesis, which we assume to be true, is a statement about the value of  $\mu$ ,

$$H_0$$
:  $\mu = 20000$ ,

and the alternative hypothesis is

$$H_a: \mu \neq 20000$$
,

since we are concerned with a deviation in either direction from  $\mu_0 = 20000$ .

Since the population is normal with known variance, we compute the test statistic:

$$z = \frac{\overline{x} - \mu_0}{\sigma / \sqrt{m}} = \frac{20992 - 20000}{17000 / \sqrt{400}} = 1.167.$$

That is, the observed sample mean  $\bar{x}$  is a little more than 1 standard deviation above what we would expect under  $H_0$ . Consulting 3.2, we see that a two-tailed test is indicated for this particular  $H_a$  (i.e., containing " $\neq$ "). The *P*-value is the area, <sup>1</sup>

$$P = 2(1 - \Phi(1.167)) = 2(0.1216052) = 0.2432.$$

Thus, since  $P = 0.2432 > 0.05 = \alpha$ , we fail to reject  $H_0$  at the level 0.05. The data does not support the claim that the true population mean differs from the value 20000 at the 0.05 level.  $\Diamond$ 

#### 3.1.2 Mean of a population with unknown variance (large-sample)

Consider samples  $X_1, \ldots, X_m$  from a population with mean  $\mu$  and variance  $\sigma^2$ . Provided that m is large enough, the Central Limit Theorem implies that the estimator for the sample mean  $\overline{X}$  in (2.1) has *approximately* a normal distribution. Then

$$P\left(-z_{\alpha/2} < \frac{\overline{X} - \mu}{\sigma/\sqrt{m}} < z_{\alpha/2}\right) \approx 1 - \alpha, \qquad (3.5)$$

since the transformed variable has approximately a standard normal distribution. Thus, computing a point estimate based on a large m of samples yields a CI for the population parameter  $\mu$  at an *approximate* confidence level  $\alpha$ . However, it is often the case that the variance is unknown. When m is large, replacing the population variance  $\sigma^2$  by the sample variance  $S^2$  in (2.2) will not typically introduce too much additional variability.

**Proposition 3.3.** For large sample size m, an approximate  $100(1 - \alpha)\%$  confidence interval for the mean  $\mu$  of any population when the variance is uknown is given by

$$\left(\overline{x} - z_{\alpha/2} \cdot \frac{s}{\sqrt{m}}, \overline{x} + z_{\alpha/2} \cdot \frac{s}{\sqrt{m}}\right),\tag{3.6}$$

or 
$$\overline{x} \pm z_{\alpha/2} \cdot s/\sqrt{m}$$
.

The CI for the mean (3.6) applies regardless of the shape of the population distribution so long as the number of samples is large. A rule of thumb is that m > 40 is sufficient.<sup>2</sup> In words, the CI (3.6) can be expressed (cf. (3.1)) as

point estimate  $\mu \pm (z \text{ critical value}) \cdot (\text{estimated standard error of mean})$ .

point estimate 
$$\pm 2 \text{ sd}$$

has 95% coverage and is surprisingly robust, i.e. applies to a wide variety of population distributions including the normal. However, this rule of thumb won't apply if you want to consider some different level, say 80% [van Belle, 2008, §1].

<sup>&</sup>lt;sup>1</sup>Note  $\Phi(z) = P(Z \le z)$  is found by calling pnorm(z) in r or by looking up the value in a Z table.

<sup>&</sup>lt;sup>2</sup>For m > 20, the interval estimate

Typically, a large-sample CI for a general parameter  $\theta$  holds that is similar in nature to (3.6) for any estimator  $\hat{\theta}$  that satisfies: (1) approximately normal in distribution, (2) approximately unbiased, and (3) an expression for the standard error is available.

To conduct a large-sample hypothesis test regarding the population mean  $\mu$ , we consider the test statistic

$$Z = \frac{\overline{X} - \mu_0}{S/\sqrt{m}}$$

under the null hypothesis, i.e., we replace the population standard deviation  $\sigma$  by the sample standard deviation S. When the number of samples m is large (say m > 40) then Z will be approximately normal. Substituting this test statistic Z for (3.4), we follow Proposition 3.2 to determine how to calculate the P-value.

**Example 3.4.** Consider the **Iris Data** from 2.1 and use the infer package to make inferences. In particular, consider whether the true mean petal length of Iris flowers exceeds 3.5 cm at the 0.05 level.

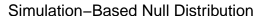
Recall that the **Iris Data** contains m=150 measurements of petal length across three species of Iris flowers and that the true variance is unknown. We are interested in testing the null hypothesis  $H_0: \mu \le 3.5$  against the alternative  $H_a: \mu > 3.5$  (i.e., a one-sided test).

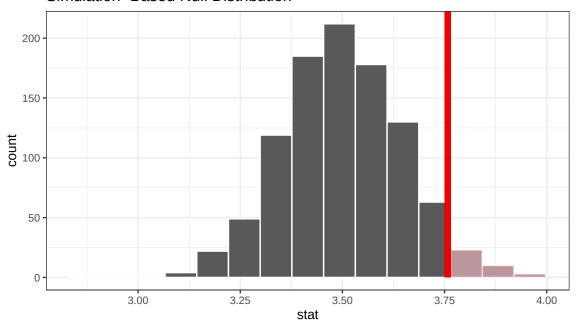
We begin by first computing the observed statistic (sample mean)  $\hat{\mu}$ . We use the infer package to construct a null distribution *computationally* for the response variable (petal length). We specifying that the hypothesis test is for parameter based on a point estimate and that we are testing for equality with the value  $\mu_0 = 3.5$ . The null distribution is generated by computing 1000 bootstrap replications of the sample mean, i.e., the sample mean is generated 1000 times by drawing 150 values at random with replacement from the original corpus of m = 150 samples. (Note that here, we obtain the null distribution computationally, so we do not need to standardize to Z.)

```
mu_hat <- mean(iris$Petal.Length)

null_dist <- iris %>%
    specify(response = Petal.Length) %>%
    hypothesise(null = "point", mu = 3.5) %>%
    generate(reps = 1000, type = "bootstrap") %>%
    calculate(stat = "mean")

null_dist %>%
    visualise() +
    shade_p_value(obs_stat = mu_hat, direction = "greater")
```





```
p_val <- null_dist %>%
  get_p_value(obs_stat = mu_hat, direction = "greater")
p_val
```

The bootstrapped null distribution is plotted using the visualize command and the regions of the null distribution that are as (or more) extreme than the observed statistic  $\hat{\mu}$  can be highlighted using the shade\_p\_value command. P=0.041 is found which is quite small; if  $\mu \leq 3.5$ , then the probability of obtaining the sample mean value  $\hat{\mu}=3.758$  is only 0.041! Thus, the data provides sufficient evidence at the 0.05 level against the hypothesis that the true mean petal length is at most 3.5 cm.  $\Diamond$ 

#### 3.1.3 Mean of a normal population with unknown variance

In Section 3.1.1, we considered samples  $X_1, \ldots, X_m$  from a normal population with a known  $\mu$  and  $\sigma^2$ . In contrast, here we consider samples from a normal population and assume the population parameters  $\mu$  and  $\sigma^2$  are unknown. If the number of samples is large, the discussion in Section 3.1.2 indicates that the rv  $Z = (\overline{X} - \mu)\sqrt{m}/S$  has approximately a standard normal distribution. However, if m is not sufficiently large<sup>3</sup> then the transformed variable will be more spread out than a standard normal distribution.

**Theorem 3.1.** For the sample mean  $\overline{X}$  based on m samples from a normal distribution with mean  $\mu$ , the rv

$$T = \frac{\overline{X} - \mu}{S/\sqrt{m}} \sim \mathsf{t}(m-1), \tag{3.7}$$

that is, T has Student's t distribution with v = m - 1 df.

This leads us to consider a CI for the population parameter  $\mu$  that is based on critical values of the t distribution.

 $<sup>^{3}</sup>$ Recall that we would consider m > 40 to be large.

**Proposition 3.4.** A  $100(1-\alpha)\%$  confidence interval for the mean  $\mu$  of a normal population, when  $\sigma^2$  is unknown, is given by

$$\left(\overline{x} - t_{\alpha/2, m-1} \cdot \frac{s}{\sqrt{m}}, \overline{x} + t_{\alpha/2, m-1} \cdot \frac{s}{\sqrt{m}}\right),\tag{3.8}$$

or  $\overline{x} \pm t_{\alpha/2,m-1} \cdot s/\sqrt{m}$ . Here  $\overline{x}$  and s are the sample mean and sample standard deviation, respectively.

**Example 3.5.** Let us return to the height of 31 felled black cherry trees from the **Cherry Tree Data** in Table 2.1. Give a 99% CI for the population mean  $\mu$ .

For m = 31, the critical value of the reference distribution is  $t_{0.005,30} \approx 2.7499$ , which can looked up in a table of critical values for t(v = m - 1) or found using the r command qt(1-0.01/2, df = 31-1). The sample mean  $\overline{x} = 76$  (computed in Example 2.1) is combined with the sample standard deviation,

$$s = \sqrt{\frac{1}{m-1} \sum_{i=1}^{m} (x_i - \overline{x})^2}$$
$$= \sqrt{\frac{1}{30} \left( (63 - 76)^2 + \dots + (87 - 76)^2 \right)}$$
$$= 6.372.$$

to form the interval estimate

$$\left(\overline{x} - t_{\alpha/2, m-1} \cdot \frac{s}{\sqrt{m}}, \overline{x} + t_{\alpha/2, m-1} \cdot \frac{s}{\sqrt{m}}\right)$$

$$= \left(76 - 2.750 \cdot \frac{6.372}{\sqrt{31}}, 76 + 2.750 \cdot \frac{6.372}{\sqrt{31}}\right)$$

$$= (72.85, 79.15).$$

For comparison, the critical value  $t_{.01/2,\nu}$  for  $\nu = 13, ..., 30$ 

```
qt(1-0.01/2, df = seq(12:39))
```

[1] 63.656741 9.924843 5.840909 4.604095 4.032143 3.707428 3.499483 3.355387 [9] 3.249836 3.169273 3.105807 3.054540 3.012276 2.976843 2.946713 2.920782 [17] 2.898231 2.878440 2.860935 2.845340 2.831360 2.818756 2.807336 2.796940 [25] 2.787436 2.778715 2.770683 2.763262

can deviate significantly from the corresponding  $z_{0.01/2} = 2.575829$ . In particular, if we had erroneously used the large sample estimate (3.6) then we would have obtained 99% CI (73.05, 78.95) which might give us a false sense of security as it is narrower.  $\Diamond$ 

In contrast to Proposition 3.1, it is difficult to select the sample size m to control the width of the t-based CI as the width involves the unknown (before the sample is acquired) s and because m also enters through  $t_{\alpha/2,m-1}$ . A one-sample t test based on (3.7) can be used to test hypothesis about the population mean when the population is normal and  $\sigma^2$  is unknown.

**Proposition 3.5.** Assume that we sample  $X_1, \ldots, X_m$  from a normal population with mean  $\mu$  and unknown variance  $\sigma^2$ .

Consider  $H_0$ :  $\mu = \mu_0$ . The test statistic is

$$T = \frac{\overline{X} - \mu_0}{S/\sqrt{m}} \,.$$

For a hypothesis test at level  $\alpha$ , we use the following procedure:

If  $H_a: \mu > \mu_0$ , then P-value is the area under t(m-1) to the right of t.

If  $H_a: \mu < \mu_0$ , then P-value is the area under t(m-1) to the left of t.

If  $H_a: \mu \neq \mu_0$ , then P-value is twice the area under t(m-1) to the right of |t|.

**Example 3.6.** From the **Cherry Tree Data**, let's look at the average volume of timber, given in Table 3.1. The distribution for this data is approximately normal.<sup>4</sup> We might ask if the data provides compelling evidence, say at level 0.05, for concluding that the true average volume of timber exceeds 21.3 cubic feet.<sup>5</sup>

Table 3.1: Observations of m = 31 felled black cherry trees.

Volume [cu ft]	
10.2, 10.3, 10.3, 15.6, 16.4, 18.2, 18.8, 19.1, 19.7, 19.9, 21.0, 21.3, 25.7, 27.4, 31.7, 33.8, 34.5, 36.3, 38.3, 42.6, 51.0, 51.5, 55.4, 55.7	

Let's carry out a significance test for the true average volume of timber  $\mu$  at level  $\alpha = 0.05$ . We assume the null hypothesis

$$H_0: \mu = 21.3$$
.

An appropriate null hypothesis is

$$H_a: \mu > 21.3$$
,

that is, we will adopt the stance that the true average exceeds  $\mu_0 = 21.3$  only if the null is rejected.

From our m=31 samples, we find that  $\overline{x}=30.17$  and that s=16.44. The computed value of the one-sample t-statistic is given by

$$t = \frac{\overline{x} - \mu_0}{s/\sqrt{m}}$$

$$= \frac{30.17 - 21.3}{16.44/\sqrt{31}}$$

$$= 3.$$
(3.9)

The test is based on v = 31 - 1 df, and P = 0.002663. This is the upper-tail area, i.e. the area to the right of t (see Figure 3.2).

Since  $P \ll \alpha$ , we reject the null hypothesis that the population mean is 21.3. The data provides sufficient evidence that the population mean is different from 21.3.  $\Diamond$ 

#### 3.2 Estimating proportions

Consider a population of size M in which a proportion p of the population satisfies a given property. The  $p \in (0, 1)$  is a parameter characterizing the population, with distribution F(p), that we might be interested

<sup>&</sup>lt;sup>4</sup>After looking at the normal quantile-quantile plot, I decided to test a hypothesis. For level 0.01, I ran a Kolmogorov–Smirnov test for the null hypothesis that the data is consistent with  $N(\overline{x}, s^2)$  vs the alternative that the data is not consistent with the specified reference distribution. The *P*-value attained was P = 0.2532 > 0.10, and therefore I fail to reject the null hypothesis. The data is consistent with being drawn from a normal population.

<sup>&</sup>lt;sup>5</sup>How much wood is that? About a sixth of a cord. A full cord of chopped fire wood in the US is 124 cu ft; about enough to keep you warm through a New England winter (according to my inlaws).

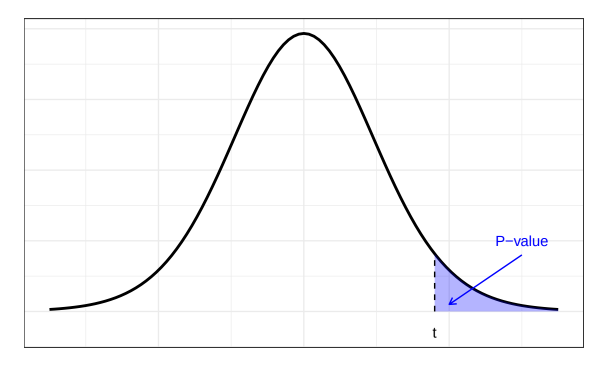


Figure 3.2: For this test, the *P*-value is the upper-tail area, i.e., to the right of the computed statistic t.

in estimating. A sample,  $X_1, \dots, X_m \sim F(p)$ , from the population contains a proportion,

$$\hat{p} = \frac{1}{m} \sum_{i=1}^{m} X_i \,, \tag{3.10}$$

satisfying the given property. The estimator  $\hat{p}$  varies with the sample and for large m it's sampling distribution has the following properties:

$$\mu_{\hat{p}} = \mathbf{E}[X_i] = p$$

and

$$\sigma_{\hat{p}}^2 = \frac{\text{Var}[X_i]}{m} = \frac{p(1-p)}{m},$$
(3.11)

provided that m is small relative to M (a rule of thumb is  $m \le 0.05M$ ). Moreover, by invoking the Central Limit Theorem we have the distribution of  $\hat{p}$  is approximately normal for sufficiently large m as (3.10) is a sample mean. Indeed, this normal approximation works well for moderately large m as long as p is not too close to zero or one; a rule of thumb is that mp > 5 and m(1-p) > 5.

**Proposition 3.6.** For large samples n, a  $100(1-\alpha)\%$  confidence interval for the parameter p is given by

$$\hat{p} \pm z_{\alpha/2} \sqrt{\frac{\hat{p}(1-\hat{p})}{m}}$$
 (3.12)

This follows from Proposition 3.3 by observing that (3.10) is a sample mean and replacing the standard

$$\sigma_{\hat{p}}^2 = \frac{p(1-p)}{m} \frac{M-m}{M-1} \,,$$

where for fixed *m* the factor converges to 1 as  $M \to \infty$ .

<sup>&</sup>lt;sup>6</sup>Here we write F for a general distribution, but what special distribution might this be?

<sup>&</sup>lt;sup>7</sup>Note that if m is large relative to M (m > 0.05M) then the variance (3.11) must be adjusted by a factor (related to the hypergeometric distribution):

error  $\sigma_{\hat{p}}$  from (3.11) by the estimated standard error,

$$\widehat{\mathsf{se}}(\widehat{p}) = \sqrt{\frac{\widehat{p}(1-\widehat{p})}{m}}\,;$$

recall the s in (3.6) is the sample variance for the *population* and  $s/\sqrt{m} = se$  is the standard error of the point estimator.

**Proposition 3.7.** Let X be the count of members with a given property based on a sample of size m from a population where a proportion p share the property. Then  $\hat{p} = X/m$  is an estimator of p. Assume  $mp_0 \ge 10$  and  $m(1 - p_0) \ge 10$ .

Consider  $H_0$ :  $p = p_0$ . The test statistic is

$$Z = \frac{\hat{p} - p_0}{\sqrt{p_0(1 - p_0)/m}} \,.$$

For a hypothesis test at level  $\alpha$ , we use the following procedure:

If  $H_a: p > p_0$ , then P-value is the area under N(0, 1) to the right of z.

If  $H_a$ :  $p < p_0$ , then P-value is the area under N(0, 1) to the left of z.

If  $H_a: p \neq p_0$ , then P-value is twice the area under N(0,1) to the right of |t|.

**Example 3.7.** Let us revisit Example 2.2, where we were considering Churchill's claim that he will receive half the votes for the House of Commons seat for the constituency of Dundee. We are skeptical that he is as popular as he says. Suppose 116 out of 263 Dundonians polled claimed that they intended to vote for Churchill. Can it be concluded at significance level 0.10 that more than half of all eligible Dundonains will vote for Churchill?

The parameter of interest is p, the proportion of votes for Churchill. The null hypothesis is  $H_0$ : p=0.5. The alternative hypothesis is  $H_a$ : p<0.5, since we . Since 263(0.5)=131.5>10, we satisfy the assumptions stated in Proposition 3.7.

Based on the sample,  $\hat{p} = 116/263 = 0.4411$ . The test statistic value is

$$z = \frac{\hat{p} - p_0}{\sqrt{p_0(1 - p_0)/m}}$$
$$= \frac{0.4411 - 0.5}{\sqrt{0.5(1 - 0.5)/263}}$$
$$= -1.91.$$

The *P*-value for this lower-tailed *z* test is  $P = \Phi(-1.91) = 0.028$ . Since  $P < 0.10 = \alpha$ , we reject the null hypothesis at the 0.05 level. The evidence for concluding that the true proportion is different from  $p_0 = 0.5$  at the 0.10 level is compelling.<sup>8</sup>  $\Diamond$ 

# 3.3 Estimating variances

Next we consider estimates of the population variance (and standard deviation) when the population is assumed to have a normal distribution. In this case, the sample variance  $S^2$  in (2.2) provides the basis

<sup>&</sup>lt;sup>8</sup>Churchill took ca. 44% of the vote in the 1908 by-election to become MP for Dundee [https://www.wikiwand.com/en/19 08\_Dundee\_by-election].

for inferences. Consider iid samples  $X_1, \dots, X_m \sim N(\mu, \sigma^2)$ . We provide the following theorem without proof.

**Theorem 3.2.** For the sample variance  $S^2$  based on m samples from a normal distribution with variance  $\sigma^2$ , the rv

$$V = \frac{(m-1)S^2}{\sigma^2} = \frac{\sum_i (X_i - \overline{X})^2}{\sigma^2} \sim \chi_{m-1}^2,$$

that is, V has a  $\chi^2$  distribution with  $\nu = m - 1$  df.

Based on Theorem 3.2,

$$P\left(\chi_{1-\alpha/2,m-1}^2 < \frac{(m-1)S^2}{\sigma^2} < \chi_{\alpha/2,m-1}^2\right) = 1 - \alpha,$$

i.e., the area captured between the right and left tail critical  $\chi^2$  values is  $1 - \alpha$ . The expression above can be further manipulated to obtain an interval for the unknown parameter  $\sigma^2$ :

$$P\left(\frac{(m-1)s^2}{\chi^2_{\alpha/2,m-1}} < \sigma^2 < \frac{(m-1)s^2}{\chi^2_{1-\alpha/2,m-1}}\right) = 1 - \alpha,$$

where we substitute the computed value of the point estimate  $s^2$  for the estimator into the limits to give a CI for  $\sigma^2$ . If we take square roots of the inequality above, we obtain a CI for the population standard deviation  $\sigma$ .

**Proposition 3.8.** A  $100(1-\alpha)\%$  confidence interval for the variance of a normal population is

$$\left((m-1)s^2/\chi^2_{\alpha/2,m-1}\,,(m-1)s^2/\chi^2_{1-\alpha/2,m-1}\right)\,.$$

A  $100(1-\alpha)\%$  confidence interval for the standard deviation  $\sigma$  of a normal population is given by taking the square roots of the lower and upper limits in (3.8).

**Example 3.8.** For the **Cherry Tree Data** in Table 3.1 concerning the timber volume of 31 felled black cherry trees, give a 95 CI for the variance.

We are interested in estimating the true variance  $\sigma^2$  of the volume of timber based on m=31 samples. Recall that the mean of our data is  $\overline{x}=30.17$  cu ft and that the sample variance is  $s^2=270.2$  using the estimator (2.2). The critical values for the  $\chi^2_{.975,30}=16.7908$  and  $\chi^2.025,30=46.9792$  can be found by checking a table of critical values of the  $\chi^2(v=30)$  distribution or by using the r code qchisq(1-0.05/2, df=30, lower.tail = FALSE) and qchisq(0.05/2, df=df, lower.tail = FALSE), respectively (see 3.3).

Pulling everything together, a 95% CI for the population variance is given by

$$\left( (m-1)s^2/\chi_{\alpha/2,m-1}^2, (m-1)s^2/\chi_{1-\alpha/2,m-1}^2 \right) 
= ((30)270.2/46.9792, (30)270.2/16.7908) 
= (172.5, 482.8) .$$

Note the position of the critical values—don't swap them round. ◊

**Example 3.9.** Revisit Example 3.8 and using the infer package to construct a 95% confidence interval for the true standard deviation of the timber volume of black cherry trees based on the available mesurements in the **Cherry Tree Data**, Table 3.1.

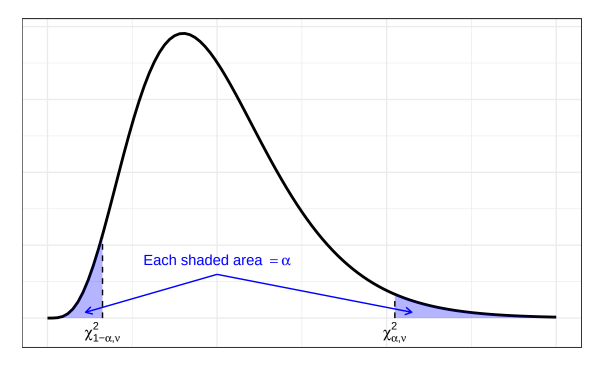


Figure 3.3: As the  $\chi^2$  distribution is not symmetric, the upper and lower critical values will not be the same (the shaded areas are equal).

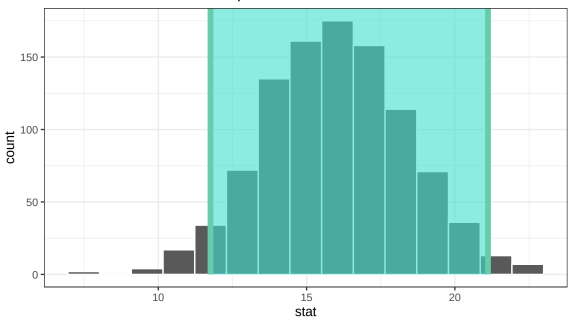
```
s <- sd(trees$Volume)

null_dist <- trees %>%
    specify(response = Volume) %>%
    generate(reps = 1000, type = "bootstrap") %>%
    calculate(stat = "sd")

ci <- null_dist %>%
    get_confidence_interval(point_estimate = s, level = 0.95, type = "se")

null_dist %>%
    visualise() + shade_ci(ci)
```

## Simulation-Based Bootstrap Distribution



We plot the 95% confidence interval for the standard deviation based on the computational null distribution obtained using 1000 bootstrap replications; note the interval estimate

ci<sup>2</sup>

lower\_ci upper\_ci 1 138.2487 445.9603

is in good agreement with the values obtained Example 3.8. Due to the computational nature, the bootstrapped interval estimate is not precisely the same as the theoretical interval estimate and moreover rerunning the code will yield a slightly different interval.  $\Diamond$ 

# **Topic 4**

# Inferences based on two samples

We consider inferences—estimators, confidence intervals, and hypothesis testing—for comparing means, proportions, and variances based on two independent samples from different populations, respectively, in Sections 4.1, 4.3, 4.4. We also consider inferences when the samples are not independent, so-called paired samples, in Section 4.2.

## 4.1 Comparing means

Let us assume that we have two normal populations with iid samples

$$X_1, \dots, X_m \sim \mathsf{N}(\mu_X, \sigma_X^2)$$

and

$$Y_1, \dots, Y_n \sim \mathsf{N}(\mu_Y, \sigma_Y^2)$$

and, moreover, that the X and Y samples are independent of one another. When comparing the means of two populations, the quantity of interest is the difference:  $\mu_X - \mu_Y$ .

**Proposition 4.1.** If we consider the sample means  $\overline{X}$  and  $\overline{Y}$ , then the mean of the variable  $\overline{X} - \overline{Y}$  is,

$$\mu_{\overline{X}-\overline{Y}} = \mathbf{E}\left[\overline{X} - \overline{Y}\right] = \mu_X - \mu_Y,$$

and the variance is,

$$\sigma_{\overline{X}-\overline{Y}}^2 = \operatorname{Var}\left[\overline{X} - \overline{Y}\right] = \frac{\sigma_X^2}{m} + \frac{\sigma_Y^2}{n}$$
.

Proposition 4.1 follows directly from the definition of the sample mean in (2.1) and properties of expectation and variance. If our parameter of interest is

$$\theta = \mu_1 - \mu_2 \,,$$

then its estimator,

$$\hat{\theta} = \overline{X} - \overline{Y}$$

is normally distributed with mean and variance given by Proposition 4.1. If the samples sizes m and n are large, then the estimator is approximately normally distributed by the Central Limit Theorem regardless of the population. We now discuss CIs and hypothesis tests for comparing population means  $\theta = \mu_X - \mu_Y$ . We consider three cases when comparing means:

- 1. normal populations when the variances  $\sigma_X^2$  and  $\sigma_Y^2$  are known,
  2. any populations with unknown variances  $\sigma_X^2$  and  $\sigma_Y^2$ , when the sample sizes m and n are large,
  3. normal populations when the variances  $\sigma_X^2$  and  $\sigma_Y^2$  are unknown, when the sample sizes m and nare small,

noting that the development largely reflects that of Section 3.1.

### Comparing means of normal populations when variances are known

When  $\sigma_X^2$  and  $\sigma_Y^2$  are known, standardizing  $\overline{X}-\overline{Y}$  yields the standard normal variable:

$$Z = \frac{\overline{X} - \overline{Y} - (\mu_X - \mu_Y)}{\sqrt{\frac{\sigma_X^2}{m} + \frac{\sigma_Y^2}{n}}} \sim N(0, 1).$$
(4.1)

Inferences proceed by treating the parameter of interest  $\theta$  as in the single sample case using the test statistic (4.1).

**Proposition 4.2.** A 100(1 –  $\alpha$ )% CI for the parameter  $\theta = \mu_X - \mu_Y$  based on samples of size m from a normal population  $N(\mu_X, \sigma_X^2)$  and of size n from  $N(\mu_Y, \sigma_Y^2)$  with known variances, is given by

$$(\overline{x} + \overline{y}) \pm z_{\alpha/2} \cdot \sqrt{\frac{\sigma_X^2}{m} + \frac{\sigma_Y^2}{n}}$$
.

**Proposition 4.3.** Assume that we sample iid  $X_1, \dots, X_m \sim \mathsf{N}(\mu_X, \sigma_X^2)$  and iid  $Y_1, \dots, Y_n \sim \mathsf{N}(\mu_Y, \sigma_Y^2)$  and that the X and Y samples are independent.

Consider  $H_0: \mu_X - \mu_Y = \theta_0$ . The test statistic is

$$Z = \frac{\overline{X} - \overline{Y} - \theta_0}{\sqrt{\frac{\sigma_X^2}{m} + \frac{\sigma_Y^2}{n}}}.$$
 (4.2)

For a hypothesis test at level  $\alpha$ , we use the following procedure:

If  $H_a: \mu_X - \mu_Y > \theta_0$ , then  $P = 1 - \Phi(z)$ , i.e., upper-tail  $R = \{z > z_\alpha\}$ .

If  $H_a$ :  $\mu_X - \mu_Y < \theta_0$ , then  $P = \Phi(z)$ , i.e., lower-tail  $R = \{z < -z_\alpha\}$ .

If  $H_a: \mu_X - \mu_Y \neq \theta_0$ , then  $P = 2(1 - \Phi(|z|))$ , i.e., two-tailed  $R = \{|z| > z_{\alpha/2}\}$ .

#### Comparing means when the sample sizes are large

When the samples are large, then the assumptions about normality of the populations and knowledge of the variances  $\sigma_X^2$  and  $\sigma_Y^2$  can be relaxed. For sufficiently large m and n, the difference of the sample means,  $\overline{X} - \overline{Y}$ , has approximately a normal distribution for any underlying population distributions by the Central Limit Theorem. Moreover, if m and n are large enough, then replacing the population variances with the sample variances  $S_X^2$  and  $S_Y^2$  will not increase the variability of the estimator or the test statistic too much.

**Proposition 4.4.** For m and n sufficiently large, an approximate  $100(1-\alpha)\%$  CI for  $\mu_X - \mu_Y$  for two samples from populations with any underlying distribution is given by

$$(\overline{x} + \overline{y}) \pm z_{\alpha/2} \cdot \sqrt{\frac{s_X^2}{m} + \frac{s_Y^2}{n}}$$

**Proposition 4.5.** Under the same assumptions and procedures as in Proposition 4.3, a large-sample, i.e., m > 40 and n > 40, test statistic,

$$Z = \frac{\overline{X} - \overline{Y} - \theta_0}{\sqrt{\frac{S_X^2}{m} + \frac{S_Y^2}{n}}},$$

can be used in place of (4.2) for hypothesis testing.

# 4.1.3 Comparing means of normal populations when variances are unknown and the sample size is small

If  $\sigma_X$  and  $\sigma_Y$  are unknown and either sample is small (e.g., m < 30 or n < 30), but both populations are normally distributed, then we can use Student's t distribution to make inferences. We provide the following theorem without proof.

**Theorem 4.1.** When both population distributions are normal, the standardized variable

$$T = \frac{\overline{X} - \overline{Y} - (\mu_X - \mu_Y)}{\sqrt{\frac{S_X^2}{m} + \frac{S_Y^2}{n}}} \sim t(v)$$

where the df  $\nu$  is estimated from the data. Namely,  $\nu$  is given by (round  $\nu$  down to the nearest integer):

$$v = \frac{\left(\frac{s_X^2}{m} + \frac{s_Y^2}{n}\right)^2}{\frac{(s_X^2/m)^2}{m-1} + \frac{(s_Y^2/n)^2}{n-1}} = \frac{\left(\frac{s_X^2 + s_Y^2}{Y}\right)^2}{\frac{s_X^4}{m-1} + \frac{s_Y^4}{n-1}}$$
(4.3)

where  $s_X^2$  and  $s_Y^2$  are point estimators of the sample variances; alternatively, we see that the formula (4.3) can also be written in terms of the standard error of the sample means:

$$s_{\overline{X}} = \frac{s_X}{\sqrt{m}}$$
 and  $s_{\overline{Y}} = \frac{s_Y}{\sqrt{n}}$ .

The formula (4.3) for the data-driven choice of  $\nu$  calls for the computation of the standard error of the sample means.

**Proposition 4.6.** A  $100(1-\alpha)\%$  CI for  $\mu_X - \mu_Y$  for two samples of size m and n from normal populations where the variances are unknown is given by

$$(\overline{x} - \overline{y}) \pm t_{\alpha/2,\nu} \sqrt{\frac{s_X^2}{m} + \frac{s_Y^2}{n}},$$

where we recall that  $t_{\alpha/2,\nu}$  is the  $\alpha/2$  critical value of  $t(\nu)$  with  $\nu$  given by (4.3).

**Proposition 4.7.** Assume that we sample iid  $X_1, ..., X_m$  and iid  $Y_1, ..., Y_n$  from normal populations with with unknown variances and means  $\mu_X$  and  $\mu_Y$ , respectively, and that the X and Y samples are independent.

Consider  $H_0: \mu_X - \mu_Y = \theta_0$ . The test statistic is

$$T = \frac{\overline{X} - \overline{Y} - \theta_0}{\sqrt{\frac{S_X^2}{m} + \frac{S_Y^2}{n}}}.$$
 (4.4)

For a hypothesis test at level  $\alpha$ , we use the following procedure:

If  $H_a: \mu_X - \mu_Y > \theta_0$ , then P-value is the area under t(v) to the right of t, i.e., upper-tail  $R = \{t > t_{\alpha,v}\}$ .

If  $H_a: \mu_X - \mu_Y < \theta_0$ , then P-value is the area under t(v) to the left of t, i.e., lower-tail  $R = \{t < -t_{\alpha,v}\}$ .

If  $H_a: \mu_X - \mu_Y \neq \theta_0$ , then P-value is twice the area under t(v) to the right of |t|, i.e., two-tailed  $R = \{|t| > t_{\alpha/2...}\}.$ 

Here v is given by (4.3).

If the variances of the normal populations are unknown but are the same,  $\sigma_X^2 = \sigma_Y^2$ , then deriving CIs and test statistics for comparing the means can be simplified by considering a combined or pooled estimator for the single parameter  $\sigma^2$ . If we have two samples from populations with variance  $\sigma^2$ , then each sample provides an estimate for  $\sigma^2$ . That is,  $S_X^2$ , based on the *m* observations of the first sample, is one estimator for  $\sigma^2$  and another is given by  $S_Y^2$ , based on *n* observations of the second sample. The correct way to combine these two estimators into a single estimator for the sample variance is to consider the **pooled estimator** of  $\sigma^2$ ,

$$S_{p}^{2} = \frac{m-1}{m+n-2} S_{X}^{2} + \frac{n-1}{m+n-2} S_{Y}^{2}.$$
 (4.5)

The pooled estimator is a weighted average that adjusts for differences between the sample sizes m and n.

**Proposition 4.8.** A  $100(1-\alpha)\%$  CI for  $\mu_X - \mu_Y$  for two samples of size m and n from normal populations where the variance  $\sigma^2$  is unknown is given by

$$(\overline{x} - \overline{y}) \pm t_{\alpha/2, m+n-2} \cdot \sqrt{s_p^2 \left(\frac{1}{m} + \frac{1}{n}\right)},$$

where we recall that  $t_{\alpha/2,m+n-2}$  is the  $\alpha/2$  critical value of the t(v) with v=m+n-2 df.

Similarly, one can consider a pooled t test, i.e., a hypothesis test based on the pooled estimator for the variance as opposed to the two-sample t test in Proposition 4.7. In the case of a pooled t test, the test statistic

$$T = \frac{\overline{X} - \overline{Y} - \theta_0}{\sqrt{S_p^2 \left(\frac{1}{m} + \frac{1}{n}\right)}},$$

with the pooled estimator of the variance, replaces (4.4) in Proposition 4.7 and the same procedures are followed for determining the *P*-value with v = m + n - 2 in place of (4.3). If you have reasons to believe that  $\sigma_X^2 = \sigma_Y^2$ , these pooled t procedures are appealing because v is very easy to compute.



Pooled t procedures are not robust if the assumption of equalized variance is violated. Theoretically, you could first carry out a statistical test  $H_0$ :  $\sigma_X^2 = \sigma_Y^2$  on the equality of variances and then use a pooled t procedure if the null hypothesis is not rejected. However, there is no free lunch: the typical F test for equal variances (see Section 4.4) is sensitive to normality assumptions. The two sample t procedures, with the data-driven choice of v in (4.3), are therefore recommended unless, of course, you have a very compelling reason to believe  $\sigma_X^2 = \sigma_Y^2$ .

<sup>&</sup>lt;sup>1</sup>If  $m \neq n$ , then the estimator with *more* samples will contain *more* information about the parameter  $\sigma^2$ . Thus, the simple average  $(S_X^2 + S_Y^2)/2$  wouldn't really be fair, would it?

## 4.2 Comparing paired samples

The preceding analysis for comparing population means was based on the assumption that a random sample  $X_1, \ldots, X_m$  is drawn from a distribution with mean  $\mu_X$  and that a completely independent random sample  $Y_1, \ldots, Y_n$  is drawn from a distribution with mean  $\mu_Y$ . Some situations, e.g., comparing observations before and after a treatment or exposure, necessitate the consideration of paired values.

Consider a random sample of iid pairs

$$(X_1, Y_1), \ldots, (X_n, Y_n)$$

with  $\mathbf{E}[X_i] = \mu_X$  and  $\mathbf{E}[Y_i] = \mu_Y$ . If we are interested in making inferences about the difference  $\mu_X - \mu_Y$  then the paired differences

$$D_i = X_i - Y_i \,, \quad i = 1, \dots, n \,,$$

constitute a sample with mean  $\mu_D = \mu_X - \mu_Y$  that can be treated using single-sample CIs and tests, e.g., see Section 3.1.3.

# 4.3 Comparing proportions

Consider a population containing a proportion  $p_X$  of individuals satisfying a given property. For a sample of size m from this population, we denote the sample proportion by  $\hat{p}_X$ . Likewise, we consider a population containing a proportion  $p_Y$  of individuals satisfying the same given property. For a sample of size n from this population, we denote the sample proportion by  $\hat{p}_Y$ . We assume the samples from the X and Y populations are independent. The natural estimator for the difference in population proportions  $p_X - p_Y$  is the difference in the sample proportions  $\hat{p}_X - \hat{p}_Y$ .

Provided the samples are much smaller than the population sizes (i.e., the populations are about 20 times larger than the samples),

$$\mu_{(\hat{p}_X-\hat{p}_Y)} = \mathbf{E}[\hat{p}_X - \hat{p}_Y] = p_X - p_Y \,,$$

and

$$\sigma_{(\hat{p}_X - \hat{p}_Y)}^2 = \text{Var}[\hat{p}_X - \hat{p}_Y] = \frac{p_X(1 - p_X)}{m} + \frac{p_Y(1 - p_Y)}{n} \,,$$

by considering the fact that the count of individuals satisfying the given property in each population will be independent draws from  $Binom(m, p_X)$  and  $Binom(n, p_Y)$ , respectively. Further, if m and n are large (e.g.,  $m \ge 30$  and  $n \ge 30$ ), then  $\hat{p}_X$  and  $\hat{p}_Y$  are (approximately) normally distributed. Standardizing  $\hat{p}_X - \hat{p}_Y$ ,

$$Z = \frac{\hat{p}_X - \hat{p}_Y - (p_X - p_Y)}{\sqrt{\frac{p_X(1 - p_X)}{m} + \frac{p_Y(1 - p_Y)}{n}}} \sim \mathsf{N}(0, 1) \,.$$

A CI for  $\hat{p}_X - \hat{p}_Y$  then follows from the large-sample CI considered in Section 3.1.2.

**Proposition 4.9.** An approximate  $100(1-\alpha)\%$  CI for  $p_X - p_Y$  is given by

$$\hat{p}_{X} - \hat{p}_{Y} \pm z_{\alpha/2} \sqrt{\frac{\hat{p}_{X}(1 - \hat{p}_{X})}{m} + \frac{\hat{p}_{Y}(1 - \hat{p}_{Y})}{n}} \,,$$

and, as a rule of thumb, can be reliably used if  $m\hat{p}_X$ ,  $m(1-\hat{p}_X)$ ,  $n\hat{p}_Y$ , and  $n(1-\hat{p}_Y)$  are greater than or equal to 10.

Proposition 4.9 does not pool the estimators for the population proportions. However, if we are considering a hypothesis test concerning the equality of the population proportions with the null hypothesis

$$H_0: p_X - p_Y = 0$$
,

then we assume  $p_X = p_Y$  as our default position. Therefore, as a matter of consistency, we should replace the standard error in (4.9) with a pooled estimator for the standard error of the population proportion,

$$\hat{p} = \frac{m}{m+n} \hat{p}_X + \frac{n}{m+n} \hat{p}_Y \,.$$

**Proposition 4.10.** Assume that  $m\hat{p}_X$ ,  $m(1-\hat{p}_X)$ ,  $n\hat{p}_Y$ ,  $n(1-\hat{p}_Y)$  are all greater than 10.

Consider  $H_0: p_X - p_Y = 0$ . The test statistic is

$$Z = \frac{\hat{p}_X - \hat{p}_Y}{\sqrt{\hat{p}(1-\hat{p})\left(\frac{1}{m} + \frac{1}{n}\right)}}.$$

For a hypothesis test at level  $\alpha$ , we use the following procedure:

If  $H_a: p_X - p_Y > 0$ , then  $P = 1 - \Phi(z)$ , i.e., upper-tail  $R = \{z > z_\alpha\}$ .

If  $H_a: p_X - p_Y < 0$ , then  $P = \Phi(z)$ , i.e., lower-tail  $R = \{z < -z_\alpha\}$ .

If  $H_a: p_X - p_Y \neq 0$ , then  $P = 2(1 - \Phi(|z|))$ , i.e., two-tailed  $R = \{|z| > z_{\alpha/2}\}$ .

# 4.4 Comparing variances

For a random sample

$$X_1, \dots, X_m \sim \mathsf{N}(\mu_X, \sigma_X^2)$$

and an independent random sample

$$Y_1, \ldots, Y_n \sim \mathsf{N}(\mu_Y, \sigma_Y^2),$$

the rv

$$F = \frac{S_X^2 / \sigma_X^2}{S_Y^2 / \sigma_Y^2} \sim F(m - 1, n - 1), \tag{4.6}$$

that is, F has an F distribution with df  $v_1 = m - 1$  and  $v_2 = n - 1$ . The statistic F in (4.6) comprises the *ratio* of variances  $\sigma_X^2/\sigma_Y^2$  and not the difference; therefore, the plausibility of  $\sigma_X^2 = \sigma_Y^2$  will be based on how much the ratio differs from 1.

**Proposition 4.11.** For the null hypothesis  $H_0: \sigma_X^2 = \sigma_Y^2$ , the test statistic to consider is:

$$f = \frac{s_X^2}{s_Y^2}$$

and the P-values are determined by the F(m-1, n-1) curve where m and n are the respective sample sizes.

A  $100(1-\alpha)\%$  CI for the ratio  $\sigma_X^2/\sigma_Y^2$  is based on forming the probability,

$$P(F_{1-\alpha/2,\nu_1,\nu_2} < F < F_{\alpha/2,\nu_1,\nu_2}) = 1 - \alpha$$

where  $F_{\alpha/2,\nu_1,\nu_2}$  is the  $\alpha/2$  critical value from the  $F(\nu_1 = m - 1, \nu_2 = n - 1)$  distribution. Substituting (4.6) with point estimates for F and manipulating the inequalities it is possible to isolate the ratio  $\sigma_X^2/\sigma_Y^2$ ,

$$P\left(\frac{1}{F_{\alpha/2,\nu_1,\nu_2}}\frac{s_X^2}{s_V^2} < \frac{\sigma_X^2}{\sigma_V^2} < \frac{1}{F_{1-\alpha/2,\nu_1,\nu_2}}\frac{s_X^2}{s_V^2}\right) = 1 - \alpha.$$

**Proposition 4.12.** A 100(1 –  $\alpha$ )% CI for the ratio of population variances  $\sigma_X^2/\sigma_Y^2$  is given by

$$\left(F_{\alpha/2,m-1,n-1}^{-1}s_X^2/s_Y^2\,,\,F_{1-\alpha/2,m-1,n-1}^{-1}s_X^2/s_Y^2\right)\,.$$

**Proposition 4.13.** Assume the population distributions are normal and the random samples are both independent of one another.

Consider  $H_0: \sigma_X^2 = \sigma_Y^2$ . The test statistic is

$$F = S_Y^2 / S_Y^2.$$

For a hypothesis test at level  $\alpha$ , we use the following procedure:

If  $H_a:\sigma_X^2>\sigma_Y^2$ , then P-value is  $A_R=$  area under the F(m-1,n-1) curve to the right of f.

If  $H_a:\sigma_X^2<\sigma_Y^2$ , then P-value is  $A_L=$  area under the  $\mathsf{F}(m-1,n-1)$  curve to the left of f.

If  $H_a: \sigma_X^2 \neq \sigma_Y^2$ , then P-value is  $2 \cdot \min(A_R, A_L)$ .

# **Topic 5**

# **Analysis of variance (ANOVA)**

Analysis of variance, shortened as ANOVA, is a collection of statistical models and estimation procedures for analyzing the variation among different groups. In particular, a single-factor ANOVA provides a hypothesis test regarding the equality of two or more population means, thereby generalizing the one-sample and two-sample t tests considered in Sections 3.1.3 and 4.1.3.

## 5.1 Single factor ANOVA test

Suppose that we have k normally distributed populations<sup>1</sup> with different means  $\mu_1, \ldots, \mu_k$  and equal variances  $\sigma^2$ . We denote the rv for the jth measurement taken from the ith population by  $X_{ij}$  and the corresponding sample observation by  $x_{ij}$ . For samples of size  $m_1, \ldots, m_k$ , we denote the sample means

$$\overline{X}_i = \frac{1}{m_i} \sum_{j=1}^{m_i} X_{ij} \,,$$

and sample variances

$$S_i^2 = \frac{1}{m_i - 1} \sum_{i=1}^{m_i} (X_{ij} - \overline{X}_i)^2,$$

for each i = 1, ..., k; likewise, we denote the associated point estimates for the sample means  $\overline{x}_1, ..., \overline{x}_k$  and the sample variances  $s_1^2, ..., s_k^2$ . The average over all observations  $m = \sum m_i$ , called the **grand mean**, is denoted by

$$\overline{X} = \frac{1}{m} \sum_{i=1}^{k} \sum_{j=1}^{m_i} X_{ij}.$$

The sample variances  $s_i^2$ , and hence the sample standard deviations, will generally vary even when the k populations share the same variance; a rule of thumb is that the equality of variances is reasonable if the largest  $s_i$  is not much more than two times the smallest.

We wish to test the equality of the population means, given by the null hypothesis,

$$H_0: \mu_1 = \mu_2 = \dots = \mu_k$$

versus the alternative hypothesis,

$$H_a$$
: at least two  $\mu_i$  differ.

 $<sup>^{1}</sup>$ In the context of ANOVA, these k populations are often referred to as **treatment distributions**.

Note that if k=3 then  $H_0$  is true only if all three means are the same, i.e.,  $\mu_1=\mu_1=\mu_3$ , but there are a number of ways which the alternative might hold:  $\mu_1\neq\mu_2=\mu_3$  or  $\mu_1=\mu_2\neq\mu_3$  or  $\mu_1=\mu_3\neq\mu_2$  or  $\mu_1\neq\mu_2\neq\mu_3$ .

The test procedure is based on comparing a measure of difference in variation among the sample means, i.e., the variation between  $x_i$ 's, to a measure of variation within each sample.

#### **Definition 5.1.** The mean square for treatments is

$$\mathsf{MSTr} = \frac{1}{k-1} \sum_{i=1}^k m_i (\overline{X}_i - \overline{X})^2 \,,$$

and the mean square error is

MSE = 
$$\frac{1}{m-k} \sum_{i=1}^{k} (m_i - 1) S_i^2$$
.

The MSTr and MSE are statistics that measure, respectively, the variation among sample means and the variation within samples. We will also use MSTr and MSE to denote calculated values of these statistics.

#### **Proposition 5.1.** The test statistic

$$F = \frac{MSTr}{MSF}$$

is the appropriate test statistic for the single-factor ANOVA problem involving k populations (or treatments) with a random sample of size  $m_1, \ldots, m_k$  from each. When  $H_0$  is true,

$$F \sim F(v_1 = k - 1, v_2 = m - k)$$
.

In the present context a large test statistic value is more constradictory to  $H_0$  than a smaller value, therefore the test is upper-tailed, i.e., consider the area  $F_{\alpha}$  to the right of the critical value  $F_{\alpha,\nu_1,\nu_2}$ . We reject  $H_0$  if the value of the test statistic  $F > F_{\alpha}$ .

**Example 5.1.** Consider the average salary data from lcocal councils in Table 5.1. Is the expected average salary in each nation the same at the 5% level?

Table 5.1: \*\*Average Salary Data\*\* reported from 20 local councils.

Nation	Average salaries ('000 £)	Size $(m_i)$	Sample Mean $(\overline{x}_i)$	Sample SD $(s_i)$
England	17, 12, 18, 13, 15, 12	6	14.5	2.588
N Ireland	11, 7, 9, 13	4	10.0	2.582
Scotland	15, 10, 13, 14, 13	5	13.0	1.871
Wales	10, 12, 8, 7, 9	5	9.2	1.924



Table 5.1 presents the **Average Salary Data** (in thousands of pounds) reported from 20 local councils classified by nation (England, N Ireland, Scotland, and Wales). The sample means together with the sample standard deviations are summarized in the table as well as presented using using box plots in Figure 5.1.

For  $\alpha = 0.05$ , we compute the upper-tail area  $F_{0.05}$  i.e. to the right of the critical value  $F_{0.05,3,16}$  by consulting a statistical table or by using r to find  $F_{0.05} = 3.2388715$ .

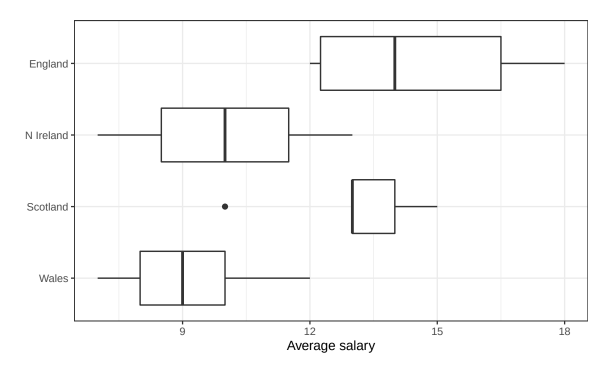


Figure 5.1: Box plots of the average mean salary data in Table 5.1 indicate five summary statistics: the median, two hinges (first and third quartiles) and two whiskers (extending from the hinge to the most extreme data point within  $1.5 \cdot IQR$ ).

[1] 3.238872

The grand mean is

$$\overline{x} = \frac{17 + 12 + 18 + \dots + 8 + 7 + 9}{20} = 11.9,$$

and hence the variation among sample means is given by,

$$\begin{aligned} \mathsf{MSTr} &= \frac{1}{4-1} \left( m_1 (\overline{x}_1 - \overline{x})^2 + \dots + m_4 (\overline{x}_4 - \overline{x})^2 \right) \\ &= \left( 6(14.5 - 11.9)^2 + 4(10.0 - 11.9)^2 + 5(13.0 - 11.9)^2 + 5(9.2 - 11.9)^2 \right) / 3 \\ &= 32.5 \ . \end{aligned}$$

The mean square error is

$$\begin{aligned} \mathsf{MSE} &= \frac{1}{20-4} \left( (m_1 - 1) s_1^2 + \cdots (m_4 - 1) s_4^2 \right) \\ &= \frac{5(2.588)^2 + 3(2.582)^2 + 4(1.871)^2 + 4(1.924)^2}{16} \\ &= 5.14366 \end{aligned}$$

yielding the test statistic value

$$F = \frac{\text{MSTr}}{\text{MSE}} = \frac{32.5}{5.14366} = 6.3184581.$$

Since  $F > F_{\alpha}$  we reject  $H_0$ . The data does not support they hypothesis that the mean salaries in each nation are identical at the 5% level.  $\Diamond$ 

### **5.2** Confidence intervals

In Section 4.1 we gave a CI for comparing population means that involved the difference  $\mu_X - \mu_Y$ . In some settings, we would like to give CIs for more complicated functions of population means  $\mu_i$ . Let

$$\theta = \sum_{i=1}^k c_i \mu_i \,,$$

for constants  $c_i$ . As we assume the  $X_i j$  are normally distributed with  $\mathbf{E}[X_{ij}] = \mu_i$  and  $\mathrm{Var}[X_{ij}] = \sigma^2$ , the estimator

$$\hat{\theta} = \sum_{i=1}^{k} c_i \overline{X}_i,$$

is normally distributed with

$$\operatorname{Var}[\hat{\theta}] - \sum_{i=1}^{k} c_i^2 \operatorname{Var}[\overline{X}_i] = \sigma^2 \sum_{i=1}^{k} \frac{c_i}{m_i}.$$

Estimating  $\sigma^2$  by the MSE and forming  $\hat{\sigma}_{\hat{\theta}}$  results in a t variable

$$\frac{\hat{\theta}-\theta}{\hat{\sigma}_{\hat{\theta}}}$$
.

**Proposition 5.2.** A  $100(1-\alpha)\%$  CI for  $\sum c_i \mu_i$  is given by

$$\sum_{i=1}^k c_i \overline{x}_i \pm t_{\alpha/2,m-k} \sqrt{\text{MSE} \sum_{i=1}^k \frac{c_i}{m_i}}.$$

**Example 5.2.** Determine a 90% CI for the difference in mean average salary for councils in Scotland and England, based on the data available in Table 5.1

For  $\alpha = 0.10$ , the critical value  $t_{0.05,16} = 1.7458837$  is found by looking in a table of t critical values or by using r:

```
# alt: qt(0.1/2, 16, lower.tail = FALSE)
qt(1-0.1/2, df = 20 - 4)
```

[1] 1.745884

Then for the function  $\overline{x}_2 - \overline{x_1}$ ,

$$(\overline{x}_{Eng} - \overline{x}_{Sco}) \pm t_{0.05,16} \sqrt{\text{MSE}} \sqrt{\frac{1}{m_{Eng}} + \frac{1}{m_{Sco}}}$$

$$= (14.5 - 13.0) \pm 1.7458837 \sqrt{5.14366} \sqrt{\frac{1}{6} + \frac{1}{5}}$$

$$= 1.5 \pm 2.3976575.$$

Thus a 90% confidence interval for  $\mu_{Eng} - \mu_{Sco}$  is (-0.8977, 3.898).  $\Diamond$ 



How does the result in Example 5.2 compare to the t method in Section 4.1.3?

# **Topic 6**

# Linear regression

**Regression analysis** allows us to study the relationship among two or more rvs. Typically, we are interested in the relationship between a **response** or **dependent** rv Y and a **covariate** X. The relationship between X and Y will be explained through a **regression function**,

$$r(x) = \mathbf{E}[Y \mid X = x] = \int y f(y \mid x) dy.$$

In particular, we shall assume that r is linear,

$$r(x) = \beta_0 + \beta_1 x, \qquad (6.1)$$

and estimate the intercept  $\beta_0$  and slope  $\beta_1$  of this linear model from sample data

$$(Y_1, X_1), \ldots, (Y_m, X_m) \sim F_{Y,X}$$
.

## 6.1 Simple linear regression models

The simplest regression is when  $X_i$  is one-dimensional and r(x) is linear as in (6.1). A linear regression posits the expected value of  $Y_i$  is a linear function of the data  $X_i$ , but that Y deviates from its expected value by a random amount for fixed  $x_i$ .

**Definition 6.1.** The **simple linear regression model** relates a random response  $Y_i$  to a set of independent variables  $X_i$ ,

$$Y_i = \beta_0 + \beta_1 X_i + \epsilon_i \,, \tag{6.2}$$

where the intercept  $\beta_0$  and slope  $\beta_1$  are unknown parameters and the **random deviation** or **random error**  $\epsilon_i$  is a rv assumed to satisfy:

- 1.  $\mathbf{E}[\epsilon_i \mid X_i = x_i] = 0$ ,
- 2.  $Var[\epsilon_i \mid X_i = x_i] = \sigma^2$  does not depend on  $x_i$ ,
- 3.  $\epsilon_i$  and  $\epsilon_j$  are independent for i, j = 1, ..., m.

From the assumptions on  $\epsilon_i$ , the linear model (6.2) implies

$$\mathbf{E}[Y_i \mid X_i = x_i] = \beta_0 + \beta_1 x_i$$
.

 $<sup>^{1}</sup>$ The covariates X are also called **predictor variables**, **explanatory variables**, **independent variables**, and/or **features** depending on who you are talking to.

Thus, if  $\hat{\beta}_0$  and  $\hat{\beta}_1$  are estimators of  $\beta_0$  and  $\beta_1$ , then the **fitted line** is

$$\hat{r}(x) = \hat{\beta}_0 + \hat{\beta}_1 x$$

and the **predicted** or **fitted value**  $\hat{Y}_i = \hat{r}(X_i)$  is an estimator for  $\mathbf{E}[Y_i \mid X_i = x_i]$ . The **residuals** are defined to be

$$\hat{\epsilon}_i = Y_i - \hat{Y}_i = Y_i - (\hat{\beta}_0 + \hat{\beta}_1 X_i) . \tag{6.3}$$

The residual sums of squares,<sup>2</sup>

$$RSS = \sum_{i=1}^{m} \hat{\epsilon}_i^2, \qquad (6.4)$$

measures how well the regression line  $\hat{r}$  fits the data  $(Y_1, X_1), \dots, (Y_m, X_m)$ . The **least squares estimates** of  $\hat{\beta}_0$  and  $\hat{\beta}_1$  are the values that minimize the RSS in (6.4).

**Theorem 6.1.** The least squares estimates for  $\hat{\beta}_1$  and  $\hat{\beta}_0$  are given by, respectively,

$$\hat{\beta}_1 = \frac{\sum_{i=1}^m (X_i - \overline{X})(Y_i - \overline{Y})}{\sum_{i=1}^m (X_i - \overline{X})^2} = \frac{S_{xy}}{S_{xx}},$$
(6.5)

and

$$\hat{\beta}_0 = \overline{Y} - \hat{\beta}_1 \overline{X} \,. \tag{6.6}$$

Equation (6.4) is a function of  $\hat{\beta}_0$  and  $\hat{\beta}_1$  from the definition of the residuals (6.3). Then (6.5) and (6.6) follow by equating the partial derivatives of (6.4) to zero. The  $\hat{\beta}_0$  and  $\hat{\beta}_1$  are the unique solution to this linear system.

**Example 6.1.** In Figures 6.1 and 6.2, we consider the **Cherry Tree Data** (see Table 2.1 and discussion). We fit a least squares regression of the volume of timber (response variable) to the diameter of the tree (independent variable). As you would expect, the timber yield increases with diameter.

The r code below can be used to calculate the least squares regression and residuals.

```
data(trees)
y <- trees$Volume
x <- trees$Girth # NB: this is diameter; data mislabeled!
fit <- lm(y ~ x)
e <- resid(fit)
yhat <- predict(fit)</pre>
```

The fit data frame contains the estimates for  $\hat{\beta}_0$  and  $\hat{\beta}_1$ :

#### fit\$coefficients

```
(Intercept) x -36.943459 5.065856
```

Both Figures 6.1 and 6.2 are scatter plots of the observed values y. In Figure 6.1, the regression line  $\hat{y}$  is plotted along with the residuals  $\hat{\epsilon}$ . In Figure 6.2, the sample mean  $\overline{y}$  is plotted together with the deviations  $y - \overline{y}$ .  $\diamondsuit$ 

<sup>&</sup>lt;sup>2</sup>The RSS is sometimes referred to as the **error sum of squares** and abbreviated SSE (no, the order is not a typo a typo).

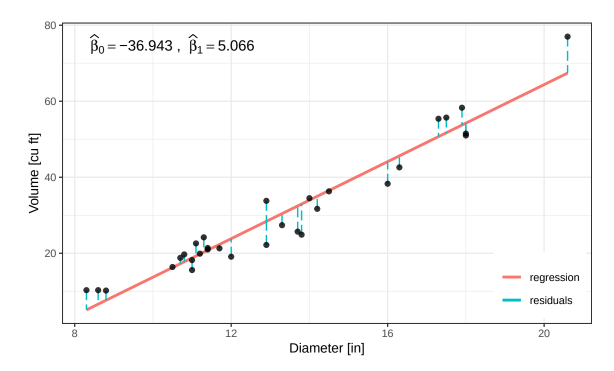


Figure 6.1: Linear regression (or least squares fit) of Volume to Diameter from the **Cherry Tree Data**. The vertical bars between the observed data point and the regression line indicate the error in the fit (the least squares residual). The residuals are squared and summed to yield the RSS (alt: SSE).

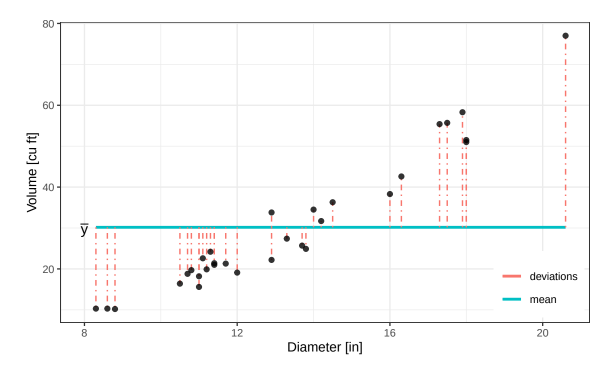


Figure 6.2: The deviations about the sample mean  $\overline{y}$ . The sum of the squared deviations or SST (total sum of squares) is a measure of the total variation in the observations.

# **6.2** Estimating $\sigma^2$ for linear regressions

The parameter  $\sigma^2$  (the variance of the random deviation) determines the variability in the regression model.

**Theorem 6.2.** An unbiased estimate of  $\sigma^2$  is given by

$$\hat{\sigma}^2 = s^2 = \frac{\text{RSS}}{m-2} = \frac{1}{m-2} \sum_{i=1}^m (y_i - \hat{y}_i)^2.$$
 (6.7)

In Figure 6.3, we present a least squares regression of timber volume on both tree diameter and height (for the **Cherry Tree Data**). As expected, the regressions indicate the volume increases with both covariates. Estimates for the variance of the random deviation (6.7) in both regression models,  $\sigma_D^2$  and  $\sigma_H^2$ , respectively, are computed to be  $s_D^2 = 18.08$  and  $s_H^2 = 179.48$ . Thus, we see that small variances lead to observations of  $(x_i, y_i)$  that sit tightly around the regression line, in contrast to large variances that lead to a large cloud of points.

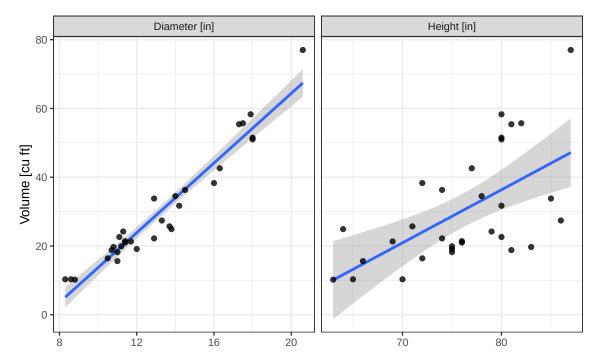


Figure 6.3: For the **Cherry Tree Data**, we estimate the variance to be  $s_D^2 = 18.08$  (for Diameter) and  $s_H^2 = 179.48$  (for Height); small variances lead to observations of  $(x_i, y_i)$  that sit tightly around the regression line, in contrast to large variances that lead to a large cloud of points.



In Theorem 6.2, the number in the denominator is the df associated with the RSS and  $s^2$ . To calculate RSS you must estimate two parameters  $\beta_0$  and  $\beta_1$  which results in the loss of two df. Hence the m-2.

We note for the purposes of making inferences, the statistic

$$S^2 = \frac{\text{RSS}}{m-2}$$

is an unbiased estimator or  $\sigma^2$  and the random variable

$$\frac{(m-2)S^2}{\sigma^2} \sim \chi^2(m-2).$$

Moreover, the statistic  $S^2$  is independent of both  $\hat{\beta}_0$  and  $\hat{\beta}_1$ .

#### 6.3 **Inferences for least-squares parameters**

If  $\epsilon_i$  in (6.2) is assumed to be normally distributed, then we can derive the sampling distributions of the estimators  $\hat{\beta}_0$  and  $\hat{\beta}_1$ . Hence, we can use these sampling distributions to make inferences about the parameters  $\beta_0$  and  $\beta_1$ .

Provided iid  $\epsilon_i \mid X_i \sim N(0, \sigma^2)$ , the least-squares estimators possess the following properties.

- 1. Both  $\hat{\beta}_0$  and  $\hat{\beta}_1$  are normally distributed.
- 2. Both  $\hat{\beta_0}$  and  $\hat{\beta_1}$  are unbiased, i.e.,  $\mathbf{E}[\hat{\beta_i}] = \beta_i$  for i = 0, 1. 3.  $\mathrm{Var}[\hat{\beta_0}] = c_{00}\sigma^2$  where  $c_{00} = \sum_{i=1}^m x_i^2/(mS_{xx})$ .
- 4.  $Var[\hat{\beta}_1] = c_{11}\sigma^2$  where  $c_{11} = 1/S_{xx}$ .
- 5.  $\text{Cov}[\hat{\beta}_0, \hat{\beta}_1] = c_{01}\sigma^2 \text{ where } c_{01} = -\overline{x}/S_{xx}.$

These properties can be determined by working directly from (6.5) and (6.6).

**Proposition 6.1.** Consider  $H_0: \beta_i = \beta_{i0}$ . The test statistic is

$$T = \frac{\hat{\beta}_i - \beta_{i0}}{S\sqrt{c_{ii}}} \, .$$

For a hypothesis test at level  $\alpha$ , we use the following procedure:

If  $H_a: \beta_i > \beta_{i0}$ , then P-value is the area under t(m-2) to the right of t.

If  $H_a: \beta_i < \beta_{i0}$ , tthen P-value is the area under t(m-2) to the left of t.

If  $H_a: \beta_i \neq \beta_{i0}$ , then P-value is twice the area under t(m-2) to the right of |t|.

A confidence interval for  $\beta_i$ , based on the statistic (6.1), can be given following the procedures in 3.

**Proposition 6.2.** A  $100(1 - \alpha)\%$  CI for  $\beta_i$  is given by

$$\hat{\beta}_i \pm t_{\alpha/2, m-2} S \sqrt{c_{ii}} .$$

#### 6.4 Correlation

Let  $(X_1, Y_1), \dots, (X_m, Y_m)$  denote a random sample from a bivariate normal distribution with  $\mathbf{E}[X_i] = \mu_X$ ,  $\mathbf{E}[Y_i] = \mu_Y$ ,  $\mathrm{Var}[X_i] = \sigma_X^2$ ,  $\mathrm{Var}[Y_i] = \sigma_Y^2$ , and correlation coefficient  $\rho$ . The sample correlation coefficient is given by,

$$r = \frac{\sum_{i=1}^{m} (X_i - \overline{X})(Y_i - \overline{Y})}{\sqrt{\sum_{i=1}^{m} (X_i - \overline{X})^2 \sum_{i=1}^{m} (Y_i - \overline{Y})^2}},$$
(6.8)

which can be rewritten in terms of  $S_{xx}$ ,  $S_{xy}$ , and  $S_{yy}$ :

$$r = \frac{S_{xy}}{\sqrt{S_{xx}S_{yy}}} = \hat{\beta}_1 \sqrt{\frac{S_{xx}}{S_{yy}}},$$

using (6.5) and we see that r and  $\hat{\beta}_1$  have the same sign. A |r| close to 1 means that the regression line is a good fit to the data and, similarly, an |r| close to 0 means a poor fit to the data. Note that the correlation coefficient (and the least squares regression) are only suitable for describing *linear* relationships; a nonlinear relationship can also yield r near zero (see Figure 6.4).

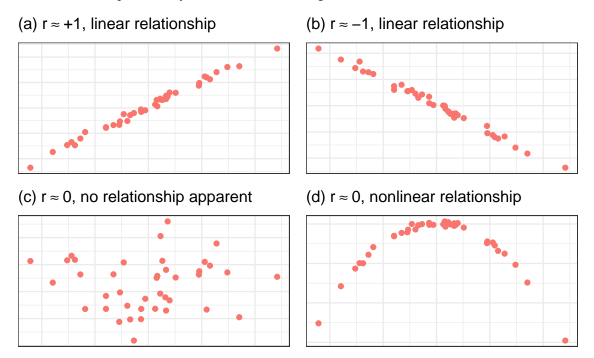


Figure 6.4: Correlations range from -1 to 1 with |r| = 1 indicating a strong linear relationship and r near zero indicating the absence of a linear relationship.

Once a model is fit, it can be used to predict a value of y for a given x. However, the model only gives the most likely value of y; a corresponding **prediction interval** is usually more appropriate.

**Proposition 6.3.** A 100(1 –  $\alpha$ )% prediction interval for an actual value of Y when  $x = x^*$  is given by

$$(\hat{\beta}_0 + \hat{\beta}_1 x^*) \pm t_{\alpha/2, m-2} S \sqrt{1 + \frac{1}{n} + \frac{(x^* - \overline{x})^2}{S_{xx}}}$$
.



The prediction interval is not the same as the confidence interval for expected Y. Note that the length of the *confidence interval* for  $\mathbf{E}[Y]$  when  $x = x^*$  is given by

$$2 \cdot t_{\alpha/2} S \sqrt{\frac{1}{n} + \frac{(x^* - \overline{x})^2}{S_{xx}}}$$

whereas the length for the prediction interval of Y is

$$2 \cdot t_{\alpha/2} S \sqrt{1 + \frac{1}{n} + \frac{(x^* - \overline{x})^2}{S_{xx}}}$$
.

Thus the prediction intervals for an actual value of Y are longer than the confidence intervals for  $\mathbf{E}[Y]$  if both are determined for the same value  $x^*$ .

The linear model

$$\mathbf{E}[Y\mid X=x]=\beta_0+\beta_1 x\,,$$

assumes that the conditional expectation of Y for a fixed value of X is a linear function of the x value. If we assume that (X,Y) has a bivariate normal distribution, then

$$\beta_1 = \frac{\sigma_Y}{\sigma_X} \rho \,,$$

and thus, for the simple hypothesis tests we have considered (Table 2.2), statistical tests for  $\beta$  and  $\rho$  are equivalent.

# **Topic 7**

# Categorical data

# 7.1 Multinomial experiments

Suppose we have a population that is divided into k > 2 distinct categories. We consider an experiment where we select m individuals (or objects) from the population and categorize each one. We denote the proportion of the population in the ith category by  $p_i$ . If the sample size m is much smaller than the population size M (so that the m trials are independent), this experiment will be approximately multinomial with success probability  $p_i$  for each category, i = 1, ..., k.

Before the experiment is performed, we denote the number (or count) of the trials resulting in category i by the rv  $N_i$ . The expected number of trails that result in category i is given by

$$\mathbf{E}[N_i] = mp_i \,, \quad i = 1, \dots, k \,. \tag{7.1}$$

After the experiment is performed, we denote the corresponding observed value by  $n_i$ . Since the trials result in distinct categories,

$$\sum_{i=1}^{k} N_i = \sum_{i=1}^{k} n_i = m,$$

which indicates that, for a given m, we only need to observe k-1 of the variables to be able to work out what the kth variable should be.

# 7.2 Goodness-of-fit for a single factor

We are interested in making inferences about the proportion parameters  $p_i$ . Specifically, we will consider the null hypothesis,

$$H_0: p_1 = p_{10}, p_2 = p_{20}, \cdots, p_k = p_{k0},$$
 (7.2)

that completely specifies a value  $p_{i0}$  for each  $p_i$ . The alternative hypothesis  $H_a$  will state that  $H_0$  is not true, i.e., that at least one  $p_i$  is different from the value  $p_{i0}$  claimed under the null  $H_0$ .

Provided the null hypothesis in (7.2) is true, the expected values (7.1) can be written in terms of the expected frequencies,

$$\mathbf{E}[N_i] = mp_{i0}, \quad i = 1, \dots, k.$$

Often the  $n_i$ , referred to as the **observed cell counts**, and the corresponding  $mp_{i0}$ , referred to as the **expected cell counts**, are tabulated, for example, as in Table 7.1.

<sup>&</sup>lt;sup>1</sup>Here for i = 1, ..., k we use the notation  $p_{i0}$  to denote the value of  $p_i$  claimed under the null hypothesis.

Table 7.1: Observed and expected cell counts.

Category	i = 1	i = 2	•••	i = k	Row total
Observed	$n_1$	$n_2$	•••	$n_k$	m
Expected	$mp_{10}$	$mp_{20}$	•••	$mp_{k0}$	m

The test procedure assesses the discrepancy between the value of the observed and expected cell counts. This discrepancy, or **goodness of fit**, is measured by the squared deviations divided by the expected count.<sup>2</sup>

**Theorem 7.1.** For  $mp_i \ge 5$  for i = 1, ..., k, the rv

$$V = \sum_{i=1}^{k} \frac{(N_i - mp_i)^2}{mp_i} \sim \chi^2(k-1),$$

that is, V has approximately a  $\chi^2$  distribution with v = k - 1 df.

**Proposition 7.1.** Consider the null

$$H_0: p_1 = p_{10}, p_2 = p_{20}, \cdots, p_k = p_{k0},$$

and the alternative

 $H_a: p_i \neq p_{i0}$  for at least one i.

The test statistic is

$$V = \sum_{i=1}^k \frac{(N_i - mp_{i0})^2}{mp_{i0}} \,.$$

As a rule of thumb, provided  $mp_{i0} \ge 5$  for all i = 1, ..., k, then the P-value is the area under  $\chi^2(k-1)$  to the right of v.

If  $mp_{i0} < 5$  for some *i* then it may be possible to combine the categories such that the new categorizations satisfy the assumptions of Proposition 7.1.



Things are much more complicated if the category probabilities are not completely specified.

# 7.3 Test for independence of factors

In Section 7.2 we considered the categorization of a population into a single factor. We now consider a single population where each individual is categorized into two factors with I distinct categories for the first factor and J distinct categories for the second factor. That is, each individual from the population belongs to exactly one of the I categories of the first factor and exactly one of the J categories of the second factor. We want to determine whether or not there is any dependency between the two factors.

For a sample of m individuals, we denote by  $n_{ij}$  the count of the m samples that fall both in category i of the first factor and category j of the second factor, for i = 1, ..., I and j = 1, ..., J. A **contingency (data)** 

<sup>&</sup>lt;sup>2</sup>The division by the expected cell counts is to account for possible differences in the relative magnitude of the observed/expected counts.

**table** with I rows and J columns (i.e., IJ cells) will be used to record the  $n_{ij}$  counts (in the obvious way).<sup>3</sup> Let  $p_{ij}$  be the proportion of individuals in the population who belong in category i of factor 1 and category j of factor 2. Then, the probability that a randomly selected individual falls in category i of factor 1 is found by summing over all j:

$$p_i = \sum_{i=1}^J p_{ij} \,,$$

and likewise, the probability that a randomly selected individual falls in category j of factor 2 is found by summing over all i:

$$p_j = \sum_{i=1}^I p_{ij} .$$

The null hypothesis that we will be interested in adopting is

$$H_0: p_{ij} = p_i \cdot p_j \ \forall (i,j),$$

that is, that an individual's category in factor 1 is independent of the category in factor 2.

Following the same program as for the single category goodness-of-fit test, we note that, assuming the null hypothesis (7.3) is true, then the expected count in cell i, j is

$$\mathbf{E}[N_{ii}] = mp_{ii} = mp_ip_i;$$

and we estimate  $p_i$  and  $p_j$  by the appropriate sample proportion:

$$\hat{p}_i = \frac{n_i}{m}$$
,  $n_i = \sum_i n_{ij}$  (row totals),

and

$$\hat{p}_j = \frac{n_j}{m}$$
,  $n_j = \sum_i n_{ij}$  (column totals).

Thus, the expected cell count is given by

$$\hat{e}_{ij} = m\hat{p}_i\hat{p}_j = \frac{n_i n_j}{m} \,,$$

and we assess the goodness of fit between the observed cell count  $n_i j$  and the expected cell count  $\hat{e}_i j$ .

**Proposition 7.2.** Assume the null hypothesis

$$H_0: p_{ij} = p_i p_j \text{ for all } i = 1, ..., I, j = 1, ..., J,$$

against the alternative hypothesis

$$H_a: H_0$$
 is not true.

The test statistic is

$$V = \sum_{i=1}^{I} \sum_{j=1}^{J} \frac{(N_{ij} - \hat{e}_{ij})^2}{\hat{e}_{ij}}.$$

As a rule of thumb, provided  $\hat{e}_{ij} \geq 5$  for all i, j and when  $H_0$  is true, then the test statistic has approximately a  $\chi^2(v)$  distribution with v = (I-1)(J-1) df. For a hypothesis test at level  $\alpha$ , the procedure is upper-tailed and the P-value is the area under  $\chi^2(v)$  to the right of v.

<sup>&</sup>lt;sup>3</sup>Contingency is another word for dependency in this context.

# **Topic 8**

# **Quality Control**

Quality control is an area of applies statistics that seeks to make interventions to maintain or improve the outcome of industrial processes. Random variations in output processes might negatively impact the quality of a product. We would like to identify the sources of random variations in output processes that might have *assignable causes*. **Control charts** are a tool that help us to recognize when industrial processes are no longer controlled so that one might then seek to identify assignable causes.

The basic elements of control charting involve specifying a control region and then analyzing time-series data. We will specify a base-line value along with an upper and lower limit of control and assume that a process is under control unless a test statistic suggests otherwise.<sup>1</sup> To construct a control chart, one collects data about a process at fixed points of time and calculates the running value of quality statistic. If the quality statistic exceeds the upper or lower control limits, the process is deemed to out of control and the quality of the product is assumed to be negatively impacted.

The process of creating a control chart is best illustrated through an extended example, like Example 8.1 provided below.

**Example 8.1.** Here we consider the typical  $3\sigma$  control charting for a process mean  $\overline{X}$  based on estimated parameters. That is, we assume the generating process X is normally distributed with unknown parameters  $\mu$  and  $\sigma^2$ . We seek to estimate the mean  $\overline{X}$ . Our control region is specified to be three standard deviations; that is, the process is in-control if it remains within three standard devaitions of a baseline value.

The **Beer Data** contains measurements of the features OG, ABV, pH, and IBU for 50 batches of each of three types of product (Premium Lager, IPA, and Light Lager). We are interested the IPA's pH value, which influences saccharification. We assume that 3 batches of IPA are produced per day are we prepare the data as follows.

```
ipa <- beer %>%
select(Batch_Id, pH, Beer) %>%
filter(Beer == "IPA") %>%
rename(Day = Batch_Id)
ipa$Day[1:48] <- rep(1:16, each = 3)
ipa <- ipa[1:48,]
m <- 3  # three batches per day
k <- 16  # number of days</pre>
```

<sup>&</sup>lt;sup>1</sup>The default position here will be reminiscent of hypothesis testing.

4.0

3.5

We first observe that the pH measurements are (at least approximately) normal, as can be seen from the quantile-quantile plot in Figure 8.1.

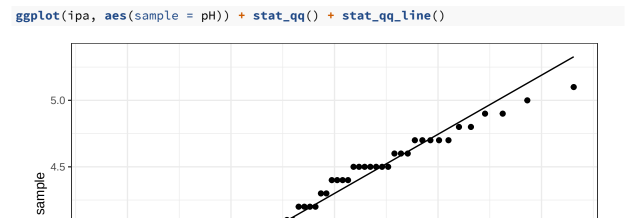


Figure 8.1: Normal quantile-quantile plot of observed pH measurements of the IPA batches in Table 8.1.

theoretical

We consider the data for pH readings from three batches of IPA taken over sixteen days (k = 16) presented in Table 8.1. The Table includes the sample mean per day  $\bar{x}$ , the sample standard deviation s, and the range of values max  $x_i - \min x_i$  per day (each based on m = 3 batches).

We estimate of the mean

$$\widehat{\mu} = \frac{1}{k} \sum_{i=1}^{k} \overline{x}_i \,,$$

by averaging the means found for the k days and, similarly, estimate the mean of the sample standard deviation,

$$\overline{s} = \frac{1}{k} \sum_{i=1}^{k} s_i \,,$$

Day	pH Observations	$\overline{x}$	S	Range
1	4.7, 4.5, 4.9	4.700	0.20000	0.4
2	4.0, 4.6, 4.5	4.367	0.32150	0.6
3	4.7, 3.3, 4.6	4.200	0.78100	1.4
4	3.9, 3.5, 4.2	3.867	0.35120	0.7
5	4.0, 4.7, 3.6	4.100	0.55680	1.1
6	4.4, 4.5, 4.1	4.333	0.20820	0.4
7	4.5, 3.9, 4.8	4.400	0.45830	0.9
8	4.0, 4.9, 4.7	4.533	0.47260	0.9
9	4.3, 4.4, 4.8	4.500	0.26460	0.5
10	5.0, 4.5, 3.5	4.333	0.76380	1.5
11	3.8, 3.7, 3.9	3.800	0.10000	0.2
12	5.1, 4.5, 4.5	4.700	0.34640	0.6
13	4.7, 4.4, 4.1	4.400	0.30000	0.6
14	4.0, 4.4, 4.6	4.333	0.30550	0.6
15	4.0, 3.3, 4.2	3.833	0.47260	0.9
16	4.2, 4.2, 4.3	4.233	0.05774	0.1

Table 8.1: Observations and summary statistics for the \*\*Beer Data\*\*.

by averaging the sample standard deviations for the k days. It can be shown that

$$\widehat{\sigma} = \frac{\overline{S}}{a_m}$$

is an unbiased estimator of  $\sigma$  where

$$a_m = \frac{\sqrt{2}\Gamma(m/2)}{\sqrt{m-1}\Gamma((n-1)/2)}.$$

Thus, we compute the  $3\sigma$  upper and lower control limits, respectively,

$$UCL = \hat{\mu} + 3 \frac{\overline{s}}{a_m \sqrt{m}}$$

and

$$LCL = \hat{\mu} - 3 \frac{\overline{s}}{a_m \sqrt{m}}.$$

The computations in r follow, along with the resulting **control chart** in Figure 8.2.

```
a <- function(m){ sqrt(2) * gamma(m/2) / (sqrt(m-1) * gamma((m-1)/2)) }
muhat = sum(ipa_stat$mean) / k
sbar = sum(ipa_stat$sd) / k
lcl = muhat - 3*sbar / (a(m) * sqrt(m))
ucl = muhat + 3*sbar / (a(m) * sqrt(m))

ggplot(ipa_stat, aes(x = Day)) + geom_point(aes(y = mean)) +
    geom_hline(aes(yintercept = muhat, color = "Mean"), size = lsz) +
    geom_hline(aes(yintercept = lcl, color = "LCL"), size = lsz*1.5) +</pre>
```

```
geom_hline(aes(yintercept = ucl, color = "UCL"), size = lsz*1.5) + ylab("pH") +
    theme(legend.justification = c(1,1), legend.position = c(1,1),
        legend.title = element_blank(),
        legend.box.margin = margin(c(4, 4, 4, 4), unit = "pt"))
```

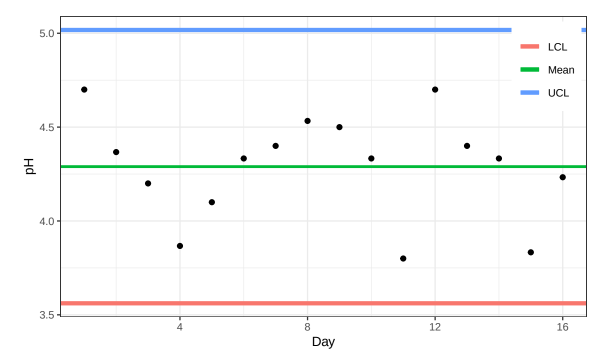


Figure 8.2: The  $3\sigma$  **control chart** illustrates that with respect to pH the brewing process is in-control over the selected timeframe as the observations fall within the (LCL, UCL) control interval.

From Figure 8.2, we observe for each day the process is in-control as the observed mean pH values fall within the control limits (LCL, UCL). If this were not the case, then our initial assumption, that process is in-control, would be violated. The violation of the assumption would require that we seek to identify an assignable cause for the variation. If a cause could be identified, then the we would need to recompute our control limits with the observations that were out of control removed.  $\Diamond$ 

# Appendix

# **Curated Content**

Below we provide links to supplementary online material. Hopefully some of the items will inspire you view the course material in a broader context and lead to further investigations.

## **Investigation 0**

What is Statistics?

#### • Cambridge Ideas - Professor Risk

https://www.youtube.com/watch?v=a1PtQ67urG4

Prof David Spiegelhalter (Cambridge University) discusses public understanding of risk.

#### • The Joy of Statistics

https://www.youtube.com/watch?v=jbkSRLYSojo

Prof Hans Rosling (Karolinska Institute and Gapminder Foundation) analyzes data from 200 Countries over 200 Years in 4 Minutes - The Joy of Stats - BBC Four.

#### • Teach statistics before calculus!

https://www.ted.com/talks/arthur\_benjamin\_teach\_statistics\_before\_calculus

Prof Arthur Benjamin (Harvey Mudd College) argues that the pinnacle of math education be probability and statistics — not calculus.

# **Investigation 1**

Defense against the dark arts.

#### • Three ways to spot bad statistics

https://www.ted.com/talks/mona\_chalabi\_3\_ways\_to\_spot\_a\_bad\_statistic

Mona Chalabi (Data Journalist) discusses three ways to spot bad statistics.

#### • No. 10 daily Covid-19 briefings

https://www.youtube.com/watch?v=QIw2l-trRXc

Prof David Spiegelhalter (Cambridge University) claims that No. 10 is misrepresenting Covid-19 data in a discussion with Andrew Marr (BBC).

#### • Statistics Done Wrong

https://www.statisticsdonewrong.com/

A book by Dr Alex Reinhart (Carnegie Mellon University).

#### • How to defend yourself against misleading statistics in the news

https://www.youtube.com/watch?v=mJ63-bQc9Xg

Sanne Blauw (Journalist) discusses how the presentation of statistics can mislead.

CURATED CONTENT 68

### **Investigation 2**

Data analysis and vizualization.

#### • The Grammar of Graphics

https://www.youtube.com/watch?v=h-62NwWUI5c

What Makes A Good Visualisation? Rhys Jackson from RocketMill, a UK Digital Marketing Agency, gives a perspective on visualizing data from a marketing perspective.

https://www.youtube.com/watch?v=kepKM7Z2O54

David Keyes (RStudio) discusses how the grammar of graphics underpins the ggplot2 data vizualization package in R.

### • Same Stats, Different Graphs

https://www.autodeskresearch.com/publications/samestats

Generating Datasets with Varied Appearance and Identical Statistics through Simulated Annealing (ACM SIGCHI Conference on Human Factors in Computing Systems) by Justin Matejka, George Fitzmaurice.

#### • Why do we so often use 0.05 for hypothesis testing?

https://www.openintro.org/book/stat/why05/

In this online exercise, you will gain an improved understanding of what a significance level is, and why a value in the neighborhood of 0.05 is reasonable as a default.

#### • Data visualizations

https://flowingdata.com/

FlowingData blog by Nathan Yau.

https://fivethirtyeight.com/

FiveThirtyEight blog by Nate Silver.

## **Investigation 3**

Statistical paradoxes.

#### • How statistics can be misleading (TED-Ed)

https://www.ted.com/talks/mark\_liddell\_how\_statistics\_can\_be\_misleading Mark Liddell (Educator) discusses Simpson's Paradox in this TED-Ed animation.

#### • Low birth-weight paradox

https://www.wikiwand.com/en/Low\_birth-weight\_paradox

#### • Gambler's Fallacy

https://www.youtube.com/watch?v=4eVluL-idkM

Prof Kelly Shue (Chicago Booth) discusses the gambler's fallacy.

### **Investigation 4**

The law and interpreting statistics.

#### • How stats fool juries.

https://ed.ted.com/lessons/peter-donnelly-shows-how-stats-fool-juries

Prof Peter Donnelly (Oxford University) discusses common mistakes in interpreting statistics.

CURATED CONTENT 69

#### • Better Data in Forensic Science

https://www.dundee.ac.uk/leverhulme/projects/details/better-data-in-forensic-science.php Dr Christian Cole (Dundee) is leading a data-focused project as part of the Leverhulme Research Centre for Forensic Science right here at Dundee.

#### • Prosecutor's fallacy

https://www.wikiwand.com/en/Prosecutor%27s\_fallacy

A fallacy of statistical reasoning, typically used by a prosecutor to exaggerate the likelihood of guilt: because  $P(\text{hypothesis} \mid \text{evidence}) \neq P(\text{evidence} \mid \text{hypothesis})!$ 

## **Investigation 5**

Data-driven decision making in epidemiology.

#### • Project Tycho

https://www.tycho.pitt.edu/

Digitized archival epidemiological data for the United States and the world.

https://www.youtube.com/watch?v=Kn9OJy1BPDo

An overview of the origins of project Tycho.

#### • Public Health Scotland COVID-19 Dashboard

https://public.tableau.com/shared/N2M72MH2B

The official COVID-19 dashboard of Public Health Scotland.

#### • Our World in Data

https://ourworldindata.org/

A project of the Oxford Martin School to make public health data, including progress in UN Sustainable Development Goals, available and accessible.

#### • Demographic Party Trick

https://www.youtube.com/watch?v=2nDh8MQuS-Y

Prof Hans Rosling (Karolinska Institute and Gapminder Foundation) and Bill Gates seek to shed light on the true statistics of childhood vaccinations.

# **Investigation 6**

Spurious correlations!

#### • The danger of mixing up causality and correlation

https://www.youtube.com/watch?v=8B271L3NtAw

Prov Ionica Smeets (University of Leiden) discusses causality and correlation.

#### • Spurious correlations

https://tylervigen.com/spurious-correlations

Tyler Vigen's site dedicated to spurious correlations.

#### • Cause & Effect

https://www.youtube.com/watch?v=lbODqslc4Tg

Correlation vs. causality from the Clip from the 2010 documentary "Freakonomics: The Movie".

CURATED CONTENT 70

### **Investigation 7**

Data and society: can data-driven and predictive modeling lead to a better world? What are the ethics of mass data collection?

#### • Science behind the news: Predictive Policing

https://www.youtube.com/watch?v=U0gX\_z0V0nE

The Los Angeles Police Department is using a new tactic in their fight against crime called "predictive policing." It's a computer program that was originally developed by a team at UCLA, including mathematician Andrea Bertozzi and anthropologist Jeff Brantingham. "Science Behind the News" is produced in partnership with NBC Learn. (Provided by the National Science Foundation & NBC Learn)

#### • You should get paid for your data

https://www.nytimes.com/video/opinion/100000006678020/data-privacy-jaron-lanier-2.html

Jaron Lanier (Computer Scientist and Author) discusses a compensation plan and data dignity. https://www.ted.com/talks/jennifer\_zhu\_scott\_why\_you\_should\_get\_paid\_for\_your\_data
Jennifer Zhu Scott (Computer Scientist) also thinks you should get paid for your data.

#### • How tech companies deceive you into giving up your data and privacy

https://www.ted.com/talks/finn\_lutzow\_holm\_myrstad\_how\_tech\_companies\_deceive\_you \_into\_giving\_up\_your\_data\_and\_privacy

Finn Lützow-Holm Myrstad (Norwegian Consumer Council) discusses consumer protections and data collection.

#### • Your company's data could help end world hunger

https://www.ted.com/talks/mallory\_freeman\_your\_company\_s\_data\_could\_help\_end\_world\_hunger

Mallory Freeman (Data Scientist) discusses how to do the most good with data.

# **Investigation 8**

Machine learning / big data.

#### • What is Machine Learning?

https://www.youtube.com/watch?v=f uwKZIAeM0

OxfordSparks discusses the topic of supervised learning algorithms and how machine learning is used all around us.

#### • Big Data (TED-Ed)

https://www.youtube.com/watch?v=j-0cUmUyb-Y

Tim Smith (educator) discusses the historical arc of big data in this TED-Ed animation.

#### • The human insights missing from big data

https://www.ted.com/talks/tricia\_wang\_the\_human\_insights\_missing\_from\_big\_data Tricia Wang (Ethnographer) discusses the human insights missing from big data.

#### • How we can find ourselves in data

https://www.ted.com/talks/giorgia\_lupi\_how\_we\_can\_find\_ourselves\_in\_data Giorgia Lupi (Designer) discusses a humanistic approach to data and data visualization.

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