

Simple statistical tests

::: callout-tip ## Extended Materials

You can find the original, extended version of this chapter [here](#). ::

This page demonstrates how to conduct simple statistical tests using **base R**, **rstatix**, and **gtsummary**.

- T-test
- Shapiro-Wilk test
- Wilcoxon rank sum test
- Kruskal-Wallis test
- Chi-squared test
- Correlations between numeric variables

...many other tests can be performed, but we showcase just these common ones and link to further documentation.

Each of the above packages bring certain advantages and disadvantages:

- Use **base R** functions to print a statistical outputs to the R Console
- Use **rstatix** functions to return results in a data frame, or if you want tests to run by group
- Use **gtsummary** if you want to quickly print publication-ready tables

Load packages

```
pacman::p_load(
  rio,          # File import
  here,         # File locator
  skimr,        # get overview of data
  tidyverse,    # data management + ggplot2 graphics,
  gtsummary,    # summary statistics and tests
  rstatix,      # statistics
  corrr,        # correlation analysis for numeric variables
  janitor,      # adding totals and percents to tables
  flextable     # converting tables to HTML
)
```

An example statistical test

Let's look at a classic example in statistics: flipping a coin to see if it is fair. We flip the coin 100 times and each time record whether it came up heads or tails. So, we have a record that could look something like HHTHTHTT...

Let's simulate the experiment in R, using a biased coin:

```
set.seed(0xdada)
numFlips = 100
probHead = 0.6
# Sample is a function in base R which lets us take a random sample from a vector, with or without replacement.
# This line is sampling numFlips times from the vector ['H','T'] with replacement, with the probability of each item in the vector being defined in the prob argument as [probHead, 1-probHead]
coinFlips = sample(c("H", "T"), size = numFlips,
  replace = TRUE, prob = c(probHead, 1 - probHead))
# Thus, coinFlips is a character vector of a random sequence of 'T' and 'H'.
head(coinFlips)
```

```
[1] "T" "T" "H" "T" "H" "H"
```

Now, if the coin were fair, we would expect half of the time to get heads. Let's see.

```
table(coinFlips)
```

```
coinFlips
```

H T
59 41

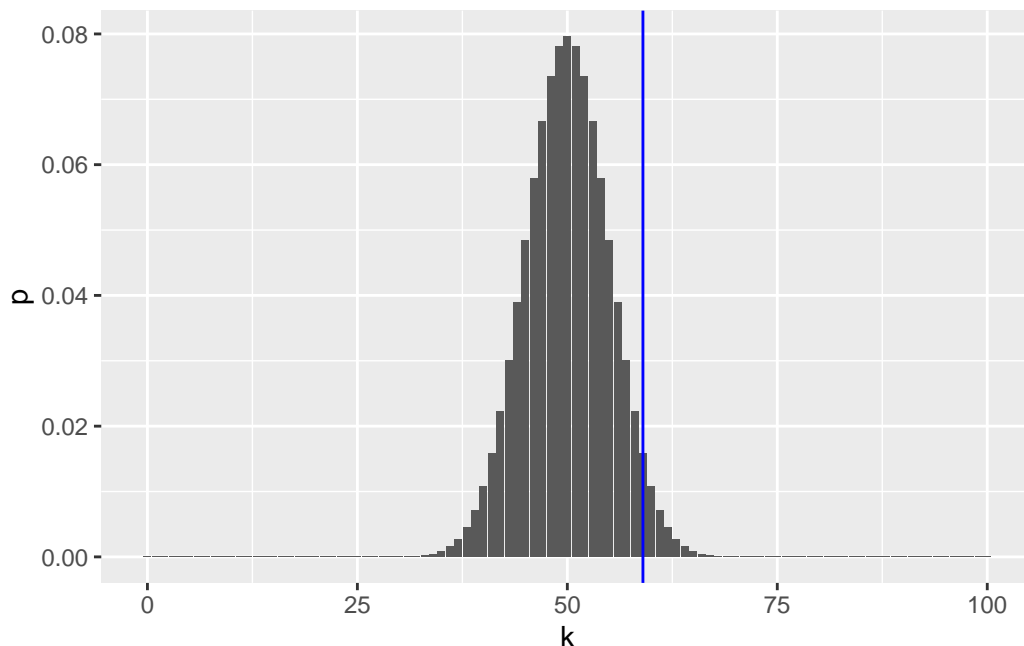
That is different from 50/50. However, does the data deviates strong enough to conclude that this coin isn't fair? We know that the total number of heads seen in 100 coin tosses for a fair coin follows $B(100, 0.5)$, making it a suitable test statistic.

To decide, let's look at the sampling distribution of our test statistic – the total number of heads seen in 100 coin tosses – for a fair coin. As we learned, we can do this with the binomial distribution. Let's plot a fair coin and mark our observation with a blue line:

```
# This line sets k as the vector [0, 1, 2,...,numFlips]
k <- 0:numFlips
# Recall that binary variables (TRUE and FALSE) are interpreted as 1 and 0, so we can use
# to count the number of heads in coinFlips. We practice these kinds of operations in sess
numHeads <- sum(coinFlips == "H")
# We use dbinom here to get the probability mass at every integer from 1-numFlips so that
p <- dbinom(k, size = numFlips, prob = 0.5)
# We then convert it into a dataframe for easier plotting.
binomDensity <- data.frame(k = k, p = p)
head(binomDensity)
```

	k	p
1	0	7.888609e-31
2	1	7.888609e-29
3	2	3.904861e-27
4	3	1.275588e-25
5	4	3.093301e-24
6	5	5.939138e-23

```
# Here, we are plotting the binomial distribution, with a vertical line representing
# the number of heads we actually observed. We will learn how to create plots in session 4
# Thus, to complete our test we simply need to identify whether or not the blue line
# is in our rejection region.
ggplot(binomDensity) +
  geom_bar(aes(x = k, y = p), stat = "identity") +
  geom_vline(xintercept = numHeads, col = "blue")
```



How do we quantify whether the observed value is among those values that we are likely to see from a fair coin, or whether its deviation from the expected value is already large enough for us to conclude with enough confidence that the coin is biased?

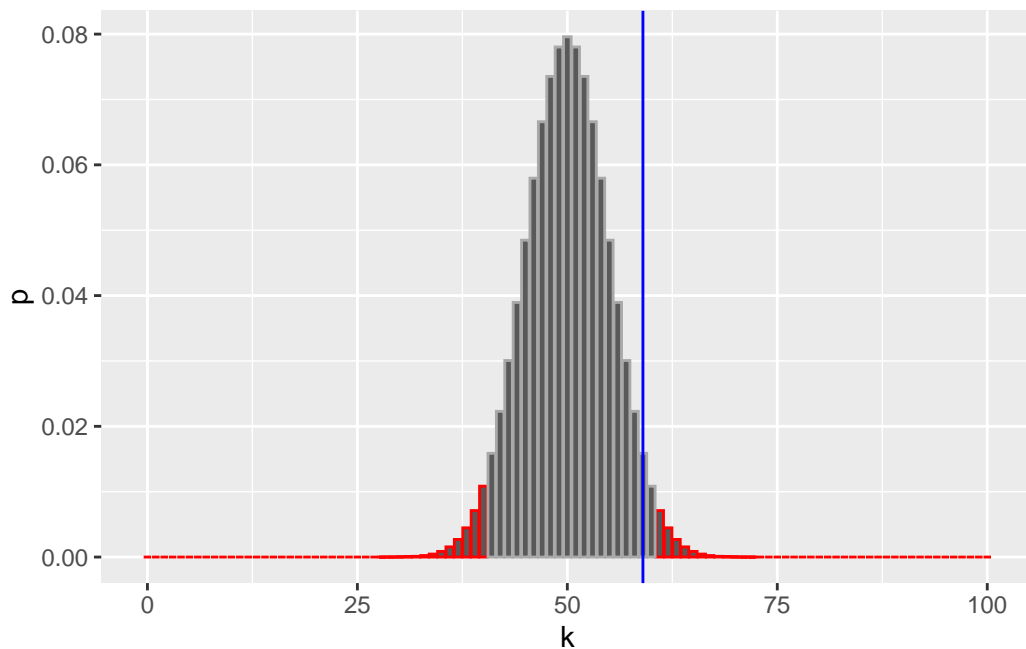
We divide the set of all possible $k(0-100)$ in two complementary subsets, the **rejection region** and the region of no rejection. We want to make the rejection region as large as possible while keeping their total probability, assuming the null hypothesis, below some threshold α (say, 0.05).

```
alpha <- 0.05
# We get the density of our plot in sorted order, meaning that we'll see binomDensity
# jump back and forth between the distribution's tails as p increases.
binomDensity <- binomDensity[order(p),]
# We then manually calculate our rejection region by finding where the cumulative sum in t
# is less than or equal to our chosen alpha level.
binomDensity$reject <- cumsum(binomDensity$p) <= alpha
head(binomDensity)
```

	k	p	reject
1	0	7.888609e-31	TRUE
101	100	7.888609e-31	TRUE
2	1	7.888609e-29	TRUE

100	99	7.888609e-29	TRUE
3	2	3.904861e-27	TRUE
99	98	3.904861e-27	TRUE

```
# Now we recreate the same plot as before, but adding red borders around the parts of our
# in the rejection region.
ggplot(binomDensity) +
  geom_bar(aes(x = k, y = p, col = reject), stat = "identity") +
  scale_colour_manual(
    values = c(`TRUE` = "red", `FALSE` = "darkgrey")) +
  geom_vline(xintercept = numHeads, col = "blue") +
  theme(legend.position = "none")
```



We sorted the p -values from lowest to highest (**order**), and added a column **reject** by computing the cumulative sum (**cumsum**) of the p -values and thresholding it against **alpha**.

The logical column **reject** therefore marks with **TRUE** a set of k s whose total probability is less than α .

The rejection region is marked in red, containing both very large and very small values of k , which can be considered unlikely under the null hypothesis.

R provides not only functions for the densities (e.g., `dbinom`) but also for the cumulative distribution functions (`pbinom`). Those are more precise and faster than `cumsum` over the probabilities.

The (cumulative) *distribution function* is defined as the probability that a random variable X will take a value less than or equal to x .

$$F(x) = P(X \leq x)$$

We have just gone through the steps of a **binomial test**. This is a frequently used test and therefore available in R as a single function.

We have just gone through the steps of a binomial test. In fact, this is such a frequent activity in R that it has been wrapped into a single function, and we can compare its output to our results.

```
binom.test(x = numHeads, n = numFlips, p = 0.5)
```

```
Exact binomial test
```

```
data: numHeads and numFlips
number of successes = 59, number of trials = 100, p-value = 0.08863
alternative hypothesis: true probability of success is not equal to 0.5
95 percent confidence interval:
 0.4871442 0.6873800
sample estimates:
probability of success
               0.59
```

Hypothesis Tests

We can summarize what we just did with a series of steps:

1. Decide on the effect that you are interested in, design a suitable experiment or study, pick a data summary function and test statistic.
2. Set up a null hypothesis, which is a simple, computationally tractable model of reality that lets you compute the null distribution, i.e., the possible outcomes of the test statistic and their probabilities under the assumption that the null hypothesis is true.
3. Decide on the rejection region, i.e., a subset of possible outcomes whose total probability is small.
4. Do the experiment and collect the data; compute the test statistic.
5. Make a decision: reject the null hypothesis if the test statistic is in the rejection region.

Types of Error



Having set out the mechanics of testing, we can assess how well we are doing. The following table, called a **confusion matrix**, compares reality (whether or not the null hypothesis is in fact true) with our decision whether or not to reject the null hypothesis after we have seen the data.

Test vs reality	Null is true	Null is false
Reject null	Type I error (false positive)	True positive
Do not reject null	True negative	Type II error (false negative)

It is always possible to reduce one of the two error types at the cost of increasing the other one. The real challenge is to find an acceptable trade-off between both of them. We can always decrease the **false positive rate** (FPR) by shifting the threshold to the right. We can become more “conservative”. But this happens at the price of higher **false negative rate** (FNR). Analogously, we can decrease the FNR by shifting the threshold to the left. But then again, this happens at the price of higher FPR. The FPR is the same as the probability α that we mentioned above. $1 - \alpha$ is also called the **specificity** of a test. The FNR is sometimes also called β , and $1 - \beta$ the **power, sensitivity** or **true positive rate** of a test. The power of a test can be understood as the likelihood of it “catching” a true positive, or correctly rejecting the null hypothesis.

Generally, there are three factors that can affect statistical power:

- Sample size: Larger samples provide greater statistical power
- Effect size: A given design will always have greater power to find a large effect than a small effect (because finding large effects is easier)
- Type I error rate: There is a relationship between Type I error and power such that (all else being equal) decreasing Type I error will also decrease power.

Preparation

Import data

We import the dataset of cases from a simulated Ebola epidemic. If you want to follow along, click to download the “clean” linelist (as .rds file).

```
# import the linelist
linelist <- import("linelist_cleaned.rds")
```

base R

You can use **base** R functions to conduct statistical tests. The commands are relatively simple and results will print to the R Console for simple viewing. However, the outputs are usually lists and so are harder to manipulate if you want to use the results in subsequent operations.

T-tests

A [t-test](#), also called “Student’s t-Test”, is typically used to determine if there is a significant difference between the means of some numeric variable between two groups. Here we’ll show the syntax to do this test depending on whether the columns are in the same data frame.

Syntax 1: This is the syntax when your numeric and categorical columns are in the same data frame. Provide the numeric column on the left side of the equation and the categorical column on the right side. Specify the dataset to `data =`. Optionally, set `paired = TRUE`, and `conf.level = (0.95 default)`, and `alternative =` (either “two.sided”, “less”, or “greater”). Enter `?t.test` for more details.

```
## compare mean age by outcome group with a t-test
t.test(age_years ~ gender, data = linelist)
```

Welch Two Sample t-test


```

data:  age_years by gender
t = -21.344, df = 4902.3, p-value < 2.2e-16
alternative hypothesis: true difference in means between group f and group m is not equal to
95 percent confidence interval:
  -7.571920 -6.297975
sample estimates:
mean in group f mean in group m
    12.60207      19.53701

```

Syntax 2: You can compare two separate numeric vectors using this alternative syntax. For example, if the two columns are in different data sets.

```
t.test(df1$age_years, df2$age_years)
```

You can also use a t-test to determine whether a sample mean is significantly different from some specific value. Here we conduct a one-sample t-test with the known/hypothesized population mean as `mu =`:

```
t.test(linelist$age_years, mu = 45)
```

Shapiro-Wilk test

The [Shapiro-Wilk test](#) can be used to determine whether a sample came from a normally-distributed population (an assumption of many other tests and analysis, such as the t-test). However, this can only be used on a sample between 3 and 5000 observations. For larger samples a [quantile-quantile plot](#) may be helpful.

```
shapiro.test(linelist$age_years)
```

Wilcoxon rank sum test

The Wilcoxon rank sum test, also called the [Mann-Whitney U test](#), is often used to help determine if two numeric samples are from the same distribution when their populations are not normally distributed or have unequal variance.

```

## compare age distribution by outcome group with a wilcox test
wilcox.test(age_years ~ outcome, data = linelist)

```

Wilcoxon rank sum test with continuity correction

```
data: age_years by outcome
W = 2501868, p-value = 0.8308
alternative hypothesis: true location shift is not equal to 0
```

Kruskal-Wallis test

The [Kruskal-Wallis test](#) is an extension of the Wilcoxon rank sum test that can be used to test for differences in the distribution of more than two samples. When only two samples are used it gives identical results to the Wilcoxon rank sum test.

```
## compare age distribution by outcome group with a kruskal-wallis test
kruskal.test(age_years ~ outcome, linelist)
```

Kruskal-Wallis rank sum test

```
data: age_years by outcome
Kruskal-Wallis chi-squared = 0.045675, df = 1, p-value = 0.8308
```

Chi-squared test

[Pearson's Chi-squared test](#) is used in testing for significant differences between categorical groups.

```
## compare the proportions in each group with a chi-squared test
chisq.test(linelist$gender, linelist$outcome)
```

Pearson's Chi-squared test with Yates' continuity correction

```
data: linelist$gender and linelist$outcome
X-squared = 0.0011841, df = 1, p-value = 0.9725
```

Correlations

Correlation between numeric variables can be investigated using the **tidyverse** **corr** package. It allows you to compute correlations using Pearson, Kendall tau or Spearman rho. The package creates a table and also has a function to automatically plot the values.

```
correlation_tab <- linelist %>%  
  select(generation, age, ct_blood, days_onset_hosp, wt_kg, ht_cm) %>% # keep numeric variables  
  correlate() # create correlation table (using default pearson)
```

```
correlation_tab # print
```

A tibble: 6 x 7

term <chr>	generation <dbl>	age <dbl>	ct_blood <dbl>	days_onset_hosp <dbl>	wt_kg <dbl>	ht_cm <dbl>
1 generation	NA	-2.22e-2	0.179	-0.288	-0.0302	-0.00942
2 age	-0.0222	NA	0.00849	-0.000635	0.833	0.877
3 ct_blood	0.179	8.49e-3	NA	-0.600	-0.00636	0.0181
4 days_onset_hosp	-0.288	-6.35e-4	-0.600	NA	0.0153	-0.00953
5 wt_kg	-0.0302	8.33e-1	-0.00636	0.0153	NA	0.884
6 ht_cm	-0.00942	8.77e-1	0.0181	-0.00953	0.884	NA

```
## remove duplicate entries (the table above is mirrored)
```

```
correlation_tab <- correlation_tab %>%  
  shave()
```

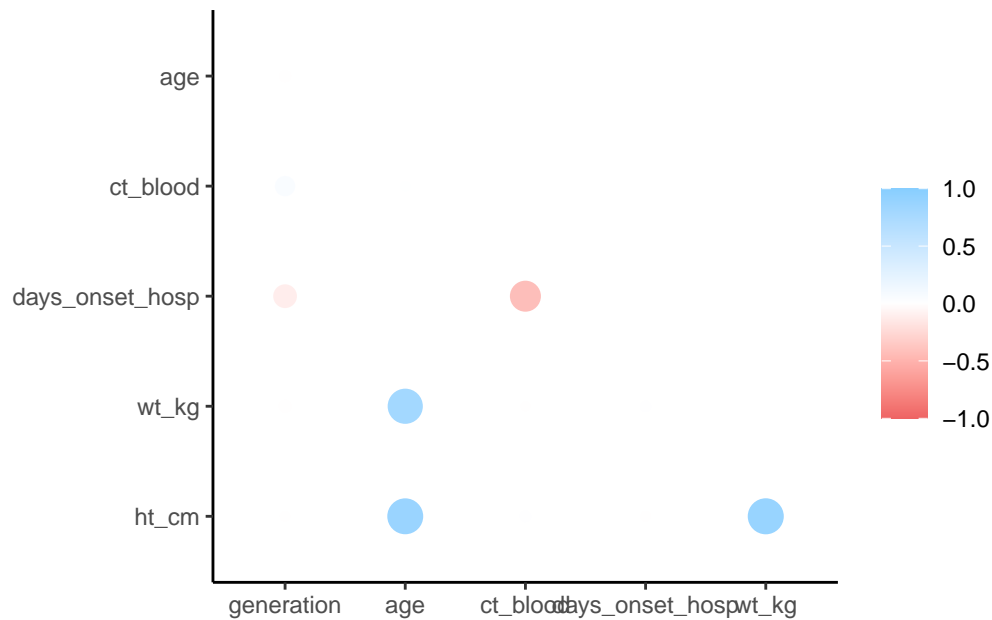
```
## view correlation table
```

```
correlation_tab
```

A tibble: 6 x 7

term <chr>	generation <dbl>	age <dbl>	ct_blood <dbl>	days_onset_hosp <dbl>	wt_kg <dbl>	ht_cm <dbl>
1 generation	NA	NA	NA	NA	NA	NA
2 age	-0.0222	NA	NA	NA	NA	NA
3 ct_blood	0.179	0.00849	NA	NA	NA	NA
4 days_onset_hosp	-0.288	-0.000635	-0.600	NA	NA	NA
5 wt_kg	-0.0302	0.833	-0.00636	0.0153	NA	NA
6 ht_cm	-0.00942	0.877	0.0181	-0.00953	0.884	NA

```
## plot correlations
rplot(correlation_tab)
```



Resources

Much of the information in this page is adapted from these resources and vignettes online:

- [gtsummary](#)
- [dplyr](#)
- [corr](#)
- [sthda correlation](#) - *Statistical Thinking for the 21st Century* by Russell A. Poldrack. This work is distributed under the terms of the [Attribution-NonCommercial 4.0 International \(CC BY-NC 4.0\)](#), which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited and the material is used for noncommercial purposes.
- [Modern Statistics for Modern Biology](#) by Susan Holmes and Wolfgang Huber. This work is distributed under the terms of the [Attribution-NonCommercial-ShareAlike 2.0 Generic \(CC BY-NC-SA 2.0\)](#), which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited, the material is used for noncommercial purposes, and the same license is used for any derivative material.