# **Logistic regression - introduction**

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#### Introduction

THESE NOTES are a basic introduction to binary logistic regression used to analyse data with binary outcomes. In these notes we'll aim to cover the following learning outcomes:

- · Refresher on logs, odds, probability and linear regression
- · Understand why linear regression not sensible for binary data
- · Explain how logit and binomial model let us extend linear regression
- Be able to run a simple logistic regression in R
- · Be able to explain basic R glm output
- Be able to explain estimates with categorical and continuous variables
- · Explain significance test results on variables
- Introduce some basic ideas for selecting variables and models
- · Things to watch out for!
- · Know where to go next!

## **Prerequisites**

We'll cover a couple of background topics but to follow these notes you'll need to be able to load packages in R, run simple code in R and have a basic understanding of linear regression and statistical hypothesis tests.

### Running the example code

To run the R code in these notes you'll need to start an rstudio project, load in the example data-set and have a few packages downloaded and loaded into your R session.

To download the packages (if you don't have them already) use...

```
install.packages("tidyverse")
install.packages("boot")
install.packages("broom")
install.packages("skimr")
install.packages("sjPlot")
install.packages("kableExtra")
```

Start a new project in restudio and in an recript use the floowing code to load the libraries you need and to import/load the data...

You can also use the install button in the RStudio packages tab to install these

Loading the packages...

```
library(tidyverse)
library(boot)
library(broom)
library(skimr)
library(sjPlot)
library(kableExtra)
```

Loading the data (from the csv file on Basecamp)...

```
dat <- read_csv("logreg_data_01_20190530.csv")</pre>
```

This dataset describes 500 animals giving their weight in kg, age in days, supplement levels (mg), sex, region where they liver (A, B, C or D) and whether they were treated with anthelmintics or not. It's a dataset made up for this course by the way!

#### Revision / background topics

Logarithms ('logs')

Skip this if you are happy with logs (including base 'e')

'Logs' are a mathematical function that changes a number. They take the form  $log_b(x) = y$ . What this means is b to the power y will give us x. It's easier to understand with examples. Let's start with base 10....

$$log_{10}(10) = 1$$

$$log_{10}(1000) = 3$$

$$log_{10}(0.01) = -2$$

So the logs of all the numbers in brackets are the number you'd need to raise 10 to to get them.

We can have other bases e.g. e., e is a special mathematical constant that feautres a lot behind the scene in statistics it's roughly 2.718...

$$log_e(2.718) \simeq 1$$

'Inverse logs' let us turn logs back into the original number. We simply raise the 'base' of our logs to the number we what to invert and we end up with the original number. So  $log_{10}(1000) = 3$  and  $10^3 = 1000...$ 

One feature of logs is that adding the logs of two numbers is equivalent to multiplying the numbers...

Once loaded you should see the a 'dat' object in the RStudio environment tab (usually in the top-right of your screen)

$$100 \times 1000 = 100000$$

$$log_{10}(100) + log_{10}(1000) = log_{10}(100000)$$

Because  $log_{10}(100) = 2$ ,  $log_{10}(1000) = 3$  and  $log_{10}(100000) = 5$ 

If this all seems a bit too much don't worry. Just remember that adding logs is like multiplying numbers and you'll be fine!

## Odds and probability

Probabilities have values from 0 ('never happens') to 1 ('always happens')

'events of interest' ÷ 'all events'.

What is the probability that a fair coin lands on heads?

$$1/2 = 0.5$$

What is the probability that a 6 sided die lands on 4?

$$1/6 \simeq 0.166$$

**Odds** have values from 0 ('never happen') to infinity ('always happens')

'events of interest' ÷ 'other events'.

What are the odds that a fair coin lands on heads?

$$1/1 = 1$$

What are the odds that a 6 sided die lands on 4?

$$1/5 = 0.2$$

#### Linear regression

Remember that linear regression is a statistical method that lets us understand and predict numerical outcomes using one or more predictor variables. The predictor variables may be numerical or categorical. Linear regression also assumes there's a linear i.e. straight line relationship between the predictor numerical variable sand the outcome. Using the data set we have loaded we can plot weight against age and sex and see that there looks to be a linear relationship between age and weight for each sex...

```
ggplot(dat) +
  aes(age, weight, colour = sex) +
  geom_point(shape = 1) +
  geom_smooth(method = "lm", se = FALSE) +
  theme_bw() +
  labs(x = "Age (days)",
       y = "Weight (Kg)",
       colour = "Sex")
```

In R we can use the lm() function to fit a linear model to this data. Normally we store the results of using the function in an R object and look at it using 'summary(). We can also get a tidier output using get\_model\_data() from the sjPlot package. Here we make a linear model predicting the animal's weight from their age (a numerical variable) and their sex (a categorical variable).

```
mod1 <- lm(weight ~ age + sex, data = dat)</pre>
```

Using summary() from base-R...

```
summary (mod1)
```

```
Call:
lm(formula = weight ~ age + sex, data = dat)
Residuals:
   Min
            1Q Median
                           3Q
                                 Max
-2.3506 -0.4732 -0.0509 0.4572 2.0223
Coefficients:
          Estimate Std. Error t value Pr(>|t|)
(Intercept) 49.93305
                      0.40486 123.33 <2e-16 ***
           0.12045
                      0.00191
                               63.07 <2e-16 ***
sexmale
            3.51683
                      0.06541
                               53.76
                                     <2e-16 ***
Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' '1
```

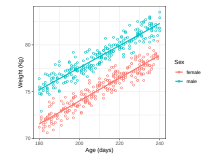


Figure 1: Weight vs Age of the animals

```
Residual standard error: 0.7312 on 497 degrees of freedom
Multiple R-squared: 0.932, Adjusted R-squared: 0.9317
F-statistic: 3404 on 2 and 497 DF, p-value: < 2.2e-16
```

Using get\_model\_data() from sjPlot package and selecting the output columns we want...

```
get_model_data(mod1) %>%
 select(term:p.stars) %>%
 print()
# A tibble: 2 x 8
 term estimate std.error statistic p.value conf.low conf.high p.stars
          <dbl> <dbl> <dbl> <dbl> <dbl>
                                                    <dbl> <chr>
 <fct>
1 age
           0.120 0.00191 63.1 2.50e-239 0.117
                                                    0.124 ***
2 sexmale
           3.52 0.0654
                           53.8 2.87e-209 3.39
                                                    3.65 ***
```

The analysis suggestes that both age and sex are significant predictors of weight. Weight increasing by about 0.12 Kg per day of age and that males are about 3.5 Kg heavier than females. Right - now to look at binary (yes/no) outcome data and logistic regression...

#### **Analysing binary data**

Binary data is common in epidemiological studies e.g. disease status (diseased or healthy), life (alive or dead) etc. In the example dataset we have the column status where animals can be either healthy or diseased.

ID	treatment	age	region	supp	sex	weight	status
A0049	control	221	D	7.891	male	80.9	diseased
A0485	control	220	Α	2.651	male	79.5	diseased
A0321	treated	238	В	1.671	male	82.2	diseased
A0153	treated	183	С	6.346	female	71.7	healthy
A0074	treated	187	Α	9.655	male	77.0	diseased
A0228	control	206	С	8.046	female	74.7	diseased

As we know that some animals were treated with an anthelmintic and some were not it may be interesting to look at the number of healthy and diseased animals in the treated and control (untreated) groups.

treatment	diseased	healthy
control	154	44
treated	209	93

Approximately 78% of the untreated (control) animals were diseased and 69% of the treated animals were diseased. We can use a 'Fisher's Exact test' to test if the proportion of diseased anaimals in the treatment vs control groups are significantly different. This test also estimates the odds ratio of disease between the two groups.

```
with(dat,
     {{fisher.test(status, treatment)}})
```

Fisher's Exact Test for Count Data

```
data: status and treatment
p-value = 0.04032
alternative hypothesis: true odds ratio is not equal to 1
95 percent confidence interval:
1.010694 2.419528
sample estimates:
odds ratio
  1.556051
```

The Fisher's exact test reports a p-value, in this case 0.04. If this is less

You can use ?fisher.test in the RStudio console to get help on this function. Here we have also used the with() function to make it easier to use the 'dat' dataset and refer to the columns in the fisher.test() code

than your chosen critical p-value you have evidence that the proprtion of diaased animals is significantly different between the treated and untreated (control) animal groups. Fisher's Exact test is useful for comaprions between a binary outcome and a single, binary predciotr. However we may want to look at the association between several predcitors and a binary outcome in one go. This is called multivariable analysis and you'll have met it in linear regression. We can't look at this with a simple test so need to consider regression approaches...

#### Multivariable analysis

We could convert out status column to a number coding healhty animals as 0 and diseased animals as 1 and then use conventional linear regression with the variables we want to consider as predictors. The following code will make a new column, status01 recoding status to a number.

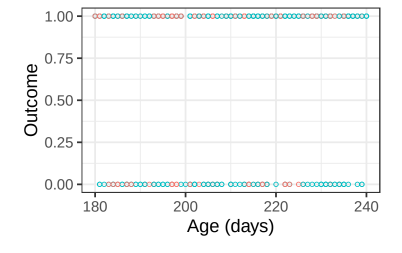
```
dat <- dat %>%
 mutate(status01 = as.numeric(status == "diseased"))
```

To understand how this works remember that == in R compares things and if they are the same returns TRUE andf if they aren't returns FALSE. Then as.numeric() will convert FALSE to a 0 and TRUE to 1. It's a handy way of making binary text /factor data into 0's and

ID	treatment	age	region	supp	sex	weight	status	status01
A0001	treated	219	D	6.731	female	76.2	diseased	1
A0002	treated	218	С	7.950	female	76.0	diseased	1
A0003	control	214	Α	5.617	female	75.3	healthy	0
A0004	control	194	Α	8.490	female	72.1	diseased	1

Now we have converted the binary data into a number (0 or 1) we **could** try using linear regression. First, to help understand what we are trying here, we can plot the outcome against the treatment group and age of the animals we see this...

```
ggplot(dat) +
  aes(age, status01, colour = treatment) +
  geom_point(shape = 1) +
  theme_bw(base_size = 16) +
  labs(x = "Age (days)",
       y = "Outcome",
       colour = "Treatment")
```



#### Treatment

- control
- treated

Figure 2: Status vs Age and treatment

<dbl> <chr> 0.00430 ""

-0.00796 \*

Linear regression will attempt to fit straight lines of 'prediction' to the control and treatment data.

<dbl>

0.00203 0.00116

<fct>

2 treatmenttreated -0.0876

1 age

```
mod_lin <- lm(status01 ~ age + treatment, data = dat)</pre>
get_model_data(mod_lin) %>%
  select(term:p.stars)
# A tibble: 2 x 8
  term
                   estimate std.error statistic p.value conf.low conf.high p.stars
```

<dbl>

0.0406

<dbl>

<dbl>

1.75 0.0811 -0.000246

-2.16 0.0316 -0.167

<dbl>

Before we get too excited by this let's look at the fitted prediction line on the original plot.

```
dat %>%
  modelr::add_predictions(mod_lin) %>%
  ggplot() +
  aes(age, status01, colour = treatment) +
  geom_point(shape = 1) +
  geom_line(aes(y = pred)) +
  theme_bw(base_size = 16) +
  labs(x = "Age (days)",
       y = "Outcome",
       colour = "Treatment")
```

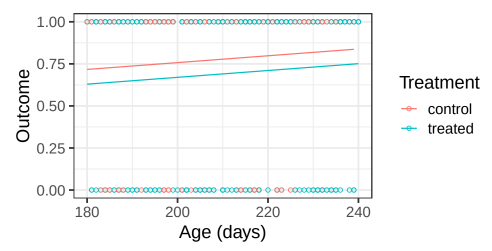


Figure 3: Status vs Age and treatment (with linear model)

Our outomce can only be 0 i.e. healhty or 1 i.e. diseased. Howvere our linear mode is predicting numbers inbetween. In some circumstance trying to use a linear model to predict binary 0/1/ data will even give values less than zero and more than one. We need to re-think the regression approach so that our model makes sense with an outcomethat can only be 0 or 1 (or healhty / diseased etc)