Class 3: Linear Regression and GLMs

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Learning outcomes

- ► Know how to fit and interpret a linear regression
- ▶ Know the difference between a linear regression model and a generalised linear model (GLM)
- ► Know what a link function is and why it is used
- ▶ Be able to interpret the output of a simple GLM

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Using more of the data

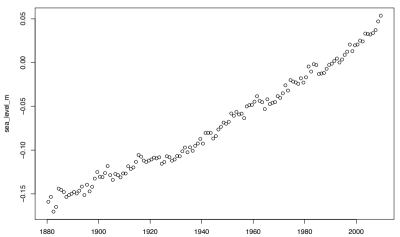
- ▶ It's very rare that we are given a data set with just a single
- ▶ More often we're given multiple variables and asked to predict one or more of the variables from the others
- ► This is an example of *conditional* inference
- ▶ It might look more complicated, but this is still just fitting a probability distribution to some data

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Linear regression example

Here is some data on sea level rise:

```
sl = read.csv('../../data/tide_gauge.csv')
with(sl, plot(year_AD, sea_level_m))
```



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Linear regression models

- ▶ The simplest version of a linear regression model has:
 - ► A response variable (y) which is what we are trying to predict/understand
 - ► An explanatory variable or covariate (x) which is what we are trying to predict the response variable from
 - ▶ Some *residual uncertainty* (ϵ) which is the leftover uncertainty that is not accounted for by the explanatory variable
- ▶ Our goal is to predict the response variable from the explanatory variable, *or* to try and discover if the explanatory variable *causes* some kind of change in the response

The linear models in maths

▶ We write the linear model as:

$$y_i = \alpha + \beta x_i + \epsilon_i$$

where α is the intercept, β the slope, and $i=1,\ldots,N$ represents each of the N observations

- ▶ Usually we make the additional assumption that $\epsilon_i \sim N(0, \sigma^2)$ where σ^2 is the residual standard deviation
- Writing this in probability distributions:

$$v_i|x_i,\alpha,\beta,\sigma\sim N(\alpha+\beta x_i,\sigma^2)$$

.

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Fiting linear regression models

► We can create a likelihood as before by guessing some values of the parameters and then using the dnorm function to compute the likelihood value

```
alpha = 2
beta = 1.5
sigma = 0.6
y = sl$sea_level_m
x = sl$year_AD
sum(dnorm(y, mean = alpha + beta*x, sd = sigma, log = TRUE)
```

[1] -1539607458

▶ Not a very high value of the likelihood!

Finding the best values

► Luckily, R has the lm function to find the best fitting values of the parameters

```
summary(lm(y ~ x))
## Call:
## lm(formula = y ~ x)
## Residuals:
                     1Q
                           Median
   -0.0167787 -0.0051874 -0.0003646
## 0.0063022 0.0252621
## Coefficients:
                Estimate Std. Error
##
## (Intercept) -3.062e+00 3.937e-02
## v
               1.538e-03 2.024e-05
##
              t value Pr(>|t|)
## (Intercept) -77.78 <2e-16 ***
                75.99 <2e-16 ***
## Signif. codes:
## 0 '***' 0.001 '**' 0.01 '*' 0.05
    '.' 0.1 ' ' 1
## Residual standard error: 0.008659 on 128 degrees of freedom
## Multiple R-squared: 0.9783, Adjusted R-squared: 0.9781
## F-statistic: 5775 on 1 and 128 DF, p-value: < 2.2e-16
```

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Checking the likelihood

```
## [1] 433.9335
```

A much higher value of the likelihood!

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Checking the model

- ▶ Just finding the best values of the parameters and their uncertainty is not the whole story
- ▶ We need to check the fit of the model
- ► We can do this by analysing whether the assumed probability distribution is correct or not
- ► Can look at the probability distribution we have fitted to the data, but most people just look at the leftover bits the residuals

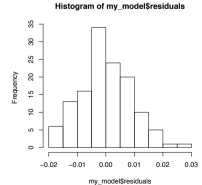
Other notes about 1m

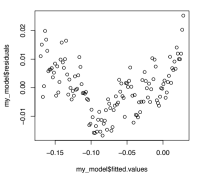
- Usually we would store the output from lm in another object to
 allow us to manipulate the output, e.g. my_model = lm(y ~
 x)
- ► We can use the confint function to get confidence intervals on the parameters
- ► We can predict future values of sea level from the model by giving it new x values, e.g.

Residual plot

► Common to plot a histogram of the residuals, and a scatter plot of the residuals vs the fits

```
par(mfrow=c(1, 2))
hist(my_model$residuals)
plot(my_model$fitted.values, my_model$residuals)
```





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Transforming the data

- ► Sometimes the residuals of a linear regression look a little bit mis-shapen
- ▶ We might improve the fit by adding more covariates, or by transforming the data (the response and/or the covariates)
- ▶ If your variables have very large values then you might get better results by standardising your data (subtracting the mean and dividing by the standard deviation)
- ► Common transformations include the log or square root
- ► A common transformation in time series data is the *Box-Cox* transformation...

Box-Cox

► The Box-Cox transformation is:

$$f(x; \lambda) = \frac{x^{\lambda} - 1}{\lambda}$$
 if $\lambda \neq 0$

or

$$f(x; \lambda) = \log(x)$$
 if $\lambda = 0$

- ► The usual reason to use it is when the data are *skewed* and we want it to look more symmetrical
- ▶ You need to choose the value of λ ; usually trial and error

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From LMs to GLMs

- ▶ If a normal distribution is not suitable for the residuals we need to choose another probability distribution
- ▶ Here is some data from an experiment on whitefly:

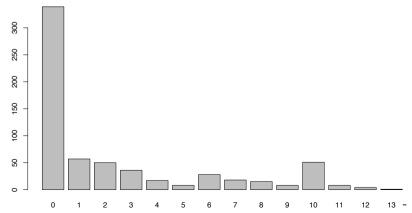
```
whitefly = read.csv('.../.../data/whitefly.csv')
head(whitefly, 4)
```

imm week block trt n live plantid ## 1 0 5 3 5 6 6 6 3 ## 2 0 1 2 3 10 2 4 ## 3 0 3 2 3 10 0 4 ## 4 0 4 2 3 10 6

- ► The live and n columns indicate how many whitefly survived and were used in the experiment respectively
- ▶ We have a fixed total and a number of surviving whitefly out of this total. Which probability distribution might be appropriate?

Plotting the whitefly data

barplot(table(whitefly\$live))

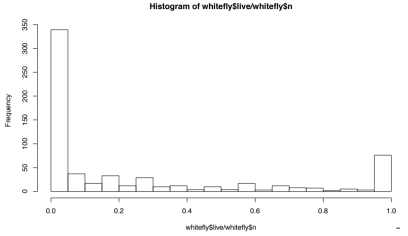


Doesn't really show the whole story as we're ignoring the total

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Second go

hist(whitefly\$live/whitefly\$n, breaks = 30)



Better, but actually there is also a covariate in the number of immature whitefly that were included (variable imm)

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Fitting a model

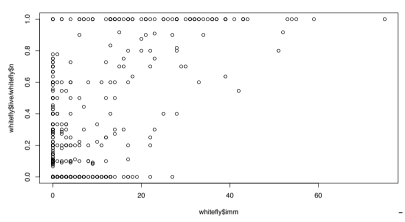
- ► Let's call *y* the number of live whitefly at the end of the experiment, and *n* the number of whitefly used in the experiment
- ► Let's fit a binomial distribution:

$$y \sim Bin(n, p)$$

- ► We know the value of *n* so it is not really a parameter but a fixed part of the data
- ▶ We need to estimate *p*
- We could use method of moments or maximum likelihood. If we use method of moments we get $\hat{p} = 0.245$

Third go

plot(whitefly\$imm, whitefly\$live/whitefly\$n)



Looks like the more immature whitefly there were, the more likely they were to die

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Fitting a better model

- ▶ What if we wanted to include imm as a covariate?
- ▶ One way would be to let $p = \alpha + \beta x$ where x is the number of immature whitefly
- \blacktriangleright We could fit this using maximum likelihood to get estimates of $\hat{\alpha}$ and $\hat{\beta}$
- ► This is now a Generalised Linear Model (GLM)
- ► The likelihood would be:

```
y = whitefly$live
n = whitefly$n
x = whitefly$imm
alpha = 0.5
beta = 0.0001
sum(dbinom(y, n, alpha + beta * x, log = TRUE))
```

```
## [1] -3330.495
```

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Fitting an even better model

▶ Suppose after using maximum likelihood we got $\hat{\alpha} = 0.2$ and $\hat{\beta} = 0.01$. Then the likelihood would be

```
alpha = 0.2
beta = 0.01
sum(dbinom(y, n, alpha + beta * x, log = TRUE))
```

```
## [1] -1971.889
```

Can anyone see any problems with this model? (Hint: suppose I wanted to predict what proportion would die when x = 100)

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The logit function

- To stop the lines going out of the range (0, 1) people often use the *logit* transformation:

$$\log \left[\frac{p}{1-p} \right] = \alpha + \beta x \text{ or } p = \frac{e^{\alpha + \beta x}}{e^{\alpha + \beta x} + 1}$$

- The latter is known as the inverse logit function
 - ▶ We now maximise the likelihood:

```
p = \exp(alpha + beta * x) / (exp(alpha + beta * x) + 1)

sum(dbinom(y, n, p, log = TRUE))
```

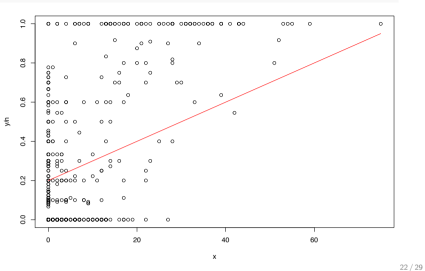
[1] -3630.322

▶ These logit and inverse logit functions are in the boot package:

```
library(boot)
sum(dbinom(y, n, inv.logit(alpha + beta * x), log = TRUE))
## [1] -3630.322
```

Plotting the fitted model

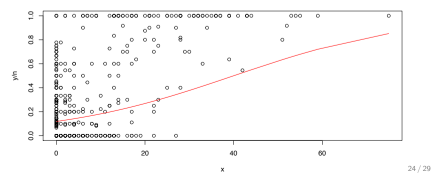
```
plot(x, y/n)
lines(x, alpha + beta * x, col = 'red')
```



Plotting the fit

- ▶ Suppose under this method we got maximum likelihood estimates of $\hat{\alpha} = -2$ and $\hat{\beta} = 0.05$
- ► A plot of the fitted values is now:

```
alpha = -2
beta = 0.05
plot(x, y/n)
lines(x, inv.logit(alpha + beta * x), col = 'red')
```



Finding the maximum likelihood values

- ► R has a function called glm to find the maximum likelihood values for us
- ▶ For binomial model with a logit *link function* we would type:

Other glms

- ► Another common glm is the Poisson, useful for count data
- ► For example, suppose we treated the number of whitefly who died as a count, and temporarily ignored the *n* values
- ▶ We could fit:

Further details about the fit

- ► The interpretation of the $\hat{\beta}$ value (the coefficient of x) is in terms of $\log odds$. A unit increase in x gives a $\exp(\hat{\beta}) \approx 1.06$ times increase in the probability of a whitefly survivign
- ➤ You'll see amongst the output something called the *deviance*. This is minus twice the log-likelihood
- ▶ It's a common measure used to compare models as the deviance for a linear regression model is just the mean square error
- Next to it you'll see the Akaike Information Criterion or AIC value, which penalises the deviance by adding on twice the number of parameters (i.e. a measure of the complexity of the model)
- ► Often, you would fit multiple models with different covariates and choose the one(s) with the smallest AIC

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A final word on glms

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- ► There are lots of different types of GLMs We can do Gamma, Negative Binomial, Beta, Inverse Gaussian, . . .
- ► Each has a link function which transforms the main parameter into an unrestricted range through which we can include covariates
- ▶ It's also simple to include extra covariates or interactions:

```
glm(y \sim x1 + x2 + x1:x2, family = poisson(link = log))
```

- Residual checks are still important, and R will create them for us
- ▶ We can get at them via e.g.

```
my_model = glm(cbind(y, n) ~ x, family = binomial(link = lot
plot(my_model)
```

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Summary

- ► Linear Regression and Generalised Linear Models are two common ways to extend standard probability distributions to include covariates
- ► We estimate the parameters via maximum likelihood using e.g. lm or glm
- ► We sometimes need to include a link function which transforms the parameters into an unrestricted range
- ► There are lots of different types of GLM for every flavour of probability distribution