Quantitative Analysis in Ecology and Evolution

The Linear Model 1: Linear regression

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4. The linear model I: Introduction and linear regression

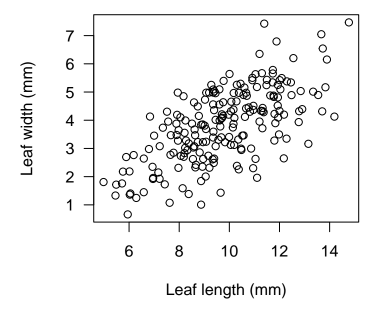
Nearly all the statistical models we will discuss in this text are forms of the linear model

$$y_i = \beta_0 + \Sigma_j x_{ij} \beta_j + \epsilon_i$$

The term β_0 (sometimes denoted α) is the *intercept*, which in the context of a linear regression gives the value of the response variable y when the predictor variable x is zero. The β_j are the coefficients ('slopes') for the predictor variables x, and the ϵ represent the *residuals*, the deviations of each data point from it's expected value based on the fitted model. In a regression, the residual is the perpendicular distance from the fitted regression line to each data point. The linear model assumes that the residuals (not the data!) are normally distributed, though minor deviations from this is not generally a problem.

In the following, we will consider a series of examples of linear models fitted to simulated data. After simulating some values of the predictor x, we define y as a function of x and add some residual variance to the data (i.e. we simulate data from the same linear model that we will eventually fit to the data.) The advantage of starting from simulated data is that we know the true values of the parameters we will try to estimate. This is very useful when we want to check that our analysis is doing what we think it is doing.

As noted above, R functions know the order of arguments. Note how the second incidence of rnorm below skips the formalities.



At this point let's talk about how to write R code. In the R chunks above you may start to notice some 'style conventions'. When we are new to R and coding, most of us write really messy code. It is now becoming mandatory to publish our code alongside our papers, and we thus have to learn to write code that is easy to read and pleasent to look at. I don't follow all the 'rules' around this, but at least I try to be consistent. For example, the strict R convention has been to use the <- operator for assignments, but I find the = easier. Note though the spaces to either side of = that makes the code easier to read. Within functions, add a space after commas.

In scatterplots, a useful change from the default is to make the y-axis labels horizontal by setting las=1. There are hundreds of ways to change the appearence of R-plots, see the wonderful world of par(). If you prefer, you can choose to learn alternative plotting frameworks such as ggplot, but these lecture notes will use R packages only when strictly needed.

The aim of regression analysis is to estimate the linear relationship between a response variable (or 'dependent variable') and one or more predictor variables ('independent variables'). The most common form of regression analysis is so-called ordinary least-square (OLS) regression, in which the regression parameters are estimated so as to minimize the square deviations of the data points from the estimated regression line. The deviations are termed *residuals*, and are assumed to be normally distributed.

```
m = lm(y\sim x)
```

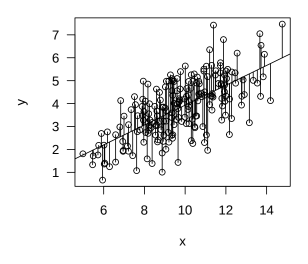
The object m now holds the fitted model. Let's extract the model coefficients and produce some plots of the residuals.

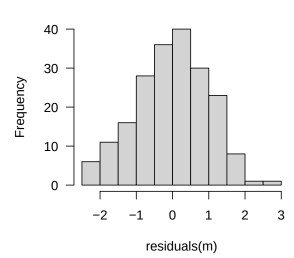
```
cf = summary(m)$coef
predvals = cf[1,1] + cf[2,1]*x

par(mfrow=c(1,2))
plot(x, y, las=1)
abline(m)
```

```
segments(x, y, x, predvals)
hist(residuals(m), las=1)
```

Histogram of residuals(m)





These residuals are fine, as is of course fully expected given that we simulated the data from the (gaussian) linear model. There are many other ways of assessing whether the model assumptions are met, see for example what happens if you call plot(m) after setting par(mfrow=c(2,2)). Notice that the plot function is *generic*, it produces a different result depending on what is fed to it. If we call plot(x,y) when both x and y are continuous variables, we get a scatterplot. If x is a factor, we get a boxplot.

Let's now have a look at the results of our linear model fit.

summary(m)

```
##
## Call:
##
  lm(formula = y \sim x)
##
##
  Residuals:
##
        Min
                   1Q
                        Median
                                              Max
    -2.45122 -0.68319
                       0.02913
##
                                0.69861
##
##
   Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
   (Intercept) -0.40114
                            0.35186
                                       -1.14
                                                0.256
##
##
   х
                 0.43330
                            0.03538
                                       12.25
                                               <2e-16 ***
##
##
                            0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' '1
##
## Residual standard error: 0.9912 on 198 degrees of freedom
## Multiple R-squared: 0.4311, Adjusted R-squared: 0.4282
## F-statistic:
                   150 on 1 and 198 DF, p-value: < 2.2e-16
```

This summary contains a lot of information. First, we can see some quantiles of the residual distribution, which confirms what we have already seen from the histogram: the residuals are fine because the median is

close to zero, the 1st and 3rd quartile are symmetrical, and the min and max values are nearly symmetrical too.

Next we see the model parameter estimates, their standard errors, a test statistic (t), and a P-value. It is tempting to look first at the P-value, the magic measure of significance and, to some, 'importance'. Before going on, let's take a moment to recall what the P-value means and how it is obtained. In the context of the linear regression above, the test statistic t is given by $t = \frac{\hat{\beta}}{SE(\beta)}$. We already know that the standard error $SE = \frac{\sigma}{\sqrt{n}}$, thus

$$t = \frac{\hat{\beta}}{\frac{\sigma(\hat{\beta})}{\sqrt{n}}}$$

The most important thing to notice here is that the sample size n is in the denominator of the expression for the standard error, so that larger sample size will lead to a smaller standard error, and thus a greater t-value.

The *P*-value is the probability of observing the observed value of the test statistic given that the null hypothesis (here, a slope of zero) is true, or $P_{obs} = Pr(T > t_{obs} = t(X_{obs}|H_0))$. In other words, it represents the probability that we would have obtained our results by chance.

Because the P-value is obtained by comparing our observed test statistic t to its known distribution, and t increases with sample size, it follows that when the sample size increases, anything will at some point be statistically significant. This is the reason why there are now increasing calls for abandoning P-values as the standard measure of statistical significance. That being said P-values do provide a 'quick and dirty' way of assessing statistical support, and can help guide our interpretation of the results. We will later return to alternative methods of evaluating statistical support, but for now we focus on the more important point: interpretation of the results needs to be done in light of the parameter estimates, their units, and their consequences within the context of the analysis/study.

EXERCISE: Use the parameter estimates to draw (by hand) the scatterplot of y vs x with the regression line, indicating the location of the intercept and roughly the correct slope.

EXERCISE: Use non-parametric bootstrapping to derive a standard error for the slope of the linear regression above. To do so, sample from the data set, fit the model, and save the sampling distribution for the slope of y on x.

[1] 0.03579301

Now, let us return to how we interpret the results of our linear regression. The slope of y on x is about 0.43. Although regression slopes are very often reported without any units, it is important to remember that the slopes in fact carry the units of both the response and predictor variables. In our example the response and predictor are both measured in mm, and the slope is therefore 0.43 mm/mm. When we report this in the text, we generally want also to report the standard error, i.e. $slope = 0.43 \pm 0.04 \ mm/mm$. Thus, in our example, the response variable increases by 0.43 mm per mm increase in the predictor. The small standard error (relative to the slope estimate) directly indicates the strong statistical support.

To facilitate further interpretation, we can also report the consequences of a realistic change in the predictor variable. Let's say that we want to know how much y changes for a one standard deviation change in x.

```
coefs = summary(m)$coef
(coefs[2,1]*(mean(x) + sd(x))) - (coefs[2,1]*mean(x))
```

[1] 0.8606095

Here, we could write in the results section that 'In the study population, corolla width increased by $0.80 \ mm$ per standard deviation increase in corolla diameter'.

As a special case, a regression where both the response and predictor variable are natural log-transformed will have a slope interpretable as an *elasticity*, which describes the % change in the response per % change in the predictor. This is another example of the nice proportional properties of the natural log.

Another important parameter in the summary table is the coefficient of determination, the r^2 . In our simple univariate regression, the r^2 is simply the square of the Pearson correlation coefficient r between the response and predictor.

```
cor(x,y)^2
```

[1] 0.4310643

The r^2 of our model is 0.431, which means that 43.1% of the variance in y is explained by x. In general, it is often nice to report the r^2 directly as a percent (i.e. ×100). To understand why the r^2 gives the % variance explained, note that the r^2 can be computed as the variance in the predicted values \hat{y} ,

$$V(\hat{y}) = X\beta$$

divided by the total variance in the response varible V(y).

```
y_hat = coefs[1,1] + coefs[2,1]*x
var(y_hat)
```

[1] 0.7406487

```
var(y_hat)/var(y)
```

[1] 0.4310643

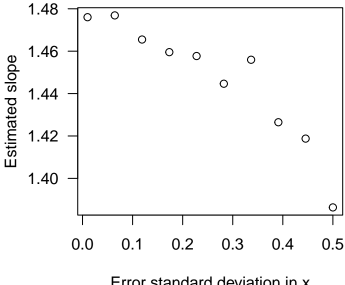
Another way to compute the variance explained by a predictor is $V(x) = \beta_x^2 \sigma_x^2$, where β_x is the parameter estimate (regression slope) for predictor x, and σ_x^2 is the variance of the predictor.

```
coefs[2,1]^2*var(x)
```

[1] 0.7406487

Exercise: How error in x- and y-variables affect the slope

The standard linear model assumes that the predictor variable is measured without error. When there is measurement error, this can lead to a bias in the estimated slope. Simulate data with measurement error in the predictor, and produce a plot showing the effect on the estimated slope.



Error standard deviation in x

For a simple model like this, the expected attenuation bias (downward bias) in the slope can be estimated by the reliability ratio

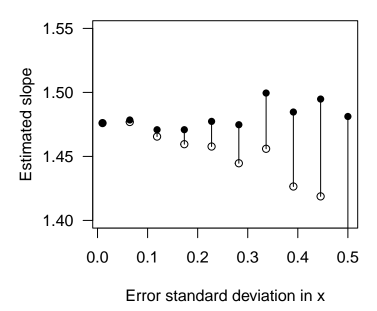
$$K = 1 - \frac{\sigma_{me}}{\sigma_x}$$

where σ_{me} is the measurement error variance and σ_x is the variance in the predictor x. We can thus obtain a corrected slope as

$$\beta' = \frac{\beta}{K}$$

Try to correct your estimated slopes in this way, and a produce a plot showing both the estimated and the corrected slope connected by line segments.

- ## [1] 2.562152e-05 1.064084e-03 3.621491e-03 7.697843e-03 1.329314e-02
- [6] 2.040738e-02 2.904057e-02 3.919270e-02 5.086378e-02 6.405380e-02 ##



Exercise: fitting a linear regression to real data

Choose any dataset you may have involving a continuous response variable and an continuous predictor. Fit a simple linear regression, interpret the results, produce a nice figure including the fitted regression line, and write simple methods and results presenting the analysis and results.

If you don't have any data, use the dataset bird_allometry in the datasets folder. This dataset contains body mass and brain mass for males and females of different bird species. The scaling of brain size (or other body parts) with body size is referred to as the study of allometry, and you may want to read about these analyses before fitting your models. As a hint, the scaling of parts of a body with body size is expected to follow a power-law relationship on the form $y = ax^b$, which can be linearized through the logarithmic transformation $log(y) = log(a) + b \times log(x)$.

```
birds = read.csv("datasets/allometry/bird_allometry.csv")
head(birds)
```

```
##
                  Genus_Species Sex brain_mass body_mass
## 1
             Accipiter_gentilis
                                        7.686143 1049.1571
## 2
                                        7.618500
             Accipiter_gentilis
                                                  678.2833
## 3
                Accipiter_nisus
                                   f
                                        3.112797
                                                  252.1263
## 4
                Accipiter_nisus
                                                  136.1441
                                   \mathbf{m}
                                        2.637390
             Accipiter_striatus
                                   f
                                        5.700000
                                                  520.0000
## 6 Acridotheres_cristatellus
                                        2.310000
                                                  122.3800
```