Date: 16 lectures, MT09.
1. Course

The course develops the theory of statistical methods, and introduces students to the analysis of data using a statistical package. The main topics of the MT09 course are: Simulation based inference, Practical aspects of linear models, and Logistic regression and generalized linear models.

Reading (Michaelmas Term):

Further Reading:
D. Lunn, Lecture notes http://www.stats.ox.ac.uk/~dlunn/BS1_05/BS1_MT05.htm

1.1. Website. Goto the Statistics Department website and access
http://www.stats.ox.ac.uk/current_students/bammath/course_material
Select BS1a Applied Statistics I, or straight to
http://www.stats.ox.ac.uk/~nicholls/bs1a/
Lecture notes, problem sheets, etc. are linked from that page.

1.2. Classes. Bookings for classes will be made in the first lecture. There will be 6 problems classes this term, each lasting for 1 hour. They will be held in weeks 3 to 8. Details of classes will be linked from the website.

1.3. Assessment and Practicals. There are three assessed BS1a practicals and one assessed BS1b practical. These contribute 8.5% each of a total 34% practical component in your BS1 mark. The BS1 paper in finals is a 2 hour paper worth 66% of your BS1 mark.

The practical problems for you to solve will be made available in the practical teaching labs. The deadlines for handing in your answers are as follows:

First assessed practical - 12 noon Tuesday, week 8 Michaelmas Term 2009
Second assessed practical - 12 noon, Tuesday, week 2 Hilary Term 2010
Third assessed practical - 12 noon, Tuesday , week 7 Hilary Term 2010
Fourth assessed practical - 12 noon, Friday, week 1 Trinity Term 2010

In order to solve the practical problems you need to learn R. We will have a session introducing R itself, in the lab, in week 3, and then, for each of the four assessed practicals above, a session where we show you some code relevant to the
particular practical. Charalambos Loizides is running these teaching-practicals and will provide information linked from

http://www.stats.ox.ac.uk/~loizides

Each session is presented on two evenings (in the OUCS). You may choose which evening you wish to attend. The BS1 practical teaching sessions are held 5:30-7:30 Tuesday and Wednesday evenings, this term in weeks 3, 5 and 8 and and next term in weeks 2 and 7.

1.4. The R package. The R package is available free of charge on the web. Go straight to

www.stats.bris.ac.uk/R/

and download and install R. There is a FAQ, linked from the left margin, full of useful information. Once installed, An Introduction to R is a good place to start. Again, from the left margin,

Manuals → An Introduction to R.

1.5. Guide.

- Read the lecture notes and interleave your own notes.
- We will begin by reviewing and extending some of the Part A material on Linear Models.
- Linear models lead naturally to looking at Diagnostics. Is the model plausible? If not, why not?
- The next topic is Generalised Linear Models. In a normal linear model the linear predictor $X^T\beta$ is the mean. We keep this model element, but link it to the mean of some other distribution. This greatly extends the application domain of the linear model. We introduce the theory and look at practical problems.
- The final topic is Monte Carlo testing. This is a framework that allows us to test relatively complex models.
2. Linear Regression

2.1. Normal linear models. Does $y$ get bigger when there is more $x'$? Normal linear models are attractive because they are simple, and in some respects easy to interpret. If, for a given observation process, they happen to give a correct or near-correct description of the distribution of the response $y$, and its dependence on predictive factors $x$, then they are hard to beat.

We choose to model a randomly variable response $Y = y$ as a linear function of explanatory variables $x_1, x_2, ..., x_p$. At the $i$th observation, we set the explanatory variables to values $x_i = (x_{i1}, x_{i2}, ..., x_{ip})$ (with $x_i$ a row vector), and measure a response $Y_i = y_i$. Under a normal linear model, the expected response is given as a linear combination of the explanatory variables, weighted by parameters $\beta = (\beta_1, \beta_2, ..., \beta_p)^T$,

$$y_i = \sum_{j=1}^{p} \beta_j x_{ij} + \epsilon_i,$$

with $\epsilon_i \sim N(0, \sigma^2)$ iid normal errors for $i = 1, 2, ..., n$. If $\epsilon = (\epsilon_1, \epsilon_2, ..., \epsilon_p)^T$ then $y_i = x_i \beta + \epsilon_i$. If $y = (y_1, y_2, ..., y_n)^T$, and $X$ is the $n \times p$ design matrix with rows $x_i$, then our linear model has matrix form

$$Y = X \beta + \epsilon.$$

We will write $X_j$, $j = 1, 2, ..., p$ for the columns of $X = (X_1, X_2, ..., X_p)$. We will use $y$ and $\epsilon$ to denote both a column vector of responses, as above, and a single generic realization of the scalar response $Y = y$ etc. For example if we write down a model in terms of scalars $y$ and $\epsilon$,

$$y = \alpha + \gamma_1 x_1 + ... + \gamma_m x_m + \epsilon$$

omitting the $i = 1, 2, ..., n$ subscript, we have in mind $y = X \beta + \epsilon$ (now vectors) with $\beta_1 = \alpha$ and $\beta_i = \gamma_{i+1}$ and $p = m + 1$. In this example, $X_1 = 1_n 1$ that is, the first column if $X$ is a column of ones and corresponds to the explanatory variable for the intercept parameter $\alpha$.

The nomenclature assumes that we set the values of the explanatory variables and measure the response. When we choose the $x_i$, $i = 1, 2, ..., n$ we are designing the experiment. We will see that not all designs are equally good. The subject of experimental design is one we will just touch on.

From time to time, we will assume that the columns of $X$ are linearly independent vectors, that is, the explanatory variables are not linearly dependent. If they are, we can throw out linearly dependent columns till we have a linearly independent set; the discarded columns tell us nothing new about the measurement context. This issue comes up, for example, in Section ???. We are, as a consequence, often assuming $p \leq n$, that is, we have more measurements than parameters.

The number one problem for interpreting linear models arises from correlation between explanatory variables. You can think of this as a kind of weak linear dependence between variables, and groups of variables.

Example 2.1. The dataset cig contains measurements of the carbon-monoxide (variable CO), tar and nicotine content and tobacco weight for $n = 25$ cigarettes. The data are plotted in Figure 1. In the normal linear model $CO \sim 1 + \text{Nicotine} + \text{Tar} + \text{Weight}$ the response is CO and all the other variables (including intercept) are
explanatory. This notation (which comes from R) means we are fitting the model

\[ y = \beta_1 + x_2\beta_2 + x_3\beta_3 + x_4\beta_4 + \epsilon \]

with \( y = \text{CO} \) output for one cigarette, \( x_1 = 1 \) and \( x_2, x_3 \) and \( x_4 \) respectively the measured nicotine and tar content and weight of the cigarette.

```r
> loc<-’http://www.stats.ox.ac.uk/~nicholls/bs1a/data/cigarettes.txt’
> cig<-read.table(loc,header=T) #load the data from a file
> names(cig) #inspect the data
[1] "Brand"  "Tar"  "Nicotine" "Weight" "CO"
> dim(cig) #number of observations by number of variables
[1] 25 5
> head(cig)
   Brand   Tar Nicotine  Weight   CO
1   Alpine 14.1   0.86  0.9853 13.6
2 Benson&Hedges 16.0   1.06  1.0938 16.6
3    BullDurham 29.8   2.03  1.1650 23.5
4     CamelLights 8.0   0.67  0.9280 10.2
5        Carlton 4.1   0.40  0.9462  5.4
6  Chesterfield 15.0  1.04  0.8885 15.0
> pairs(cig[,c("Tar","Nicotine","Weight","CO")])
```
The following partial R-output gives the fitted MLE parameter values for this model.

```R
> cig.lm<-lm(CO~Nicotine+Tar+Weight,data=cig) #ignoring brand for now
> summary(cig.lm)

Coefficients:
                   Estimate Std. Error t value  Pr(>|t|)
(Intercept)       3.2022     3.4618   0.925  0.365464
Nicotine          -2.6317     3.9006  -0.675  0.507234
Tar                0.9626     0.2422   3.974  0.000692 ***
Weight             -0.1305     3.8853  -0.034  0.973527

The columns give estimates \( \hat{\beta} \), standard errors (ie \( \text{estimates of var}(\hat{\beta}) \)), \( t \)-values for the test that the parameter is zero (Estimate/Std. Error) and \( p \) values for those test statistics. Notice that the parameter for the variable Nicotine is negative. Look at the pairs plot. How does this make sense? The problem is that tar and nicotine are correlated. Tar is explaining the bulk of the variation in CO making the contribution from nicotine hard to interpret. Which variables are explanatory? Is there a minimal set? My guess would be that Tar gives rise to CO and Tar separately predicts Nicotine so Nicotine is only indirectly linked to CO. We will return to this sort of problem later in the course.

For the linear model theory to go through, the response must be a linear function of the parameters \( \beta \). It need not be a linear function of the explanatory variables. Also, we may find that some function of the response is a linear function of the explanatory variables.

**Example 2.2.** Consider the trees data. What variables, and what functions of those variables are important on physical grounds? A lattice plot of the logged trees data is shown in Figure 2.

```R
> data(trees) #bring the data into the workspace
> names(trees) #what are the variable names?
[1] "Girth" "Height" "Volume"
> dim(trees) #n=31 observations
[1] 31 3
> pairs(log(trees),main='logged trees data') #Make the lattice plot
```

If \( v \) is the volume, and \( h \) and \( g \) are the height and girth, a natural model on physical grounds would be

\[
v = \eta h^{1+\beta_2} g^{2+\beta_3} \gamma
\]

with \( \eta \) a fixed constant and \( \gamma \) varying randomly about one. The idea of using a multiplying error \( \gamma \) here is that large volume trees have a higher volume variance than lower volume trees. We will investigate the linear model

\[
y = \beta_1 + \beta_2 x_2 + \beta_3 x_3 + \epsilon
\]

with \( y = \log(v/hg^2) \), \( \beta_1 = \log(\eta) \), \( x_2 = \log(h) \), \( x_3 = \log(g) \) and \( \epsilon = \log(\gamma) \). In the lattice plot of Figure 2, the logged data has little curvature, skew or uneven distribution of \( X \)-values (bunching by height or girth) so a linear model seems acceptable.
The model we are fitting has \( n = 31 \) observations and \( p = 3 \) parameters. Fitting the model we obtain

\[
\text{trees.lm1} <- \text{lm}(\log(\text{Volume}/(\text{Height} \times \text{Girth}^2))^2 + \log(\text{Height}) + \log(\text{Girth}), \text{data=trees})
\]

```
> names(trees.lm1)
[1] "coefficients" "residuals" "effects" "rank"
[5] "fitted.values" "assign" "qr" "df.residual"
[9] "xlevels" "call" "terms" "model"
```

```
> summary(trees.lm1)
```

Coefficients:

|                | Estimate | Std. Error | t value | Pr(>|t|)  |
|----------------|----------|------------|---------|-----------|
| (Intercept)    | -6.63162 | 0.79979    | -8.292  | 5.06e-09  *** |
| log(Height)    | 0.11712  | 0.20444    | 0.573   | 0.571     |
| log(Girth)     | -0.01735 | 0.07501    | -0.231  | 0.819     |

```

Reading off the estimated parameters, \( \beta = (\log(\eta), \beta_2, \beta_3) \), so \( \tilde{\eta} = \exp(-6.6) \) etc, we arrive at the model

\[
v = \exp(-6.6)h^{1.12}g^{1.98}\gamma,
\]

with \( \log(\gamma) \sim N(0, 0.08^2) \).
2.2. **Estimators.** Given a normal linear model with data $y$ and an $n \times p$ design $X$, the log-likelihood for $\beta$ is

$$\ell(\beta, \sigma^2; y) = -\frac{n}{2} \log(\sigma^2) - \frac{1}{2\sigma^2} \sum_{i=1}^{n} (y_i - x_i \beta)^2.$$ 

Let $\text{RSS}(\beta) = (y - X\beta)^T (y - X\beta)$ denote the residual sum of squares. The maximum likelihood estimator $\hat{\beta}$ for $\beta$ minimises the RSS, for any fixed $\sigma$. Denote by

$$\text{col}(X) = \{ z \in \mathbb{R}^n : z = X\beta, \beta \in \mathbb{R}^p \}$$

the column span of $X$. We suppose to begin with that $X$ is rank $p$, so $\text{col}(X)$ is a $p$-dimensional linear subspace of $\mathbb{R}^n$. Since $\hat{\beta}$ minimises the RSS, $X\hat{\beta}$ is that point in $\text{col}(X)$ lying closest to $y$. The point $\hat{y} = X\hat{\beta}$ therefore lies at the orthogonal projection of $y$ into $\text{col}(X)$. Since $y - \hat{y}$ is orthogonal to all vectors in $\text{col}(X)$, we have the $p$ normal equations

$$X^T (y - X\hat{\beta}) = 0$$

which fix the values of the $p$ parameters $\beta$. Now if $X$ has $p$ linearly independent columns then the $p \times p$ matrix $X^T X$ is rank $p$ and invertible. It follows that

$$\hat{\beta} = (X^T X)^{-1} X^T y$$

gives the MLE in terms of the design matrix and observations. This is also the least-squares estimator for $\beta$, since it minimizes the RSS.

We will shortly derive an unbiased estimator for the error variance $\sigma^2$. However, recall that when we make likelihood ratio tests we substitute parameter MLEs into the likelihood, and it is for this reason that we will later need the MLE for $\sigma^2$ and the value of the maximized log-likelihood. Since $\hat{\beta}$ is the MLE for all $\sigma^2$, the MLE $\hat{\sigma}^2_{\text{MLE}}$ for the error variance maximises

$$\ell(\hat{\beta}, \sigma^2; y) = -\frac{n}{2} \log(\sigma^2) - \frac{1}{2\sigma^2} (y - X\hat{\beta})^T (y - X\hat{\beta})$$

so the MLE is

$$\hat{\sigma}^2_{\text{MLE}} = \frac{\text{RSS}}{n}$$

This is a biased estimator (RSS means RSS($\hat{\beta}$) from here on). The value of the log-likelihood at the joint MLE, is

$$\ell(\hat{\beta}, \sigma^2_{\text{MLE}}; y) = -\frac{n}{2} \log(\text{RSS}/n) - n/2$$

since the two factors of $(y - X\hat{\beta})^T (y - X\hat{\beta})$ cancel.

2.3. **Properties of Estimators.** Let $\tilde{y} = X\hat{\beta}$ give the estimated response. Define the $n \times n$ hat matrix $H$

$$H = X (X^T X)^{-1} X^T$$

so that $\tilde{y} = Hy$. The hat matrix is a projection operator, projecting $y$ into the column space of $X$, so $H = HH$ and $H$ is symmetric. Define the vector of residuals, $e = y - \tilde{y}$. The residual sum of squares, is the squared norm of the residuals, $\text{RSS}(\hat{\beta}) = e^T e$. Under the normal linear model, the residuals $e$ and the estimated response $\tilde{y}$ are actually independent. We begin by showing that they are uncorrelated.

**Exercise**: Show that $e^T e + \tilde{y}^T \tilde{y} = y^T y$ ($y, \tilde{y}$ and $e$ form a right-angle triangle).
Exercise: Show that $H$ has $p$ eigenvalues equal one and $n - p$ equal zero.

For generic random vectors $U = (U_1, ..., U_p)^T$ and $W = (W_1, ..., W_n)^T$ with means $\mu_U = \mathbb{E}(U)$ and $\mu_W = \mathbb{E}(W)$ denote by

$$\text{cov}(U, W) = \mathbb{E}((U - \mu_U)(W - \mu_W)^T)$$

the $p \times n$ covariance matrix with entries $\text{cov}(U, W)_{i,j} = \text{cov}(U_i, W_j)$. Notice that $(U - \mu_U)(W - \mu_W)^T$ is an outer product.

Exercise Let $C, C'$ be constant (i.e., non-random) $p, n$-component vectors. Show that $\text{cov}(U + C, W + C') = \text{cov}(U, W)$.

The variance matrix

$$\text{var}(U) = \mathbb{E}((U - \mu_U)(U - \mu_U)^T)$$

is a symmetric $p \times p$ matrix with entries $\text{var}(U)_{i,j} = \text{var}(U_i, U_j)$. The variance matrix of $Y$ is $\text{var}(Y) = \text{var}(X\beta + \epsilon)$, $X\beta$ is a constant and $\text{var}(\epsilon) = \sigma^2I_n$, so $\text{var}(Y) = \sigma^2I_n$.

Exercise Let $L$ and $M$ be matrices of suitable dimension. Show that $\text{cov}(LU, MW) = L\text{cov}(U, W)M^T$ and $\text{var}(LU) = L\text{var}(U)L^T$.

Under the normal linear model, the estimated responses $\hat{Y}$ and residuals $e$ are uncorrelated: $\text{cov}(\hat{Y}, Y - \hat{Y}) = \text{cov}(HY, (I_n - H)Y)$ and the RHS there is

$$H\text{cov}(Y, Y)(I_n - H)^T = \sigma^2I_n(H - HH^T)$$

which is zero, that is, $\text{cov}(\hat{Y}, Y - \hat{Y}) = 0_{n,n}$ where $0_{n,n}$ is an $n \times n$ matrix of zeros. In fact, as we will see, $\hat{Y}$ and $e$ are independent. This is sometimes asserted on the basis that $\hat{Y}$ and $e$ are normal, with zero covariance, so they are independent. Beware: this kind of argument works when the two quantities are jointly normal. It is easy to see that $\hat{Y}$ and $e$ are not jointly normal (the covariance matrix for each is singular). The conclusion is nevertheless correct in this case. [END L1]

We next compute the distribution of our parameter estimate $\hat{\beta}$. The MLE $\hat{\beta} = (X^TX)^{-1}X^TY$ is a linear combination of the normal random variables $Y = (Y_1, ..., Y_n)$. In general, if $W \sim N(\mu_W, \Sigma)$, so that $W$ is an $n$-component multivariate normal (MVN) random vector (r.v.) with positive definite $n \times n$ variance matrix $\text{var}(W) = \Sigma$, and $L$ is a $p \times n$ matrix with $p \leq n$ linearly independent rows, then the $p$-component r.v. $LW \sim N(L\mu_W, L\Sigma L^T)$. The conditions are then to ensure $L\Sigma L^T$ is positive definite, and hence invertible.

Exercise Show that $\mathbb{E}(\hat{\beta}) = \beta$ and $\text{var}(\hat{\beta}) = \sigma^2(X^TX)^{-1}$, so that $\hat{\beta} \sim N(\beta, \sigma^2(X^TX)^{-1})$.

Ans: This is the generic case with $L = (X^TX)^{-1}X^T$ and $W = Y$, so $\text{var}(\hat{\beta}) = (X^TX)^{-1}X^T\text{var}(Y)((X^TX)^{-1}X^T)$, or $\text{var}(\hat{\beta}) = \sigma^2(X^TX)^{-1}X^TX(X^TX)^{-1}$ which is $\text{var}(\hat{\beta}) = \sigma^2(X^TX)^{-1}$. The result for the MVN distribution follows from the result for $LW$ above.

Let us now show that $e$ and $\hat{Y}$ are independent. It follows that $\hat{\beta}$ and RSS are independent, since the estimator $\hat{\beta} = (X^TX)^{-1}X^T\hat{Y}$ and RSS = $e^Te$. We use the independence properties when we construct test statistics in the next section.
Let $e_1, ..., e_p$ be a fixed orthonormal basis for the column space of $X$. Extend this basis to $e_1, ..., e_p, e_{p+1}, ..., e_n$, an orthonormal basis for $R^n$. The vectors $e_{p+1}, ..., e_n$ are orthogonal to the column vectors of $X$. Expand the $n$-component random vector $Y$ in this basis,

$$Y = Z_1 e_1 + ... + Z_n e_n.$$  

We should think of $Z_i$ as a random function of $Y$, with $Z_i = e_i^T Y$. Now $He_i = e_i$ for $i = 1, 2, ..., p$ since these vectors are in $\text{col}(X)$. On the other hand $He_i = 0$ for $i = p+1, ..., n$ (since $X^T e_i = 0$ for $i > p$). It follows that

$$\hat{Y} = Z_1 e_1 + ... + Z_p e_p,$$

since $HY = H(Z_1 e_1 + ... + Z_p e_p)$. Now $e = Y - \hat{Y}$ so 

$$e = Z_{p+1} e_{p+1} + ... + Z_n e_n.$$ 

The weights $Z_i = e_i^T Y$ are distributed $Z_i \sim N(e_i^T E(Y), \sigma^2)$. For $i = p+1, p+2, ..., n$, they are mean zero, since $E(Y) = X\beta$, and $e_i^T X = 0_{1,p}$. They are uncorrelated, since $\text{cov}(Z_i, Z_j) = e_i^T \text{cov}(Y,Y)e_j$ which is zero for $i \neq j$, since $\text{var}(Y) \propto I_n$. Taking $i = j$ we have $\text{var}(Z_i) = \sigma^2$. Since they are also jointly normal, they are independent.

The estimated response $\hat{Y}$ and the residuals $e = Y - \hat{Y}$ are functions of the two non-overlapping sets, $(Z_1, ..., Z_p)$ and $(Z_{p+1}, ..., Z_n)$, of mutually independent random variables, so $\hat{Y}$ and $e$ are independent under the normal linear model.

We can now read off the distribution of RSS, and get an unbiased estimator for $\sigma$. Since $RSS = e^T e$,

$$RSS = Z_{p+1}^2 + ... + Z_n^2.$$ 

Since $Z_i/\sigma \sim N(0,1)$ for $i = p+1, p+2, ..., n$, and $RSS/\sigma^2 = (Z_{p+1}/\sigma)^2 + ... + (Z_n/\sigma)^2$, with $(Z_i/\sigma)^2 \sim \chi^2(1)$ mutually independent rv each having a chi-squared distribution with one degree of freedom, it follows that

$$RSS/\sigma^2 \sim \chi^2(n-p),$$

under $H0$. Now if $A \sim \chi^2(r)$ then $E(A) = r$ so $E(RSS/\sigma^2) = n - p$ and

$$s^2 = \frac{RSS}{n-p}$$

is an unbiased estimator for $\sigma^2$. It follows that $\hat{\sigma}_{MLE}^2 = \frac{RSS}{n}$ is biased (but it is also asymptotically unbiased, as it is a MLE).

2.4. Tests. We would like now to consider a collection of tests on the parameters of a normal linear regression. We would like to test for the significance of a parameter, of a group of parameters, test for parameters to be equal, or greater than one another, and for properties of linear combinations of parameters.

The test for significance of a single parameter, we know. Suppose we want to test $H0 : \beta_k = 0$ against $H1 : \beta_k \neq 0$ for some particular $k$ from 1 to $p$. Under $H0$,

$$\hat{\beta}_k \sim N(0,1)$$

$$\sqrt{\hat{s}^2 (X^T X)^{-1}} \sim N(0,1)$$

where

$$\hat{s}^2 = \frac{RSS}{n-p}.$$
and $\text{RSS}/\sigma^2 \sim \chi^2(n-p)$, so writing $s^2 = \text{RSS}/(n-p)$,

$$t = \frac{\hat{\beta}_k}{\sqrt{s^2(X^TX)^{-1}_{k,k}}} \times \sqrt{\frac{\sigma^2(n-p)}{\text{RSS}}}$$

$$= \frac{\hat{\beta}_k}{s\sqrt{(X^TX)^{-1}_{k,k}}}$$

is a suitably scaled ratio of independent standard normal and $\chi^2$ random variates. Under the null, $t$ is a realisation of a Student’s-$t$ distributed random variable $T$,$$
T \sim t(n-p).
$$

The $p$-value for a two sided test is $2(1 - \Pr(T < |t|))$.

When we test for the significance of a group of parameters we use a test called an $F$-test. If there is just one parameter in the group, then the $F$-test reduces to the $T$-test we just described. Suppose we have a conjecture that there is no linear relation between the response $y$ and the last $k$ explanatory variables, $x_{p-k+1}, \ldots, x_p$. We want to test $H_0 : \beta_{p-k+1} = 0, \beta_{p-k+2} = 0, \ldots, \beta_p = 0$ against $H_1 :$ at least one of the last $k$ parameters is non-zero. Under $H_0$, with $\beta = \beta^{(0)}$ say, we are fitting the normal linear model

$$y = \sum_{i=1}^{p-k} \beta^{(0)}_i x_i + \epsilon.$$

Let $\tilde{X}$ be a matrix made up of the first $p-k$ columns of $X$. Under $H_0$, $y = \tilde{X}\beta^{(0)} + \epsilon$ with $\beta^{(0)}$ a $(p-k) \times 1$ vector, $\beta^{(0)} = (\beta_1, \ldots, \beta_{p-k})$. When we fit this model we get $\hat{\beta}^{(0)} = (\tilde{X}^T \tilde{X})^{-1} \tilde{X}^T y$. Let $H^{(0)} = \tilde{X}(\tilde{X}^T \tilde{X})^{-1} \tilde{X}^T$ be the hat matrix for the linear model with design matrix $\tilde{X}$. Let $\tilde{Y}^{(0)} = H^{(0)} Y$ and

$$\text{RSS}^{(0)} = (Y - \tilde{Y}^{(0)})^T (Y - \tilde{Y}^{(0)}).$$

Under $H_0$, the MLE for $\sigma^2$ is

$$\hat{\sigma}^2_{MLE,0} = \frac{\text{RSS}^{(0)}}{n}.$$ 

The dimension of parameter space under the null is $p-k+1$ (ie, $\beta_1, \ldots, \beta_{p-k}, \sigma^2$).

Under $H_1$, with $\beta = \beta^{(1)}$, we are fitting the normal linear model

$$y = \sum_{i=1}^{p} \beta^{(1)}_i x_i + \epsilon.$$

This is the usual setup, with $Y = X \beta^{(1)} + \epsilon$, $\hat{\beta}^{(1)} = (X^T X)^{-1} X^T y$, $\tilde{Y}^{(1)} = HY$,

$$\text{RSS}^{(1)} = (Y - \tilde{Y}^{(1)})^T (Y - \tilde{Y}^{(1)}),$$

and

$$\hat{\sigma}^2_{MLE,1} = \frac{\text{RSS}^{(1)}}{n}.$$ 

The dimension of parameter space under the alternative is $p+1$.

We can now give the Likelihood Ratio Test (LRT) statistic $\Lambda$ for $H_0$. Substituting the local values into the expression for $\ell(\hat{\beta}, \hat{\sigma}^2_{MLE}; y)$, which we gave at the
end of Section 2.2, we get
\[ \Lambda(Y) = -2(\ell(\hat{\beta}(0), \hat{\sigma}^2_{MLE,0}; Y) - \ell(\hat{\beta}(1), \hat{\sigma}^2_{MLE,1}; Y)) = n \log(RSS(0)) - n \log(RSS(1)). \]

We reject \( H_0 \) when the likelihood ratio statistic \( \Lambda \) falls in the critical region. We know from the Neyman-Pearson theorem that this region has the form \( C_1 = \{y : \Lambda(y) > C \} \). Asymptotically in \( n \), \( \Lambda \) has as \( \chi^2 \) distribution with \( k \) degrees of freedom, and this would give an approximate test for \( H_0 \). However, we will see that the LR T statistic is a strictly increasing function of another statistic, \( F(y) \), for which we possess an exact distribution. This leads to an exact test with the same critical region as the LR T: if \( C' \) is chosen so that, under \( H_0 \), \( F(Y) > C' \) with probability \( 1 - \alpha \), and \( C \) is chosen so that \( \Lambda(Y) > C \) with probability \( 1 - \alpha \), then \( F(Y) > C' \) if and only if \( \Lambda(Y) > C \) (imagine sorting the states \( y \) by \( F(y) \) and by \( \Lambda(y) \) - you get the same order so the threshold is set at the same states).

Consider the \( F \)-statistic,
\[ F(y) = \frac{(RSS(0) - RSS(1))}{RSS(1)/(n - p)}. \]

The corresponding random variable \( F(Y) \) has a \( \chi^2 \) distribution under the null hypothesis in which the last \( k \) parameters are zero. The \( F \)-distribution is new to us. If \( A \sim \chi^2(a) \) and \( B \sim \chi^2(b) \) are two independent \( \chi^2 \) r.v.'s with \( a \) and \( b \) degrees of freedom respectively, then the new r.v.
\[ F = \frac{(A/a)}{(B/b)} \]
has a \( F(a,b) \)-distribution on \( F > 0 \). This property defines the distribution. The mean of \( F \sim F(a,b) \) is \( b/(b - 2) \) so, under the null, the mean of \( F \sim F(k,n - p) \) is \( (n - p)/(n - p - 2) \) for \( n - p > 2 \), or about 1 for \( n \gg p \). The quantiles of an \( F \)-distribution with \( k \) numerator and \( n - p \) denominator degrees of freedom are known. Let \( F_{1-\alpha}(k,n - p) \) be the \( 1 - \alpha \) quantile of \( F(k,n - p) \). We reject \( H_0 \) at significance level \( \alpha \) if \( F(y) > F_{1-\alpha}(k,n - p) \).

How do we know \( F \) has the properties we claim for it? First, it is easy to check that
\[ \Lambda = n \log \left( 1 + \frac{k}{(n - p)F} \right) \]
with \( n > p > k > 0 \), so \( \Lambda \) is a strictly increasing function of \( F \), and this test based on \( F \) is indeed a LRT. Secondly,
\[ \frac{RSS(1)}{\sigma^2} \sim \chi^2(n - p) \]
and
\[ \frac{RSS(0) - RSS(1)}{\sigma^2} \sim \chi^2(k). \]
The former we know, and the latter we demonstrate shortly. Thirdly, \( RSS(1) \) and \( RSS(0) - RSS(1) \) are independent, again, something we will verify. Finally, since \( F \) is the ratio of suitably scaled and independent \( \chi^2(n - p) \) and \( \chi^2(k) \) r.v. it follows that \( F \) has an \( F(k,n - p) \) distribution, by the definition of this distribution given above.
What is the intuition here? The question which a LRT answers is, is there evidence that the data fits the $H_1$ model $Y = X\beta^{(1)} + \epsilon$ better than the $H_0$ model $Y = \tilde{X}\beta^{(0)} + \epsilon$? Tests which look for significant changes in the RSS, such as the $F$-test above, are called ANOVA, short for Analysis of Variance. When we fit the more complex model, $H_1$ above, there will be a reduction in the residual sum of squares compared to the residual sum of squares we get when we fit the simpler model $H_0$. If $H_0$ is good, then the fractional improvement $(\text{RSS}^{(0)} - \text{RSS}^{(1)})/\text{RSS}^{(1)}$ in the RSS is slight. However, if we add lots of explanatory variables in $H_1$, so that $\dim(\text{col}(X))$ approaches $n$, then we will see a big drop in the RSS, towards zero. In order to account for this, the ratio $(\text{RSS}^{(0)} - \text{RSS}^{(1)})/\text{RSS}^{(1)}$ must be weighted by the fractional change $(n - p)/k$ in the number of degrees of freedom. Now large values of $F$ are a sign that that the added parameters in $H_1$ are reducing the estimated variance by an amount which is too great to be put down to chance.

We need to verify the second and third properties above. We are interested in the distribution of $F$ under $H_0$, where $E(Y) = \tilde{X}\beta^{(0)}$. We will modify the expansion

$$Y = Z_1\epsilon_1 + ... + Z_n\epsilon_n.$$  

As before, $\epsilon_1, ..., \epsilon_n$ are an orthonormal basis for $R^n$, and $e_1, ..., e_p$ are an orthonormal basis for $\text{col}(X)$. We can choose this basis so that there is a first group $e_1, ..., e_{p-k}$ of vectors spanning the first $p - k$ columns of $X$ and a second group $e_{p-k+1}, ..., e_p$ completing the basis $e_1, ..., e_p$ for $\text{col}(X)$ (and notice $e_{p-k+1}, ..., e_p$ are not a basis for $\text{span}(X_{p-k+1}, ..., X_p)$ unless $\text{span}(X_{p-k+1}, ..., X_p) \perp \text{span}(X_1, ..., X_{p-k})$). Since $\hat{Y}^{(0)} = H^{(0)}Y$ $H^{(0)}$ projects into space spanned by the first $p - k$ columns of $X$, we must have $He_j = 0$ for $j > p - k$, so

$$\hat{Y}^{(0)} = Z_1\epsilon_1 + ... + Z_{p-k}\epsilon_{p-k}.$$  

Similarly, $\hat{Y}^{(1)} = HY$ with $He_j = 0$ for $j > p$, so

$$\hat{Y}^{(1)} = Z_1\epsilon_1 + ... + Z_p\epsilon_p.$$  

Now $Y - \hat{Y}^{(1)} = Z_{p+1}\epsilon_{p+1} + ... + Z_n\epsilon_n$, so

$$\text{RSS}^{(1)} = Z_{p+1}^2 + ... + Z_n^2.$$  

Similarly, $\text{RSS}^{(0)} = Z_{p-k+1}^2 + ... + Z_n^2$. It follows that

$$\text{RSS}^{(0)} - \text{RSS}^{(1)} = Z_{p-k+1}^2 + ... + Z_n^2.$$  

The weights $Z_i$, $i = 1, 2, ..., n$ are mutually independent, since they are jointly normal, with zero covariance, exactly as before. They are distributed as $Z_i \sim N(e_i^T E(Y), \sigma^2)$ with $E(Y) = [X_1, ..., X_{p-k}]\beta$ so $Z_i \sim N(0, \sigma^2)$ for $i = p - k + 1, ..., n$ (i.e., not just $i = p + 1, ..., n$ as before).

We can see that $\text{RSS}^{(0)} - \text{RSS}^{(1)}$ and $\text{RSS}^{(1)}$ are independent rv, since they are functions of disjoint sets of independent rv. Also, since there are $k$ terms in the last sum above, $(\text{RSS}^{(0)} - \text{RSS}^{(1)})/\sigma^2$ has a $\chi^2(k)$ distribution and we are done demonstrating the second and third properties.

We often test for two parameters $\beta_1$ and $\beta_2$ to be equal, so $H_0 : \beta_1 = \beta_2 = 0$. The MLE is $\hat{\beta}_1 - \hat{\beta}_2$ with variance

$$\text{var}(\hat{\beta}_1 - \hat{\beta}_2) = \text{var}(\hat{\beta}_1) + \text{var}(\hat{\beta}_2) - 2\text{cov}(\hat{\beta}_1, \hat{\beta}_2) = \sigma^2(X^T X)^{-1}_{1,1} + \sigma^2(X^T X)^{-1}_{2,2} - 2\sigma^2(X^T X)^{-1}_{1,2}.$$
so the test statistic is

\[ s \sqrt{(X^TX)^{-1}_{1,1} + (X^TX)^{-1}_{2,2} - 2(X^TX)^{-1}_{1,2}} \sim t(n - p) \]

This works for linear combinations of parameters. If \( v \) is \( p \times 1 \) and we want to test for \( v^T \beta = 0 \) then the MLE is \( v^T \hat{\beta} \), and

\[ \text{var}(v^T \hat{\beta}) = v^T \text{var}(\beta)v \]

so the test statistic is

\[ \frac{v^T \hat{\beta}}{s \sqrt{(v^T (X^TX)^{-1}v)}} \sim t(n - p). \]

The quantities \( s^2 \) and \( v \hat{\beta} \) are independent, since \( s^2 \) and \( \hat{\beta} \) are independent.

**Exercise** verify that \( v = (1, -1, 0, \ldots, 0)^T \) gives the test for \( \hat{\beta}_1 - \hat{\beta}_2 = 0 \).

There are some shortcuts. For example, in a test for \( \beta_1 = \beta_2 \) the reduced model, with \( \beta'_1 = \beta_1 = \beta_2 \) is

\[ y = \beta'_1(x_1 + x_2) + \beta_3 x_3 + \ldots \]

and the full model, \( Y = X \beta + \epsilon \), can be written

\[ y = \beta'_1(x_1 + x_2) + \beta'_2(x_1 - x_2) + \beta_3 x_3 + \ldots \]

so the test for \( \beta_1 = \beta_2 \) can be framed as a test \( \beta'_2 = 0 \). We can run a T or F test to drop \( \beta'_2 \), with design matrix \( X = [X_1 + X_2, X_1 - X_2, X_3, \ldots, X_p] \) and parameter vector \( \beta = (\beta'_1, \beta'_2, \beta_3, \ldots, \beta_p) \).

2.5. **ANOVA, and an example.** Because ANOVA tests are used so frequently, the important numbers in the test are laid out in a standard way, to facilitate reading. There is a little variation (the default R table doesn’t exactly follow my rules), but just a little.

The ANOVA table sets out the numbers we need to make tests for dropping certain collections of variables. Suppose the variables \( x_1, \ldots, x_p \) come in \( m = 3 \) groups (typically just 2 or 3 groups) and are ordered so that the groups are

\[ \{1\}, \{2, 3, \ldots, p-k\}, \{p-k+1, p-k+2, \ldots, p\}. \]

This splits off the variables \( x_{p-k+1} \) to \( x_p \). This is the variable grouping relevant for the hypothesis \( H_0 : \beta_{p-k+1} = \beta_{p-k+2} = \ldots = \beta_p = 0 \), which we test with an F-test.

A typical ANOVA table gives fitting information for each of the models starting from a simplest model with just intercept \( \beta_1 \), adding the groups one at a time, up to the model with all \( p \) variables. For the \( m = 3 \)-group case, testing \( H_0 : \beta_{p-k+1} = \beta_{p-k+2} = \ldots = \beta_p = 0 \), the model sequence is

\[ y = \beta_1 + \epsilon, \]

\[ y = \beta_1 + \beta_2 x_2 + \ldots + \beta_{p-k} x_{p-k} + \epsilon, \]

\[ y = \beta_1 + \beta_2 x_2 + \ldots + \beta_{p-k} x_{p-k} + \ldots + \beta_p x_p + \epsilon, \]

If we order the variables in the right way, we can sometimes do model selection at a glance, as we read down the table from top to bottom. If \( X_{1,i} = [x_1, x_2, \ldots, x_i] \),
then the design matrices build from $X_1$ to $X = X_{1:p}$. Let $RSS_{1,i}$ be the residual sum of squares for the fit with design matrix $X_{1:i}$. The decrease in the residual sum of squares when we add the variables $x_{i+1}, \ldots, x_{i+k}$ to a model that already has the variables $x_1, x_2, \ldots, x_i$ is $RSS_{1,i} - RSS_{1,i+k}$. The number of residual degrees of freedom in the fit for the model with design matrix $X_{1:i}$ is $n - i$ (assuming the columns of $X_{1:i}$ are linearly independent).

The layout of an ANOVA table for the three groups \{1\}, \{2, ..., $p - k$\}, \{$p - k + 1, ..., p$\} is shown in Table 1. $TSS = (y - \bar{y})^2 (y - \bar{y})$ is the residual sum of squares for a model with just intercept, in other words, the total sum of squares adjusted for intercept. $RSS_{1:p}$ is the residual sum of squares for the full model.

The $F$-statistic in row two of Table 1 is the $F$-test statistic for the test to add the variables \{$x_{p-k+1}, \ldots, x_p$\} to a model with variables \{$x_1, \ldots, x_{p-k}$\}, which is the test we set up in Section 2.4.

The $F$-statistic in row one of Table 1 is an $F$-test statistic for the test to add the variables \{$x_2, \ldots, x_{p-k}$\} to a model with just $x_1$, the intercept variable. It might seem natural to use the divisor $RSS_{1:p}/(n - (p - k))$, for an $F$ with $p - k - 1$ numerator and $n - (p - k)$ denominator degrees of freedom. However, (i) the divisor $RSS_{1:p}/(n - p)$ is “just as good” as $RSS_{1:p}/(n - (p - k))$, since it too is independent of $TSS - RSS_{1:p-k}$, so we can see $(TSS - RSS_{1:p-k})(n - p)/RSS_{1:p}(p - k - 1)$ has an $F(p - k - 1, n - p)$ distribution under the null, and (ii) it is better, as $RSS_{1:p}/(n - p)$ is an estimate of $\sigma^2$ which is not biased if the variables $x_{p-k+1}, \ldots, x_p$ added in the row below turn out to be explanatory. You might possibly add (iii) the divisor $RSS_{1:p}/(n - p)$ has a higher variance than $RSS_{1:p-k}/(n - (p - k))$, if variables in the rows below really were not related to the response, so in that case we would do better to drop them from the ANOVA. That is equivalent to using the $RSS_{1:p-k}/(n - (p - k))$ divisor. Another way to make point (iii) is that the $RSS_{1:p-k}/(n - (p - k))$ divisor is the one given by the LRT, where as $RSS_{1:p}/(n - p)$ is just some statistic with a distribution we happen to know under the null. On balance item (ii) controls our choice of test statistic, so the table opts for higher variance in return for lower bias.

<table>
<thead>
<tr>
<th>Terms added</th>
<th>Degrees Freedom</th>
<th>Reduction in RSS</th>
<th>Mean Square</th>
<th>$F$ statistic</th>
</tr>
</thead>
<tbody>
<tr>
<td>$X_{2:(p-k)}$</td>
<td>$p - k - 1$</td>
<td>$TSS - RSS_{1:(p-k)}$</td>
<td>$\frac{TSS - RSS_{1:(p-k)}}{p - k - 1}$</td>
<td>$\frac{(TSS - RSS_{1:(p-k)})/(p - k - 1)}{RSS_{1:p}/(n - p)}$</td>
</tr>
<tr>
<td>$X_{(p-k+1):p}$</td>
<td>$k$</td>
<td>$RSS_{1:(p-k)} - RSS_{1:p}$</td>
<td>$\frac{RSS_{1:(p-k)} - RSS_{1:p}}{k}$</td>
<td>$\frac{(RSS_{1:(p-k)} - RSS_{1:p})/k}{RSS_{1:p}/(n - p)}$</td>
</tr>
<tr>
<td>Residual</td>
<td>$n - p$</td>
<td>$RSS_{1:p}$</td>
<td>$\frac{RSS_{1:p}}{n - p}$</td>
<td></td>
</tr>
</tbody>
</table>

Table 1. ANOVA table for the groups \{$x_2, \ldots, x_{p-k}$\} and \{$x_{p-k+1}, \ldots, x_p$\} added incrementally to the intercept group \{$x_1$\}. In some tables a final column giving the $p$-value is included.
Table 1 is the table we might set out if we were carrying out the \( F \)-test for \( H_0: \beta_{p-k+1} = \beta_{p-k+2} = \ldots = \beta_p = 0 \) against \( H_1: \beta \in \mathbb{R}^p \), though we could omit the first row.

Other relevant test statistics can be computed from the numbers in an ANOVA table like Table 1. For example, the test for the model hypothesis \( H_0': \beta_2 = \ldots = \beta_p = 0 \) against the full model \( H_1: \beta \in \mathbb{R}^p \), is the test for no linear relation to the variables \( x_2, \ldots, x_p \). The test statistic is

\[
F' = \frac{TSS - RSS_{1:p}}{p-1} \times \frac{n-p}{RSS_{1:p}},
\]

since \( k = p-1 \) here. The second factor is given at the bottom of the table. The first element \( TSS - RSS_{1:p} \) is the sum of the terms in the third column of the table,

\[
TSS - RSS_{1:p} = (RSS_{1:(p-k)} - RSS_{1:p}) + (TSS - RSS_{1:(p-k)}),
\]

so we can form \( F' \) by taking appropriate sums and ratios of table elements. Note that the statistic \( F' \) replaces the widely used statistic \( R^2 \) as a measure of fit quality (or the lack of it). While \( R^2 \) runs from zero to one, we have no absolute scale for quality of fit (how close to one is acceptable), \( F' \) runs from 0 to infinity (and big \( F' \) is poor fit) and does give a direct test for significant linear dependence. Note that R outputs both \( F' \), the test statistic for no linear relation, and the \( p \)-value for this test, automatically, when we make a \texttt{summary()} of a \texttt{lm()} output. This is more useful to us than \( R^2 \), though R gives this as well.

Again, these numbers are useful for calculating other tests. The test for no linear dependence on the variables \( x_2, \ldots, x_p \), has \( F \) statistic

\[
F = \frac{(TSS - RSS_{1:p})/(p-1)}{RSS_{1:p}/(n-p)}
\]

with \( (p-1) \) numerator and \( (n-p) \) denominator degrees of freedom. The quantity \( TSS - RSS_{1:p} \) is the sum of all the entries in the third column, bar the last,

\[
TSS - RSS_{1:p} = (RSS_{1:i_{m-1}} - RSS_{1:p}) + (RSS_{1:i_{m-2}} - RSS_{1:i_{m-1}}) + \ldots + (RSS_{1:i_2} - RSS_{1:i_3}) + (TSS - RSS_{1:i_2}).
\]

**Example 2.3.** Consider variable selection for the trees data. We looked at this in Example 2.2. When we have many variables, we can’t test all possible combinations. Physical considerations (aka common sense) are always important, but especially so, when we are setting up the hypotheses. We considered the linear model

\[
y = \beta_1 + \beta_2 x_2 + \beta_3 x_2 + \epsilon
\]

with \( y = \log(v/h^2) \), \( \beta_1 = \log(\eta) \), \( x_2 = \log(h) \), \( x_3 = \log(g) \) and \( \epsilon = \log(\gamma) \) (and \( v \), \( g \) and \( h \) the volume, girth and height).

Look at the R-output for this model in Example 2.2. The quoted residual standard error is \( s^2 = RSS/(n-p) \): the RSS is \( (Y - \hat{Y})^2 \), or in R,

\[
> (rss<-sum(trees.lm1$residuals^2))
> (sqrt(rss/28))
\]

so estimated error variance aka the Residual standard error \( s^2 \) is

\[
> (sqrt(rss/28))
\]
Figure 3. (LEFT) A qqplot of the residuals of the logged-model fit to the trees data looks healthy, and (RIGHT) no sign of correlation between residuals and fitted values (though a possible funnel, increasing variance with fitted value).

You might like to check the elements in the summary(trees.lm1) table.

Later on we look at checks on the fit. Under the model we have just estimated (which is $H_1$, with the full parameter set), the residuals are given by $e = (I_n - H)e$, so that each residual $e_i$ is normally distributed and the residuals $e$ and fitted values $\hat{Y} = HY$ are independent. We can make a qqplot of the sample quantiles against the normal quantiles and look for normality. Also, we can plot fitted values against residuals and look for a trend. The qqplot is obviously important here, as our $e = \log(\gamma)$ is the log of a multiplicative error, so this is an area we might expect to see departures from the model. The qqplot in Figure 3 is very well behaved, except possibly in the upper tail.

We now come to the test for $\beta_2 = \beta_3 = 0$. R gives us the ANOVA tables. Here are several ways to do this. We begin by fitting the reduced, $H_0$, model.

```r
> trees.lm0<-lm(log(Volume/(Height*Girth^2))~1)
```

The fields of `trees.lm0` are special cases. There is just an intercept, so $\tilde{X} = 1_{n,1}$, $X^T X = n$, $(X^T X)^{-1} = 1/n$, and $(X^T X)^{-1} X^T y = \bar{y}$.

Now we use `ANOVA` to see if $\beta_2 = \beta_3 = 0$. We can do this ‘by hand’.

```r
> (rss0<-sum(trees.lm0$residuals^2))
[1] 0.1876858
> (rss1<-sum(trees.lm1$residuals^2))
[1] 0.1854634
> k<-2
> n.minus.p<-31-3
```
The $p$-value, $p = 0.85$, is not significant, so the LRT supports $H_0 : \beta_2 = \beta_3 = 0$ (that is, there is no evidence for dependence of $y$ on $x_2$ and $x_3$).

We can get R to form something like a regular ANOVA table, Table ???. The default R ANOVA adds one variable at a time, so the groups of variables are \{\{x_1\}\}, \{\{x_2\}\} \{x_3\} (since $F$-tests for many other grouping can be computed form this 'finest resolution' table). Because the change in the number of degrees of freedom is one at each row, columns three and four of Table ?? are identical.

\[ \text{anova(trees.lm1)} \]

Analysis of Variance Table

<table>
<thead>
<tr>
<th>Response: log(Volume/(Height * Girth^2))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Df</td>
</tr>
<tr>
<td>-----</td>
</tr>
<tr>
<td>log(Height)</td>
</tr>
<tr>
<td>log(Girth)</td>
</tr>
<tr>
<td>Residuals</td>
</tr>
</tbody>
</table>

In this table Sum Sq corresponds to “Reduction in RSS” in Table ???. Thus the residual sum of squares $\text{RSS}_{1:p}$ for the full model is $\text{RSS}_{1:3} = 0.185463$, and $\text{RSS}_{1:p}/(n - p) = 0.185463/28 = 0.006624$. The other quantity we need, for the test $\beta_3 = \beta_2 = 0$ is $\text{TSS} - \text{RSS}_{1:3}$, which is

\[ \text{RSS}_{1:2} - \text{RSS}_{1:3} + \text{TSS} - \text{RSS}_{1:2} = 0.000354 + 0.001868 = 0.002222. \]

So

\[ F = \frac{(\text{TSS} - \text{RSS}_{1:3})/2}{\text{RSS}_{1:p}/(n - p)} = \frac{0.001111/0.006624}{0.1677} \approx 0.1677 \]

and the $F$-test proceeds as before.

The ANOVA table we got from R didn’t quite have the variable grouping we wanted - we got the default grouping, which just put every explanatory variable in a separate group, so we had to do some arithmetic to get our test statistic. Alternatively, we can tell R which specific models we want to compare, and get R to form a table

\[ \text{anova(trees.lm0,trees.lm1)} \]

Analysis of Variance Table

<table>
<thead>
<tr>
<th>Model 1: log(Volume/(Height * Girth^2)) ~ 1</th>
</tr>
</thead>
<tbody>
<tr>
<td>Res.Df</td>
</tr>
<tr>
<td>--------</td>
</tr>
<tr>
<td>1</td>
</tr>
<tr>
<td>2</td>
</tr>
</tbody>
</table>

which we see is not a standard ANOVA table, but is nevertheless easy enough to read. We read this $\text{RSS}_{1:3} = 0.185463$, $\text{TSS} = 0.187686$, $\text{TSS} - \text{RSS}_{1:3} = 0.002222$, with $k = 2$ for the two variables we set to zero, and if $F = 28(\text{TSS} - \text{RSS}_{1:3})/2\text{RSS}_{1:3}$ then $F = 0.1678$. Finally, if $F_{2,28} \sim F(2, 28)$ is an $F$-distributed
rv with 2 numerator and 28 denominator degrees of freedom, then we read off \( Pr(F_{2,28} > F) \approx 0.85 \), so the p-value shows no evidence for a departure from \( H_0 \). This time, there was no arithmetic needed.

We could get this directly, also, from the `summary(trees.lm1)` output above. Recall that this output automatically gives the \( F \)-statistic for the reduced model with no explanatory variables except the intercept, \( \beta_1 \) here. That is just the model reduction we are considering, so the final line

\[
F \text{-statistic: 0.1678 on 2 and 28 DF, p-value: 0.8464}
\]

gives us the same elements, \( F = 0.1678 \) and \( Pr(F > f) \approx 0.85 \). This part of the output would not be relevant if we were considering dropping a smaller subset of the explanatory variables.

2.6. **Categorical variables.** So far we have treated continuous explanatory variables. However, explanatory variables may be categorical. The values taken by a categorical variable are called its **levels**, and the levels may be ordered or unordered. We will discuss unordered categorical variables.

A categorical explanatory variable \( x_k^{\text{cat}} \) takes \( c \) levels, equivalent to \( c \) binary indicator variables \( g_{k,a} = 1 \) for \( a = 1, 2, \ldots, c \) the level index, so the \( k \)'th response \( y_k \) has one explanatory variable \( g_{k,a} \) for each level of the original categorical variable. Suppose we want to allow the response to have a mean which depends on the level of \( x_k^{\text{cat}} \), and suppose that, for the \( k \)'th response \( y_k \), there are \( m \) other explanatory variables \( x_{k,1}, x_{k,2}, \ldots, x_{k,m} \) including an intercept, \( x_{k,1} = 1 \).

The model

\[
y_k = \alpha + \alpha_2 g_{k,2} + \ldots + \alpha_c g_{k,c} + \gamma_2 x_{k,2} + \ldots + \gamma_m x_{k,m} + \epsilon_k
\]

allows the mean of \( y_k \) to vary with the level of \( x_k^{\text{cat}} \). What happened to \( \alpha_1 \)g_{k,1}\? If we include it, then our model is over-parameterized. If the level for response \( k \) is \( x_k^{\text{cat}} = a \), then \( g_{k,1} = 1 \) and \( g_{k,2} = \ldots = g_{k,c} = 0 \), so

\[
E(Y_k) = \alpha + \gamma_2 x_{k,2} + \ldots + \gamma_m x_{k,m}.
\]

If the level is \( x_k^{\text{cat}} = a \), then \( g_{k,a} = 1 \) and the others are zero, so

\[
E(Y_k) = \alpha + \alpha_a + \gamma_2 x_{k,2} + \ldots + \gamma_m x_{k,m}.
\]

We see that \( \alpha_a \) is the offset in the intercept of the level-\( a \) samples relative to the intercept \( \alpha \) of the level-1 samples. We are using level 1 as the **baseline** level.

If \( G_a \) is the binary column vector \( G_a = (g_{1,a}, \ldots, g_{n,a})^T \) for the level-\( a \) indicator, and \( X_1, X_2, \ldots, X_m \) are column vectors for other variables, then the design matrix for the model above is \( X = (X_1, G_2, \ldots, G_c, X_2, \ldots, X_m) \) (so \( p = m + c - 1 \) here, and columns are in no particular order). The model itself is \( Y = X \beta + \epsilon \) with \( \beta = (\alpha, \alpha_2, \ldots, \alpha_c, \gamma_2, \ldots, \gamma_m) \). We left out \( G_1 \) when we formed the new design matrix because \( X \) has a first column of ones, corresponding to the intercept. But then \( X_1 = \sum_{a=1}^c G_j \) since each observation must have its categorical variable in one of the levels 1, 2, \ldots, \( c \), and so the columns of \( (X_1, G_1, G_2, \ldots, G_c, X_2, \ldots, X_m) \) are not linearly independent, and again, the model with all \( c \) columns, \( G_1, \ldots, G_c \), is over-parameterized. The variables \( G_1, \ldots, G_c \) are sometimes called a **dummy variables** for the level and the matrix \( (G_2, \ldots, G_c) \) is called a **contrast** matrix.
Example 2.4. The data depicted in Figure 4 shows average Oxford house prices (in thousands of pounds) for 100 months starting April 2000 ending July 2008 for Detached, Semi-Detached and Terraced houses and Flats. The website [www.home.co.uk](http://www.home.co.uk) displays data of this kind. The Flats and Detached properties clearly have lower and higher variance respectively, than the two other classes. We will deal with them later. We start with an analysis of the two-hundred Semi-Detached and Terraced prices. Here are some rows of data:

```r
> ohp[1:3,]
price  type  month  sales
329  Semi-Detached  100  19
276  Semi-Detached  99   37
300  Semi-Detached  98   45
```

The prices `price` are average figures for the month, in thousands of pounds, while the number of sales `sales` gives the number of individual sale prices which were averaged to form the price reported for that month. We will leave a discussion of column four for the moment. The variable `type` is now a two-level categorical variable.

Is there any difference between the price trends for the two types of houses? The lattice plot, Figure 4, supports a linear model for `price` as a function of `month`. We have omitted the two-level categorical variable `type` in Figure 4. We will begin by fitting a model with a different offset for the two levels - we assume prices grow at the same rate, but there is an offset in the price for Terraced relative to Semi-Detached properties. Let \( y_k = \text{price}[k], x_{k,1} = 1, x_{k,M} = \text{month}[k] \) and \( g_{k,T} = I_{\text{type}[k] = \text{Terraced}} \), so that

\[
y_k \sim \alpha + \alpha_T g_{k,T} + \gamma_M x_{k,M} + \epsilon_k, \quad \epsilon_k \sim N(0, \sigma^2).
\]

In this model \( \alpha \) is the price of a Semi-Detached house in month 0, and \( \alpha + \alpha_T \) is the price of a Terraced house in month 0. In R, the intercept is included by default, so `price~month+type` and `price~1+month+type` specify the same model. Also, R will automatically construct dummy variables for the levels of a categorical variable (such as `type`, which has levels Semi-Detached and Terraced).
How does R code the categorical variable type?

```r
> X <- model.matrix(price ~ month + type, data = ohp)
> X[1:3,]
       (Intercept) month typeTerraced
 1         1  100        0
 2         1   99        0
 3         1   98        0
> X[99:102,]
       (Intercept) month typeTerraced
 1         1    2        0
 2         1    1        0
 3         1  100        1
 4         1    99        1
> X[198:200,]
       (Intercept) month typeTerraced
 1         1    3        1
 2         1    2        1
 3         1    1        1
```

Figure 4. Lattice scatter plot of monthly average prices, against month and number of sales. (solid/black circle) Semi-Detached, (open/red circle) Terraced, (solid/green square) Flat, (open/blue square) Detached.
The rightmost column of the design matrix in this R implementation is $G_T = (g_1,T, ..., g_n,T)^T$. The baseline level is **Semi-Detached** and the variable mapping for the design matrix above is $(\alpha, \gamma, \alpha_T) = (\beta_1, \beta_2, \beta_3)$. You need to check you know which variable R is using as the baseline, though you wouldn’t use `model.matrix()` to do that: the baseline level is simply the level omitted in the `summary()` output (see below). The offset in the mean for a house with **type** equal **Terraced** is $\beta_3$, the parameter for column three in $X$, the R the design matrix above.

OK, so let’s fit the model `price~month+type`.

```r
> ohp.lm<-lm(price~month+type,data=ohp)
> summary(ohp.lm)
...
Coefficients:
            Estimate Std. Error t value Pr(>|t|)
(Intercept) 158.09727  3.59152 44.020  < 2e-16 ***
  month       1.69887  0.05531 30.716  < 2e-16 ***
  typeTerraced -2.58000  3.19306  -0.808   0.42

Residual standard error: 22.58 on 197 degrees of freedom
Multiple R-Squared: 0.8274, Adjusted R-squared: 0.8256
F-statistic: 472.1 on 2 and 197 DF, p-value: < 2.2e-16
```

We don’t need an $F$-test to test $\alpha_T = 0$ (**ie**, $\beta_3 = 0$) here: for a single parameter, the $t$-test is equivalent. We have $n = 200$ data and $p = 3$. Reading off the table, $\hat{\alpha}_T = -2.58$, and the $t$-statistic for the test $\alpha_T = 0$,

$$\hat{\alpha}_T / s\sqrt{(X^TX)^{-1}_{33}} = -2.58 / 3.19$$

is equal to $-0.81$, with $n-p=197$ degrees of freedom. The $p$-value $2(1-Pr(T < |t|))$ is $2 \times (1 - pt(0.81, 197)) \approx 0.42$ (which we can read in the right column) and this shows that the Terraced/Semi-Detached distinction is not significant. The two regressions are plotted in Figure 5. There is clearly little difference. Note that this is not the same as making two regressions and using a $t$-test for equality of intercepts, as we are imposing (i) equal slopes, and (ii) equal error variance $\sigma^2$.

**Exercise** Can you see any sign of model-mispecification in Figure 5?

### 2.7. Variable interactions.

We can form new explanatory variables from old by taking functions of explanatory variables. Interactions of the form $\beta_i x_i + \beta_j x_j + \beta_I x_i x_j$ are particularly common, and mean something like “variable $j$ has more impact on the response when variable $i$ is large” and *vice versa*. If $x_i$ is a binary dummy variable for some level $a$ of a categorical variable $x_{\text{cat}}$, then the slope with increasing $x_j$ is $\beta_j$ for observations with $x_{\text{cat}} \neq a$ (where $x_i = 0$) and the slope is $\beta_j + \beta_I$ for observations with $x_{\text{cat}} = a$ (where $x_i = 1$).

When we have interactions we often include lower order terms in the model, though we might have no ‘physical’ use for them. Suppose $y = \alpha + \beta_1 x_1 x_2$ with $x_1$ in Celsius. If we switch to Farenheit, $x_1 = mx_1' + d$ with $m = 5/9$ and $d = 160/9$, then $y = \alpha + d\beta_2 x_2 + m\beta_1 x_1 x_2$. Now we have a new kind of term $d\beta_2 x_2$. We may dislike the idea that the kinds of terms in our model (rather than just the parameter values) are dependent on the zero-location for interacting variables, and instead at least begin our modeling with $y = \alpha + \beta_1 x_1 + \beta_2 x_2 + \beta_I x_1 x_2$. Faraway (2004) Chapter 8 page 122 has more on this.
Example 2.5. We looked in Example 2.4 at prices for Terraced and Semi-Detached houses in Oxford. Let us see if Terraced and Semi-Detached houses have grown at different rates. In order to make the whole thing a little more interesting, I will add Flats to the picture. I am ignoring the somewhat lower variance of the Flats data, and more on this anon. Have Flats increased in price at the same rate as Terraced and Semi-Detached properties? Is there any difference in the rates or intercepts for the latter house types?

With Flat providing the baseline, we fit the model

\[ y_k \sim \alpha + \alpha_T g_{k,T} + \alpha_{SD} g_{k,SD} + \gamma_M x_{k,M} + \gamma_{MT} x_{k,M} g_{k,T} + \gamma_{MSD} x_{k,M} g_{k,SD} + \epsilon_k \]

with \( \epsilon_k \sim N(0, \sigma^2) \) and, for observation \( k = 1, 2, ..., n \), we have \( y_k = \text{price}[k] \), \( g_{k,T} \) is the dummy indicator variable for \( \text{type}[k] = \text{Terraced} \), \( g_{k,SD} \) is the dummy indicator variable for \( \text{type}[k] = \text{Semi-Detached} \) and \( x_{k,M} \) is the value of \( \text{month}[k] \). In month 0 (so, at the intercept), the Flat price is \( \alpha \), the Semi-Detached price is \( \alpha + \alpha_{SD} \), and the Terraced price is \( \alpha + \alpha_T \) thousands of pounds. Flat prices go up at rate \( \gamma_M \), Semi-Detached up at rate \( \gamma_M + \gamma_{MSD} \), and Terraced up at rate \( \gamma_M + \gamma_{MT} \) thousands of pounds per month. In R the model \( \text{price} \sim \text{month} \times \text{type} \) is expanded as the model \( \text{price} \sim 1 + \text{month} + \text{type} + \text{month}:\text{type} \), so that the : notation gives the product term by itself, with no lower order terms, while the * notation includes lower order terms by default.

```r
> ohp.lm<-lm(price~month*type,data=ohp)
> (ohp.lms<-summary(ohp.lm))

Call:
  lm(formula = price ~ month * type, data = ohp)
```
Residuals:

<table>
<thead>
<tr>
<th></th>
<th>Min</th>
<th>1Q</th>
<th>Median</th>
<th>3Q</th>
<th>Max</th>
</tr>
</thead>
</table>

Coefficients:

| Term                      | Estimate | Std. Error | t value | Pr(>|t|) |
|---------------------------|----------|------------|---------|----------|
| (Intercept)               | 126.50182 | 4.14322 | 30.532 | < 2e-16 *** |
| month                     | 1.21838 | 0.07123 | 17.105 | < 2e-16 *** |
| typeSemi-Detached         | 29.61091 | 5.85940 | 5.054 | 7.63e-07 *** |
| typeTerraced              | 31.00000 | 5.85940 | 5.291 | 2.39e-07 *** |
| month:typeSemi-Detached   | 0.51978 | 0.10073 | 5.160 | 4.55e-07 *** |
| month:typeTerraced        | 0.44119 | 0.10073 | 4.380 | 1.65e-05 *** |

Signif. codes: * * * 0.001 ** 0.01 * 0.05 . 0.1 ' 1

Residual standard error: 20.56 on 294 degrees of freedom
Multiple R-Squared: 0.8661, Adjusted R-squared: 0.8638
F-statistic: 380.3 on 5 and 294 DF, p-value: < 2.2e-16

The variable mapping is

$$(\alpha, \alpha_{SD}, \alpha_T, \gamma_M, \gamma_{M SD}, \gamma_{MT}) = (\beta_1, \beta_2, \beta_3, \beta_4, \beta_5, \beta_6).$$

We see that Flat price grows at a lower rate than Semi or Terraced (since the offsets $\alpha_{SD} \simeq 0.52, \alpha_T \simeq 0.44$ are positive, and significant). We can test for differing rates between Terraced and Semi-Detached. The test statistic is

$$T = \frac{\beta_5 - \beta_6}{\sqrt{s^2(X^TX)_{5,5}^{-1} + s^2(X^TX)_{6,6}^{-1} - 2s^2(X^TX)_{5,6}^{-1}}} \simeq \frac{0.52 - 0.44}{\sqrt{0.10073^2 + 0.10073^2 - 2 \times 0.005074}} \simeq 0.78$$

Now if $T_{n-p} \sim t(n-p)$ with $n = 300$ and $p = 6$ here, then the p-value $2(1 - \Pr(T_{n-p} > T))$ is $2 \times (1 - pt(T, 294)) \simeq 0.436$. Note that the matrix $(X^TX)^{-1}$ is part of the output of `summary()`, so the code I used to make this test was

```r
> (beta<-ohp.lm$coefficients)
(Intercept) month typeSemi-Detached
126.5018182 1.2183798 29.6109091

typeTerraced month:typeSemi-Detached month:typeTerraced
31.0000000 0.5197840 0.4411881

> (s<-ohp.lms$sigma)
[1] 20.56092
> XTXi<-ohp.lms$cov.unscaled
> (T<-abs( (beta[5]-beta[6])/(s*sqrt(XTXi[5,5]+XTXi[6,6]-2*XTXi[5,6])) ))
0.780243
> 2*(1-pt(T,ohp.lms$df.residual))
0.4358756
```

We conclude that there is no evidence for different rates, when the intercepts are unequal. We can see the fitted lines in Figure 6. There is clearly little in it.
We now want to make the combined test, dropping the intercept and slope distinction between Terraced and Semi-Detached. Since we now make no distinction between Terraced and Semi-Detached, we are effectively merging the levels Terraced and Semi-Detached within the categorical variable type. Let \( y_k,TSD = g_k,T + g_k,SD \). With level Flat again providing the baseline, the reduced model, with \( \alpha_T = \alpha_{SD}, \gamma_T = \gamma_{SD} \) is
\[
y_k \sim \alpha + \alpha_{TSD}g_k,TSD + \gamma_Mx_k,M + \gamma_{MTSD}x_k,Mg_k,TSD + \epsilon_k
\]
One easy way to implement this in R is to re-level the categorical variable, but otherwise proceed as before.

```r
> # last two levels Semi-Detached and Terraced are over-written with 'T.or.SD'
> ohpr<-ohp
> levels(ohpr$type)<-c('Flat','T.or.SD','T.or.SD')
> # fit the reduced model
> ohpr.lm<-lm(price~month*type,data=ohpr)
> # calculate residual sums of squares for full (rss1) and reduced (rss0) models
> rss0<-sum(ohpr.lm$residuals^2)
> rss1<-sum(ohpr.lm$residuals^2)
```
> # form the F statistic and calculate a p-value
> F<-(rss0-rss1)/2/(rss1/294)
> (pval<-1-pf(F,2,294))
> [1] 0.4983896

At around 0.5, the p-value for the LRT is not significant, so the test favors the reduced model. It seems that both prices and trends in prices for Terraced and Semi-Detached types are the same, as you might guess from Figure 6.

You can check, from the `summary(ohpr.lm)` output (not shown) that \( \hat{\gamma}_{MTSD} \) is non-zero and positive (the p-value for \( \hat{\gamma}_{MTSD} > 0 \) is tiny), so Flat prices have increased at a rate which is significantly lower than the rate of increase for Terraced and Semi-Detached properties. Note that this is still not the same as making two regressions, as we are imposing equal error variance \( \sigma^2 \) for the response under the two types. Note also that our conclusions are based on gross data for Oxford, and the data have been slightly jittered, so local trends could differ. [End L5]

2.8. Blocks, Treatments and Designs. Chapters 14-16 of 'Linear Models with R' by J. Faraway covers this material at about the right level for us. See the discussion in Davison (2003) for more detail.

One important kind of categorical variable arises when the data have been gathered from blocks, \( y_{i,j}, i = 1, 2, ..., b, j = 1, 2, ..., n_i \), corresponding to groups of subjects that are expected to have similar response to the explanatory variables. Imagine collecting observations of the body mass index (BMI) of eighty five-year-old children from different schools. In one design we measure the BMI of eight randomly selected five-year-olds in each of 10 schools. For each child we record the BMI and the school. In another design, we might do exactly the same, but fail to record the school. The first data set has a block structure, with 10 blocks. The block index is often explanatory. If, for example, there is a correlation between parent income-level, school and incidence of obesity, then 'school' will be explanatory for 'BMI'. In such cases we code the block index \( i \) as a categorical explanatory variable in the design matrix.

Besides taking subjects from distinct groups, and distinguishing group responses, we may also give subjects different treatments, and distinguish responses to different treatments. If there is a block structure to the population, with subjects in different blocks having different treatment responses, and we ignore it, then this will tend to inflate the estimated error variance \( \sigma^2 \), and real differences in the treatment response may not be detected.

"Treatment factors are those for which we wish to determine if there is an effect. Blocking factors are those for which we believe there is an effect. We wish to prevent a presumed blocking effect from interfering with our measurement of the treatment effect", a neat encapsulation I found in Heiberger and Holland ‘Statistical Analysis and Data Display’, Springer (2004).

Example 2.6. The following example is taken from Dr D. Lunn’s previous lecture notes for this course. Twelve piglets were fed three different diets (A, B and C) in order to see which diet led to the greatest weight gain. The piglets came from four different litters (I, II, III and IV). We are looking for an effect (a mean shift in weight gain) due to diet, and we want to allow for variation (another mean shift) due to litter. If \( y_{i,j} \) is the response in row \( i \) and column \( j \) of Table 2 then the model
we want to fit is
\[
\begin{align*}
y_{1,j} &= \alpha + \epsilon_{1,j} \\
y_{i,j} &= \alpha + \tau_j + \epsilon_{i,j} \\
y_{i,1} &= \alpha + \gamma_i + \epsilon_{i,1} \\
y_{i,j} &= \alpha + \gamma_i + \tau_j + \epsilon_{i,j}
\end{align*}
\]
for \(i = 2, 3, 4\) and \(j = 2, 3\). Notice that \(i = 1\) (Litter I) and \(j = 1\) (Diet A) are the baseline levels for Litter and Diet. The blocks (Litter) and treatments (Diet) are categorical, and can be coded using dummy variables. Let \(k = 1, 2, ..., n = 12\) run over subjects, as appears in the margin of the following output.

```r
```
In matrix notation the model we are fitting is $Y = X\beta + \epsilon$ with $eta = (\alpha, \gamma_2, \gamma_3, \gamma_4, \tau_2, \tau_3)$.

Now, let's fit the model.

```r
> pigs.lm<-lm(Gain~Litter+Diet,data=pigs)
> summary(pigs.lm)

Call:
  lm(formula = Gain ~ Litter + Diet, data = pigs)

Residuals:
  Min     1Q Median     3Q    Max
-8.250 -4.937 -0.375  2.938  12.750

Coefficients:  Estimate Std. Error t value Pr(>|t|)
(Intercept)  86.250     5.763   14.97   5.6e-06 ***
LitterII    -7.000     6.654    -1.05   0.3333    
LitterIII   21.000     6.654    3.16    0.0197    *
LitterIV     1.000     6.654    0.15    0.8855    
DietB       -21.750    5.763   -3.77    0.0092    **
DietC       -18.000    5.763   -3.12    0.0205    *
---
Signif. codes:  0 '***'  0.001 '**'  0.01 '*'  0.05 '.'  0.1 ' ' 1

Residual standard error: 8.15 on 6 degrees of freedom
Multiple R-Squared: 0.8569,   Adjusted R-squared: 0.7376
F-statistic: 7.184 on 5 and 6 DF,  p-value: 0.01623
```

It is clear from the parameter estimates that Litter I and Diet A have been taken as the baseline levels for the respective categorical variables.

Is Diet predictive for Gain? The ANOVA table for variable groups

```r
> anova(pigs.lm)

Analysis of Variance Table

Response: Gain
  Df Sum Sq Mean Sq  F value  Pr(>F)
Litter  3 1304.25  434.75  6.5458   0.0255 **
Diet   2 1081.50  540.75  8.1418   0.0195 *
Residuals  6  398.50  66.42
---
Signif. codes:  0 '***'  0.001 '**'  0.01 '*'  0.05 '.'  0.1 ' ' 1
```

so Diet is predictive at 5% significance.

**Example 2.7.** What happens if the data did not record the block variable Litter? The variation within each diet is inflated by variation due to Litter. If the Litter variable is not there, this variation increases the residuals, so the estimated variance,
\[ s^2 = \frac{\text{RSS}}{(n - p)} \]

This in turn pulls down the \( t \) and \( F \) test statistics, which both have RSS in the denominator, so \( p \) values tend to increase. The outcome of ignoring variation due to block variables is that the response due to treatment variables which are marginally significant can be explained away as noise, as their parameters are no longer significantly different from zero.

```r
> anova(lm(Gain~Diet,data=pigs))
Analysis of Variance Table

Response: Gain

<table>
<thead>
<tr>
<th>Df</th>
<th>Sum Sq</th>
<th>Mean Sq</th>
<th>F value</th>
<th>Pr(&gt;F)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diet</td>
<td>2</td>
<td>1081.50</td>
<td>540.75</td>
<td>2.8582</td>
</tr>
<tr>
<td>Residuals</td>
<td>9</td>
<td>1702.75</td>
<td>189.19</td>
<td>1.1367</td>
</tr>
</tbody>
</table>
```

Now the \( F \)-test for a linear relation between Gain and Diet shows Diet is no longer explanatory - the effect is lost. Notice that the error variance estimate \( s^2 \) has jumped. In the `summary()` output above, for the model \( \text{Gain} \sim 1 + \text{Litter} + \text{Diet} \), we see \( s = 8.15 \) where the model \( \text{Gain} \sim 1 + \text{Diet} \) with no block structure has \( s = \sqrt{\frac{\text{RSS}}{(n - p)}} = \sqrt{1702.75/9} = 13.75 \). This jump in RSS, from 398 (in the \( \text{Gain} \sim 1 + \text{Litter} + \text{Diet ANOVA} \)) to 1702 (in the \( \text{Gain} \sim 1 + \text{Diet ANOVA} \)), is due to the extra variation across litter being treated as variation across treatment.

When we set up a design, we may have some choice in the assignment of treatments to subjects. The point of the trial is to see if the response depends on the treatment. Suppose the ‘treatment’ levels are ‘Give drug’ and ‘Give placebo’ and the response is some index of health. If we are allowed to choose which subject gets which treatment, we could distort the trial outcome by, for example, choosing to give the drug to subjects who for some reason are more likely to get better anyway. Wonderdrug! In order to avoid all traps of this kind, we typically assign the treatments to subjects completely at random. A design with everyone in one block, and treatments assigned to subjects at random, is called \textit{completely randomised}. If the subjects are in blocks, with \( m \) subjects in block \( a \), and we apply treatment \( t = 1, 2, ..., T \) to \( m_{a,t} \) different subjects in block \( a = 1, 2, ..., b \), then, for each treatment, we choose \( m_{a,t} \) subjects independently at random and without replacement from the \( m_a \) subjects in the \( a \)’th block. If \( m_{a,t} = m \), so that each treatment is applied to the same number of subjects in each block, then the design is called a \textit{randomised complete block design}. The treatments are distributed in a balanced way through the blocks.

The piglets data is balanced, as each of the four litters contains one piglet on each of the three diets, so \( b = 4, T = 3 \) and \( m_{a,t} = 1 \) for all \( a = 1, ..., 4 \), and \( t = 1, 2, 3 \). A \textit{randomised complete block design} is balanced in such a way that the block and treatment parameter estimates are independent, and so the treatments can be analyzed separately from blocks. Informally, in a completely balanced design the treatments are all tested under the same conditions, so when we compare treatments, it doesn’t matter what those conditions were. You have another example of a completely balanced design in the last problem of Problem sheet 2.

Sometimes the number of subjects \( m_a \) in a block is smaller than \( T \) the number of treatments. In that case the design will be \textit{incomplete}, since some blocks will have no instances of some treatments. An incomplete block design may still be balanced.
3. Model Checking and Model Selection

3.1. Model Checking. We are fitting a normal linear model \( y = X \beta + \epsilon \) with \( \epsilon \sim N(0, \sigma^2) \), and \( X \) a \( n \times p \) design matrix with one column \( X_i = (x_{1,i}, x_{2,i}, ..., x_{n,i})^T \) for each of the \( i = 1, 2, ..., p \) explanatory variables. Under this model the vector \( y \) of \( n \) responses is a realisation of the rv \( Y \sim N(X \beta, \sigma^2 I_n) \).

Model violations take many forms. A misspecified model is structurally inappropriate for essentially all responses. For example, the response \( y \) may not be a linear function of the linear predictors \( X \beta \), and we need to transform the response. This is discussed in a section below on the Box-Cox family of transformations. We may choose to work with a misspecified model if we have reason to believe that the biases in fitted values or parameter estimates are not large enough to invalidate the inference, for parameters of interest. On the other hand the problem may lie with the data. The normal linear framework may be good for most data points, but a few of the responses may have quite different causative factors. Such data points are called outliers. We try to identify them and then typically remove them from further analysis.

We have a range of validatory checks for model misspecification and outlier detection. The most straightforward check for linearity is to plot the explanatory variables against the response, as we do in a lattice plot, such as Figure 1. Variation in the response caused by variation in other variables may obscure the linear response to any single variable. Added variable plots (Problem sheet 1 Q1) help by focusing on the relation between the response and a single variable, removing variation due to other fitted variables. These check the linearity of the running mean. We have checks also on the independence and constant variance of \( \epsilon \), the errors. We saw that the fitted values \( \hat{y} = Hy \) (with hat matrix \( H = X(X^T X)^{-1}X^T \)) and the \( n \)-component vector of residuals \( e = y - \hat{y} \) are independent under the model, so a plot of residuals against fitted values should show no correlation.

Misfit. One weakness of the residuals/fitted-values plot, as a diagnostic tool, is that the residuals may have unequal variance under the model. A large residual \( e_k \) could be a sign that the \( k \)th data point is an outlier, but it might have a large variance as a consequence of the experimental design. The individual residuals \( e_i, i = 1, 2, ..., n \) are all mean zero normal rv, with variance matrix

\[
\text{var}(e) = \text{var}((I - H)y) = \sigma^2 (I - H)(I - H)^T = \sigma^2 (I - H),
\]

since \( H^2 = H = H^T \). The diagonal entries \( h_{kk}, k = 1, 2, ..., n \) of \( H \) control the variances of the components of \( e \) and \( \hat{y} \). The residual variances can be unequal, since \( \text{var}(e_k) = \sigma^2 (1 - h_{kk}) \).

**Exercise** Show that \( \text{var}(\hat{y}) = \sigma^2 H \).

The standardised residuals \( r_k, k = 1, 2, ..., n \) have zero mean and equal variance. They are obtained by scaling the residuals:

\[
r_k = \frac{e_k}{s\sqrt{1 - h_{kk}}},
\]

with \( s^2 \) an unbiased estimate of \( \sigma^2 \), under the normal linear model. They (the \( r_k \)) have approximately unit variance, and can be compared with standard normal
variables, an approximation which will be good at large \( n - p \). Because \( s \) and \( e \) are correlated, we cannot easily compute the distribution of the standardised residuals.

**Exercise** Show that the standardised residuals \( r \) are (like the residuals \( e \)) independent of the fitted values, \( \hat{y} \).

We can make normal qqplots of standardised residuals, and plot them (the \( r_k \)) against fitted values \( \hat{y} \). We often mistakenly fit a model of constant variance to data in which the variance of the response increases with the underlying mean. This model mispecification is witnessed by a trend to increasing variance in \( e' \) with \( \hat{y} \).

A response \( y_k \) which generates a relatively large residual \( e_k \) need not be an outlier, since \( \text{var}(e_k) = \sigma^2(1 - h_{kk}) \), and \( 1 - h_{kk} \) may be relatively large. We might expect the standardised residuals \( r \) to be a good basis for outlier detection, since these should have variance about one under the normal linear model. Standardised residuals exceeding two (standard deviations) are large. The problem here is that \( s^2 = e'r/e \), the denominator in the expression for \( r_k \), is computed from the residuals \( e \) themselves. A response \( y_k \), which is truly outlying, may have a large residual \( e_k \), but this will inflate our estimate \( s^2 \) for \( \sigma \), and we may end up with a moderate standardised residual \( r_k \). A bad response with an \( h_{kk} \) value close to one has a low variance. It ‘pulls’ the fitted surface towards itself, so that \( e_k \) is small, and again \( r_k \) is not obviously large. What to do?

We can treat this problem using an idea related to ‘cross-validation’. We remove response \( y_k \) and row \( k \), \( \mathbf{x}_k = (X_{k,1}, ..., X_{k,p})^T \) say, from the data and compare the fitted values \( \hat{y} \) we got with all the data with the fitted values \( \hat{y}_{-k} \) we get when \( \mathbf{x}_k, y_k \) are removed. Denote by \( y_{-k} \) and \( X_{-k} \) the remaining response and design data with \( y_k \) and \( \mathbf{x}_k \) removed. Let \( \beta_{-k} = (X_{-k}^TX_{-k})^{-1}X_{-k}^Ty_{-k} \) give the new parameter estimates. The \( k \)th deletion, or studentised residual is

\[
r_k' = \frac{y_k - \mathbf{x}_k\hat{\beta}_{-k}}{\text{std.err}(y_k - \mathbf{x}_k\hat{\beta}_{-k})}.
\]

It may be shown (Problem Sheet 3 Q2) that \( r_k' \sim t(n - p - 1) \) and \( r' \) and \( \hat{y} \) are independent. Our studentised residuals have equal variance, known distribution, and can be compared to a standard normal (using for example a qqplot). We can plot \( r' \) against \( \hat{y} \). Any visible correlation is a sign of model mispecification, and data points with \( |r_k'| > 2 \) show misfit, and are possible outliers.

We will now derive a useful auxiliary formula for the studentised residuals. It may be shown that

\[
\hat{\beta}_{-k} = \hat{\beta} - (X^TX)^{-1}X^T\frac{e_k}{1 - h_{kk}}.
\]

**Exercise** Verify this expression for \( \hat{\beta}_{-k} = (X_{-k}^TX_{-k})^{-1}X_{-k}^Ty_{-k} \). Quote the Woodbury formula

\[
(X_{-k}X_{-k})^{-1} = (X^TX)^{-1} + (X^TX)^{-1}\mathbf{x}_k(1 - \mathbf{x}_k(X^TX)^{-1}\mathbf{x}_k)X_k^T(X^TX)^{-1}
\]

without proof, and use the fact that \( X_{-k}^Ty_{-k} = X^Ty - X^T(0, ..., 0, y_k, 0, ..., 0)^T \).

**Exercise** Using the formula above for \( \hat{\beta}_{-k} \), show that

\[
y_k - \mathbf{x}_k\hat{\beta}_{-k} = \frac{e_k}{1 - h_{kk}}.
\]
and hence show
\[ \text{var}(y_k - x_k \hat{\beta}_{-k}) = \sigma^2 / (1 - h_{kk}). \]

Estimating \( \sigma^2 \) as \( s_{-k}^2 \), the residual standard error with the \( k \)th observation removed, we have studentised residuals
\[ r'_k = \frac{e_k}{s_k \sqrt{1 - h_{kk}}}. \]

**Exercise** The residual variance is \( (n - p)s^2 = y^T y - y^T X \hat{\beta} \). The \( k \)th deletion residual-variance is \( (n - p - 1)s^2_{-k} = y^T y - y_k^2 - (y^T X - y_k x_k) \hat{\beta}_{-k} \). Using the formulae above, show that
\[ (n - p - 1)s^2_{-k} = (n - p - r^2_k)s^2 \]
and hence obtain the useful form
\[ r'_k = r_k \sqrt{\frac{n - p - 1}{n - p - r^2_k}}. \]

This formula is important computationally, since it shows that we can compute the studentised residuals without making \( n \) linear regressions for the \( n \) deletions, just using the results of the primary regression.

3.1.1. **Leverage.** The diagonal entries \( h_{kk} = H_{kk} \) of the hat matrix are called the leverage components. Since \( h_{kk} = \text{var}(y_k)/\sigma^2 \) we have \( 0 \leq h_{kk} \leq 1 \). Since \( \text{var}(e_k) = \sigma^2(1 - h_{kk}) \), a point with leverage \( h_{kk} \) close to one has low variance: since \( E(e_k) = 0 \) the fitted surface \( x \hat{\beta} \) must be pulled close to the \( k \)th response \( y_k \). If \( x_k, y_k \) is an outlier with high leverage, then predictions \( x \hat{\beta} \) for \( x \) near \( x_k \) will be poor.

How big is big, when it comes to leverage? The ’average’ leverage for a \( n \times p \) design is \( \bar{h} = p/n \), as we will see shortly, so points with leverage values above \( 2p/n \) get special attention. Since they are having more impact on the final fit than other points, it is important that they are not outliers. The average leverage is given by the trace of the hat matrix \( H \), so
\[ \sum_{k=1}^{n} h_{kk} = \text{trace}(H) \]
\[ = \text{trace}(X(X^T X)^{-1} X^T) \]
\[ = \text{trace}(X^T X(X^T X)^{-1}), \]
by the cyclic permutation property of the trace \( \text{trace}(ABC) = \text{trace}(CAB) \). It follows that
\[ \frac{1}{n} \sum_{k=1}^{n} h_{kk} = \frac{\text{trace}(I_p)}{n} \]
\[ = \frac{p}{n}. \]
Alternatively, \( \text{trace}(H) = p \), as \( H \) has the eigenvalue \( \lambda = 1 \) repeated \( p \) times, and the eigenvalue \( \lambda = 0 \) repeated \( n - p \) times, since there are \( p \) linearly independent vectors \( v = e_1, ..., e_p \) satisfying \( H v = v \) and \( n - p \) vectors \( u = e_{p+1}, ..., e_n \) satisfying \( H u = 0 \). We saw this when we made the expansion of \( y \) in \( e_1, ..., e_n \) in Section 2.
3.1.2. Influence. A point with high leverage need not do much damage, if it lies close to the fitted surface through other points - the surface would have gone close to it anyway. Such a point is said to have high leverage but low influence. Highly influential points shift the fitted surface far from where it would have lain if the influential point were not included. The Cook’s distance for a point \( \mathbf{x}_k, y_k \) is a measure of influence given by the sum of the squares of the shift in fitted values when point \( k \) is removed. Now the Cook’s distance for the \( k \)th data point is defined to be

\[
C_k = \frac{(\hat{y} - \bar{y}_k)^T (\hat{y} - \bar{y}_k)}{p s^2}.
\]

It may be shown, using the formula in Section 3.1 above, that

\[
C_k = \frac{r^2_k h_k}{p(1 - h_{kk})}.
\]

Our intuition is that high influence occurs where there is misfit and large leverage. The factor \( h_{kk}/(1 - h_{kk}) \) rises with increasing leverage. The factor \( r^2_k \) (standardised residual squared) is related to misfit. A rough rule of thumb is that \(|r_k| > 2\) and \( h_{kk} \) are separate causes for concern, so points with Cook’s distance exceeding

\[
C_k \gtrsim \frac{8}{n - 2p}
\]

have high influence. Such points are generally outliers, and are removed from the analysis.

3.1.3. Model checking, example.

Example 3.1. The `data(swiss)` dataset is described in the R documentation as follows.

The data give a standardized fertility measure and socio-economic indicators for each of 47 French-speaking provinces of Switzerland at about 1888.

Switzerland, in 1888, was entering a period known as the Demographic Transition, its fertility was beginning to fall from the high level typical of underdeveloped countries. The data come from Mosteller, F. and Tukey, J. W. (1977) who give the original source.

The `swiss` data frame has 47 observations on 6 variables, each of which is given as a percentage. The columns are:

- **Fertility**: a standardized fertility measure
- **Agriculture**: % of males involved in agriculture as occupation
- **Examination**: % draftees receiving highest mark on army examination
- **Education**: % draftees receiving education beyond primary school
- **Catholic**: % catholic (ie, not protestant)
- **Infant.Mortality**: % live births who live less than 1 year

Which variables are explanatory for Fertility?

We can map the percentages (0, 100) into \((-\infty, \infty)\) using the logistic transformation \( x \leftarrow \log(x/(100 - x)) \). Since 0 and 100 may appear, we modify this to something like \( x \leftarrow \log((1 + x)/(101 - x)) \), checking for linearity between mapped variable and response.

\[
> \text{data(swiss)}
\]

```r
> head(swiss) #first few rows
```

Fertility  Agriculture  Examination  Education  Catholic  Infant.Mortality
Notice that each row (i.e., each data point) is named according to the Swiss province that data point measures. The transformed data are displayed in Figure 7. We will (i) fit a normal linear model, (ii) look for outliers, (iii) remove them, refit, and look again for outliers, (iv) make a model reduction to something simple which nevertheless predicts the response and finally (v) check again for outliers in the reduced model.

We can find the points of high influence and display them as in Figure 7. The R code to do this is available in the course website in the file L7.R. We apply the `cooks.distance()` function to the `lm()` output (and `round(.,3)` to 3dp). See the top panel of Figure 8. We see that that the data points for V. De Geneve and La Vallee have high influence. They must have high misfit (as measured by standardised residual) or high leverage, or both. The leverages are the diagonal entries in the hat matrix (rounded to 3 dp). See the centre panel of Figure 8. The data points for V. De Geneve and La Vallee have high leverage in the sense that they exceed twice the mean leverage \( p/n \). How about misfit? The numbers are displayed in the bottom panel of Figure 8. V. De Geneve and Rive Gauche are above threshold (which equals 2) for high misfit. Rive Droite and La Vallee have highish misfit. However, Rive Gauche and Rive Droite do not have high enough leverages (0.1 and 0.178) to make them a problem. On the other hand La Vallee has a particularly high leverage (0.379), so its milder misfit (1.7) is still a cause for concern. There are two diagnostic functions we haven’t mentioned: `rstudent()` and `fitted.values` act on the `lm()` output to give studentised residuals \( r' \) and fitted
Figure 7. Pairs plot for Swiss fertility data. Percentage data has been mapped monotonically to the interval $(-\infty, \infty)$. 
> #V. De Geneve and La Vallee are points of high influence
> cooks.distance(sw1.lm)

<table>
<thead>
<tr>
<th></th>
<th>Courtelary</th>
<th>Delemont</th>
<th>Franches-Mnt</th>
<th>Moutier</th>
<th>Neuveville</th>
<th>Porrentruy</th>
<th>Broye</th>
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<tbody>
<tr>
<td>Cook's Distance</td>
<td>0.021</td>
<td>0.002</td>
<td>0.043</td>
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<td>0.001</td>
<td>0.005</td>
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<td>La Chauxfnd</td>
<td>Le Locle</td>
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<tr>
<td>Cook's Distance</td>
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<td>0.082</td>
<td>0.035</td>
<td>0.007</td>
<td>0.003</td>
<td>0.013</td>
<td>0.012</td>
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<td>Rive Droite</td>
<td>Rive Gauche</td>
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<td></td>
</tr>
<tr>
<td>Cook's Distance</td>
<td>0.007</td>
<td>0.000</td>
<td>0.464</td>
<td>0.118</td>
<td>0.081</td>
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> \frac{8}{(n-2*p)}
[1] 0.229

> #V. De Geneve and La Vallee are points of high leverage
> hatvalues(sw1.lm)

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<th>Courtelary</th>
<th>Delemont</th>
<th>Franches-Mnt</th>
<th>Moutier</th>
<th>Neuveville</th>
<th>Porrentruy</th>
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<tbody>
<tr>
<td>Hat Values</td>
<td>0.118</td>
<td>0.160</td>
<td>0.177</td>
<td>0.052</td>
<td>0.084</td>
<td>0.164</td>
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<td>0.128</td>
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<td>0.145</td>
<td>0.094</td>
<td>0.092</td>
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<td>Lavaux</td>
<td>Morges</td>
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<tr>
<td>Hat Values</td>
<td>0.114</td>
<td>0.178</td>
<td>0.056</td>
<td>0.087</td>
<td>0.379</td>
<td>0.118</td>
<td>0.072</td>
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<td>Paysd'enhaut</td>
<td>Rolle</td>
</tr>
<tr>
<td>Hat Values</td>
<td>0.105</td>
<td>0.068</td>
<td>0.117</td>
<td>0.183</td>
<td>0.114</td>
<td>0.219</td>
<td>0.080</td>
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<td>Martigwy</td>
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<tr>
<td>Hat Values</td>
<td>0.052</td>
<td>0.069</td>
<td>0.223</td>
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<td>0.138</td>
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<tr>
<td>Hat Values</td>
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<tr>
<td>Hat Values</td>
<td>0.057</td>
<td>0.149</td>
<td>0.332</td>
<td>0.178</td>
<td>0.100</td>
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> \frac{2*p}{n}
[1] 0.255

> #V. De Geneve and Rive Gauche have higher misfit than Rive Droite and La Vallee.
> rstandard(sw1.lm)

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<tr>
<td>R Standard Deviation</td>
<td>0.967</td>
<td>0.281</td>
<td>1.099</td>
<td>1.380</td>
<td>1.485</td>
<td>-1.396</td>
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<td>1.651</td>
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<td>0.771</td>
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<td>Morges</td>
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<tr>
<td>R Standard Deviation</td>
<td>-0.469</td>
<td>-1.124</td>
<td>0.329</td>
<td>-0.386</td>
<td>1.700</td>
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<td>Rolle</td>
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<tr>
<td>R Standard Deviation</td>
<td>-1.656</td>
<td>-0.324</td>
<td>-0.459</td>
<td>-1.052</td>
<td>0.033</td>
<td>-0.567</td>
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<tr>
<td>R Standard Deviation</td>
<td>-0.581</td>
<td>-0.882</td>
<td>-0.776</td>
<td>-1.112</td>
<td>-0.778</td>
<td>-0.522</td>
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<tr>
<td>R Standard Deviation</td>
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<td>1.447</td>
<td>1.455</td>
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<td>-0.272</td>
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<tr>
<td>R Standard Deviation</td>
<td>0.938</td>
<td>-0.109</td>
<td>-2.365</td>
<td>-1.806</td>
<td>-2.097</td>
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</table>

**Figure 8.** (Top) Influence, (Centre) Leverage and (Bottom) Misfit.
The theoretical quantiles are compared with the sample quantiles of the residuals.

The residuals are displayed as a QQ-plot (quantile-quantile plot) to assess the normality of the residuals. Outliers can be identified in this plot.

The Cook's distance is calculated to identify influential observations. Points with high Cook's distance are considered potential outliers.

The standardized residuals are also plotted against the fitted values to check for patterns or trends that might indicate model misspecification.

If any points are identified as outliers, the model can be refitted without these points to see if the model improves.

### Code

```r
> i <- cooks.distance(sw1.lm) > (8/(n-2*p))
> swr <- sw[-which(i),]
> nr <- dim(swr)[1];
```
```r
> swr1.lm<-lm(Fertility~Infant.Mortality+Examination+Education+Catholic+Agriculture, + data=swr)
> summary(swr1.lm)
...  
Coefficients:  
   Estimate Std. Error t value Pr(>|t|)  
(Intercept) 82.5002 12.3752 6.667 6.17e-08 ***  
Infant.Mortality 26.9630 8.2567 3.266 0.00228 **  
Examination -6.7927 3.5219 -1.929 0.06107 .  
Education -5.9604 2.5509 -2.337 0.02469 *  
Catholic 1.0270 0.6005 1.710 0.09514 .  
Agriculture -2.8355 1.7661 -1.606 0.11645  
Residual standard error: 7.866 on 39 degrees of freedom  
F-statistic: 10.41 on 5 and 39 DF, p-value: 2.139e-06  
```

Considering we removed just two of 47 data points, the parameters and significance levels have changed a fair bit. For example, Examination and Catholic are now marginally significant. Before we continue, let's repeat the diagnostics. Outliers can mask outliers, so we may find new points of high influence in the reduced data set. The new diagnostics are plotted in Figure 10. The qqplot was already acceptable, and is no worse. The plot of studentised residuals against fitted values is improved, with no discernable trend. No points of high influence have been exposed in the reanalysis. La Chauxdfnd has now a slightly elevated leverage, but its misfit is mild so its influence is not large. Conversely there are some points with studentised residuals exceeding two, however we can assume their influence remains slight.

Notice that Examination and Education have similar parameter values (around -6 to -7). Looking at the pairs plot above they are clearly correlated (as you might expect, thinking about the variables). This is an instance of variance inflation due to correlation (ie, near linear dependence (Section ?? and Problem Sheet 3 Q3)). If we remove one of these variables it is likely that the Std. Error of the other will drop substantially. I will go straight to the simplest model that seems to work, and make an F-test to drop Education + Catholic + Agriculture from the model.
```
> swr0.lm<-lm(Fertility~Infant.Mortality+Examination,data=swr)
> summary(swr0.lm)
...  
Coefficients:  
   Estimate Std. Error t value Pr(>|t|)  
(Intercept) 95.133 10.871 8.751 5.15e-11 ***  
Infant.Mortality 32.982 8.009 4.118 0.000175 ***  
Examination -11.811 2.100 -5.624 1.38e-06 ***  
Residual standard error: 8.181 on 42 degrees of freedom  
F-statistic: 21.1 on 2 and 42 DF, p-value: 4.545e-07  
```

```
shows that there are at least no points of high influence.

```r
> any(cooks.distance(swr0.lm) > (2*pr)/nr) # there are 2 parameters in the reduced model
[1] FALSE
```

Finally, does the reduced model have any outliers? A quick check:

```r
> pr <- 2 # there is no reference to support the more complex model. Notice that the p-value for Examination above is now far more significant.
```

The p-value comparing the reduced model (Model 1, above) to the full model (Model 2, above) is 0.11 so there is no evidence to support the more complex Model 2. Fertility ~ Infant.Mortality + Examination + Education + Catholic + Agriculture +

The p-value comparing the reduced model (Model 1, above) to the full model (Model 2. Fertility ~ Infant.Mortality + Examination + Education + Catholic + Agriculture +)

Finally, does the reduced model have any outliers? A quick check:

```r
> pr <- 3 # there are 3 parameters in the reduced model
```

```r
> any(cooks.distance(swr0.lm) > (2*pr)/nr)
[1] FALSE
```

shows that there are at least no points of high influence.
2009 notes updated to here
3.2. **Model Selection.** In some settings we aim to filter out from some original, possibly large, set of explanatory variables the ones that are explanatory for the response. Where variables are correlated, this may not be possible - which of a group of near linearly dependent variables do we drop? Physical considerations are very important. Does the model make sense? We have to go some way down the road to understanding the application domain in order to make sense of the variables and their interactions, and make sensible selections of subsets of variables.

3.2.1. *Model choice v. Exploratory data analysis.* The easiest case is where we go into the analysis with some preconceived hypothesis. The trees example, Example 2.2, worked that way. We modeled the tree as something like a cylinder, and that gave us a clear hypothesis about the relation between volume, girth and height. We fit the model and test the hypothesis.

More commonly we look at the parameters in the fitted model, observe that some are not significant, fit a range of reduced and added-variable plots, and formulate some hypothesis about the relations between variables. This leads to a test for the significance of some subset of explanatory variables which is informed by the data. We have in effect made many tests, but report just the final one. This introduces a hazard for multiple testing, which can sometimes be corrected, see Ripley (2002) section 6.10. If we don’t correct, and we hardly every do, then, paraphrasing Davison (2003) section 8.7 in the subsection ‘Inference after model selection’, “…the only covariates for which subsequent inference using the standard confidence intervals is reliable are those for which the evidence for inclusion is overwhelming”. It sometimes happens that we frame a hypothesis by looking at the data, but then realise that the hypothesis has some natural physical meaning. I can tell you that the cylinder model in the trees example (Example 2.2) was not the first model I tried for that data, but emerged in the analysis. However, the final model is so natural, that we could easily imagine a scientist going into the analysis with precisely this hypothesis (we just did).

Sometimes we use *automatic* variable selection. This means, essentially, any model selection procedure not guided by physical considerations for variable meaning. We might search over all subsets of variables for the ‘best’ reduced model. One choice is to look for the largest fully significant model. No variable, or set of variables, can be dropped, and if we add variables we have a model with some non-significant variables. There may be many such models. When the model space is very large we may search using stepwise methods. In *Backwards elimination* we start with the full set of variables, and successively drop the least significant, until we have a fully significant set. There is a *Forwards selection* scheme, which adds the next most significant variable. We sometimes sort the variables in an ANOVA table so that the $p$-values on the RHS of the table show in effect the steps of forward selection. One advantage of backwards selection, is that the initial estimator for $s^2$ is from the fit for all the variables, so it is not biased (upwards) by significant variables which are not included. If forwards selection started with an estimate of $s^2$ based on just the one or two variables in the initial model, then it might be a large overestimate, since variation in the response $y$ due to variations in significant explanatory variables would inflate $s^2$, and levels of significance would suffer accordingly. This is the reason we use the same $\text{RSS}_{1:p}/(n - p)$ divisor in every row of the $F$-column of an ANOVA, rather than $\text{RSS}_{1:k}/(n - (k - k_{-1}))$. 

One strategy is to search over all models and optimise some measure of the relative worth of models. The measure must somehow penalise models which are too complex or too simple, since both are poor for prediction. The AIC is just such a model choice criterion. The AIC-approach is set out in the next section.

Automatic methods are exposed to the hazard for multiple testing mentioned above. If we make a lot of tests on a large number of non-significant variables we may find marginally significant variables where there are none. The approach is justified as part of exploratory data analysis. The hope is that the method will turn up some physically natural set of explanatory variables.

3.2.2. The AIC. One idea (there are lots, but this is core) is to consider what happens when new data \(Y'\) come along. We have parameter MLE’s \(\hat{\beta}(Y)\) and \(\hat{\sigma}_{\text{MLE}}(Y)\) computed using the old data \(Y\) and some particular design \(X\). If the model is a good model, then the loglikelihood \(\ell(\beta(Y), \hat{\sigma}_{\text{MLE}}(Y); Y')\) for \(\hat{\beta}\) and \(\hat{\sigma}_{\text{MLE}}\) in the new data should be large. This should “usually” hold, i.e., the average over \(Y\) and \(Y'\) of \(\ell(\hat{\beta}(Y), \hat{\sigma}_{\text{MLE}}(Y); Y')\) should be large. If

\[
C = -2\ell(\hat{\beta}(Y), \hat{\sigma}_{\text{MLE}}(Y); Y')
\]

then we like models that make \(\mathbb{E}(C)\) small. The expectation here is over realisations of the old \(Y\) and new \(Y'\) data. Davison (2003) section 8.7.3, page 402, gives a somewhat better motivation on this crucial point, linking the analysis to ideas of cross-validation.

In an earlier version of these notes I set the following as an exercise. Here is the detail. Omit this at first reading, and jump to the definition of the AIC.

We will now derive a consistent estimator, AIC, for \(\mathbb{E}(C)\). In the following, \(v^2\) means the scalar inner product \(v^T v\). We will need one auxiliary result, namely, that if \(Z \sim \chi^2(\nu)\) with \(\nu > 2\) then \(\mathbb{E}(1/Z) = 1/(\nu - 2)\). Looking at \(\ell(\hat{\beta}(Y), \hat{\sigma}_{\text{MLE}}(Y); Y)\) in Section 2.2, we see we have here

\[
-2\mathbb{E}(\ell(\hat{\beta}(Y), \hat{\sigma}_{\text{MLE}}(Y); Y')) = \mathbb{E} \left( n \log (\hat{\sigma}_{\text{MLE}}^2(Y)) + \frac{(Y' - X\hat{\beta}(Y))^2}{\hat{\sigma}_{\text{MLE}}^2(Y)} \right).
\]

We will deal with the second term first.

\[
\mathbb{E} \left( \frac{(Y' - X\hat{\beta}(Y))^2}{\hat{\sigma}_{\text{MLE}}^2(Y)} \right) = \mathbb{E} \left( \frac{(Y' - X\beta + X\beta - X\hat{\beta}(Y))^2}{\hat{\sigma}_{\text{MLE}}^2(Y)} \right)
\]

\[
= \mathbb{E}_Y \left( \frac{n\sigma^2}{\hat{\sigma}_{\text{MLE}}^2(Y)} \right) + \mathbb{E}_Y \left( \frac{(X\beta - X\hat{\beta}(Y))^2}{\hat{\sigma}_{\text{MLE}}^2(Y)} \right) + \text{constant terms},
\]

since \(\mathbb{E}_Y((Y' - X\beta)^2) = n\sigma^2\) and

\[
\mathbb{E} \left( 2(Y' - X\hat{\beta})^T (X\beta - X\hat{\beta}(Y)) \right) = 0
\]

using the \(Y'\) expectation. Recall that \(\hat{\sigma}_{\text{MLE}}^2 = \text{RSS}/n\). Now \(X\beta - X\hat{\beta}(Y) = \mathbb{E}(\hat{Y}) - \hat{Y}\) and \(\hat{Y}\) and RSS are independent, so

\[
\mathbb{E} \left( \frac{(X\beta - X\hat{\beta}(Y))^2}{\hat{\sigma}_{\text{MLE}}^2} \right) = \mathbb{E} \left( (\mathbb{E}(\hat{Y}) - \hat{Y})^2 \right) \mathbb{E} \left( \frac{n}{\text{RSS}(Y)} \right).
\]
Using \( \text{var}(\hat{Y}) = \sigma^2H \) we have

\[
E \left( (X\beta - X\hat{\beta}(Y))^2 \right) = \sum_{k=1}^{n} \sigma^2 h_{kk}
\]

\[
= \sigma^2 p.
\]

Also, \( \text{RSS}/\sigma^2 \) has a \( \chi^2(n - p) \) distribution, so \( E(\sigma^2/\text{RSS}) = 1/(n - p - 2) \), using our auxiliary result, so

\[
E \left( \frac{(Y' - X\hat{\beta}(Y))^2}{\hat{\sigma}^2_{\text{MLE}(Y)}} \right) = \frac{n^2}{n - p - 2} + \frac{np}{n - p - 2}.
\]

We need now to deal with the first term in the expression above for \( E(C) \). If \( D \) is the deviance

\[
D(Y) = -2\ell(\hat{\beta}(Y), \hat{\sigma}^2_{\text{MLE}(Y)}; Y)
\]

then

\[
E(D) = E \left( n \log(\hat{\sigma}^2_{\text{MLE}(Y)}) + n \right)
\]

from Section 2.2. We have then

\[
E(C) = E(D) + \frac{n^2}{n - p - 2} + \frac{np}{n - p - 2} + \text{constant terms}.
\]

Expanding the last two terms in \( p/n \) about zero, we have

\[
E(C) = E(D) + 2p + O(p/n) + \text{terms not depending on } p.
\]

Now \( D(y) \) is an unbiased estimator for \( E(D) \), and \( 2p \) is, up to a constant term, twice the number of parameters, so the objective function we seek to minimise in our search for the optimal normal linear model is

\[
\text{AIC} = -2\ell(\hat{\beta}, \hat{\sigma}^2_{\text{MLE}}; y) + 2 \times \text{number of parameters}.
\]

We derived this for normal linear models only, but it is in fact the relevant criterion for model choice in a much wider setting. In our setting

\[
\text{AIC} = n \log(\text{RSS}/n) + 2p
\]

where we have dropped constants, such as \( n \), which are fixed for given data. There is another, closely related criterion, called Mallow’s \( C_p \), which is relevant for regression. It may be shown that

\[
C_p = \frac{\text{RSS}}{\sigma^2} + 2p
\]

is an approximation to the AIC (itself an approximation to \( E(C) \)). We do not in general know \( \sigma^2 \), so it is estimated using the full model (and thereby avoiding bias from any significant variables in the full model). Mallows \( C_p \) has motivation beyond simply approximating the AIC.

**Example 3.2.** Returning to the trees data in Example 2.2, I mentioned in the text above that I tried a few models before settling on the (admittedly fairly obvious) product form in Example 2.2. Here is part of the model exploration I did. Let \( y_i \) be the volume, \( x_{i,1} \) be the girth, and \( x_{i,2} \) the height. Denote by \( \text{RSS}(\alpha, \beta_1, \beta_2, \gamma) \) the residual sum of squares for the linear model

\[
Y_i = \alpha + \beta_1 x_{i,1} + \beta_2 x_{i,2} + \gamma x_{i,1}^2 x_{i,2} + \epsilon_i.
\]
with $\epsilon \sim N(0, \sigma^2)$. The following results were obtained from regression of four different models: $\text{RSS}(\alpha, \beta_1, \beta_2, \gamma) = 179.3$, $\text{RSS}(\alpha, 0, 0, \gamma) = 180.2$, $\text{RSS}(\alpha, \beta_1, \beta_2, 0) = 219.4$, $\text{RSS}(0, 0, 0, \gamma) = 181$. Carry out variable selection.

Here we have data for $n = 31$ trees, so computing the AIC’s, 

```r
> n<-31; p<-c(4,2,3,1); RSS<-c(179.3,180.2,219.4,181.0);
> n*log(RSS/n)+2*p
```

we can make a table of AIC’s by model

<table>
<thead>
<tr>
<th>Model</th>
<th>p</th>
<th>RSS</th>
<th>AIC</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\alpha, \beta_1, \beta_2, \gamma$</td>
<td>4</td>
<td>179.3</td>
<td>62.4</td>
</tr>
<tr>
<td>$\alpha, \gamma$</td>
<td>2</td>
<td>180.2</td>
<td>58.56</td>
</tr>
<tr>
<td>$\alpha, \beta_1, \beta_2$</td>
<td>3</td>
<td>219.4</td>
<td>66.66</td>
</tr>
<tr>
<td>$\gamma$</td>
<td>1</td>
<td>181.0</td>
<td>56.70</td>
</tr>
</tbody>
</table>

and the winner is... the model in the last line with the least AIC, $E(Y_i) = \gamma x_{i,1}^2 x_{i,2}$.

Although we did this by hand, this is in effect automatic variable selection, with no regard for the physical content of the model. The comments above about multiple testing apply here. An undesirable feature of the final model, is that we end up with the interactions without the lower order terms. The product form in the original example deals with this issue.

Note that the `step()` function in R, applied to the output of `lm()`, will carry out automatic backwards variable selection using the AIC, and report all these numbers. See the last sections of `L7.R` for another example of guided, and automatic variable selection, using the AIC, applied to the `swiss` fertility data.

3.3. **Two model revision strategies.** The following two subsections treat a couple of discrete topics concerning model revision. Suppose that, in the course of our diagnostic analysis we find that the errors are non-normal, or correlated. It may be possible to make a linear transformation of the model to get iid normal mean zero errors $\epsilon$. We transform the data, fit, and then invert the transformation to get results for the original model. This is weighted regression, which we look at in the next part section.

We may find that the response $y_i$ is not linearly related to the linear predictor $X\beta$. Can we find a transform which restores linearity? It would be convenient to take a family of transformations, and choose the transform which best restores linearity. This is the Box-Cox approach, presented in the second part section.

3.3.1. **Weighted Regression.** We get linear normal data with variance varying from observation to observation. If $\sigma_i^2$ is the variance for the $i$th observation, then

$$Y_i = X\beta + \epsilon_i \quad \text{for } i = 1, 2, ..., n$$

as before, but this time the independent errors $\epsilon_i$ are distributed $\epsilon_i \sim N(0, \sigma_i^2)$. There are three cases to consider: $\sigma_i^2$ unequal and unknown, $\sigma_i^2$ unequal and known, and $\sigma_i^2 = \sigma^2/\omega_i$, with $\omega_i$ known, but $\sigma^2$ unknown.

The first case cannot be treated without some assumption on the joint distribution of the $\sigma_i$, since we have a variance parameter for each datum, which we cannot estimate. The second case can be treated in the same framework as the third (see the 2nd exercise in Problem sheet 1 where $\sigma^2$ is known).

What is the motivation for considering the third case? If, for $i = 1, 2, ..., n$, $Y_i$ is actually the outcome of $n_i$ independent measurements, $Y_{i,j} \sim N(x_i\beta, \sigma^2)$, then we
may pool the data, so \( Y_i = n_i^{-1} \sum_j Y_{i,j} \) so that \( \text{var}(Y_i) = \sigma^2/n_i \). This works for the more general case where \( Y_{i,j} \) are iid for \( j = 1, 2, ..., n_i \), with mean \( x_i \beta \) and variance \( \text{var}(Y_{i,j}) = \sigma^2 \), but not normal. By the CLT, the approximation \( Y_i \sim N(x_i \beta, \sigma^2/n_i) \) may be good.

We can map the weighted variance problem onto our original problem. If \( W = \text{diag}(w_1, ..., w_n) \) (i.e., an \( n \times n \) matrix with square-roots of \( w_i \)'s on the diagonal, and otherwise zero) and we define 'data' \( Y' = W^{1/2}Y \) and a 'design' \( X' = W^{1/2}X \) then

\[
y' = X' \beta + \epsilon'
\]

with \( \epsilon' \sim N(0, \sigma^2 I_n) \). We are back to our standard problem. The weighted least squares estimators for \( \beta \) and \( \sigma^2 \) are

\[
\hat{\beta} = (X'W^{-1}X')^{-1}X'W^{-1}Y',
\]

and

\[
s^2 = \frac{(Y' - X' \hat{\beta})'W(Y' - X' \hat{\beta})}{n - p}.
\]

Exercise suppose \( Y = X \beta + \epsilon \), and the errors are correlated, with \( \epsilon \sim N(0, \Sigma) \) with \( \Sigma \) a \( n \times n \) positive definite covariance matrix. Give formulae for the weighted least squares estimators \( \hat{\beta} \) and \( s^2 \) for \( \beta \) and \( \sigma^2 \). Ans: Suppose there is a transformation \( L \) such that \( \text{var}(L \epsilon) = \sigma^2 I_n \) for some \( \sigma^2 > 0 \). Now it is the same as the last problem. Is there such a transformation?

Example 3.3. The monthly prices quoted in Example 2.4 and Example 2.5 are averages. The number of houses used to form the averages are given (as a scatterplot) in the lattice plot in Figure 4. The numbers are quite large (60 is typical) so we might expect the central limit theorem to give us averages which are approximately normally distributed, scattered about a running (straight-line regression) mean. Let \( w_i = \text{sales}[i] \). I return to the house types Semi-Detached and Terraced and the model of Example 2.4, but weighted,

\[
y_k \sim \alpha + \alpha_T g_k, T + \gamma_M x_k, M + \epsilon_k, \quad \epsilon_k \sim N(0, \sigma^2/w_i),
\]

and I am looking to remove all the variance variation by house type and month by taking into account the sales weighting. We can fit this model in R using the weights option in the \texttt{lm()} function. We set weights = \texttt{sales}, and otherwise the variables are set up as before.

```r
> #weighted by number of sales
> ohp.wlm<-lm(price~month+type,weights=sales,data=ohp)
> summary(ohp.wlm)
```

Coefficients:

|            | Estimate | Std. Error | t value | Pr(>|t|) |
|-------------|----------|------------|---------|----------|
| (Intercept) | 157.94603| 3.29306    | 47.963  | <2e-16 *** |
| month       | 1.70720  | 0.05509    | 30.987  | <2e-16 *** |
| typeTerraced| -1.95419 | 3.06955    | -0.637  | 0.525    |

In this fit Semi-Detached gives the baseline. So, has our weighting corrected the variance? We have been using R to plot residuals against fitted values, as a check for goodness of fit. Looking at Figure 11. We do see an improvement in the pattern of residuals, so we have a 'better' fit. The residuals and fitted values should be independent and the residuals should have homogeneous variance. There is a funnel-shape structure to the scatter plot at left in Figure 11. The scatter plot at right is somewhat more even. The weighting brings the outlier into the
3.4. The Box-Cox family of transformations. Suppose we have data \(y, X_1, \ldots, X_p\) with \(y_k \geq 0\). The response may not be a linear function of the linear predictor. A simple transformation is enough to set things right. The family of transformations 

\[ y'_k = \left( y_k^\lambda - 1 \right) / \lambda \]

for \(k = 1, 2, \ldots, n\) can fit in your pocket, and includes many of the

normal range (sales have fallen lately - the weighting corrects the recent part of the variance trend - but there is still some increase in variance with fitted values at small weighted fitted values - this is the trend to increased variance with price, which we haven’t treated).

Note the scaling on the \(x\) and \(y\) axes at right in Figure 11. We have \(\hat{Y}' = W^{1/2}X\hat{\beta}\) so the weighted residuals \(e' = Y' - \hat{Y}' = W^{1/2}(Y - X\hat{\beta})\) have variance matrix \(\sigma^2(I_n - H)\) with \(H = W^{1/2}X(X^TWX)^{-1}X^TW^{1/2}\). When we make a weighted regression, \(\hat{\beta}\) determines the fitted values in both weighted and unweighted coordinate systems. R returns the fitted values \(\hat{Y} = X\hat{\beta}\) and residuals \(e = (Y - X\hat{\beta})\) in the unweighted coordinates rather than \(\hat{Y}'\) and \(e'\). However, the standardised and studentised residuals are the same in the two coordinate systems. We plot the studentised residuals \(r'(e)\) against the fitted values \(\hat{y}\) since they have the same diagnostic properties (independence and unit variance) as plotting \(r'(e')\) against \(\hat{y}'\)

![Figure 11. LEFT: studentised residuals against fitted values and qqplot for the unweighted fit. RIGHT the fit weighted by sales. Weighting by sales improves the variance of the terraced and semi-detached properties.](image-url)
transformations modeling actually leads us to use. The idea is to find a \( \lambda \) value that linearises the data.

**Exercise** Let \( g(z; \lambda) = (z^\lambda - 1)/\lambda \) for \( z \geq 0 \). Show that \( g(z) = 1 - z^{-1}, 2z^{1/2} - 2, z - 1, (z^2 - 1)/2 \) and \( g(z) \to \log(z) \) for \( \lambda = -1, 1/2, 1, 2 \) and as \( \lambda \to 0 \) respectively. Notice that, if \( X_1 = 1 \) so \( \beta_1 \) is the intercept, then the transform \( y_k' = y_k^\lambda \) (as opposed to \( y_k' = (y_k^\lambda - 1)/\lambda \)) leads to MLE’s for the parameters of \( y' = X\beta + \epsilon \) which differ from those above by a scale and shift, \( \beta'_1 = \beta_1\lambda + 1, \beta'_2 = \lambda\beta_2, i = 2, \ldots, p \).

If we fit \( y' = X\beta + \epsilon \) for \( \epsilon \sim N(0, \sigma^2) \), with \( x_k \) the \( k \)th row \( X \), then the likelihood is

\[
L(\beta, \sigma^2, \lambda; y') = (2\pi\sigma^2)^{-n/2} \exp \left( -\frac{1}{2\sigma^2} \sum_k (y_k' - x_k'\beta)^2 \right).
\]

To get the likelihood for \( \lambda \) in terms of \( y \), we make the change of variables \( y_k' = g(y_k; \lambda) \) with Jacobian \( g_k(z; \lambda) = z^\lambda - 1 \).

**Exercise** Make the change of variables, and calculate the log-likelihood, \( \ell(\beta, \sigma^2, \lambda; y) \) in terms of \( y \). Show that the MLE’s for \( \beta \) and \( \sigma^2 \) are \( \hat{\beta} = (X^T X)^{-1}X^T y' \) and \( \hat{\sigma}_\text{MLE}^2 = (y' - H y')^T (y' - H y')/n \), with \( H = X(X^T X)^{-1}X^T \) and \( y' = y'(y, \lambda) \) (ie all as usual, but the ‘data’ \( y' \) depends on \( \lambda \)). Substitute these into the likelihood, and show that the MLE for \( \lambda \) is the argument maximising

\[
\ell(\hat{\beta}, \hat{\sigma}^2, \lambda; y) = -\frac{n}{2} \log(\hat{\sigma}^2_{\text{MLE}}(y; \lambda)) + (\lambda - 1) \log((y_1, y_2, \ldots, y_n)).
\]

In order to maximise \( \ell(\hat{\beta}, \hat{\sigma}^2_{\text{MLE}}, \lambda; y) \) as a function over \( \lambda \), we simply evaluate it at a range of values. In the 4th problem sheet you will compute a confidence interval for \( \lambda \) at level \( \alpha \).

Having computed \( \hat{\lambda} \) and a confidence interval for \( \lambda \) we usually fix on the nearest readily interpreted \( \lambda \)-value in the interval. We then recompute the fit (for \( \hat{\beta} \) and \( s^2 \) etc) conditioned on this estimate.

**Example 3.4**. The putting data records the fraction of successful putts as a function of distance in feet. Gelman and Nolan (2001) model these data. See http://www.stat.berkeley.edu/users/nolan/Papers/golfnew.ps.

We often transform proportion data as \( \log(p/(1 - p)) \) since this (the log-odds) is the link function for a Bernoulli rv. It is a monotone map from \( (0, 1) \) to \( (-\infty, \infty) \). In this case the odds of failure \( (1 - p)/p \) is the natural object (it increases with distance). A log turns out to be the wrong transform for linearity. Box-Cox gives a linear response.

```r
> putts<-data.frame(2:20,
+ c(0.93,0.83,0.74,0.59,0.55,0.53,0.46,0.32,0.34,0.32,
+ 0.26,0.24,0.31,0.17,0.13,0.16,0.17,0.14,0.16)
+ )
> names(putts)<-c('Dist','Prop')
> head(putts)
Dist  Prop
1     2  0.93
2  3.0  0.83
3  4.0  0.74
```
The data are plotted in the top left panel of Figure 12. The $\lambda$ value was estimated by maximising the likelihood, as above.

```r
> y <- (1 - putts$Prop)/putts$Prop
> x <- putts$Dist
> putts.bc <- boxcox(y ~ x)
```

This generates the graph in Figure 13, and from the graph we see that the MLE is at around 0.46 but the CI covers $\lambda = 0.5$, which is easier to deal with in later modeling. We transform the data. If distance is $x = \text{Dist}$ and the response is $y = (1 - p)/p$ for $p = \text{Prop}$ then we want

$$\sqrt{y} = \beta_1 + \beta_2 x + \epsilon$$

(2$\sqrt{y} - 2 = ...$ is not materially different).

```r
> putts.lm <- lm(sqrt(y) ~ x)
> summary(putts.lm)
```

...  

**Coefficients:**

| Estimate | Std. Error | t value | Pr(>|t|) |
|----------|------------|---------|----------|
| (Intercept) | 0.14342 | 0.09818 | 1.461 | 0.162 |
The resulting fit is plotted in the lower left panel of Figure 12. The lower right panel shows the log-transformed response, which is clearly not suitable. Notice that $\beta_1$ (the (Intercept) row in the summary() output) is not significant. Enforcing $\beta_1 = 0$ is natural on physical grounds also, as the odds of failure should go to zero for very short puts. We conclude that the odds of putt-failure increase as the square of the distance.
4. Generalised Linear Models

We have tried just about every possible transformation of the response and explanatory variables in order to get a normal linear model. However, we encounter data for which the response is binary valued, categorical, or in some other way incorrigibly non-normal. How should we set up a model, estimate parameters and confidence intervals, and carry out variable selection? The GLM framework generalises normal linear models in a natural way. Inference for GLM’s has some familiar stages.

4.1. Exponential Families. A probability mass function for a scalar rv $y$ is an natural linear exponential family of order one if it can be written

$$f(y|\theta) = \exp(y\theta - \kappa(\theta) + c(y))$$

for $y \in \Omega$ the support of $f$. In this pmf, $\kappa$ is a normalising constant, since

$$\sum_{y \in \Omega} f(y|\theta)$$

The parameter space, $\theta \in \Xi$ is the set

$$\Xi = \{\theta; \kappa(\theta) < \infty\}$$

of values which make $f$ a probability. The same definition applies for a probability density, with

$$\exp(\kappa(\theta)) = \int_{y \in \Omega} \exp(y\theta + c(y)) \, dy.$$

The function $\kappa(\theta)$ is the cumulant generating function for a probability density proportional to $\exp(c(y))$.

For the record, the exponential family of order $q > 1$ has the form

$$f(y|\theta) = \exp(g(y)^T \theta(w) - \kappa(w) + c(y)),$$

for $y \in \Omega$ possibly a vector, $g: \Omega \rightarrow \mathbb{R}^q$ a vector of $q$ linearly independent functions, and $\theta(w)$ a vector of $q$ functions of the basic parameter vector $w$ of dimension less than or equal $q$. The random vector $g(y)$ is called the natural observation for the family. The parameterisation $\theta(w)$ might disallow $\theta$ that give finite $\kappa$. If the parameterisation is simply $\exp(g(y)^T \theta - \kappa(\theta) + c(y))$, for all $\theta$ that make this function a pmf (or pdf) then we have a natural exponential family. If the parameter space $\Xi$ is open, it is regular.

There are many useful general results for inference under exponential family models. Most of the models you know fit within this framework.

Example 4.1. In the Binomial distribution Binomial$(m, \pi)$ the pmf $\Pr(Y = y)$ is

$$C^m_y \pi^y (1-\pi)^{m-y} = \exp(y \log(\pi/(1-\pi)) + m \log(1-\pi) + \log(C^m_y)).$$

This is a natural exponential family of order one, with natural parameter the log odds $\theta = \log(\pi/(1-\pi))$, natural observation $y \in \Omega = \{0, 1, 2, \ldots\}$, $\kappa = -\log(1-\pi)$ and $c(y) = \log(C^m_y)$.

Example 4.2. In the Normal distribution $N(\mu, \sigma^2)$ the pdf $f(y|\mu, \sigma^2)$ is

$$(2\pi\sigma^2)^{-1/2} \exp(-(y - \mu)^2/2\sigma^2) = \exp(y\mu/\sigma^2 - \mu^2/2\sigma^2 - (1/2) \log(2\pi\sigma^2) - y^2/2\sigma^2).$$

If $\sigma^2$ is known (so, it is not a parameter indexing members of the family) then this is a natural exponential family of order one, with natural parameter the $\theta = \mu/\sigma^2$,
natural observation \( y \in R \), \( \kappa = \mu^2/2\sigma^2 - (1/2) \log(2\pi\sigma^2) \) and \( c(y) = -y^2/2\sigma^2 \). If \( \sigma^2 \) is unknown we have a natural exponential family of order two, with \( g(y)^T \theta = (y, -y^2/2)(\mu/\sigma^2, 1/\sigma^2)^T \) which is natural since \( \theta \in R \times (0, \infty) \).

4.2. GLM, setup. A normal linear model has three parts: a deterministic part \( \eta = X\beta \), a stochastic part \( Y \sim N(\mu, \sigma^2) \), and a link between the stochastic and deterministic parts, \( \mu = \eta \). This kind of language is overkill for such a simple model.

The GLM has a stochastic part, \( Y_i \sim f \), for \( i = 1, 2, ..., n \) which we insist has the form

\[
(f(y_i|\theta_i, \phi) = \exp \left( \frac{y_i\theta_i - \kappa(\theta_i)}{\phi} + c(y_i; \phi) \right).
\]

If the scale parameter \( \phi \) is known, this is a natural exponential family (with natural parameter \( \theta/\phi \)).

It has a deterministic part \( \eta_i = X_i^T\beta \) (as before \( x_i \) is the \( p \times 1 \) vector of explanatory variables for the \( i \)th response and \( \beta \) is a \( p \times 1 \) parameter vector). The linear function \( \eta_i(\beta) \) is called the linear predictor. If \( \eta = (\eta_1, ..., \eta_n)^T \) and \( X \) is a \( n \times p \) matrix with rows \( X_i^T \), then \( \eta = X\beta \).

A GLM has a third part, which links the stochastic and deterministic parts. If \( \mu_i = E(Y_i) \), then \( g(\mu_i) = \eta_i \). Here \( g \), the link function, is a smooth invertible function of the mean.

When we find a GLM to modl given data, we make these three modeling choices. The purpose of the link function is to map the linear predictor into the scale of the response. For example, if \( Y_i \) is a binary rv, then \( E(Y_i) \) is in fact a probability, so we consider invertible (i.e. monotone) link functions \( g \) that map \( R \to (0, 1) \).

4.2.1. Moments. The mean \( \mu_i \) is a function of \( \theta_i \). Since

\[
\int (y/\phi)^n \exp(y\theta_i/\phi + c(y; \phi)) \, dy = \frac{d^n}{d\theta_i^n} \exp(\kappa(\theta_i)/\phi)
\]

we have

\[
E(Y_i) = \exp(-\kappa(\theta_i)/\phi) \int y \exp(y\theta_i + c(y; \phi)) \, dy
\]

\[
= \exp(-\kappa(\theta_i)/\phi) \frac{\kappa'(\theta_i)}{\phi} \exp(\kappa(\theta_i)/\phi)
\]

\[
= \kappa'(\theta_i),
\]

so \( \mu_i = \kappa'(\theta_i) \). For the variance,

\[
\text{var}(Y) = E(Y_i^2) - E(Y_i)^2
\]

\[
= \exp(-\kappa(\theta_i)/\phi) \phi^2 \frac{d^2}{d\theta_i^2} \exp(\kappa(\theta_i)/\phi) - (\kappa')^2
\]

\[
= \exp(-\kappa(\theta_i)/\phi) \phi^2 \frac{d}{d\theta_i} (\kappa' \exp(\kappa(\theta_i)/\phi) - (\kappa')^2
\]

\[
= \phi \kappa''(\theta_i),
\]

and so \( \text{var}(Y) = \phi \kappa''(\theta_i) \). Since \( \text{var}(Y) > 0 \), we have \( \kappa''(\theta_i) > 0 \), so \( d\mu_i/d\theta_i > 0 \) and \( \mu_i \) is a strictly increasing function of \( \theta_i \).

We have seen that the mean of \( Y_i \) increases with \( \theta_i \). Since both \( \mu_i = \kappa'(\theta_i; \phi) \) and \( g(\mu_i) = \eta_i \) are monotone functions, we must have an invertible relation \( \theta_i = \theta(\eta_i) \) between the original parameter \( \theta_i \) and the linear predictor.
We define a variance function \( V(\mu_i) \) relating the variance \( \text{var}(Y_i) \) and the mean, \( \mu_i \) as
\[
\text{var}(Y_i) = \phi V(\mu_i).
\]

**Exercise** Show that \( \text{var}(Y_i) = \phi \kappa''(\theta_i) \) and hence \( V(\mu_i) = \kappa''(\kappa^{-1}(\mu_i)) \).

**Exercise** Show that \( d\theta_k/d\mu_k = 1/V(\mu_k) \).

One particularly simple possibility is that \( \theta_i = \eta_i \). This arises when \( \kappa^{-1}(g^{-1}(\eta_i); \phi) = \eta_i \), that is, if \( g^{-1}(x) = \kappa'(x) \), corresponding to the canonical choice of link function
\[
g(\mu_i) = \kappa^{-1}(\mu_i).
\]

### 4.3. Inference for GLM’s.

#### 4.3.1. Likelihood.

One of the key ideas of modeling with GLM’s is that we start with one parameter \( \theta_i \) for each observation \( y_i \), \( i = 1, 2, ..., n \). We have knitted them all together by modeling \( \theta_i = \theta(x_i, \beta) \) in terms of the set of parameters, \( \beta = (\beta_1, ..., \beta_p)^T \). There are now just \( p \) parameters (with \( p \ll n \) often, as in Section 2) in the GLM. Since \( E(Y_i) = g^{-1}(x_i, \beta) \) is a monotone function of \( x_i, \beta \), we are back to the situation where we have \( p \) explanatory variables \( x_i = (x_{i1}, ..., x_{ip})^T \) for the \( i \)th response and, for positive \( \beta_j \), the mean response \( Y_i \) increases as \( x_{i1} \) goes up. Significant explanatory variables generate significant shift in the mean response.

Since the rv \( Y_i \) are iid, the log-likelihood for \( \beta \) is
\[
\ell(\beta) = \sum^n_{i=1} y_i \frac{\theta_i}{\phi} + c(y; \phi),
\]
with \( \theta_i = \theta_i(\beta) \). The MLE for \( \beta \) is the solution to the \( p \) equations \( df/d\beta_j = 0 \). There is no general convenient closed form for the MLE. However, it may be computed numerically, via the iteratively re-weighted least squares algorithm.

#### 4.3.2. Iteratively Re-weighted Least Squares.

Suppose we want to find \( z^* \in R \) such that \( f(z^*) = 0 \) for some continuously differentiable function \( f : R \rightarrow R \). The Newton Raphson algorithm is an iteration \( z_0, z_1, ... \) which, under certain conditions, converges to \( z^* \). We start with \( z = z^{(0)} \) and approximate \( f(z) \simeq f(z^{(0)}) + (df/dz|_{z^{(0)}})(z - z^{(0)}) \). We seek \( z = z^{(1)} \) so that \( f(z^{(1)}) = 0 \). The local linear approximation for \( z^{(1)} \) is \( z^{(1)} = z^{(0)} - (df/dz|_{z^{(0)}})^{-1}f(z^{(0)}) \), and we can iterate this to improve the approximation,
\[
z^{(i+1)} = z^{(i)} - (df/dz|_{z^{(i)}})^{-1}f(z^{(i)}), \quad i = 0, 1, 2, ...
\]
In order to solve \( df/dz = 0 \) numerically, we replace \( f \) by \( df/dz \). The N.R. iteration becomes
\[
z^{(i+1)} = z^{(i)} - (d^2 f/dz^2|_{z^{(i)}})^{-1}df/dz(z^{(i)}), \quad i = 0, 1, 2, ...
\]
This sequence may fail to converge. If \( f \) is convex then the sequence converges for some \( z_0 \) sufficiently close to \( z^* \). The multivariate case \( z = (z_1, z_2, ..., z_p)^T \) is given in terms of the Hessian of \( f : R^p \rightarrow R \). Let \( \partial f/\partial z^T = (\partial f/\partial z_1, \partial f/\partial z_2, ..., \partial f/\partial z_p) \).
be the row vector of derivatives. Let
\[
\frac{\partial^2 f}{\partial z \partial z^T} = \begin{pmatrix}
\frac{\partial^2 f}{\partial z_1 \partial z_1} & \frac{\partial^2 f}{\partial z_1 \partial z_2} & \cdots & \frac{\partial^2 f}{\partial z_1 \partial z_p} \\
\frac{\partial^2 f}{\partial z_2 \partial z_1} & \frac{\partial^2 f}{\partial z_2 \partial z_2} & \cdots & \frac{\partial^2 f}{\partial z_2 \partial z_p} \\
\vdots & \vdots & \ddots & \vdots \\
\frac{\partial^2 f}{\partial z_p \partial z_1} & \frac{\partial^2 f}{\partial z_p \partial z_2} & \cdots & \frac{\partial^2 f}{\partial z_p \partial z_p}
\end{pmatrix}
\]
give the Hessian for \( f(z) \). In order to find an extremum we seek \( z^* \in \mathbb{R}^p \) to solve the \( p \) equations \( \partial f / \partial z = 0 \). The local linear approximation for \( \partial f / \partial z \) at \( z = z^{(0)} \) is
\[
\frac{\partial f}{\partial z} \approx \left. \frac{\partial f}{\partial z} \right|_{z^{(0)}} + \left. \frac{\partial^2 f}{\partial z \partial z^T} \right|_{z^{(0)}} (z - z^{(0)}). 
\]
Notice that if \( f \) is a quadratic in \( z \) then “\( \approx \)” is “=”. If \( \frac{\partial^2 f}{\partial z \partial z^T} \) is invertible, then our multivariate Newton Raphson iteration for an extremum of \( f \) is
\[
\beta^{(i+1)} = \beta^{(i)} - \left( \left. \frac{\partial^2 \ell}{\partial \beta \partial \beta^T} \right|_{\beta^{(i)}} \right)^{-1} \left. \frac{\partial \ell}{\partial \beta} \right|_{\beta^{(i)}}.
\]
This iteration might be applied to any twice differentiable log-likelihood. It (ie \( \beta^{(i)} \)) will converge to the MLE if started sufficiently close to the MLE (since the likelihood is quadratic in the neighborhood of the MLE). Notice that, if the log-likelihood is a quadratic function function of the parameter \( \beta \) as is the case for regression with a normal linear model (see Section 2.1) then this iteration must converge in a single step.

The quantity
\[
J(y) = -\frac{\partial^2 \ell}{\partial \beta \partial \beta^T}
\]
is the observed information, whilst
\[
I = -\mathbb{E}\left(\frac{\partial^2 \ell}{\partial \beta \partial \beta^T}\right)
\]
is the expected information. For a GLM the expected information is often more convenient to work with. It has a special role in the inference, since the difference \( \beta - \hat{\beta} \) between the true parameters vector and the MLE becomes normal,
\[
(\beta - \hat{\beta}) \overset{D}{\to} \mathcal{N}(0, I^{-1}(\beta)),
\]
asymptotically in \( n \). Estimating the variance \( I(\beta) \) using \( I(\hat{\beta}) \) or \( J(y) \) are both valid, since both converge to \( I \) (see Davison (2003) Section 4.4.2). The revised iteration is
\[
\beta^{(i+1)} = \beta^{(i)} + I^{-1}(\beta^{(i)}) \frac{\partial \ell}{\partial \beta} \bigg|_{\beta^{(i)}}.
\]
For a GLM we can simplify things a bit. In that case

\[
\frac{\partial \ell}{\partial \beta_i} = \sum_k \frac{\partial \ell}{\partial \eta_k} \frac{\partial \eta_k}{\partial \beta_i} x_{k,i}
\]

since \(g(\mu_k) = \eta_k\) and using the exercise above for \(d\theta_k/d\mu_k\). In vector notation, with \(u = \partial \ell/\partial \eta\), so \(u = (u_1, ..., u_n)^T\) with \(u_k = (y_k - \mu_k)/g'(\mu_k)\phi V(\mu_k)\),

\[
\frac{\partial \ell}{\partial \beta} = X^T u.
\]

Also,

\[
\frac{\partial^2 \ell}{\partial \beta \partial \beta^T} = \frac{\partial}{\partial \beta \partial \eta^T} X = X^T \frac{\partial^2 \ell}{\partial \eta \partial \eta^T} X,
\]

so the expected information is

\[
I = X^T W X,
\]

where \(W = -E(\partial^2 \ell/\partial \eta \partial \eta^T)\) is a diagonal matrix. We can see it is diagonal, as \(u_k\) is a function of \(\mu_k\) only, and \(\mu_k = g^{-1}(\eta_k)\) is a function of \(\eta_k\) only. The diagonal entries are

\[
W_{kk} = -E \left( \frac{\partial^2 \ell}{\partial \eta_k^2} \right) = \frac{1}{g'(\mu_k)^2 \phi V(\mu_k)}.
\]

**Exercise** show that \(E(\partial \ell/\partial \theta_k) = 0\) and use it to show that

\[
E \left( \frac{\partial^2 \ell}{\partial \eta_k^2} \right) = E \left( \frac{d \theta_k}{d \eta_k} \right)^2 \frac{\partial^2 \ell}{\partial \theta_k^2}.
\]

Now show that

\[
-E \left( \frac{\partial^2 \ell}{\partial \theta_k^2} \right) = E \left( \frac{\partial \ell}{\partial \theta_k} \right)^2,
\]

and use this result to calculate \(W_{kk}\) as above.

Now the Newton Raphson iteration is

\[
\beta^{(i+1)} = \beta^{(i)} + (X^T W^{(i)} X)^{-1} X^T u^{(i)},
\]
with $W^{(i)}$ and $u^{(i)}$ evaluated at $\beta^{(i)}$. This can be written
\[ \beta^{(i+1)} = (X^TW^{(i)}X)^{-1}X^TW^{(i)}(X\beta^{(i)} + (W^{(i)})^{-1}u^{(i)}) \]
\[ = (X^TW^{(i)}X)^{-1}X^TW^{(i)}z^{(i)} \]
where $z^{(i)} = (z_1^{(i)}, ..., z_n^{(i)})^T$ is the iterated ‘data vector’,
\[ z_k^{(i)} = [X\beta^{(i)}]_k + g'(\mu_k^{(i)})(y_k - \mu_k^{(i)}). \]
This is iteratively reweighted least squares. If you look back at Section 3.3.1 you will see that if $y = X\beta + \epsilon$ with $\epsilon \sim N(0, W^{-1})$ then $\hat{\beta} = (X^TWX)^{-1}X^TWy$. The IRLS algorithm for estimation of the MLE for a GLM via the sequence $\beta^{(i)} \to \hat{\beta}$ as $i \to \infty$ is:

Start with $\mu^{(0)} = y$ so $X\beta^{(0)} = \eta^{(0)} = g(\mu^{(0)}) = g(y)$, and $z^{(0)} = g(y)$ and $W^{(0)} = \text{diag}(g'(y)^2\phiV(y))^{-1}$. For $i = 0, 1, 2, ...$
(a) set $\beta^{(i+1)} = (X^TW^{(i)}X)^{-1}X^TW^{(i)}z^{(i)}$ (ie $\beta^{(i+1)}$ are the MLE parameter values in the weighted regression of $z^{(i)}$ on $X$).
(b) $\eta^{(i+1)} = X\beta^{(i+1)}$, $\mu^{(i+1)} = g^{-1}(\eta^{(i+1)})$,
\[ z^{(i+1)} = \eta^{(i+1)} + g'(\mu^{(i+1)})(y - \mu^{(i+1)}) \]
and
\[ W^{(i+1)} = \text{diag}\left(\frac{1}{g'(\mu^{(i+1)})^2\phiV(\mu^{(i+1)})}\right). \]

**Exercise** Show that if the link function is the canonical link function then $u_k = (y_k - \mu_k)/\phi$, $W_{kk} = V(\mu_k)/\phi$ and
\[ z_k = [X\beta]_k + \frac{(y_k - \mu_k)}{V(\mu_k)}. \]

**Exercise** Compute these quantities for the case of regression with a normal linear model with $\sigma^2$ known. Show that $\phi = \sigma^2$, $V(\mu) = 1$, $W = I_n/\sigma^2$ and $z = y$, and verify that IRLS converges in one step for regression with a normal linear model.

### 4.3.3. Variance of MLE estimates.
Recall that if $\hat{\beta}$ is an MLE then, asymptotically in $n$,
\[ (\beta - \hat{\beta}) \overset{D}{\to} N(0, I^{-1}), \]
with $I$ the expected information matrix, $I = -E(\partial\ell/\partial\beta\partial\beta^T)$. We have seen that, for a GLM, $I = X^TWX$. This is handy, as $(X^TWX)^{-1}$ is computed in the course of the IRLS algorithm. We have variances $\text{var}(\hat{\beta}_i) \approx (X^TWX)^{-1}$ which are good at large $n$, with $\hat{W} = W(\hat{\mu})$ computed in the IRLS.

**Exercise** Show that if the link function is the canonical link function then
\[ \hat{\beta} \overset{D}{\to} N(\beta, \phi(X^T\text{diag}(V(\mu_1), ..., V(\mu_n))X)^{-1}). \]

**Exercise** Check this reduces to something familiar for the case of regression with a normal linear model.
This gives us a test for $\beta_i = 0$, the significance of a single GLM parameter, as

$$\frac{\hat{\beta}_i}{\sqrt{\hat{I}_{ii}^{-1}}} \xrightarrow{D} N(0, 1)$$

gives us

$$\frac{\hat{\beta}_i}{\sqrt{\hat{I}_{ii}^{-1}}} \xrightarrow{D} N(0, 1)$$

at large $n$, with $\hat{I}_{ii}^{-1}$ estimated using $\hat{I} = I(\hat{\mu})$ or $\hat{I} = J(y)$. The std.dev, Z-value and p-value reported by R in the following GLM, logistic regression, are computed using the approximate standard normal distribution for $Z = \hat{\beta}_i / \sqrt{(X^T W X)_{ii}^{-1}}$ with $W$ the final weight matrix of the converged IRLS iteration.

4.4. Logistic regression.


```r
> # From Casella and Berger Statistical Inference (2002)
> # Challenger O-ring failures (fail=1, OK=0, temp in deg. F)
> ch.dat<-data.frame(
+ fail=c(1,1,1,0,0,0,0,0,0,0,0,0,1,1,0,0,0,1,0,0,0,0,0),
+ temp=c(53,57,58,63,66,67,67,67,68,69,70,70,70,70,72,73,75,76,76,76,78,79,81))
> head(ch.dat)

fail temp
1 1 53
2 1 57
3 1 58
4 1 63
5 0 66
6 0 67
```

Now if $y_i = \text{fail}[i]$ and $x_i = (x_{i,1}, x_{i,2}) = (1, \text{temp}[i])$ (so we have an intercept), then we model $y_i \sim \text{Bernoulli}(\pi_i)$ with $\pi_i$ the probability for failure a function of temperature. The observation model for $y_i$ has pmf

$$\pi_i^{y_i} (1 - \pi_i)^{1-y_i} = \exp(y_i \log(\pi_i/(1 - \pi_i)) + \log(1 - \pi_i)),$$

so $\theta_i = \log(\pi_i/(1 - \pi_i))$ is the natural parameter, $\phi = 1$ and $\kappa(\theta_i) = \log(1 + \exp(\theta_i))$. Since $E(Y_i) = \pi_i$, we have $\mu_i = \pi_i$.

The linear predictor is $\eta_i = \beta_1 + \beta_2 x_{i,2}$. We need to link this to the mean. The logistic link

$$\log(\mu_i/(1 - \mu_i)) = \eta_i$$

is natural here, since this is equivalent to

$$\pi_i = \frac{\exp(\beta_1 + \beta_2 x_{i,2})}{1 + \exp(\beta_1 + \beta_2 x_{i,2})}.$$

The probability for failure is a logistic function of temperature. As we vary $\beta_1$ and $\beta_2$ we move the threshold for failure to different temperatures, and change how sharply the onset occurs. We can thereby choose $\beta_1$ and $\beta_2$ to fit the binary responses (see Figure 14 below). This is logistic regression, modeling the probability for success for a binary response as logistic function of a linear predictor.
Since \( \mu_i = \kappa'(\theta_i) \) and \( \kappa'(\theta_i) = e^{\theta_i}/(1 + e^{\theta_i}) \), we have

\[
\kappa^{-1}(\mu_i) = \log(\mu_i/(1 - \mu_i)),
\]

so the link function \( g(z) = \log(z/(1 - z)) \) that we chose when we set up the logistic regression was in fact the canonical link. The question “is temperature explanatory for O-ring failure” is answered by a test for \( \beta_2 \neq 0 \) significant (in fact the conjecture is that failure probability increases with decreasing temperature, so the test will check for \( \beta_2 < 0 \) significant).

**Example 4.4. Fitting a logistic model.** We now fit the logistic regression model we proposed in to the Challenger data. R uses IRLS to compute \( \hat{\beta} \) and reports the number of (score) iterations it took to get convergence.

The log-likelihood for \( \beta \) in the example above, with our choice of the canonical link, is

\[
\ell(\beta) = \sum_{i=1}^{n} y_i \eta_i - \log(1 + e^{\eta_i})
\]

with \( n = 23 \) and \( \eta_i = \beta_1 + \beta_2 x_{i,2} \). Fitting,

```r
> ch.glm<-glm(fail~temp, data=ch.dat, family=binomial())
> summary(ch.glm)
```

Deviance Residuals:

```text
Min 1Q Median 3Q Max
-1.0611 -0.7613 -0.3783 0.4524 2.2175
```

Coefficients:

```text
Estimate Std. Error z value Pr(>|z|)
(Intercept) 15.0429 7.3786 2.039 0.0415 *
temp -0.2322 0.1082 -2.145 0.0320 *
---
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

(Dispersion parameter for binomial family taken to be 1)

Null deviance: 28.267 on 22 degrees of freedom
Residual deviance: 20.315 on 21 degrees of freedom
AIC: 24.315

Number of Fisher Scoring iterations: 5

we see that the probability for failure decreases with increasing temperature. The number of scoring iterations is the number of iterations of of the IRLS algorithm we described above. We will define residual deviance shortly.

The values of the probability for failure at the observation points \( x_i \) are in effect the “fitted values”, \( \hat{\mu}_i = g^{-1}(x_i, \hat{\beta}) \), rather like \( \hat{y}_i \) in our normal linear model. Since
\hat{\pi}(\eta) = \pi(\hat{\eta})$, we can evaluate the function

$$\hat{\pi}(t) = \frac{\exp(\hat{\beta}_1 + \hat{\beta}_2 t)}{1 + \exp(\hat{\beta}_1 + \hat{\beta}_2 t)}.$$ 

to get the estimated probability for failure as a function of temperature. The `predict()` function takes the `glm()` output, and some x values and returns $x^T \hat{\beta}$. With the option `type='response'` we get $g^{-1}(x^T \hat{\beta})$, or in this case $\hat{\pi}(x, \beta)$. We can evaluate this function at other x-values using the `newdata` option. We take a sequence of values of the explanatory variable `temp`, and plot $\pi$ against temperature in Figure 14.

```r
> ch.prob <- predict(ch.glm, newdata=data.frame(temp=40:90), type='response')
> plot(40:90, ch.prob, type='l'); #the solid line
```

and I omit the R for the plotting of the points, see L10.R.

![Figure 14. O-ring failure data (points) with fitted probability for failure (curve).](image)

The $p$-value for $\beta_2 = 0$ is 0.032, supporting the view that temperature is explanatory for O-ring failure. How is this computed? The reported standard errors are $\text{var}(\hat{\beta}_i) \approx (X^T \hat{W} X)^{-1}_{ii}$ for $i = 1, 2, ..., p$. The quoted $Z$ value is $\hat{\beta}_i / \sqrt{(X^T \hat{W} X)^{-1}_{ii}}$ which has an approximate $N(0, 1)$ distribution. The $p$-value for the test for $\beta_2 < 0$ is 0.016.
Exercise Verify that \( V(\mu_i) = \kappa''(\bar{\theta}_i)/\phi = \mu_i(1 - \mu_i) \), and hence

\[
W = \text{diag}(\hat{\mu}_1(1 - \hat{\mu}_1), ..., \hat{\mu}_n(1 - \hat{\mu}_n))
\]
is the last weight matrix of the IRLS. Check the quoted std errs. Ans:

> # check we understand what the reported std errs are
> p<-predict(ch.glm,type='response')
> X<-model.matrix(fail~temp, data=ch.dat)
> sqrt(diag(solve(t(X)%*%diag(p*(1-p))%*%X)))

7.378636 0.1082365

Example 4.5. Odds and the odds ratio. The quantity \( \pi/(1 - \pi) \) is the odds of success (O-ring failure, but Bernoulli success, i.e. \( Y = 1 \)). For a single generic binary response \( Y \) with explanatory variables \( x \) with \( \mu = \mathbb{E}(Y) \) and \( \pi = \Pr(Y = 1) = \mu \), the canonical link \( g(\mu) = \eta \) for logistic regression gives \( \log(\pi/(1 - \pi)) = x^T \beta \), and so the odds of success, \( \hat{O} = \pi/(1 - \pi) \), say, are estimated as \( \hat{O} = O(x^T \hat{\beta}) \), so we have \( \hat{O} = \exp(x^T \hat{\beta}) \). This gives the interpretation of \( \hat{\beta}_i \) in logistic regression: \( \hat{\beta}_i \) gives the change in log odds for success (the outcome \( Y = 1 \)) when the corresponding explanatory variable changes \( x_i \rightarrow x_i + 1 \). For example, for the O-ring data, if \( x = (x_1, x_2)^T \) and \( x' = (x_1, x_2 + 1) \), so the temperature goes up a degree, and \( \hat{O} = O(x^T \hat{\beta}) \) and \( \hat{O}' = O(x'^T \hat{\beta}) \), then

\[
\frac{\hat{O}'}{\hat{O}} = \exp(\hat{\beta}_2).
\]

In this example we estimated \( \hat{\beta}_2 \approx -0.2321627 \) so the odds of O-ring failure go down by around \( \exp(-0.2321627) \approx 0.7928171 \) for an increase in the temperature by one deg. F.

On the day of the disaster it was unusually cold, 31 deg. F. What is our predicted probability for O-ring failure that day?

> #launch at temp=31 deg F - estimate for the probability of failure:
> predict(ch.glm,newdata=data.frame(temp=31),type='response')

[1] 0.9996088

> 

The estimated odds of O-ring failure are 1:1 at a temperature around 64.79465 deg F (since \( \hat{O} = 1 \) when \( \hat{\beta}_1 + \hat{\beta}_2 x_2 = 0 \) and solve for \( x_2 \)). How much higher are the odds of failure at 31 deg F? This must be \( 0.9996088/(1 - 0.9996088) \approx 2555 \), or \( \exp(\hat{\beta}_2 \Delta \text{temp}) \) which is 0.7928171\(^{64.79465}\).

Example 4.6. Binomial data in a 2 way table The data in Table 3 are taken from Dr Lunn’s 2007 lecture notes, and give the number of men and women smokers and non-smokers in a particular industry. Table 3 gives smoking information for 2916 men and 2503 women. In a binomial model with \( p \) the probability for a man to be a smoker and \( q \) the probability for a woman to be a smoker, the MLE’s for \( p \) and \( q \) are \( \hat{p} = n_{11}/(n_{11} + n_{12}) \approx 0.71 \) and \( \hat{q} = n_{21}/(n_{21} + n_{22}) \approx 0.45 \). How much higher are the odds for a man to be a smoker (smoker given man) than for a woman to be a smoker (smoker given woman)? The MLE for the odds ratio is a function of the MLEs for \( p \) and \( q \), i.e \( \hat{p}/(1 - \hat{p})]/[\hat{q}/(1 - \hat{q})] = (n_{11}/n_{12})/(n_{21}/n_{22}) \approx 2.92 \).

We will use these data to illustrate the Binomial GLM. We have two observations, \( y_1 = 2059 \) ‘successes’ from \( m_1 = 2916 \) trials and \( y_2 = 1130 \) successes from \( m_2 = \)
Smoking
Sex
<table>
<thead>
<tr>
<th>Y</th>
<th>N</th>
<th>Tot</th>
</tr>
</thead>
<tbody>
<tr>
<td>M</td>
<td>2059</td>
<td>857</td>
</tr>
<tr>
<td>F</td>
<td>1130</td>
<td>1373</td>
</tr>
<tr>
<td>Tot</td>
<td>3189</td>
<td>2230</td>
</tr>
</tbody>
</table>

Table 3. Two way table for smoking and gender, giving the number of subjects in each category.

2503 trials. Let $\pi_i$ give probability for probability for ‘success’ (here, smoker) for the $i$th group of trials (so $\pi_1 = p$ and $\pi_2 = q$). The likelihood for a single observation is

$$f(y_i|m_i) = \exp(y_i \log(\pi_i/(1 - \pi_i)) + m_i \log(1 - \pi_i) + \log(C_{y_i}^{m_i}))$$

The natural observation is $y_i$, the number of successes in the $i$th group of trials, the natural parameter is again $\theta_i = \log(\pi_i/(1 - \pi_i))$. $\kappa(\theta_i) = m_i \log(1 + e^{\theta_i})$, and $\phi = 1$.

We are interested in the explanatory variable ‘Sex’. Let $x_{1,2} = 1$ and $x_{2,2} = 0$, so $x_2$ is the dummy indicator variable for the M level of the categorical variable Sex. Level F is the baseline. The linear predictor is $\eta_k = \beta_1 + \beta_2 x_{k,2}$ and the design matrix is

$$X = \begin{pmatrix} 1 & 1 & 1 \\ 1 & 0 & 0 \end{pmatrix}.$$ 

As for the binary data, model the probability as a logistic function of the indicator variable. Here $\pi_k = \mu_k/m_k$, so

$$\log(\mu_k/(m_k - \mu_k)) = \beta_1 + \beta_2 x_{k,2}.$$ 

Since $\theta_k = \eta_k$, this is again the canonical link function, with

$$\pi_1 = \exp(\beta_1 + \beta_2)/(1 + \exp(\beta_1 + \beta_2))$$

$$\pi_2 = \exp(\beta_1)/(1 + \exp(\beta_1 + \beta_2)).$$

We can fit these data in R using glm() again.

```r
> # smoking data from Dunn 07
> smk<-data.frame(S=c(2059,857),NS=c(1130,1373),Sex=c('M','F'))
> # response is two columns '#successes' S and '#failures' NS
> smk
   S NS Sex
1 2059 1130 M
2  857 1373 F
> #The design matrix has an intercept and an indicator for Sex==M
> model.matrix(cbind(smk$S,smk$NS)~Sex,data=smk)
   (Intercept) SexM
1         1      1
2         1      0
> #fit response as binomial with Sex explanatory
> smk.glm<-glm(cbind(smk$S,smk$NS)~Sex,data=smk,family=binomial)
```
The estimated change in the log odds when we move from \( F \) to \( M \) (so \( x_{k,2} \to x_{k,2}+1 \)) is \( \beta_2 = 1.07 \) which is an increase by a factor of \( \exp(1.07132) \approx 2.92 \).

4.5. Model choice. We have seen how to test for the significance of a single GLM parameter \( \beta_i \), using the asymptotically standard normal distribution of \( Z = \hat{\beta}_i / \sqrt{(X^TWX)^{-1}} \).

How do we test for the significance of a group of variables? We use the likelihood ratio test. For regression with the normal linear model we managed to get an exact test, by writing the LRT statistic as a monotone function of a statistic, \( F \), with a distribution we have exactly. For regression with GLM’s, we stick to the LRT statistic, and accept a test based on the asymptotic distribution of this statistic.

There are two important models ‘above’ and ‘below’ our model. The saturated model unlinks the \( \theta_i \). They were bound by the constraint \( g(\mu_i) = x_i^T \beta \) on the means. In the saturated model we have one parameter \( \theta_i \) for each response \( y_i \). The MLE \( \hat{\theta}_i^{(s)} \) for \( \theta_i \) in the saturated model is just \( \hat{\theta}_i^{(s)} = \arg \max_{\theta} f(y_i|\theta) \).

Exercise Suppose \( Y_i \sim \text{Binomial}(\pi_i, m_i) \). Show that \( \hat{\pi}_i^{(s)} = y_i/m_i \) so that \( \hat{\theta}_i^{(s)} = \log(y_i/(m_i - y_i)) \) and the log-likelihood for the saturated model is

\[
\ell(\hat{\theta}_i^{(s)}; y) = \sum_i y_i \log(y_i/(m_i - y_i)) + m_i \log(1 - y_i/m_i).
\]

The other model of interest is the null model, in which \( \beta_2 = \ldots = \beta_p = 0 \), with \( g(\mu_i) = \beta_1 \) so we just have the intercept in the linear predictor. The means are all equal so there is a single common natural parameter \( \theta \). Let \( \hat{\theta}^{(0)} \) be the MLE for the natural parameter of the null model.

Exercise Show that, for the binomial model, \( \hat{\pi}^{(0)} = (\sum y_i)/(\sum m_i) = \bar{y}/\bar{m} \) so that \( \hat{\theta}^{(0)} = \log(\bar{y}/(\bar{m} - \bar{y})) \) and the log-likelihood for the null model is

\[
\ell(\hat{\theta}^{(0)}; y) = n \bar{y} \log(\bar{y}/(\bar{m} - \bar{y})) + n \bar{m} \log(1 - \bar{y}/\bar{m}).
\]

4.5.1. Deviance. The scaled deviance \( D(y) \) for our GLM is simply related to the log-likelihood,

\[
D(y) = -2\ell(\hat{\beta}; y) + 2\ell(\hat{\theta}^{(s)}; y).
\]

This is of course the LRT statistic for a test comparing the saturated model with the GLM of interest. The deviance itself is the scaled deviance at scale parameter \( \phi = 1 \). Since the parameter space of \( \hat{\theta}^{(s)} \) includes the parameter space of \( \theta(\beta) \) as a subspace, \( D(y) \geq 0 \). The null deviance is

\[
D^{(0)} = -2\ell(\hat{\theta}^{(0)}; y) + 2\ell(\hat{\theta}^{(s)}; y).
\]
4.5.2. Goodness of fit. Since $D$ is the LRT statistic for a test with null parameter space of dimension $p$ and alternative of dimension $n$, we expect $D(Y) \sim \chi^2(n-p)$ approximately, under the hypothesis that our GLM model includes all the factors generating variation in the response. If $D(y)$ is large on the scale of a $\chi^2(n-p)$ rv, then we question our model.

This model check is not always applicable. The problem is that $D(Y) \sim \chi^2(n-p)$ holds asymptotically, yes, but not asymptotically in $n$. We achieve the asymptotic distribution for a LRT statistic as our MLE’s converge to their limiting values. Since, in the saturated model, we have one parameter for each observation, increasing $n$ doesn’t add to the precision of our estimates of these parameters (increasing $n$ does increase the precision of our estimate of $\beta$, but that is just half the LRT statistic).

So, when is this distributional assumption good? Suppose we had multiple replicates for each observation $y_{ij}$, $j = 1, \ldots, m_i$, for given explanatory variables $x_i$, and, in the saturated model, just the one parameter $\theta_i^{(s)}$ for each batch of $m_i$ replicates. Now $\hat{\theta}_i^{(s)} \rightarrow \theta_i^{(s)}$ as $m_i \rightarrow \infty$. This is the limit where the approximation $D(Y) \sim \chi^2(n-p)$ holds good. So, where do we have replicates?

Exercise Show that a binomial response $Y_i \sim \text{Binomial}(m_i, \pi_i)$ is $m_i$ replicates of a Bernoulli response, and hence that $\hat{\theta}_i^{(s)}$ converges to $\log(\pi_i^{(s)}/(1 - \pi_i^{(s)}))$ and $D(Y) \overset{D}{\rightarrow} \chi^2(n-p)$ as $m_i \rightarrow \infty$ for each $i = 1, 2, \ldots, n$.

Bernoulli data has none of this structure. In this case $D(Y) \sim \chi^2(n-p)$ will not hold and any inference that depends on this approximation (residual analysis below) is not useful.

4.5.3. Model choice. A test comparing two nested models $(Q)$ with dimension $q$ and $(P)$ with dimension $p < q$ has LRT statistic $\Lambda = D^{(P)}(y) - D^{(Q)}(y)$, so

$$D^{(P)}(y) - D^{(Q)}(y) \sim \chi^2(q-p).$$

The test for no relation between the mean response $\mu_i$ and the explanatory variables $\eta_i = x_i \beta$ in the model $(Q)$ is the test for $\beta_2 = \ldots = \beta_p = 0$ (if $\beta_1$ is the intercept) has test statistic

$$D^{(0)}(y) - D^{(Q)}(y) \sim \chi^2(q-1).$$

All the remarks we made about model choice for a LM, in Section 3.2 apply equally to GLMs. The problem has a similar structure. We want to check that we have all the important variables in our model (using a Goodness of fit test, as above), and then check we have no redundant variables. The tools available which help us organise the search for a set of explanatory variables are like those for LM’s. We can make forwards addition and backwards elimination, one variable at a time. We can identify a reduced model by these or physical considerations, and test to drop the corresponding complementary group of variables. We can make automatic model choice using the AIC.

Example 4.7. In the following example, quoted from Venables and Ripley (2002), based on data from Collet (1991, see V&R for reference), 12 batches of 20 tobacco budworm moths were exposed for 3 days to different dose levels of a toxin. The numbers dead or disabled were recorded. The data are displayed in Table 4. We have a categorical variable $\text{sex} \in \{M, F\}$ and a real ordinal variable $\text{dose}$, which we

will code as \( \text{ldose} = \log_2(\text{dose}) \), as suggested by the scale of the variable. We look for an effect due to \( \text{ldose} \) (fairly obvious!) and allow for possibly different intercepts and slopes in the linear predictors for the two sexes. Our linear predictors are, for \( i = 1, 2, ...n \) with \( n = 12 \),

\[
\eta_i = \beta_1 + \beta_2 g_{M,i} + \beta_3 x_{d,i} + \beta_4 g_{M,i} x_{d,i}
\]

with \( g_{M,i} = 1 \) if \( \text{sex}[i] = M \) and zero otherwise, and \( x_{d,i} = \text{ldose}[i] \). If \( y_i \) are the cell counts, then R will fit for the scaled response \( y_i/m_i \) (with \( m_i = 20 \) for \( i = 1, 2, ...n \) here). The expected response is then \( \mu_i = E(Y_i/m_i) \) which is \( \mu_i = \pi_i \) with \( \pi_i \) the probability of success in the \( i \)th Binomial trial, so \( y_i \sim \text{Binomial}(m_i, \pi_i) \). In the notation of Example 4.6, we have the logistic link function

\[
\log(\mu_i/(1 - \mu_i)) = \eta_i
\]

modeling the proportion \( \mu_i \) of ‘successes’, which are Moth-deaths in this case.

```r
> #Example from Venables and Ripley (2002) Sec 7.2
> ldose<-rep(0:5,2) #the log-dose
> numdead<-c(1,4,9,13,18,20,0,2,6,10,12,16)
> sex<-c(rep("M",6),rep("F",6))
>
> #gather data in a data-frame
> bw$sex is categorical with levels M,F
> bw<-data.frame(numdead,numalive=20-numdead,ldose,sex)
>
> #to use glm we have some options how to enter the response
> #here we are using the [successes,failures] format
> #logistic regression
> bw.glm<-glm(cbind(numdead,numalive)~sex*ldose,data=bw,family=binomial)
> summary(bw.glm)
```
Residual deviance: 4.9937 on 8 degrees of freedom AIC: 43.104

Number of Fisher Scoring iterations: 4

Are we missing sources of variation? The test for goodness-of-fit is \( D \sim \chi^2(n-p) \) with \( n = 12 \) and \( p = 4 \). Since \( m_i = 20 \) is acceptably large, we can expect the asymptotics for \( D \) to hold good. Comparing \( D(y) = 4.9937 \) to a \( \chi^2(8) \) random variable,

\[
> 1-\text{pchisq}(4.9937, 8)
\]

[1] 0.7582493

we get a \( p - \text{value} \) of around 0.76, so we see no evidence for misfit in this respect. Can we drop any terms from this model. The two variables \texttt{sexM} and \texttt{sexM:ldose} are obvious candidates. In order to get the deviance for the reduced model, we refit using the reduced model.

\[
> \texttt{bw.glm<-glm(cbind(numdead,numalive)~ldose,data=bw,family=binomial)}
> \texttt{summary(bw.glm)}
\]

... Coefficients:

| Estimate | Std. Error | z value | Pr(>|z|) |
|---------|------------|---------|----------|
| (Intercept) | -2.7661 | 0.3701 | -7.473 | 7.82e-14 *** |
| ldose | 1.0068 | 0.1236 | 8.147 | 3.74e-16 *** |

Residual deviance: 16.984 on 10 degrees of freedom AIC: 51.094

The change in the deviance on dropping \texttt{sex} and \texttt{sex:ldose} is 16.984 − 4.9937 = 11.9903 with \( n = 12, q = 4, p = 2 \) and since \( D' - D \sim \chi^2(q-p) \) under the null model without \texttt{sex} and \texttt{sex:ldose}, the test to drop the two variables has \( p \)-value

\[
> 1-\text{pchisq}(11.99, 2)
\]

[1] 0.002491177

strong evidence for some role for at least one of these variables.

As Venables and Ripley (2202) point out, this \( p \)-value might be a bit surprising given the apparent low significance of the individual variables. As they note, the fact that the offset \( \beta_2 \) in the intercept is not significant is not surprising. This is the difference in mean response between sexes at \( \texttt{dose} = 1 \) (so \( \texttt{ldose} = 0 \)) and there are few deaths on either side at low dose. The intercept and slope offset estimates are quite significantly (anti) correlated (inspect \( I^{-1} = \text{summary(bw.glm)}$\texttt{cov.scaled} \) for detail) so we have here the kind of significance-masking due to near-linear dependence that we have discussed for linear models. If we shift the intercept to a central dose value, say \( \texttt{dose} = 8 \) or \( \texttt{ldose} = 3 \) we reduce this correlation (without changing the meaning of the other parameters). This can be thought of as a step towards orthogonalising the original variables \( g_M \) in order to reduce correlation.

\[
> \texttt{bw.glm<-glm(cbind(numdead,numalive)^*sex*I(ldose-3),data=bw,family=binomial)}
> \texttt{summary(bw.glm)}
\]

... Coefficients:

| Estimate | Std. Error | z value | Pr(>|z|) |
|---------|------------|---------|----------|
| (Intercept) | -0.2754 | 0.2305 | -1.195 | 0.23215 |
| sexM | 1.2337 | 0.3770 | 3.273 | 0.00107 ** |
| I(ldose - 3) | 0.9060 | 0.1671 | 5.422 | 5.89e-08 *** |
We can now see that there is no evidence for different slopes for the two sexes, but the intercepts are not equal. Venables and Ripley (2002) go on to test for curvature, looking for dependence on \((\text{ldose} - 3)^2\).

Note that we can form an analysis of deviance table for a GLM fit. Like the default R ANOVA table, this looks at the nested sequence of models obtained by dropping the variables one at a time. The analysis of deviance table gives, in each row, the increase in the deviance as the model variables in the corresponding row are removed from the model. For example, here is the analysis of deviance table for the test on the original fit with intercept as \(\text{ldose} = 0\).

\[
\begin{array}{lcccc}
\text{Df} & \text{Deviance} & \text{Resid. Df} & \text{Resid. Dev} & \text{P(>|Chi|)} \\
\text{NULL} & 11 & 124.876 & & \\
\text{ldose} & 1 & 107.892 & 10 & 16.984 \times 10^{-25} \\
\text{sex} & 1 & 10.227 & 9 & 6.757 \times 10^{-1} \\
\text{ldose:sex} & 1 & 1.763 & 8 & 4.994 \times 10^{-1} \\
\end{array}
\]

The first row gives the null deviance, \(D(0)\) (called here the residual deviance). The second row gives the p-value for testing for an effect due to \(\text{ldose}\) in a model with just intercept. The column headed Deviance is what we would call the change in deviance, and the column headed Resid. Dev is what we would call the deviance. The levels of significance depend on the order in which the variables are added. We can see now that \(\text{sex}\) was significant from the first, but in the original \text{summary()} output its significance was masked by correlation with the slope offset.

4.6. Further GLM diagnostics. In order to identify outliers, we can make an analysis of residuals, just as for normal linear models. There are a number of different ways to define residuals for GLM’s.

A poorly fitting point will make a large contribution to the deviance. Let

\[
\ell_i(\theta; y_i) = \frac{y_i \theta - s(\theta_i)}{\phi} + c(y_i; \phi),
\]

and let

\[
d_i = -2\ell_i(\hat{\theta}; y_i) + 2\ell_i(\hat{\theta}_i^{(s)}; y_i),
\]

so that

\[
D(y) = \sum_{i=1}^{n} d_i.
\]

Now \(d_i \geq 0\) and data with relatively larger \(d_i\) are data in greater conflict with the model and the rest of the data. The deviance residuals are defined to be

\[
r_i = \text{sign}(y_i - \hat{\mu}_i)d_i.
\]

The function \(\text{sign}(y_i - \hat{\mu}_i)\) is +1 or −1 depending on the sign of \(y_i - \hat{\mu}_i\) and shows us whether \(y_i\) is in conflict by being relatively too large or too small compared to the fitted mean.
Another way to think about residuals is to consider the misfit in the space of the linear predictor. Since \( g \) maps \( \mu \) to \( \eta \), we might compare \( g(y) \) and \( \hat{\eta} \). Expanding \( g(y) \) about \( y = \mu \), \( g(y) \approx g(\mu) + g'(\mu)(y - \mu) \), an object which appears in our IRLS, \( z = \eta + g'(\mu)(y - \mu) \). This gives us an interpretation of \( z \) in that algorithm. The residuals \( z - \hat{\eta} = g'(\hat{\mu})(y - \hat{\mu}) \) are called the working residuals (in R, for example). They are exactly the residuals of the linear regression made in the IRLS. The Hat matrix for that regression is

\[
H = W^{1/2}X(X^TX)^{-1}X^TW^{1/2},
\]

and the leverages \( h_{ii} \) have the same role as before in the analysis of the working residuals.

The standardised deviance residuals are

\[
r'_i = \frac{d_i}{\sqrt{1 - h_{ii}}} \text{sign}(y_i - \hat{\mu}_i)
\]

It can be shown that if the original approximation \( D(y) \sim \chi^2(n - p) \) is good then the \( r' \) usually have roughly unit variance and distributions close to normal. This leads to a check for misfit.

Example 4.8. For the budworm moth analysis of Example 4.7, the standardised deviance residuals are shown in Figure 15. These are given for the final model, with equal slopes for the two sexes.

```r
> bw.glm<-glm(cbind(numdead,numalive)~sex+I(ldose-3),data=bw,family=binomial)
> summary(bw.glm)
```

... 

Deviance Residuals:

<table>
<thead>
<tr>
<th></th>
<th>Min</th>
<th>1Q</th>
<th>Median</th>
<th>3Q</th>
<th>Max</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>-1.10540</td>
<td>-0.65343</td>
<td>-0.02225</td>
<td>0.48471</td>
<td>1.42944</td>
</tr>
</tbody>
</table>

Coefficients:

|                | Estimate | Std. Error | z value | Pr(>|z|) |
|----------------|----------|------------|---------|----------|
| (Intercept)    | -0.2805  | 0.2431     | -1.154  | 0.24854  |
| sexM           | 1.1007   | 0.3558     | 3.093   | 0.00198 ** |
| I(ldose - 3)   | 1.0642   | 0.1311     | 8.119   | 4.7e-16 *** |

... 

Null deviance: 124.876 on 11 degrees of freedom
Residual deviance: 6.757 on 9 degrees of freedom
AIC: 42.867

Number of Fisher Scoring iterations: 4

```r
> plot(fitted.values(bw.glm),rstandard(bw.glm))
```

There is no evidence for misfit, as there are no large (compared to one) standardised deviance residuals. There is no obvious sign of a trend in the residuals. This is often a sign of a problem with the link function (see problem 2 of PS5 - for an example - try computing residuals for the log-link, and compare them with the residuals for the sqrt link).

4.7. Dispersion and the scale parameter. When we model using a GLM based on a one parameter family of distributions, such as Binomial and Poisson distributions, the scale parameter \( \phi \) is equal one. When we model using other distributions
(Gamma, Normal etc) \( \phi \) is not one, and may be unknown. In these cases we might allow for over-dispersion, and extend the model, allowing \( \phi \neq 1 \). We need an estimate for \( \phi \).

Notice that the IRLS algorithm doesn’t need \( \phi \)! It follows that, operationally, we can fit our GLM at \( \phi = 1 \), and then look for estimates of \( \phi \) based on the fitted values of \( \hat{\beta} \).

**Exercise** verify that \( \phi \) cancels out in each step of the IRLS squares iteration.

Since \( \text{var}(Y_i) = \phi V(\mu_i) \) we can form an estimate

\[
\hat{\phi} = \frac{1}{n - p} \sum_{i=1}^{n} \frac{(y_i - \hat{\mu}_i)^2}{V(\hat{\mu}_i)},
\]

with \( \hat{\mu}_i = g^{-1}(x_i^T \hat{\beta}) \).

When we fit a simple model with \( \phi = 1 \), or otherwise known, and we suspect over- (or who knows? under-) dispersion, we may wish to compute \( \hat{\phi} \). If this differs from one, the estimated confidence intervals for \( \hat{\beta} \), which depend on \( \phi \), as in Section 4.3.3, will be wrong, and the estimated variance should now be multiplied by \( \hat{\phi} \).
The picture cannot be so simple for Binomial and Poisson models. We can’t simply scale the \( y\theta - \kappa(\theta) \) term by \( \phi \) in these models and expect the exponential family to remain the same. The requirement that the likelihood be normalised over \( y \) (\( \int \exp (y\theta + c(y; \phi)) \, dy = \exp(\kappa(\theta_i)/\phi) \)) is no longer satisfied within the original family. If we define \( \phi = \text{var}(Y)/V(\mu) \), yielding the estimator given above, then we can get an ad-hoc adjustment for over-dispersion.

**Exercise** see related exercise on cloth fault data in PS6.

In either of these cases (\( \text{i.e. when } \phi \text{ is a parameter of the likelihood, and when it is an ad-hoc scale parameter adjusting the variance of estimates} \)), the test for model comparison based on the deviance is adjusted. If \( \phi \) is estimated then the scaled deviance is the ratio of the deviance at \( \phi = 1 \) divided by \( \hat{\phi} \). Since these quantities have distributions which are approximated by \( \chi^2 \) distributions you will often see an \( F \)-statistic used in GLM analysis to carry out this test. [End L14]
5. Monte Carlo Tests

Some properties of random variables are hard to calculate in closed form, but straightforward to estimate by simulation.

Suppose we want to make a hypothesis test by comparing the distribution of some test statistic under the null hypothesis to the value of that statistic computed from the data. It often happens that the test statistic is easy to compute from the data, but the distribution of the test statistic is hard to compute. For example, the distribution of the Likelihood Ratio Test statistic \( \Lambda(Y) \) is known only asymptotically as \( \chi^2 \). For small sample sizes this may be a poor approximation, and the exact distribution of \( \Lambda \) may be hard to compute.

If we can simulate synthetic data under the null hypothesis, then we can see how often the test statistic computed on synthetic data simulated under the null exceeds that computed on the real data. This gives us an estimate of the \( p \)-value.

The errors that arise from estimating a \( p \)-value using a finite number of simulated data sets may be small compared to the errors which might arise if we used an inexact asymptotic approximation.

5.1. Monte Carlo simulation. Before we look at Monte Carlo tests, let us briefly review simulation methods. Much of this material has been moved into the Part A simulation course. In this course we will simply use the R simulation tools to simulate readily available rv.

R has simulation tools to simulate all the well known distributions. So `runif(n, min=0, max=1)`, `rnorm(n, mean=0, sd=1)` and `rpois(n, lambda)` simulate \( n \) pseudo-random iid Uniform, Normal and Poisson distributed rv respectively.

Simulation builds up from simple to more complex distributions, by taking functions of easily simulated rv in order to simulate more complicated distributions. At the bottom of all this are the pseudo-random uniform variates, \( U \sim U(0,1) \).

The method of inversion is one of the base simulation methods. Let \( X \) be a scalar random variable with cumulative distribution function (cdf) \( F(x) = \Pr(X \leq x) \) at \( X = x \) in some space of states \( \Omega \). Let \( F^{-1}(u) \) be the smallest value of \( x \) such that \( F(x) \) is greater than or equal to \( u \). If \( F \) is continuous and strictly increasing then this is just the inverse of \( F \). Simulation by the method of inversion exploits the fact that if \( U \sim U(0,1) \) and \( X = F^{-1}(U) \) then \( X \sim F \) (meaning, the rv \( X \) is distributed with cdf \( F \)). This follows (for the simple case) by \( \Pr(X \leq x) = \Pr(F^{-1}(U) \leq x) = \Pr(U \leq F(x)) = F(x) \).

For example, if \( X \sim \text{Exp}(\lambda) \), then \( F(x) = 1 - \exp(-\lambda x) \) and if \( U \sim U(0,1) \) and \( X = -\log(U)/\lambda \) then \( X \) is exponentially distributed with mean \( 1/\lambda \). Here are 4 \( \text{Exp}(1/10) \) rv.

\[
\lambda = 1/10; \quad (Y <- -\log(runif(n=4))/\lambda)
\]

\[
[1] \quad 0.0007818083 \quad 6.7491588149 \quad 9.2983470373 \quad 6.8312430345
\]

In part A probability you learnt about some combinations of rv that generate other rv. If for \( i = 1, 2, 3, ..., a, Y_i \) are iid rv distributed \( Y_i \sim \text{Exp}(\lambda) \) and

\[
X = \sum_{i=1}^{a} Y_i
\]

then \( X \sim \text{Gamma}(a, \beta) \) (the Gamma(\( a, \beta \)) density is \( f_X(x) \propto x^{a-1}\exp(-x/\beta) \)). If \( Y \) is the vector of four rv simulated above, then here is a realisation of a Gamma(4,0.1) rv.
Deaths          0  1  2  3  4  ≥ 5  
Observed Frequency 109  65  22  3  1  0  
Expected Frequency  108.67 66.29 20.22 4.11 0.63 0.08  

Table 5. Frequency for numbers of deaths totaled over years and corps. Expected frequency is MLE for Poisson

> sum(Y)  
[1] 22.87953  
and here is another  
> rgamma(n=1,shape=4,rate=1/10)  
[1] 17.93972

We can build up realisations of rv with other familiar distributions in a similar way. Notice that in the example above we are restricted to simulating Gamma-distributions with integer shape parameter. Also, we may have a random variable with some non-standard density. In the part A simulation course we looked at methods which allow us to simulate fairly general probability distributions. In this course, we will assume a function simulating the rv we need is available, or can be constructed, by elementary operations such as those above, using readily available functions.

5.2. Monte Carlo and the Likelihood ratio test. Many tests can be Monte-Carlo’ed, and the LRT is an obvious candidate. We can replace the asymptotic $\chi^2$ approximation, which is hard to check, with a Monte-Carlo approximation.

Example 5.1. The data in Table 5 give the number of deaths by Horse-kicks in the Prussian cavalry between 1875 and 1894. There are 10 years and 20 corps. In each year and each corps the number of deaths by horse-kick is recorded. Let \( i = 1, 2, \ldots, n \), \( n = 200 \) be the year-corps (block) index, and let \( y_i \) give the number of deaths in the \( i \)th block. Table 5 is a frequency table for these data, giving the number of 0’s, 1’s etc in the data. Are the \( y_i \) Poisson with a common mean? Denote by \( O_j, j = 0, 1, 2, \ldots, k \) the observed frequency for \( j \) deaths per corps per month, so \( O_0 = 109 \) etc and \( k = 5 \). Let \( \pi_j(\mu) = \mu^j \exp(-\mu)/j! \) give the probability \( \Pr(Y_i = j) \) to see \( j \) deaths, under \( H_0 \), the proposed model (with \( \pi_5 \equiv \Pr(Y_i \geq 5) \) so \( \pi_5(\mu) = 1 - \sum_{j=0}^{4} \pi_j(\mu) \)). If the distribution is not Poisson, then it is multinomial, the largest class of distributions we can allow. Let \( \pi_j, j = 0, 1, \ldots, k \) with \( k = 5 \) give the probabilities, under this multinomial distribution \( H_1 \). The MLE for \( \mu \) under \( H_0 \) is \( \hat{\mu} = (\sum_j jO_j)/\sum O_j = 0.61 \),  
> O<-c(109,65,22,3,1,0)  
> n<-sum(O)  
> (mu<-sum((0:5)*O)/n)  
[1] 0.61

so the MLE for \( \pi_j(\mu) \) is \( \pi_j(\hat{\mu}) \). The MLE for \( \pi_j \) is \( \hat{\pi}_j = O_j/n \). The likelihood under \( H_0 \) is  
\[
L(\mu; y) = \prod_j \frac{n!}{O_j!} \prod_j \pi_j(\mu)^{O_j}
\]
so the likelihood ratio test statistic is

\[ \Lambda(y) = 2 \sum_{j=0}^{k} O_j \left( -\log(\pi_j(\hat{\mu})) + \log(\hat{\pi}_j) \right) \]

\[ = 2 \sum_{j} O_j \log(O_j/n\pi(\hat{\mu})) \]

\[ = 2 \sum_{j} O_j \log(O_j/E_j), \]

where \( E_j = n\pi_j(\hat{\mu}) \) is the expected frequency under \( H_0 \). The Pearson statistic \( P(y) \) approximates \( \Lambda \) but is more robust for small sample sizes where the logarithm in the LRT statistic can cause problems. The Pearson statistic is given by

\[ P(y) = \sum_{j} \frac{(O_j - E_j)^2}{E_j}. \]

What are the dimensions of the parameter spaces under \( H_0 \) and \( H_1 \)? Under \( H_0 \), we have one parameter, \( \mu \), so the dimension is \( p = 1 \). Under \( H_1 \) we have \( k + 1 \) parameters \( \pi_j j = 0, 1, \ldots, k \) subject to \( \sum_j \pi_j = 1 \), so the dimension is \( q = k \). Asymptotically,

\[ \Lambda(y) \xrightarrow{D} \chi^2(q - p) \]

with \( k = 5 \), \( q - p = 5 - 1 = 4 \) so \( \Lambda(y) \xrightarrow{D} \chi^2(4) \) and similarly \( P(y) \).

Now it is easy enough formally to make this test and obtain a \( p \)-value. Comparing \( P \) with a \( \chi^2(4) \) distribution we obtain \( p \simeq 0.94 \).

\[
> \text{pr0<-dpois(0:4,lambda=mu)}
> (\text{pr0<-c(pr0,1-sum(pr0)))})
> [1] 0.5433508691 0.3314440301 0.1010904292 0.0205550539 0.0031346457
> [6] 0.0004249719
> \text{E<-n*pr0}
> (\text{P<-sum( (0-E)^2/E ))}
> [1] 0.7903603
> (\text{1-pchisq(P,4))})
> [1] 0.9397357
\]

However, the cell counts for 3, 4 and \( \geq 5 \) deaths are too small for the asymptotics to be reliable. You had in Part A a rule of thumb of \( E_i \geq 5 \), and the last three cells violate this condition.

What we need in the example about is the probability that a random \( P(Y) \), computed from data \( Y \) simulated under \( H_0 \) exceeds \( p(y) \simeq 0.79 \), the test statistic computed on the data.

Suppose the data were Poisson with mean \( \hat{\mu} \). We can make a Monte Carlo test as follows.

1. Compute the MLE \( \hat{\mu} \) and the value of the test statistic \( P \) as above.
2. Simulate \( M \) iid data sets \( y^{(m)} = (y_1^{(m)}, \ldots, y_N^{(m)}) \) \( m = 1, 2, \ldots, M \), with

\[
\Pr(Y_i^{(m)} = j) = \pi_j(\hat{\mu}) \text{ independent of all other draws, using the MLE for } \mu \text{ under } H_0 \text{ to fix the data distribution under } H_0. \]

For each data set compute \( O^{(m)} \).
(3) Form the parameter estimate \( \hat{\mu}^{(m)} \), compute \( \pi_j(\hat{\mu}^{(m)}) \) and \( E_j^{(m)} = n \pi_j(\hat{\mu}^{(m)}) \) for \( j = 1, 2, ..., k \). Calculate
\[
P^{(m)} = \sum_j \frac{(O_j^{(m)} - E_j^{(m)})^2}{E_j^{(m)}}
\]
for \( m = 1, 2, ..., M \).

(4) Now \( p = \Pr(P(Y) > P(y)) \). Estimate this \( p \)-value as
\[
\hat{p} = \frac{1}{M} \sum_{m=1}^M I_{P^{(m)}>P}
\]
with estimated variance \( \hat{p}(1-\hat{p})/M \) for the estimator.

**Example 5.2.** We can apply this procedure to Example 5.1.

```r
> M<-1000
> Pm<-rep(0,M)
> for (m in 1:M) {
+   #Step 2 - simulate synthetic data
+   ym<-rpois(n=n,lambda=mu)
+   #count the number in each cell from 0 to 4 and >=5
+   Om<-hist(ym,breaks=c(0:5,max(ym)+1)-0.5,plot=FALSE)$counts
+   #step 3 compute the MLE and Pearson stat for the synthetic data
+   mum<-mean(ym) #MLE
+   pr0m<-dpois(0:4,lambda=mum)
+   pr0m<-c(pr0m,1-sum(pr0m))
+   Em<-n*pr0m
+   Pm[m]<-sum( (Om-Em)^2/Em )
+ }
> #step 4 estimate the p-value
> (p<-mean(Pm>P))
[1] 0.916
> (p.std<-sqrt(p*(1-p)/M))
[1] 0.008771773
```

Compare this \( \hat{p} \simeq 0.91(1) \) with \( p_{ASY} \simeq 0.94 \) above. It seems that, although the conditions for the use of the asymptotic result did not hold, the estimate we obtained was quite OK, and our Monte-Carlo test has confirmed the inference that there is no evidence for non-Poisson structure in the counts in Table 5.

5.3. **Goodness of fit for continuous rv.** The material in this section closely follows that in section 4.5 of Dr Lunn’s lecture notes on this topic from 06-07.

In the previous section we used a LRT to make a goodness of fit test. We parameterised the alternative as the most general distribution possible in the context. In this section we look at a goodness of fit test in which the alternative is simply “not-H0”.

It is easy to make up tests for deviation from a given null distribution. We can choose essentially any function of the data we like, and look for evidence that our data has a ‘strange’ value of this function. The function, and the critical region, can be chosen so the test is sensitive to particular kinds of departure from the null, though these choices must of course be made prior to seeing the data.
In the Kolmogorov-Smirnov one-sample test we look for evidence that given data are not distributed according to a given distribution. In the two sample test we take two data sets and look for evidence that they are not distributed according to a common but unknown underlying distribution. The idea is that, if two distributions of scalar rv are unequal, then their cdf’s will be unequal. The test statistic is a measure of the difference between two cdf’s. The method applies naturally to continuously distributed observations. This is just one function of the data, in the sense of the previous paragraph. However, it is a natural and sensitive check, since it measures misfit over the entire cdf.

Denote by $F(x) = \Pr(X \leq x)$ the cdf for the rv $X$. Let data $y_i, i = 1, 2, ..., n$ be given, and denote by $y(i)$ the sorted data. Denote by $F_e(x; Y)$ the empirical cdf,

$$F_e(x; Y) = \frac{1}{n} \sum_{i=1}^{n} \mathbb{1}_{y_i \leq x}.$$  

Notice that

$$F_e(y(i); y) = i/n.$$  

**Example 5.3.** Bat detection distances for 11 successful insect catches.

$$23 \ 27 \ 34 \ 40 \ 42 \ 45 \ 52 \ 56 \ 62 \ 68 \ 83$$

Is the distribution of the detection-distance exponential? The empirical cdf is $F_e(x) = 0$ for $x < 23$, $F_e(x) = 1/11$ for $23 \leq x < 27$, $F_e(x) = 2/11$ for $27 \leq x < 34$, and so on in steps of $1/n$ up to $F_e(x) = 10/11$ for $68 \leq x < 83$, and $F_e(x) = 1$ for $83 \leq x$.

Denote by $F$ the underlying cdf. The idea of the test is that, if the data do have cdf $F$, then the empirical distribution should closely follow $F$. Consider the statistic

$$d = \max_x |F(x) - F_e(x)|.$$  

The steps in $F_e$ occur at $x = y_i$, and since $F$ is increasing, the maxima in $F(x) - F_e(x)$ occur at data points, and are equal to $F_e(y(i)) - F(y(i)) = i/n - F(y(i))$. The maxima in $F(x) - F_e(x)$ occur just before a data point, and are equal to $F(y(i)) - F_e(y(i)) = F(y(i)) - (i - 1)/n$ (and notice $F_e(x) = 0$ for $x < y_1$). It follows that

$$d(y) = \max_{i=1, \ldots, n} \left\{ \frac{i}{n} - F(y(i)) - (i - 1)/n \right\}.$$  

The $p$-value for the one-sample KS test is $\Pr(d(Y) \geq d(F))$.

We will see that, if the parameters of the cdf $F$ under $H_0$ are all known, then the distribution of $F$ is something simple. However, the distribution of $d(Y)$ will otherwise be hard to compute, and this is the case of interest. We can use Monte Carlo to avoid the need to calculate a closed form for $\Pr(d(Y) \geq d(F))$. The procedure is fundamentally the same as before, but the test statistic is a little more complicated to calculate than $P$, the Pearson test statistic.

Let data $y = y_1, ..., y_n$ and a null cdf $F$ with unknown parameters $\theta$ be given.

**Step 1** Calculate the MLE $\hat{\theta}$ for any parameters of $F$.

**Step 2** Calculate $d(y)$ using $F$ evaluated with the data $y$ and MLE $\hat{\theta}$.

**Step 3** Simulate $M$ iid datasets $y^{(1)}, ..., y^{(M)}$ with $y^{(m)} = (y_1^{(m)}, ..., y_n^{(m)})$.  

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Step 4 For each set of simulated data $y^{(m)}$, compute the MLE $\hat{\theta}^{(m)}$, and calculate $d(y^{(m)})$ using $F$ evaluated with the $m$'th simulated data $y^{(m)}$ and derived MLE $\hat{\theta}^{(m)}$.

Step 5 Estimate the $p$-value $Pr(d(Y) \geq d(y)|F)$ as

$$\hat{p} = \frac{1}{M} \sum_{m=1}^{M} I\{d(y^{(m)}) > d(y)\}.$$ 

**Example 5.4.** Let us apply this to the Bat-detection data.

```r
> #Step 1. Notice the data are already sorted
> y<-c(23,27,34,40,42,45,52,56,62,68,83)
> (mu<-mean(y))
[1] 48.36364
> n<-11
> #Step 2. The cdf at the data, pexp(y,lambda) is the Exp(lambda) cdf
> F<-pexp(y,rate=1/mu)
> #the empirical cdf is $F_e(y_{(i)}; y) = i/n$ so $F_e - F$ at i=1…n is
> (1:n)/n-F
[1] -0.2875564  -0.2459841  -0.2321787  -0.1990338  -0.1258409  -0.0601697
[7] -0.0224028   0.0414196   0.0956761   0.1542090   0.1797541
> #and $F - F_e(y_{(i-1)})$ at i=1…n is
> F-(0:(n-1))/n
[1] 0.3784655 0.3368932 0.3230878 0.2899429 0.2167506 0.1542090
[7] 0.1133194 0.0494895 -0.004767 -0.0632999
[11] -0.08884502
> #the KS statistic $d$ is the largest difference
> (d<-max(c((1:n)/n-F,F-(0:(n-1))/n)))
[1] 0.3784655
> #Step 3 (each i=1…M) - and remember to sort!
> Ym<-sort(rexp(n,1/mu))
> mum<-mean(Ym) #Step 4.1 MLE of ith data set
> Fm<-pexp(Ym,rate=1/mum)
> d.sim[i]<-max(c((1:n)/n-Fm,Fm-(0:(n-1))/n)) #Step 4.2 KS for ith data
> (p<-mean(d.sim>d)) #step 5 p-hat
[1] 0.0058
```

The chances of getting such a poor fit to an exponential if the data were truly exponentially distributed are only $p \approx 0.006$, so we should look for some other shape for this distribution.

5.4. **Monte Carlo test, simple hypothesis.** In both Example 5.1 and Example 5.4 we were making tests for distributions with an unknown parameter. The null hypothesis is a composite hypothesis, in which the value of the parameter is not
fixed to a single value. In Example 5.1 the question was ‘are these data Poisson’, not ‘are these data Poisson with mean such-and-such’. Similarly, in the second example, these question was ‘are these data Exp-distributed’ and we didn’t have any fixed value for the mean.

Because we didn’t want to specify the parameters of the distribution under the null, we used the MLE for the parameter calculated from the data. When we came to do the Monte Carlo tests we calculated the parameter MLE for each synthetic data set, so that the process generating the synthetic-data test statistic mimicked the process generating the real data test statistic (and therefore gave the synthetic-data test-statistic the same distribution).

If we did know the value of a parameter, then we would use it when we calculate the test statistic (Pearson, or KS) for the real data. Since our Monte Carlo test aims to mimic the process that gave rise to the real-data test statistic, we would use the known value of the parameter when we simulate the synthetic data, and we would use the known value when we compute the test statistic for the synthetic data. In other words, if we know the value of a parameter we use the known value and not the MLE at all points in the Monte-Carlo significance test.

Exercise Look at Example 5.1. Suppose $\mu = 0.61$ was the exact known value of $\mu$, not simply the MLE as above. How would you modify the Monte-Carlo test code to make a Monte Carlo test for goodness of fit to Poisson for known $\mu$? Show that the lines

for (m in 1:M) {
    #step 3 compute the MLE and Pearson stat for the synthetic data
    mum<-mean(ym) #MLE
    pr0m<-dpois(0:4,lambda=mum)
}

in Example 5.1 become

for (m in 1:M) {
    #step 3 compute the Pearson stat for the synthetic data
    pr0m<-dpois(0:4,lambda=mu)
}

so that we compute the Pearson test statistic using the known fixed $\mu$-value.

The KS-test has a particularly simple form when the parameters of the comparison distribution (that is, the null distribution for the data, with cdf $F$) are known. In this case the distribution of the KS-test statistic $d(Y)$ has a distribution which doesn’t depend on $F$. Once we have $d(y)$ we can look up a $p$-value in standard tables. We don’t need to make Monte-Carlo tests to get the distribution of $d(Y)$ for $Y \sim F$.

Recall (see above) that the KS-test statistic computed on the data $y$ is

$$d(y) = \max_x |F(x) - F_e(x)|.$$
Suppose $F$ is strictly increasing (as is usually the case) so $a < b \Rightarrow F(a) < F(b)$. Then (following Dr Lunn’s lecture notes from 2007)

$$
\Pr(d(Y) < d(y)|F) = \Pr \left( \max_x \left\{ \frac{1}{n} \sum_{i=1}^{n} \mathbb{I}_{Y_i \leq x} - F(x) \right\} \right)
= \Pr \left( \max_x \left\{ \frac{1}{n} \sum_{i=1}^{n} \mathbb{I}_{F(Y_i) \leq F(x)} - F(x) \right\} \right).
$$

But if $U_i = F(Y_i)$ then $U_i \sim U(0, 1)$ (under the null) since $F$ is the cdf of $Y_i$. Let $z = F(x)$ and replace the search for maxima over $x$ with a search over $z \in (0, 1)$. The distribution of the KS-statistic for a simple hypothesis is

$$
\Pr(d(Y) < d(y)|F) = \Pr \left( \max_{z \in (0,1)} \left\{ \frac{1}{n} \sum_{i=1}^{n} \mathbb{I}_{U_i \leq z} - z \right\} \right),
$$

and, since the $U_i$ are simply iid uniform rv, this doesn’t depend on $F$. We could estimate this probability accurately for a range of $d(y)$-values from zero up to one, and record them. Then when we make future tests for goodness of fit, and get a $d(y)$ value, we can look up the $p$ value in our table, since that $p$ value doesn’t depend on any other details of the test.

The KS-test can be used to compare two data sets. Dr Lunn, (2007) lecture notes, gives the example of two collections of silver coins. The percentages of silver in the first collection are 5.9, 6.8, 6.4, 7.0, 6.6, 7.7, 7.2, 6.9, 6.2. The percentages of silver in the second collection are 5.3, 5.6, 5.5, 5.1, 6.2, 5.8, 5.8. Let $A \sim F_A$ be the percent silver in a coin in the first collection, and $B \sim F_B$ be the percent silver in a coin in the second collection. Are the two distributions the same, that is, is it the case that $F_A = F_B$? We would like to make this test without specifying a family of distributions for $A$ and $B$. The two sample test is based on the distance between the two empirical cdf’s.

**Exercise** Let $F_c(A)$ and $F_c(B)$ denote the empirical cdf’s for the percent silver in the two collections of coins. Show that $\max_x |F_c(A)(x) - F_c(B)(x)| = 6/7 - 1/9 = 47/63$. Look up the $p$-value in a KS table (for a pair of samples of size 9 and 7) and verify that $p \simeq 0.02$, so it is very unlikely that the simple picture of a common silver standard holds.

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