

Chapter 7: Models with Multiple Continuous Predictors

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In this chapter, we consider **multiple regression** and other models in which there are more than one predictor (or X) variable. This extends what we covered in the last chapter where we examined one predictor to multiple predictors. Once again our focus is on finding the estimates of coefficients or parameters in multiple linear regression models by the method of **least squares**. For this we assume that

1. The predictor (or explanatory or controlled or covariate) variables X_i ($i = 1, 2, \dots, p$) are known without error.
2. The mean or expected value of the dependent (or response) variable Y is related to the X_i ($i = 1, 2, \dots, p$) according to a linear expression

$$E(y | x) = a + b_1x_1 + b_2x_2 + \dots + b_p x_p$$

i.e. a straight line (for one X variable), a plane (for two X variables) or a hyperplane (for more than two X variables). This means that the fitted model can be written as

$$\text{fit} = a + b_1x_1 + b_2x_2 + \dots + b_p x_p.$$

3. There is random (unpredictable, unexplained) variability of Y about the fitted model. That is,

$$y = \text{fit} + \text{residual}.$$

4. In order to apply statistical inferences to a model, a number of assumptions need to be made. To be able to form t and F statistics, we assume that
5. The variability in Y about the line (plane etc) is constant and independent of the X variables.
6. The variability of Y follows a Normal distribution. That is, the distribution of Y (given certain values of the X_i variables) is Normal.
7. Given (different) outcomes of the X variables, the corresponding Y variables are independent of one another.

We will continue to use the data set **horsehearts** of the weights of horses' hearts and other related measurements.

1 Full Regression

With one explanatory variable scatterplots and correlation coefficients provided good starting points for exploring relationships between the explanatory and response variables. This is even more relevant with two or more explanatory variables. For the horses' hearts data, there are six potential explanatory variables; namely `EXTDIA`, `EXTSYS`, `INNERDIA`, `INNERSYS`, `OUTERDIA` and `OUTERSYS`. These measurements of heart width are made of the exterior width, inner wall and outer wall at two different phases, the diastole phase and the systole phase. So a matrix of scatter plots (or matrix plot) of these variables will be useful for exploratory analysis.

It is also a good idea to form the simple correlation coefficients between each pair of explanatory variables and between each explanatory variable and the response variable Y , the weight of the horse's heart. These correlation coefficients can be displayed in a **correlation matrix** as shown in Figure 1.1.

```
library(tidyverse)
library(tidymodels)
library(kableExtra)
theme_set(theme_minimal())

load("../data/horsehearts.RData")

library(GGally)

Registered S3 method overwritten by 'GGally':
  method from
  +.gg   ggplot2

ggpairs(horsehearts)
```

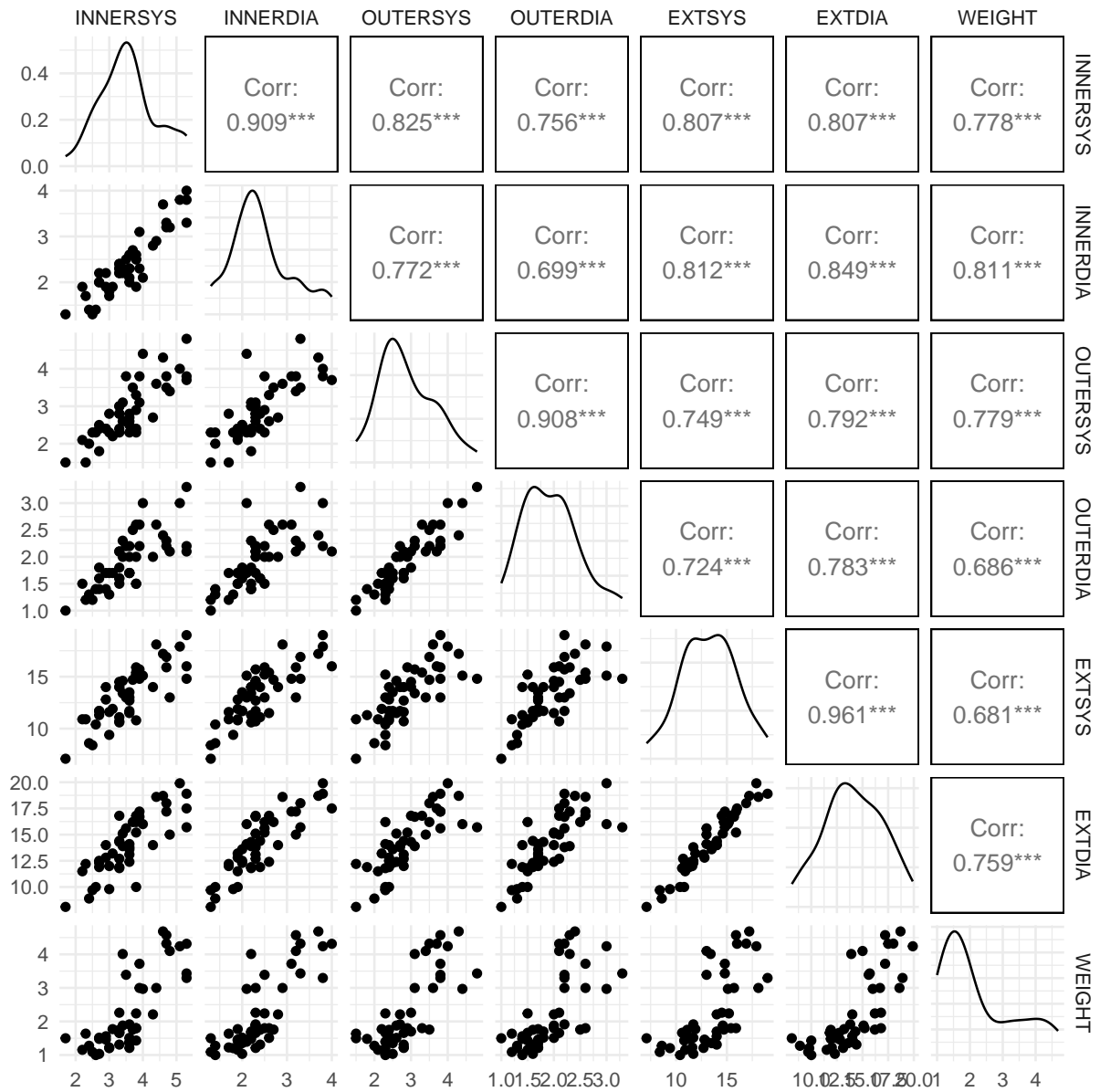


Figure 1.1: Scatter plot and correlation matrix

It is also possible to obtain the p -values for all the simple correlation coefficients displayed above and test whether these are significantly different from zero.

A number of facts about the data emerge from our EDA. All of the correlation coefficients are positive and reasonably large which indicates that with large hearts all the lengths increase in a fairly uniform manner. The predictor variables are also highly inter-correlated. **This suggests that not all of these variables are needed but only a subset of them.**

The usual `tidy()` function output of multiple regression weight on all of the available (six) predictors is shown in Table 1.1.

```
full.reg <- lm(WEIGHT ~ ., data=horsehearts)
tidy(full.reg) # or summary(full.reg)
```

Table 1.1: Full Regression `tidy()` output

term	estimate	std.error	statistic	p.value
(Intercept)	-1.631	0.488	-3.343	0.002
INNERSYS	0.232	0.308	0.753	0.456
INNERDIA	0.520	0.395	1.314	0.197
OUTERSYS	0.711	0.329	2.164	0.037
OUTERDIA	-0.557	0.451	-1.236	0.224
EXTSYS	-0.300	0.135	-2.227	0.032
EXTDIA	0.339	0.148	2.296	0.027

This regression model is known as the **full regression** because we have included all the predictors in our model. The R syntax `~.` means that we are placing all variables in the dataframe except the one selected as the response variable. We note that the slope coefficients of the predictors `INNERDIA`, `INNERSYS`, and `OUTERDIA` are not significant at 5% level. This confirms that we do not need to place all six predictors in the model but only a subset of them.

The highly correlated predictor `INNERDIA` (see Figure 1.1) is also found to have a insignificant coefficient in Table 1.1. This is somewhat surprising and casts doubts on the suitability of the full regression fit. If two or more explanatory variables are very highly correlated (i.e. almost collinear), then we deal with **multicollinearity**. The estimated standard errors of the regression coefficients will be inflated in the presence of multicollinearity. As result, the *t*-value will become small leading to a model with many insignificant coefficients. Multicollinearity does not affect the residual standard error much. The obvious remedy for multicollinearity is that one or more of the highly correlated variables can be dropped. Measures such as the Variance Inflation factor (**VIF**) are available to study the effect of multicollinearity. A VIF factor of more than 5 for a coefficient means that its variance is artificially inflated by the high correlation among the predictors. For the full regression model, the VIF factors are obtained using the `car` package function `vif()` and shown as Table 1.2.

```
car::vif(full.reg)
```

Table 1.2: Variance Inflation Factors

	x
INNERSYS	8.77
INNERDIA	8.60
OUTERSYS	7.71
OUTERDIA	6.66
EXTSYS	16.05
EXTDIA	22.00

All the VIF values are over 5, and hence the full regression model must be simplified dropping one or more predictors.

Let us now compare the fit and summary measures of the simple regression `lm(WEIGHT ~ INNERDIA, data=horsehearts)` with the full regression `lm(WEIGHT ~ ., data=horsehearts)`. Figure 1.2 compares the actual and fitted Y values for these two models.

```
library(modelr)

full.reg <- lm(WEIGHT ~ ., data=horsehearts)

simple.reg <- lm(WEIGHT ~ INNERDIA, data=horsehearts)

hhpred <- horsehearts |>
  gather_predictions(full.reg, simple.reg) |>
  mutate(residuals=WEIGHT-pred)

hhpred |>
  ggplot() +
  aes(x=WEIGHT, y=pred, colour=model) +
  geom_point() +
  geom_abline(slope=1, intercept = 0, alpha=.5) +
  theme(aspect.ratio = 1) +
  ylab("Predicted WEIGHT") +
  ggtitle("Comparison of model predictions")
```

Comparison of model predictions

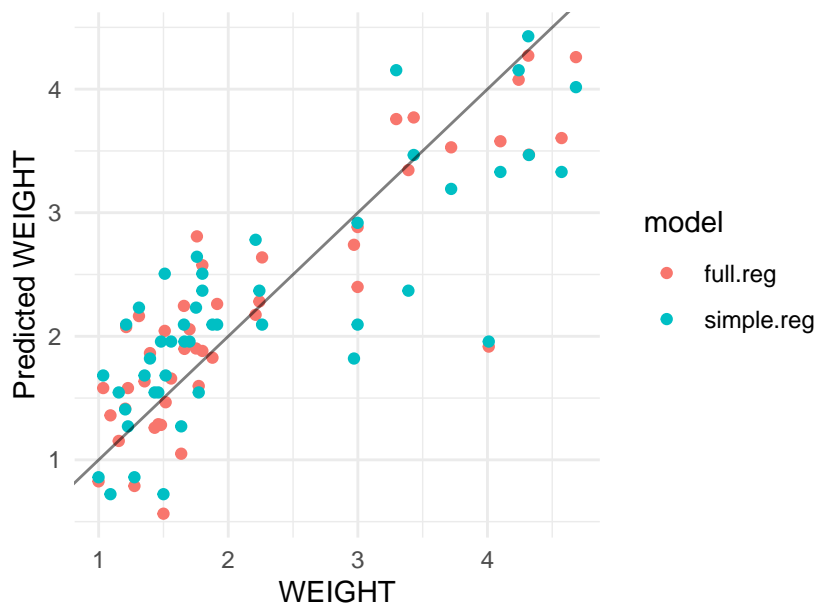


Figure 1.2: Comparison of Multiple Regression and Simple Regression

Both the simple and full regression models give similar predictions when the horses heart weight is below 2.5 kg, but the simple regression residuals are bit bigger for larger hearts.

We are rather hesitant to make unique claims about any particular subset of predictors based on the correlation matrix or based on the significance of the coefficients from the multiple regression output. In forthcoming sections, methods to decide on a subset of these variables will be explained, but first we look at the issues involved when predictor variables are correlated to each other.

2 Measuring Variation Explained by Predictors

The variation in a variable can be measured by its sum of squares. In this section, we illustrate this variation by the area of a circle. For brevity, we denote the **Total Sum of Squares**, the **Regression Sum of Squares** and the **Error or residual Sum of Squares** by **SST**, **SSR** and **SSE** respectively. In Figure 2.1, the circle is labelled y and represents the Sum of Squares for all the y observations, that is, SST.

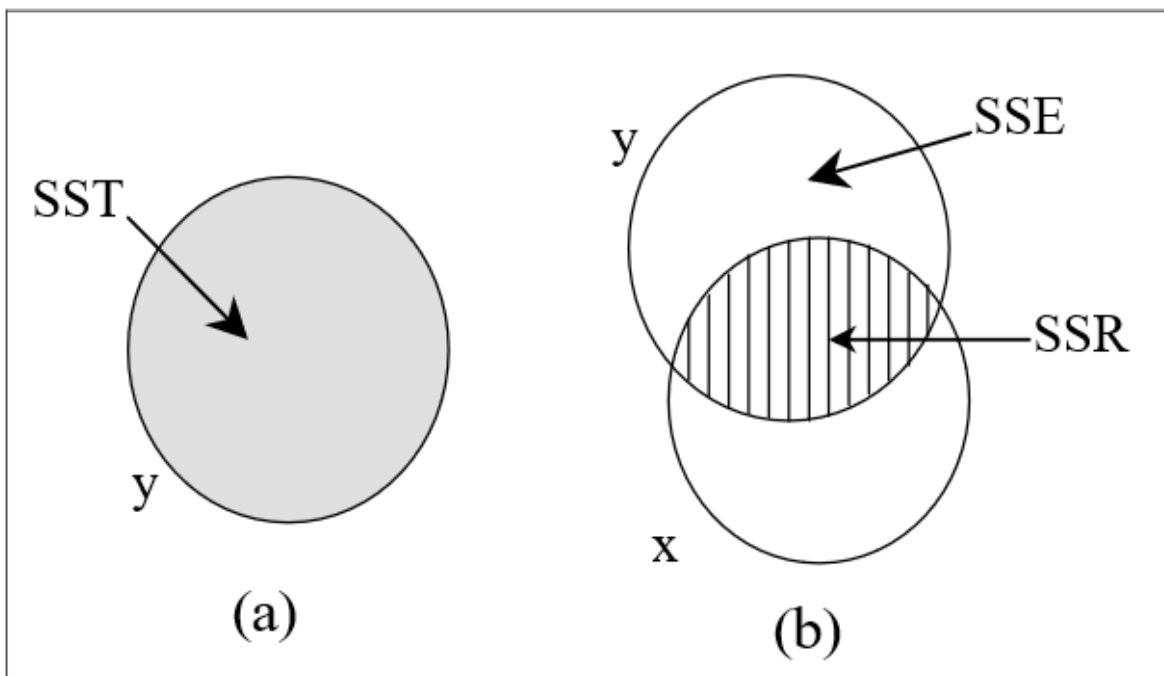


Figure 2.1: Effect of predictor correlation with the response

For the full regression of horse heart weight, we obtain the ANOVA output using the command

```
anova(full.reg)
```

Let's take a look at the sums of squares table.

```

SS <- anova(full.reg) |>
  tidy() |>
  select(term:sumsq) |>
  janitor::adorn_totals()

```

Table 2.1: Sums of squares for the full regression of horsehearts data

term	df	sumsq
INNERSYS	1	34.40
INNERDIA	1	3.53
OUTERSYS	1	2.76
OUTERDIA	1	0.13
EXTSYS	1	0.06
EXTDIA	1	1.90
Residuals	39	14.07
Total	45	56.85

Now let's calculate the Sums of Squares Total (SST), Error (SSE), and Regression (SSR).

```

tibble(
  SST = SS |> filter(term=="Total") |> pull(sumsq),
  SSE = SS |> filter(term=="Residuals") |> pull(sumsq),
  SSR = SST - SSE
)

# A tibble: 1 x 3
  SST    SSE    SSR
  <dbl> <dbl> <dbl>
1 56.8  14.1  42.8

```

In Figure 2.1(a), $SST = SSR + SSE = 32.731 + 24.115 = 56.845$. Consider now the straight line relationship between y and one response variable x . This situation is illustrated in Figure 2.1(b). The shaded overlap of the two circles illustrates the variation in y about the mean explained by the variable x , and this shaded area represents the regression sum of squares SSR. The remaining area of the circle for y represents the unexplained variation in y or the residual sum of squares SSE. Note that the circle or Venn diagrams represent SS only qualitatively (not to scale). The variation in y is thus separated into two parts, namely $SST = SSR + SSE$.

Notice that we are not very interested in the unshaded area of the circle representing the explanatory variable, x ; *it is the variation in the response variable, y* , which is important.

Also notice that the overlapping circles indicate that the two variables are correlated, that is the correlation coefficient, r_{xy} , is not zero. The shaded area is related to

$$R^2 = \text{proportion of the variation of } y \text{ explained by } x = \text{SSR/SST} = 32.731/56.845 = 0.576 = r_{xy}^2.$$

Note from Figure 1.1 that the correlation between WEIGHT and EXTDIA is 0.759 and 0.759 squared equals 0.576.

The situation becomes more interesting when a second explanatory variable is added to the model as illustrated by Figure 2.2.

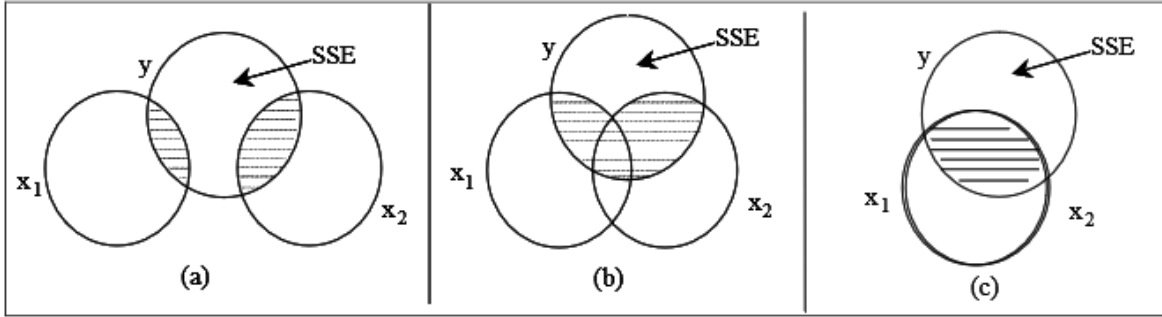


Figure 2.2: Effect of adding two predictors

In the following discussion, the variable EXTDIA is denoted by x_1 and OUTERDIA as x_2 . The total overlap of $(x_1$ and $x_2)$ and y will depend on the relationship of y with x_1 , y with x_2 , and the correlation of x_1 and x_2 .

In Figure 2.2(a), as the circles for x_1 and x_2 do not overlap, this represents a correlation coefficient between these two variables of zero. In this special case,

$$R^2 = \frac{\text{SSR}(x_1) + \text{SSR}(x_2)}{\text{SST}} = r_{x_1 y}^2 + r_{x_2 y}^2.$$

Here, $\text{SSR}(x_1)$ represents the Regression SS when y is regressed on x_1 only. $\text{SSR}(x_2)$ represents the Regression SS when y is regressed on x_2 only. The unshaded area of y represents SSE, the residual sum of squares, which is the sum of squares of y **unexplained** by x_1 or x_2 . The **special case of uncorrelated** explanatory variables is in many ways ideal but it usually only occurs when x_1 and x_2 are constructed to have zero correlation (which means that the situation, known as orthogonality, is usually confined to experimental designs). There is an added bonus when x_1 and x_2 have zero correlation. In this situation the fitted model is

$$\hat{y} = a + b_1 x_1 + b_2 x_2$$

where b_1 and b_2 take the same values as in the separate straight line models $\hat{y} = a + b_1 x_1$ and $\hat{y} = a + b_2 x_2$.

However in observational studies the correlations between predictor variables will usually be nonzero. The circle diagram shown in Figure 2.2(b) illustrates the case when x_1 and x_2 are correlated. In this case

$$R^2 < \frac{\text{SSR}(x_1) + \text{SSR}(x_2)}{\text{SST}}$$

and the slope coefficients for both x_1 and x_2 change when both these variables are included in the regression model.

Figure 2.2(c) gives the extreme case when x_1 and x_2 have nearly perfect correlation. If the correlation between x_1 and x_2 is perfect, then the two variables will be said to be collinear. If two or more explanatory variables are very highly correlated (i.e. almost collinear), then we deal with **multicollinearity**.

From Figure 2.1 and Figure 2.2, it is clear that for correlated variables, the variation (SS) explained by a particular predictor cannot be independently extracted (due to the commonly shared variation). Hence, we consider how much a predictor explains **additionally** given that there are already certain predictors in the model. The additional overlap due to x_2 with y **after** x_1 , known as the **additional SSR** or **Sequential SS** is an important idea in model building. The additional SSR is known as **Type I sums of squares** in the statistical literature.

Note that we can also define the additional variation in y explained by x_1 after x_2 . It is important to note that in general the additional SSR depends on the **order** of placing the predictors. **This order does not have any effect on the coefficient estimation, standard errors etc.**

2.1 Significance testing of Type I SS

The significance of the additional variation explained by a predictor can be tested using a t or F statistic. Consider the simple regression model of **WEIGHT** on **EXTDIA**. Suppose we decided to add the explanatory variable **OUTERDIA** to the model, i.e. regress **WEIGHT** on two explanatory variables **EXTDIA** and **OUTERDIA**. Is this new model a significant improvement on the existing one? For testing the null hypothesis that the true slope coefficient of **OUTERDIA** in this model is zero, the t -statistic is 1.531 (see output below).

```
twovar.model <- lm(WEIGHT ~ EXTDIA+OUTERDIA, data=horsehearts)

twovar.model |> tidy()

# A tibble: 3 x 5
  term      estimate std.error statistic p.value
  <chr>      <dbl>     <dbl>      <dbl>    <dbl>
1 (Intercept)  100.       10.        10.0     0.000
2 EXTDIA      15.3       1.53      10.0     0.000
3 OUTERDIA     1.53       1.53      1.00     0.312
```

1	(Intercept)	-1.97	0.551	-3.57	0.000885
2	EXTDIA	0.226	0.0614	3.68	0.000637
3	OUTERDIA	0.522	0.341	1.53	0.133

The t and F distributions are related by the equation $t^2 = F$ when the numerator df is just one for the F statistic. Hence $1.532 = 2.34$ is the F value for testing the significance of the additional SSR due to `OUTERDIA`. In other words, the addition of `OUTERDIA` to the simple regression model does not result in a significant improvement in the sense that the reduction in residual SS ($= 1.247$) as measured by the F value of 2.34 is not significant (p -value being 0.133).

```
onevar.model <- lm(WEIGHT~ EXTDIA, data=horsehearts)

twovar.model <- lm(WEIGHT~ EXTDIA+OUTERDIA, data=horsehearts)

anova(onevar.model, twovar.model)
```

Analysis of Variance Table

```
Model 1: WEIGHT ~ EXTDIA
Model 2: WEIGHT ~ EXTDIA + OUTERDIA
  Res.Df   RSS Df Sum of Sq    F Pr(>F)
1     44 24.115
2     43 22.867  1    1.2472 2.3453  0.133
```

Although `OUTERDIA` is correlated with `WEIGHT`, it also has high correlation with `EXTDIA`. In other words, the correlation matrix gives us some indication of how many variables might be needed in a multiple regression model, although by itself it cannot tell us what combination of predictor variables is good or best.

Figure 2.3 and Figure 2.4 summarise the following facts:

- When there is only **one** explanatory variable, $R^2 = \text{SSR/SST}$ equals the square of the correlation coefficient between that variable and the dependent variable. Therefore if only one variable is to be chosen, it should have the highest correlation with the response variable, Y .
- When variables are added to a model, the regression sum of squares SSR will increase and the residual or error sum of squares SSE will reduce. The opposite is true if variables are dropped from the model. This fact follows from Figure 2.4.
- The other side of the coin to the above remark is that as additional variables are added, the Sums of Squares for residuals, SSE, will decrease towards zero as also shown in Figure 2.4(c).

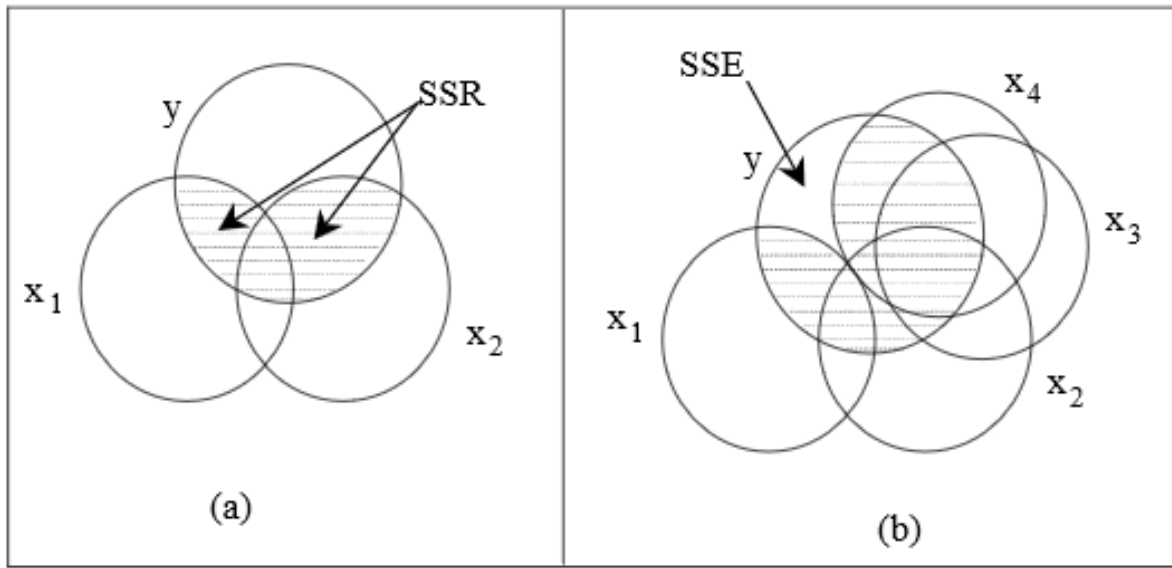


Figure 2.3: Issues with multiple predictors

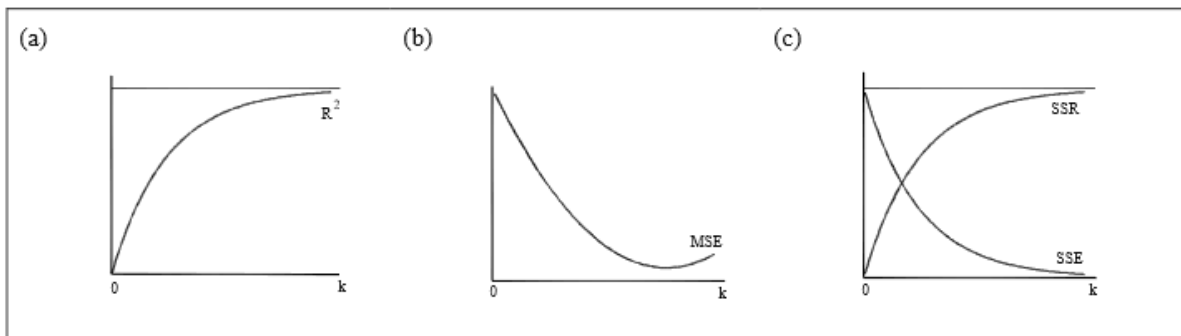


Figure 2.4: Effect of multiple predictors on model summaries

4. The overlap of circles in suggests that these changes in both SSR and SST will lessen as more variables are added, see Figure 2.3(b).
5. Following on from the last two notes, as $R^2 = \text{SSR}/\text{SST}$, R^2 will increase monotonically towards 1 as additional variables are added to the model. (monotonically increasing means that it never decreases although it could remain the same). This is indicated by Figure 2.4(a). If variables are dropped, then R^2 will monotonically decrease.
6. Against the above trends, the graph of residual mean square in Figure 2.4(b) reduces to a *minimum* but may eventually start to increase if enough variables are added. The residual sum of squares SSE decreases as variables are added to the model (see Figure 2.3(b)). However, the associated df values also decrease so that the residual standard deviation decreases at first and then starts to increase as shown in Figure 2.4(b). (Note that the residual standard error s_e is the square root of the residual mean square

$$s_e^2 = \frac{\text{SSE}}{\text{error degrees of freedom}},$$

denoted as MSE in Figure 2.3(b)). After a number of variables have been entered, the additional amount of variation explained by them slows down but the degrees of freedom continues to change by 1 for every variable added, resulting in the eventual increase in residual mean square. Note that the graphs in Figure 2.3 are idealised ones. For some data sets, the behaviour of residual mean square may not be monotone.

7. Notice that the above trends will occur even if the variables added are **garbage**. For example, you could generate a column of random data or a column of birthdays of your friends, and this would improve the R^2 **but not the adjusted R^2** . The adjusted R^2 makes adjustment for the degrees of freedom for the SSR and SSE, and hence reliable when compared to the unadjusted or multiple R^2 . The residual mean square error also partly adjusts for the drop in the degrees of freedom for the SSE and hence becomes an important measure. The addition of unimportant variables will not improve the adjusted R^2 and the mean square error s_e^2 .

2.2 Other SS types

The R `anova` function `anova()` calculates sequential or Type-I SS values.

Type-II sums of squares is based on the principle of marginality. Type II SS correspond to the R convention in which each variable effect is adjusted for all other *appropriate* effects.

Type-III sums of squares is the SS added to the regression SS after ALL other predictors including an intercept term. This SS however creates theoretical issues such as violation of marginality principle and we should avoid using this SS type for hypothesis tests.

The R package `car` has the function `Anova()` to compute the Type II and III sums of squares.
Try-

```
full.model <- lm(WEIGHT~ ., data=horsehearts)

anova(full.model)

library(car)

Anova(full.model, type=2)
Anova(full.model, type=3)
```

For the `horsehearts` data, a comparison of the Type I and II sums squares is given below:

```
full.model <- lm(WEIGHT~ ., data=horsehearts)

anova1 <- full.model |>
  anova() |>
  tidy() |>
  select(term, "Type I SS" = sumsq)

anova2 <- full.model |>
  Anova(type=2) |>
  tidy() |>
  select(term, "Type II SS" = sumsq)

type1and2 <- full_join(anova1, anova2, by="term")

type1and2
```

term	Type I SS	Type II SS
INNERSYS	34.40	0.20
INNERDIA	3.53	0.62
OUTERSYS	2.76	1.69
OUTERDIA	0.13	0.55
EXTSYS	0.06	1.79
EXTDIA	1.90	1.90
Residuals	14.07	14.07

When predictor variables are correlated, it is difficult to assess their absolute importance and the importance of a variable can be assessed only relatively. This is not an issue with the most

highly correlated predictor in general.

3 Regression Fitting with Fewer Predictors

The first step before selection of the best subset of predictors is to study the correlation matrix. For horses' heart data, the explanatory variable which is most highly correlated with y (**WEIGHT**) is x_2 (**INNERDIA**) having a correlation coefficient of 0.811 (see Figure 1.1). This means that **INNERDIA** should be the single best predictor. We may guess that the next best variable to join **INNERDIA**. This would be x_3 (**OUTERSYS**) but the correlations between x_3 and the other explanatory variables clouds the issue. In other words, the significance or otherwise of a variable in a multiple regression model depends on the other variables in the model.

Consider the regression of horses' heart **WEIGHT** on **INNERDIA**, **OUTERSYS**, and **EXTSYS**.

```
threevar.model <- lm(WEIGHT ~ INNERDIA + OUTERSYS + EXTSYS,  
                      data=horsehearts)  
  
threevar.model |> tidy()
```

term	estimate	std.error	statistic	p.value
(Intercept)	-1.341	0.474	-2.828	0.007
INNERDIA	0.958	0.262	3.649	0.001
OUTERSYS	0.597	0.203	2.940	0.005
EXTSYS	-0.034	0.063	-0.534	0.596

The coefficient of **EXTSYS** is not significant at 5% level. However coefficient of **EXTSYS** was found to be significant in the full regression. The significance of **INNERDIA** coefficient has also changed. This example shows that *we cannot fully rely on the t-test* and discard a variable because its coefficient is insignificant.

There are various search methods for finding the best subset of explanatory variables. We will consider **stepwise procedures**, namely algorithms that follow a series of steps to find a good set of predictors. At each step, the current regression model is compared with competing models in which one variable has either been added (*forward selection* procedures) or removed (*backward elimination* procedures). Some measure of goodness is required so that the variable selection procedure can decide whether to switch to one of the competing models or to stop at the current best model. Of the two procedures, backward elimination has two advantages. One is computational: step 2 of the forward selection requires calculation of a large number of

competing models whereas step 2 of the backward elimination only requires one. The other is statistical and more subtle. Consider two predictor variables x_i and x_j and suppose that the forward selection procedure does not add either because their individual importance is low. It may be that their joint influence is important, but the forwards procedure has not been able to detect this. In contrast, the backward elimination procedure starts with all variables included and so is able to delete one and keep the other.

A stepwise regression algorithm can also combine *both* the backward elimination and forward selection procedures. The procedure is the same as forward selection, but immediately after each step of the forward selection algorithm, a step of backward elimination is carried out.

Variable selection solely based p values is preferred only for certain applications such as analysis of factorial type experimental data where response surfaces are fitted. The base R does model selection based on *AIC* which has to be as minimum as possible for a good model. We shall now discuss the concept of *AIC* and other model selection criteria.

One way to balance model fit with model complexity (number of parameters) is to choose the model with the **minimal** value of Akaike Information Criterion (**AIC** for short, derived by Prof. Hirotugu Akaike as the minimum information theoretic criterion):

$$AIC = n \log\left(\frac{SSE}{n}\right) + 2p$$

Here n is the size of the data set and p is the number of variables in the model. A model with more variables (larger value of p) will produce a smaller residual sum of squares SSE but is penalised by the second term.

Bayesian Information Criterion (*BIC*) (or also called Schwarz's Bayesian criterion, *SBC*) places a higher penalty that depends on n , the number of observations. As a result *BIC* fares well for selecting a model that explains the relationships well while *AIC* fares well when selecting a model for prediction purposes.

A number of corrections to *AIC* and *BIC* have been proposed in the literature depending on the type of model fitted. We will not study them in this course.

An alternative measure called Mallow's C_p index is also available using which we may judge whether the variables at the current step (smaller model) are excessive or short. If unimportant variables are added to the model, then the variance of the fitted values will increase. Similarly if important variables are added, then the bias of the fitted values will decrease. The C_p index, which balances the variance and bias, is given by the formula

$$C_p = \frac{\text{SS Error for Smaller Model}}{\text{Mean Square Error for full regression}} - (n - 2p)$$

where p = no. of estimated coefficients (including the intercept) in the smaller model and n = total number of observations. The most desired value for the C_p index is the number

of parameters (including the y -intercept) or just smaller. If $C_p \gg p$, the model is biased. On the other hand, if $C_p \ll p$, the model associated variability is too large. The trade-off between bias and variance is best when $C_p = p$. But the C_p index is not useful in judging the adequacy of the full regression model because it requires an assumption on what constitutes the full regression. This is not an issue with the *AIC* or *BIC* criterion.

For prediction modelling, the following three measures are popular and the `modelr` package will extract these prediction accuracy measures and many more.

Mean Squared Deviation (MSD):

MSD is the mean of the squared errors (i.e., deviations).

$$MSD = \frac{\sum (\text{observation-fit})^2}{\text{number of observations}},$$

MSD is also sometimes called the Mean Squared Error (MSE). Note that while computing the MSE, the divisor will be the degrees of freedom and not the number of observations. The square-root of MSE is abbreviated as *RMSE*, and commonly employed as a measure of prediction accuracy.

Mean Absolute Percentage Error (MAPE):

MAPE is the average percentage relative error per observation. MAPE is defined as

$$MAPE = \frac{\sum \frac{|\text{observation-fit}|}{\text{observation}}}{\text{number of observations}} \times 100.$$

Note that MAPE is unitless.

Mean Absolute Deviation (MAD):

MAD is the average absolute error per observation and also known as MAE (mean absolute error). MAD is defined as

$$MAD = \frac{\sum |\text{observation-fit}|}{\text{number of observations}}.$$

For the horsehearts data, stepwise selection can be implemented using many R packages including `MASS`, `car`, `leaps` `HH` `caret`, and `SignifReg`. Examples given below are based on the horses hearts data.

1. The `step()` function performs a combination of both forward and backward regression. This method favours a model with four variables: `WEIGHT ~ INNERDIA + OUTERSYS + EXTSYS + EXTDIA`

```

full.model <- lm(WEIGHT ~ ., data = horsehearts)

stats::step(full.model)
# or MASS::stepAIC(full.model)
# or step(full.model, trace = FALSE)

```

2. The `stepAIC()` function from the MASS package can also be used instead of the `step()` function. Try-

```

library(MASS, exclude="select")

stepAIC(full.reg, direction="backward", trace = FALSE)
stepAIC(full.reg, direction="both", trace = FALSE)

null.model <- lm(WEIGHT~ 1, data=horsehearts)

stats::step(
  full.reg,
  scope = list(
    lower = null.model,
    upper = ~INNERSYS+INNERDIA+OUTERSYS+OUTERDIA+EXTSYS+EXTDIA
  ),
  direction = "forward")

```

3. The `SignifReg` package allows variable selection under various criteria. Try-

```

library(SignifReg)

SignifReg(full.reg,
          direction = "backward",
          criterion = "BIC",
          adjust.method = "none")

SignifReg(full.reg,
          direction = "backward",
          criterion = "r-adj",
          adjust.method = "none")

SignifReg(full.reg,
          direction = "backward",
          criterion = "p-value",

```

```

adjust.method = "none")

SignifReg(full.reg,
           direction = "both",
           criterion = "BIC",
           adjust.method = "none")

SignifReg(full.reg,
           direction = "both",
           criterion = "r-adj",
           adjust.method = "none")

SignifReg(full.reg,
           direction = "both",
           criterion = "p-value",
           adjust.method = "none")

```

The forward selection procedure also picks only just two variables as seen from the following output:

```

full.model <- lm(WEIGHT ~ ., data=horsehearts)

stmdl <- SignifReg(full.reg,
                     direction = "both",
                     criterion = "AIC",
                     adjust.method = "none")

stmdl |> tidy()

```

term	estimate	std.error	statistic	p.value
(Intercept)	-1.512	0.468	-3.230	0.002
INNERDIA	0.799	0.267	2.987	0.005
OUTERSYS	0.493	0.204	2.415	0.020
EXTSYS	-0.236	0.122	-1.938	0.059
EXTDIA	0.250	0.130	1.920	0.062

For the full regression model, the *AIC* is -40.5 and it drops to -42.35 for the four variable model. That is, according to the AIC criterion, a further reduction in model size does not compensate for the decline in model fit as measured by the AIC. The C_p index also recommends the four variable model because for the C_p value of 4.9 is closer to 5, the number of model coefficients.

4. Step-wise selection of predictors can also be done along with cross validation in each step. The R package *caret* enables this. For the horses heart data, the following codes perform the backward regression.

```
library(caret)
library(leaps)

set.seed(123)

fitControl <- trainControl(
  method = "repeatedcv",
  number = 5,
  repeats = 100
)

leapBackwardfit <- train(
  WEIGHT ~ .,
  data = horsehearts,
  trControl = fitControl,
  method = "leapBackward"
)

summary(leapBackwardfit)
```

Note that an asterisk in the row means that a particular variable is included in the step. The model in the last step excludes `INNERSYS` and `OUTERDIA`. On the other hand, the forward regression includes only two variables namely `INNERDIA` and `OUTERSYS`. We can also directly use the `leaps` package without cross validation.

```
fitControl <- trainControl(
  method = "repeatedcv",
  number = 5,
  repeats = 100
)

leapForwardfit <- train(
  WEIGHT ~ .,
  data = horsehearts,
  trControl = fitControl,
  method = "leapForward"
)
```

```

summary(leapForwardfit)

Subset selection object
6 Variables (and intercept)
  Forced in Forced out
INNERSYS FALSE FALSE
INNERDIA FALSE FALSE
OUTERSYS FALSE FALSE
OUTERDIA FALSE FALSE
EXTSYS FALSE FALSE
EXTDIA FALSE FALSE
1 subsets of each size up to 2
Selection Algorithm: forward
      INNERSYS INNERDIA OUTERSYS OUTERDIA EXTSYS EXTDIA
1 ( 1 ) " "    "*"    " "    " "    " "    " "
2 ( 1 ) " "    "*"    "*"    " "    " "    " "

```

3.1 Best Subsets Selection

An exhaustive screening of all possible regression models (and hence the name **best subsets** regression) can also be done using software. For example, there are 6 predictor variables in the horses' hearts data. If we fix the number of predictors as 3, then $\binom{6}{3} = 20$ regression models are possible. One may select the 'best' 3-variable model based on criteria such as AIC, C_p , R^2_{adj} etc. Software must be employed to perform the conventional stepwise regression procedures. Software algorithms give one or more best candidate models fixing the number of variables in each step.

On the basis of our analysis on the horses' hear data, we might decide to recommend the model with predictor variables EXTDIA, EXTSYS, INNERDIA and OUTERSYS. In particular if the model is to be used for describing relationships then we would tend to include more variables. For prediction purposes, however, a simpler feasible model is preferred and in this case we may opt for the smaller model with only INNERDIA and OUTERSYS. See Table 3.3 produced using the following R codes:

```

library(leaps)
library(HH)
library(kableExtra)

b.model <- regsubsets(WEIGHT ~ ., data = horsehearts) |>

```

```

summaryHH()

b.model |>
  kable(digits = 3) |>
  kable_styling(bootstrap_options = "basic", full_width = F)

```

Table 3.3: Subset selection

model	p	rsq	rss	adjr2	cp	bic	stderr
INNERD	2	0.658	19.450	0.650	11.923	-41.677	0.665
INNERD-OUTERS	3	0.715	16.173	0.702	4.838	-46.335	0.613
INNERD-OUTERS-OUTERD	4	0.718	16.043	0.698	6.477	-42.878	0.618
INNERD-OUTERS-EXTS-EXTD	5	0.741	14.739	0.715	4.862	-42.949	0.600
INNERD-OUTERS-OUTERD-EXTS-EXTD	6	0.749	14.272	0.718	5.566	-40.602	0.597
INNERS-INNERD-OUTERS-OUTERD-EXTS-EXTD	7	0.753	14.067	0.714	7.000	-37.437	0.601

Sometimes theory may indicate that a certain explanatory variable should be included in the model (e.g. due to small sample size). If this variable is found to make an insignificant contribution to the model, then one should exclude the variable when the model is to be used for prediction but if the model is to be used for explanation purposes only then the variable should be included. Other considerations such as cost and time may also be taken into account. For every method or algorithm, one could find peculiar data sets where it fouls up. The moral – be alert and don't automatically accept models thrown up by a program. Note there is **never one right answer** as different methods and different criteria lead to different models.

Variable selection procedures can be a valuable tool in data analysis, particularly in the early stages of building a model. At the same time, they present certain dangers. There are several reasons for this:

1. These procedures automatically snoop through many models and may select ones which, by chance, happen to fit well.
2. These forward or backward stepwise procedures are *heuristic* (i.e., shortcut) algorithms, which often work very well but which may not always select the best model for a given number of predictors (here best may refer to adjusted R^2 -values, or AIC or some other criterion).
3. Automatic procedures cannot take into account special knowledge the analyst may have about the data. Therefore, the model selected may not be the best (or make sense) from a practical point of view.
4. Methods are available that *shrink* coefficients towards zero. The least squares approach minimises the residual sums of squares or RSS without placing any constraint on the

coefficients. The shrinkage methods, which place a constraint on the coefficients, work well when there are large numbers of predictors. A *ridge regression* shrinks the coefficients towards zero but in relation to each other. On the other hand, (Least Absolute Selection and Shrinkage Operator) *lasso* regression shrinks some of coefficients to zero which means these predictors can be dropped. Note that the ridge regression does not completely remove predictors. By shrinking large coefficients, we obtain a model with higher bias but lower variance. This process is known as *regularisation* in the literature (not covered in this course).

4 Polynomial Models

Consider the **pinetree** data set which contains the circumference measurements of pine trees at four positions. The simple regression of the top circumference on the first (bottom) circumference, the fit is $\text{Top} = -6.33 + 0.763 \text{ First}$. This fit is satisfactory on many counts (highly significant t and F values, high R^2 etc); see Table 4.1 and Table 4.2.

```
load("../data/pinetree.RData")  
  
pine1 <- lm(Top ~ First, data = pinetree)  
  
pine1 |> tidy()  
  
# A tibble: 2 x 5  
  term      estimate std.error statistic p.value  
  <chr>    <dbl>     <dbl>     <dbl>     <dbl>  
1 (Intercept) -6.33     0.765    -8.28 2.10e-11  
2 First       0.763     0.0240    31.8  2.20e-38
```

Table 4.1: `tidy()` output of `lm(Top~First, data=pinetree)`

term	estimate	std.error	statistic	p.value
(Intercept)	-6.334	0.765	-8.278	0
First	0.763	0.024	31.779	0

```
pine1 |>  
glance() |>  
select(adj.r.squared, sigma, statistic, p.value, AIC, BIC)
```

Table 4.2: `glance()` output of `lm(Top~First, data=pinetree)`

adj.r.squared	sigma	statistic	p.value	AIC	BIC
0.945	1.291	1009.896	0	204.85	211.133

The residual plot, shown as Figure 4.1, still provides an important clue that we should try a polynomial (cubic) model.

```
library(ggfortify)
```

```
Registered S3 method overwritten by 'ggfortify':
  method      from
  autoplot.glmnet parsnip
```

```
autoplot(pine1, which=1, ncol=1)
```

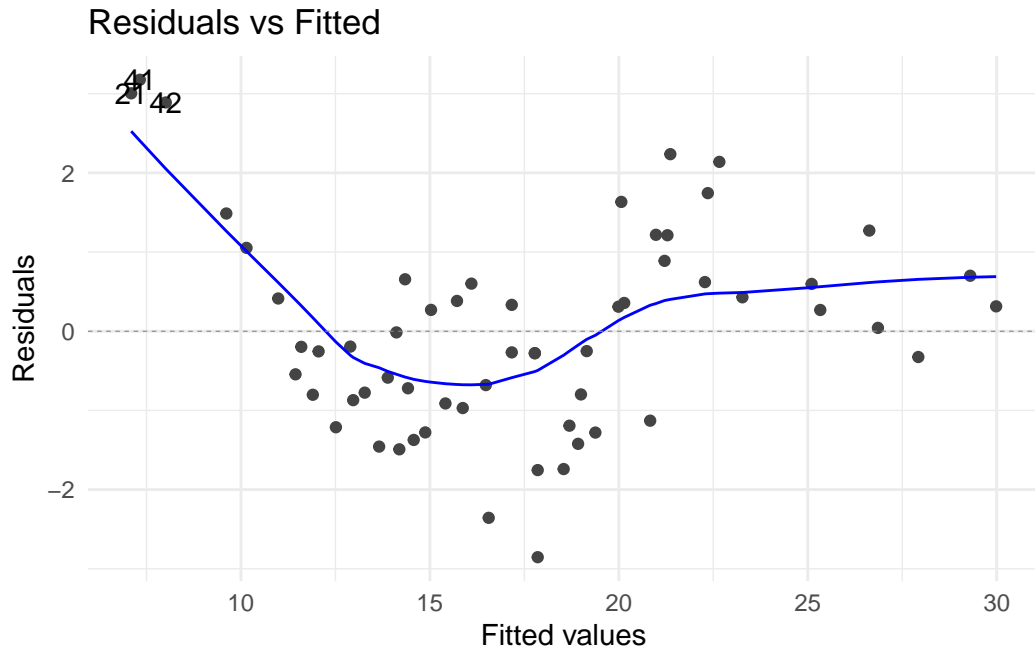


Figure 4.1: Residuals vs fits plot

```
pine3 <- lm(Top ~ poly(First, 3, raw=TRUE),
             data = pinetree)
```

Table 4.3 shows the significance results for the polynomial model $\text{Top} = 44.1 - 3.97\text{First} + 0.142(\text{First})^2 - 0.00135(\text{First})^3$. This model has achieved a good reduction in the residual standard error and improved AIC and BIC (see Table 4.4). The residual diagnostic plots are somewhat satisfactory. The Scale-Location plot suggests that there may be a subgrouping variable. The fitted model can be further improved using the Area categorical factor. This topic, known as the analysis of covariance will be covered later on. Note that both models are satisfactory in terms of Cook's distance. A few leverage or h_{ii} values cause concern seen in Figure 4.2 but we will ignore them given the size of the data set.

```
pine3 |> tidy()
```

Table 4.3: tidy() output of lm(Top~poly(First, 3, raw=TRUE), data=pinetree)

term	estimate	std.error	statistic	p.value
(Intercept)	44.1213	7.0391	6.2680	0
poly(First, 3, raw = TRUE)1	-3.9716	0.6951	-5.7141	0
poly(First, 3, raw = TRUE)2	0.1416	0.0221	6.3939	0
poly(First, 3, raw = TRUE)3	-0.0014	0.0002	-5.9510	0

```
pine3 |>
  glance() |>
  select(adj.r.squared, sigma, statistic, p.value, AIC, BIC)
```

Table 4.4: glance() output of lm(Top~poly(First,3, raw=TRUE), data=pinetree)

adj.r.squared	sigma	statistic	p.value	AIC	BIC
0.97	0.89	725.98	0	162.46	172.93

```
autoplot(pine3, which=5, ncol=1)
```

Residuals vs Leverage

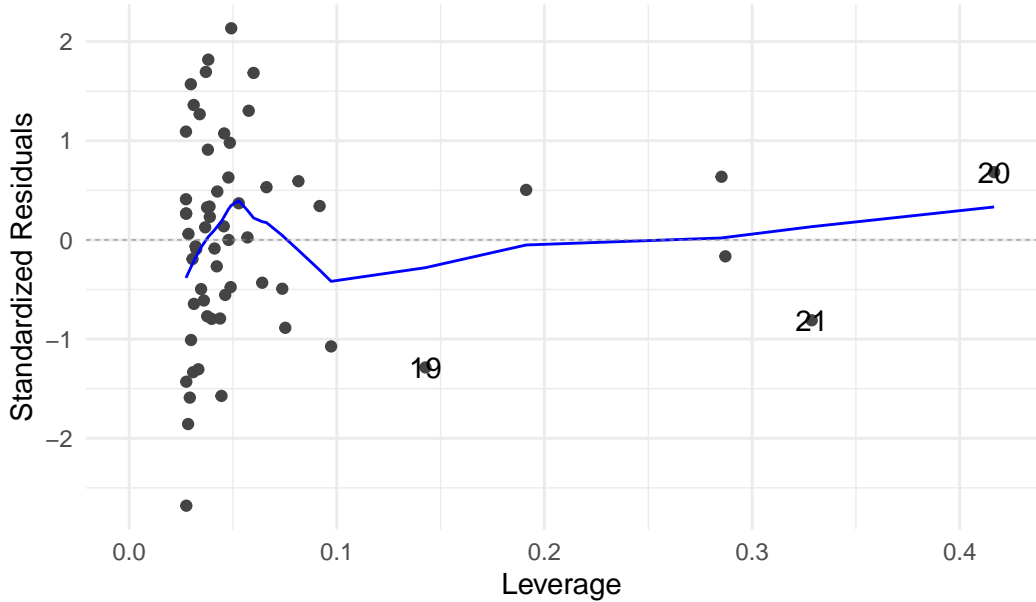


Figure 4.2: Residual vs Leverage plot

How quadratic and quartic models fare compared to the cubic fit is also of interest. Try the R code given below and compare the outputs:

```
summary(lm(WEIGHT ~ OUTERDIA, data = horsehearts))
summary(lm(WEIGHT ~ poly(OUTERDIA, 2, raw=T), data = horsehearts))
summary(lm(WEIGHT ~ poly(OUTERDIA, 3, raw=T), data = horsehearts))
summary(lm(WEIGHT ~ poly(OUTERDIA, 4, raw=T), data = horsehearts))
```

The key model summary measures of the four polynomial models are shown in Table 4.5. As expected, the R^2 value increases, although not by much in this case, as polynomial terms are added. Note that the multicollinearity among the polynomial terms renders all the coefficients of the quadratic regression insignificant at 5% level. For the cubic regression model, all the coefficients are significant. It is usual to keep adding the higher order terms until there is no significant increase in the additional variation explained (measured by the t or F statistic). Alternatively we may use the AIC criterion. In the above example, when the quartic term OUTERDIA^4 is added, the AIC slightly increases to 114.99 (from 114.65) suggesting that we may stop with the cubic regression.

```
modstats <- list(
  straight.line = lm(WEIGHT ~ OUTERDIA,
    data = horsehearts),
```

```

quadratic = lm(WEIGHT ~ poly(OUTERDIA,2,raw=T),
               data=horsehearts),
cubic = lm(WEIGHT~ poly(OUTERDIA,3, raw=T),
            data=horsehearts),
quartic = lm(WEIGHT~ poly(OUTERDIA,4, raw=T),
             data=horsehearts)
) |>
enframe(
  name = "model",
  value = "fit"
) |>
mutate(glanced = map(fit, glance)) |>
unnest(glanced) |>
select(model, r.squared, adj.r.squared, sigma,
       statistic, AIC, BIC)

modstats

```

Table 4.5: Glancing Polynomial models

model	r.squared	adj.r.squared	sigma	statistic	AIC	BIC
straight.line	0.47	0.46	0.83	39.13	117.01	122.50
quadratic	0.48	0.46	0.83	19.87	118.17	125.49
cubic	0.54	0.51	0.79	16.38	114.65	123.79
quartic	0.56	0.51	0.79	12.81	114.99	125.96

It is desirable to keep the coefficients the same when higher order polynomial terms are added. This can be done using orthogonal polynomial coefficients (we will skip the theory) for which we will avoid the argument `raw` within the function `poly()`. Try-

```

lm(Top ~ poly(First,1), data=pinetree)
lm(Top ~ poly(First,2), data=pinetree)
lm(Top ~ poly(First,3), data=pinetree)

```

Stepwise methods are not employed for developing polynomial models as it would not be appropriate (say) to have the linear and cubic terms but drop the quadratic one. The coefficient terms for the higher order terms become very small. It is also possible that the coefficient estimation may be incorrect due to ill conditioning of the data matrix which is used obtain the model coefficients. Some authors recommend appropriate rescaling of the polynomial terms (such as subtracting the mean etc) to avoid such problems.

The use of polynomials greater than second-order is discouraged. Higher-order polynomials are known to be extremely volatile; they have high variance, and make bad predictions. If you need a more flexible model, then it is generally better to use some sort of smoother than a high-order polynomial.

In fact, a popular method of smoothing is known as “local polynomial fitting”, or **spline smoothing**. Local polynomials are sometimes preferred to a single polynomial regression model for the whole data set. An example based on the `pinetree` data is shown in Figure 4.3 which uses the `bs()` function from the *splines* package.

```
pinetree |>
  ggplot() +
  aes(First, Top) +
  geom_point() +
  geom_smooth(method = lm,
              formula = y ~ splines::bs(x, 3))
```

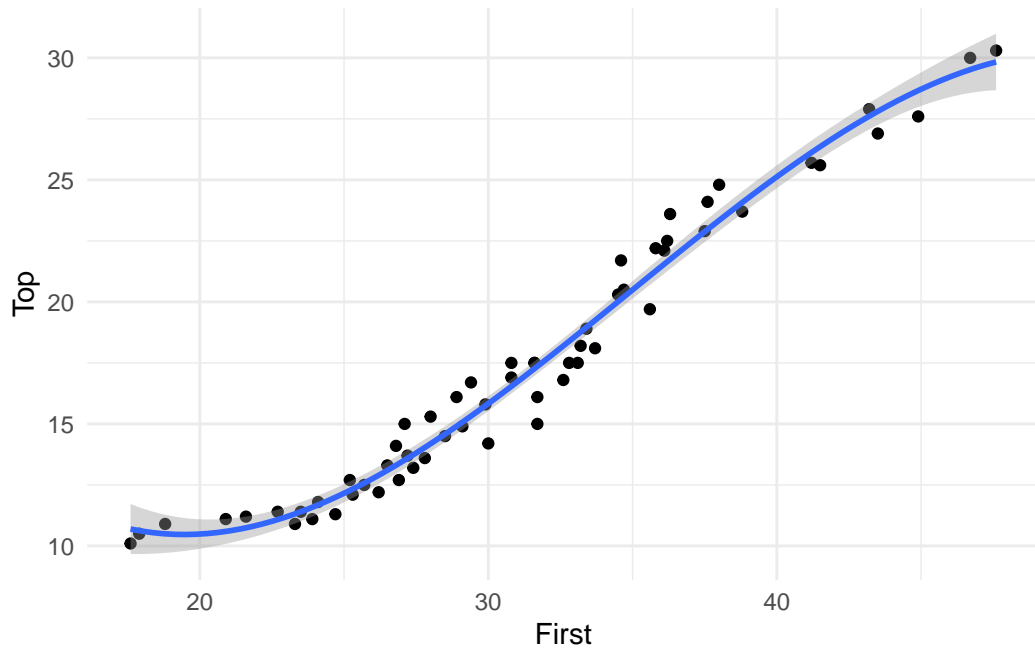


Figure 4.3: Polynomial Spline smoothing

5 Model structure and other issues

The difficult task in statistical modelling is the assessment of the underlying model structure or alternatively knowing the true form of the relationship. For example, the true relationship between Y and X variables may be nonlinear. If we incorrectly assume a multiple linear relationship instead, a good model may not result. The interaction between the explanatory variables is also important and this topic is covered in a different section. We may also fit a robust linear model in order to validate the ordinary least squares fit. For the horses heart data, OUTERSYS and EXTDIA were short-listed as the predictors of WEIGHT using the AIC criterion. This least squares regression model can be compared to the robust versions as shown in Figure 5.1.

```
hh_lm <- lm(WEIGHT ~ OUTERSYS + EXTDIA, data=horsehearts)

hh_rlm <- MASS::rlm(WEIGHT ~ OUTERSYS + EXTDIA, data=horsehearts)

hh_lmrob <- robustbase::lmrob(WEIGHT ~ OUTERSYS + EXTDIA, data=horsehearts)

horsehearts |>
  gather_predictions(hh_lm, hh_rlm, hh_lmrob) %>%
  ggplot(aes(x=WEIGHT, y=pred, colour=model)) +
  geom_point() +
  geom_abline(slope=1, intercept = 0) +
  ylab("Predicted WEIGHT") + theme_minimal() +
  ggtitle("Comparison of model predictions")
```

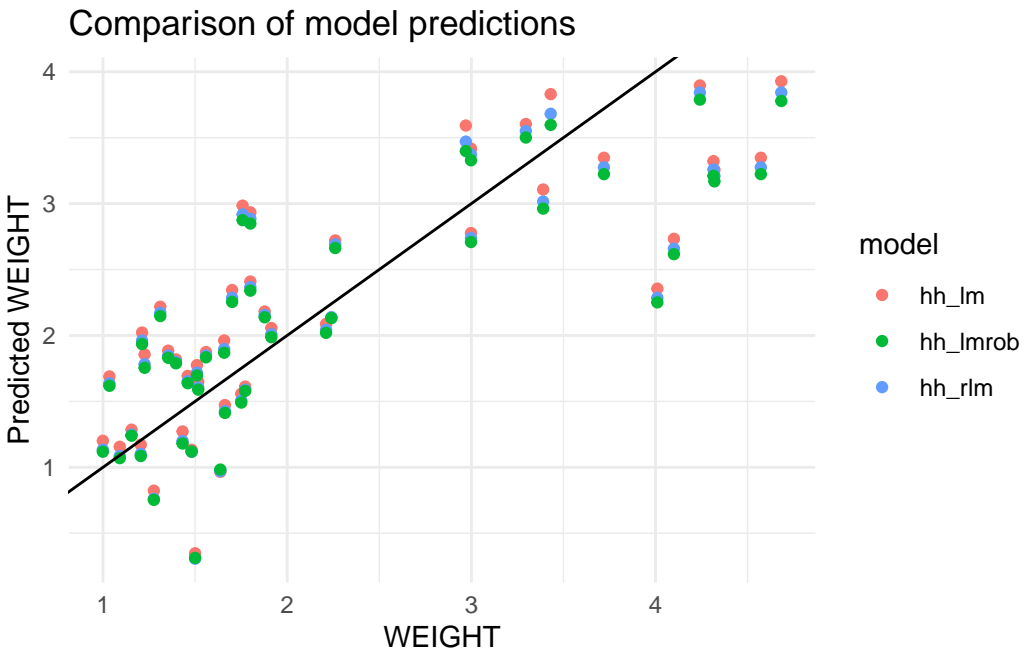


Figure 5.1: Comparison of predictions

Figure 5.1 shows that the scatter plot of predicted versus actual Y values for the three fitted models which confirms that these models perform very similarly. We may also extract measures such as mean absolute percentage error (MAPE) or residual SD or root mean square error (RMSE) for the three models using `modelr` package; see the code shown below:

```
list(hh_lm = hh_lm,
     hh_rlm = hh_rlm,
     hh_lmrob = hh_lmrob) |>
enframe("method", "fit") |>
mutate(
  MAPE = map_dbl(fit, \((x)\){mape(x, horsehearts)}),
  RMSE = map_dbl(fit, \((x)\){rmse(x, horsehearts)})
)

# A tibble: 3 x 4
  method   fit      MAPE    RMSE
  <chr>   <list>  <dbl>  <dbl>
1 hh_lm   <lm>    0.255  0.648
2 hh_rlm  <rlm>   0.247  0.651
3 hh_lmrob <lmrob> 0.243  0.655
```

Measures such as AIC or BIC need corrections when the normality assumption does not hold but the above summary measures do not require such a distributional assumption to hold.

If a large dataset is in hand, a part of the data (training data) can be used to fit the model and then we can see how well the fitted model works for the remaining data.

6 Summary

Regression methods are the most commonly used of statistics techniques. The main aim is to fit a model by least squares to explain the variation in the response variable Y by using one or more explanatory variables X_1, X_2, \dots, X_k . The correlation coefficient r_{xy} measures the strength of the linear relationship between Y and X ; R^2 measures the strength of the linear relationship between Y and X_1, X_2, \dots, X_k . When $k=1$, then $r_{xy}^2 = R^2$. Scatter plots and correlation coefficients provide important clues to the inter-relationships between the variables.

In building up a model by adding new variables, the correlation (or overlap) with y is important but the correlations between a new explanatory variable and each of the existing explanatory variables also determine how effective the addition of the variable will be.

Stepwise regression procedures identify potentially good regression models by repeatedly comparing an existing model with other models in which an explanatory variable has been either deleted or added, using some criterion such as significance of the deleted or added term (as measured by the p -value of the relevant t or F statistic) or the AIC of the model. Polynomial regression models employ the square, cube etc terms of the original variables as additional predictors.

When at least two explanatory variables are highly correlated, we have multicollinear data. The effect is that the variance of least square estimators will be inflated rendering the coefficients insignificant and hence we may need to discard one or more of the highly correlated variables.

EDA plots of residuals help to answer the question as to whether the fit is good or whether a transformation may help or whether other variables (including square, cubic etc) should be added. If residuals are plotted against fitted Y or X variables no discernible pattern should be observed. Estimated regression coefficients may be affected by leverage points, and hence influence diagnostics are performed.

7 What you should know

By the end of this chapter you should be able to:

- Fit and display a multiple regression (2 or more predictor variables).
- Check the assumptions of a multiple regression using residual diagnostic plots, tests for assumptions, and examine multicollinearity.
- Measure variability explained by the model and predictors.
- Use a fitted regression to predict new data.
- Explain the significance of your model and interpret findings in context of the data and hypotheses.
- Describe model comparison/selection and polynomial terms.