Data Science for Biological, Medical and Health Research: Notes for 432

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Introduction

These Notes provide a series of examples using R to work through issues that are likely to come up in PQHS/CRSP/MPHP 432.

While these Notes share some of the features of a textbook, they are neither comprehensive nor completely original. The main purpose is to give students in 432 a set of common materials on which to draw during the course. In class, we will sometimes:

- reiterate points made in this document,
- amplify what is here,
- simplify the presentation of things done here,
- use new examples to show some of the same techniques,
- refer to issues not mentioned in this document,

but what we don't (always) do is follow these notes very precisely. We assume instead that you will read the materials and try to learn from them, just as you will attend classes and try to learn from them. We welcome feedback of all kinds on this document or anything else. Just email us at 431-help at case dot edu, or submit a pull request. Note that we still use 431-help even though we're now in 432.

What you will mostly find are brief explanations of a key idea or summary, accompanied (most of the time) by R code and a demonstration of the results of applying that code.

Everything you see here is available to you as HTML or PDF. You will also have access to the R Markdown files, which contain the code which generates everything in the document, including all of the R results. We will demonstrate the use of R Markdown (this document is generated with the additional help of an R package called bookdown) and R Studio (the "program" which we use to interface with the R language) in class.

To download the data and R code related to these notes, visit the Data and Code section of the 432 course website.

R Packages used in these notes

Here, we'll load in the packages used in these notes.

```
library(tableone)
library(skimr)
library(ggridges)
library(magrittr)
library(arm)
library(rms)
library(leaps)
library(leaps)
library(Epi)
library(pROC)
library(ROCR)
library(simputation)
library(modelr)
library(broom)
library(tidyverse)
```

Data used in these notes

Here, we'll load in the data sets used in these notes.

```
fakestroke <- read.csv("data/fakestroke.csv") %>% tbl_df
bloodbrain <- read.csv("data/bloodbrain.csv") %>% tbl_df
smartcle1 <- read.csv("data/smartcle1.csv") %>% tbl_df
bonding <- read.csv("data/bonding.csv") %>% tbl_df
cortisol <- read.csv("data/cortisol.csv") %>% tbl_df
emphysema <- read.csv("data/emphysema.csv") %>% tbl_df
prost <- read.csv("data/prost.csv") %>% tbl_df
pollution <- read.csv("data/pollution.csv") %>% tbl_df
resect <- read.csv("data/resect.csv") %>% tbl_df
```

Special Functions used in these notes

Chapter 1

Building Table 1

Many scientific articles involve direct comparison of results from various exposures, perhaps treatments. In 431, we studied numerous methods, including various sorts of hypothesis tests, confidence intervals, and descriptive summaries, which can help us to understand and compare outcomes in such a setting. One common approach is to present what's often called Table 1. Table 1 provides a summary of the characteristics of a sample, or of groups of samples, which is most commonly used to help understand the nature of the data being compared.

1.1 Two examples from the New England Journal of Medicine

1.1.1 A simple Table 1

Table 1 is especially common in the context of clinical research. Consider the excerpt below, from a January 2015 article in the New England Journal of Medicine (Tolaney et al., 2015).

Table 1. Baseline Characteristics of the Patients.*				
Characteristic	Patients (N=406)			
	no. (%)			
Age group				
<50 yr	132 (32.5)			
50–59 yr	137 (33.7)			
60–69 yr	96 (23.6)			
≥70 yr	41 (10.1)			
Sex				
Female	405 (99.8)			
Male	1 (0.2)			
Race†				
White	351 (86.5)			
Black	28 (6.9)			
Asian	11 (2.7)			
Other	16 (3.9)			

This (partial) table reports baseline characteristics on age group, sex and race, describing 406 patients with

HER2-positive¹ invasive breast cancer that began the protocol therapy. Age, sex and race (along with severity of illness) are the most commonly identified characteristics in a Table 1.

In addition to the measures shown in this excerpt, the full Table also includes detailed information on the primary tumor for each patient, including its size, nodal status and histologic grade. Footnotes tell us that the percentages shown are subject to rounding, and may not total 100, and that the race information was self-reported.

1.1.2 A group comparison

A more typical Table 1 involves a group comparison, for example in this excerpt from Roy et al. (2008). This Table 1 describes a multi-center randomized clinical trial comparing two different approaches to caring for patients with heart failure and atrial fibrillation².

Table 1. Baseline Characteristics of the Patients.*					
Variable	Rhythm-Control Group (N = 682)	Rate-Control Group (N = 694)			
Male sex (%)	78	85			
Age (yr)	66±11	67±11			
Body-mass index†	27.8±5.4	28.0±5.1			
Nonwhite race (%)‡	16	13			
NYHA class III or IV (%)					
At baseline	32	31			
During previous 6 mo	76	76			
Predominant cardiac diagnosis (%)∫					
Coronary artery disease	48	48			
Valvular heart disease	5	5			
Nonischemic cardiomyopathy	36	39			
Congenital heart disease	1	1			
Hypertensive heart disease	10	7			

The article provides percentages, means and standard deviations across groups, but note that it does not provide p values for the comparison of baseline characteristics. This is a common feature of NEJM reports on randomized clinical trials, where we anticipate that the two groups will be well matched at baseline. Note that the patients in this study were *randomly* assigned to either the rhythm-control group or to the rate-control group, using blocked randomizations stratified by study center.

1.2 The MR CLEAN trial

Berkhemer et al. (2015) reported on the MR CLEAN trial, involving 500 patients with acute ischemic stroke caused by a proximal intracranial arterial occlusion. The trial was conducted at 16 medical centers in the Netherlands, where 233 were randomly assigned to the intervention (intraarterial treatment plus usual care) and 267 to control (usual care alone.) The primary outcome was the modified Rankin scale score at 90 days; this categorical scale measures functional outcome, with scores ranging from 0 (no symptoms) to 6 (death). The fundamental conclusion of Berkhemer et al. (2015) was that in patients with acute ischemic stroke

¹HER2 = human epidermal growth factor receptor type 2. Over-expression of this occurs in 15-20% of invasive breast cancers, and has been associated with poor outcomes.

²The complete Table 1 appears on pages 2668-2669 of Roy et al. (2008), but I have only reproduced the first page and the footnote in this excerpt.

caused by a proximal intracranial occlusion of the anterior circulation, intraarterial treatment administered within 6 hours after stroke onset was effective and safe.

Here's the Table 1 from Berkhemer et al. (2015).

Characteristic	Intervention (N = 233)	Control (N = 267)
Age — yr		
Median	65.8	65.7
Interquartile range	54.5-76.0	55.5-76.4
Male sex — no. (%)	135 (57.9)	157 (58.8)
NIHSS score†		
Median (interquartile range)	17 (14–21)	18 (14-22)
Range	3-30	4-38
Location of stroke in left hemisphere — no. (%)	116 (49.8)	153 (57.3)
History of ischemic stroke — no. (%)	29 (12.4)	25 (9.4)
Atrial fibrillation — no. (%)	66 (28.3)	69 (25.8)
Diabetes mellitus — no. (%)	34 (14.6)	34 (12.7)
Prestroke modified Rankin scale score — no. (%)‡		
0	190 (81.5)	214 (80.1)
1	21 (9.0)	29 (10.9)
2	12 (5.2)	13 (4.9)
>2	10 (4.3)	11 (4.1)
Systolic blood pressure — mm Hg∫	146±26.0	145±24.4
Treatment with IV alteplase — no. (%)	203 (87.1)	242 (90.6)
Time from stroke onset to start of IV alteplase — min		
Median	85	87
Interquartile range	67-110	65-116
ASPECTS — median (interquartile range)¶	9 (7-10)	9 (8-10)
Intracranial arterial occlusion — no./total no. (%)		
Intracranial ICA	1/233 (0.4)	3/266 (1.1)
ICA with involvement of the M1 middle cerebral artery segment	59/233 (25.3)	75/266 (28.2)
M1 middle cerebral artery segment	154/233 (66.1)	165/266 (62.0)
M2 middle cerebral artery segment	18/233 (7.7)	21/266 (7.9)
A1 or A2 anterior cerebral artery segment	1/233 (0.4)	2/266 (0.8)
Extracranial ICA occlusion — no./total no. (%) **	75/233 (32.2)	70/266 (26.3)
Time from stroke onset to randomization — min††		
Median	204	196
Interquartile range	152-251	149–266
Time from stroke onset to groin puncture — min		
Median	260	NA
Interquartile range	210-313	

The Table was accompanied by the following notes.

- * The intervention group was assigned to intraarterial treatment plus usual care, and the control group was assigned to usual care alone. Plus-minus values are means ±SD. ICA denotes internal carotid artery, IV intravenous, and NA not applicable.
- † Scores on the National Institutes of Health Stroke Scale (NIHSS) range from 0 to 42, with higher scores indicating more severe neurologic deficits. The NIHSS is a 15-item scale, and values for 30 of the 7500 items were missing (0.4%). The highest number of missing items for a single patient was 6.
- Scores on the modified Rankin scale of functional disability range from 0 (no symptoms) to 6 (death). A score of 2 or less indicates functional independence.
- Data on systolic blood pressure at baseline were missing for one patient assigned to the control group.
- The Alberta Stroke Program Early Computed Tomography Score (ASPECTS) is a measure of the extent of stroke. Scores ranges from 0 to 10, with higher scores indicating fewer early ischemic changes. Scores were not available for four patients assigned to the control group: noncontrast computed tomography was not performed in one patient, and three patients had strokes in the territory of the anterior cerebral artery.
- Vessel imaging was not performed in one patient in the control group, so the level of occlusion was not known.
- ** Extracranial ICA occlusions were reported by local investigators.
- †† Data were missing for two patients in the intervention group.

1.3 Simulated fakestroke data

Consider the simulated data, available on the Data and Code page of our course website in the fakestroke.csv file, which I built to let us mirror the Table 1 for MR CLEAN (Berkhemer et al., 2015). The fakestroke.csv file contains the following 18 variables for 500 patients.

Variable	Description
studyid	Study ID # (z001 through z500)
trt	Treatment group (Intervention or Control)
age	Age in years
sex	Male or Female
nihss	NIH Stroke Scale Score (can range from 0-42; higher scores
	indicate more severe neurological deficits)
location	Stroke Location - Left or Right Hemisphere
hx.isch	History of Ischemic Stroke (Yes/No)
afib	Atrial Fibrillation $(1 = Yes, 0 = No)$
dm	Diabetes Mellitus $(1 = Yes, 0 = No)$
mrankin	Pre-stroke modified Rankin scale score $(0, 1, 2 \text{ or } > 2)$
	indicating functional disability - complete range is 0 (no
	symptoms) to 6 (death)
sbp	Systolic blood pressure, in mm Hg
iv.altep	- ' ' '
time.iv	Time from stroke onset to start of IV alteplase (minutes) if
	iv.altep=Yes
aspects	Alberta Stroke Program Early Computed Tomography
	score, which measures extent of stroke from 0 - 10 ; higher
	scores indicate fewer early ischemic changes
ia.occlus	Intracranial arterial occlusion, based on vessel imaging -
	five categories ³
extra.ica	Extracranial ICA occlusion $(1 = Yes, 0 = No)$
time.rand	Time from stroke onset to study randomization, in minutes
time.punc	Time from stroke onset to groin puncture, in minutes (only
	if Intervention)

Here's a quick look at the simulated data in fakestroke.

³The five categories are Intracranial ICA, ICA with involvement of the M1 middle cerebral artery segment, M1 middle cerebral artery segment, M2 middle cerebral artery segment, A1 or A2 anterior cerebral artery segment

fakestroke

```
# A tibble: 500 x 18
   studyid trt
                      age sex
                                nihss location hx.isch afib
                                                                 dm mrankin
           <fct>
   <fct>
                    <dbl> <fct> <int> <fct>
                                                <fct>
                                                        <int> <int> <fct>
 1 z001
           Control 53.0 Male
                                                            0
                                                                  0 2
                                   21 Right
                                                No
 2 z002
                                                                  0 0
           Interve~ 51.0 Male
                                   23 Left
                                                No
                                                            1
                     68.0 Fema~
 3 z003
                                                            0
                                                                  0 0
           Control
                                   11 Right
                                                No
 4 z004
           Control
                     28.0 Male
                                   22 Left
                                                No
                                                            0
                                                                  0 0
                                                            0
 5 z005
           Control
                     91.0 Male
                                   24 Right
                                                No
                                                                  0 0
 6 z006
           Control
                     34.0 Fema~
                                   18 Left
                                                No
                                                                  0 2
 7 z007
                                   25 Right
                                                            0
                                                                  0 0
           Interve~ 75.0 Male
                                                No
 8 z008
           Control
                     89.0 Fema~
                                   18 Right
                                                No
                                                            0
                                                                  0 0
9 z009
           Control
                     75.0 Male
                                   25 Left
                                                No
                                                            1
                                                                  0 2
10 z010
           Interve~ 26.0 Fema~
                                   27 Right
                                                            0
                                                                  0 0
                                                No
# ... with 490 more rows, and 8 more variables: sbp <int>, iv.altep <fct>,
   time.iv <int>, aspects <int>, ia.occlus <fct>, extra.ica <int>,
   time.rand <int>, time.punc <int>
```

1.4 Building Table 1 for fakestroke: Attempt 1

Our goal, then, is to take the data in fakestroke.csv and use it to generate a Table 1 for the study that compares the 233 patients in the Intervention group to the 267 patients in the Control group, on all of the other variables (except study ID #) available. I'll use the tableone package of functions available in R to help me complete this task. We'll make a first attempt, using the CreateTableOne function in the tableone package. To use the function, we'll need to specify:

- the vars or variables we want to place in the rows of our Table 1 (which will include just about everything in the fakestroke data except the studyid code and the trt variable for which we have other plans, and the time.punc which applies only to subjects in the Intervention group.)
 - A useful trick here is to use the dput function, specifically something like dput (names (fakestroke)) can be used to generate a list of all of the variables included in the fakestroke tibble, and then this can be copied and pasted into the vars specification, saving some typing.
- the strata which indicates the levels want to use in the columns of our Table 1 (for us, that's trt)

Stratified by trt Control Intervention test 267 233 age (mean (sd)) 65.38 (16.10) 63.93 (18.09) 0.343 sex = Male (%) 157 (58.8) 135 (57.9) 0.917 nihss (mean (sd)) 18.08 (4.32) 17.97 (5.04) 0.787 117 (50.2) location = Right (%) 114 (42.7) 0.111

hx.isch = Yes (%)	25	(9.4)	29	(12.4)	0.335
afib (mean (sd))	0.26	(0.44)	0.28	(0.45)	0.534
dm (mean (sd))	0.13	(0.33)	0.12	(0.33)	0.923
mrankin (%)					0.922
> 2	11	(4.1)	10	(4.3)	
0	214	(80.1)	190	(81.5)	
1	29	(10.9)	21	(9.0)	
2	13	(4.9)	12	(5.2)	
sbp (mean (sd))	145.00	(24.40)	146.03	(26.00)	0.647
iv.altep = Yes (%)	242	(90.6)	203	(87.1)	0.267
time.iv (mean (sd))	87.96	(26.01)	98.22	(45.48)	0.003
aspects (mean (sd))	8.65	(1.47)	8.35	(1.64)	0.033
ia.occlus (%)					0.795
A1 or A2	2	(0.8)	1	(0.4)	
ICA with M1	75	(28.2)	59	(25.3)	
Intracranial ICA	3	(1.1)	1	(0.4)	
M1	165	(62.0)	154	(66.1)	
M2	21	(7.9)	18	(7.7)	
extra.ica (mean (sd))	0.26	(0.44)	0.32	(0.47)	0.150
time.rand (mean (sd))	213.88	(70.29)	202.51	(57.33)	0.051

1.4.1 Some of this is very useful, and other parts need to be fixed.

- 1. The 1/0 variables (afib, dm, extra.ica) might be better if they were treated as the factors they are, and reported as the Yes/No variables are reported, with counts and percentages rather than with means and standard deviations.
- 2. In some cases, we may prefer to re-order the levels of the categorical (factor) variables, particularly the mrankin variable, but also the ia.occlus variable. It would also be more typical to put the Intervention group to the left and the Control group to the right, so we may need to adjust our trt variable's levels accordingly.
- 3. For each of the quantitative variables (age, nihss, sbp, time.iv, aspects, extra.ica, time.rand and time.punc) we should make a decision whether a summary with mean and standard deviation is appropriate, or whether we should instead summarize with, say, the median and quartiles. A mean and standard deviation really only yields an appropriate summary when the data are least approximately Normally distributed. This will make the p values a bit more reasonable, too. The test column in the first attempt will soon have something useful to tell us.
- 4. If we'd left in the time.punc variable, we'd get some warnings, having to do with the fact that time.punc is only relevant to patients in the Intervention group.

1.4.2 fakestroke Cleaning Up Categorical Variables

Let's specify each of the categorical variables as categorical explicitly. This helps the CreateTableOne function treat them appropriately, and display them with counts and percentages. This includes all of the 1/0, Yes/No and multi-categorical variables.

Then we simply add a factorVars = fs.factorvars call to the CreateTableOne function.

We also want to re-order some of those categorical variables, so that the levels are more useful to us. Specifically, we want to:

- place Intervention before Control in the trt variable,
- reorder the mrankin scale as 0, 1, 2, > 2, and

• rearrange the ia.occlus variable to the order⁴ presented in Berkhemer et al. (2015).

To accomplish this, we'll use the fct_relevel function from the forcats package (loaded with the rest of the core tidyverse packages) to reorder our levels manually.

1.5 fakestroke Table 1: Attempt 2

Stratified by trt						
	Interve	ention	Control	L	p	test
n	233		267			
age (mean (sd))	63.93	(18.09)	65.38	(16.10)	0.343	
sex = Male (%)		(57.9)		(58.8)		
nihss (mean (sd))	17.97	(5.04)	18.08	(4.32)	0.787	
<pre>location = Right (%)</pre>	117	(50.2)	114	(42.7)	0.111	
hx.isch = Yes (%)	29	(12.4)	25	(9.4)	0.335	
afib = 1 (%)	66	(28.3)	69	(25.8)	0.601	
dm = 1 (%)	29	(12.4)	34	(12.7)	1.000	
mrankin (%)					0.922	
0	190	(81.5)	214	(80.1)		
1	21	(9.0)	29	(10.9)		
2	12	(5.2)	13	(4.9)		
> 2	10	(4.3)	11	(4.1)		
sbp (mean (sd))	146.03	(26.00)	145.00	(24.40)	0.647	
iv.altep = Yes (%)	203	(87.1)	242	(90.6)	0.267	
time.iv (mean (sd))	98.22	(45.48)	87.96	(26.01)	0.003	
aspects (mean (sd))	8.35	(1.64)	8.65	(1.47)	0.033	
ia.occlus (%)					0.795	
Intracranial ICA	1	(0.4)	3	(1.1)		
ICA with M1	59	(25.3)	75	(28.2)		
M1	154	(66.1)	165	(62.0)		
M2	18	(7.7)	21	(7.9)		
A1 or A2	1	(0.4)	2	(0.8)		
extra.ica = 1 (%)	75	(32.2)	70	(26.3)	0.179	
<pre>time.rand (mean (sd))</pre>	202.51	(57.33)	213.88	(70.29)	0.051	

The categorical data presentation looks much improved.

⁴We might also have considered reordering the ia.occlus factor by its frequency, using the fct_infreq function

1.5.1 What summaries should we show?

Now, we'll move on to the issue of making a decision about what type of summary to show for the quantitative variables. Since the fakestroke data are just simulated and only match the summary statistics of the original results, not the details, we'll adopt the decisions made by Berkhemer et al. (2015), which were to use medians and interquartile ranges to summarize the distributions of all of the continuous variables except systolic blood pressure.

- Specifying certain quantitative variables as *non-normal* causes R to show them with medians and the 25th and 75th percentiles, rather than means and standard deviations, and also causes those variables to be tested using non-parametric tests, like the Wilcoxon signed rank test, rather than the t test. The test column indicates this with the word nonnorm.
 - In real data situations, what should we do? The answer is to look at the data. I would not make the decision as to which approach to take without first plotting (perhaps in a histogram or a Normal Q-Q plot) the observed distributions in each of the two samples, so that I could make a sound decision about whether Normality was a reasonable assumption. If the means and medians are meaningfully different from each other, this is especially important.
 - To be honest, though, if the variable in question is a relatively unimportant covariate and the
 p values for the two approaches are nearly the same, I'd say that further investigation is rarely
 important,
- Specifying *exact* tests for certain categorical variables (we'll try this for the location and mrankin variables) can be done, and these changes will be noted in the test column, as well.
 - In real data situations, I would rarely be concerned about this issue, and often choose Pearson (approximate) options across the board. This is reasonable so long as the number of subjects falling in each category is reasonably large, say above 10. If not, then an exact test may be a tiny improvement.
 - Paraphrasing Rosenbaum (2017), having an exact rather than an approximate test result is about as valuable as having a nice crease in your trousers.

To finish our Table 1, then, we need to specify which variables should be treated as non-Normal in the print statement - notice that we don't need to redo the CreateTableOne for this change.

	Stratif	ied by trt		
	Interve	ention	Control	L
n	233		267	
age (median [IQR])	65.80	[54.50, 76.00]	65.70	[55.75, 76.20]
sex = Male (%)	135	(57.9)	157	(58.8)
nihss (median [IQR])	17.00	[14.00, 21.00]	18.00	[14.00, 22.00]
location = Right (%)	117	(50.2)	114	(42.7)
hx.isch = Yes (%)	29	(12.4)	25	(9.4)
afib = 1 (%)	66	(28.3)	69	(25.8)
dm = 1 (%)	29	(12.4)	34	(12.7)
mrankin (%)				
0	190	(81.5)	214	(80.1)
1	21	(9.0)	29	(10.9)
2	12	(5.2)	13	(4.9)
> 2	10	(4.3)	11	(4.1)
sbp (mean (sd))	146.03	(26.00)	145.00	(24.40)
<pre>iv.altep = Yes (%)</pre>	203	(87.1)	242	(90.6)
time.iv (median [IQR])	85.00	[67.00, 110.00]	87.00	[65.00, 116.00]
aspects (median [IQR])	9.00	[7.00, 10.00]	9.00	[8.00, 10.00]
ia.occlus (%)				

```
Intracranial ICA
                               1 (0.4)
                                                        3 (1.1)
   ICA with M1
                              59 (25.3)
                                                       75 (28.2)
   M1
                             154 (66.1)
                                                      165 (62.0)
   M2
                              18 (7.7)
                                                       21 (7.9)
   A1 or A2
                               1 (0.4)
                                                        2 (0.8)
extra.ica = 1 (\%)
                              75 (32.2)
                                                       70 (26.3)
time.rand (median [IQR]) 204.00 [152.00, 249.50] 196.00 [149.00, 266.00]
                         Stratified by trt
                                 test
n
age (median [IQR])
                           0.579 nonnorm
                           0.917
sex = Male (%)
nihss (median [IQR])
                           0.453 nonnorm
location = Right (%)
                           0.106 exact
hx.isch = Yes (%)
                           0.335
afib = 1 (\%)
                           0.601
dm = 1 (\%)
                           1.000
mrankin (%)
                           0.917 exact
   0
   1
   2
   > 2
sbp (mean (sd))
                           0.647
iv.altep = Yes (%)
                           0.267
time.iv (median [IQR])
                           0.596 nonnorm
aspects (median [IQR])
                           0.075 nonnorm
ia.occlus (%)
                           0.795
   Intracranial ICA
   ICA with M1
   M1
   M2
   A1 or A2
extra.ica = 1 (%)
                           0.179
time.rand (median [IQR])
                          0.251 nonnorm
```

1.6 Obtaining a more detailed Summary

If this was a real data set, we'd want to get a more detailed description of the data to make decisions about things like potentially collapsing categories of a variable, or whether or not a normal distribution was useful for a particular continuous variable, etc. You can do this with the $\mathtt{summary}$ command applied to a created Table 1, which shows, among other things, the effect of changing from normal to non-normal p values for continuous variables, and from approximate to "exact" p values for categorical factors.

Again, as noted above, in a real data situation, we'd want to plot the quantitative variables (within each group) to make a smart decision about whether a t test or Wilcoxon approach is more appropriate.

Note in the summary below that we have some missing values here. Often, we'll present this information within the Table 1, as well.

```
summary(att2)
```

trt:	Interver	ntion
------	----------	-------

	n	miss	p.miss	${\tt mean}$	sd	${\tt median}$	p25	p75	\min	${\tt max}$	skew	kurt
age	233	0	0.0	64	18	66	54	76	23	96	-0.34	-0.52
nihss	233	0	0.0	18	5	17	14	21	10	28	0.48	-0.74
sbp	233	0	0.0	146	26	146	129	164	78	214	-0.07	-0.22
time.iv	233	30	12.9	98	45	85	67	110	42	218	1.03	0.08
aspects	233	0	0.0	8	2	9	7	10	5	10	-0.56	-0.98
time.rand	233	2	0.9	203	57	204	152	250	100	300	0.01	-1.16

trt: Control

	n	miss	p.miss	mean	sd	${\tt median}$	p25	p75	${\tt min}$	max	skew	kurt
age	267	0	0.0	65	16	66	56	76	24	94	-0.296	-0.28
nihss	267	0	0.0	18	4	18	14	22	11	25	0.017	-1.24
sbp	267	1	0.4	145	24	145	128	161	82	231	0.156	0.08
time.iv	267	25	9.4	88	26	87	65	116	44	130	0.001	-1.32
aspects	267	4	1.5	9	1	9	8	10	5	10	-1.071	0.36
time.rand	267	0	0.0	214	70	196	149	266	120	360	0.508	-0.93

p-values

	pNormal	pNonNormal
age	0.342813660	0.57856976
nihss	0.787487252	0.45311695
sbp	0.647157646	0.51346132
time.iv	0.003073372	0.59641104
aspects	0.032662901	0.07464683
$\verb time.rand $	0.050803672	0.25134327

Standardize mean differences

1 vs 2 age 0.08478764 nihss 0.02405390 sbp 0.04100833 time.iv 0.27691223 aspects 0.19210662

time.rand 0.17720957

Summary of categorical variables

trt: Intervention

var	n	miss	p.miss	level	freq	percent	cum.percent
sex	233	0	0.0	Female	98	42.1	42.1
				Male	135	57.9	100.0
location	233	0	0.0	Left	116	49.8	49.8
				Right	117	50.2	100.0
hx.isch	233	0	0.0	No	204	87.6	87.6
				Yes	29	12.4	100.0
afib	233	0	0.0	0	167	71.7	71.7
				1	66	28.3	100.0

dm	233	0	0.0	0 1	204 29		87.6 100.0
mrankin	233	0	0.0	0 1 2		9.0 5.2	81.5 90.6 95.7
iv.altep	233	0	0.0	No	30 203	12.9	100.0 12.9 100.0
ia.occlus	233	0	0.0	Intracranial ICA ICA with M1	1 59 154	0.4 25.3 66.1	0.4
extra.ica	233	0	0.0		1 158 75	67.8	67.8 100.0
trt: Contro	 ol						
var			p.miss				cum.percent
sex	267	0	0.0	Female Male	157		
location	267	0	0.0	Left Right			
hx.isch	267	0	0.0	No Yes	242 25		90.6 100.0
afib	267	0	0.0	0 1			74.2 100.0
dm	267	0	0.0	0 1	233 34	87.3 12.7	87.3 100.0
mrankin	267	0	0.0	0 1 2 > 2	214 29 13 11	80.1 10.9 4.9 4.1	80.1 91.0 95.9 100.0
iv.altep	267	0	0.0	No Yes	25 242	9.4 90.6	9.4 100.0
ia.occlus	267	1	0.4	Intracranial ICA ICA with M1 M1 M2 A1 or A2	3 75 165 21 2	1.1 28.2 62.0 7.9 0.8	1.1 29.3 91.4 99.2 100.0
extra.ica	267	1	0.4	0 1	196 70	73.7 26.3	73.7 100.0

```
p-values
           pApprox
                      pExact
         0.9171387 0.8561188
location 0.1113553 0.1056020
hx.isch 0.3352617 0.3124683
afib
         0.6009691 0.5460206
dm
         1.0000000 1.0000000
mrankin 0.9224798 0.9173657
iv.altep 0.2674968 0.2518374
ia.occlus 0.7945580 0.8189090
extra.ica 0.1793385 0.1667574
Standardize mean differences
               1 vs 2
         0.017479025
sex
location 0.151168444
hx.isch 0.099032275
         0.055906317
         0.008673478
mrankin 0.062543164
iv.altep 0.111897009
ia.occlus 0.117394890
extra.ica 0.129370206
```

In this case, I have simulated the data to mirror the results in the published Table 1 for this study. In no way have I captured the full range of the real data, or any of the relationships in that data, so it's more important here to see what's available in the analysis, rather than to interpret it closely in the clinical context.

1.7 Exporting the Completed Table 1 from R to Excel or Word

Once you've built the table and are generally satisfied with it, you'll probably want to be able to drop it into Excel or Word for final cleanup.

1.7.1 Approach A: Save and open in Excel

One option is to save the Table 1 to a .csv file within our data subfolder (note that the data folder must already exist), which you can then open directly in Excel. This is the approach I generally use. Note the addition of some quote, noSpaces and printToggle selections here.

When I then open the fs-table1.csv file in Excel, it looks like this:

4	Α	В	С	D	E
1		Intervention	Control	р	test
2	n	233	267		
3	age (median [IQR])	65.80 [54.50, 76.00]	65.70 [55.75, 76.20]	0.579	nonnorm
4	sex = Male (%)	135 (57.9)	157 (58.8)	0.917	
5	nihss (median [IQR])	17.00 [14.00, 21.00]	18.00 [14.00, 22.00]	0.453	nonnorm
6	location = Right (%)	117 (50.2)	114 (42.7)	0.111	
7	hx.isch = Yes (%)	29 (12.4)	25 (9.4)	0.335	
8	afib = 1 (%)	66 (28.3)	69 (25.8)	0.601	
9	dm = 1 (%)	29 (12.4)	34 (12.7)	1	
10	mrankin (%)			0.922	
11	0	190 (81.5)	214 (80.1)		
12	1	21 (9.0)	29 (10.9)		
13	2	12 (5.2)	13 (4.9)		
14	>2	10 (4.3)	11 (4.1)		
15	sbp (mean (sd))	146.03 (26.00)	145.00 (24.40)	0.647	
16	iv.altep = Yes (%)	203 (87.1)	242 (90.6)	0.267	
17	time.iv (median [IQR])	85.00 [67.00, 110.00]	87.00 [65.00, 116.00]	0.596	nonnorm
18	aspects (median [IQR])	9.00 [7.00, 10.00]	9.00 [8.00, 10.00]	0.075	nonnorm
19	ia.occlus (%)			0.795	
20	Intracranial ICA	1 (0.4)	3 (1.1)		
21	ICA with M1	59 (25.3)	75 (28.2)		
22	M1	154 (66.1)	165 (62.0)		
23	M2	18 (7.7)	21 (7.9)		
24	A1 or A2	1 (0.4)	2 (0.8)		
25	extra.ica = 1 (%)	75 (32.2)	70 (26.3)	0.179	
26	time.rand (median [IQR])	204.00 [152.00, 249.50]	196.00 [149.00, 266.00]	0.251	nonnorm
27	time.punc (median [IQR])	260.00 [212.00, 313.00]	NA [NA, NA]	NA	nonnorm
28					

And from here, I can either drop it directly into Word, or present it as is, or start tweaking it to meet formatting needs.

1.7.2 Approach B: Produce the Table so you can cut and paste it

This will look like a mess by itself, but if you:

- 1. copy and paste that mess into Excel
- 2. select Text to Columns from the Data menu
- 3. select Delimited, then Space and select Treat consecutive delimiters as one

you should get something usable again.

Or, in Word,

1. insert the text

- 2. select the text with your mouse
- 3. select Insert ... Table ... Convert Text to Table
- 4. place a quotation mark in the "Other" area under Separate text at ...

After dropping blank columns, the result looks pretty good.

1.8 A Controlled Biological Experiment - The Blood-Brain Barrier

My source for the data and the following explanatory paragraph is page 307 from Ramsey and Schafer (2002). The original data come from Barnett et al. (1995).

The human brain (and that of rats, coincidentally) is protected from the bacteria and toxins that course through the bloodstream by something called the blood-brain barrier. After a method of disrupting the barrier was developed, researchers tested this new mechanism, as follows. A series of 34 rats were inoculated with human lung cancer cells to induce brain tumors. After 9-11 days they were infused with either the barrier disruption (BD) solution or, as a control, a normal saline (NS) solution. Fifteen minutes later, the rats received a standard dose of a particular therapeutic antibody (L6-F(ab')2. The key measure of the effectiveness of transmission across the brain-blood barrier is the ratio of the antibody concentration in the brain tumor to the antibody concentration in normal tissue outside the brain. The rats were then sacrificed, and the amounts of antibody in the brain tumor and in normal tissue from the liver were measured. The study's primary objective is to determine whether the antibody concentration in the tumor increased when the blood-barrier disruption infusion was given, and if so, by how much?

1.9 The bloodbrain.csv file

Consider the data, available on the Data and Code page of our course website in the bloodbrain.csv file, which includes the following variables:

Variable	Description
case	identification number for the rat (1 - 34)
brain	an outcome: Brain tumor antibody count (per gram)
liver	an outcome: Liver antibody count (per gram)
tlratio	an outcome: tumor / liver concentration ratio
solution	the treatment: BD (barrier disruption) or NS (normal saline)
sactime	a design variable: Sacrifice time (hours; either 0.5, 3, 24 or 72)
postin	covariate: Days post-inoculation of lung cancer cells (9, 10 or
	11)
sex	covariate: M or F
wt.init	covariate: Initial weight (grams)
wt.loss	covariate: Weight loss (grams)
wt.tumor	covariate: Tumor weight (10 ⁻⁴ grams)

And here's what the data look like in R.

bloodbrain

```
# A tibble: 34 x 11

case brain liver tlratio solution sactime postin sex wt.init
<int> <int> <int> <int> <fot> <fot> <fot> 
1 41081 1456164 0.0282 BD 0.500 10 F 239
```

```
2 44286 1602171 0.0276 BD
                                           0.500
                                                     10 F
                                                                  225
 3
      3 102926 1601936 0.0642 BD
                                           0.500
                                                     10 F
                                                                  224
                                                     10 F
 4
      4 25927 1776411 0.0146 BD
                                           0.500
                                                                  184
 5
      5 42643 1351184 0.0316 BD
                                           0.500
                                                     10 F
                                                                  250
 6
      6
         31342 1790863 0.0175 NS
                                           0.500
                                                     10 F
                                                                  196
7
                                           0.500
      7 22815 1633386 0.0140 NS
                                                     10 F
                                                                  200
                                           0.500
8
        16629 1618757 0.0103 NS
                                                     10 F
                                                                  273
                                                     10 F
                                                                  216
9
      9
         22315 1567602 0.0142 NS
                                           0.500
10
     10
         77961 1060057 0.0735 BD
                                           3.00
                                                     10 F
                                                                  267
# ... with 24 more rows, and 2 more variables: wt.loss <dbl>,
   wt.tumor <int>
```

1.10 A Table 1 for bloodbrain

Barnett et al. (1995) did not provide a Table 1 for these data, so let's build one to compare the two solutions (BD vs. NS) on the covariates and outcomes, plus the natural logarithm of the tumor/liver concentration ratio (tlratio). We'll opt to treat the sacrifice time (sactime) and the days post-inoculation of lung cancer cells (postin) as categorical rather than quantitative variables.

Summary of continuous variables

```
solution: BD
                                              p25
         n miss p.miss
                         mean
                                 sd median
                                                    p75
                                                           min
                                                                 max
wt.init
       17
              0
                          243 3e+01
                                     2e+02
                                            2e+02 3e+02
                                                         2e+02 3e+02
wt.loss 17
              0
                     0
                            3 5e+00 4e+00
                                            1e+00 6e+00 -5e+00 1e+01
wt.tumor 17
              0
                     0
                          157 8e+01
                                     2e+02
                                            1e+02 2e+02
                                                         2e+01 4e+02
              0
                     0 56043 3e+04 5e+04 4e+04 8e+04
                                                         6e+03 1e+05
brain
         17
                     0 672577 7e+05 6e+05 2e+04 1e+06
                                                         2e+03 2e+06
liver
         17
              0
                            2 3e+00 1e-01 6e-02 3e+00 1e-02 9e+00
tlratio
        17
              0
                     0
logTL
              0
                           -1 2e+00 -2e+00 -3e+00 1e+00 -4e+00 2e+00
         17
         skew kurt
wt.init -0.39 0.7
wt.loss -0.10 0.2
```

```
wt.tumor 0.53 1.0
brain 0.29 -0.6
                0.35 - 1.7
liver
tlratio 1.58 1.7
logTL 0.08 -1.7
 -----
solution: NS
                   n miss p.miss mean sd median p25 p75 min max
wt.init 17 0 0 240 3e+01 2e+02 2e+02 3e+02 2e+02 3e+02

      wt.lnit
      17
      0
      0
      240 Se+01
      2e+02
      2e+02
      3e+02
      2e+02
      3e+02
      2e+02
      3e+02
      2e+02
      3e+02
      3e+04
      1e+03
      3e+0
                   skew kurt
wt.init 0.33 -0.48
wt.loss -0.09 0.08
wt.tumor 0.63 0.77
brain 0.30 -0.35
liver
                0.40 - 1.56
tlratio 2.27 4.84
logTL
                0.27 - 1.61
p-values
                         pNormal pNonNormal
wt.init 0.807308940 0.641940278
wt.loss 0.683756156 0.876749808
wt.tumor 0.151510151 0.190482094
brain 0.001027678 0.002579901
liver
                 0.974853609 0.904045603
tlratio 0.320501715 0.221425879
logTL
                0.351633525 0.221425879
Standardize mean differences
                         1 vs 2
wt.init 0.08435244
wt.loss 0.14099823
wt.tumor 0.50397184
brain 1.23884159
liver 0.01089667
tlratio 0.34611465
logTL 0.32420504
          ### Summary of categorical variables ###
solution: BD
          var n miss p.miss level freq percent cum.percent
```

sactime 17 0 0.0 0.5 5 29.4 29.4

3 4 23.5

24 4 23.5

72 4 23.5

52.9

76.5

100.0

postin	17	0	0.0	9	1	5.9	5.9
				10	14	82.4	88.2
				11	2	11.8	100.0
sex	17	0	0.0	F	13	76.5	76.5
				M	4	23.5	100.0
solution	: NS	3					
var	n	miss	p.miss	level	freq	percent	cum.percent
sactime	17	0	0.0	0.5	4	23.5	23.5
				3	5	29.4	52.9
				24	4	23.5	76.5
				72	4	23.5	100.0
postin	17	0	0.0	9	2	11.8	11.8
				10	13	76.5	88.2
				11	2	11.8	100.0
sex	17	0	0.0	F	13	76.5	76.5
				M	4	23.5	100.0

p-values

pApprox pExact sactime 0.9739246 1 postin 0.8309504 1 sex 1.0000000 1

Standardize mean differences

1 vs 2 sactime 0.1622214 postin 0.2098877 sex 0.0000000

Note that, in this particular case, the decisions we make about normality vs. non-normality (for quantitative variables) and the decisions we make about approximate vs. exact testing (for categorical variables) won't actually change the implications of the p values. Each approach gives similar results for each variable. Of course, that's not always true.

1.10.1 Generate final Table 1 for bloodbrain

I'll choose to treat tlratio and its logarithm as non-Normal, but otherwise, use t tests, but admittedly, that's an arbitrary decision, really.

print(bb.att1, nonnormal = c("tlratio", "logTL"))

	Stratified by solution	on
	BD	NS
n	17	17
<pre>sactime (%)</pre>		
0.5	5 (29.4)	4 (23.5)
3	4 (23.5)	5 (29.4)
24	4 (23.5)	4 (23.5)

```
4 (23.5)
   72
                                4 (23.5)
postin (%)
   9
                                1 (5.9)
                                                         2 (11.8)
   10
                               14 (82.4)
                                                        13 (76.5)
   11
                                2 (11.8)
                                                         2 (11.8)
sex = M (\%)
                                4 (23.5)
                                                         4 (23.5)
wt.init (mean (sd))
                           242.82 (27.23)
                                                    240.47 (28.54)
wt.loss (mean (sd))
                                                      3.94 (3.88)
                             3.34 (4.68)
wt.tumor (mean (sd))
                           157.29 (84.00)
                                                    208.53 (116.68)
brain (mean (sd))
                         56043.41 (33675.40)
                                                  23887.18 (14610.53)
liver (mean (sd))
                        672577.35 (694479.58)
                                                 664975.47 (700773.13)
tlratio (median [IQR])
                             0.12 [0.06, 2.84]
                                                      0.05 [0.03, 0.94]
logTL (median [IQR])
                            -2.10 [-2.74, 1.04]
                                                     -2.95 [-3.41, -0.07]
                       Stratified by solution
                               test
sactime (%)
                         0.974
   0.5
   3
   24
   72
postin (%)
                         0.831
   9
   10
   11
sex = M (\%)
                         1.000
wt.init (mean (sd))
                         0.807
wt.loss (mean (sd))
                         0.684
wt.tumor (mean (sd))
                         0.152
brain (mean (sd))
                         0.001
liver (mean (sd))
                         0.975
tlratio (median [IQR])
                        0.221 nonnorm
logTL (median [IQR])
                         0.221 nonnorm
```

Or, we can get an Excel-readable version placed in a data subfolder, using

4	Α	В	С	D	E
1		BD	NS	р	test
2	n	17	17		
3	sex = M (%)	4 (23.5)	4 (23.5)	1	
4	sactime (%)			0.974	
5	0.5	5 (29.4)	4 (23.5)		
6	3	4 (23.5)	5 (29.4)		
7	24	4 (23.5)	4 (23.5)		
8	72	4 (23.5)	4 (23.5)		
9	postin (%)			0.831	
10	9	1 (5.9)	2 (11.8)		
11	10	14 (82.4)	13 (76.5)		
12	11	2 (11.8)	2 (11.8)		
13	wt.init (mean (sd))	242.82 (27.23)	240.47 (28.54)	0.807	
14	wt.loss (mean (sd))	3.34 (4.68)	3.94 (3.88)	0.684	
15	wt.tumor (mean (sd))	157.29 (84.00)	208.53 (116.68)	0.152	
16	brain (mean (sd))	56043.41 (33675.40)	23887.18 (14610.53)	0.001	
17	liver (mean (sd))	672577.35 (694479.58)	664975.47 (700773.13)	0.975	
18	tlratio (median [IQR])	0.12 [0.06, 2.84]	0.05 [0.03, 0.94]	0.221	nonnorm
19	logTL (median [IQR])	-2.10 [-2.74, 1.04]	-2.95 [-3.41, -0.07]	0.221	nonnorm
20					

One thing I would definitely clean up here, in practice, is to change the presentation of the p value for sex from 1 to > 0.99, or just omit it altogether. I'd also drop the computer-ese where possible, add units for the measures, round a lot, identify the outcomes carefully, and use notes to indicate deviations from the main approach.

1.10.2 A More Finished Version (after Cleanup in Word)

Table 1. Comparing Rats Receiving BD to those Receiving NS on Available Covariates and Design Variables, and Key Outcomes

	Barrier Disruption	Normal Saline	
	(BD: treatment)	(NS: control)	р
# of Rats	17	17	
Sex = Male	4 (23.5)	4 (23.5)	-
Sacrifice Time (hours)			0.97
0.5	5 (29.4)	4 (23.5)	
3	4 (23.5)	5 (29.4)	
24	4 (23.5)	4 (23.5)	
72	4 (23.5)	4 (23.5)	
Days post-inoculation of			0.83
lung cancer cells			0.03
9	1 (5.9)	2 (11.8)	
10	14 (82.4)	13 (76.5)	
11	2 (11.8)	2 (11.8)	
Initial Weight (g)	243 (27)	240 (29)	0.81
Weight Loss (g)	3.3 (4.7)	3.9 (3.9)	0.68
Tumor Weight (10 ⁻⁴ g)	157.3 (84.0)	208.5 (116.7)	0.15
Key Outcomes: mean (sd) unless otherw	ise indicated		
Brain Tumor Antibody Count (per g)	56,043 (33,675)	23,887 (14,611)	0.001
Liver Antibody Count (per g)	672,577 (694,480)	664,975 (700,773)	0.98
Tumor/Liver Ratio	0.12	0.05	0.22
(median [Q25, Q75])	[0.06, 2.84]	[0.03, 0.94]	0.22
Natural Log of Tumor/Liver Ratio	-2.10	-2.95	0.22
(median [Q25, Q75])	[-2.74, 1.04]	[-3.41, -0.07]	0.22

Table 1 Notes:

- Categorical variables are summarized with counts, percentages and p values based on approximate chi-square tests.
- Continuous variables, unless otherwise indicated, are summarized with means, standard deviations and p values based on t tests.
- The Tumor / Liver ratio and its natural logarithm are summarized with the median and quartiles and a p value from a non-parametric (Wilcoxon signed rank) test.

Chapter 2

Linear Regression on a small SMART data set

2.1 BRFSS and SMART

The Centers for Disease Control analyzes Behavioral Risk Factor Surveillance System (BRFSS) survey data for specific metropolitan and micropolitan statistical areas (MMSAs) in a program called the Selected Metropolitan/Micropolitan Area Risk Trends of BRFSS (SMART BRFSS.)

In this work, we will focus on data from the 2016 SMART, and in particular on data from the Cleveland-Elyria, OH, Metropolitan Statistical Area. The purpose of this survey is to provide localized health information that can help public health practitioners identify local emerging health problems, plan and evaluate local responses, and efficiently allocate resources to specific needs.

2.1.1 Key resources

- the full data are available in the form of the 2016 SMART BRFSS MMSA Data, found in a zipped SAS Transport Format file. The data were released in August 2017.
- the MMSA Variable Layout PDF which simply lists the variables included in the data file
- the Calculated Variables PDF which describes the risk factors by data variable names there is also an online summary matrix of these calculated variables, as well.
- the lengthy 2016 Survey Questions PDF which lists all questions asked as part of the BRFSS in 2016
- the enormous Codebook for the 2016 BRFSS Survey PDF which identifies the variables by name for

Later this term, we'll use all of those resources to help construct a more complete data set than we'll study today. I'll also demonstrate how I built the smartcle1 data set that we'll use in this Chapter.

2.2 The smartcle1 data: Cookbook

The smartcle1.csv data file available on the Data and Code page of our website describes information on 11 variables for 1036 respondents to the BRFSS 2016, who live in the Cleveland-Elyria, OH, Metropolitan Statistical Area. The variables in the smartcle1.csv file are listed below, along with (in some cases) the BRFSS items that generate these responses.

Vaniable	Description
variable	Description
SEQNO	respondent identification number (all begin with 2016)

Variable	Description
physhealth	Now thinking about your physical health, which includes physical illness and injury, for how many days during the past 30 days was your physical health not good?
menthealth	Now thinking about your mental health, which includes stress, depression, and problems with emotions, for how many days during the past 30 days was your mental health not good?
poorhealth	During the past 30 days, for about how many days did poor physical or mental health keep you from doing your usual activities, such as self-care, work, or recreation?
genhealth	Would you say that in general, your health is (five categories: Excellent, Very Good, Good, Fair or Poor)
bmi	Body mass index, in kg/m ²
female	Sex, $1 = \text{female}$, $0 = \text{male}$
internet30	Have you used the internet in the past 30 days? $(1 = yes, 0 = no)$
exerany	During the past month, other than your regular job, did you participate in any physical activities or exercises such as running, calisthenics, golf, gardening, or walking for exercise? $(1 = yes, 0 = no)$
sleephrs	On average, how many hours of sleep do you get in a 24-hour period?
alcdays	How many days during the past 30 days did you have at least one drink of any alcoholic beverage such as beer, wine, a malt beverage or liquor?

str(smartcle1)

```
Classes 'tbl_df', 'tbl' and 'data.frame':
                                          1036 obs. of 11 variables:
           : num 2.02e+09 2.02e+09 2.02e+09 2.02e+09 2.02e+09 ...
$ physhealth: int  0 0 1 0 5 4 2 2 0 0 ...
 $ menthealth: int  0 0 5 0 0 18 0 3 0 0 ...
 $ poorhealth: int NA NA O NA O 6 O O NA NA ...
 $ genhealth : Factor w/ 5 levels "1_Excellent",..: 2 1 2 3 1 2 3 3 2 3 ...
            : num 26.7 23.7 26.9 21.7 24.1 ...
 $ bmi
            : int 1001001100...
 $ internet30: int 1 1 1 1 1 1 1 1 1 1 ...
          : int 1 1 0 1 1 1 1 1 1 0 ...
 $ exerany
$ sleephrs : int 6 6 8 9 7 5 9 7 7 7 ...
 $ alcdays
           : int 1 4 4 3 2 28 4 2 4 25 ...
```

2.3 smartcle2: Omitting Missing Observations: Complete-Case Analyses

For the purpose of fitting our first few models, we will eliminate the missingness problem, and look only at the *complete cases* in our **smartcle1** data. We will discuss methods for imputing missing data later in these Notes.

To inspect the missingness in our data, we might consider using the skim function from the skimr package. We'll exclude the respondent identifier code (SEQNO) from this summary as uninteresting.

```
skim_with(numeric = list(hist = NULL), integer = list(hist = NULL))
## above line eliminates the sparkline histograms
## it can be commented out when working in the console,
## but I need it to produce the Notes without errors right now
```

```
smartcle1 %>%
   skim(-SEQNO)
Skim summary statistics
n obs: 1036
n variables: 11
Variable type: factor
 variable missing complete
                              n n_unique
genhealth
                      1033 1036
                3
                            top_counts ordered
2_V: 350, 3_G: 344, 1_E: 173, 4_F: 122
Variable type: integer
  variable missing complete
                                        sd p0 p25 median p75 p100
                               n mean
   alcdays
              46
                      990 1036 4.65 8.05
   exerany
                3
                       1033 1036 0.76 0.43 0
                                                1
                                                               1
    female
                 0
                       1036 1036 0.6 0.49 0
                                                0
                                                          1
                                                               1
                6
                       1030 1036 0.81 0.39 0
internet30
                                               1
                                                      1
                                                          1
                                                               1
menthealth
                11
                       1025 1036 2.72 6.82 0
                                                              30
                       1019 1036 3.97 8.67 0
                                                          2
physhealth
               17
                                                      0
                                                              30
                                                0
poorhealth
               543
                       493 1036 4.07 8.09 0
                                                              30
  sleephrs
                 8
                       1028 1036 7.02 1.53 1
                                                              20
Variable type: numeric
variable missing complete
                             n mean
                                       sd
                                             p0 p25 median
                                                             p75 p100
                      952 1036 27.89 6.47 12.71 23.7 26.68 30.53 66.06
     bmi
```

Now, we'll create a new tibble called smartcle2 which contains every variable except poorhealth, and which includes all respondents with complete data on the variables (other than poorhealth). We'll store those observations with complete data in the smartcle2 tibble.

```
smartcle2 <- smartcle1 %>%
    select(-poorhealth) %>%
    filter(complete.cases(.))
smartcle2
```

A tibble: 896 x 10

	SEQNO	physhealth	${\tt menthealth}$	genhealth	bmi	${\tt female}$	internet30	exerany
	<dbl></dbl>	<int></int>	<int></int>	<fct></fct>	<dbl></dbl>	<int></int>	<int></int>	<int></int>
1	2.02e9	0	0	2_VeryGo~	26.7	1	1	1
2	2.02e9	0	0	1_Excell~	23.7	0	1	1
3	2.02e9	1	5	2_VeryGo~	26.9	0	1	0
4	2.02e9	0	0	3_Good	21.7	1	1	1
5	2.02e9	5	0	1_Excell~	24.1	0	1	1
6	2.02e9	4	18	2_VeryGo~	27.6	0	1	1
7	2.02e9	2	0	3_Good	25.7	1	1	1
8	2.02e9	2	3	3_Good	28.5	1	1	1
9	2.02e9	0	0	2_VeryGo~	28.6	0	1	1
10	2.02e9	0	0	3_Good	23.1	0	1	0

... with 886 more rows, and 2 more variables: sleephrs <int>,

alcdays <int>

Note that there are only 896 respondents with **complete** data on the 10 variables (excluding **poorhealth**) in the **smartcle2** tibble, as compared to our original **smartcle1** data which described 1036 respondents and

11 variables, but with lots of missing data.

2.4 Summarizing the smartcle2 data numerically

2.4.1 The New Toy: The skim function

```
skim(smartcle2, -SEQNO)
Skim summary statistics
n obs: 896
n variables: 10
Variable type: factor
 variable missing complete
                             n n_unique
genhealth
                       896 896
                            top_counts ordered
2_V: 306, 3_G: 295, 1_E: 155, 4_F: 102
Variable type: integer
  variable missing complete
                                       sd p0 p25 median p75 p100
                              n mean
               0
                        896 896 4.83 8.14
   alcdays
                                               0
                 0
   exerany
                        896 896 0.77 0.42 0
                                               1
                                                      1
    female
                 0
                        896 896 0.58 0.49 0
 internet30
                 0
                        896 896 0.81 0.39 0
                                               1
                                                               1
menthealth
                 0
                        896 896 2.69 6.72 0
                                                      0
                                                              30
                 0
                        896 896 3.99 8.64 0
                                                      0 2
                                                              30
physhealth
                        896 896 7.02 1.48 1
  sleephrs
                 0
Variable type: numeric
                                            p0 p25 median
variable missing complete
                            n mean
                                      sd
                                                             p75 p100
                      896 896 27.87 6.33 12.71 23.7
                                                      26.8 30.53 66.06
     bmi
```

2.4.2 The usual summary for a data frame

Of course, we can use the usual summary to get some basic information about the data.

summary(smartcle2)

```
SEQNO
                     physhealth
                                    menthealth
                                                        genhealth
Min.
      :2.016e+09
                  Min. : 0.00
                                  Min. : 0.000 1_Excellent:155
                   1st Qu.: 0.00
                                  1st Qu.: 0.000
1st Qu.:2.016e+09
                                                  2_VeryGood:306
Median :2.016e+09
                   Median: 0.00
                                  Median : 0.000
                                                  3_{Good}
                                                             :295
     :2.016e+09
                   Mean
                        : 3.99
                                  Mean
                                        : 2.693
                                                  4_Fair
                                                             :102
                                                             : 38
3rd Qu.:2.016e+09
                   3rd Qu.: 2.00
                                  3rd Qu.: 2.000
                                                  5_Poor
      :2.016e+09
                  Max. :30.00
                                  Max.
                                         :30.000
Max.
                   female
                                 internet30
    bmi
                                                  exerany
               Min. :0.0000 Min. :0.0000
                                                      :0.0000
Min.
      :12.71
                                               Min.
1st Qu.:23.70 1st Qu.:0.0000
                              1st Qu.:1.0000
                                               1st Qu.:1.0000
Median :26.80
               Median :1.0000
                               Median :1.0000
                                               Median :1.0000
Mean :27.87
               Mean :0.5848
                               Mean :0.8147
                                               Mean
                                                      :0.7667
3rd Qu.:30.53
               3rd Qu.:1.0000
                               3rd Qu.:1.0000
                                               3rd Qu.:1.0000
Max. :66.06
                     :1.0000
                              Max. :1.0000
                                                      :1.0000
               Max.
                                               {\tt Max.}
```

```
      sleephrs
      alcdays

      Min. : 1.000
      Min. : 0.000

      1st Qu.: 6.000
      1st Qu.: 0.000

      Median : 7.000
      Median : 1.000

      Mean : 7.022
      Mean : 4.834

      3rd Qu.: 8.000
      3rd Qu.: 5.000

      Max. : 20.000
      Max. : 30.000
```

2.4.3 The describe function in Hmisc

Or we can use the describe function from the Hmisc package.

```
Hmisc::describe(select(smartcle2, bmi, genhealth, female))
select(smartcle2, bmi, genhealth, female)
                 896 Observations
 3 Variables
      n missing distinct Info Mean Gmd .05 .10
896 0 467 1 27.87 6.572 20.06 21.23
.25 .50 .75 .90 .95
      . 25
   23.70 26.80 30.53 35.36 39.30
lowest: 12.71 13.34 14.72 16.22 17.30, highest: 56.89 57.04 60.95 61.84 66.06
______
genhealth
        n missing distinct
      896 0 5

        Value
        1_Excellent
        2_VeryGood
        3_Good
        4_Fair

        Frequency
        155
        306
        295
        102

        Proportion
        0.173
        0.342
        0.329
        0.114

                                                                              38
                                                                               0.042
      n missing distinct Info Sum Mean Gmd 896 0 2 0.728 524 0.5848 0.4862
```

2.5 Counting as exploratory data analysis

Counting things can be amazingly useful.

2.5.1 How many respondents had exercised in the past 30 days? Did this vary by sex?

```
7.14
1
                 0
                       64
2
        0
                 1
                      308
                             34.4
3
        1
                 0
                      145
                             16.2
4
                             42.3
        1
                 1
                      379
```

so we know now that 42.3% of the subjects in our data were women who exercised. Suppose that instead we want to find the percentage of exercisers within each sex...

```
smartcle2 %>%
    count(female, exerany) %>%
   group_by(female) %>%
   mutate(prob = 100*n / sum(n))
# A tibble: 4 x 4
# Groups:
            female [2]
  female exerany
                     n prob
   <int>
           <int> <int> <dbl>
               0
                       17.2
1
       0
                    64
2
                   308 82.8
       0
               1
3
               0
                   145
                        27.7
       1
                   379 72.3
```

and now we know that 82.8% of the males exercised at least once in the last 30 days, as compared to 72.3% of the females.

What's the distribution of sleephrs?

We can count quantitative variables with discrete sets of possible values, like sleephrs, which is captured as an integer (that must fall between 0 and 24.)

```
smartcle2 %>% count(sleephrs)
```

```
# A tibble: 14 x 2
   sleephrs
                  n
       <int> <int>
 1
           1
                  5
           2
 2
                  1
 3
           3
                  6
 4
           4
                 20
 5
           5
                 63
 6
           6
                192
 7
           7
                276
 8
           8
                266
9
           9
                 38
10
          10
                 22
                  2
          11
11
12
          12
                  2
13
          16
                  2
14
          20
```

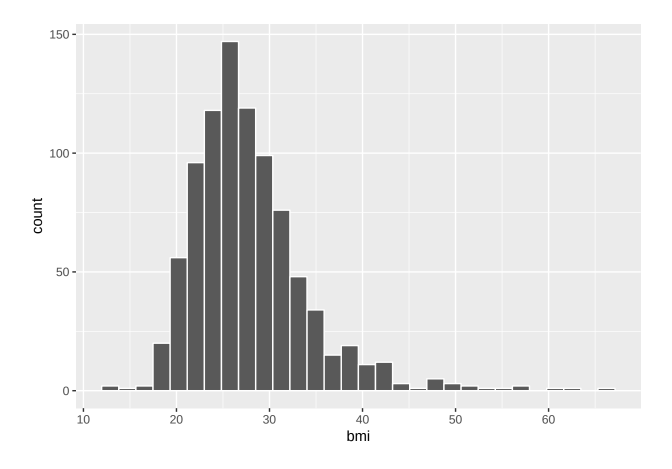
Of course, a natural summary of a quantitative variable like this would be graphical.

```
ggplot(smartcle2, aes(sleephrs)) +
    geom_histogram(binwidth = 1, fill = "dodgerblue", col = "darkred")
```



2.5.3 What's the distribution of BMI?

```
ggplot(smartcle2, aes(bmi)) +
  geom_histogram(bins = 30, col = "white")
```



2.5.4 How many of the respondents have a BMI below 30?

2.5.5 How many of the respondents who have a BMI < 30 exercised?

```
smartcle2 %>% count(exerany, bmi < 30) %>%
    group_by(exerany) %>%
    mutate(percent = 100*n/sum(n))
# A tibble: 4 x 4
# Groups: exerany [2]
  exerany `bmi < 30`
                        n percent
    <int> <lgl>
                     <int>
                             <dbl>
1
       0 F
                       88
                              42.1
2
       0 T
                       121
                              57.9
3
       1 F
                      165
                              24.0
4
       1 T
                      522
                             76.0
```

2.5.6 Is obesity associated with sex, in these data?

```
smartcle2 %>% count(female, bmi < 30) %>%
    group_by(female) %>%
    mutate(percent = 100*n/sum(n))
# A tibble: 4 x 4
# Groups: female [2]
  female `bmi < 30`</pre>
                         n percent
   <int> <lgl>
                             <dbl>
                    <int>
       0 F
                       105
                              28.2
1
2
       0 T
                       267
                              71.8
3
       1 F
                       148
                              28.2
4
       1 T
                       376
                              71.8
```

2.5.7 Comparing sleephrs summaries by obesity status

Can we compare the sleephrs means, medians and 75th percentiles for respondents whose BMI is below 30 to the respondents whose BMI is not?

2.5.8 The skim function within a pipe

The **skim** function works within pipes and with the other **tidyverse** functions.

```
smartcle2 %>%
   group_by(exerany) %>%
   skim(bmi, sleephrs)
Skim summary statistics
n obs: 896
n variables: 10
group variables: exerany
Variable type: integer
 exerany variable missing complete n mean sd p0 p25 median p75 p100
      0 sleephrs
                       0
                          209 209 7
                                           1.85 1
                                                            7
                                                                    20
                       0
                              687 687 7.03 1.34 1
      1 sleephrs
                                                            7
                                                                    16
Variable type: numeric
 exerany variable missing complete
                                                         p25 median
                                    n mean
                                              sd
                                                    p0
      0
             bmi
                       0
                              209 209 29.57 7.46 18
                                                       24.11 28.49 33.13
      1
             bmi
                       0
                              687 687 27.35 5.84 12.71 23.7
                                                              26.52 29.81
 p100
```

66.06 60.95

2.6 First Modeling Attempt: Can bmi predict physhealth?

We'll start with an effort to predict physhealth using bmi. A natural graph would be a scatterplot.

```
ggplot(data = smartcle2, aes(x = bmi, y = physhealth)) +
   geom_point()
```



A good question to ask ourselves here might be: "In what BMI range can we make a reasonable prediction of physhealth?"

Now, we might take the plot above and add a simple linear model ...

```
ggplot(data = smartcle2, aes(x = bmi, y = physhealth)) +
   geom_point() +
   geom_smooth(method = "lm", se = FALSE)
```



which shows the same least squares regression model that we can fit with the ${\tt lm}$ command.

2.6.1 Fitting a Simple Regression Model

```
Residuals:
    Min    1Q Median    3Q    Max
-9.171 -4.057 -3.193 -1.576 28.073
```

Coefficients:

```
Estimate Std. Error t value Pr(>|t|)
(Intercept) -1.45143
                       1.29185 -1.124
bmi
            0.19527
                        0.04521
                                  4.319 1.74e-05 ***
Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
Residual standard error: 8.556 on 894 degrees of freedom
Multiple R-squared: 0.02044,
                               Adjusted R-squared: 0.01934
F-statistic: 18.65 on 1 and 894 DF, p-value: 1.742e-05
confint(model_A, level = 0.95)
                 2.5 %
                          97.5 %
(Intercept) -3.9868457 1.0839862
            0.1065409 0.2840068
```

The model coefficients can be obtained by printing the model object, and the summary function provides several useful descriptions of the model's residuals, its statistical significance, and quality of fit.

2.6.2 Model Summary for a Simple (One-Predictor) Regression

The fitted model predicts physhealth with the equation -1.45 + 0.195*bmi, as we can read off from the model coefficients.

Each of the 896 respondents included in the smartcle2 data makes a contribution to this model.

2.6.2.1 Residuals

Suppose Harry is one of the people in that group, and Harry's data is bmi = 20, and physhealth = 3.

- Harry's *observed* value of physhealth is just the value we have in the data for them, in this case, observed physhealth = 3 for Harry.
- Harry's fitted or predicted physhealth value is the result of calculating -1.45 + 0.195*bmi for Harry. So, if Harry's BMI was 20, then Harry's predicted physhealth value is -1.45 + (0.195)(20) = 2.45.
- The residual for Harry is then his observed outcome minus his fitted outcome, so Harry has a residual of 3 2.45 = 0.55.
- Graphically, a residual represents vertical distance between the observed point and the fitted regression line.
- Points above the regression line will have positive residuals, and points below the regression line will have negative residuals. Points on the line have zero residuals.

The residuals are summarized at the top of the summary output for linear model.

- The mean residual will always be zero in an ordinary least squares model, but a five number summary of the residuals is provided by the summary, as is an estimated standard deviation of the residuals (called here the Residual standard error.)
- In the smartcle2 data, the minimum residual was -9.17, so for one subject, the observed value was 9.17 days smaller than the predicted value. This means that the prediction was 9.17 days too large for that subject.
- Similarly, the maximum residual was 28.07 days, so for one subject the prediction was 28.07 days too small. Not a strong performance.
- In a least squares model, the residuals are assumed to follow a Normal distribution, with mean zero, and standard deviation (for the smartcle2 data) of about 8.6 days. Thus, by the definition of a Normal distribution, we'd expect
- about 68% of the residuals to be between -8.6 and +8.6 days,

- about 95% of the residuals to be between -17.2 and +17.2 days,
- about all (99.7%) of the residuals to be between -25.8 and +25.8 days.

2.6.2.2 Coefficients section

The summary for a linear model shows Estimates, Standard Errors, t values and p values for each coefficient fit.

- The Estimates are the point estimates of the intercept and slope of bmi in our model.
- In this case, our estimated slope is 0.195, which implies that if Harry's BMI is 20 and Sally's BMI is 21, we predict that Sally's physhealth will be 0.195 days larger than Harry's.
- The Standard Errors are also provided for each estimate. We can create rough 95% confidence intervals by adding and subtracting two standard errors from each coefficient, or we can get a slightly more accurate answer with the confint function.
- Here, the 95% confidence interval for the slope of bmi is estimated to be (0.11, 0.28). This is a good measure of the uncertainty in the slope that is captured by our model. We are 95% confident in the process of building this interval, but this doesn't mean we're 95% sure that the true slope is actually in that interval.

Also available are a t value (just the Estimate divided by the Standard Error) and the appropriate p value for testing the null hypothesis that the true value of the coefficient is 0 against a two-tailed alternative.

- If a slope coefficient is statistically significantly different from 0, this implies that 0 will not be part of the uncertainty interval obtained through confint.
- If the slope was zero, it would suggest that bmi would add no predictive value to the model. But that's unlikely here.

If the bmi slope coefficient is associated with a small p value, as in the case of our model_A, it suggests that the model including bmi is statistically significantly better at predicting physhealth than the model without bmi.

• Without bmi our model_A would become an *intercept-only* model, in this case, which would predict the mean physhealth for everyone, regardless of any other information.

2.6.2.3 Model Fit Summaries

The summary of a linear model also displays:

- The residual standard error and associated degrees of freedom for the residuals.
- For a simple (one-predictor) least regression like this, the residual degrees of freedom will be the sample size minus 2.
- The multiple R-squared (or coefficient of determination)
- This is interpreted as the proportion of variation in the outcome (physhealth) accounted for by the model, and will always fall between 0 and 1 as a result.
- Our model_A accounts for a mere 2% of the variation in physhealth.
- The Adjusted R-squared value "adjusts" for the size of our model in terms of the number of coefficients included in the model.
- The adjusted R-squared will always be less than the Multiple R-squared.
- We still hope to find models with relatively large adjusted R² values.
- In particular, we hope to find models where the adjusted R² isn't substantially less than the Multiple R-squared.
- The adjusted R-squared is usually a better estimate of likely performance of our model in new data than is the Multiple R-squared.
- The adjusted R-squared result is no longer interpretable as a proportion of anything in fact, it can fall below 0.

• We can obtain the adjusted \mathbb{R}^2 from the raw \mathbb{R}^2 , the number of observations N and the number of predictors p included in the model, as follows:

$$R_{adj}^2 = 1 - \frac{(1 - R^2)(N - 1)}{N - p - 1},$$

- The F statistic and p value from a global ANOVA test of the model.
 - Obtaining a statistically significant result here is usually pretty straightforward, since the comparison is between our model, and a model which simply predicts the mean value of the outcome for everyone.
 - In a simple (one-predictor) linear regression like this, the t statistic for the slope is just the square root of the F statistic, and the resulting p values for the slope's t test and for the global F test will be identical.
- To see the complete ANOVA F test for this model, we can run anova(model_A).

```
anova(model_A)
```

Analysis of Variance Table

```
Response: physhealth

Df Sum Sq Mean Sq F value Pr(>F)

bmi 1 1366 1365.5 18.655 1.742e-05 ***

Residuals 894 65441 73.2

---

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

2.6.3 Using the broom package

The broom package has three functions of particular use in a linear regression model:

2.6.3.1 The tidy function

tidy builds a data frame/tibble containing information about the coefficients in the model, their standard errors, t statistics and p values.

```
tidy(model_A)
```

```
term estimate std.error statistic p.value
1 (Intercept) -1.4514298 1.29185199 -1.123526 2.615156e-01
2 bmi 0.1952739 0.04521145 4.319125 1.741859e-05
```

2.6.3.2 The glance function

glance' builds a data frame/tibble containing summary statistics about the model, including

- the (raw) multiple R^2 and adjusted R^2
- sigma which is the residual standard error
- the F statistic, p.value model df and df.residual associated with the global ANOVA test, plus
- several statistics that will be useful in comparing models down the line:
- the model's log likelihood function value, logLik
- the model's Akaike's Information Criterion value, AIC
- the model's Bayesian Information Criterion value, BIC
- and the model's deviance statistic

glance(model_A)

```
r.squared adj.r.squared sigma statistic p.value df logLik
1 0.02044019 0.01934449 8.555737 18.65484 1.741859e-05 2 -3193.723
AIC BIC deviance df.residual
1 6393.446 6407.84 65441.36 894
```

2.6.3.3 The augment function

augment builds a data frame/tibble which adds fitted values, residuals and other diagnostic summaries that describe each observation to the original data used to fit the model, and this includes

- .fitted and .resid, the fitted and residual values, in addition to
- .hat, the leverage value for this observation
- .cooksd, the Cook's distance measure of influence for this observation
- .stdresid, the standardized residual (think of this as a z-score a measure of the residual divided by its associated standard deviation .sigma)
- and se.fit which will help us generate prediction intervals for the model downstream

Note that each of the new columns begins with . to avoid overwriting any data.

```
head(augment(model_A))
```

```
physhealth
               bmi
                   .fitted
                              .se.fit
                                                                 .sigma
                                            .resid
                                                          .hat
           0 26.69 3.760430 0.2907252 -3.76043009 0.001154651 8.559600
           0 23.70 3.176561 0.3422908 -3.17656119 0.001600574 8.559865
2
           1 26.92 3.805343 0.2890054 -2.80534308 0.001141030 8.560010
           0 21.66 2.778202 0.4005101 -2.77820248 0.002191352 8.560020
           5 24.09 3.252718 0.3329154 1.74728200 0.001514095 8.560326
5
           4 27.64 3.945940 0.2860087 0.05405972 0.001117490 8.560526
                 .std.resid
       .cooksd
1 1.117852e-04 -0.439775451
2 1.106717e-04 -0.371575999
3 6.147744e-05 -0.328077528
4 1.160381e-04 -0.325074461
5 3.167016e-05 0.204378225
6 2.235722e-08 0.006322069
```

For more on the broom package, you may want to look at this vignette.

2.6.4 How does the model do? (Residuals vs. Fitted Values)

• Remember that the R^2 value was about 2%.

```
plot(model_A, which = 1)
```



This is a plot of residuals vs. fitted values. The goal here is for this plot to look like a random scatter of points, perhaps like a "fuzzy football", and that's **not** what we have. Why?

If you prefer, here's a ggplot2 version of a similar plot, now looking at standardized residuals instead of raw residuals, and adding a loess smooth and a linear fit to the result.

```
ggplot(augment(model_A), aes(x = .fitted, y = .std.resid)) +
   geom_point() +
   geom_smooth(method = "lm", se = FALSE, col = "red", linetype = "dashed") +
   geom_smooth(method = "loess", se = FALSE, col = "navy") +
   theme_bw()
```



The problem we're having here becomes, I think, a little more obvious if we look at what we're predicting. Does physhealth look like a good candidate for a linear model?

```
ggplot(smartcle2, aes(x = physhealth)) +
geom_histogram(bins = 30, fill = "dodgerblue", color = "royalblue")
```



No matter what model we fit, if we are predicting physhealth, and most of the data are values of 0 and 30, we have limited variation in our outcome, and so our linear model will be somewhat questionable just on that basis.

A normal Q-Q plot of the standardized residuals for our model_A shows this problem, too.

plot(model_A, which = 2)



We're going to need a method to deal with this sort of outcome, that has both a floor and a ceiling. We'll get there eventually, but linear regression alone doesn't look promising.

All right, so that didn't go anywhere great. Let's try again, with a new outcome.

2.7 A New Small Study: Predicting BMI

We'll begin by investigating the problem of predicting bmi, at first with just three regression inputs: sex, exerany and sleephrs, in our new smartcle2 data set.

- The outcome of interest is bmi.
- Inputs to the regression model are:
 - female = 1 if the subject is female, and 0 if they are male
 - exerany = 1 if the subject exercised in the past 30 days, and 0 if they didn't
 - sleephrs = hours slept in a typical 24-hour period (treated as quantitative)

2.7.1 Does female predict bmi well?

2.7.1.1 Graphical Assessment

```
ggplot(smartcle2, aes(x = female, y = bmi)) +
   geom_point()
```



Not so helpful. We should probably specify that female is a factor, and try another plotting approach.

```
ggplot(smartcle2, aes(x = factor(female), y = bmi)) +
   geom_boxplot()
```



The median BMI looks a little higher for males. Let's see if a model reflects that.

2.8 c2_m1: A simple t-test model

Residuals:

Coefficients:

Min 1Q Median 3Q Max -15.650 -4.129 -1.080 2.727 38.546

```
2.5 % 97.5 % (Intercept) 27.717372 29.00262801 female -1.686052 -0.00539878
```

The model suggests, based on these 896 subjects, that

- our best prediction for males is $BMI = 28.36 \text{ kg/m}^2$, and
- our best prediction for females is BMI = $28.36 0.85 = 27.51 \text{ kg/m}^2$.
- the mean difference between females and males is -0.85 kg/m^2 in BMI
- \bullet a 95% confidence (uncertainty) interval for that mean female male difference in BMI ranges from -1.69 to -0.01
- the model accounts for 0.4% of the variation in BMI, so that knowing the respondent's sex does very little to reduce the size of the prediction errors as compared to an intercept only model that would predict the overall mean (regardless of sex) for all subjects.
- the model makes some enormous errors, with one subject being predicted to have a BMI 38 points lower than his/her actual BMI.

Note that this simple regression model just gives us the t-test.

t.test(bmi ~ female, var.equal = TRUE, data = smartcle2)

```
Two Sample t-test

data: bmi by female

t = 1.9752, df = 894, p-value = 0.04855

alternative hypothesis: true difference in means is not equal to 0

95 percent confidence interval:
0.00539878 1.68605160

sample estimates:
mean in group 0 mean in group 1
28.36000 27.51427
```

2.9 c2_m2: Adding another predictor (two-way ANOVA without interaction)

When we add in the information about exerany to our original model, we might first picture the data. We could look at separate histograms,

```
ggplot(smartcle2, aes(x = bmi)) +
  geom_histogram(bins = 30) +
  facet_grid(female ~ exerany, labeller = label_both)
```



```
or maybe boxplots?
```

```
ggplot(smartcle2, aes(x = factor(female), y = bmi)) +
    geom_boxplot() +
    facet_wrap(~ exerany, labeller = label_both)
```



```
ggplot(smartcle2, aes(x = female, y = bmi))+
  geom_point(size = 3, alpha = 0.2) +
  theme_bw() +
  facet_wrap(~ exerany, labeller = label_both)
```



OK. Let's try fitting a model.

```
c2_m2 <- lm(bmi ~ female + exerany, data = smartcle2)</pre>
c2_m2
```

Call:

lm(formula = bmi ~ female + exerany, data = smartcle2)

Coefficients:

(Intercept) female exerany 30.334 -1.095 -2.384

This new model predicts only four predicted values:

- bmi = 30.334 if the subject is male and did not exercise (so female = 0 and exerany = 0)
- bmi = 30.334 1.095 = 29.239 if the subject is female and did not exercise (female = 1 and exerany = 0)
- bmi = 30.334 2.384 = 27.950 if the subject is male and exercised (so female = 0 and exerany = 1), and, finally
- bmi = 30.334 1.095 2.384 = 26.855 if the subject is female and exercised (so both female and exerany = 1).

For those who did not exercise, the model is:

• bmi = 30.334 - 1.095 female

and for those who did exercise, the model is:

• bmi = 27.95 - 1.095 female

Only the intercept of the bmi-female model changes depending on exerany.

```
summary(c2_m2)
lm(formula = bmi ~ female + exerany, data = smartcle2)
Residuals:
            1Q Median
   Min
                            3Q
                                   Max
-15.240 -4.091 -1.095
                         2.602 36.822
Coefficients:
           Estimate Std. Error t value Pr(>|t|)
(Intercept) 30.3335
                        0.5231
                                 57.99 < 2e-16 ***
            -1.0952
                        0.4262
                                -2.57
                                        0.0103 *
female
exerany
            -2.3836
                        0.4965
                                -4.80 1.86e-06 ***
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
Residual standard error: 6.239 on 893 degrees of freedom
                              Adjusted R-squared: 0.02722
Multiple R-squared: 0.02939,
F-statistic: 13.52 on 2 and 893 DF, p-value: 1.641e-06
confint(c2 m2)
```

```
2.5 % 97.5 % (Intercept) 29.306846 31.3602182 female -1.931629 -0.2588299 exerany -3.358156 -1.4090777
```

The slopes of both female and exerany have confidence intervals that are completely below zero, indicating that both female sex and exerany appear to be associated with reductions in bmi.

The R² value suggests that just under 3% of the variation in bmi is accounted for by this ANOVA model.

In fact, this regression (on two binary indicator variables) is simply a two-way ANOVA model without an interaction term.

```
anova(c2_m2)
```

Analysis of Variance Table

```
Response: bmi

Df Sum Sq Mean Sq F value Pr(>F)

female 1 156 155.61 3.9977 0.04586 *

exerany 1 897 896.93 23.0435 1.856e-06 ***

Residuals 893 34759 38.92

---

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

2.10 c2_m3: Adding the interaction term (Two-way ANOVA with interaction)

Suppose we want to let the effect of female vary depending on the exerany status. Then we need to incorporate an interaction term in our model.

```
c2_m3 <- lm(bmi ~ female * exerany, data = smartcle2)</pre>
c2_m3
```

Call:

lm(formula = bmi ~ female * exerany, data = smartcle2)

Coefficients:

(Intercept) female exerany female:exerany -0.8104 -2.1450-0.359230.1359

So, for example, for a male who exercises, this model predicts

• bmi = 30.136 - 0.810(0) - 2.145(1) - 0.359(0)(1) = 30.136 - 2.145 = 27.991

And for a female who exercises, the model predicts

•
$$bmi = 30.136 - 0.810$$
 (1) - 2.145 (1) - 0.359 (1)(1) = 30.136 - 0.810 - 2.145 - 0.359 = 26.822

For those who did not exercise, the model is:

• bmi = 30.136 - 0.81 female

But for those who did exercise, the model is:

- bmi = (30.136 2.145) + (-0.810 + (-0.359)) female, or ",
- $\bullet \ \operatorname{bmi} = 27.991 1.169 \ \operatorname{female}$

Now, both the slope and the intercept of the bmi-female model change depending on exerany.

```
summary(c2_m3)
```

Call:

lm(formula = bmi ~ female * exerany, data = smartcle2)

Residuals:

```
1Q Median
                          3Q
   Min
                                Max
-15.281 -4.101 -1.061
                       2.566 36.734
```

Coefficients:

```
Estimate Std. Error t value Pr(>|t|)
(Intercept)
             30.1359 0.7802 38.624
                                          <2e-16 ***
female
              -0.8104
                          0.9367 - 0.865
                                          0.3872
              -2.1450
                          0.8575 - 2.501
                                          0.0125 *
exerany
female:exerany -0.3592
                                          0.7328
                          1.0520 -0.341
```

Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1

Residual standard error: 6.242 on 892 degrees of freedom Multiple R-squared: 0.02952, Adjusted R-squared: 0.02625 F-statistic: 9.044 on 3 and 892 DF, p-value: 6.669e-06

confint(c2 m3)

```
2.5 %
                            97.5 %
              28.604610 31.6672650
(Intercept)
female
              -2.648893 1.0280526
exerany
              -3.827886 -0.4620407
female:exerany -2.423994 1.7055248
```

In fact, this regression (on two binary indicator variables and a product term) is simply a two-way ANOVA model with an interaction term.

```
anova(c2_m3)
```

Analysis of Variance Table

```
Response: bmi
               Df Sum Sq Mean Sq F value
                                            Pr(>F)
female
                     156 155.61 3.9938
                                           0.04597 *
exerany
                     897
                          896.93 23.0207 1.878e-06 ***
female: exerany
                       5
                            4.54
                                 0.1166
                                           0.73283
Residuals
              892
                  34754
                           38.96
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

The interaction term doesn't change very much here. Its uncertainty interval includes zero, and the overall model still accounts for just under 3% of the variation in bmi.

2.11 c2_m4: Using female and sleephrs in a model for bmi

```
ggplot(smartcle2, aes(x = sleephrs, y = bmi, color = factor(female))) +
    geom_point() +
    guides(col = FALSE) +
    geom_smooth(method = "lm", se = FALSE) +
    facet_wrap(~ female, labeller = label_both)
```



Does the difference in slopes of bmi and sleephrs for males and females appear to be substantial and important?

```
c2_m4 <- lm(bmi ~ female * sleephrs, data = smartcle2)
summary(c2_m4)</pre>
```

Call:

```
lm(formula = bmi ~ female * sleephrs, data = smartcle2)
```

Residuals:

```
Min 1Q Median 3Q Max
-15.498 -4.179 -1.035 2.830 38.204
```

Coefficients:

```
Estimate Std. Error t value Pr(>|t|)
(Intercept)
                 27.2661
                             1.6320 16.707
                                               <2e-16 ***
female
                  2.5263
                              2.0975
                                      1.204
                                                0.229
                  0.1569
                             0.2294
                                      0.684
                                                0.494
sleephrs
female:sleephrs
                -0.4797
                             0.2931 - 1.636
                                                0.102
```

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

```
Residual standard error: 6.31 on 892 degrees of freedom Multiple R-squared: 0.008341, Adjusted R-squared: 0.005006 F-statistic: 2.501 on 3 and 892 DF, p-value: 0.05818
```

Does it seem as though the addition of sleephrs has improved our model substantially over a model with female alone (which, you recall, was c2_m1)?

Since the c2_m4 model contains the c2_m1 model's predictors as a subset and the outcome is the same for each model, we consider the models *nested* and have some extra tools available to compare them.

• I might start by looking at the basic summaries for each model.

```
glance(c2_m4)
```

```
r.squared adj.r.squared sigma statistic p.value df logLik
1 0.004345169 0.003231461 6.31531 3.901534 0.04854928 2 -2921.675
AIC BIC deviance df.residual
1 5849.35 5863.744 35655.53 894
```

- The R² is twice as large for the model with sleephrs, but still very tiny.
- The p value for the global ANOVA test is actually less significant in c2_m4 than in c2_m1.
- Smaller AIC and smaller BIC statistics are more desirable. Here, there's little to choose from, but c2_m1 is a little better on each standard.
- We might also consider a significance test by looking at an ANOVA model comparison. This is only
 appropriate because c2_m1 is nested in c2_m4.

```
anova(c2_m4, c2_m1)
```

Analysis of Variance Table

```
Model 1: bmi ~ female * sleephrs

Model 2: bmi ~ female

Res.Df RSS Df Sum of Sq F Pr(>F)

1 892 35512

2 894 35656 -2 -143.11 1.7973 0.1663
```

The addition of the sleephrs term picked up 143 in the sum of squares column, at a cost of two degrees of freedom, yielding a p value of 0.166, suggesting that this isn't a significant improvement over the model that just did a t-test on female.

2.12 Making Predictions with a Linear Regression Model

Recall model 4, which yields predictions for body mass index on the basis of the main effects of sex (female) and hours of sleep (sleephrs) and their interaction.

2.12.1 Fitting an Individual Prediction and 95% Prediction Interval

What do we predict for the bmi of a subject who is female and gets 8 hours of sleep per night?

```
c2_new1 <- data_frame(female = 1, sleephrs = 8)
predict(c2_m4, newdata = c2_new1, interval = "prediction", level = 0.95)

fit lwr upr
1 27.21065 14.8107 39.6106</pre>
```

The predicted bmi for this new subject is 27.61. The prediction interval shows the bounds of a 95% uncertainty interval for a predicted bmi for an individual female subject who gets 8 hours of sleep on average per evening. From the predict function applied to a linear model, we can get the prediction intervals for any new data points in this manner.

2.12.2 Confidence Interval for an Average Prediction

• What do we predict for the **average body mass index of a population of subjects** who are female and sleep for 8 hours?

```
predict(c2_m4, newdata = c2_new1, interval = "confidence", level = 0.95)

fit lwr upr
1 27.21065 26.57328 27.84801
```

• How does this result compare to the prediction interval?

2.12.3 Fitting Multiple Individual Predictions to New Data

• How does our prediction change for a respondent if they instead get 7, or 9 hours of sleep? What if they are male, instead of female?

```
c2_new2 <- data_frame(subjectid = 1001:1006, female = c(1, 1, 1, 0, 0, 0), sleephrs = c(7, 8, 9, 7, 8,
pred2 <- predict(c2_m4, newdata = c2_new2, interval = "prediction", level = 0.95) %% tbl_df
result2 <- bind_cols(c2_new2, pred2)
result2
# A tibble: 6 x 6
 subjectid female sleephrs
                             fit
                                   lwr
                                         upr
      <int> <dbl>
                     <dbl> <dbl> <dbl> <dbl> <
      1001
             1.00
                      7.00 27.5 15.1 39.9
1
      1002
                      8.00 27.2 14.8 39.6
2
             1.00
3
                            26.9 14.5 39.3
      1003
             1.00
                      9.00
4
      1004
                      7.00
                            28.4 16.0 40.8
             0
      1005
                      8.00 28.5 16.1 40.9
5
             0
      1006
             0
                      9.00 28.7 16.2 41.1
```

The result2 tibble contains predictions for each scenario.

• Which has a bigger impact on these predictions and prediction intervals? A one category change in female or a one hour change in sleephrs?

2.12.4 Simulation to represent predictive uncertainty in Model 4

Suppose we want to predict the bmi of a female subject who sleeps for eight hours per night. As we have seen, we can do this automatically for a linear model like this one, using the predict function applied to the linear model, but a simulation prediction can also be done. Recall the detail of c2_m4:

```
Call:
lm(formula = bmi ~ female * sleephrs, data = smartcle2)
Coefficients:
    (Intercept)
                                          sleephrs female:sleephrs
                          female
        27.2661
                          2.5263
                                            0.1569
                                                            -0.4797
glance(c2_m4)
   r.squared adj.r.squared
                               sigma statistic
                                                   p.value df
                                                                 logLik
1 0.008341404
                0.005006229 6.309685
                                        2.50104 0.05818038 4 -2919.873
       AIC
                BIC deviance df.residual
```

We see that the residual standard error for our bmi predictions with this model is 6.31.

For a female respondent sleeping eight hours, recall that our point estimate (predicted value) of bmi is 27.21

```
predict(c2_m4, newdata = c2_new1, interval = "prediction", level = 0.95)
```

```
fit lwr upr
1 27.21065 14.8107 39.6106
```

1 5849.747 5873.736 35512.42

 $c2_m4$

The standard deviation is 6.31, so we could summarize the predictive distribution with a command that tells R to draw 1000 random numbers from a normal distribution with mean 27.21 and standard deviation 6.31. Let's summarize that and get a quick picture.

```
set.seed(432094)
pred.sim <- rnorm(1000, 27.21, 6.31)
hist(pred.sim, col = "royalblue")</pre>
```

Histogram of pred.sim



```
mean(pred.sim)

[1] 27.41856

quantile(pred.sim, c(0.025, 0.975))

2.5% 97.5%
14.48487 40.16778
```

How do these results compare to the prediction interval of (14.81, 39.61) that we generated earlier?

2.13 Centering the model

Our model c2_m4 has four predictors (the constant, sleephrs, female and their interaction) but just two inputs (female and sleephrs.) If we center the quantitative input sleephrs before building the model, we get a more interpretable interaction term.

```
smartcle2_c <- smartcle2 %>%
  mutate(sleephrs_c = sleephrs - mean(sleephrs))
```

```
c2_m4_c <- lm(bmi ~ female * sleephrs_c, data = smartcle2_c)
summary(c2_m4_c)</pre>
```

Call:

```
lm(formula = bmi ~ female * sleephrs_c, data = smartcle2_c)
```

Residuals:

```
Min 1Q Median 3Q Max
-15.498 -4.179 -1.035 2.830 38.204
```

Coefficients:

```
Estimate Std. Error t value Pr(>|t|)
(Intercept)
                  28.3681
                              0.3274 86.658
                                               <2e-16 ***
female
                  -0.8420
                              0.4280 - 1.967
                                               0.0495 *
                   0.1569
                              0.2294
                                       0.684
                                               0.4940
sleephrs_c
female:sleephrs_c -0.4797
                              0.2931 -1.636
                                              0.1021
```

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Residual standard error: 6.31 on 892 degrees of freedom Multiple R-squared: 0.008341, Adjusted R-squared: 0.005006

F-statistic: 2.501 on 3 and 892 DF, p-value: 0.05818

What has changed as compared to the original c2 m4?

- Our original model was bmi = 27.26 + 2.53 female + 0.16 sleephrs 0.48 female x sleephrs
- Our new model is bmi = 28.37 0.84 female + 0.16 centered sleephrs 0.48 female x centered sleephrs.

So our new model on centered data is:

- 28.37 + 0.16 centered sleephrs_c for male subjects, and
- (28.37 0.84) + (0.16 0.48) centered sleephrs_c, or 27.53 0.32 centered sleephrs_c for female subjects.

In our new (centered sleephrs_c) model,

- the main effect of female now corresponds to a predictive difference (female male) in bmi with sleephrs at its mean value, 7.02 hours,
- the intercept term is now the predicted bmi for a male respondent who sleeps an average number of hours, and
- the product term corresponds to the change in the slope of centered sleephrs_c on bmi for a female rather than a male subject, while
- the residual standard deviation and the R-squared values remain unchanged from the model before centering.

2.13.1 Plot of Model 4 on Centered sleephrs: c2_m4_c

```
ggplot(smartcle2_c, aes(x = sleephrs_c, y = bmi, group = female, col = factor(female))) +
    geom_point(alpha = 0.5, size = 2) +
    geom_smooth(method = "lm", se = FALSE) +
    guides(color = FALSE) +
    labs(x = "Sleep Hours, centered", y = "Body Mass Index",
```

```
title = "Model `c2_m4` on centered data") +
facet_wrap(~ female, labeller = label_both)
```

Model `c2_m4` on centered data



2.14 Rescaling an input by subtracting the mean and dividing by 2 standard deviations

Centering helped us interpret the main effects in the regression, but it still leaves a scaling problem.

- The female coefficient estimate is much larger than that of sleephrs, but this is misleading, considering that we are comparing the complete change in one variable (sex = female or not) to a 1-hour change in average sleep.
- Gelman and Hill (2007) recommend all continuous predictors be scaled by dividing by 2 standard deviations, so that:
 - a 1-unit change in the rescaled predictor corresponds to a change from 1 standard deviation below the mean, to 1 standard deviation above.
 - an unscaled binary (1/0) predictor with 50% probability of occurring will be exactly comparable to a rescaled continuous predictor done in this way.

```
smartcle2_rescale <- smartcle2 %>%
mutate(sleephrs_z = (sleephrs - mean(sleephrs))/(2*sd(sleephrs)))
```

2.14.1 Refitting model c2_m4 to the rescaled data

```
c2_m4_z <- lm(bmi ~ female * sleephrs_z, data = smartcle2_rescale)
summary(c2_m4_z)
lm(formula = bmi ~ female * sleephrs_z, data = smartcle2_rescale)
Residuals:
   Min
            10 Median
                             3Q
                                   Max
-15.498 -4.179 -1.035
                          2.830 38.204
Coefficients:
                 Estimate Std. Error t value Pr(>|t|)
(Intercept)
                   28.3681
                              0.3274 86.658
                                                <2e-16 ***
                   -0.8420
                               0.4280 - 1.967
female
                                                0.0495 *
sleephrs_z
                   0.4637
                               0.6778 0.684
                                                0.4940
female:sleephrs_z -1.4173
                              0.8661 - 1.636
                                               0.1021
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 6.31 on 892 degrees of freedom
Multiple R-squared: 0.008341, Adjusted R-squared: 0.005006
F-statistic: 2.501 on 3 and 892 DF, p-value: 0.05818
```

2.14.2 Interpreting the model on rescaled data

What has changed as compared to the original c2_m4?

- Our original model was bmi = 27.26 + 2.53 female + 0.16 sleephrs 0.48 female x sleephrs
- Our model on centered sleephrs was bmi = 28.37 0.84 female + 0.16 centered sleephrs_c 0.48 female x centered sleephrs_c.
- Our new model on rescaled sleephrs is bmi = 28.37 0.84 female + 0.46 rescaled sleephrs_z 1.42 female x rescaled sleephrs_z.

So our rescaled model is:

- 28.37 + 0.46 rescaled sleephrs_z for male subjects, and
- (28.37 0.84) + (0.46 1.42) rescaled sleephrs_z, or 27.53 0.96 rescaled sleephrs_z for female subjects.

In this new rescaled (sleephrs_z) model, then,

- the main effect of female, -0.84, still corresponds to a predictive difference (female male) in bmi with sleephrs at its mean value, 7.02 hours,
- the intercept term is still the predicted bmi for a male respondent who sleeps an average number of hours, and
- the residual standard deviation and the R-squared values remain unchanged,

as before, but now we also have that:

• the coefficient of sleephrs_z indicates the predictive difference in bmi associated with a change in sleephrs of 2 standard deviations (from one standard deviation below the mean of 7.02 to one standard deviation above 7.02.)

- Since the standard deviation of sleephrs is 1.48, this corresponds to a change from 5.54 hours per night to 8.50 hours per night.
- the coefficient of the product term (-1.42) corresponds to the change in the coefficient of sleephrs_z for females as compared to males.

2.14.3 Plot of model on rescaled data

Model `c2_m4_z` on rescaled data



2.15 c2_m5: What if we add more variables?

We can boost our R² a bit, to over 5%, by adding in two new variables, related to whether or not the subject (in the past 30 days) used the internet, and on how many days the subject drank alcoholic beverages.

sleephrs

1

Residuals 890 33928

39

38.12

38.83 1.0186 0.3131176

```
Call:
lm(formula = bmi ~ female + exerany + sleephrs + internet30 +
   alcdays, data = smartcle2)
Residuals:
            10 Median
   Min
                            30
                                   Max
-16.147 -3.997 -0.856
                         2.487 35.965
Coefficients:
           Estimate Std. Error t value Pr(>|t|)
(Intercept) 30.84066 1.18458 26.035 < 2e-16 ***
female
           -1.28801
                       0.42805 -3.009 0.0027 **
           -2.42161
exerany
                       0.49853 -4.858 1.40e-06 ***
                       0.13988 -1.009
sleephrs
           -0.14118
                                        0.3131
internet30 1.38916
                       0.54252
                                2.561
                                        0.0106 *
          -0.10460
                       0.02595 -4.030 6.04e-05 ***
alcdays
---
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 6.174 on 890 degrees of freedom
Multiple R-squared: 0.05258,
                              Adjusted R-squared: 0.04726
F-statistic: 9.879 on 5 and 890 DF, p-value: 3.304e-09
  1. Here's the ANOVA for this model. What can we study with this?
anova(c2_m5)
Analysis of Variance Table
Response: bmi
           Df Sum Sq Mean Sq F value
                                       Pr(>F)
female
               156 155.61 4.0818
                                      0.04365 *
            1
                 897 896.93 23.5283 1.453e-06 ***
exerany
            1
                      32.90 0.8631
sleephrs
            1
                 33
                                      0.35313
internet30
            1
                 178 178.33 4.6779
                                      0.03082 *
                 619 619.26 16.2443 6.044e-05 ***
alcdavs
            1
Residuals 890 33928
                       38.12
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
  2. Consider the revised output below. Now what can we study?
anova(lm(bmi ~ exerany + internet30 + alcdays + female + sleephrs,
        data = smartcle2))
Analysis of Variance Table
Response: bmi
           Df Sum Sq Mean Sq F value
                                        Pr(>F)
                795 795.46 20.8664 5.618e-06 ***
            1
exerany
                 212 211.95 5.5599 0.0185925 *
internet30 1
                 486 486.03 12.7496 0.0003752 ***
alcdays
            1
female
                 351 350.75 9.2010 0.0024891 **
            1
```

2.16 c2_m6: Would adding self-reported health help?

And we can do even a bit better than that by adding in a multi-categorical measure: self-reported general health.

```
Call:
lm(formula = bmi ~ female + exerany + sleephrs + internet30 +
    alcdays + genhealth, data = smartcle2)
Residuals:
```

Min 1Q Median 3Q Max -16.331 -3.813 -0.838 2.679 34.166

Coefficients:

```
Estimate Std. Error t value Pr(>|t|)
(Intercept)
              26.49498 1.31121 20.206 < 2e-16 ***
female
               -1.61968 0.50541 -3.205 0.001400 **
exerany
               -0.12719 0.13613 -0.934 0.350368
sleephrs
internet30
               alcdays
                       0.59408 3.544 0.000415 ***
genhealth2_VeryGood 2.10537
genhealth3_Good 4.08245 0.60739 6.721 3.22e-11 ***
genhealth4_Fair 4.99213 0.80178 6.226 7.37e-10 *** genhealth5_Poor 3.11025 1.12614 2.762 0.005866 **
                         0.80178 6.226 7.37e-10 ***
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

Residual standard error: 5.993 on 886 degrees of freedom

```
Multiple R-squared: 0.1115, Adjusted R-squared: 0.1024 F-statistic: 12.35 on 9 and 886 DF, p-value: < 2.2e-16
```

- 1. If Harry and Marty have the same values of female, exerany, sleephrs, internet30 and alcdays, but Harry rates his health as Good, and Marty rates his as Fair, then what is the difference in the predictions? Who is predicted to have a larger BMI, and by how much?
- 2. What does this normal probability plot of the residuals suggest?

```
plot(c2_m6, which = 2)
```



Theoretical Quantiles
Im(bmi ~ female + exerany + sleephrs + internet30 + alcdays + genhealth)

2.17 c2_m7: What if we added the menthealth variable?

```
Call:
lm(formula = bmi ~ female + exerany + sleephrs + internet30 +
    alcdays + genhealth + physhealth + menthealth, data = smartcle2)
Residuals:
    Min    1Q    Median    3Q    Max
```

```
-16.060 -3.804 -0.890 2.794 33.972
```

Coefficients:

```
Estimate Std. Error t value Pr(>|t|)
(Intercept)
                    25.88208
                               1.31854 19.629 < 2e-16 ***
female
                    -0.96435
                                0.41908 -2.301 0.021616 *
                                0.50635 -2.828 0.004797 **
exerany
                    -1.43171
sleephrs
                    -0.08033
                                0.13624 -0.590 0.555583
internet30
                     2.00267
                                0.53759
                                         3.725 0.000207 ***
alcdays
                    -0.07997
                                0.02528 -3.163 0.001614 **
genhealth2_VeryGood 2.09533
                                0.59238
                                         3.537 0.000425 ***
genhealth3_Good
                     3.90949
                                0.60788
                                          6.431 2.07e-10 ***
genhealth4_Fair
                     4.27152
                                0.83986
                                          5.086 4.47e-07 ***
genhealth5_Poor
                                         0.958 0.338361
                     1.26021
                                1.31556
physhealth
                     0.06088
                                0.03005
                                          2.026 0.043064 *
menthealth
                     0.06636
                                0.03177
                                          2.089 0.037021 *
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

Residual standard error: 5.964 on 884 degrees of freedom Multiple R-squared: 0.1219, Adjusted R-squared: 0.111 F-statistic: 11.16 on 11 and 884 DF, p-value: < 2.2e-16

2.18 Key Regression Assumptions for Building Effective Prediction Models

- 1. Validity the data you are analyzing should map to the research question you are trying to answer.
 - The outcome should accurately reflect the phenomenon of interest.
 - The model should include all relevant predictors. (It can be difficult to decide which predictors are necessary, and what to do with predictors that have large standard errors.)
 - The model should generalize to all of the cases to which it will be applied.
 - Can the available data answer our question reliably?
- 2. Additivity and linearity most important assumption of a regression model is that its deterministic component is a linear function of the predictors. We often think about transformations in this setting.
- 3. Independence of errors errors from the prediction line are independent of each other
- 4. Equal variance of errors if this is violated, we can more efficiently estimate paramaters using *weighted* least squares approaches, where each point is weighted inversely proportional to its variance, but this doesn't affect the coefficients much, if at all.
- 5. Normality of errors not generally important for estimating the regression line

2.18.1 Checking Assumptions in model c2_m7

1. How does the assumption of linearity behind this model look?

 $plot(c2_m7, which = 1)$



Im(bmi ~ female + exerany + sleephrs + internet30 + alcdays + genhealth + p ...

We see no strong signs of serious non-linearity here. There's no obvious curve in the plot, for example.

2. What can we conclude from the plot below?

 $plot(c2_m7, which = 5)$



This plot can help us identify points with large standardized residuals, large leverage values, and large influence on the model (as indicated by large values of Cook's distance.) In this case, I see no signs of any points used in the model with especially large influence, although there are some poorly fitted points (with especially large standardized residuals.)

Chapter 3

Analysis of Variance

3.1 The bonding data: A Designed Dental Experiment

The bonding data describe a designed experiment into the properties of four different resin types (resin = A, B, C, D) and two different curing light sources (light = Halogen, LED) as they relate to the resulting bonding strength (measured in MPa¹) on the surface of teeth. The source is Kim (2014).

The experiment involved making measurements of bonding strength under a total of 80 experimental setups, or runs, with 10 runs completed at each of the eight combinations of a light source and a resin type. The data are gathered in the bonding.csv file.

bonding

```
# A tibble: 80 x 4
   run_ID light
                  resin strength
   <fct> <fct>
                   <fct>
                            <dbl>
 1 R101
          LED
                  В
                             12.8
 2 R102
          Halogen B
                             22.2
 3 R103
          Halogen B
                             24.6
 4 R104
          LED
                             17.0
 5 R105
          LED
                   C
                             32.2
 6 R106
          Halogen B
                             27.1
7 R107
                             23.4
          LED
                   Α
8 R108
                             23.5
          Halogen A
9 R109
          Halogen D
                             37.3
10 R110
          Halogen A
                             19.7
# ... with 70 more rows
```

3.2 A One-Factor Analysis of Variance

Suppose we are interested in the distribution of the strength values for the four different types of resin.

```
bonding %>% group_by(resin) %>% summarize(n = n(), mean(strength), median(strength))
```

¹The MPa is defined as the failure load (in Newtons) divided by the entire bonded area, in mm².

2 B	20	22.2	22.7
3 C	20	25.2	25.7
4 D	20	32.1	35.3

I'd begin serious work with a plot.

3.2.1 Look at the Data!

```
ggplot(bonding, aes(x = resin, y = strength)) +
   geom_boxplot()
```



Another good plot for this purpose is a ridgeline plot.

```
ggplot(bonding, aes(x = strength, y = resin, fill = resin)) +
   geom_density_ridges2() +
   guides(fill = FALSE)
```

Picking joint bandwidth of 3.09



3.2.2 Table of Summary Statistics

bonding %>% group_by(resin) %>% skim(strength)

With the small size of this experiment (n = 20 for each resin type), graphical summaries may not perform as well as they often do. We'll also produce a quick table of summary statistics for strength within each resin type, with the skim() function.

```
Skim summary statistics
n obs: 80
n variables: 4
group variables: resin
Variable type: numeric
resin variable missing complete n mean
                                           sd
                                               p0
                                                     p25 median
    A strength
                     0
                             20 20 18.41 4.81 9.3 15.73 17.95 20.4
    B strength
                     0
                             20 20 22.23 6.75 11.8 18.45
                                                         22.7
                                                                25.75
    C strength
                     0
                             20 20 25.16 6.33 14.5 20.65
                                                         25.7
                                                                30.7
    D strength
                             20 20 32.08 9.74 17.3 21.82 35.3 40.15
p100
28
35.2
34.5
47.2
```

Since the means and medians within each group are fairly close, and the distributions (with the possible exception of resin D) are reasonably well approximated by the Normal, I'll fit an ANOVA model².

```
anova(lm(strength ~ resin, data = bonding))
```

```
Analysis of Variance Table
```

```
Response: strength

Df Sum Sq Mean Sq F value Pr(>F)

resin 3 1999.7 666.57 13.107 5.52e-07 ***

Residuals 76 3865.2 50.86

---

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

It appears that the resin types have a significant association with mean strength of the bonds. Can we identify which resin types have generally higher or lower strength?

```
TukeyHSD(aov(lm(strength ~ resin, data = bonding)))
```

```
Tukey multiple comparisons of means
95% family-wise confidence level

Fit: aov(formula = lm(strength ~ resin, data = bonding))

$resin
diff lwr upr p adj
B-A 3.815 -2.1088676 9.738868 0.3351635
C-A 6.740 0.8161324 12.663868 0.0193344
D-A 13.660 7.7361324 19.583868 0.0000003
C-B 2.925 -2.9988676 8.848868 0.5676635
D-B 9.845 3.9211324 15.768868 0.0002276
D-C 6.920 0.9961324 12.843868 0.0154615
```

Based on these confidence intervals (which have a family-wise 95% confidence level), we see that D is associated with significantly larger mean strength than A or B or C, and that C is also associated with significantly larger mean strength than A. This may be easier to see in a plot of these confidence intervals.

```
plot(TukeyHSD(aov(lm(strength ~ resin, data = bonding))))
```

²If the data weren't approximately Normally distributed, we might instead consider a rank-based alternative to ANOVA, like the Kruskal-Wallis test.

95% family-wise confidence level



Differences in mean levels of resin

3.3 A Two-Way ANOVA: Looking at Two Factors

Now, we'll now add consideration of the light source into our study. We can look at the distribution of the strength values at the combinations of both light and resin, with a plot like this one...

```
ggplot(bonding, aes(x = resin, y = strength, color = light)) +
    geom_point(size = 2, alpha = 0.5) +
    facet_wrap(~ light) +
    guides(color = FALSE) +
    scale_color_manual(values = c("purple", "darkorange")) +
    theme_bw()
```



3.4 A Means Plot (with standard deviations) to check for interaction

Sometimes, we'll instead look at a plot simply of the means (and, often, the standard deviations) of strength at each combination of light and resin. We'll start by building up a data set with the summaries we want to plot.

```
bond.sum <- bonding %>%
    group_by(resin, light) %>%
    summarize(mean.str = mean(strength), sd.str = sd(strength))
bond.sum
# A tibble: 8 x 4
# Groups:
            resin [?]
  resin light
                mean.str sd.str
  <fct> <fct>
                   <dbl>
                          <dbl>
1 A
        Halogen
                    17.8
                           4.02
2 A
        LED
                    19.1
                           5.63
3 B
        Halogen
                    19.9
                           5.62
4 B
        LED
                    24.6
                           7.25
5 C
        Halogen
                    22.5
                           6.19
6 C
        LED
                    27.8
                           5.56
7 D
        Halogen
                    40.3
                           4.15
                    23.8
8 D
        LED
                           5.70
```

Now, we'll use this new data set to plot the means and standard deviations of strength at each combination of resin and light.

Observed Means (+/- SD) of Bonding Strength



Is there evidence of a meaningful interaction between the resin type and the light source on the bonding strength in this plot?

- Sure. A meaningful interaction just means that the strength associated with different resin types depends on the light source.
 - With LED light, it appears that resin C leads to the strongest bonding strength.
 - With Halogen light, though, it seems that resin D is substantially stronger.
- Note that the lines we see here connecting the light sources aren't in parallel (as they would be if we had zero interaction between resin and light), but rather, they cross.

3.4.1 Skimming the data after grouping by resin and light

We might want to look at a numerical summary of the strengths within these groups, too.

```
bonding %>%
   group_by(resin, light) %>%
   skim(strength)
Skim summary statistics
n obs: 80
n variables: 4
group variables: resin, light
Variable type: numeric
resin
      light variable missing complete n mean sd p0 p25 median
    A Halogen strength
                      0 10 10 17.77 4.02 9.3 15.75 18.35
         LED strength
                           0
                                  10 10 19.06 5.63 11.6 16.18 17.8
    B Halogen strength
                           0
                                  10 10 19.9 5.62 11.8 14.78 21.75
         LED strength
                          0
                                  10 10 24.56 7.25 12.8 20.45 24.45
    C Halogen strength
                          0
                                  10 10 22.54 6.19 14.5 18.85 21.3
                         0
          LED strength
                                 10 10 27.77 5.56 16.5 24.7
                                                              28.45
                        0
                                10 10 40.3 4.15 35.5 36.55 40.4
    D Halogen strength
    D
          LED strength
                          0
                                 10 10 23.85 5.7 17.3 19.75 21.45
  p75 p100
 20
      23.5
22.5 28
24.12 27.1
27.87 35.2
25.8 33
31.83 34.5
43.62 47.2
28.2 35.1
```

3.5 Fitting the Two-Way ANOVA model with Interaction

```
c3_m1 <- lm(strength ~ resin * light, data = bonding)</pre>
summary(c3_m1)
lm(formula = strength ~ resin * light, data = bonding)
Residuals:
   Min
            1Q Median
                            3Q
                                  Max
-11.760 -3.663 -0.320
                        3.697 11.250
Coefficients:
               Estimate Std. Error t value Pr(>|t|)
(Intercept)
                 17.770
                            1.771 10.033 2.57e-15 ***
resinB
                  2.130
                            2.505 0.850 0.3979
resinC
                 4.770
                            2.505 1.904 0.0609 .
                            2.505 8.995 2.13e-13 ***
                 22.530
resinD
```

```
lightLED
                   1.290
                              2.505
                                      0.515
                                              0.6081
                                              0.3446
resinB:lightLED
                                      0.951
                   3.370
                              3.542
resinC:lightLED
                  3.940
                              3.542
                                      1.112
                                              0.2697
resinD:lightLED
                -17.740
                              3.542
                                    -5.008 3.78e-06 ***
Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
Residual standard error: 5.601 on 72 degrees of freedom
Multiple R-squared: 0.6149,
                                Adjusted R-squared: 0.5775
F-statistic: 16.42 on 7 and 72 DF, p-value: 9.801e-13
```

3.5.1 The ANOVA table for our model

In a two-way ANOVA model, we begin by assessing the interaction term. If it's important, then our best model is the model including the interaction. If it's not important, we will often move on to consider a new model, fit without an interaction.

The ANOVA table is especially helpful in this case, because it lets us look specifically at the interaction effect.

```
anova(c3_m1)
```

Analysis of Variance Table

```
Response: strength
```

3.5.2 Is the interaction important?

In this case, the interaction:

- is evident in the means plot, and
- is highly statistically significant, and
- accounts for a sizeable fraction (27%) of the overall variation

$$\eta^2_{interaction} = \frac{\text{SS(resin:light)}}{SS(Total)} = \frac{1571.96}{1999.72 + 34.72 + 1571.96 + 2258.52} = 0.268$$

If the interaction were *either* large or significant we would be inclined to keep it in the model. In this case, it's both, so there's no real reason to remove it.

3.5.3 Interpreting the Interaction

Recall the model equation, which is:

```
c3_m1
```

Call:

```
lm(formula = strength ~ resin * light, data = bonding)
```

Coefficients:

```
      (Intercept)
      resinB
      resinC
      resinD

      17.77
      2.13
      4.77
      22.53

      lightLED
      resinB:lightLED
      resinC:lightLED
      resinD:lightLED

      1.29
      3.37
      3.94
      -17.74
```

so we have:

strength = 17.77 + 2.13 resinB + 4.77 resinC + 22.53 resinD + 1.29 lightLED + 3.37 resinB * lightLED + 3.94 resinC * li

So, if light = Halogen, our equation is:

$$strength = 17.77 + 2.13resinB + 4.77resinC + 22.53resinD$$

And if light = LED, our equation is:

```
strength = 19.06 + 5.50resinB + 8.71resinC + 4.79resinD
```

Note that both the intercept and the slopes change as a result of the interaction. The model yields a different prediction for every possible combination of a resin type and a light source.

3.6 Comparing Individual Combinations of resin and light

To make comparisons between individual combinations of a resin type and a light source, using something like Tukey's HSD approach for multiple comparisons, we first refit the model using the aov structure, rather than lm.

```
c3m1_aov <- aov(strength ~ resin * light, data = bonding)
summary(c3m1_aov)</pre>
```

```
Df Sum Sq Mean Sq F value
                                        Pr(>F)
             3 1999.7
                        666.6 21.250 5.79e-10 ***
resin
light
             1
                 34.7
                         34.7
                               1.107
                                         0.296
resin:light 3 1572.0
                        524.0 16.704 2.46e-08 ***
Residuals
           72 2258.5
                         31.4
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

And now, we can obtain Tukey HSD comparisons (which will maintain an overall 95% family-wise confidence level) across the resin types, the light sources, and the combinations, with the TukeyHSD command. This approach is only completely appropriate if these comparisons are pre-planned, and if the design is balanced (as this is, with the same sample size for each combination of a light source and resin type.)

```
TukeyHSD(c3m1_aov)
```

```
Tukey multiple comparisons of means 95% family-wise confidence level
```

Fit: aov(formula = strength ~ resin * light, data = bonding)

```
$resin
      diff
                lwr
                          upr
                                  p adj
B-A 3.815 -0.843129 8.473129 0.1461960
C-A 6.740 2.081871 11.398129 0.0016436
D-A 13.660 9.001871 18.318129 0.0000000
C-B 2.925 -1.733129 7.583129 0.3568373
D-B 9.845 5.186871 14.503129 0.0000026
D-C 6.920 2.261871 11.578129 0.0011731
$light
              diff
                         lwr
                                          p adj
                                  upr
LED-Halogen -1.3175 -3.814042 1.179042 0.2963128
```

\$`resin:light`

	diff	lwr	upr	p adj
B:Halogen-A:Halogen	2.13	-5.68928258	9.949283	0.9893515
${\tt C:Halogen-A:Halogen}$	4.77	-3.04928258	12.589283	0.5525230
D:Halogen-A:Halogen	22.53	14.71071742	30.349283	0.0000000
A:LED-A:Halogen	1.29	-6.52928258	9.109283	0.9995485
B:LED-A:Halogen	6.79	-1.02928258	14.609283	0.1361092
C:LED-A:Halogen	10.00	2.18071742	17.819283	0.0037074
D:LED-A:Halogen	6.08	-1.73928258	13.899283	0.2443200
C:Halogen-B:Halogen	2.64	-5.17928258	10.459283	0.9640100
D:Halogen-B:Halogen	20.40	12.58071742	28.219283	0.0000000
A:LED-B:Halogen	-0.84	-8.65928258	6.979283	0.9999747
B:LED-B:Halogen	4.66	-3.15928258	12.479283	0.5818695
C:LED-B:Halogen	7.87	0.05071742	15.689283	0.0473914
D:LED-B:Halogen	3.95	-3.86928258	11.769283	0.7621860
D:Halogen-C:Halogen	17.76	9.94071742	25.579283	0.0000000
A:LED-C:Halogen	-3.48	-11.29928258	4.339283	0.8591455
B:LED-C:Halogen	2.02	-5.79928258	9.839283	0.9922412
C:LED-C:Halogen	5.23	-2.58928258	13.049283	0.4323859
D:LED-C:Halogen	1.31	-6.50928258	9.129283	0.9995004
A:LED-D:Halogen	-21.24	-29.05928258	-13.420717	0.0000000
B:LED-D:Halogen	-15.74	-23.55928258	-7.920717	0.000006
C:LED-D:Halogen	-12.53	-20.34928258	-4.710717	0.0001014
D:LED-D:Halogen	-16.45	-24.26928258	-8.630717	0.0000002
B:LED-A:LED	5.50	-2.31928258	13.319283	0.3665620
C:LED-A:LED	8.71	0.89071742	16.529283	0.0185285
D:LED-A:LED	4.79	-3.02928258	12.609283	0.5471915
C:LED-B:LED	3.21	-4.60928258	11.029283	0.9027236
D:LED-B:LED	-0.71	-8.52928258	7.109283	0.9999920
D:LED-C:LED	-3.92	-11.73928258	3.899283	0.7690762

One conclusion from this is that the combination of D and Halogen is significantly stronger than each of the other seven combinations.

3.7 The bonding model without Interaction

It seems incorrect in this situation to fit a model without the interaction term, but we'll do so just so you can see what's involved.

```
c3_m2 <- lm(strength ~ resin + light, data = bonding)
summary(c3_m2)</pre>
```

Call:

lm(formula = strength ~ resin + light, data = bonding)

Residuals:

Min 1Q Median 3Q Max -14.1163 -4.9531 0.1187 4.4613 14.4663

Coefficients:

Estimate Std. Error t value Pr(>|t|)
(Intercept) 19.074 1.787 10.676 < 2e-16 ***
resinB 3.815 2.260 1.688 0.09555 .
resinC 6.740 2.260 2.982 0.00386 **
resinD 13.660 2.260 6.044 5.39e-08 ***
lightLED -1.317 1.598 -0.824 0.41229

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Residual standard error: 7.147 on 75 degrees of freedom Multiple R-squared: 0.3469, Adjusted R-squared: 0.312 F-statistic: 9.958 on 4 and 75 DF, p-value: 1.616e-06

In the no-interaction model, if light = Halogen, our equation is:

$$strength = 19.07 + 3.82 resinB + 6.74 resinC + 13.66 resinD$$

And if light = LED, our equation is:

$$strength = 17.75 + 3.82 resinB + 6.74 resinC + 13.66 resinD$$

So, in the no-interaction model, only the intercept changes.

anova(c3_m2)

Analysis of Variance Table

```
Response: strength
```

```
Df Sum Sq Mean Sq F value Pr(>F)
resin 3 1999.7 666.57 13.0514 6.036e-07 ***
light 1 34.7 34.72 0.6797 0.4123
Residuals 75 3830.5 51.07
---
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

And, it appears, if we ignore the interaction, then resin type has a significant impact on strength but light source doesn't. This is clearer when we look at boxplots of the separated light and resin groups.

```
p1 <- ggplot(bonding, aes(x = light, y = strength)) +
    geom_boxplot()
p2 <- ggplot(bonding, aes(x = resin, y = strength)) +
    geom_boxplot()</pre>
```



3.8 cortisol: A Hypothetical Clinical Trial

156 adults who complained of problems with a high-stress lifestyle were enrolled in a hypothetical clinical trial of the effectiveness of a behavioral intervention designed to help reduce stress levels, as measured by salivary cortisol.

The subjects were randomly assigned to one of three intervention groups (usual care, low dose, and high dose.) The "low dose" subjects received a one-week intervention with a follow-up at week 5. The "high dose" subjects received a more intensive three-week intervention, with follow up at week 5.

Since cortisol levels rise and fall with circadian rhythms, the cortisol measurements were taken just after rising for all subjects. These measurements were taken at baseline, and again at five weeks. The difference (baseline - week 5) in cortisol level (in micrograms / l) serves as the primary outcome.

3.8.1 Codebook and Raw Data for cortisol

The data are gathered in the cortisol data set. Included are:

Variable	Description
U	subject identification code intervention group (UC = usual care, Low, High) waist circumference at baseline (in inches)

Variable	Description
	male or female
	salivary cortisol level (microg/l) week 1 salivary cortisol level (microg/l) week 5

```
cortisol
# A tibble: 156 x 6
   subject interv waist sex
                                cort.1 cort.5
     <int> <fct>
                   <dbl> <fct>
                                <dbl>
                                        <dbl>
      1001 UC
                    48.3 M
                                 13.4
                                         13.3
 1
 2
      1002 Low
                    58.3 M
                                 17.8
                                         16.6
 3
      1003 High
                    43.0 M
                                 14.4
                                         12.7
 4
      1004 Low
                    44.9 M
                                  9.00
                                         9.80
 5
      1005 High
                    46.1 M
                                 14.2
                                         14.2
 6
      1006 UC
                    41.3 M
                                 14.8
                                         15.1
 7
      1007 Low
                    51.0 F
                                 13.7
                                        16.0
 8
      1008 UC
                    42.0 F
                                 17.3
                                        18.7
9
      1009 Low
                    24.7 F
                                        15.8
                                 15.3
10
                    59.4 M
                                 12.4
      1010 Low
                                        11.7
# ... with 146 more rows
```

3.9 Creating a factor combining sex and waist

Next, we'll put the waist and sex data in the cortisol example together. We want to build a second categorical variable (called fat_est) combining this information, to indicate "healthy" vs. "unhealthy" levels of fat around the waist.

- Male subjects whose waist circumference is 40 inches or more, and
- Female subjects whose waist circumference is 35 inches or more, will fall in the "unhealthy" group.

```
subject
                interv
                              waist
                                                      cort.1
                                          sex
                                 :20.80
                                                         : 6.000
Min.
       :1001
               High:53
                          Min.
                                          F:83
                                                 Min.
1st Qu.:1040
               Low :52
                          1st Qu.:33.27
                                          M:73
                                                 1st Qu.: 9.675
Median:1078
               UC :51
                          Median :40.35
                                                 Median :12.400
Mean
      :1078
                          Mean
                                 :40.42
                                                         :12.686
                                                 Mean
3rd Qu.:1117
                          3rd Qu.:47.77
                                                 3rd Qu.:16.025
       :1156
                                 :59.90
                                                         :19.000
Max.
                          Max.
                                                 Max.
    cort.5
                    fat_est
                                  cort_diff
     : 4.2
               healthy: 56
                                       :-2.3000
Min.
                                Min.
1st Qu.: 9.6
               unhealthy:100
                                1st Qu.:-0.5000
Median:12.6
                                Median : 0.2000
Mean
     :12.4
                                Mean
                                      : 0.2821
```

```
3rd Qu.:15.7 3rd Qu.: 1.2000 Max. :19.7 Max. : 2.0000
```

3.10 A Means Plot for the cortisol trial (with standard errors)

Again, we'll start by building up a data set with the summaries we want to plot.

```
cort.sum <- cortisol %>%
   group_by(interv, fat_est) %>%
   summarize(mean.cort = mean(cort_diff),
        se.cort = sd(cort_diff)/sqrt(n()))
```

Now, we'll use this new data set to plot the means and standard errors.

Observed Means (+/- SE) of Salivary Cortisol



3.11 A Two-Way ANOVA model for cortisol with Interaction

```
c3_m3 <- lm(cort_diff ~ interv * fat_est, data = cortisol)
anova(c3_m3)</pre>
```

Analysis of Variance Table

```
Response: cort_diff

Df Sum Sq Mean Sq F value Pr(>F)

interv 2 7.847 3.9235 4.4698 0.01301 *

fat_est 1 4.614 4.6139 5.2564 0.02326 *

interv:fat_est 2 0.943 0.4715 0.5371 0.58554

Residuals 150 131.666 0.8778
```

100 101.000 0.0110

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Does it seem like we need the interaction term in this case?

```
summary(c3_m3)
```

```
Call:
lm(formula = cort_diff ~ interv * fat_est, data = cortisol)
```

Residuals:

```
1Q
                Median
                              3Q
-2.62727 -0.75702 0.08636 0.84848 2.12647
```

Coefficients:

```
Estimate Std. Error t value Pr(>|t|)
                        (Intercept)
intervLow
                       -0.1950 0.3001 -0.650 0.51689
                       -0.3479 0.3091 -1.126 0.26206
intervUC
                       -0.3435 0.2655 -1.294 0.19774
fat_estunhealthy
intervLow:fat_estunhealthy 0.1708 0.3785 0.451 0.65256
intervUC:fat_estunhealthy -0.2300 0.3846 -0.598 0.55068
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 0.9369 on 150 degrees of freedom
```

Multiple R-squared: 0.0924, Adjusted R-squared: 0.06214

F-statistic: 3.054 on 5 and 150 DF, p-value: 0.01179

How do you reconcile the apparent difference in significance levels between this regression summary and the ANOVA table above?

A Two-Way ANOVA model for cortisol without Interaction 3.12

3.12.1 The Graph

```
p1 <- ggplot(cortisol, aes(x = interv, y = cort_diff)) +</pre>
    geom_boxplot()
p2 <- ggplot(cortisol, aes(x = fat_est, y = cort_diff)) +</pre>
    geom_boxplot()
gridExtra::grid.arrange(p1, p2, nrow = 1)
```



3.12.2 The ANOVA Model

```
c3_m4 <- lm(cort_diff ~ interv + fat_est, data = cortisol)
anova(c3_m4)</pre>
```

Analysis of Variance Table

```
Response: cort_diff
```

Df Sum Sq Mean Sq F value Pr(>F)
interv 2 7.847 3.9235 4.4972 0.01266 *
fat_est 1 4.614 4.6139 5.2886 0.02283 *

Residuals 152 132.609 0.8724

Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1

How do these results compare to those we saw in the model with interaction?

3.12.3 The Regression Summary

```
summary(c3_m4)
```

```
Call:
lm(formula = cort_diff ~ interv + fat_est, data = cortisol)
```

Residuals:

```
Min 1Q Median 3Q Max -2.55929 -0.74527 0.05457 0.86456 2.05489
```

Coefficients:

```
Estimate Std. Error t value Pr(>|t|)

(Intercept) 0.70452 0.16093 4.378 2.22e-05 ***

intervLow -0.08645 0.18232 -0.474 0.63606

intervUC -0.50063 0.18334 -2.731 0.00707 **

fat_estunhealthy -0.35878 0.15601 -2.300 0.02283 *
---
```

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 0.934 on 152 degrees of freedom

Multiple R-squared: 0.0859, Adjusted R-squared: 0.06785

F-statistic: 4.761 on 3 and 152 DF, p-value: 0.00335

3.12.4 Tukey HSD Comparisons

Without the interaction term, we can make direct comparisons between levels of the intervention, and between levels of the fat_est variable. This is probably best done here in a Tukey HSD comparison.

```
TukeyHSD(aov(cort_diff ~ interv + fat_est, data = cortisol))
```

```
Tukey multiple comparisons of means 95% family-wise confidence level
```

Fit: aov(formula = cort_diff ~ interv + fat_est, data = cortisol)

\$interv

```
diff lwr upr p adj
Low-High -0.09074746 -0.5222655 0.34077063 0.8724916
UC-High -0.51642619 -0.9500745 -0.08277793 0.0150150
UC-Low -0.42567873 -0.8613670 0.01000948 0.0570728
```

\$fat_est

What conclusions can we draw, at a 5% significance level?

Chapter 4

Analysis of Covariance

4.1 An Emphysema Study

My source for this example is Riffenburgh (2006), section 18.4. Serum theophylline levels (in mg/dl) were measured in 16 patients with emphysema at baseline, then 5 days later (at the end of a course of antibiotics) and then at 10 days after baseline. Clinicians anticipate that the antibiotic will increase the theophylline level. The data are stored in the emphysema.csv data file, and note that the age for patient 5 is not available.

4.1.1 Codebook

Variable	Description
patient	ID code
age	patient's age in years
sex	patient's sex (F or M)
st_base	patient's serum theophylline at baseline (mg/dl)
st_day5	patient's serum theophylline at day 5 (mg/dl)
st_day10	patient's serum theophylline at day 10 (mg/dl)

We're going to look at the change from baseline to day 5 as our outcome of interest, since the clinical expectation is that the antibiotic (azithromycin) will increase theophylline levels.

```
emphysema <- emphysema %>%
    mutate(st_delta = st_day5 - st_base)
emphysema
```

```
# A tibble: 16 x 7
              age sex
                         st_base st_day5 st_day10 st_delta
     <int>
            <int> <fct>
                            <dbl>
                                     <dbl>
                                               <dbl>
                                                         <dbl>
          1
               61 F
                            14.1
                                      2.30
                                               10.3
                                                       -11.8
 1
 2
          2
               70 F
                             7.20
                                      5.40
                                                7.30
                                                      - 1.80
 3
          3
               65 M
                            14.2
                                     11.9
                                               11.3
                                                       - 2.30
 4
          4
               65 M
                            10.3
                                     10.7
                                               13.8
                                                         0.400
 5
          5
               NA M
                             9.90
                                     10.7
                                               11.7
                                                         0.800
 6
          6
               76 M
                             5.20
                                      6.80
                                                4.20
                                                         1.60
 7
          7
               72 M
                                               14.1
                                                         4.20
                            10.4
                                     14.6
 8
          8
               69 F
                            10.5
                                      7.20
                                                5.40
                                                      - 3.30
```

```
9
        9
             66 M
                        5.00
                                5.00
                                        5.10
                                                0
10
       10
             62 M
                        8.60
                               8.10
                                        7.40 - 0.500
11
       11
             65 F
                       16.6
                               14.9
                                       13.0
                                              - 1.70
12
       12
             71 M
                       16.4
                               18.6
                                       17.1
                                                2.20
13
       13
             51 F
                       12.2
                               11.0
                                       12.3
                                              - 1.20
14
       14
            71 M
                       6.60
                                        4.50 - 2.90
                               3.70
          64 F
                       15.4
                               15.2
                                       13.6 - 0.200
15
       15
          50 M
                      10.2
                                       11.2
16
       16
                               10.8
                                                0.600
```

4.2 Does sex affect the mean change in the ophylline?

```
emphysema %>% skim(st_delta)
Skim summary statistics
n obs: 16
n variables: 7
Variable type: numeric
variable missing complete n mean sd p0 p25 median p75 p100
                       16 16 -0.99 3.48 -11.8 -1.92 -0.35 0.65 4.2
emphysema %>% group_by(sex) %>% skim(st_delta)
Skim summary statistics
n obs: 16
n variables: 7
group variables: sex
Variable type: numeric
sex variable missing complete n mean sd p0 p25 median
  F st_delta
                   0
                         6 6 -3.33 4.27 -11.8 -2.92 -1.75 -1.32 -0.2
  M st_delta
                   0
                          10 10 0.41 2.07 -2.9 -0.38
                                                        0.5
                                                              1.4
```

Overall, the mean change in the ophylline during the course of the antibiotic is -0.99, but this is -3.33 for female patients and 0.41 for male patients.

A one-way ANOVA model looks like this:

The ANOVA F test finds a statistically significant difference between the mean st_delta among males and the mean st_delta among females. But is there more to the story?

4.3 Is there an association between age and sex in this study?

```
emphysema %>% group_by(sex) %>% skim(age)
Skim summary statistics
n obs: 16
n variables: 7
group variables: sex
Variable type: integer
 sex variable missing complete n mean sd p0 p25 median p75 p100
  F
          age
                    0
                             6 6 63.33 6.89 51 61.75
                                                        64.5
                                                              68
                                                                    70
  М
          age
                             9 10 66.44 7.57 50 65
                                                        66
                                                              71
                                                                    76
```

But we note that the male patients are also older than the female patients, on average (mean age for males is 66.4, for females 63.3)

- Does the fact that male patients are older affect change in the ophylline level?
- And how should we deal with the one missing age value (in a male patient)?

4.4 Adding a quantitative covariate, age, to the model

We could fit an ANOVA model to predict st_delta using sex and age directly, but only if we categorized age into two or more groups. Because age is not categorical, we cannot include it in an ANOVA. But if age is an influence, and we don't adjust for it, it may well bias the outcome of our initial ANOVA. With a quantitative variable like age, we will need a method called ANCOVA, for analysis of covariance.

4.4.1 The ANCOVA model

ANCOVA in this case is just an ANOVA model with our outcome (st_delta) adjusted for a continuous covariate, called age. For the moment, we'll ignore the one subject with missing age and simply fit the regression model with sex and age.

```
summary(lm(st_delta ~ sex + age, data = emphysema))
lm(formula = st_delta ~ sex + age, data = emphysema)
Residuals:
   Min
             1Q Median
                             3Q
                                    Max
-8.3352 -0.4789 0.6948 1.5580 3.5202
Coefficients:
            Estimate Std. Error t value Pr(>|t|)
(Intercept) -6.90266
                        7.92948
                                 -0.871
                                          0.4011
             3.52466
                        1.75815
                                  2.005
                                          0.0681
sexM
             0.05636
                        0.12343
                                  0.457
                                          0.6561
age
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 3.255 on 12 degrees of freedom
  (1 observation deleted due to missingness)
```

```
Multiple R-squared: 0.2882, Adjusted R-squared: 0.1696 F-statistic: 2.43 on 2 and 12 DF, p-value: 0.13
```

This model assumes that the slope of the regression line between st_delta and age is the same for both sexes.

Note that the model yields $st_delta = -6.9 + 3.52$ (sex = male) + 0.056 age, or

- st delta = -6.9 + 0.056 age for female patients, and
- $st_delta = (-6.9 + 3.52) + 0.056$ age = -3.38 + 0.056 age for male patients.

Note that we can test this assumption of equal slopes by fitting an alternative model (with a product term between sex and age) that doesn't require the assumption, and we'll do that later.

4.4.2 The ANCOVA Table

First, though, we'll look at the ANCOVA table.

When we tested sex without accounting for age, we found a p value of 0.032, which is less than our usual cutpoint of 0.05. But when we adjusted for age, we find that sex loses significance, even though age is not a significant influence on st_delta by itself, according to the ANCOVA table.

4.5 Rerunning the ANCOVA model after simple imputation

We could have *imputed* the missing age value for patient 5, rather than just deleting that patient. Suppose we do the simplest potentially reasonable thing to do: insert the mean age in where the NA value currently exists.

```
emph_imp <- replace_na(emphysema, list(age = mean(emphysema$age, na.rm = TRUE)))
emph_imp</pre>
```

```
# A tibble: 16 x 7
                       st_base st_day5 st_day10 st_delta
  patient
             age sex
     <int> <dbl> <fct>
                         <dbl>
                                  <dbl>
                                           <dbl>
                                                    <dbl>
         1 61.0 F
                                   2.30
                                           10.3
 1
                         14.1
                                                  -11.8
 2
         2
            70.0 F
                          7.20
                                  5.40
                                            7.30 - 1.80
 3
         3 65.0 M
                         14.2
                                                  - 2.30
                                 11.9
                                           11.3
 4
           65.0 M
                         10.3
                                           13.8
                                                    0.400
                                 10.7
 5
                                           11.7
         5
           65.2 M
                          9.90
                                 10.7
                                                    0.800
 6
         6
            76.0 M
                          5.20
                                            4.20
                                                    1.60
                                  6.80
7
         7 72.0 M
                         10.4
                                  14.6
                                           14.1
                                                    4.20
8
           69.0 F
                         10.5
                                  7.20
                                                 - 3.30
                                            5.40
 9
           66.0 M
                          5.00
                                  5.00
                                            5.10
                                                    0
```

```
10
        10 62.0 M
                           8.60
                                    8.10
                                              7.40 - 0.500
            65.0 F
11
        11
                          16.6
                                   14.9
                                             13.0
                                                    -1.70
                                   18.6
12
        12
            71.0 M
                           16.4
                                             17.1
                                                       2.20
13
            51.0 F
                          12.2
                                   11.0
                                             12.3
                                                    - 1.20
        13
14
            71.0 M
                           6.60
                                    3.70
                                              4.50
                                                    - 2.90
            64.0 F
15
        15
                          15.4
                                   15.2
                                             13.6
                                                    - 0.200
            50.0 M
16
        16
                          10.2
                                   10.8
                                             11.2
                                                       0.600
```

More on simple imputation and missing data is coming soon.

For now, we can rerun the ANCOVA model on this new data set, after imputation...

```
anova(lm(st_delta ~ sex + age, data = emph_imp))
```

Analysis of Variance Table

```
Response: st_delta

Df Sum Sq Mean Sq F value Pr(>F)

sex 1 52.547 52.547 5.3623 0.03755 *

age 1 2.151 2.151 0.2195 0.64721

Residuals 13 127.392 9.799

---

Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

When we do this, we see that now the sex variable returns to a p value below 0.05. Our complete case analysis (which omitted patient 5) gives us a different result than the ANCOVA based on the data after mean imputation.

4.6 Looking at a factor-covariate interaction

Let's run a model including the interaction (product) term between age and sex, which implies that the slope of age on our outcome (st_delta) depends on the patient's sex. We'll use the imputed data again. Here is the new ANCOVA table, which suggests that the interaction of age and sex is small (because it accounts for only a small amount of the total Sum of Squares) and not significant (p = 0.91).

```
anova(lm(st_delta ~ sex * age, data = emph_imp))
```

Analysis of Variance Table

```
Response: st_delta
         Df Sum Sq Mean Sq F value Pr(>F)
             52.547 52.547 4.9549 0.04594 *
sex
age
              2.151
                      2.151
                            0.2028 0.66051
          1
              0.130
                      0.130
                            0.0123 0.91355
sex:age
Residuals 12 127.261
                    10.605
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

Since the interaction term is neither substantial nor significant, we probably don't need it here. But let's look at its interpretation anyway, just to fix ideas. To do that, we'll need the coefficients from the underlying regression model.

```
3 age 0.03651685 0.2113871 0.1727488 0.8657284
4 sexM:age 0.02885946 0.2603044 0.1108681 0.9135536
```

Our ANCOVA model for st_delta incorporating the age x sex product term is -5.65 + 1.72 (sex = M) + 0.037 age + 0.029 (sex = M)(age). So that means:

- our model for females is $st_delta = -5.65 + 0.037$ age
- our model for males is $st_delta = (-5.65 + 1.72) + (0.037 + 0.029)$ age, or -3.93 + 0.066 age

but, again, our conclusion from the ANCOVA table is that this increase in complexity (letting both the slope and intercept vary by sex) doesn't add much in the way of predictive value for our st_delta outcome.

4.7 Centering the Covariate to Facilitate ANCOVA Interpretation

When developing an ANCOVA model, we will often **center** or even **center and rescale** the covariate to facilitate interpretation of the product term. In this case, let's center **age** and rescale it by dividing by two standard deviations.

```
emph_imp %>% skim(age)
Skim summary statistics
n obs: 16
n variables: 7
Variable type: numeric
variable missing complete n mean
                                     sd p0 p25 median p75 p100
                        16 16 65.2 6.98 50 63.5
                                                   65.1 70.25
Note that in our imputed data, the mean age is 65.2 and the standard deviation of age is 7 years.
So we build the rescaled age variable that I'll call age z, and then use it to refit our model.
emph_imp <- emph_imp %>%
   mutate(age_z = (age - mean(age))/ (2 * sd(age)))
anova(lm(st_delta ~ sex * age_z, data = emph_imp))
Analysis of Variance Table
Response: st_delta
          Df Sum Sq Mean Sq F value Pr(>F)
           1 52.547 52.547 4.9549 0.04594 *
sex
           1
               2.151
                       2.151 0.2028 0.66051
sex:age_z 1
               0.130
                       0.130
                              0.0123 0.91355
Residuals 12 127.261
                     10.605
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
tidy(lm(st_delta ~ sex * age_z, data = emph_imp))
                                                  p.value
              estimate std.error statistic
         term
1 (Intercept) -3.2651685 1.386802 -2.3544587 0.03641637
2
         sexM 3.6019471 1.735706 2.0752055 0.06013138
        age z 0.5096337 2.950144 0.1727488 0.86572835
  sexM:age_z 0.4027661 3.632839 0.1108681 0.91355364
Comparing the two models, we have:
```

• (unscaled): $st_delta = -5.65 + 1.72 (sex = M) + 0.037 age + 0.029 (sex = M) x (age)$

• (rescaled): $st_delta = -3.27 + 3.60$ (sex = M) + 0.510 rescaled $age_z + 0.402$ (sex = M) x (rescaled age_z)

In essence, the rescaled model on age_z is:

- $st_delta = -3.27 + 0.510 age_z$ for female subjects, and
- $st_delta = (-3.27 + 3.60) + (0.510 + 0.402)$ age_z = 0.33 + 0.912 age_z for male subjects

Interpreting the centered, rescaled model, we have:

- no change in the ANOVA results or R-squared or residual standard deviation compared to the uncentered, unscaled model, but
- the intercept (-3.27) now represents the st_delta for a female of average age,
- the sex slope (3.60) represents the (male female) difference in predicted st_delta for a person of average age,
- the age_z slope (0.510) represents the difference in predicted st_delta for a female one standard deviation older than the mean age as compared to a female one standard deviation younger than the mean age, and
- the product term's slope (0.402) represents the male female difference in the slope of age_z, so that if you add the age_z slope (0.510) and the interaction slope (0.402) you see the difference in predicted st_delta for a male one standard deviation older than the mean age as compared to a male one standard deviation younger than the mean age.

Chapter 5

Missing Data Mechanisms and Single Imputation

Almost all serious statistical analyses have to deal with missing data. Data values that are missing are indicated in R, and to R, by the symbol NA.

5.1 A Toy Example

In the following tiny data set called **sbp_example**, we have four variables for a set of 15 subjects. In addition to a subject id, we have:

- the treatment this subject received (A, B or C are the treatments),
- an indicator (1 = yes, 0 = no) of whether the subject has diabetes,
- the subject's systolic blood pressure at baseline
- the subject's systolic blood pressure after the application of the treatment

3	103 C	0	150	150
4	104 A	1.00	NA	120
5	105 C	NA	155	135
6	106 A	1.00	NA	115
7	107 A	0	135	160
8	108 <na></na>	1.00	NA	150
9	109 B	NA	115	130
10	110 C	1.00	170	155
11	111 A	0	150	140
12	112 B	0	145	140
13	113 C	1.00	140	150
14	114 A	1.00	160	135
15	115 B	NA	135	120

5.1.1 How many missing values do we have in each column?

```
colSums(is.na(sbp_example))
subject treat diabetes sbp.before sbp.after
0 1 3 3 0
```

We are missing one treat, 3 diabetes and 3 sbp.before values.

5.1.2 What is the pattern of missing data?

```
mice::md.pattern(sbp_example)
  subject sbp.after treat diabetes sbp.before
9
        1
                   1
                          1
                                   1
3
                          1
                                   0
                                               1 1
2
        1
                                   1
                                               0 1
                   1
                          1
        1
                   1
                          0
                                   1
                                               0 2
                          1
                                   3
                                               3 7
```

We have nine subjects with complete data, three subjects with missing diabetes (only), two subjects with missing sbp.before (only), and 1 subject with missing treat and sbp.before.

5.1.3 How can we identify the subjects with missing data?

```
sbp_example %>% filter(!complete.cases(.))
# A tibble: 6 x 5
  subject treat diabetes sbp.before sbp.after
    <int> <fct>
                   <dbl>
                               <dbl>
                                         <dbl>
                    1.00
1
      104 A
                                 NA
                                           120
2
      105 C
                                 155
                                           135
                   NA
3
      106 A
                    1.00
                                 NA
                                           115
4
                    1.00
                                           150
      108 <NA>
                                 NA
5
      109 B
                   NA
                                 115
                                           130
6
      115 B
                                 135
                                           120
                   NA
```

5.2 Missing-data mechanisms

My source for this description of mechanisms is Chapter 25 of Gelman and Hill (2007), and that chapter is available at this link.

- 1. MCAR = Missingness completely at random. A variable is missing completely at random if the probability of missingness is the same for all units, for example, if for each subject, we decide whether to collect the diabetes status by rolling a die and refusing to answer if a "6" shows up. If data are missing completely at random, then throwing out cases with missing data does not bias your inferences.
- 2. Missingness that depends only on observed predictors. A more general assumption, called missing at random or MAR, is that the probability a variable is missing depends only on available information. Here, we would have to be willing to assume that the probability of nonresponse to diabetes depends only on the other, fully recorded variables in the data. It is often reasonable to model this process as a logistic regression, where the outcome variable equals 1 for observed cases and 0 for missing. When an outcome variable is missing at random, it is acceptable to exclude the missing cases (that is, to treat them as NA), as long as the regression controls for all the variables that affect the probability of missingness.
- 3. Missingness that depends on unobserved predictors. Missingness is no longer "at random" if it depends on information that has not been recorded and this information also predicts the missing values. If a particular treatment causes discomfort, a patient is more likely to drop out of the study. This missingness is not at random (unless "discomfort" is measured and observed for all patients). If missingness is not at random, it must be explicitly modeled, or else you must accept some bias in your inferences.
- 4. Missingness that depends on the missing value itself. Finally, a particularly difficult situation arises when the probability of missingness depends on the (potentially missing) variable itself. For example, suppose that people with higher earnings are less likely to reveal them.

Essentially, situations 3 and 4 are referred to collectively as **non-random missingness**, and cause more trouble for us than 1 and 2.

5.3 Options for Dealing with Missingness

There are several available methods for dealing with missing data that are MCAR or MAR, but they basically boil down to:

- Complete Case (or Available Case) analyses
- Single Imputation
- Multiple Imputation

5.4 Complete Case (and Available Case) analyses

In Complete Case analyses, rows containing NA values are omitted from the data before analyses commence. This is the default approach for many statistical software packages, and may introduce unpredictable bias and fail to include some useful, often hard-won information.

- A complete case analysis can be appropriate when the number of missing observations is not large, and the missing pattern is either MCAR (missing completely at random) or MAR (missing at random.)
- Two problems arise with complete-case analysis:
 - 1. If the units with missing values differ systematically from the completely observed cases, this could bias the complete-case analysis.
 - 2. If many variables are included in a model, there may be very few complete cases, so that most of the data would be discarded for the sake of a straightforward analysis.

• A related approach is *available-case* analysis where different aspects of a problem are studied with different subsets of the data, perhaps identified on the basis of what is missing in them.

5.5 Single Imputation

In **single imputation** analyses, NA values are estimated/replaced *one time* with *one particular data value* for the purpose of obtaining more complete samples, at the expense of creating some potential bias in the eventual conclusions or obtaining slightly *less* accurate estimates than would be available if there were no missing values in the data.

- A single imputation can be just a replacement with the mean or median (for a quantity) or the mode (for a categorical variable.) However, such an approach, though easy to understand, underestimates variance and ignores the relationship of missing values to other variables.
- Single imputation can also be done using a variety of models to try to capture information about the NA values that are available in other variables within the data set.
- The simputation package can help us execute single imputations using a wide variety of techniques, within the pipe approach used by the tidyverse. Another approach I have used in the past is the mice package, which can also perform single imputations.

5.6 Multiple Imputation

Multiple imputation, where NA values are repeatedly estimated/replaced with multiple data values, for the purpose of obtaining mode complete samples *and* capturing details of the variation inherent in the fact that the data have missingness, so as to obtain *more* accurate estimates than are possible with single imputation.

• We'll postpone the discussion of multiple imputation for a while.

5.7 Building a Complete Case Analysis

We can drop all of the missing values from a data set with drop_na or with na.omit or by filtering for complete.cases. Any of these approaches produces the same result - a new data set with 9 rows (after dropping the six subjects with any NA values) and 5 columns.

```
cc.1 <- na.omit(sbp_example)
cc.2 <- sbp_example %>% drop_na
cc.3 <- sbp_example %>% filter(complete.cases(.))
```

5.8 Single Imputation with the Mean or Mode

The most straightforward approach to single imputation is to impute a single summary of the variable, such as the mean, median or mode.

```
skim(sbp_example)

Skim summary statistics
  n obs: 15
  n variables: 5

Variable type: factor
  variable missing complete n n_unique top_counts ordered
```

```
14 15
                                      3 A: 6, B: 4, C: 4, NA: 1
    treat
Variable type: integer
variable missing complete
                           n mean
                                      sd p0
                                               p25 median
  subject
                        15 15
                              108 4.47 101 104.5
Variable type: numeric
   variable missing complete n
                                   mean
                                           sd
                                               p0 p25 median
   diabetes
                  3
                           12 15
                                   0.58
                                       0.51
                                                0
                                                    0
                                                            1
                                                                1
  sbp.after
                  0
                           15 15 136
                                        15.83 105 125
                                                          135 150
                                                                      160
 sbp.before
                  3
                           12 15 143.33 15.72 115 135
                                                          145 151.25
                                                                      170
```

Here, suppose we decide to impute

- sbp.before with the mean (143.33) among non-missing values,
- diabetes with its median (1) among non-missing values, and
- treat with its most common value, or mode (A)

```
# A tibble: 15 x 5
```

```
subject treat diabetes sbp.before sbp.after
     <int> <fct>
                      <dbl>
                                   <dbl>
                                              <dbl>
        101 A
                       1.00
                                     120
                                                 105
 1
 2
        102 B
                       0
                                     145
                                                 135
 3
        103 C
                       0
                                     150
                                                 150
 4
       104 A
                       1.00
                                     143
                                                 120
 5
       105 C
                       1.00
                                     155
                                                 135
 6
       106 A
                       1.00
                                     143
                                                 115
 7
       107 A
                       0
                                     135
                                                 160
 8
        108 A
                        1.00
                                     143
                                                 150
9
        109 B
                       1.00
                                     115
                                                 130
10
        110 C
                        1.00
                                     170
                                                 155
                        0
                                                 140
11
        111 A
                                     150
12
        112 B
                        0
                                     145
                                                 140
13
        113 C
                        1.00
                                     140
                                                 150
14
        114 A
                        1.00
                                     160
                                                 135
        115 B
                        1.00
15
                                     135
                                                 120
```

We could accomplish the same thing with, for example:

5.9 Doing Single Imputation with simputation

Single imputation is a potentially appropriate method when missingness can be assumed to be either completely at random (MCAR) or dependent only on observed predictors (MAR). We'll use the simputation package to accomplish it.

- The simputation vignette is available at https://cran.r-project.org/web/packages/simputation/vignettes/intro.html
- The simputation reference manual is available at https://cran.r-project.org/web/packages/simputation/simputation.pdf

5.9.1 Mirroring Our Prior Approach (imputing means/medians/modes)

Suppose we want to mirror what we did above, simply impute the mean for sbp.before and the median for diabetes again.

```
si.3 <- sbp_example %>%
    impute_lm(sbp.before ~ 1) %>%
    impute_median(diabetes ~ 1) %>%
   replace_na(list(treat = "A"))
si.3
# A tibble: 15 x 5
   subject treat diabetes sbp.before sbp.after
     <int> <fct> <dbl> <dbl>
                                         <dbl>
 1
       101 A
                     1.00
                                120
                                           105
2
       102 B
                     Ω
                                 145
                                           135
 3
       103 C
                     0
                                 150
                                           150
 4
       104 A
                     1.00
                                 143
                                           120
 5
      105 C
                     1.00
                                 155
                                           135
 6
       106 A
                     1.00
                                 143
                                           115
7
       107 A
                     0
                                 135
                                           160
8
       108 A
                     1.00
                                 143
                                           150
9
       109 B
                     1.00
                                 115
                                           130
10
       110 C
                     1.00
                                 170
                                           155
11
       111 A
                     0
                                 150
                                           140
12
       112 B
                     0
                                 145
                                           140
13
       113 C
                     1.00
                                 140
                                           150
14
       114 A
                     1.00
                                 160
                                           135
15
       115 B
                     1.00
                                 135
                                           120
```

5.9.2 Using a model to impute sbp. before and diabetes

Suppose we wanted to use:

- a robust linear model to predict sbp.before missing values, on the basis of sbp.after and diabetes status, and
- a predictive mean matching approach to predict diabetes status, on the basis of sbp.after, and
- a decision tree approach to predict treat status, using all other variables in the data

```
imp.4 <- sbp_example %>%
   impute_rlm(sbp.before ~ sbp.after + diabetes) %>%
   impute_pmm(diabetes ~ sbp.after) %>%
   impute_cart(treat ~ .)
```

A tibble: 15 x 5

	subject	treat	diabetes	sbp.before	sbp.after
*	<int></int>	<fct></fct>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>
1	101	Α	1.00	120	105
2	102	В	0	145	135
3	103	C	0	150	150
4	104	Α	1.00	139	120
5	105	C	1.00	155	135
6	106	Α	1.00	136	115
7	107	Α	0	135	160
8	108	Α	1.00	155	150
9	109	В	1.00	115	130
10	110	C	1.00	170	155
11	111	Α	0	150	140
12	112	В	0	145	140
13	113	C	1.00	140	150
14	114	Α	1.00	160	135
15	115	В	1.00	135	120

Details on the many available methods in simputation are provided in its manual. These include:

- impute_cart uses a Classification and Regression Tree approach for numerical or categorical data. There is also an impute_rf command which uses Random Forests for imputation.
- impute_pmm is one of several "hot deck" options for imputation, this one is predictive mean matching, which can be used with numeric data (only). Missing values are first imputed using a predictive model. Next, these predictions are replaced with the observed values which are nearest to the prediction. Other imputation options in this group include random hot deck, sequential hot deck and k-nearest neighbor imputation.
- impute_rlm is one of several regression imputation methods, including linear models, robust linear models (which use what is called M-estimation to impute numerical variables) and lasso/elastic net/ridge regression models.

The simputation package can also do EM-based multivariate imputation, and multivariate random forest imputation, and several other approaches.

Chapter 6

A Study of Prostate Cancer

6.1 Data Load and Background

The data in prost.csv is derived from Stamey et al. (1989) who examined the relationship between the level of prostate-specific antigen and a number of clinical measures in 97 men who were about to receive a radical prostatectomy. The prost data, as I'll name it in R, contains 97 rows and 11 columns.

prost

# A	tibble:	97 x 3	10							
s	ubject	lpsa	lcavol	lweight	age	bph	svi	lcp	gleason	pgg45
	<int></int>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>	<int></int>	<fct></fct>	<int></int>	<dbl></dbl>	<fct></fct>	<int></int>
1	1	-0.431	-0.580	2.77	50	Low	0	-1.39	6	0
2	2	-0.163	-0.994	3.32	58	Low	0	-1.39	6	0
3	3	-0.163	-0.511	2.69	74	Low	0	-1.39	7	20
4	4	-0.163	-1.20	3.28	58	Low	0	-1.39	6	0
5	5	0.372	0.751	3.43	62	Low	0	-1.39	6	0
6	6	0.765	-1.05	3.23	50	Low	0	-1.39	6	0
7	7	0.765	0.737	3.47	64	${\tt Medium}$	0	-1.39	6	0
8	8	0.854	0.693	3.54	58	High	0	-1.39	6	0
9	9	1.05	-0.777	3.54	47	Low	0	-1.39	6	0
10	10	1.05	0.223	3.24	63	Low	0	-1.39	6	0
#	. with	87 more	e rows							

Note that a related **prost** data frame is also available as part of several R packages, including the **faraway** package, but there is an error in the **lweight** data for subject 32 in those presentations. The value of **lweight** for subject 32 should not be 6.1, corresponding to a prostate that is 449 grams in size, but instead the **lweight** value should be 3.804438, corresponding to a 44.9 gram prostate¹.

I've also changed the gleason and bph variables from their presentation in other settings, to let me teach some additional details.

6.2 Code Book

Variable	Description
subject	subject number (1 to 97)

 $^{^{1}} https://statweb.stanford.edu/\sim tibs/ElemStatLearn/\ attributes\ the\ correction\ to\ Professor\ Stephen\ W.\ Link.$

Variable	Description
lpsa	log(prostate specific antigen in ng/ml), our outcome
lcavol	log(cancer volume in cm ³)
lweight	log(prostate weight, in g)
age	age
bph	benign prostatic hyperplasia amount (Low, Medium, or High)
svi	seminal vesicle invasion $(1 = yes, 0 = no)$
lcp	log(capsular penetration, in cm)
gleason	combined Gleason score $(6, 7, or > 7 here)$
pgg45	percentage Gleason scores 4 or 5

Notes:

- in general, higher levels of PSA are stronger indicators of prostate cancer. An old standard (established almost exclusively with testing in white males, and definitely flawed) suggested that values below 4 were normal, and above 4 needed further testing. A PSA of 4 corresponds to an lpsa of 1.39.
- all logarithms are natural (base e) logarithms, obtained in R with the function log()
- all variables other than subject and lpsa are candidate predictors
- the gleason variable captures the highest combined Gleason score [Scores range (in these data) from 6 (a well-differentiated, or low-grade cancer) to 9 (a high-grade cancer), although the maximum possible score is 10. 6 is the lowest score used for cancerous prostates. As this combination value increases, the rate at which the cancer grows and spreads should increase. This score refers to the combined Gleason grade, which is based on the sum of two areas (each scored 1-5) that make up most of the cancer.] in a biopsy, and higher scores indicate more aggressive cancer cells. It's stored here as 6, 7, or > 7.
- the pgg45 variable captures the percentage of individual Gleason scores [The 1-5 scale for individual biopsies are defined so that 1 indicates something that looks like normal prostate tissue, and 5 indicates that the cells and their growth patterns look very abnormal. In this study, the percentage of 4s and 5s shown in the data appears to be based on 5-20 individual scores in most subjects.] that are 4 or 5, on a 1-5 scale, where higher scores indicate more abnormal cells.

6.3 Additions for Later Use

The code below adds to the prost tibble:

- a factor version of the svi variable, called svi_f, with levels No and Yes,
- a factor version of gleason called gleason_f, with the levels ordered > 7, 7, and finally 6,
- a factor version of bph called bph_f, with levels ordered Low, Medium, High,
- a centered version of lcavol called lcavol_c,
- exponentiated cavol and psa results derived from the natural logarithms lcavol and lpsa.

```
Observations: 97
Variables: 16
$ subject <int> 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 1...
```

```
$ lpsa
          <dbl> -0.4307829, -0.1625189, -0.1625189, -0.1625189, 0.37...
          <dbl> -0.5798185, -0.9942523, -0.5108256, -1.2039728, 0.75...
$ lcavol
$ lweight
          <dbl> 2.769459, 3.319626, 2.691243, 3.282789, 3.432373, 3....
          <int> 50, 58, 74, 58, 62, 50, 64, 58, 47, 63, 65, 63, 63, ...
$ age
$ bph
          <fct> Low, Low, Low, Low, Low, Medium, High, Low, Low...
          $ svi
          <dbl> -1.3862944, -1.3862944, -1.3862944, -1.3862944, -1.3...
$ 1cp
          <fct> 6, 6, 7, 6, 6, 6, 6, 6, 6, 6, 6, 6, 7, 7, 7, 6, 7, 6...
$ gleason
          <int> 0, 0, 20, 0, 0, 0, 0, 0, 0, 0, 0, 30, 5, 5, 0, 30...
$ pgg45
$ svi_f
          $ gleason_f <fct> 6, 6, 7, 6, 6, 6, 6, 6, 6, 6, 6, 6, 7, 7, 7, 6, 7, 6...
          <fct> Low, Low, Low, Low, Low, Medium, High, Low, Low...
$ bph_f
$ lcavol_c <dbl> -1.9298281, -2.3442619, -1.8608352, -2.5539824, -0.5...
          <dbl> 0.56, 0.37, 0.60, 0.30, 2.12, 0.35, 2.09, 2.00, 0.46...
$ cavol
          <dbl> 0.65, 0.85, 0.85, 0.85, 1.45, 2.15, 2.15, 2.35, 2.85...
$ psa
```

6.4 Fitting and Evaluating a Two-Predictor Model

To begin, let's use two predictors (lcavol and svi) and their interaction in a linear regression model that predicts lpsa. I'll call this model c5_prost_A

Earlier, we centered the lcavol values to facilitate interpretation of the terms. I'll use that centered version (called lcavol_c) of the quantitative predictor, and the 1/0 version of the svi variable[^We could certainly use the factor version of svi here, but it won't change the model in any meaningful way. There's no distinction in model *fitting* via lm between a 0/1 numeric variable and a No/Yes factor variable. The factor version of this information will be useful elsewhere, for instance in plotting the model.].

```
c5_prost_A <- lm(lpsa ~ lcavol_c * svi, data = prost)
summary(c5_prost_A)</pre>
```

```
lm(formula = lpsa ~ lcavol_c * svi, data = prost)
Residuals:
   Min
             1Q Median
                             30
                                    Max
-1.6305 -0.5007 0.1266 0.4886 1.6847
Coefficients:
             Estimate Std. Error t value Pr(>|t|)
(Intercept)
              2.33134
                         0.09128 25.540 < 2e-16 ***
lcavol_c
              0.58640
                         0.08207
                                   7.145 1.98e-10 ***
svi
              0.60132
                         0.35833
                                   1.678
                                           0.0967
lcavol_c:svi 0.06479
                         0.26614
                                   0.243
                                           0.8082
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 0.7595 on 93 degrees of freedom
Multiple R-squared: 0.5806,
                                Adjusted R-squared: 0.5671
F-statistic: 42.92 on 3 and 93 DF, p-value: < 2.2e-16
```

6.4.1 Using tidy

It can be very useful to build a data frame of the model's results. We can use the tidy function in the broom package to do so.

```
tidy(c5_prost_A)
```

```
term estimate std.error statistic p.value
(Intercept) 2.33134409 0.09128253 25.5398727 8.246849e-44

cappa lcavol_c 0.58639599 0.08206929 7.1451331 1.981492e-10

svi 0.60131973 0.35832695 1.6781314 9.667899e-02

lcavol_c:svi 0.06479298 0.26614194 0.2434527 8.081909e-01
```

This makes it much easier to pull out individual elements of the model fit.

For example, to specify the coefficient for svi, rounded to three decimal places, I could use tidy(c5_prost_A) %>% filter(term == "svi") %>% select(estimate) %>% round(., 3)

- The result is 0.601.
- If you look at the Markdown file, you'll see that the number shown in the bullet point above this one was generated using inline R code, and the function specified above.

6.4.2 Interpretation

- 1. The intercept, 2.33, for the model is the predicted value of lpsa when lcavol is at its average and there is no seminal vesicle invasion (e.g. svi = 0).
- 2. The coefficient for lcavol_c, 0.59, is the predicted change in lpsa associated with a one unit increase in lcavol (or lcavol_c) when there is no seminal vesicle invasion.
- 3. The coefficient for svi, 0.60, is the predicted change in lpsa associated with having no svi to having an svi while the lcavol remains at its average.
- 4. The coefficient for lcavol_c:svi, the product term, which is 0.06, is the difference in the slope of lcavol_c for a subject with svi as compared to one with no svi.

Note: If you look at the R Markdown, you'll notice that in bullet point 3, I didn't use round to round off the estimate (as I did in the other three bullets), but instead a special function I specified at the start of the R Markdown file called specify_decimal() which uses the format function. This forces, in this case, the trailing zero in the two decimal representation of the svi coefficient to be shown. The special function, again, is:

```
specify_decimal <- function(x, k) format(round(x, k), nsmall=k)</pre>
```

6.5 Exploring Model c5_prost_A

The glance function from the broom package builds a nice one-row summary for the model.

```
glance(c5_prost_A)
```

```
r.squared adj.r.squared sigma statistic p.value df logLik
1 0.5806435 0.5671158 0.7594785 42.92278 1.678836e-17 4 -108.9077
AIC BIC deviance df.residual
1 227.8153 240.6889 53.64311 93
```

This summary includes, in order,

- the model R^2 , adjusted R^2 and $\hat{\sigma}$, the residual standard deviation,
- the ANOVA F statistic and associated p value,
- the number of degrees of freedom used by the model, and its log-likelihood ratio
- the model's AIC (Akaike Information Criterion) and BIC (Bayesian Information Criterion)

the model's deviance statistic and residual degrees of freedom

6.5.1 summary for Model c5_prost_A

If necessary, we can also run summary on this $c5_prost_A$ object to pick up some additional summaries. Since the svi variable is binary, the interaction term is, too, so the t test here and the F test in the ANOVA yield the same result.

```
summary(c5_prost_A)
```

```
Call:
lm(formula = lpsa ~ lcavol_c * svi, data = prost)
Residuals:
   Min
            1Q Median
                                   Max
-1.6305 -0.5007 0.1266 0.4886
                               1.6847
Coefficients:
            Estimate Std. Error t value Pr(>|t|)
                        0.09128 25.540 < 2e-16 ***
(Intercept)
             2.33134
lcavol_c
             0.58640
                        0.08207
                                  7.145 1.98e-10 ***
svi
             0.60132
                        0.35833
                                  1.678
                                          0.0967
lcavol_c:svi 0.06479
                        0.26614
                                  0.243
                                          0.8082
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
Residual standard error: 0.7595 on 93 degrees of freedom
Multiple R-squared: 0.5806,
                               Adjusted R-squared: 0.5671
F-statistic: 42.92 on 3 and 93 DF, p-value: < 2.2e-16
```

If you've forgotten the details of the pieces of this summary, review the Part C Notes from 431.

6.5.2 Adjusted R^2

 \mathbb{R}^2 is greedy.

- R² will always suggest that we make our models as big as possible, often including variables of dubious predictive value.
- As a result, there are various methods for penalizing R² so that we wind up with smaller models.
- The adjusted \mathbb{R}^2 is often a useful way to compare multiple models for the same response.
 - $-R_{adj}^2 = 1 \frac{(1-R^2)(n-1)}{n-k}$, where n = the number of observations and k is the number of coefficients estimated by the regression (including the intercept and any slopes).
 - So, in this case, $R_{adj}^2 = 1 \frac{(1-0.5806)(97-1)}{97-4} = 0.5671$
 - The adjusted R² value is not, technically, a proportion of anything, but it is comparable across models for the same outcome.
 - The adjusted R^2 will always be less than the (unadjusted) R^2 .

6.5.3 Coefficient Confidence Intervals

Here are the 90% confidence intervals for the coefficients in Model A. Adjust the level to get different intervals.

What can we conclude from this about the utility of the interaction term?

6.5.4 ANOVA for Model c5_prost_A

The interaction term appears unnecessary. We might wind up fitting the model without it. A complete ANOVA test is available, including a p value, if you want it.

```
anova(c5_prost_A)
```

Analysis of Variance Table

```
Response: lpsa
            Df Sum Sq Mean Sq F value
                                         Pr(>F)
             1 69.003 69.003 119.6289 < 2.2e-16 ***
lcavol c
             1 5.237
                       5.237
                               9.0801 0.003329 **
svi
lcavol_c:svi 1 0.034
                        0.034
                               0.0593 0.808191
Residuals
            93 53.643
                        0.577
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

Note that the anova approach for a lm object is sequential. The first row shows the impact of lcavol_c as compared to a model with no predictors (just an intercept). The second row shows the impact of adding svi to a model that already contains lcavol_c. The third row shows the impact of adding the interaction (product) term to the model with the two main effects. So the order in which the variables are added to the regression model matters for this ANOVA. The F tests here describe the incremental impact of each covariate in turn.

6.5.5 Residuals, Fitted Values and Standard Errors with augment

The augment function in the broom package builds a data frame including the data used in the model, along with predictions (fitted values), residuals and other useful information.

```
c5_prost_A_frame <- augment(c5_prost_A) %>% tbl_df
skim(c5_prost_A_frame)
Skim summary statistics
n obs: 97
n variables: 10
Variable type: integer
                                     sd p0 p25 median p75 p100
variable missing complete n mean
                        97 97 0.22 0.41 0
      svi
Variable type: numeric
   variable missing complete n
                                                              p25 median
                                    mean
                                             sd
                                                      p0
                                         0.02
                                                          0.00078 0.0035
    .cooksd
                  0
                          97 97 0.011
                                                 6.9e-06
    .fitted
                  0
                          97 97
                                 2.48
                                         0.88
                                                 0.75
                                                           1.84
```

```
0
                          97 97 0.041
                                          0.041
                                                   0.013
                                                            0.016
                                                                    0.025
      .hat
                  0
                                                           -0.5
                          97 97 -6.9e-17 0.75
                                                 -1.63
                                                                    0.13
    .resid
   .se.fit
                  0
                                 0.14
                                          0.061
                                                   0.087
                                                            0.095
                                                                    0.12
                  0
                          97 97
                                 0.76
                                          0.0052 0.74
                                                            0.76
                                                                    0.76
    .sigma
                                                 -2.19
.std.resid
                  0
                             97
                                 0.0012 1.01
                                                           -0.69
                                                                    0.17
  lcavol c
                          97 97
                                                                    0.097
                  0
                                 5.4e-17 1.18
                                                 -2.7
                                                           -0.84
      lpsa
                                 2.48
                                          1.15
                                                 -0.43
                                                            1.73
                                                                    2.59
  p75 p100
0.01
     0.13
3.07 4.54
0.049 0.25
0.49 1.68
0.17 0.38
0.76 0.76
0.65 2.26
0.78
      2.47
3.06 5.58
```

Elements shown here include:

- .fitted Fitted values of model (or predicted values)
- .se.fit Standard errors of fitted values
- .resid Residuals (observed fitted values)
- .hat Diagonal of the hat matrix (these indicate *leverage* points with high leverage indicate unusual combinations of predictors values more than 2-3 times the mean leverage are worth some study leverage is always between 0 and 1, and measures the amount by which the predicted value would change if the observation's y value was increased by one unit a point with leverage 1 would cause the line to follow that point perfectly)
- .sigma Estimate of residual standard deviation when corresponding observation is dropped from model
- .cooksd Cook's distance, which helps identify influential points (values of Cook's d > 0.5 may be influential, values > 1.0 almost certainly are an influential point changes the fit substantially when it is removed from the data)
- .std.resid Standardized residuals (values above 2 in absolute value are worth some study treat these as normal deviates [Z scores], essentially)

See ?augment.lm in R for more details.

6.5.6 Making Predictions with c5_prost_A

Suppose we want to predict the lpsa for a patient with cancer volume equal to this group's mean, for both a patient with and without seminal vesicle invasion, and in each case, we want to use a 90% prediction interval?

```
newdata <- data.frame(lcavol_c = c(0,0), svi = c(0,1))
predict(c5_prost_A, newdata, interval = "prediction", level = 0.90)

fit    lwr    upr
1 2.331344 1.060462 3.602226
2 2.932664 1.545742 4.319586</pre>
```

Since the predicted value in fit refers to the natural logarithm of PSA, to make the predictions in terms of PSA, we would need to exponentiate. The code below will accomplish that task.

```
pred <- predict(c5_prost_A, newdata, interval = "prediction", level = 0.90)
exp(pred)</pre>
```

```
fit lwr upr
```

```
1 10.29177 2.887706 36.67978
2 18.77758 4.691450 75.15750
```

6.6 Plotting Model c5_prost_A

6.6.0.1 Plot logs conventionally

Here, we'll use ggplot2 to plot the logarithms of the variables as they came to us, on a conventional coordinate scale. Note that the lines are nearly parallel. What does this suggest about our Model A?

```
ggplot(prost, aes(x = lcavol, y = lpsa, group = svi_f, color = svi_f)) +
    geom_point() +
    geom_smooth(method = "lm", se = FALSE) +
    scale_color_discrete(name = "Seminal Vesicle Invasion?") +
    theme_bw() +
    labs(x = "Log (cancer volume, cc)",
        y = "Log (Prostate Specific Antigen, ng/ml)",
        title = "Two Predictor Model c5_prost_A, including Interaction")
```

Two Predictor Model c5_prost_A, including Interaction



6.6.0.2 Plot on log-log scale

Another approach (which might be easier in some settings) would be to plot the raw values of Cancer Volume and PSA, but use logarithmic axes, again using the natural (base e) logarithm, as follows. If we use the default choice with 'trans = "log", we'll find a need to select some useful break points for the grid, as I've done in what follows.

Two Predictor Model c5_prost_A, including Interaction



I've used the break point of 4 on the Y axis because of the old rule suggesting further testing for asymptomatic men with PSA of 4 or higher, but the other break points are arbitrary - they seemed to work for me, and used round numbers.

6.6.1 Residual Plots of c5_prost_A

```
plot(c5_prost_A, which = 1)
```



plot(c5_prost_A, which = 5)



6.7 Cross-Validation of Model c5_prost_A

Suppose we want to evaluate whether our model c5_prost_A predicts effectively in new data.

One approach (used, for instance, in 431) would be to split our sample into a separate training (perhaps 70% of the data) and test (perhaps 30% of the data) samples, and then:

- 1. fit the model in the training sample,
- 2. use the resulting model to make predictions for lpsa in the test sample, and
- 3. evaluate the quality of those predictions, perhaps by comparing the results to what we'd get using
 a different model.

One problem with this approach is that with a small data set like this, we may be reluctant to cut our sample size for the training or the testing down because we're afraid that our model building and testing will be hampered by a small sample size. A potential solution is the idea of **cross-validation**, which involves partitioning our data into a series of training-test subsets, multiple times, and then combining the results.

The rest of this section is built on some material by David Robinson at https://rpubs.com/dgrtwo/cv-modelr.

Suppose that we want to perform what is called 10-crossfold separation. In words, this approach splits the 97 observations in our prost data frame into 10 exclusive partitions of about 90% (so about 87-88 observations) into a training sample, and the remaining 10% (9-10 observations) in a test sample². We then refit a model of interest using the training data, and fit the resulting model on the test data using the broom package's augment function. This process is then repeated (a total of 10 times) so that each observation is used 9 times in the training sample, and once in the test sample.

²If we did 5-crossfold validation, we'd have 5 partitions into samples of 80% training and 20% test samples.

To code this in R, we'll make use of a few new ideas. Our goal will be to cross-validate model c5_prost_A, which, you'll recall, uses lcavol_c, svi and their interaction, to predict lpsa in the prost data.

- 1. First, we set a seed for the validation algorithm, so we can replicate our results down the line.
- 2. Then we use the <code>crossv_kfold</code> function from the modelr package to split the <code>prost</code> data into ten different partitions, and then use each partition for a split into training and test samples, which the machine indexes with train and test.
- 3. Then we use some magic and the map function from the purrr package (part of the core tidyverse) to fit a new lm(lpsa ~ lcavol_c * svi) model to each of the training samples generated by crossv_kfold.
- 4. Finally, some additional magic with the unnest and map2 functions applies each of these new models to the appropriate test sample, and generate predictions (.fitted) and standard errors for each prediction (.se.fit).

```
set.seed(4320308)

prost_models <- prost %>%
    crossv_kfold(k = 10) %>%
    mutate(model = map(train, ~ lm(lpsa ~ lcavol_c * svi, data = .)))

prost_predictions <- prost_models %>%
    unnest(map2(model, test, ~ augment(.x, newdata = .y)))

head(prost_predictions)
```

```
# A tibble: 6 x 19
  .id
       subject
                         lcavol lweight
                                                               lcp gleason
                  lpsa
                                          age bph
                                                       svi
  <chr>
          <int>
                <dbl>
                          <dbl>
                                  <dbl> <int> <fct>
                                                     <int>
                                                           <dbl> <fct>
1 01
              3 -0.163 -0.511
                                   2.69
                                           74 Low
                                                         0 -1.39 7
2 01
             12 1.27 -1.35
                                   3.60
                                           63 Medium
                                                         0 -1.39
3 01
                                                         0 -1.39 6
                                   3.06
             16 1.45
                        1.54
                                           66 Low
4 01
             18 1.49
                        2.29
                                   3.65
                                           66 Low
                                                         0 0.372 6
             30 1.89
                                                         0 1.62 6
5 01
                        2.41
                                   3.38
                                           65 Low
             34 2.02
                        0.00995
                                   3.27
                                           54 Low
                                                         0 -1.39 6
 ... with 9 more variables: pgg45 <int>, svi f <fct>, gleason f <fct>,
   bph f <fct>, lcavol c <dbl>, cavol <dbl>, psa <dbl>, .fitted <dbl>,
    .se.fit <dbl>
```

The results are a set of predictions based on the splits into training and test groups (remember there are 10 such splits, indexed by .id) that describe the complete set of 97 subjects again.

6.7.1 Cross-Validated Summaries of Prediction Quality

Now, we can calculate the root Mean Squared Prediction Error (RMSE) and Mean Absolute Prediction Error (MAE) for this modeling approach (using lcavol_c and svi to predict lpsa) across these observations.

For now, we'll compare our model to the "intercept only" model that simply predicts the mean lpsa across all patients.

So our model looks meaningfully better than the "intercept only" model, in that both the RMSE and MAE are much lower (better) with our model.

Another thing we could do with this tibble of predictions we have created is to graph the size of the prediction errors (observed lpsa minus predicted values in .fitted) that our modeling approach makes.

```
prost_predictions %>%
    mutate(errors = lpsa - .fitted) %>%
    ggplot(., aes(x = errors)) +
    geom_histogram(bins = 30, fill = "darkviolet", col = "yellow") +
    labs(title = "Cross-Validated Errors in Prediction of log(PSA)",
        subtitle = "Using a model (`c5_prostA`) including lcavol_c and svi and their interaction",
        x = "Error in predicting log(PSA)")
```

Cross–Validated Errors in Prediction of log(PSA)

Using a model ('c5_prostA') including lcavol_c and svi and their interaction



This suggests that some of our results are off by quite a bit, on the log(PSA) scale, which is summarized for the original data below.

```
prost %>% skim(lpsa)
```

Skim summary statistics

```
n obs: 97
n variables: 16

Variable type: numeric
variable missing complete n mean sd p0 p25 median p75 p100
lpsa 0 97 97 2.48 1.15 -0.43 1.73 2.59 3.06 5.58
```

If we like, we could transform the predictions and observed values back to the scale of PSA (unlogged) and then calculate and display errors, as follows:

```
prost_predictions %>%
    mutate(err.psa = exp(lpsa) - exp(.fitted)) %>%
    ggplot(., aes(x = err.psa)) +
    geom_histogram(bins = 30, fill = "darkorange", col = "yellow") +
    labs(title = "Cross-Validated Errors in Prediction of PSA",
        subtitle = "Using a model (`c5_prostA`) including lcavol_c and svi and their interaction",
        x = "Error in predicting PSA")
```

Cross-Validated Errors in Prediction of PSA

Using a model (`c5_prostA`) including lcavol_c and svi and their interaction



This suggests that some of our results are off by quite a bit, on the original scale of PSA, which is summarized below.

```
prost %>% mutate(psa = exp(lpsa)) %>% skim(psa)

Skim summary statistics
  n obs: 97
  n variables: 16

Variable type: numeric
  variable missing complete n mean sd p0 p25 median p75 p100
```

psa 0 97 97 23.74 40.83 0.65 5.65 13.35 21.25 265.85

We'll return to the notion of cross-validation again, but for now, let's consider the problem of considering adding more predictors to our model, and then making sensible selections as to which predictors actually should be incorporated.

Chapter 7

Stepwise Variable Selection

7.1 Strategy for Model Selection

Ramsey and Schafer (2002) suggest a strategy for dealing with many potential explanatory variables should include the following elements:

- 1. Identify the key objectives.
- Screen the available variables, deciding on a list that is sensitive to the objectives and excludes obvious redundancies.
- 3. Perform exploratory analysis, examining graphical displays and correlation coefficients.
- 4. Perform transformations, as necessary.
- 5. Examine a residual plot after fitting a rich model, performing further transformations and considering outliers.
- 6. Find a suitable subset of the predictors, exerting enough control over any semi-automated selection procedure to be sensitive to the questions of interest.
- 7. Proceed with the analysis, using the selected explanatory variables.

The Two Key Aspects of Model Selection are:

- 1. Evaluating each potential subset of predictor variables
- 2. Deciding on the collection of potential subsets

7.1.1 How Do We Choose Potential Subsets of Predictors?

Choosing potential subsets of predictor variables usually involves either:

- 1. Stepwise approaches
- 2. All possible subset (or best possible subset) searches

Note that the use of any variable selection procedure changes the properties of ...

- the estimated coefficients, which are biased, and
- the associated tests and confidence intervals, which are overly optimistic.

Leeb and Potscher (2005) summarize the key issues:

1. Regardless of sample size, the model selection step typically has a dramatic effect on the sampling properties of the estimators that cannot be ignored. In particular, the sampling properties of post-model-selection estimators are typically significantly different from the nominal distributions that arise if a fixed model is supposed.

2. As a consequence, use of inference procedures that do not take into account the model selection step (e.g. using standard t-intervals as if the selected model has been given prior to the statistical analysis) can be highly misleading.

7.2 A "Kitchen Sink" Model (Model c5_prost_ks)

Suppose that we now consider a model for the prost data we have been working with, which includes main effects (and, in this case, only the main effects) of all eight candidate predictors for lpsa, as follows.

```
c5_prost_ks <- lm(lpsa ~ lcavol + lweight + age + bph_f + svi_f +
                lcp + gleason f + pgg45, data = prost)
tidy(c5_prost_ks)
          term
                   estimate
                              std.error statistic
                                                        p.value
   (Intercept) 0.169937821 0.931332512
1
                                        0.1824674 8.556454e-01
2
       lcavol 0.544313829 0.087979210 6.1868461 2.010505e-08
3
       lweight 0.702237531 0.203013089 3.4590751 8.455164e-04
4
           age -0.023857982 0.011081414 -2.1529727 3.412099e-02
5
  bph fMedium 0.364036274 0.182575941
                                        1.9938896 4.933267e-02
6
    bph_fHigh 0.248789989 0.195975792 1.2694935 2.076898e-01
7
      svi fYes 0.710949408 0.241990241 2.9379259 4.240326e-03
8
           lcp -0.119311781 0.089458946 -1.3337043 1.858223e-01
9
    gleason_f7  0.220746268  0.343065609  0.6434520  5.216430e-01
   gleason_f6 -0.053096704 0.430098039 -0.1234526 9.020368e-01
         pgg45 0.003984574 0.004146495 0.9609499 3.392714e-01
glance(c5_prost_ks)
  r.squared adj.r.squared
                              sigma statistic
                                                   p.value df
                                                                 logLik
1 0.6790343
                0.6417127 0.6909479 18.19414 2.373796e-17 11 -95.93939
       AIC
                BIC deviance df.residual
1 215.8788 246.7753 41.05718
```

We'll often refer to this (all predictors on board) approach as a "kitchen sink" model [This refers to the English idiom "... everything but the kitchen sink" which describes, essentially, everything imaginable. A "kitchen sink regression" is often used as a pejorative term, since no special skill or insight is required to identify it, given a list of potential predictors. For more, yes, there is a Wikipedia page.].

7.3 Sequential Variable Selection: Stepwise Approaches

- Forward Selection
 - We begin with a constant mean and then add potential predictors one at a time according to some criterion (R defaults to minimizing the Akaike Information Criterion) until no further addition significantly improves the fit.
 - Each categorical factor variable is represented in the regression model as a set of indicator variables.
 In the absence of a good reason to do something else, the set is added to the model as a single unit, and R does this automatically.
- Backwards Elimination
 - Start with the "kitchen sink" model and then delete potential predictors one at a time.
 - Backwards Elimination is less likely than Forward Selection, to omit negatively confounded sets of variables, though all stepwise procedures have problems.
- Stepwise Regression can also be done by combining these methods.

7.3.1 The Big Problems with Stepwise Regression

There is no reason to assume that a single best model can be found.

- The use of forward selection, or backwards elimination, or stepwise regression including both procedures, will NOT always find the same model.
- It also appears to be essentially useless to try different stepwise methods to look for agreement.

Users of stepwise regression frequently place all of their attention on the particular explanatory variables included in the resulting model, when there's **no reason** (in most cases) to assume that model is in any way optimal.

Despite all of its problems, let's use stepwise regression to help predict lpsa given a subset of the eight predictors in c5_prost_ks.

7.4 Forward Selection with the step function

- 1. Specify the null model (intercept only)
- 2. Specify the variables R should consider as predictors (in the scope element of the step function)
- 3. Specify forward selection only
- 4. R defaults to using AIC as its stepwise criterion

```
Df Sum of Sq
                             {\tt RSS}
                                     AIC
                 69.003
                         58.915 -44.366
+ lcavol
            1
+ svi_f
             1
                  41.011 86.907 -6.658
                  38.528 89.389
                                -3.926
+ lcp
                  30.121 97.796
+ gleason_f 2
                                 6.793
+ lweight
            1
                 24.019 103.899 10.665
+ pgg45
             1
                 22.814 105.103 11.783
+ age
             1
                  3.679 124.239 28.007
                         127.918 28.838
<none>
+ bph_f
                  4.681 123.237 29.221
```

Step: AIC=-44.37
lpsa ~ lcavol

```
Df Sum of Sq
                            RSS
                                     AIC
                  7.1726 51.742 -54.958
+ lweight
             1
+ svi_f
             1
                  5.2375 53.677 -51.397
+ bph_f
             2
                  3.2994 55.615 -45.956
                  1.6980 57.217 -45.203
+ pgg45
             1
                  2.7834 56.131 -45.061
+ gleason_f 2
<none>
                         58.915 -44.366
+ lcp
             1
                  0.6562 58.259 -43.452
                  0.0025 58.912 -42.370
+ age
             1
```

```
Step: AIC=-54.96
lpsa ~ lcavol + lweight
                           RSS
           Df Sum of Sq
                                   AIC
+ svi_f
            1
               5.1737 46.568 -63.177
               1.8158 49.926 -56.424
            1
+ pgg45
+ gleason_f 2 2.6770 49.065 -56.111
<none>
                        51.742 -54.958
               0.8187 50.923 -54.506
+ lcp
            1
+ age
            1
               0.6456 51.097 -54.176
+ bph_f
            2
               1.4583 50.284 -53.731
Step: AIC=-63.18
lpsa ~ lcavol + lweight + svi_f
            Df Sum of Sq
                           RSS
                                   AIC
                        46.568 -63.177
<none>
+ gleason_f 2 1.60467 44.964 -62.579
           1 0.62301 45.945 -62.484
+ age
            2 1.50046 45.068 -62.354
+ bph_f
+ pgg45
            1 0.50069 46.068 -62.226
            1 0.06937 46.499 -61.322
+ lcp
lm(formula = lpsa ~ lcavol + lweight + svi_f)
Coefficients:
                             lweight
(Intercept)
                                         svi_fYes
                 lcavol
   -0.7772
                 0.5259
                              0.6618
                                           0.6657
The resulting model, arrived at after three forward selection steps, includes lcavol, lweight and svi_f.
model.fs <- lm(lpsa ~ lcavol + lweight + svi_f,
               data=prost)
summary(model.fs)$adj.r.squared
[1] 0.6242063
extractAIC(model.fs)
[1]
      4.00000 -63.17744
```

The adjusted R^2 value for this model is 0.624, and the AIC value used by the stepwise procedure is -63.18, on 4 effective degrees of freedom.

7.5 Backward Elimination using the step function

In this case, the backward elimination approach, using reduction in AIC for a criterion, comes to the same conclusion about the "best" model.

Start: AIC=-61.4

```
lpsa ~ lcavol + lweight + age + bph_f + svi_f + lcp + gleason_f +
   pgg45
           Df Sum of Sq RSS
- gleason_f 2 1.1832 42.240 -62.639
- pgg45 1 0.4409 41.498 -62.359
          1 0.8492 41.906 -61.409
- lcp
<none>
                       41.057 -61.395
- bph_f 2 2.0299 43.087 -60.714
          1 2.2129 43.270 -58.303
- age
- svi_f 1 4.1207 45.178 -54.118
- lweight 1 5.7123 46.769 -50.760
- lcavol 1 18.2738 59.331 -27.683
Step: AIC=-62.64
lpsa ~ lcavol + lweight + age + bph_f + svi_f + lcp + pgg45
         Df Sum of Sq RSS
          1 0.8470 43.087 -62.713
- lcp
<none>
                  42.240 -62.639
- pgg45 1 1.2029 43.443 -61.916
- bph_f 2 2.2515 44.492 -61.602
        1 2.0730 44.313 -59.992
- age
- svi_f 1 4.6431 46.884 -54.523
- lweight 1 5.5988 47.839 -52.566
- lcavol 1 21.4956 63.736 -24.736
Step: AIC=-62.71
lpsa ~ lcavol + lweight + age + bph_f + svi_f + pgg45
         Df Sum of Sq
                      RSS
                               AIC
          1 0.5860 43.673 -63.403
- pgg45
<none>
                     43.087 -62.713
          2 2.0214 45.109 -62.266
- bph_f
- age 1 1.7101 44.798 -60.938
- svi_f 1 3.7964 46.884 -56.523
- lweight 1 5.6462 48.734 -52.769
- lcavol 1 22.5152 65.603 -23.936
Step: AIC=-63.4
lpsa ~ lcavol + lweight + age + bph_f + svi_f
         Df Sum of Sq
                     RSS
                               AIC
                     43.673 -63.403
<none>
            2.2720 45.945 -62.484
- bph_f
        1 1.3945 45.068 -62.354
- age
- svi_f 1 5.2747 48.948 -54.343
- lweight 1 5.3319 49.005 -54.230
- lcavol 1 25.5538 69.227 -20.720
Call:
lm(formula = lpsa ~ lcavol + lweight + age + bph_f + svi_f)
```

```
Coefficients:
(Intercept) lcavol lweight age bph_fMedium
0.14329 0.54022 0.67283 -0.01819 0.37607
bph_fHigh svi_fYes
0.27216 0.68174
```

The backwards elimination approach in this case lands on a model with five inputs (one of which includes two bph indicators,) eliminating only gleason_f, pgg45 and lcp.

7.6 Allen-Cady Modified Backward Elimination

Ranking candidate predictors by importance in advance of backwards elimination can help avoid false-positives, while reducing model size. See Vittinghoff et al. (2012), Section 10.3 for more details.

- 1. First, force into the model any predictors of primary interest, and any confounders necessary for face validity of the final model.
 - "Some variables in the hypothesized causal model may be such well-established causal antecedents of the outcome that it makes sense to include them, essentially to establish the face validity of the model and without regard to the strength or statistical significance of their associations with the primary predictor and outcome ..."
- 2. Rank the remaining candidate predictors in order of importance.
- 3. Starting from an initial model with all candidate predictors included, delete predictors in order of ascending importance until the first variable meeting a criterion to stay in the model hits. Then stop.

Only the remaining variable hypothesized to be least important is eligible for removal at each step. When we are willing to do this sorting before collecting (or analyzing) the data, then we can do Allen-Cady backwards elimination using the drop1 command in R.

7.6.1 Demonstration of the Allen-Cady approach

Suppose, for the moment that we decided to fit a model for the log of psa and we decided (before we saw the data) that we would:

```
lcavol + lweight + svi_f + age + bph_f + gleason_f + lcp + pgg45
```

- force the gleason_f variable to be in the model, due to prior information about its importance,
- and then rated the importance of the other variables as lcavol (most important), then svi_f then age, and then bph_f, then lweight and lcp followed by pgg45 (least important)

When we are willing to do this sorting before collecting (or analyzing) the data, then we can do Allen-Cady backwards elimination using the drop1 command in R.

Step 1. Fit the full model, then see if removing pgg45 improves AIC...

Single term deletions

```
Model:

lpsa ~ gleason_f + lcavol + svi_f + age + bph_f + lweight + lcp +

pgg45

Df Sum of Sq RSS AIC

<none> 41.057 -61.395

pgg45 1 0.44085 41.498 -62.359
```

Since -62.3 is smaller (i.e. more negative) than -61.4, we delete pgg45 and move on to assess whether we can remove the variable we deemed next least important (1cp)

Step 2. Let's see if removing 1cp from this model improves AIC...

Single term deletions

```
Model:
```

Again, since -63.0 is smaller than -62.4, we delete 1cp and next assess whether we should delete 1weight.

Step 3. Does removing lweight from this model improves AIC...

Single term deletions

Model:

Since the AIC for the model after the removal of lweight is larger (i.e. less negative), we stop, and declare our final model by the Allen-Cady approach to include gleason_f, lcavol, svi_f, age, bph_f and lweight.

7.7 Summarizing the Results

Method	Suggested Predictors
Backwards elimination	<pre>lcavol, lweight, svi_f lcavol, lweight, svi_f, age, bph_f lcavol, lweight, svi_f, age, bph_f, gleason_f</pre>

7.7.1 In-Sample Testing and Summaries

Since these models are nested in each other, let's look at the summary statistics (like R², and AIC) and also run an ANOVA-based comparison of these nested models to each other and to the model with the intercept alone, and the kitchen sink model with all available predictors.

7.7.1.1 Model Fit Summaries (in-sample) from glance

Here are the models, at a glance from the broom package.

```
names r.squared adj.r.squared
                                                        AIC
                                                                BIC sigma
1
                               0.000
                                             0.000 306.112 311.261 1.154
                  intercept
                                              0.624 214.097 226.970 0.708
    lcavol + lweight + svi
                                0.636
3 ... + age + bhp + gleason
                                0.671
                                              0.641 214.233 239.980 0.691
4
                  ... + lcp
                                0.676
                                              0.642 214.915 243.237 0.691
5
                ... + pgg45
                                0.679
                                              0.642 215.879 246.775 0.691
 df df.residual
1 1
              96
2 4
              93
3 9
              88
4 10
              87
              86
5 11
```

From these summaries, it looks like:

- the adjusted R² is essentially indistinguishable between the three largest models, but a bit less strong with the three-predictor (4 df) model, and
- the AIC and BIC are (slightly) better (lower) with the three-predictor model (4 df) than any other.

So we might be motivated by these summaries to select any of the three models we're studying closely here.

7.7.1.2 Model Testing via ANOVA (in-sample)

To obtain ANOVA-based test results, we'll run...

```
anova(prost_m_int, prost_m_fw, prost_m_bw, prost_m_ac, prost_m_ks)
```

Analysis of Variance Table

```
Model 1: lpsa ~ 1

Model 2: lpsa ~ lcavol + lweight + svi_f

Model 3: lpsa ~ lcavol + lweight + svi_f + age + bph_f + gleason_f

Model 4: lpsa ~ lcavol + lweight + svi_f + age + bph_f + gleason_f + lcp

Model 5: lpsa ~ lcavol + lweight + svi_f + age + bph_f + gleason_f + lcp +

pgg45

Res.Df RSS Df Sum of Sq F Pr(>F)

1 96 127.918
2 93 46.568 3 81.349 56.7991 <2e-16 ***
```

```
3 88 42.066 5 4.503 1.8863 0.1050

4 87 41.498 1 0.568 1.1891 0.2786

5 86 41.057 1 0.441 0.9234 0.3393

---
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

What conclusions can we draw on the basis of these ANOVA tests?

- There is a statistically significant improvement in predictive value for Model 2 (the forward selection approach) as compared to Model 1 (the intercept only.)
- The ANOVA test comparing Model 5 (kitchen sink) to Model 4 (Allen-Cady result) shows no statistically significant improvement in predictive value.
- Neither does the ANOVA test comparing Model 3 to Model 2 or Model 4 to Model 3.

This suggests that, if we are willing to let the ANOVA test decide our best model than that would be the model produced by forward selection, with predictors lcavol, lweight and svi_f. But we haven't validated the models.

- 1. If the purpose of the model is to predict new data, some sort of out-of-sample or cross-validation approach will be necessary, and
- 2. Even if our goal isn't prediction but merely description of the current data, we would still want to build diagnostic plots to regression assumptions in each model, and
- 3. There is no reason to assume in advance that any of these models is in fact correct, or that any one of these stepwise approaches is necessarily better than any other, and
- 4. The mere act of running a stepwise regression model, as we'll see, can increase the bias in our findings if we accept the results at face value.

So we'll need some ways to validate the results once we complete the selection process.

7.7.2 Validating the Results of the Various Models

We can use a 5-fold cross-validation approach to assess the predictions made by our potential models and then compare them. Let's compare our three models:

- the three predictor model obtained by forward selection, including lcavol, lweight, and svi_f
- the five predictor model obtained by backwards elimination, including lcavol, lweight, svi_f, and also age, and bph_f
- the six predictor model obtained by the Allen-Cady approach, adding gleason_f to the previous model.

Here's the 5-fold validation work (and resulting RMSE and MAE estimates) for the three-predictor model.

```
1 0.745 0.587
```

Now, we'll generate the RMSE and MAE estimates for the five-predictor model.

```
# A tibble: 1 x 2

RMSE_prost5 MAE_prost5

<dbl> <dbl> 1 0.750 0.581
```

And at last, we'll generate the RMSE and MAE estimates for the six-predictor model.

It appears that the six-predictor model does better than either of the other two approaches, with smaller RMSE and MAE. The three-predictor model does slightly better in terms of root mean square prediction error and slightly worse in terms of mean absolute prediction error than the five-predictor model.

OK. A mixed bag, with different conclusions depending on which summary we want to look at. But of course, stepwise regression isn't the only way to do variable selection. Let's consider a broader range of potential predictor sets.

Chapter 8

"Best Subsets" Variable Selection in our Prostate Cancer Study

A second approach to model selection involved fitting all possible subset models and identifying the ones that look best according to some meaningful criterion and ideally one that includes enough variables to model the response appropriately without including lots of redundant or unnecessary terms.

8.1 Four Key Summaries We'll Use to Evaluate Potential Models

- 1. Adjusted R^2 , which we try to maximize.
- 2. Akaike's Information Criterion (AIC), which we try to minimize, and a Bias-Corrected version of AIC due to Hurvich and Tsai (1989), which we use when the sample size is small, specifically when the sample size n and the number of predictors being studied k are such that $n/k \leq 40$. We also try to minimize this bias-corrected AIC.
- 3. Bayesian Information Criterion (BIC), which we also try to minimize.
- 4. Mallows' C_p statistic, which we (essentially) try to minimize.

Choosing between AIC and BIC can be challenging.

For model selection purposes, there is no clear choice between AIC and BIC. Given a family of models, including the true model, the probability that BIC will select the correct model approaches one as the sample size n approaches infinity - thus BIC is asymptotically consistent, which AIC is not. [But, for practical purposes,] BIC often chooses models that are too simple [relative to AIC] because of its heavy penalty on complexity.

• Source: Hastie et al. (2001), page 208.

Several useful tools for running "all subsets" or "best subsets" regression comparisons are developed in R's leaps package.

8.2 Using regsubsets in the leaps package

We can use the leaps package to obtain results in the prost study from looking at all possible subsets of the candidate predictors. The leaps package isn't particularly friendly to the tidyverse. In particular, we cannot have any character variables in our predictor set. We specify our "kitchen sink" model, and apply the regsubsets function from leaps, which identifies the set of models.

To start, we'll ask R to find the one best subset (with 1 predictor variable [in addition to the intercept], then with 2 predictors, and then with each of 3, 4, ... 8 predictor variables) according to an exhaustive search without forcing any of the variables to be in or out.

- Use the nvmax command within the regsubsets function to limit the number of regression inputs to a maximum.
- Use the nbest command to identify how many subsets you want to identify for each predictor count.
- If all of your predictors are **quantitative** or **binary** then you can skip the **preds** step, and simply place your kitchen sink model into **regsubsets**.
- But if you have multi-categorical variables (like gleason_f or svi_f in our case) then you must create a preds group, as follows.

```
preds <- with(prost, cbind(lcavol, lweight, age, bph_f,</pre>
                            svi_f, lcp, gleason_f, pgg45))
rs.ks <- regsubsets(preds, y = prost$lpsa,
                    nvmax = 8, nbest = 1)
rs.summ <- summary(rs.ks)
rs.summ
Subset selection object
8 Variables (and intercept)
          Forced in Forced out
              FALSE
                          FALSE
lcavol
lweight
              FALSE
                          FALSE
              FALSE
                          FALSE
age
bph f
              FALSE
                          FALSE
svi_f
              FALSE
                          FALSE
              FALSE
                          FALSE
lcp
gleason_f
              FALSE
                          FALSE
pgg45
              FALSE
1 subsets of each size up to 8
Selection Algorithm: exhaustive
         lcavol lweight age bph_f svi_f lcp gleason_f pgg45
                         11 11 11 11
                                   11 11
  (1)"*"
                                         11 11 11 11
   (1)"*"
                         11 11
                                          .. .. .. ..
                 "*"
2
  (1)"*"
                         "*"
                                          11 11 11 11
3
  (1)"*"
                         " " "*"
                                   "*"
                 "*"
  (1)"*"
                         "*" "*"
5
  (1)"*"
                                    "*"
6
  (1)"*"
                         "*" "*"
                                   "*"
                                          11 * 11 11 * 11
7
8
  (1)"*"
                                                        "*"
\mathrm{So}...
```

- the best one-predictor model used lcavol
- the best two-predictor model used lcavol and lweight
- the best three-predictor model used lcavol, lweight and svi_f
- the best four-predictor model added bph_f, and
- the best five-predictor model added age
- the best six-input model added gleason_f,
- the best seven-input model added lcp,
- and the eight-input model adds pgg45.

All of these "best subsets" are hierarchical, in that each model is a subset of the one below it. This isn't inevitably true.

• To determine which model is best, we can plot key summaries of model fit (adjusted \mathbb{R}^2 , Mallows' C_p ,

bias-corrected AIC, and BIC) using either base R plotting techniques (what I've done in the past) or ggplot2 (what I use now.) I'll show both types of plotting approaches in the next two sections.

8.2.1 Identifying the models with which and outmat

To see the models selected by the system, we use:

```
rs.summ$which
  (Intercept) lcavol lweight
                                age bph_f svi_f
                                                  lcp gleason_f pgg45
1
         TRUE
                TRUE
                       FALSE FALSE FALSE FALSE
                                                           FALSE FALSE
2
                TRUE
         TRUE
                        TRUE FALSE FALSE FALSE
                                                           FALSE FALSE
3
         TRUE
                TRUE
                        TRUE FALSE FALSE
                                           TRUE FALSE
                                                           FALSE FALSE
4
         TRUE
                TRUE
                        TRUE FALSE
                                     TRUE
                                           TRUE FALSE
                                                           FALSE FALSE
5
         TRUE
                TRUE
                        TRUE
                               TRUE
                                     TRUE
                                           TRUE FALSE
                                                           FALSE FALSE
6
         TRUE
                TRUE
                        TRUE
                               TRUE
                                     TRUE
                                           TRUE FALSE
                                                            TRUE FALSE
                                                            TRUE FALSE
7
         TRUE
                TRUE
                               TRUE
                                     TRUE
                                           TRUE
                        TRUE
                                                 TRUE
8
         TRUE
                TRUE
                        TRUE
                               TRUE
                                     TRUE
                                           TRUE
                                                 TRUE
                                                            TRUE
                                                                 TRUE
```

Another version of this formatted for printing is:

```
rs.summ$outmat
```

```
lcavol lweight age bph_f svi_f lcp gleason_f pgg45
         "*"
1
   (1)
         "*"
                                   .. ..
2
   (1)
         "*"
3
   (1)
         "*"
   (1
       )
                                   "*"
         "*"
   ( 1
       )
   (1
       )
         "*"
                             "*"
                                   "*"
7
   (1)
   (1)"*"
                         "*" "*"
                                          "*" "*"
                                                        "*"
```

We built one subset of each size up to eight predictors, and if we add the intercept term, this means we have models of size k = 2, 3, 4, 5, 6, 7, 8 and 9.

The models are:

Size k	Predictors included (besides intercept)
2	lcavol
3	lcavol and lweight
4	add svi_f
5	add bph_f
6	add age
7	add gleason_f
8	add lcp
9	add pgg45

8.3 Calculating bias-corrected AIC

The bias-corrected AIC formula developed in Hurvich and Tsai (1989) requires three inputs:

- the residual sum of squares for a model
- the sample size (n) or number of observations used to fit the model
- the number of regression inputs, k, including the intercept, used in the model

So, for a particular model fit to n observations, on k predictors (including the intercept) and a residual sum of squares equal to RSS, we have:

$$AIC_c = nlog(\frac{RSS}{n}) + 2k + \frac{2k(k+1)}{n-k-1}$$

Note that the corrected AIC_c can be related to the original AIC via:

$$AIC_c = AIC + \frac{2k(k+1)}{n-k-1}$$

8.3.1 Calculation of aic.c in our setting

In our case, we have n = 97 observations, and built a series of models with k = 2:9 predictors (including the intercept in each case), so we will insert those values into the general formula for bias-corrected AIC which is:

aic.c
$$\leftarrow$$
 n * log(rs.summ\$rss / n) + 2 * k + (2 * k * (k + 1) / (n - k - 1))

We can obtain the residual sum of squares explained by each model by pulling rss from the regsubsets summary contained here in rs.summ.

```
data_frame(k = 2:9, RSS = rs.summ$rss)
```

```
# A tibble: 8 x 2
      k
          RSS
  <int> <dbl>
      2 58.9
1
2
      3
        51.7
3
        46.6
      4
4
      5
        45.7
5
      6
        44.6
6
      7
        43.7
7
      8
         43.0
      9
        42.8
```

In this case, we have:

The impact of this bias correction can be modest but important. Here's a little table looking closely at the results in this problem. The uncorrected AIC are obtained using extractAIC, as described in the next section.

Size	2	3	4	5	6	7	8	9
Bias-corrected AIC	-44.2	-54.7	-62.7	-62.3	-62.3	-62.1	-61.2	-59.4
Uncorrected AIC	-44.4	-55.0	-63.2	-62.4	-63.4	-63.0	-62.4	-61.4

extractAIC(lm(lpsa ~ lcavol, data = prost))

8.3.2 The Uncorrected AIC provides no more useful information here

We could, if necessary, also calculate the *uncorrected* aic value for each model, but we won't make any direct use of that, because that will not provide any new information not already gathered by the C_p statistic for a linear regression model. If you wanted to find the uncorrected AIC for a given model, you can use the extractAIC function.

```
2.00000 -44.36603
[1]
extractAIC(lm(lpsa ~ lcavol + lweight, data = prost))
      3.00000 -54.95846
[1]
Note that:
  • these results are fairly comparable to the bias-corrected AIC we built above, and
  • the extractAIC and AIC functions look like they give very different results, but they really don't.
AIC(lm(lpsa ~ lcavol, data = prost))
[1] 232.908
AIC(lm(lpsa ~ lcavol + lweight, data = prost))
[1] 222.3156
But notice that the differences in AIC are the same, either way, comparing these two models:
extractAIC(lm(lpsa ~ lcavol, data = prost)) - extractAIC(lm(lpsa ~ lcavol + lweight, data = prost))
[1] -1.00000 10.59243
AIC(lm(lpsa ~ lcavol, data = prost)) - AIC(lm(lpsa ~ lcavol + lweight, data = prost))
[1] 10.59243
```

- AIC is only defined up to an additive constant.
- Since the difference between two models using either AIC or extractAIC is the same, this doesn't actually matter which one we use, so long as we use the same one consistently.

8.3.3 Building a Tibble containing the necessary information

Again, note the use of 2:9 for the values of k, because we're fitting one model for each size from 2 through 9.

```
best_mods_1 <- data_frame(
    k = 2:9,
    r2 = rs.summ$rsq,
    adjr2 = rs.summ$adjr2,
    cp = rs.summ$cp,
    aic.c = rs.summ$bic
)

best_mods <- cbind(best_mods_1, rs.summ$which)

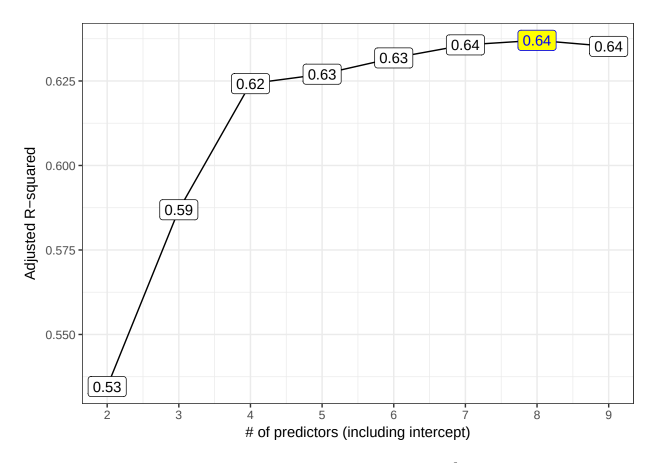
best_mods</pre>
```

```
k r2 adjr2 cp aic.c bic (Intercept) lcavol 1 2 0.5394320 0.5345839 28.213914 -44.23838 -66.05416 TRUE TRUE
```

```
2 3 0.5955040 0.5868977 15.456669 -54.70040 -74.07188
                                                          TRUE
                                                                TRUE
3 4 0.6359499 0.6242063 6.811986 -62.74265 -79.71614
                                                         TRUE
                                                                TRUE
4 5 0.6425479 0.6270065 7.075509 -62.29223 -76.91557
                                                         TRUE
                                                                TRUE
5 6 0.6509970 0.6318211 6.851826 -62.33858 -74.66120
                                                         TRUE
                                                                TRUE
6 7 0.6584484 0.6356783 6.890739 -62.10692 -72.17992
                                                         TRUE
                                                                TRUE
7 8 0.6634967 0.6370302 7.562119 -61.17338 -69.04961
                                                         TRUE
                                                                TRUE
8 9 0.6656326 0.6352355 9.000000 -59.35841 -65.09253
                                                         TRUE
                                                                TRUE
 lweight
         age bph_f svi_f lcp gleason_f pgg45
                                   FALSE FALSE
   FALSE FALSE FALSE FALSE
    TRUE FALSE FALSE FALSE
2
                                   FALSE FALSE
3
    TRUE FALSE FALSE TRUE FALSE
                                   FALSE FALSE
    TRUE FALSE TRUE TRUE FALSE
4
                                   FALSE FALSE
                                  FALSE FALSE
    TRUE TRUE TRUE TRUE FALSE
5
    TRUE TRUE TRUE FALSE
                                   TRUE FALSE
6
7
    TRUE TRUE TRUE TRUE TRUE
                                   TRUE FALSE
    TRUE TRUE TRUE TRUE TRUE
                                     TRUE TRUE
```

8.4 Plotting the Best Subsets Results using ggplot2

8.4.1 The Adjusted R^2 Plot



Models 4-9 all look like reasonable choices here. The maximum adjusted \mathbb{R}^2 is seen in the model of size 8.

8.4.2 Mallows' C_p

The C_p statistic focuses directly on the tradeoff between **bias** (due to excluding important predictors from the model) and extra **variance** (due to including too many unimportant predictors in the model.)

If N is the sample size, and we select p regression predictors from a set of K (where p < K), then the C_p statistic is

$$C_p = \frac{SSE_p}{MSE_K} - N + 2p$$

where:

- SSE_p is the sum of squares for error (residual) in the model with p predictors
- MSE_K is the residual mean square after regression in the model with all K predictors

As it turns out, this is just measuring the particular model's lack of fit, and then adding a penalty for the number of terms in the model (specifically 2p-N is the penalty since the lack of fit is measured as $(N-p)\frac{SSE_p}{MSE_K}$.

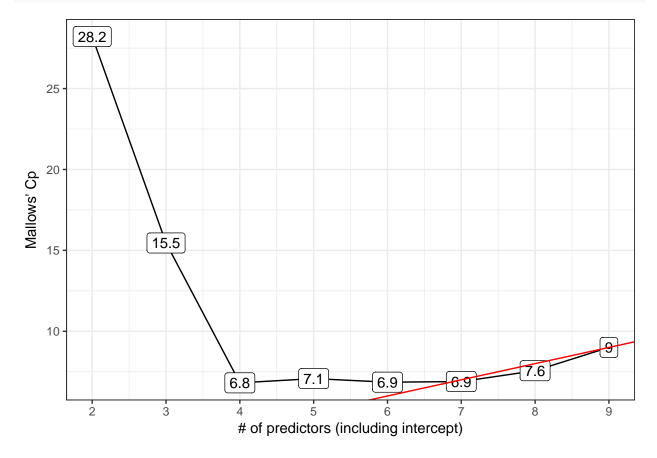
- If a model has no meaningful lack of fit (i.e. no substantial bias) then the expected value of C_p is roughly p.
- Otherwise, the expectation is p plus a positive bias term.
- In general, we want to see *smaller* values of C_p .
- We usually select a "winning model" by choosing a subset of predictors that have C_p near the value of p.

8.4.3 The C_p Plot

The C_p plot is just a scatterplot of C_p on the Y-axis, and the size of the model (coefficients plus intercept) p = k on the X-axis.

Each of the various predictor subsets we will study is represented in a single point. A model without bias should have C_p roughly equal to p, so we'll frequently draw a line at $C_p = p$ to make that clear. We then select our model from among all models with small C_p statistics.

- My typical approach is to identify the models where $C_p p \ge 0$, then select from among those models the model where $C_p p$ is minimized, and if there is a tie, select the model where p is minimized.
- Another good candidate might be a slightly overfit model (where $C_p p < 0$ but just barely.)



- Model 6 is a possibility here, with the difference $C_p p$ minimized among all models with $C_p >= p$.
- Model 7 also looks pretty good, with C_p just barely smaller than the size (p = 7) of the model.

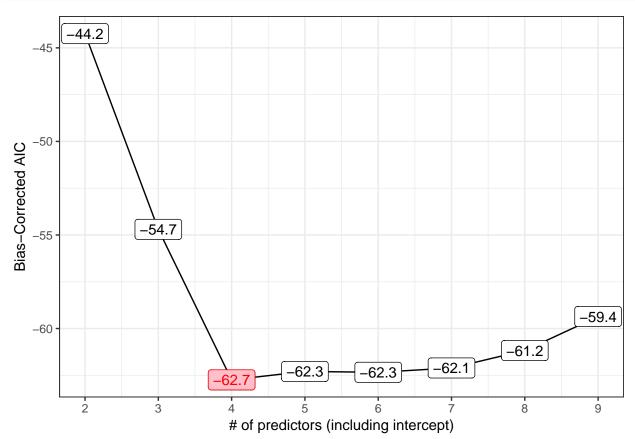
8.4.4 "All Subsets" Regression and Information Criteria

We might consider any of three main information criteria:

- the Bayesian Information Criterion, called BIC
- the Akaike Information Criterion (used by R's default stepwise approaches,) called AIC
- a corrected version of AIC due to Hurvich and Tsai (1989), called AIC $_{\rm c}$ or aic.c

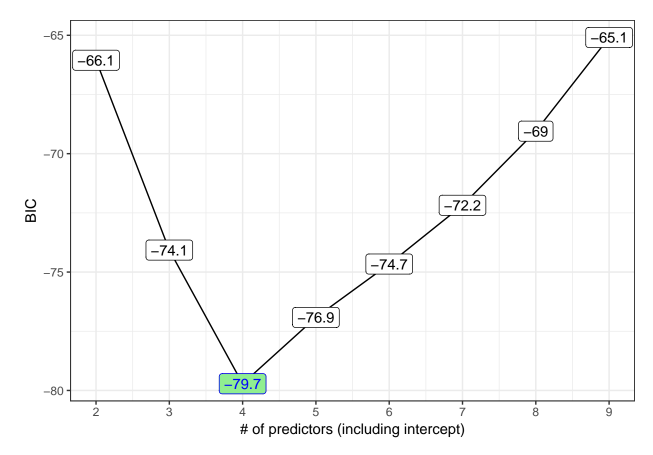
Each of these indicates better models by getting smaller. Since the C_p and AIC results will lead to the same model, I'll focus on plotting the bias-corrected AIC and on BIC.

8.4.5 The bias-corrected AIC plot



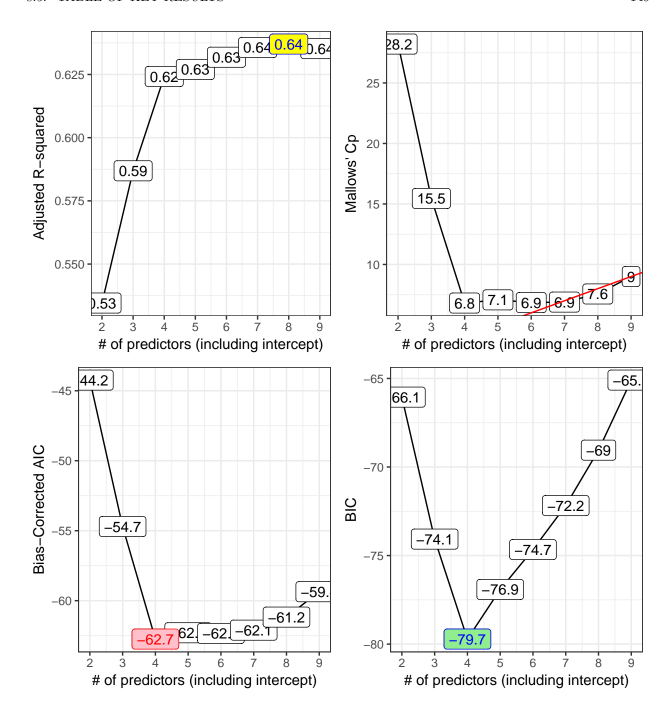
The smallest AIC_c values occur in models 4 and later, especially model 4 itself.

8.4.6 The BIC plot



8.4.7 All Four Plots in One Figure (via ggplot2)

```
gridExtra::grid.arrange(p1, p2, p3, p4, nrow = 2)
```



8.5 Table of Key Results

We can build a big table, like this:

best_mods

```
k r2 adjr2 cp aic.c bic (Intercept) lcavol
1 2 0.5394320 0.5345839 28.213914 -44.23838 -66.05416 TRUE TRUE
2 3 0.5955040 0.5868977 15.456669 -54.70040 -74.07188 TRUE TRUE
3 4 0.6359499 0.6242063 6.811986 -62.74265 -79.71614 TRUE TRUE
```

```
4 5 0.6425479 0.6270065 7.075509 -62.29223 -76.91557
                                                         TRUE
                                                                TRUE
5 6 0.6509970 0.6318211 6.851826 -62.33858 -74.66120
                                                         TRUE
                                                                TRUE
6 7 0.6584484 0.6356783 6.890739 -62.10692 -72.17992
                                                         TRUE
                                                                TRUE
7 8 0.6634967 0.6370302 7.562119 -61.17338 -69.04961
                                                         TRUE
                                                                TRUE
8 9 0.6656326 0.6352355 9.000000 -59.35841 -65.09253
                                                         TRUE
                                                                TRUE
 lweight
         age bph_f svi_f lcp gleason_f pgg45
   FALSE FALSE FALSE FALSE
                                  FALSE FALSE
    TRUE FALSE FALSE FALSE
                                   FALSE FALSE
3
    TRUE FALSE FALSE TRUE FALSE
                                   FALSE FALSE
4
    TRUE FALSE TRUE TRUE FALSE
                                   FALSE FALSE
5
    TRUE TRUE TRUE FALSE
                                   FALSE FALSE
    TRUE TRUE TRUE FALSE
6
                                    TRUE FALSE
7
    TRUE TRUE TRUE TRUE TRUE
                                    TRUE FALSE
    TRUE TRUE TRUE TRUE TRUE
8
                                    TRUE TRUE
```

8.6 Models Worth Considering?

\overline{k}	Predictors	Reason
4	<pre>lcavol lweight svi_f</pre>	minimizes BIC, AIC _c
7	+ age bph_f gleason_f	C_p near p
8	+ lcp	$\max_{i} R_{adj}^2$

8.7 Compare these candidate models in-sample?

8.7.1 Using anova to compare nested models

Let's run an ANOVA-based comparison of these nested models to each other and to the model with the intercept alone.

The models are nested because m04 is a subset of the predictors in m07, which includes a subset of the
predictors in m08.

Next, we'll run...

```
anova(m.full, m08, m07, m04, m.int)
```

Analysis of Variance Table

```
Model 1: lpsa ~ lcavol + lweight + svi_f + age + bph_f + gleason_f + lcp +
    pgg45

Model 2: lpsa ~ lcavol + lweight + svi_f + age + bph_f + gleason_f + lcp

Model 3: lpsa ~ lcavol + lweight + svi_f + age + bph_f + gleason_f

Model 4: lpsa ~ lcavol + lweight + svi_f
```

```
Model 5: lpsa ~ 1
  Res.Df
             RSS Df Sum of Sq
                                     F Pr(>F)
2
      87
          41.498 -1
                       -0.441
                                0.9234 0.3393
3
          42.066 -1
                        -0.568
                                1.1891 0.2786
                        -4.503 1.8863 0.1050
4
          46.568 -5
                       -81.349 56.7991 <2e-16 ***
5
      96 127.918 -3
Signif. codes:
                0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

What conclusions can we draw here, on the basis of these ANOVA tests?

- The first p value, of 0.3393, compares what the anova called Model 1, and what we call m.full to what the anova called Model 2, and what we call m08. So there's no significant decline in predictive value observed when we drop from the m.full model to the m08 model. This suggests that the m08 model may be a better choice.
- The second p value, of 0.2786, compares m08 to m07, and suggests that we lose no significant predictive value by dropping down to m07.
- The third p value, of 0.1050, compares m07 to m04, and suggests that we lose no significant predictive value by dropping down to m04.
- But the fourth p value, of 2e-16 (or, functionally, zero), compares m04 to m.int and suggests that we do gain significant predictive value by including the predictors in m04 as compared to a model with an intercept alone.
- So, by the significance tests, the model we'd select would be m04, but, of course, in-sample statistical significance alone isn't a good enough reason to select a model if we want to do prediction well.

8.8 AIC and BIC comparisons, within the training sample

Next, we'll compare the three candidate models (ignoring the intercept-only and kitchen sink models) in terms of their AIC values and BIC values, again using the same sample we used to fit the models in the first place.

```
AIC(m04, m07, m08)

df AIC
m04 5 214.0966
m07 10 214.2327
m08 11 214.9148

BIC(m04, m07, m08)

df BIC
m04 5 226.9702
m07 10 239.9798
m08 11 243.2366
```

- The model with the smallest AIC value shows the best performance within the sample on that measure.
- Similarly, smaller BIC values are associated with predictor sets that perform better in sample on that criterion.
- BIC often suggests smaller models (with fewer regression inputs) than does AIC. Does that happen in this case?
- Note that AIC and BIC can be calculated in a few different ways, so we may see some variation if we don't compare apples to apples with regard to the R functions involved.

8.9 Cross-Validation of Candidate Models out of Sample

8.9.1 20-fold Cross-Validation of model m04

Model m04 uses lcavol, lweight and svi_f to predict the lpsa outcome. Let's do 20-fold cross-validation of this modeling approach, and calculate the root mean squared prediction error and the mean absolute prediction error for that modeling scheme.

```
set.seed(43201)
cv_m04 <- prost %>%
    crossv_kfold(k = 20) %>%
   mutate(model = map(train,
                       ~ lm(lpsa ~ lcavol + lweight + svi_f,
                                   data = .)))
cv_m04_pred <- cv_m04 %>%
    unnest(map2(model, test, ~ augment(.x, newdata = .y)))
cv_m04_results <- cv_m04_pred %>%
    summarize(Model = "m04",
             RMSE = sqrt(mean((lpsa - .fitted) ^2)),
              MAE = mean(abs(lpsa - .fitted)))
cv_m04_results
# A tibble: 1 x 3
 Model RMSE MAE
  <chr> <dbl> <dbl>
1 m04
       0.725 0.574
```

8.9.2 20-fold Cross-Validation of model m07

Model m07 uses lcavol, lweight, svi_f, age, bph_f, and gleason_f to predict the lpsa outcome. Let's now do 20-fold cross-validation of this modeling approach, and calculate the root mean squared prediction error and the mean absolute prediction error for that modeling scheme. Note the small changes required, as compared to our cross-validation of model m04 a moment ago.

```
MAE = mean(abs(lpsa - .fitted)))

cv_m07_results

# A tibble: 1 x 3
   Model RMSE MAE
   <chr> <dbl> <dbl>
1 m07   0.730  0.556
```

8.9.3 20-fold Cross-Validation of model m08

Model m08 uses lcavol, lweight, svi_f, age, bph_f, gleason_f and lcp to predict the lpsa outcome. Let's now do 20-fold cross-validation of this modeling approach.

```
set.seed(43202)
cv_m08 <- prost %>%
    crossv_kfold(k = 20) %>%
   mutate(model = map(train,
                       ~ lm(lpsa ~ lcavol + lweight +
                                svi_f + age + bph_f +
                                gleason_f + lcp,
                                   data = .)))
cv_m08_pred <- cv_m08 %>%
   unnest(map2(model, test, ~ augment(.x, newdata = .y)))
cv_m08_results <- cv_m08_pred %>%
    summarize(Model = "m08",
              RMSE = sqrt(mean((lpsa - .fitted) ^2)),
              MAE = mean(abs(lpsa - .fitted)))
cv_m08_results
# A tibble: 1 x 3
 Model RMSE MAE
  <chr> <dbl> <dbl>
1 m08 0.729 0.557
```

8.9.4 Comparing the Results of the Cross-Validations

It appears that model m04 has the smallest RMSE and MAE in this case. So, that's the model with the strongest cross-validated predictive accuracy, by these two standards.

8.10 What about Interaction Terms?

Suppose we consider for a moment a much smaller and less realistic problem. We want to use best subsets to identify a model out of a set of three predictors for lpsa: specifically lcavol, age and svi_f, but now we also want to consider the interaction of svi_f with lcavol as a potential addition. Remember that svi is the 1/0 numeric version of svi_f. We could simply add a numerical product term to our model, as follows.

```
pred2 <- with(prost, cbind(lcavol, age, svi_f, svixlcavol = svi*lcavol))</pre>
rs.ks2 <- regsubsets(pred2, y = prost$lpsa,
                    nvmax = NULL, nbest = 1)
rs.summ2 <- summary(rs.ks2)
rs.summ2
Subset selection object
4 Variables (and intercept)
           Forced in Forced out
lcavol
               FALSE
                          FALSE
               FALSE
                          FALSE
age
               FALSE
                          FALSE
svi_f
               FALSE
svixlcavol
                          FALSE
1 subsets of each size up to 4
Selection Algorithm: exhaustive
         lcavol age svi_f svixlcavol
  (1)"*"
  (1)"*"
  (1)"*"
                          "*"
```

In this case, best subsets doesn't identify the interaction term as an attractive predictor until it has already included the main effects that go into it. So that's fine. But if that isn't the case, we would have a problem.

To resolve this, we could:

- 1. Consider interactions beforehand, and force them in if desired.
- 2. Consider interaction terms outside of best subsets, and only after the selection of main effects.
- 3. Use another approach to deal with variable selection for interaction terms.

Chapter 9

Adding Non-linear Terms to a Linear Regression Model

9.1 The pollution data

Consider the pollution data set, which contain 15 independent variables and a measure of mortality, describing 60 US metropolitan areas in 1959-1961. The data come from McDonald and Schwing (1973), and are available at http://www4.stat.ncsu.edu/~boos/var.select/pollution.html and our web site.

pollution

```
# A tibble: 60 x 16
      x1
            x2
                   xЗ
                         x4
                                x5
                                      x6
                                             x7
                                                   8x
                                                           x9
                                                                x10
   <int> <int> <int> <dbl> <dbl> <dbl> <dbl> <int>
                                                       <dbl> <dbl> <dbl>
                                                 3243
                                                       8.80
 1
                       8.10
                             3.34 11.4
                                           81.5
                                                               42.6
 2
      35
            23
                   72 11.1
                              3.14 11.0
                                           78.8
                                                 4281
                                                       3.50
                                                               50.7
 3
            29
                   74 10.4
                              3.21
                                    9.80
                                          81.6
                                                 4260
                                                       0.800
                             3.41 11.1
 4
                                                 3125 27.1
      47
            45
                   79
                       6.50
                                           77.5
                                                               50.2
 5
            35
                       7.60
                             3.44
                                   9.60
                                          84.6
                                                 6441 24.4
 6
                       7.70
                             3.45 10.2
      53
            45
                   80
                                           66.8
                                                 3325 38.5
                                                               43.1
                                                                     25.5
 7
            30
                   74 10.9
                             3.23 12.1
                                          83.9
                                                 4679
                                                       3.50
                                                               49.2
 8
      45
            30
                   73
                       9.30
                             3.29 10.6
                                           86.0
                                                 2140
                                                       5.30
                                                               40.4
                                                                     10.5
                                          83.2
 9
            24
                       9.00
                             3.31 10.5
                                                 6582
                                                       8.10
                                                               42.5 12.6
10
      36
            27
                   72 9.50
                             3.36 10.7
                                          79.3
                                                 4213
                                                       6.70
                                                               41.0 13.2
      with 50 more rows, and 5 more variables: x12 <int>, x13 <int>,
    x14 <int>, x15 <int>, y <dbl>
```

TT 1 1 1 1

Here's a codebook:

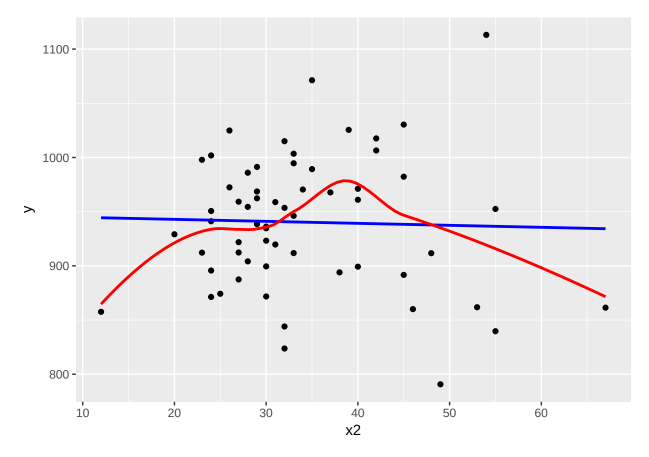
Variable	Description
У	Total Age Adjusted Mortality Rate
x1	Mean annual precipitation in inches
x2	Mean January temperature in degrees Fahrenheit
x3	Mean July temperature in degrees Fahrenheit
x4	Percent of 1960 SMSA population that is 65 years of age or over
x 5	Population per household, 1960 SMSA
x6	Median school years completed for those over 25 in 1960 SMSA
x7	Percent of housing units that are found with facilities
x8	Population per square mile in urbanized area in 1960

Variable	Description
x9	Percent of 1960 urbanized area population that is non-white
x10	Percent employment in white-collar occupations in 1960 urbanized area
x11	Percent of families with income under 3; 000 in 1960 urbanized area
x12	Relative population potential of hydrocarbons, HC
x13	Relative pollution potential of oxides of nitrogen, NOx
x14	Relative pollution potential of sulfur dioxide, SO2
x15	Percent relative humidity, annual average at 1 p.m.

9.2 Fitting a straight line model to predict y from x2

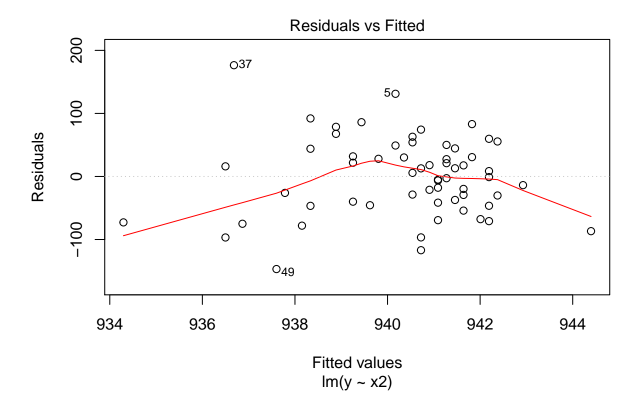
Consider the relationship between y, the age-adjusted mortality rate, and x2, the mean January temperature, across these 60 areas. I'll include both a linear model (in blue) and a loess smooth (in red.) Does the relationship appear to be linear?

```
ggplot(pollution, aes(x = x2, y = y)) +
   geom_point() +
   geom_smooth(method = "lm", col = "blue", se = F) +
   geom_smooth(method = "loess", col = "red", se = F)
```



Suppose we plot the residuals that emerge from the linear model shown in blue, above. Do we see a curve in a plot of residuals against fitted values?

```
plot(lm(y ~ x2, data = pollution), which = 1)
```



Quadratic polynomial model to predict y using x2 9.3

A polynomial in the variable x of degree D is a linear combination of the powers of x up to D.

For example:

• Linear: $y = \beta_0 + \beta_1 x$

• Quadratic: $y = \beta_0 + \beta_1 x + \beta_2 x^2$ • Cubic: $y = \beta_0 + \beta_1 x + \beta_2 x^2 + \beta_3 x^3$ • Quartic: $y = \beta_0 + \beta_1 x + \beta_2 x^2 + \beta_3 x^3 + \beta_4 x^4$ • Quintic: $y = \beta_0 + \beta_1 x + \beta_2 x^2 + \beta_3 x^3 + \beta_4 x^4 + \beta_5 x^5$

Fitting such a model creates a **polynomial regression*.

The raw quadratic model

Let's look at a quadratic model which predicts y using x2 and the square of x2, so that our model is of the form:

$$y = \beta_0 + \beta_1 x_2 + \beta_2 x_2^2 + error$$

There are several ways to fit this exact model.

- One approach is to calculate the square of x2 within our pollution data set, and then feed both x2 and x2squared to 1m.
- Another approach uses the I function within our 1m to specify the use of both x2 and its square.

• Yet another approach uses the poly function within our lm, which can be used to specify raw models including x2 and x2squared.

```
pollution <- pollution %>%
    mutate(x2squared = x2^2)

mod2a <- lm(y ~ x2 + x2squared, data = pollution)
mod2b <- lm(y ~ x2 + I(x2^2), data = pollution)
mod2c <- lm(y ~ poly(x2, degree = 2, raw = TRUE), data = pollution)</pre>
```

Each of these approaches produces the same model, as they are just different ways of expressing the same idea.

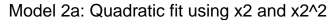
```
summary(mod2a)
```

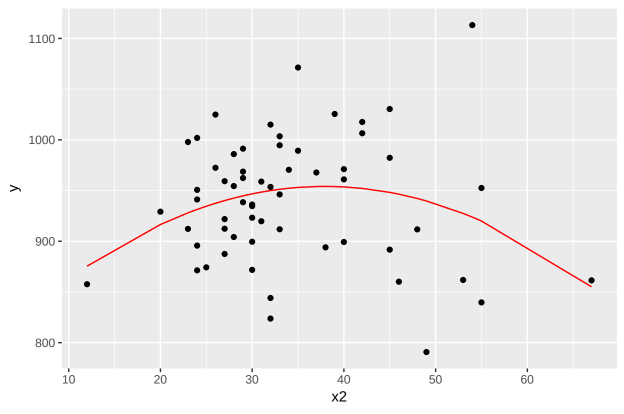
```
Call:
lm(formula = y \sim x2 + x2squared, data = pollution)
Residuals:
    Min
              1Q Median
                               ЗQ
                                       Max
-148.977 -38.651
                 6.889 35.312 189.346
Coefficients:
            Estimate Std. Error t value Pr(>|t|)
(Intercept) 785.77449 79.54086 9.879 5.87e-14 ***
            8.87640
                        4.27394 2.077 0.0423 *
            -0.11704
                        0.05429 -2.156 0.0353 *
x2squared
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 60.83 on 57 degrees of freedom
Multiple R-squared: 0.07623, Adjusted R-squared: 0.04382
F-statistic: 2.352 on 2 and 57 DF, p-value: 0.1044
```

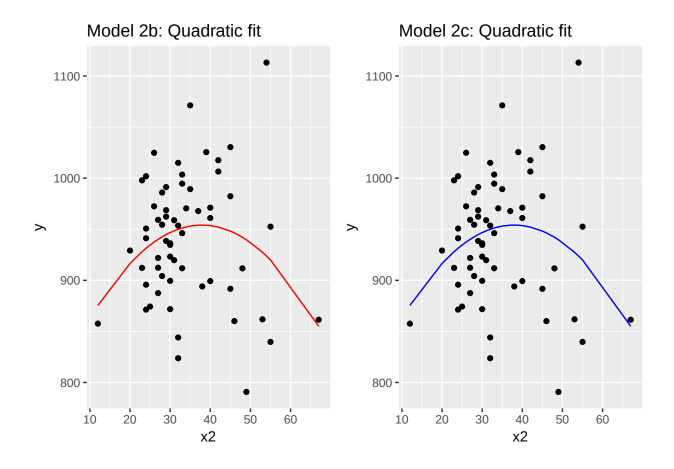
And if we plot the fitted values for this mod2 using whatever approach you like, we get exactly the same result.

```
mod2a.aug <- augment(mod2a, pollution)

ggplot(mod2a.aug, aes(x = x2, y = y)) +
    geom_point() +
    geom_line(aes(x = x2, y = .fitted), col = "red") +
    labs(title = "Model 2a: Quadratic fit using x2 and x2^2")</pre>
```







9.3.2 Raw quadratic fit after centering x2

Sometimes, we'll center (and perhaps rescale, too) the x2 variable before including it in a quadratic fit like this.

```
pollution <- pollution %>%
    mutate(x2_c = x2 - mean(x2))

mod2d <- lm(y ~ x2_c + I(x2_c^2), data = pollution)

summary(mod2d)</pre>
```

```
Call:
```

```
lm(formula = y \sim x2_c + I(x2_c^2), data = pollution)
```

Residuals:

```
Min 1Q Median 3Q Max -148.977 -38.651 6.889 35.312 189.346
```

Coefficients:

```
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

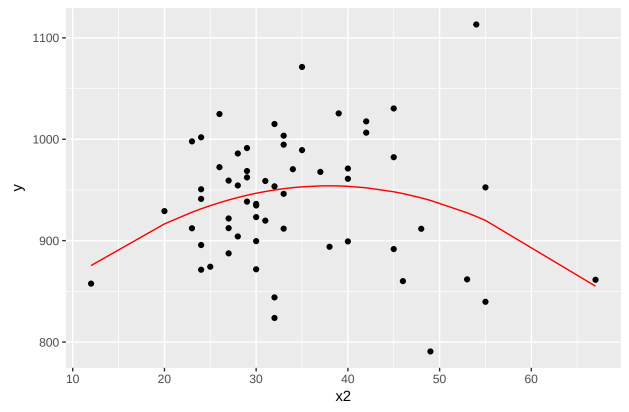
Residual standard error: 60.83 on 57 degrees of freedom Multiple R-squared: 0.07623, Adjusted R-squared: 0.04382 F-statistic: 2.352 on 2 and 57 DF, p-value: 0.1044

Note that this model looks very different, with the exception of the second order quadratic term. But, it produces the same fitted values as the models we fit previously.

```
mod2d.aug <- augment(mod2d, pollution)

ggplot(mod2d.aug, aes(x = x2, y = y)) +
    geom_point() +
    geom_line(aes(x = x2, y = .fitted), col = "red") +
    labs(title = "Model 2d: Quadratic fit using centered x2 and x2^2")</pre>
```

Model 2d: Quadratic fit using centered x2 and x2^2



Or, if you don't believe me yet, look at the four sets of fitted values another way.

mod2a.aug %>% skim(.fitted)

p100

```
Skim summary statistics
n obs: 60
n variables: 24

Variable type: numeric
variable missing complete n mean sd p0 p25 median p75
.fitted 0 60 60 940.36 17.18 855.1 936.72 945.6 950.29
```

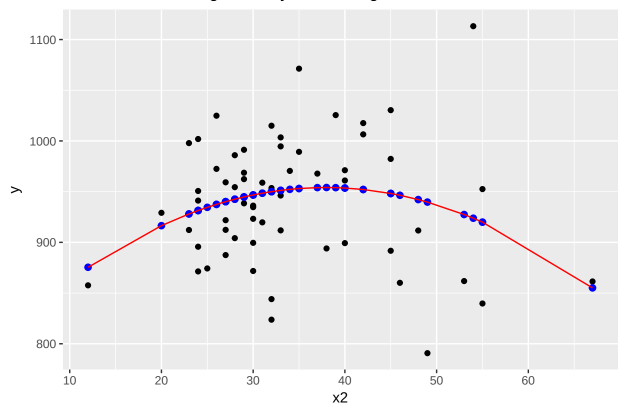
```
954.07
mod2b.aug %>% skim(.fitted)
Skim summary statistics
n obs: 60
n variables: 24
Variable type: numeric
variable missing complete n mean sd p0 p25 median
                   60 60 940.36 17.18 855.1 936.72 945.6 950.29
 .fitted
             0
  p100
954.07
mod2c.aug %>% skim(.fitted)
Skim summary statistics
n obs: 60
n variables: 24
Variable type: numeric
variable missing complete n mean sd p0 p25 median
 .fitted
             0 60 60 940.36 17.18 855.1 936.72 945.6 950.29
  p100
954.07
mod2d.aug %>% skim(.fitted)
Skim summary statistics
n obs: 60
n variables: 25
Variable type: numeric
variable missing complete n mean sd p0 p25 median
 .fitted
             0 60 60 940.36 17.18 855.1 936.72 945.6 950.29
  p100
954.07
      Orthogonal Polynomials
Now, let's fit an orthogonal polynomial of degree 2 to predict y using x2.
mod2_orth \leftarrow lm(y \sim poly(x2, 2), data = pollution)
summary(mod2_orth)
Call:
lm(formula = y \sim poly(x2, 2), data = pollution)
Residuals:
    Min
            1Q Median
                               ЗQ
                                       Max
                 6.889 35.312 189.346
-148.977 -38.651
Coefficients:
```

Estimate Std. Error t value Pr(>|t|)

```
(Intercept)
              940.358
                          7.853 119.746
                                           <2e-16 ***
poly(x2, 2)1 -14.345
                          60.829 -0.236
                                           0.8144
poly(x2, 2)2 -131.142
                          60.829
                                 -2.156
                                           0.0353 *
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 60.83 on 57 degrees of freedom
Multiple R-squared: 0.07623,
                               Adjusted R-squared:
F-statistic: 2.352 on 2 and 57 DF, p-value: 0.1044
```

Now this looks very different in the equation, but, again, we can see that this produces exactly the same fitted values as our previous models, and the same model fit summaries. Is it, in fact, the same model? Here, we'll plot the fitted Model 2a in a red line, and this new Model 2 with Orthogonal Polynomials as blue points.

Model 2 with Orthogonal Polynomial, degree 2



Yes, it is again the same model in terms of the predictions it makes for y.

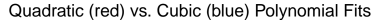
By default, with raw = FALSE, the poly() function within a linear model computes what is called an **orthogonal polynomial**. An orthogonal polynomial sets up a model design matrix using the coding we've seen previously: x2 and $x2^2$ in our case, and then scales those columns so that each column is **orthogonal** to the previous ones. This eliminates the collinearity (correlation between predictors) and lets our t tests tell us whether the addition of any particular polynomial term improves the fit of the model over the lower orders.

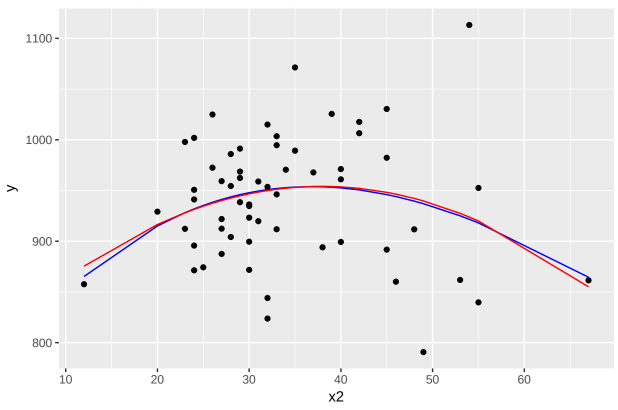
Would the addition of a cubic term help us much in predicting y from x2?

```
mod3 <- lm(y ~ poly(x2, 3), data = pollution)
summary(mod3)</pre>
```

```
Call:
lm(formula = y \sim poly(x2, 3), data = pollution)
Residuals:
    Min
               1Q
                    Median
                                 3Q
                                         Max
-146.262 -39.679
                     5.569
                             35.984
                                    191.536
Coefficients:
             Estimate Std. Error t value Pr(>|t|)
              940.358
(Intercept)
                          7.917 118.772
                                           <2e-16 ***
poly(x2, 3)1 -14.345
                          61.328
                                 -0.234
                                           0.8159
poly(x2, 3)2 -131.142
                          61.328
                                 -2.138
                                           0.0369 *
poly(x2, 3)3
               16.918
                          61.328
                                   0.276
                                           0.7837
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
Residual standard error: 61.33 on 56 degrees of freedom
Multiple R-squared: 0.07748,
                               Adjusted R-squared:
F-statistic: 1.568 on 3 and 56 DF, p-value: 0.2073
```

It doesn't appear that the cubic term adds much here, if anything. The p value is not significant for the third degree polynomial, the summaries of fit quality aren't much improved, and as we can see from the plot below, the predictions don't actually change all that much.

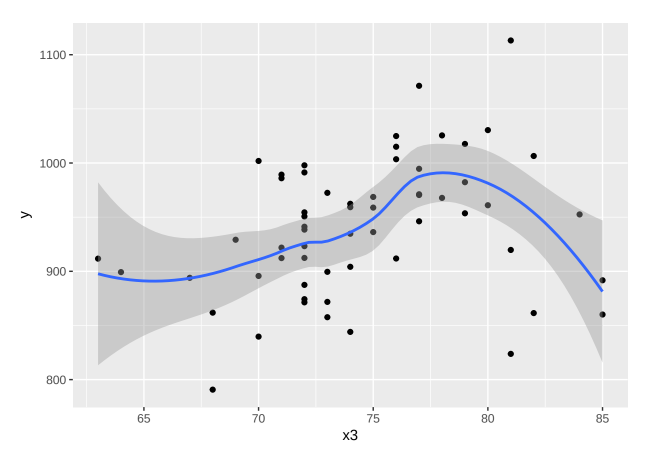




9.5 Fit a cubic polynomial to predict y from x3

What if we consider another predictor instead? Let's look at x3, the Mean July temperature in degrees Fahrenheit. Here is the loess smooth.

```
ggplot(pollution, aes(x = x3, y = y)) +
  geom_point() +
  geom_smooth(method = "loess")
```



That looks pretty curvy - perhaps we need a more complex polynomial. We'll consider a linear model (mod4_L), a quadratic fit (mod4_Q) and a polynomial of degree 3: a cubic fit (mod_4C)

```
mod4_L <- lm(y ~ x3, data = pollution)
summary(mod4_L)</pre>
```

```
Call:
```

lm(formula = y ~ x3, data = pollution)

Residuals:

Min 1Q Median 3Q Max -139.813 -34.341 4.271 38.197 149.587

Coefficients:

Estimate Std. Error t value Pr(>|t|)
(Intercept) 670.529 123.140 5.445 1.1e-06 ***
x3 3.618 1.648 2.196 0.0321 *

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

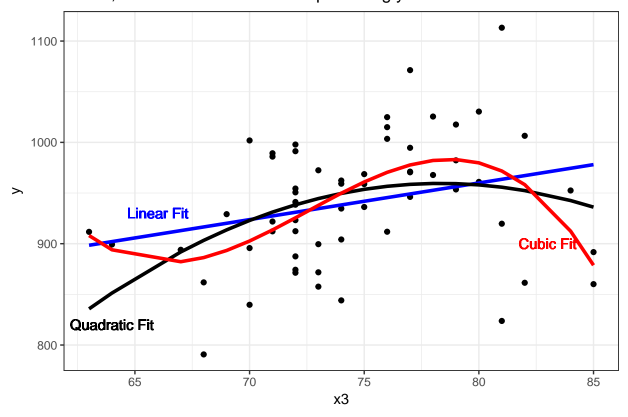
Residual standard error: 60.29 on 58 degrees of freedom Multiple R-squared: 0.07674, Adjusted R-squared: 0.06082 F-statistic: 4.821 on 1 and 58 DF, p-value: 0.03213

mod4_Q <- lm(y ~ poly(x3, 2), data = pollution)
summary(mod4_Q)</pre>

```
Call:
lm(formula = y \sim poly(x3, 2), data = pollution)
Residuals:
              1Q Median
    Min
                                3Q
                                         Max
-132.004 -42.184 4.069 47.126 157.396
Coefficients:
            Estimate Std. Error t value Pr(>|t|)
(Intercept) 940.358
                        7.553 124.503
                                          <2e-16 ***
poly(x3, 2)1 132.364
                         58.504 2.262
                                           0.0275 *
poly(x3, 2)2 -125.270
                         58.504 -2.141 0.0365 *
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 58.5 on 57 degrees of freedom
Multiple R-squared: 0.1455,
                              Adjusted R-squared: 0.1155
F-statistic: 4.852 on 2 and 57 DF, p-value: 0.01133
mod4_C \leftarrow lm(y \sim poly(x3, 3), data = pollution)
summary(mod4_C)
lm(formula = y \sim poly(x3, 3), data = pollution)
Residuals:
    Min
              10
                  Median
                                 3Q
                                         Max
-148.004 -29.998
                    1.441
                            34.579 141.396
Coefficients:
            Estimate Std. Error t value Pr(>|t|)
(Intercept) 940.358 7.065 133.095 < 2e-16 ***
poly(x3, 3)1 132.364
                         54.728 2.419 0.01886 *
poly(x3, 3)2 -125.270
                         54.728 -2.289 0.02588 *
poly(x3, 3)3 -165.439
                         54.728 -3.023 0.00377 **
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
Residual standard error: 54.73 on 56 degrees of freedom
Multiple R-squared: 0.2654,
                              Adjusted R-squared: 0.226
F-statistic: 6.742 on 3 and 56 DF, p-value: 0.0005799
It looks like the cubic polynomial term is of some real importance here. Do the linear, quadratic and cubic
model fitted values look different?
mod4_L.aug <- augment(mod4_L, pollution)</pre>
mod4_Q.aug <- augment(mod4_Q, pollution)</pre>
mod4_C.aug <- augment(mod4_C, pollution)</pre>
ggplot(pollution, aes(x = x3, y = y)) +
   geom point() +
```

geom_line(data = mod4_L.aug, aes(x = x3, y = .fitted),

Linear, Quadratic and Cubic Fits predicting y with x3



9.6 Fitting a restricted cubic spline in a linear regression

- A linear spline is a continuous function formed by connecting points (called **knots** of the spline) by line segments.
- A **restricted cubic spline** is a way to build highly complicated curves into a regression equation in a fairly easily structured way.
- A restricted cubic spline is a series of polynomial functions joined together at the knots.
 - Such a spline gives us a way to flexibly account for non-linearity without over-fitting the model.
 - Restricted cubic splines can fit many different types of non-linearities.
 - Specifying the number of knots is all you need to do in R to get a reasonable result from a restricted cubic spline.

The most common choices are 3, 4, or 5 knots. Each additional knot adds to the non-linearity, and spends an additional degree of freedom:

- 3 Knots, 2 degrees of freedom, allows the curve to "bend" once.
- 4 Knots, 3 degrees of freedom, lets the curve "bend" twice.
- 5 Knots, 4 degrees of freedom, lets the curve "bend" three times.

For most applications, three to five knots strike a nice balance between complicating the model needlessly and fitting data pleasingly. Let's consider a restricted cubic spline model for our y based on x3 again, but now with:

- in mod5a, 3 knots,
- in mod5b, 4 knots, and
- in mod5c, 5 knots

Here, for instance, is the summary of the 5-knot model:

```
summary(mod5c_rcs)
```

Call:

```
lm(formula = y \sim rcs(x3, 5), data = pollution)
```

Residuals:

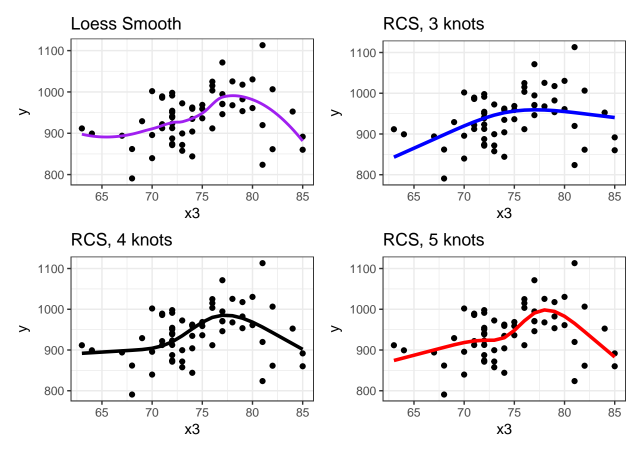
```
Min 1Q Median 3Q Max
-141.522 -32.009 1.674 31.971 147.878
```

Coefficients:

```
Estimate Std. Error t value Pr(>|t|)
(Intercept)
                468.113
                           396.319 1.181
                                             0.243
rcs(x3, 5)x3
                  6.447
                           5.749 1.121
                                             0.267
rcs(x3, 5)x3'
                -25.633
                            46.810 -0.548
                                             0.586
rcs(x3, 5)x3''
                323.137
                           293.065
                                    1.103
                                             0.275
rcs(x3, 5)x3''' -612.578
                           396.270 -1.546
                                             0.128
```

Residual standard error: 54.35 on 55 degrees of freedom Multiple R-squared: 0.2883, Adjusted R-squared: 0.2366 F-statistic: 5.571 on 4 and 55 DF, p-value: 0.0007734

We'll begin by storing the fitted values from these three models and other summaries, for plotting.



Does it seem like the fit improves markedly (perhaps approaching the loess smooth result) as we increase the number of knots?

```
anova(mod5a_rcs, mod5b_rcs, mod5c_rcs)
```

Analysis of Variance Table

Model 1: $y \sim rcs(x3, 3)$

```
Model 2: y ~ rcs(x3, 4)

Model 3: y ~ rcs(x3, 5)

Res.Df RSS Df Sum of Sq F Pr(>F)

1 57 194935

2 56 171448 1 23486.9 7.9503 0.006672 **

3 55 162481 1 8967.2 3.0354 0.087057 .

---

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Record on an ANOVA comparison, the fourth knot adds significant prodic
```

Based on an ANOVA comparison, the fourth knot adds significant predictive value (p = 0.0067), but the fifth knot is borderline (p = 0.0871). From the glance function in the broom package, we can also look at some key summaries.

```
glance(mod5a_rcs)
 r.squared adj.r.squared
                            sigma statistic
                                               p.value df
                                                             logLik
               0.1162256 58.48006 4.879558 0.01106323 3 -327.7187
1 0.146184
      AIC
               BIC deviance df.residual
1 663.4373 671.8147 194935.3
glance(mod5b rcs)
 r.squared adj.r.squared
                            sigma statistic
                                              p.value df
               0.2088274 55.33153 6.190953 0.0010423 4 -323.8671
      AIC
               BIC deviance df.residual
1 657.7342 668.2059 171448.4
glance(mod5c_rcs)
 r.squared adj.r.squared
                            sigma statistic
                                                 p.value df
                                                               logLik
```

Model	Knots	\mathbb{R}^2	$Adj. R^2$	AIC	BIC
5a	3	0.146	0.116	663.4	671.8
5b	4	0.249	0.209	657.7	668.2
5c	5	0.288	0.237	656.5	669.1

Within our sample, the five-knot RCS outperforms the 3- and 4-knot versions on adjusted R^2 and AIC (barely) and does a little worse than the 4-knot RCS on BIC.

Of course, we could also use the cross-validation methods we've developed for other linear regressions to assess predictive capacity of these models. I'll skip that for now.

To see the values of x3 where the splines place their knots, we can use the attributes function.

```
$dim
[1] 60 4

$dimnames
$dimnames[[1]]

NULL
```

```
$dimnames[[2]]
[1] "pollution" "pollution'" "pollution''" "pollution''"
```

attributes(rcs(pollution\$x3, 5))

```
$class
[1] "rms"
$name
[1] "pollution"
$label
[1] "pollution"
$assume
[1] "rcspline"
$assume.code
[1] 4
$parms
[1] 68 72 74 77 82
$nonlinear
[1] FALSE TRUE TRUE TRUE
$colnames
                                   "pollution''" "pollution''"
                   "pollution'"
[1] "pollution"
```

The knots in this particular 5-knot spline are placed by the computer at 68, 72, 74, 77 and 82, it seems.

There are two kinds of Multivariate Regression Models

- 1. [Prediction] Those that are built so that we can make accurate predictions.
- 2. [Explanatory] Those that are built to help understand underlying phenomena.

While those two notions overlap considerably, they do imply different things about how we strategize about model-building and model assessment. Harrell's primary concern is effective use of the available data for **prediction** - this implies some things that will be different from what we've seen in the past.

Harrell refers to multivariable regression modeling strategy as the process of **spending degrees of freedom**. The main job in strategizing about multivariate modeling is to

- 1. Decide the number of degrees of freedom that can be spent
- 2. Decide where to spend them
- 3. Spend them, wisely.

What this means is essentially linked to making decisions about predictor complexity, both in terms of how many predictors will be included in the regression model, and about how we'll include those predictors.

9.7 "Spending" Degrees of Freedom

- "Spending" df includes
 - fitting parameter estimates in models, or
 - examining figures built using the outcome variable Y that tell you how to model the predictors.

If you use a scatterplot of Y vs. X or the residuals of the Y-X regression model vs. X to decide whether a linear model is appropriate, then how many degrees of freedom have you actually spent?

Grambsch and O'Brien conclude that if you wish to preserve the key statistical properties of the various estimation and fitting procedures used in building a model, you can't retrieve these degrees of freedom once

they have been spent.

9.7.1 Overfitting and Limits on the # of Predictors

Suppose you have a total sample size of n observations, then you really shouldn't be thinking about estimating more than n/15 regression coefficients, at the most.

- If k is the number of parameters in a full model containing all candidate predictors for a stepwise analysis, then k should be no greater than n/15.
- k should include all variables screened for association with the response, including interaction terms.

So if you have 97 observations in your data, then you can probably just barely justify the use of a stepwise analysis using the main effects alone of 5 candidate variables (with one additional DF for the intercept term.)

Harrell (2001) also mentions that if you have a **narrowly distributed** predictor, without a lot of variation to work with, then an even larger sample size n should be required. See Vittinghoff et al. (2012), Section 10.3 for more details.

9.7.2 The Importance of Collinearity

Collinearity denotes correlation between predictors high enough to degrade the precision of the regression coefficient estimates substantially for some or all of the correlated predictors

- Vittinghoff et al. (2012), section 10.4.1
- Can one predictor in a model be predicted well using the other predictors in the model?
 - Strong correlations (for instance, $r \ge 0.8$) are especially troublesome.
- Effects of collinearity
 - decreases precision, in the sense of increasing the standard errors of the parameter estimates
 - decreases power
 - increases the difficulty of interpreting individual predictor effects
 - overall F test is significant, but individual t tests may not be

Suppose we want to assess whether variable X_j is collinear with the other predictors in a model. We run a regression predicting X_j using the other predictors, and obtain the R^2 . The VIF is defined as 1 / (1 - this R^2), and we usually interpret VIFs above 5 as indicating a serious multicollinearity problem (i.e. R^2 values for this predictor of 0.8 and above would thus concern us.)

Occasionally, you'll see the inverse of VIF reported, and this is called tolerance.

• tolerance = 1 / VIF

9.7.3 Collinearity in an Explanatory Model

2.238862 2.058731 2.153044 4.174448 3.447399 1.792996

- When we are attempting to **identify multiple independent predictors** (the explanatory model approach), then we will need to choose between collinear variables
 - options suggested by Vittinghoff et al. (2012), p. 422, include choosing on the basis of plausibility as a causal factor,
 - choosing the variable that has higher data quality (is measured more accurately or has fewer missing values.)

- Often, we choose to include a variable that is statistically significant as a predictor, and drop others, should we be so lucky.
- Larger effects, especially if they are associated with predictors that have minimal correlation with the other predictors under study, cause less trouble in terms of potential violation of the n/15 rule for what constitutes a reasonable number of predictors.

9.7.4 Collinearity in a Prediction Model

- If we are primarily building a **prediction model** for which inference on the individual predictors is not of interest, then it is totally reasonable to use both predictors in the model, if doing so reduces prediction error.
 - Collinearity doesn't affect predictions in our model development sample.
 - Collinearity doesn't affect predictions on new data so long as the new data have similar relationships between predictors.
 - If our key predictor is correlated strongly with a confounder, then if the predictor remains significant after adjustment for the confounder, then this suggests a meaningful independent effect.
 - * If the effects of the predictor are clearly confounded by the adjustment variable, we again have a clear result.
 - * If neither is statistically significant after adjustment, the data may be inadequate.
 - If the collinearity is between adjustment variables, but doesn't involve the key predictor, then
 inclusion of the collinear variables is unlikely to cause substantial problems.

9.8 Spending DF on Non-Linearity: The Spearman ρ^2 Plot

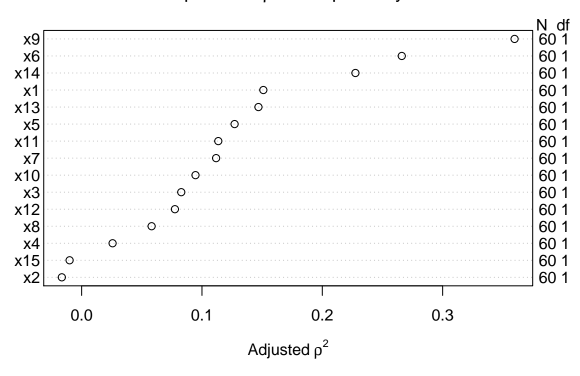
We need a flexible approach to assessing non-linearity and fitting models with non-linear predictors. This will lead us to a measure of what Harrell (2001) calls **potential predictive punch** which hides the true form of the regression from the analyst so as to preserve statistical properties, but that lets us make sensible decisions about whether a predictor should be included in a model, and the number of parameters (degrees of freedom, essentially) we are willing to devote to it.

What if we want to consider where best to spend our degrees of freedom on non-linear predictor terms, like interactions, polynomial functions or curved splines to represent our input data? The approach we'll find useful in the largest variety of settings is a combination of

- 1. a rank correlation assessment of potential predictive punch (using a Spearman ρ^2 plot, available in the Hmisc package), followed by
- 2. the application of restricted cubic splines to fit and assess models.

Suppose, for instance, that we want to create a model for y using some combination of linear and non-linear terms drawn from the complete set of 15 predictors available in the pollution data. I'd begin by running a Spearman ρ^2 plot:

```
plot(Hmisc::spearman2(y ~ x1 + x2 + x3 + x4 + x5 + x6 + x7 + x8 + x9 + x10 + x11 + x12 + x13 + x14 + x15, data = pollution))
```



Spearman ρ^2 Response: y

The variable with the largest adjusted squared Spearman ρ statistic in this setting is x9, followed by x6 and x14. With only 60 observations, we might well want to restrict ourselves to a very small model. What the Spearman plot suggests is that we focus any non-linear terms on x9 first, and then perhaps x6 and x14 as they have some potential predictive power. It may or may not work out that the non-linear terms are productive.

9.8.1 Fitting a Big Model to the pollution data

So, one possible model built in reaction this plot might be to fit:

- a restricted cubic spline with 5 knots on x9
- a restricted cubic spline with 3 knots on x6
- and a quadratic polynomial on x14
- plus a linear fit to x1 and x13

That's way more degrees of freedom (4 for x9, 2 for x6, 2 for x14 and 1 each for x1 and x13 makes a total of 10 without the intercept term) than we can really justify with a sample of 60 observations. But let's see what happens.

```
mod_big \leftarrow lm(y \sim rcs(x9, 5) + rcs(x6, 3) + poly(x14, 2) + x1 + x13, data = pollution)
anova(mod_big)
```

Analysis of Variance Table

```
Response: y

Df Sum Sq Mean Sq F value Pr(>F)

rcs(x9, 5) 4 100164 25040.9 17.8482 4.229e-09 ***
```

```
rcs(x6, 3)
             2
                38306 19152.8 13.6513 1.939e-05 ***
poly(x14, 2)
             2
                15595 7797.7 5.5579 0.006677 **
                       4787.3 3.4122 0.070759 .
                 4787
                  712
                        711.9 0.5074 0.479635
             1
x13
Residuals
            49
                68747
                       1403.0
Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
```

This anova suggests that we have at least some predictive value in each spline (x9 and x6) and some additional value in x14, although it's not as clear that the linear terms (x1 and x13) did much good.

9.8.2 Limitations of 1m for fitting complex linear regression models

We can certainly assess this big, complex model using 1m in comparison to other models:

- with in-sample summary statistics like adjusted R², AIC and BIC,
- we can assess its assumptions with residual plots, and
- we can also compare out-of-sample predictive quality through cross-validation,

But to really delve into the details of how well this complex model works, and to help plot what is actually being fit, we'll probably want to fit the model using an alternative method for fitting linear models, called ols, from the rms package developed by Frank Harrell and colleagues. That will be the focus of our next chapter.

Chapter 10

Using ols from the rms package to fit linear models

At the end of the previous chapter, we had fit a model to the pollution data that predicted our outcome y = Age-Adjusted Mortality Rate, using:

- a restricted cubic spline with 5 knots on x9
- a restricted cubic spline with 3 knots on x6
- a polynomial in 2 degrees on x14
- linear terms for x1 and x13

but this model was hard to evaluate in some ways. Now, instead of using 1m to fit this model, we'll use a new function called ols from the rms package developed by Frank Harrell and colleagues, in part to support ideas developed in Harrell (2001) for clinical prediction models.

10.1 Fitting a model with ols

We will use the datadist approach when fitting a linear model with ols from the rms package, so as to store additional important elements of the model fit.

```
library(rms)

d <- datadist(pollution)
options(datadist = "d")</pre>
```

Next, we'll fit the model using ols and place its results in newmod.

```
newmod <- ols(y ~ rcs(x9, 5) + rcs(x6, 3) + pol(x14, 2) + x1 + x13, data = pollution, x = TRUE, y = TRUE)
newmod
```

Linear Regression Model

```
ols(formula = y ~ rcs(x9, 5) + rcs(x6, 3) + pol(x14, 2) + x1 +
    x13, data = pollution, x = TRUE, y = TRUE)

    Model Likelihood Discrimination
        Ratio Test Indexes
Obs 60 LR chi2 72.02 R2 0.699
```

```
sigma37.4566 d.f. 10 R2 adj 0.637
d.f. 49 Pr(> chi2) 0.0000 g 58.961
```

Residuals

```
Min 1Q Median 3Q Max
-86.189 -18.554 -1.799 18.645 104.307
```

	Coef	S.E.	t	Pr(> t)
Intercept	796.2658	162.3269	4.91	<0.0001
x9	-2.6328	6.3504	-0.41	0.6803
x9'	121.4651	124.4827	0.98	0.3340
x9''	-219.8025	227.6775	-0.97	0.3391
x9'''	151.5700	171.3867	0.88	0.3808
x6	7.6817	15.5230	0.49	0.6229
x6'	-29.4388	18.0531	-1.63	0.1094
x14	0.5652	0.2547	2.22	0.0311
x14^2	-0.0010	0.0010	-0.96	0.3407
x1	1.0717	0.7317	1.46	0.1494
x13	-0.1028	0.1443	-0.71	0.4796

Some of the advantages and disadvantages of fitting linear regression models with ols or lm will reveal themselves over time. For now, one advantage for ols is that the entire variance-covariance matrix is saved. Most of the time, there will be some value to considering both ols and lm approaches.

Most of this output should be familiar, but a few pieces are different.

10.1.1 The Model Likelihood Ratio Test

The Model Likelihood Ratio Test compares newmod to the null model with only an intercept term. It is a goodness-of-fit test that we'll use in several types of model settings this semester.

- In many settings, the logarithm of the likelihood ratio, multiplied by -2, yields a value which can be compared to a χ^2 distribution. So here, the value 72.02 is -2(log likelihood), and is compared to a χ^2 distribution with 10 degrees of freedom. We reject the null hypothesis that newmod is no better than the null model, and conclude instead that at least one of these predictors adds statistically significant value.
 - For ols, interpret the model likelihood ratio test like the global (ANOVA) F test in lm.
 - The likelihood function is the probability of observing our data under the specified model.
 - We can compare two nested models by evaluating the difference in their likelihood ratios and degrees of freedom, then comparing the result to a χ^2 distribution.

10.1.2 The g statistic

The **g statistic** is new and is referred to as the g-index. it's based on Gini's mean difference and is purported to be a robust and highly efficient measure of variation.

- Here, g = 58.9, which implies that if you randomly select two of the 60 areas included in the model, the average difference in predicted y (Age-Adjusted Mortality Rate) using this model will be 58.9.
 - Technically, g is Gini's mean difference of the predicted values.

10.2 ANOVA for an ols model

One advantage of the ols approach is that when you apply an anova to it, it separates out the linear and non-linear components of restricted cubic splines and polynomial terms (as well as product terms, if your model includes them.)

anova (newmod)

Analysis of Variance			Response: y		
Factor	d.f.	Partial SS	MS	F	P
x9	4	35219.7647	8804.9412	6.28	0.0004
Nonlinear	3	1339.3081	446.4360	0.32	0.8121
x6	2	9367.6008	4683.8004	3.34	0.0437
Nonlinear	1	3730.7388	3730.7388	2.66	0.1094
x14	2	18679.6957	9339.8478	6.66	0.0028
Nonlinear	1	1298.7625	1298.7625	0.93	0.3407
x1	1	3009.1829	3009.1829	2.14	0.1494
x13	1	711.9108	711.9108	0.51	0.4796
TOTAL NONLINEAR	5	6656.1824	1331.2365	0.95	0.4582
REGRESSION	10	159563.8285	15956.3829	11.37	<.0001
ERROR	49	68746.8004	1402.9959		

Unlike the anova approach in 1m, in ols ANOVA, partial F tests are presented - each predictor is assessed as "last predictor in" much like the usual t tests in 1m. In essence, the partial sums of squares and F tests here describe the marginal impact of removing each covariate from newmod.

We conclude that the non-linear parts of x9 and x6 and x14 combined don't seem to add much value, but that overall, x9, x6 and x14 seem to be valuable. So it must be the linear parts of those variables within our model that are doing the lion's share of the work.

10.3 Effect Estimates

Effects

A particularly useful thing to get out of the ols approach that is not as easily available in lm (without recoding or standardizing our predictors) is a summary of the effects of each predictor in an interesting scale.

Response : y

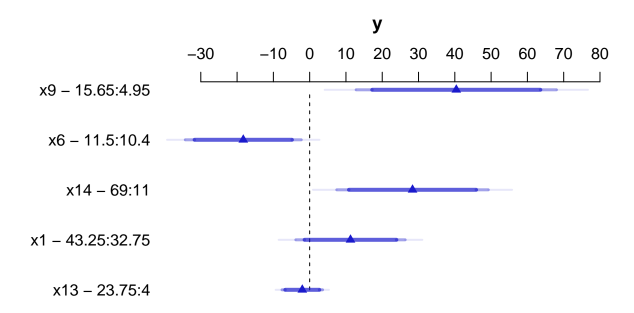
summary(newmod)

```
Factor Low
             High Diff. Effect
                                           Lower 0.95 Upper 0.95
                                  S.E.
        4.95 15.65 10.70 40.4060 14.0790
                                                      68.6990
x9
                                           12.1120
       10.40 11.50 1.10 -18.2930 8.1499 -34.6710
x6
                                                      -1.9153
       11.00 69.00 58.00
                          28.3480 10.6480
                                                      49.7460
x14
                                             6.9503
       32.75 43.25 10.50 11.2520
                                   7.6833
                                            -4.1878
                                                      26.6930
x1
        4.00 23.75 19.75
                         -2.0303
x13
                                   2.8502
                                           -7.7579
                                                       3.6973
```

This "effects summary" shows the effect on y of moving from the 25th to the 75th percentile of each variable (along with a standard error and 95% confidence interval) while holding the other variable at the level specified at the bottom of the output.

The most useful way to look at this sort of analysis is often a plot.

```
plot(summary(newmod))
```



For x9 note from the summary above that the 25th percentile is 4.95 and the 75th is 15.65. Our conclusion is that the estimated effect of moving x9 from 4.95 to 15.65 is an increase of 40.4 on y, with a 95% CI of (12.1, 68.7).

For a categorical variable, the low level is shown first and then the high level.

The plot shows the point estimate (arrow head) and then the 90% (narrowest bar), 95% (middle bar) and 99% (widest bar in lightest color) confidence intervals for each predictor's effect.

- It's easier to distinguish this in the x9 plot than the one for x13.
- Remember that what is being compared is the first value to the second value's impact on the outcome, with other predictors held constant.

10.3.1 Simultaneous Confidence Intervals

These confidence intervals make no effort to deal with the multiple comparisons problem, but just fit individual 95% (or whatever level you choose) confidence intervals for each predictor. The natural alternative is to make an adjustment for multiple comparisons in fitting the confidence intervals, so that the set of (in this case, five - one for each predictor) confidence intervals for effect sizes has a family-wise 95% confidence level. You'll note that the effect estimates and standard errors are unchanged from those shown above, but the confidence limits are a bit wider.

```
x6
       10.40 11.50 1.10 -18.2930 8.1499 -39.87400
                                                       3.2882
       11.00 69.00 58.00 28.3480 10.6480
x14
                                            0.15192
                                                     56.5440
x1
       32.75 43.25 10.50
                         11.2520
                                   7.6833
                                           -9.09340
                                                      31.5980
x13
        4.00 23.75 19.75
                          -2.0303
                                   2.8502
                                           -9.57760
                                                       5.5171
```

Remember that if you're looking for the usual lm summary for an ols object, use summary.lm, and that the display function from arm does not recognize ols objects.

10.4 The Predict function for an ols model

The Predict function is very flexible, and can be used to produce individual or simultaneous confidence limits.

```
limits.
Predict(newmod, x9 = 12, x6 = 12, x14 = 40, x1 = 40, x13 = 20) # individual limits
  x9 x6 x14 x1 x13
                       yhat
                               lower
                                       upper
1 12 12 40 40 20 923.0982 893.0984 953.098
Response variable (y): y
Limits are 0.95 confidence limits
Predict(newmod, x9 = 5:15) # individual limits
         x6 x14 x1 x13
                           yhat
                                   lower
                     9 913.7392 889.4802 937.9983
   5 11.05
            30 38
   6 11.05
             30 38
                     9 916.3490 892.0082 940.6897
3
            30 38
   7 11.05
                     9 921.3093 898.9657 943.6529
   8 11.05
            30 38
                     9 927.6464 907.0355 948.2574
   9 11.05
             30 38
                     9 934.3853 913.3761 955.3946
6
  10 11.05
            30 38
                     9 940.5510 917.8371 963.2648
7
  11 11.05
            30 38
                     9 945.2225 921.9971 968.4479
  12 11.05
            30 38
                     9 948.2885 926.4576 970.1194
                     9 950.2608 930.3003 970.2213
 13 11.05
            30 38
10 14 11.05 30 38
                     9 951.6671 932.2370 971.0971
11 15 11.05 30 38
                     9 953.0342 932.1662 973.9021
Response variable (y): y
Adjust to: x6=11.05 x14=30 x1=38 x13=9
Limits are 0.95 confidence limits
Predict(newmod, x9 = 5:15, conf.type = 'simult')
   x9
         x6 x14 x1 x13
                           yhat
                                   lower
                                             upper
   5 11.05
             30 38
                     9 913.7392 882.4311 945.0473
   6 11.05
                     9 916.3490 884.9354 947.7625
             30 38
   7 11.05
             30 38
                     9 921.3093 892.4733 950.1454
   8 11.05
             30 38
                     9 927.6464 901.0465 954.2464
   9 11.05
             30 38
                     9 934.3853 907.2713 961.4993
  10 11.05
             30 38
                     9 940.5510 911.2371 969.8649
7
  11 11.05
             30 38
                     9 945.2225 915.2484 975.1966
  12 11.05
             30 38
                     9 948.2885 920.1141 976.4629
 13 11.05
             30 38
                     9 950.2608 924.5003 976.0212
10 14 11.05 30 38
                     9 951.6671 926.5912 976.7430
```

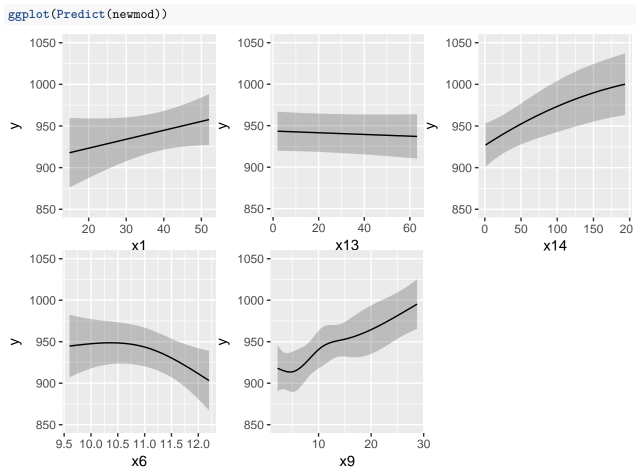
11 15 11.05 30 38 9 953.0342 926.1025 979.9658

Response variable (y): y

Adjust to: x6=11.05 x14=30 x1=38 x13=9

Limits are 0.95 confidence limits

The plot below shows the individual effects in newmod in five subpanels, using the default approach of displaying the same range of values as are seen in the data. Note that each panel shows point and interval estimates of the effects, and spot the straight lines in x1 and x13, the single bends in x14 and x6 and the wiggles in x9, corresponding to the amount of non-linearity specified in the model.



10.5 Checking Influence via dfbeta

For an ols object, we have several tools for looking at residuals. The most interesting to me is which.influence which is reliant on the notion of dfbeta.

- DFBETA is estimated for each observation in the data, and each coefficient in the model.
- The DFBETA is the difference in the estimated coefficient caused by deleting the observation, scaled by the coefficient's standard error estimated with the observation deleted.
- The which.influence command applied to an ols model produces a list of all of the predictors estimated by the model, including the intercept.
 - For each predictor, the command lists all observations (by row number) that, if removed from the

model, would cause the estimated coefficient (the "beta") for that predictor to change by at least some particular cutoff.

- The default is that the DFBETA for that predictor is 0.2 or more.

which.influence(newmod)

```
$Intercept
   2 11 28 32 37 49 59
Г17
$x9
[1]
    2 3 6 9 31 35 49 57 58
$x6
[1]
    2 11 15 28 32 37 50 56 59
$x14
    2 6 7 12 13 16 32 37
[1]
$x1
    7 18 32 37 49 57
[1]
$x13
[1] 29 32 37
```

The implication here, for instance, is that if we drop row 3 from our data frame, and refit the model, this will have a meaningful impact on the estimate of x9 but not on the other coefficients. But if we drop, say, row 37, we will affect the estimates of the intercept, x6, x14, x1, and x13.

10.5.1 Using the residuals command for dfbetas

To see the dfbeta values, standardized according to the approach I used above, you can use the following code (I'll use head to just show the first few rows of results) to get a matrix of the results.

```
head(residuals(newmod, type = "dfbetas"))
```

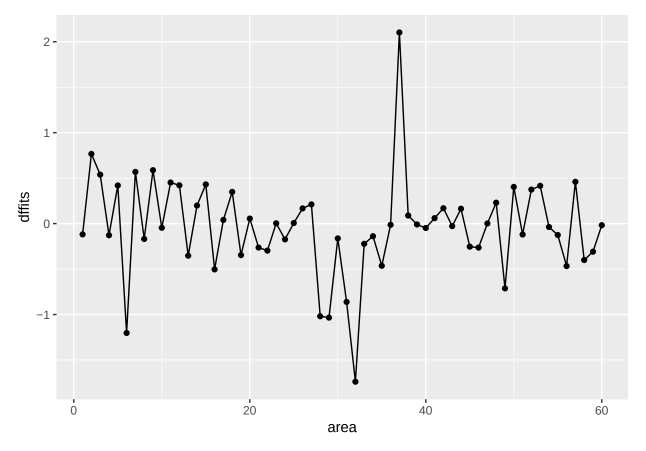
```
[,4]
           [,1]
                       [,2]
                                    [,3]
    0.03071160 -0.023775487 -0.004055111 0.01205425 -0.03260003
[2,] -0.38276573 -0.048404993 -0.142293606
                                         0.17009666 -0.22350621
[3,]
    0.17226780 -0.426153536 0.350913139 -0.32949129
                                                     0.25777913
[4,]
    0.06175110 -0.006460916 0.024828272 -0.03009337
                                                     0.04154812
[5,]
                0.039839994 -0.058178534
     0.16875200
                                         0.06449504 -0.07772208
[6,]
     0.03322073
                0.112699877 -0.203543632 0.23987378 -0.35201736
           [,6]
                      [,7]
                                  [,8]
                                              [,9]
[1,] -0.02392315
                0.01175375 -0.06494414 0.060929683 -0.011042644
[2,] 0.44737372 -0.48562818 0.19372285 -0.212186731 -0.107830147
[3,] -0.10263448
                0.05005284 -0.02049877 0.014059330 0.010793169
[4,] -0.06254145
                0.16151742 0.02723710
                                       0.065483158
[5,] -0.18058630
                                                   0.003326357
[6,] -0.04075617
                0.02900006 -0.21508009 0.171627718 0.019241676
          [,11]
    0.03425156
[1,]
[2,] -0.01503250
[3,] 0.04924166
[4,] -0.01254111
[5,] -0.05570035
[6,] 0.05775536
```

10.5.2 Using the residuals command for other summaries

The residuals command will also let you get ordinary residuals, leverage values and dffits values, which are the normalized differences in predicted values when observations are omitted. See ?residuals.ols for more details.

```
temp <- data.frame(area = 1:60)
temp$residual <- residuals(newmod, type = "ordinary")
temp$leverage <- residuals(newmod, type = "hat")
temp$dffits <- residuals(newmod, type = "dffits")
tbl_df(temp)</pre>
```

```
# A tibble: 60 x 4
   area residual leverage dffits
  <int>
          <dbl>
                    <dbl>
                            <dbl>
         -13.3
                   0.0929 -0.119
1
      1
2
      2
           81.0
                   0.0941 0.766
3
      3
           28.8
                   0.266 0.539
4
      4
          -12.5
                   0.117 -0.128
5
      5
           27.8
                   0.204
                          0.419
6
      6
          -40.4
                   0.416 - 1.20
7
      7
           37.0
                   0.207
                           0.568
8
          -14.3
                   0.145 - 0.169
9
      9
           66.6
                   0.0863 0.587
10
     10
          - 4.96
                   0.0997 -0.0460
# ... with 50 more rows
ggplot(temp, aes(x = area, y = dffits)) +
   geom_point() +
   geom_line()
```



It appears that point 37 has the largest (positive) dffits value. Recall that point 37 seemed influential on several predictors and the intercept term. Point 32 has the smallest (or largest negative) dffits, and also appears to have been influential on several predictors and the intercept.

which.max(temp\$dffits)

[1] 37

which.min(temp\$dffits)

[1] 32

10.6 Model Validation and Correcting for Optimism

In 431, we learned about splitting our regression models into **training** samples and **test** samples, performing variable selection work on the training sample to identify two or three candidate models (perhaps via a stepwise approach), and then comparing the predictions made by those models in a test sample.

At the final project presentations, I mentioned (to many folks) that there was a way to automate this process a bit in 432, that would provide some ways to get the machine to split the data for you multiple times, and then average over the results, using a bootstrap approach. This is it.

The validate function allows us to perform cross-validation of our models for some summary statistics (and then correct those statistics for optimism in describing likely predictive accuracy) in an easy way.

validate develops:

- Resampling validation with or without backward elimination of variables
- Estimates of the *optimism* in measures of predictive accuracy

• Estimates of the intercept and slope of a calibration model

```
(observed y) = Intercept + Slope (predicted y)
```

with the following code...

```
set.seed(432002); validate(newmod, method = "boot", B = 40)
```

	index.orig	training	test	optimism	${\tt index.corrected}$	n
R-square	0.6989	0.7500	0.5964	0.1536	0.5452	40
MSE	1145.7800	888.2564	1535.8277	-647.5714	1793.3514	40
g	58.9614	58.9323	55.2085	3.7238	55.2376	40
Intercept	0.0000	0.0000	86.5968	-86.5968	86.5968	40
Slope	1.0000	1.0000	0.9088	0.0912	0.9088	40

So, for R-square we see that our original estimate was 0.6989

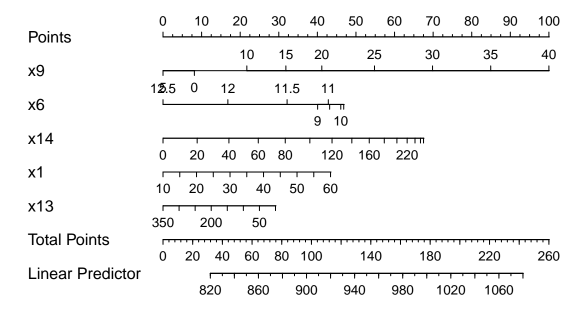
- Our estimated R-square across n = 40 training samples was 0.7500, but in the resulting tests, the average R-square was only 0.5964
- This suggests an optimism of 0.7500 0.5964 = 0.1536 (after rounding).
- We then apply that optimism to obtain a new estimate of R² corrected for overfitting, at 0.5452, which is probably a better estimate of what our results might look like in new data that were similar to (but not the same as) the data we used in building newmod than our initial estimate of 0.6989

We also obtain optimism-corrected estimates of the mean squared error (square of the residual standard deviation), the g index, and the intercept and slope of the calibration model. The "corrected" slope is a shrinkage factor that takes overfitting into account.

10.7 Building a Nomogram for Our Model

Another nice feature of an ols model object is that we can picture the model with a **nomogram** easily. Here is model **newmod**.

```
plot(nomogram(newmod))
```



For this model, we can use this plot to predict y as follows:

- 1. find our values of x9 on the appropriate line
- 2. draw a vertical line up to the points line to count the points associated with our subject
- 3. repeat the process to obtain the points associated with x6, x14, x1, and x13. Sum the points.
- 4. draw a vertical line down from that number in the Total Points line to estimate y (the Linear Predictor) = Age-Adjusted Mortality Rate.

The impact of the non-linearity is seen in the x6 results, for example, which turn around from 9-10 to 11-12. We also see non-linearity's effects in the scales of the non-linear terms in terms of points awarded.

An area with a combination of predictor values leading to a total of 100 points, for instance, would lead to a prediction of a Mortality Rate near 905. An area with a total of 140 points would have a predicted Mortality Rate of 955, roughly.

Chapter 11

Other Variable Selection Strategies

11.1 Why not use stepwise procedures?

- 1. The \mathbb{R}^2 for a model selected in a stepwise manner is biased, high.
- 2. The coefficient estimates and standard errors are biased.
- 3. The p values for the individual-variable t tests are too small.
- 4. In stepwise analyses of prediction models, the final model represented noise 20-74% of the time.
- 5. In stepwise analyses, the final model usually contained less than half of the actual number of real predictors.
- 6. It is not logical that a population regression coefficient would be exactly zero just because its estimate was not statistically significant.

This last comment applies to things like our "best subsets" approach as well as standard stepwise procedures.

Sander Greenland's comments on parsimony and stepwise approaches to model selection are worth addressing...

- Stepwise variable selection on confounders leaves important confounders uncontrolled.
- Shrinkage approaches (like ridge regression and the lasso) are far superior to variable selection.
- Variable selection does more damage to confidence interval widths than to point estimates.

If we are seriously concerned about **overfitting** - winding up with a model that doesn't perform well on new data - then stepwise approaches generally don't help.

Vittinghoff et al. (2012) suggest four strategies for minimizing the chance of overfitting

- 1. Pre-specify well-motivated predictors and how to model them.
- 2. Eliminate predictors without using the outcome.
- 3. Use the outcome, but cross-validate the target measure of prediction error.
- 4. Use the outcome, and **shrink** the coefficient estimates.

The best subsets methods we have studied either include a variable or drop it from the model. Often, this choice is based on only a tiny difference in the quality of a fit to data.

- Harrell (2001): not reasonable to assume that a population regression coefficient would be exactly zero just because it failed to meet a criterion for significance.
- Brad Efron has suggested that a stepwise approach is "overly greedy, impulsively eliminating covariates which are correlated with other covariates."

So, what's the alternative?

11.2 Ridge Regression

Ridge regression involves a more smooth transition between useful and not useful predictors which can be obtained by constraining the overall size of the regression coefficients.

Ridge regression assumes that the regression coefficients (after normalization) should not be very large. This is reasonable to assume when you have lots of predictors and you believe *many* of them have some effect on the outcome.

Pros:

- 1. Some nice statistical properties
- 2. Can be calculated using only standard least squares approaches, so it's been around for a while.
- 3. Available in the MASS package.

Ridge regression takes the sum of the squared estimated standardized regression coefficients and constrains that sum to only be as large as some value k.

$$\sum \hat{\beta_j}^2 \le k.$$

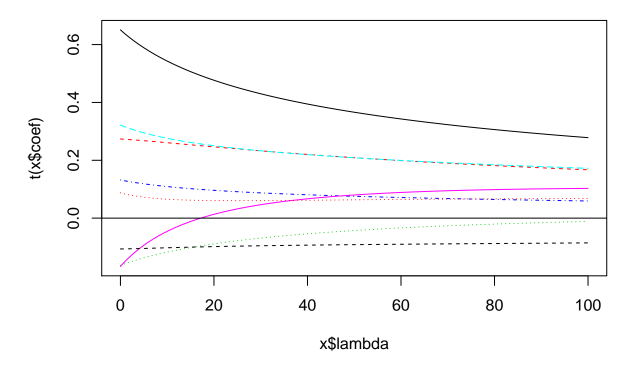
The value k is one of several available measures of the amount of shrinkage, but the main one used in the MASS package is a value λ . As λ increases, the amount of shrinkage goes up, and k goes down.

11.2.1 Assessing a Ridge Regression Approach

We'll look at a plot produced by the lm.ridge function for a ridge regression for the prostate cancer study we worked on when studying Stepwise Regression and Best Subsets methods earlier.

- Several (here 101) different values for λ , our shrinkage parameter, will be tested.
- Results are plotted so that we see the coefficients across the various (standardized) predictors.
 - Each selection of a λ value implies a different vector of covariate values across the predictors we are studying.
 - The idea is to pick a value of λ for which the coefficients seem relatively stable.

Ridge Regression for prost data



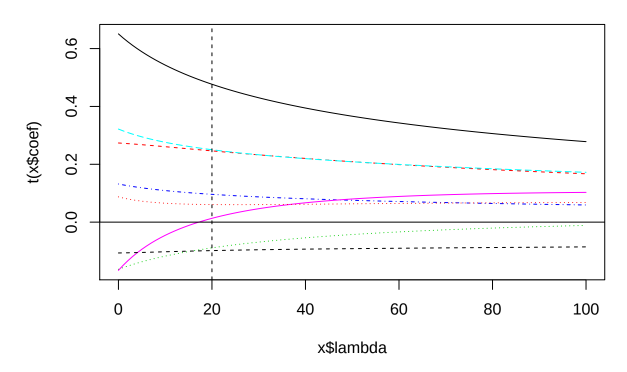
Usually, you need to use trial and error to decide the range of λ to be tested. Here, 0:100 means going from 0 (no shrinkage) to 100 in steps of 1.

11.2.2 The lm.ridge plot - where do coefficients stabilize?

```
Does \lambda = 20 seem like a stable spot here?
```

```
x <- lm.ridge(prost$lpsa ~ preds, lambda = 0:100)
plot(x)
title("Ridge Regression for prost data")
abline(h = 0)
abline(v=20, lty=2, col="black")</pre>
```





The coefficients at $\lambda=20$ can be determined from the lm.ridge output. These are fully standardized coefficients. The original predictors are centered by their means and then scaled by their standard deviations and the outcome has also been centered, in these models.

```
round(x$coef[,20],3)
```

predssvi_f	predsbph_f	predsage	predslweight	predslcavol
0.252	0.097	-0.091	0.248	0.482
		predspgg45	predsgleason_f	predslcp
		0.061	-0.099	0.009

Was an intercept used?

MASS::select(x)

x\$Inter

[1] 1

Yes, it was. There is an automated way to pick λ . Use the select function in the MASS package:

```
modified HKB estimator is 4.210238 modified L-W estimator is 3.32223 smallest value of GCV at 6
```

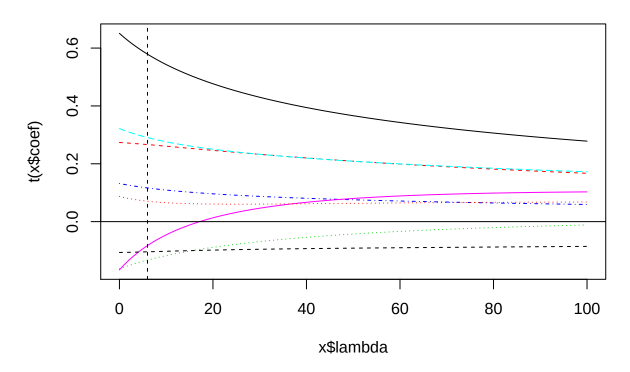
I'll use the GCV = generalized cross-validation to select $\lambda = 6$ instead.

```
x <- lm.ridge(prost$lpsa ~ preds, lambda = 0:100)
plot(x)
title("Ridge Regression for prost data")
abline(h = 0)</pre>
```

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abline(v=6, lty=2, col="black")

Ridge Regression for prost data



x\$coef[,6] predssvi_f predslcavol predslweight predsage predsbph_f 0.29491008 0.58911149 0.26773757 -0.13715070 0.11862949 predslcp predsgleason_f predspgg45 -0.09389545 -0.10477578 0.07250609

11.2.3 Ridge Regression: The Bottom Line

The main problem with ridge regression is that all it does is shrink the coefficient estimates, but it's not so useful in practical settings because it still includes all variables.

- 1. It's been easy to do ridge regression for many years, so you see it occasionally in the literature.
- 2. It leads to the **lasso**, which incorporates the positive features of shrinking regression coefficients with the ability to wisely select some variables to be eliminated from the predictor pool.

11.3 The Lasso

The lasso works by takes the sum of the absolute values of the estimated standardized regression coefficients and constrains it to only be as large as some value k.

$$\sum |\hat{\beta_j}| \le k.$$

This looks like a minor change, but it's not.

11.3.1 Consequences of the Lasso Approach

- 1. In ridge regression, while the individual coefficients shrink and sometimes approach zero, they seldom reach zero and are thus excluded from the model. With the lasso, some coefficients do reach zero and thus, those predictors do drop out of the model.
 - So the lasso leads to more parsimonious models than does ridge regression.
 - Ridge regression is a method of shrinkage but not model selection. The lasso accomplishes both tasks.
- 2. If k is chosen to be too small, then the model may not capture important characteristics of the data. If k is too large, the model may over-fit the data in the sample and thus not represent the population of interest accurately.
- 3. The lasso is far more difficult computationally than ridge regression (the problem requires an algorithm called least angle regression published in 2004), although R has a library (lars) which can do the calculations pretty efficiently.

The lasso is not an acronym, but rather refers to cowboys using a rope to pull cattle from the herd, much as we will pull predictors from a model.

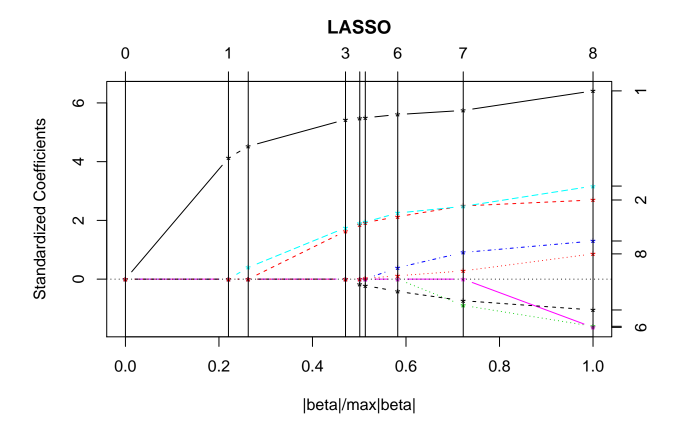
11.3.2 How The Lasso Works

The lars package lets us compute the lasso coefficient estimates and do cross-validation to determine the appropriate amount of shrinkage. The main tool is a pair of graphs.

- 1. The first plot shows what coefficients get selected as we move from constraining all of the coefficients to zero (complete shrinkage) towards fewer constraints all the way up to ordinary least squares, showing which variables are included in the model at each point.
- 2. The second plot suggests where on the first plot we should look for a good model choice, according to a cross-validation approach.

```
## requires lars package
lasso1 <- lars(preds, prost$lpsa, type="lasso")
plot(lasso1)</pre>
```

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- The y axis shows standardized regression coefficients.
 - The lars package standardizes all variables so the shrinkage doesn't penalize some coefficients because of their scale.
- The x-axis is labeled |beta|/max|beta|.
 - This ranges from 0 to 1.
 - 0 means that the sum of the $|\hat{\beta}_i|$ is zero (completely shrunk)
 - 1 means the ordinary least squares unbiased estimates.

The lasso graph starts at constraining all of the coefficients to zero, and then moves toward ordinary least squares.

Identifiers for the predictors (numbers) are shown to the right of the graph.

The vertical lines in the lasso plot show when a variable has been eliminated from the model, and in fact these are the only points that are actually shown in the default lasso graph. The labels on the top of the graph tell you how many predictors are in the model at that stage.

summary(lasso1)

```
LARS/LASSO
```

```
Call: lars(x = preds, y = prost$lpsa, type = "lasso")
  Df
         Rss
   1 127.918 168.1835
0
   2
      76.392
               64.1722
1
   3
               53.5293
2
      70.247
3
   4
      50.598
               15.1017
4
   5
      49.065
               13.9485
5
      48.550
               14.8898
   6
6
      46.284
               12.2276
```

7 8 44.002 9.5308 8 9 42.772 9.0000

Based on the C_p statistics, it looks like the improvements continue throughout, and don't really finish happening until we get pretty close to the full model with 9 df.

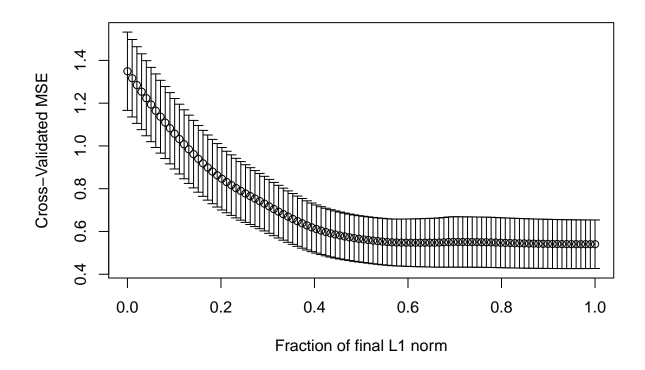
11.3.3 Cross-Validation with the Lasso

Normally, cross-validation methods are used to determine how much shrinkage should be used. We'll use the cv.lars function.

- 10-fold (K = 10) cross-validation
 - the data are randomly divided into 10 groups.
 - Nine groups are used to predict the remaining group for each group in turn.
 - Overall prediction performance is computed, and the machine calculates a cross-validation criterion (mean squared error) and standard error for that criterion.

The cross-validation plot is the second lasso plot.

```
set.seed(432)
lassocv <- cv.lars(preds, prost$lpsa, K=10)</pre>
```



default cv.lars K is 10

We're looking to minimize cross-validated mean squared error in this plot, which doesn't seem to happen until the fraction gets very close to 1.

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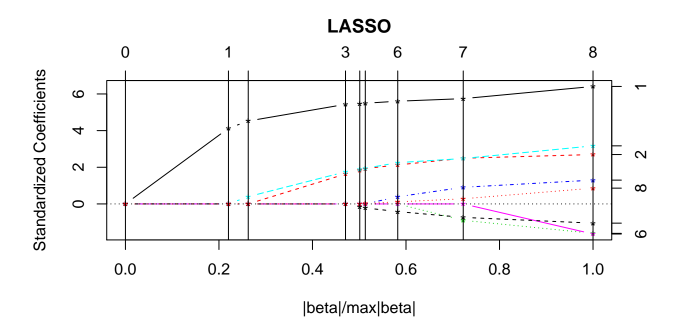
11.3.4 What value of the key fraction minimizes cross-validated MSE?

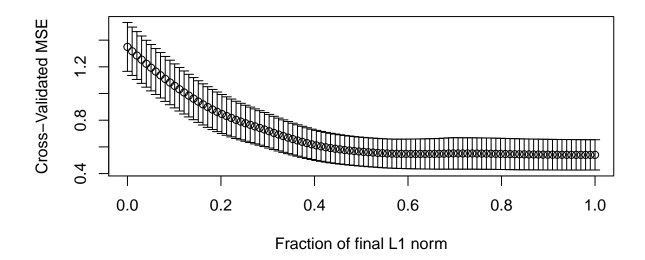
```
frac <- lassocv$index[which.min(lassocv$cv)]
frac</pre>
```

[1] 0.989899

The cross-validation plot suggests we use a fraction of about 0.3, that's suggesting a model with 4-5 predictors, based on the top LASSO plot.

```
par(mfrow=c(2,1))
lasso1 <- lars(preds, prost$lpsa, type="lasso")
plot(lasso1)
set.seed(432)
lassocv <- cv.lars(preds, prost$lpsa, K=10)</pre>
```





```
par(mfrow=c(1,1))
```

11.3.5 Coefficients for the Model Identified by the Cross-Validation

```
coef.cv <- coef(lasso1, s=frac, mode="fraction")
round(coef.cv,4)

lcavol lweight age bph_f svi_f lcp gleason_f
0.5529   0.6402  -0.0217   0.1535   0.7750  -0.1155  -0.1826
   pgg45</pre>
```

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0.0030

So the model suggested by the lasso still includes all sight of these predictors.

11.3.6 Obtaining Fitted Values from Lasso

```
fits.cv <- predict.lars(lasso1, preds, s=frac,</pre>
                        type="fit", mode="fraction")
fits.cv
$5
[1] 0.989899
$fraction
[1] 0.989899
$mode
[1] "fraction"
$fit
 [1] 0.7995838 0.7493971 0.5111634 0.6098520 1.7001847 0.8338020 1.8288518
 [8] 2.1302316 1.2487955 1.2661752 1.4704969 0.7782005 2.0755860 1.9129272
[15] 2.1533975 1.8124981 1.2713610 2.3993624 1.3232566 1.7709029 1.9757841
[22] 2.7451649 1.1658326 2.4825521 1.8036338 1.9112578 2.0144298 1.7829219
[29] 1.9706111 2.1688199 2.0377131 1.8657882 1.6955904 1.3580186 1.0516394
[36] 2.9097450 2.1898622 1.0454123 3.8896481 1.7971270 2.0932871 2.3253395
[43] 2.0809295 2.5303655 2.4451523 2.5827203 4.0692397 2.6845105 2.7034959
[50] 1.9590266 2.4522082 2.9801227 2.1902084 3.0559124 3.3447025 2.9765233
[57] 1.7620182 2.3424646 2.2856404 2.6188548 2.3056410 3.5568662 2.9756755
[64] 3.6764122 2.5097586 2.6579014 2.9482717 3.0892917 1.5113015 3.0282296
[71] 3.2887119 2.1083273 2.8889223 3.4903026 3.6959516 3.6070031 3.2749993
[78] 3.4518575 3.4049180 3.1814731 2.0496216 2.8986175 3.6743113 3.3292860
[85] 2.6965297 3.8339856 2.9892543 3.0555536 4.2903885 3.0986508 3.3784385
[92] 4.0205201 3.8309974 4.7531590 3.6290575 4.1347645 4.0982744
```

11.3.7 Complete Set of Fitted Values from the Lasso

```
round(fits.cv$fit,3)

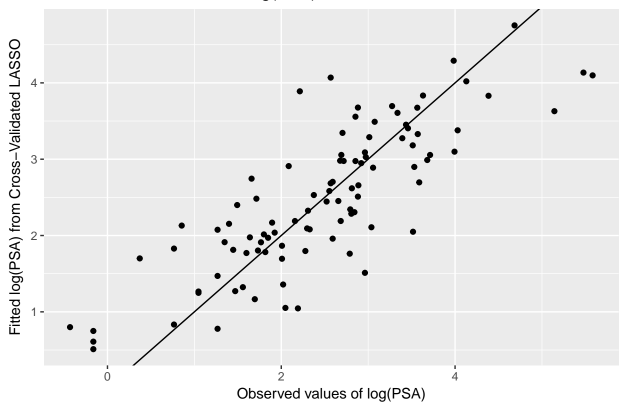
[1] 0.800 0.749 0.511 0.610 1.700 0.834 1.829 2.130 1.249 1.266 1.470 [12] 0.778 2.076 1.913 2.153 1.812 1.271 2.399 1.323 1.771 1.976 2.745 [23] 1.166 2.483 1.804 1.911 2.014 1.783 1.971 2.169 2.038 1.866 1.696 [34] 1.358 1.052 2.910 2.190 1.045 3.890 1.797 2.093 2.325 2.081 2.530 [45] 2.445 2.583 4.069 2.685 2.703 1.959 2.452 2.980 2.190 3.056 3.345 [56] 2.977 1.762 2.342 2.286 2.619 2.306 3.557 2.976 3.676 2.510 2.658 [67] 2.948 3.089 1.511 3.028 3.289 2.108 2.889 3.490 3.696 3.607 3.275 [78] 3.452 3.405 3.181 2.050 2.899 3.674 3.329 2.697 3.834 2.989 3.056 [89] 4.290 3.099 3.378 4.021 3.831 4.753 3.629 4.135 4.098
```

To assess the quality of these predictions, we might plot them against the observed values of our outcome (lpsa), or we might look at residuals vs. these fitted values.

```
resid = actual - fitted)

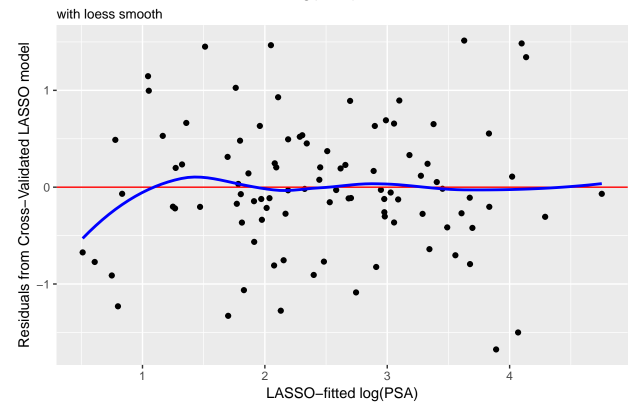
ggplot(prost_lasso_res, aes(x = actual, y = fitted)) +
    geom_point() +
    geom_abline(slope = 1, intercept = 0) +
    labs(y = "Fitted log(PSA) from Cross-Validated LASSO",
        x = "Observed values of log(PSA)",
        title = "Fitted vs. Actual Values of log(PSA)")
```

Fitted vs. Actual Values of log(PSA)



```
ggplot(prost_lasso_res, aes(x = fitted, y = resid)) +
   geom_point() +
   geom_hline(yintercept = 0, col = "red") +
   geom_smooth(method = "loess", col = "blue", se = F) +
   labs(x = "LASSO-fitted log(PSA)",
        y = "Residuals from Cross-Validated LASSO model",
        title = "Residuals vs. Fitted Values of log(PSA) from LASSO",
        subtitle = "with loess smooth")
```

Residuals vs. Fitted Values of log(PSA) from LASSO



11.3.8 When is the Lasso Most Useful?

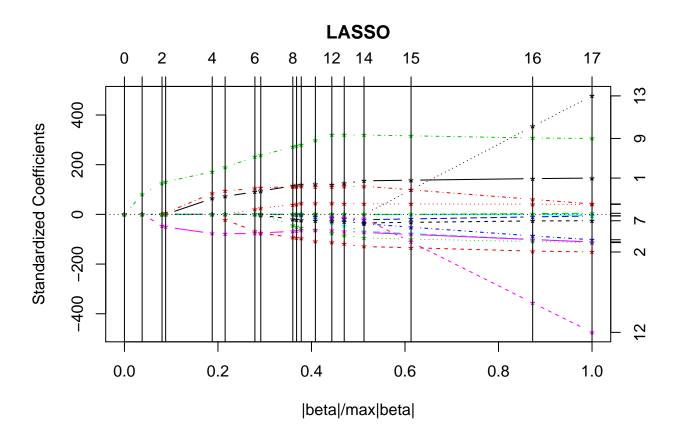
As Faraway (2015) suggests, the lasso is particularly useful when we believe the effects are sparse, in the sense that we believe that few of the many predictors we are evaluating have a meaningful effect.

Consider, for instance, the analysis of gene expression data, where we have good reason to believe that only a small number of genes have an influence on our response of interest.

Or, in medical claims data, where we can have thousands of available codes to search through that may apply to some of the people included in a large analysis relating health care costs to outcomes.

11.4 Applying the Lasso to the pollution data

Let's consider the lasso approach in application to the pollution data we've seen previously. Recall that we have 60 observations on an outcome, y, and 15 predictors, labeled x1 through x15.



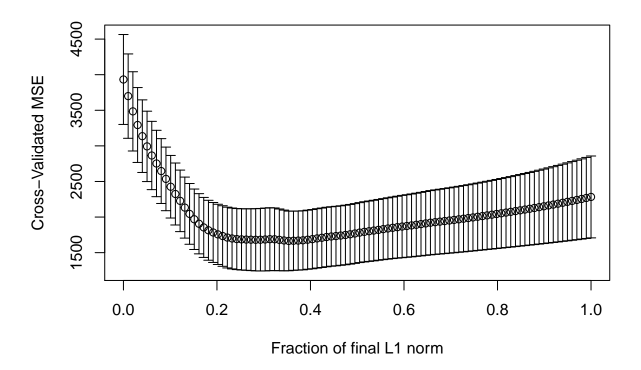
summary(lasso_p1)

```
LARS/LASSO
```

```
Call: lars(x = preds, y = pollution$y, type = "lasso")
   Df
         Rss
                    Ср
    1 228311 129.1367
    2 185419
               95.9802
1
2
    3 149370
               68.4323
3
      143812
               65.8764
4
    5
       92077
               25.4713
5
       83531
               20.4668
    6
6
    7
       69532
               10.9922
7
               11.4760
    8
       67682
8
    9
       60689
                7.7445
9
   10
       60167
                9.3163
       59609
               10.8588
10 11
11 12
       58287
               11.7757
       57266
12 13
               12.9383
13 14
       56744
               14.5107
14 13
       56159
               12.0311
15 14
       55238
               13.2765
16 15
       53847
               14.1361
17 16
       53681
```

Based on the C_p statistics, it looks like the big improvements occur somewhere around the move from 6 to 7 df. Let's look at the cross-validation

```
set.seed(432012)
pollution_lassocv <- cv.lars(preds, pollution$y, K=10)</pre>
```

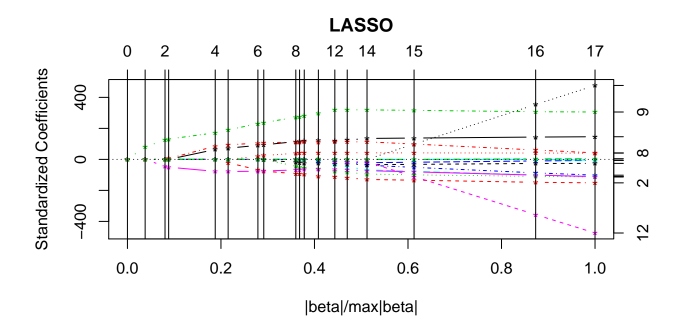


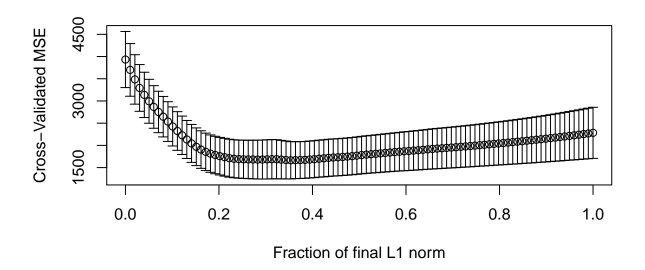
Here it looks like cross-validated MSE happens somewhere between a fraction of 0.2 and 0.4.

```
frac <- pollution_lassocv$index[which.min(pollution_lassocv$cv)]
frac</pre>
```

```
[1] 0.3535354
```

```
par(mfrow=c(2,1))
lasso_p1 <- lars(preds, pollution$y, type="lasso")
plot(lasso_p1)
set.seed(432012)
pollution_lassocv <- cv.lars(preds, pollution$y, K=10)</pre>
```





par(mfrow=c(1,1))

It looks like a model with 6-8 predictors will be the most useful. The cross-validated coefficients are as follows:

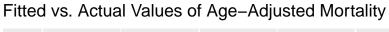
```
poll.cv <- coef(lasso_p1, s=frac, mode="fraction")
round(poll.cv,3)</pre>
```

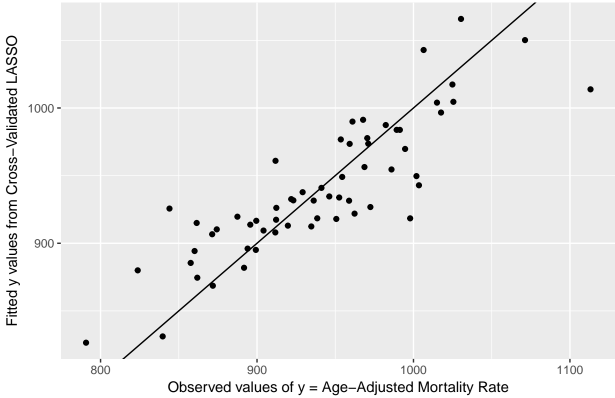
x1	x2	x3	x4	x5	x6	x7	x8	x9
1.471	-1.164	-1.102	0.000	0.000	-10.610	-0.457	0.003	3.918
x10	x11	x12	x13	x14	x15			
0.000	0.000	0.000	0.000	0.228	0.000			

Note that by this cross-validated lasso selection, not only are the coefficients for the 8 variables remaining in the model shrunken, but variables x4, x5, x10, x11, x12, x13 and x15 are all dropped from the model, and model x8 almost is, as well.

```
[1] 932.627 918.415 921.904 987.396 1050.184 1065.837 912.424 [8] 916.605 949.647 926.168 996.625 1017.362 977.730 954.550 [15] 931.455 894.263 931.551 868.599 973.471 940.937 881.867 [22] 906.666 973.609 919.640 933.821 956.352 913.018 925.650 [29] 874.528 983.829 1042.870 915.002 937.760 885.464 989.947 [36] 931.709 1013.795 969.729 1003.962 983.813 896.042 918.446 [43] 934.609 1004.565 910.273 976.747 831.132 907.996 826.485 [50] 895.082 909.398 917.969 926.777 917.381 991.266 879.972 [57] 942.867 913.737 960.952 949.030
```

Here's a plot of the actual pollution y values, against these fitted values.

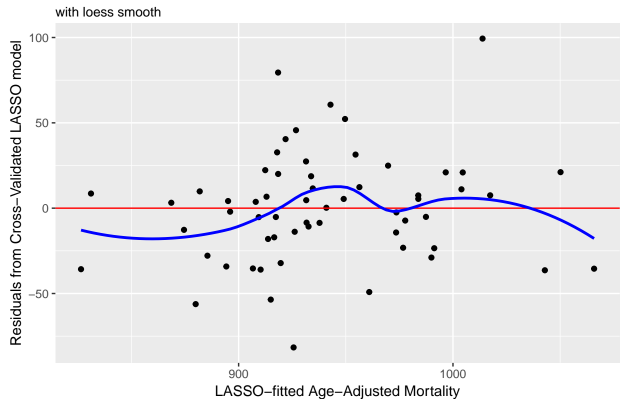




And now, here's a plot or residuals vs. fitted values.

```
ggplot(poll_lasso_res, aes(x = fitted, y = resid)) +
    geom_point() +
    geom_hline(yintercept = 0, col = "red") +
    geom_smooth(method = "loess", col = "blue", se = F) +
    labs(x = "LASSO-fitted Age-Adjusted Mortality",
        y = "Residuals from Cross-Validated LASSO model",
        title = "Residuals vs. Fitted Values of Age-Adjusted Mortality from LASSO",
        subtitle = "with loess smooth")
```

Residuals vs. Fitted Values of Age-Adjusted Mortality from LASSO



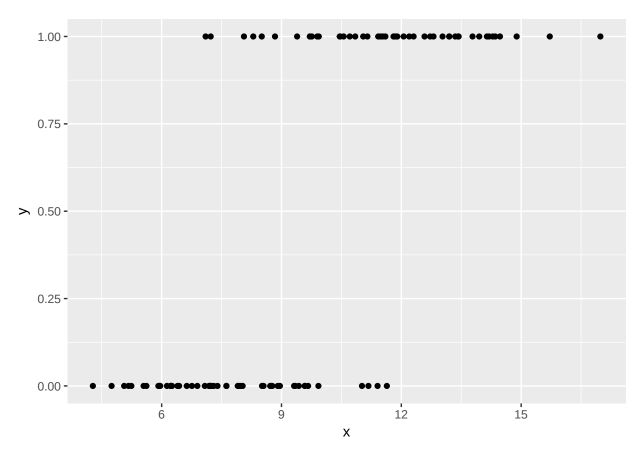
Chapter 12

Logistic Regression: The Foundations

Sources for this material include Harrell (2001), Harrell (2018), Ramsey and Schafer (2002) (chapters 20-21), Vittinghoff et al. (2012) (chapter 5) and Faraway (2006) (chapter 2).

12.1 A First Attempt: A Linear Probability Model

Suppose we want to predict a binary outcome which takes on the value 1 or 0, based on a single quantitative predictor. Let y be a 1/0 outcome, and x be a quantitative predictor in the following simulation.



Now, we want to use our variable x here to predict our variable y (which takes on the values 0 and 1).

One approach to doing this would be a linear probability model, as follows:

```
mod12a \leftarrow lm(y \sim x, data = sim12)
summary(mod12a)
```

Call:

 $lm(formula = y \sim x, data = sim12)$

Residuals:

Min 1Q Median 3Q -0.74104 - 0.23411 - 0.02894 0.23117 0.83153

Coefficients:

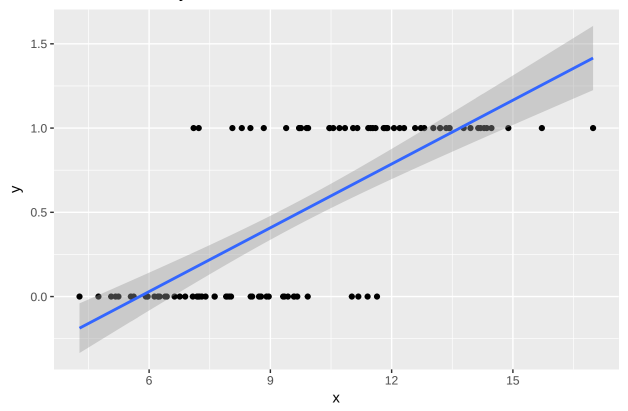
```
Estimate Std. Error t value Pr(>|t|)
                       0.12272 -5.929 4.57e-08 ***
(Intercept) -0.72761
           0.12620
                       0.01219 10.349 < 2e-16 ***
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

Residual standard error: 0.3491 on 98 degrees of freedom Multiple R-squared: 0.5222, Adjusted R-squared: 0.5173 F-statistic: 107.1 on 1 and 98 DF, p-value: < 2.2e-16

Here's a picture of this model. What's wrong here?

```
ggplot(sim12, aes(x = x, y = y)) +
   geom_point() +
   geom_smooth(method = "lm") +
   labs(title = "Linear Probability Model")
```

Linear Probability Model



If y can only take the values 0 and 1 (or, more precisely, if we're trying to predict the value $\pi = \Pr(y = 1)$) then what do we do with the predictions that are outside the range of (0, 1)?

12.2 Logistic Regression

Logistic regression is the most common model used when the outcome is binary. Our response variable is assumed to take on two values, zero or one, and we then describe the probability of a "one" response, given a linear function of explanatory predictors. We use logistic regression rather than linear regression for predicting binary outcomes. Linear regression approaches to the problem of predicting probabilities are problematic for several reasons - not least of which being that they predict probabilities greater than one and less than zero. There are several available alternatives, including probit regression and binomial regression, for the problem of predicting a binary outcome.

Logistic regression is part of a class called **generalized linear models** which extend the linear regression model in a variety of ways. There are also several extensions to the logistic regression model, including multinomial logistic regression (which is used for nominal categorical outcomes with more than two levels) and ordered logistic regression (used for ordered multi-categorical outcomes.) The methods involved in binary logistic regression may also be extended to the case where the outcomes are proportions based on counts, often through grouped binary responses (the proportion of cells with chromosomal aberrations, or the proportion of subjects who develop a particular condition.)

Although the models are different in some crucial ways, the practical use of logistic regression tracks well with much of what we've learned about linear regression.

12.3 The Logistic Regression Model

A generalized linear model (or GLM) is a probability model in which the mean of an outcome is related to predictors through a regression equation. A link function g is used to relate the mean, μ , to a linear regression of the predictors $X_1, X_2, ..., X_k$.

$$g(\mu) = \beta_0 + \beta_1 X_1 + \beta_2 X_2 + \dots + \beta_k X_k$$

In the case of a logistic regression model,

- the mean μ of our 0/1 outcome is represented by π which describes the probability of a "1" outcome.
- the linking function we use in logistic regression makes use of the logit function, which is built on the natural logarithm.

12.4 The Link Function

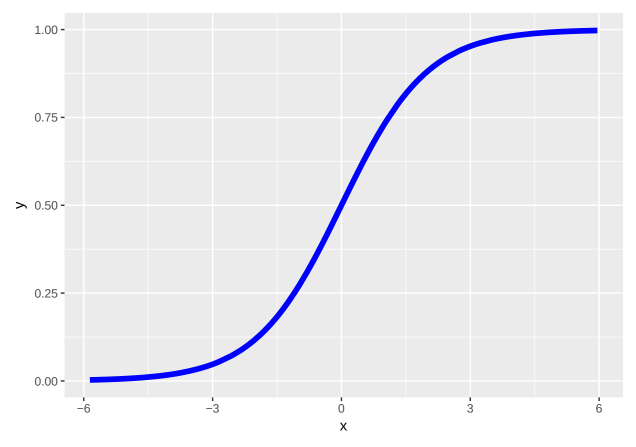
Logistic regression is a non-linear regression approach, since the equation for the mean of the 0/1 Y values conditioned on the values of our predictors $X_1, X_2, ..., X_k$ turns out to be non-linear in the β coefficients. Its nonlinearity, however, is solely found in its link function, hence the term *generalized* linear model.

The particular link function we use in logistic regression is called the **logit link**.

$$logit(\pi) = log\left(\frac{\pi}{1-\pi}\right) = \beta_0 + \beta_1 X_1 + \beta_2 X_2 + \dots + \beta_k X_k$$

The inverse of the logit function is called the **logistic function**. If $logit(\pi) = \eta$, then $\pi = \frac{exp(\eta)}{1 + exp(\eta)}$

The plot below displays the logistic function $y = \frac{e^x}{1 + e^x}$



As you can see in the figure above, the logistic function $\frac{e^x}{1+e^x}$ takes any value x in the real numbers and returns a value between 0 and 1.

12.5 The logit or log odds

We usually focus on the logit in statistical work, which is the inverse of the logistic function.

- If we have a probability $\pi < 0.5$, then $logit(\pi) < 0$.
- If our probability $\pi > 0.5$, then $logit(\pi) > 0$.
- Finally, if $\pi = 0.5$, then $logit(\pi) = 0$.

12.6 Interpreting the Coefficients of a Logistic Regression Model

The critical thing to remember in interpreting a logistic regression model is that the logit is the log odds function. Exponentiating the logit yields the odds.

So, suppose we have a yes/no outcome variable, where yes = 1, and no = 0, and $\pi = Pr(y = 1)$. Our model holds that:

$$logit(\pi) = log\left(\frac{\pi}{1-\pi}\right) = \beta_0 + \beta_1 X_1 + \beta_2 X_2 + \dots + \beta_k X_k$$

The odds of a yes response (the odds that Y = 1) at the level $X_1, X_2, ..., X_k$ are:

$$Odds(Y = 1) = exp(\beta_0 + \beta_1 X_1 + \beta_2 X_2 + ... + \beta_k X_k)$$

The **probability** of a yes response $(Pr(y = 1), or \pi)$ is just

$$\pi = Pr(Y = 1) = \frac{Odds(Y = 1)}{1 + Odds(Y = 1)} = \frac{exp(\beta_0 + \beta_1 X_1 + \beta_2 X_2 + \dots + \beta_k X_k)}{1 + exp(\beta_0 + \beta_1 X_1 + \beta_2 X_2 + \dots + \beta_k X_k)}$$

12.7 The Logistic Regression has non-constant variance

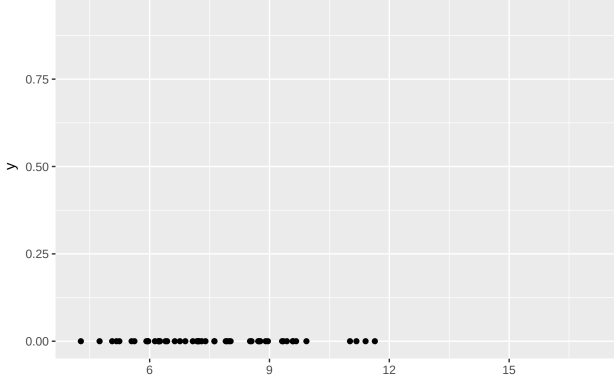
In ordinary least squares regression, the variance $Var(Y|X_1, X_2, ..., X_k) = \sigma^2$ is a constant that does not depend on the predictor values. This is not the case in logistic regression. The mean and variance specifications of the logistic regression model are quite different.

$$logit(\pi) = log\left(\frac{\pi}{1-\pi}\right) = \beta_0 + \beta_1 X_1 + \beta_2 X_2 + ... + \beta_k X_k \mu[Y|X_1,...,X_k] = \pi, Var[Y|X_1,...,X_k] = \pi(1-\pi)$$

The variance is now a function of the mean, and contains no additional parameter for us to estimate.

12.8 Fitting a Logistic Regression Model to our Simulated Data

Recall the sim12 data we built earlier.



Х

Here is the fitted logistic regression model.

```
model12b <- glm(y ~ x, data = sim12, family = binomial)
model12b</pre>
```

Call: glm(formula = y ~ x, family = binomial, data = sim12)

Coefficients:

(Intercept) x -9.1955 0.9566

Degrees of Freedom: 99 Total (i.e. Null); 98 Residual

Null Deviance: 138.6

Residual Deviance: 70.03 AIC: 74.03

The logistic regression equation is:

$$logit(Pr(y=1)) = log\left(\frac{Pr(y=1)}{1 - Pr(y=1)}\right) = -9.1955 + 0.9566x$$

We can exponentiate the results of this model to get to an equation about odds, and eventually, a prediction about probabilities. Suppose, for instance, that we are interested in the prediction when x = 12.

$$logit(Pr(y=1)|X=12) = log\left(\frac{Pr(y=1)}{1 - Pr(y=1)}\right) = -9.1955 + 0.9566 * 12 = 2.2837$$

And we can also get this from the predict function applied to our model, although the predict approach retains a few more decimal places internally:

```
predict(model12b, newdata = data.frame(x = 12))
```

1 2.284069

$$Odds(Y = 1|X = 12) = exp(-9.20 + 0.96 * 12) = exp(2.2837) = 9.812921$$

```
exp(predict(model12b, newdata = data.frame(x = 12)))
```

9.81654

The estimated **probability** of a yes response $(Pr(y = 1), or \pi)$ if x = 12 is just

$$\pi = Pr(Y = 1|X = 12) = \frac{Odds(Y = 1|X = 12)}{1 + Odds(Y = 1|X = 12)} = \frac{exp(-9.20 + 0.96x)}{1 + exp(-9.20 + 0.96x)} = \frac{9.812921}{1 + 9.812921} = 0.908$$

Does this work out?

```
exp(predict(model12b, newdata = data.frame(x = 12))) /
    (1 + exp(predict(model12b, newdata = data.frame(x = 12))))
```

0.907549

which is also directly available by running predict with type = "response".

```
predict(model12b, newdata = data.frame(x = 12), type = "response")

1
0.907549
```

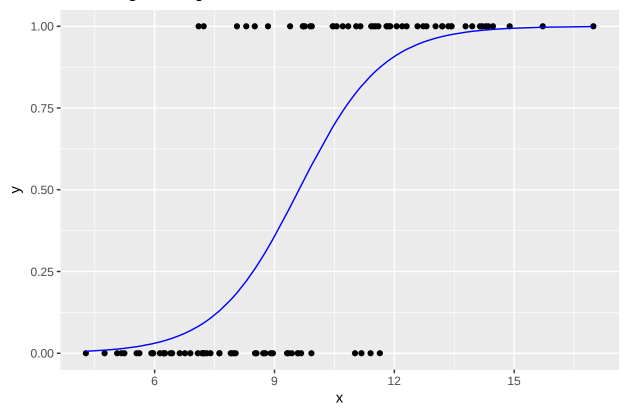
12.9 Plotting the Logistic Regression Model

We can use the augment function from the broom package to get our fitted probabilities included in the data.

```
m12b.aug <- augment(model12b, sim12, type.predict = "response")

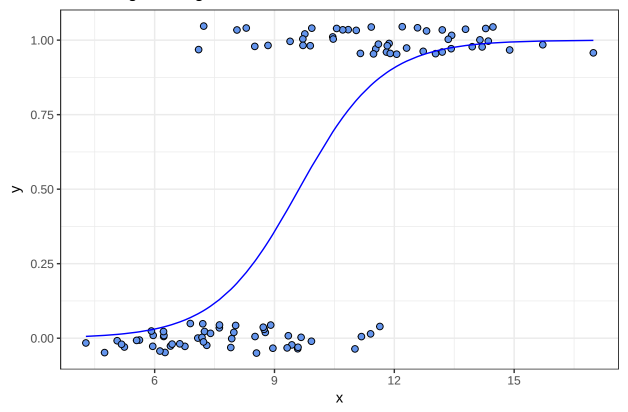
ggplot(m12b.aug, aes(x = x, y = y)) +
    geom_point() +
    geom_line(aes(x = x, y = .fitted), col = "blue") +
    labs(title = "Fitted Logistic Regression Model for sim12")</pre>
```

Fitted Logistic Regression Model for sim12



I'll add a little jitter on the vertical scale to the points, so we can avoid overlap, and also make the points a little bigger.

Fitted Logistic Regression for sim12



All right, it's time to move on to fitting models. We'll do that in the next Chapter.

Chapter 13

Logistic Regression and the resect data

13.1 The resect data

My source for these data was Riffenburgh (2006). The data describe 134 patients who had undergone resection of the tracheal carina (most often this is done to address tumors in the trachea), and the resect.csv data file contains the following variables:

```
• id = a patient ID \#,
```

- age = the patient's age at surgery,
- prior = prior tracheal surgery (1 = yes, 0 = no),
- resection = extent of the resection (in cm),
- intubated = whether intubation was required at the end of surgery (1 = yes, 0 = no), and
- died = the patient's death status (1 = dead, 0 = alive).

```
resect %>% group_by(died) %>% skim(-id)
```

Skim summary statistics

n obs: 134
n variables: 6
group variables: died

Variable type: integer

```
died variable missing complete
                                                sd p0 p25 median p75 p100
                                        mean
                                   n
                             117 117 48.05 16.01
                                                                   62
                                                                         80
   0
           age
                      0
   0 intubated
                      0
                             117 117 0.068 0.25
                                                                         1
                      0
   0
         prior
                             117 117
                                      0.24
                                              0.43
                                                                    0
                                                                         1
   1
           age
                      0
                              17
                                  17 46.41
                                             14.46 26
                                                       33
                                                               46
                                                                   60
                                                                        66
   1 intubated
                      0
                                  17
                                      0.65
                                              0.49
                                                                         1
                              17
                                  17 0.35
                                              0.49 0
   1
                                                                         1
         prior
```

```
Variable type: numeric
```

```
died variable missing complete    n mean    sd p0 p25 median p75 p100
    0 resection    0    117 117 2.82 1.21 1 2    2.5 3.5 6
    1 resection    0    17 17 3.97 1    2 3.5 4 4.5 6
```

We have no missing data, and 17 of the 134 patients died. Our goal will be to understand the characteristics of the patients, and how they relate to the binary outcome of interest, death.

13.2 Running A Simple Logistic Regression Model

In the most common scenario, a logistic regression model is used to predict a binary outcome (which can take on the values 0 or 1.) We will eventually fit a logistic regression model in two ways.

- 1. Through the glm function in the base package of R (similar to lm for linear regression)
- 2. Through the lrm function available in the rms package (similar to ols for linear regression)

We'll focus on the glm approach first, and save the lrm ideas for later in this Chapter.

13.2.1 Logistic Regression Can Be Harder than Linear Regression

- Logistic regression models are fitted using the method of maximum likelihood in glm, which requires multiple iterations until convergence is reached.
- Logistic regression models are harder to interpret (for most people) than linear regressions.
- Logistic regression models don't have the same set of assumptions as linear models.
- Logistic regression outcomes (yes/no) carry much less information than quantitative outcomes. As a result, fitting a reasonable logistic regression requires more data than a linear model of similar size.
 - The rule I learned in graduate school was that a logistic regression requires 100 observations to fit an intercept plus another 15 observations for each candidate predictor. That's not terrible, but it's a very large sample size.
 - Frank Harrell recommends that 96 observations + a function of the number of candidate predictors (which depends on the amount of variation in the predictors, but 15 x the number of such predictors isn't too bad if the signal to noise ratio is pretty good) are required just to get reasonable confidence intervals around your predictions.
 - * In a twitter note, Frank suggests that 96 + 8 times the number of candidate parameters might be reasonable so long as the smallest cell of interest (combination of an outcome and a split of the covariates) is 96 or more observations.
 - Peduzzi et al. (1996) suggest that if we let π be the smaller of the proportions of "yes" or "no" cases in the population of interest, and k be the number of inputs under consideration, then $N = 10k/\pi$ is the minimum number of cases to include, except that if N < 100 by this standard, you should increase it to 100, according to Long (1997).
 - * That suggests that if you have an outcome that happens 10% of the time, and you are running a model with 3 predictors, then you could get away with $(10 \times 3)/(0.10) = 300$ observations, but if your outcome happened 40% of the time, you could get away with only $(10 \times 3)/(0.40) = 75$ observations, which you'd round up to 100.

13.3 Logistic Regression using glm

resection

0.7417

(Intercept)

-4.4337

We'll begin by attempting to predict death based on the extent of the resection.

Degrees of Freedom: 133 Total (i.e. Null); 132 Residual

Null Deviance: 101.9

Residual Deviance: 89.49 AIC: 93.49

Note that the logit link is the default approach with the binomial family, so we could also have used:

which yields the same model.

13.3.1 Interpreting the Coefficients of a Logistic Regression Model

Our model is:

$$logit(died = 1) = log\left(\frac{Pr(died = 1)}{1 - Pr(died = 1)}\right) = \beta_0 + \beta_1 x = -4.4337 + 0.7417 \times resection$$

The predicted log odds of death for a subject with a resection of 4 cm is:

$$log\left(\frac{Pr(died=1)}{1 - Pr(died=1)}\right) = -4.4337 + 0.7417 \times 4 = -1.467$$

The predicted odds of death for a subject with a resection of 4 cm is thus:

$$\frac{Pr(died=1)}{1 - Pr(died=1)} = e^{-4.4337 + 0.7417 \times 4} = e^{-1.467} = 0.2306$$

Since the odds are less than 1, we should find that the probability of death is less than 1/2. With a little algebra, we see that the predicted probability of death for a subject with a resection of 4 cm is:

$$Pr(died = 1) = \frac{e^{-4.4337 + 0.7417 \times 4}}{1 + e^{-4.4337 + 0.7417 \times 4}} = \frac{e^{-1.467}}{1 + e^{-1.467}} = \frac{0.2306}{1.2306} = 0.187$$

In general, we can frame the model in terms of a statement about probabilities, like this:

$$Pr(died = 1) = \frac{e^{\beta_0 + \beta_1 x}}{1 + e^{\beta_0 + \beta_1 x}} = \frac{e^{-4.4337 + 0.7417 \times resection}}{1 + e^{-4.4337 + 0.7417 \times resection}}$$

and so by substituting in values for resection, we can estimate the model's fitted probabilities of death.

13.3.2 Using predict to describe the model's fits

To obtain these fitted odds and probabilities in R, we can use the **predict** function.

- The default predictions are on the scale of the log odds. These predictions are also available through the type = "link" command within the predict function for a generalized linear model like logistic regression.
- Here are the predicted log odds of death for a subject (Sally) with a 4 cm resection and a subject (Harry) who had a 5 cm resection.

```
predict(res_modA, newdata = data_frame(resection = c(4,5)))
```

• We can also obtain predictions for each subject on the original response (here, probability) scale, backing out of the logit link.

So the predicted probability of death is 0.187 for Sally, the subject with a 4 cm resection, and 0.326 for Harry, the subject with a 5 cm resection.

13.3.3 Odds Ratio interpretation of Coefficients

Often, we will exponentiate the estimated slope coefficients of a logistic regression model to help us understand the impact of changing a predictor on the odds of our outcome.

```
exp(coef(res_modA))
```

```
(Intercept) resection 0.01186995 2.09947754
```

To interpret this finding, suppose we have two subjects, Harry and Sally. Harry had a resection that was 1 cm larger than Sally. This estimated coefficient suggests that the estimated odds for death associated with Harry is 2.099 times larger than the odds for death associated with Sally. In general, the odds ratio comparing two subjects who differ by 1 cm on the resection length is 2.099.

To illustrate, again let's assume that Harry's resection was 5 cm, and Sally's was 4 cm. Then we have:

$$log\left(\frac{Pr(Harrydied)}{1 - Pr(Harrydied)}\right) = -4.4337 + 0.7417 \times 5 = -0.7253, log\left(\frac{Pr(Sallydied)}{1 - Pr(Sallydied)}\right) = -4.4337 + 0.7417 \times 4 = -1.4667.$$

which implies that our estimated odds of death for Harry and Sally are:

$$Odds(Harrydied) = \frac{Pr(Harrydied)}{1 - Pr(Harrydied)} = e^{-4.4337 + 0.7417 \times 5} = e^{-0.7253} = 0.4842 Odds(Sallydied) = \frac{Pr(Sallydied)}{1 - Pr(Sallydied)} = e^{-4.4337 + 0.7417 \times 5} = e^{-0.7253} = 0.4842 Odds(Sallydied) = \frac{Pr(Sallydied)}{1 - Pr(Sallydied)} = e^{-4.4337 + 0.7417 \times 5} = e^{-0.7253} = 0.4842 Odds(Sallydied) = \frac{Pr(Sallydied)}{1 - Pr(Sallydied)} = e^{-4.4337 + 0.7417 \times 5} = e^{-0.7253} = 0.4842 Odds(Sallydied) = \frac{Pr(Sallydied)}{1 - Pr(Sallydied)} = e^{-4.4337 + 0.7417 \times 5} = e^{-0.7253} = 0.4842 Odds(Sallydied) = \frac{Pr(Sallydied)}{1 - Pr(Sallydied)} = e^{-4.4337 + 0.7417 \times 5} = e^{-0.7253} = 0.4842 Odds(Sallydied) = \frac{Pr(Sallydied)}{1 - Pr(Sallydied)} = e^{-4.4337 + 0.7417 \times 5} = e^{-0.7253} = 0.4842 Odds(Sallydied) = \frac{Pr(Sallydied)}{1 - Pr(Sallydied)} = e^{-4.4337 + 0.7417 \times 5} = e^{-0.7253} = 0.4842 Odds(Sallydied) = \frac{Pr(Sallydied)}{1 - Pr(Sallydied)} = e^{-4.4337 + 0.7417 \times 5} = e^{-0.7253} = 0.4842 Odds(Sallydied) = \frac{Pr(Sallydied)}{1 - Pr(Sallydied)} = e^{-4.4337 + 0.7417 \times 5} = e^{-0.7253} = 0.4842 Odds(Sallydied) = \frac{Pr(Sallydied)}{1 - Pr(Sallydied)} = e^{-4.4337 + 0.7417 \times 5} = e^{-0.7253} = 0.4842 Odds(Sallydied) = \frac{Pr(Sallydied)}{1 - Pr(Sallydied)} = e^{-4.4337 + 0.7417 \times 5} = e^{-0.7253} = 0.4842 Odds(Sallydied) = \frac{Pr(Sallydied)}{1 - Pr(Sallydied)} = e^{-4.4337 + 0.7417 \times 5} = e^{-0.7253} = 0.4842 Odds(Sallydied) = \frac{Pr(Sallydied)}{1 - Pr(Sallydied)} = e^{-0.7253} = e^{-0.$$

and so the odds ratio is:

$$OR = \frac{Odds(Harrydied)}{Odds(Sallydied)} = \frac{0.4842}{0.2307} = 2.099$$

- If the odds ratio was 1, that would mean that Harry and Sally had the same estimated odds of death, and thus the same estimated probability of death, despite having different sizes of resections.
- Since the odds ratio is greater than 1, it means that Harry has a higher estimated odds of death than Sally, and thus that Harry has a higher estimated probability of death than Sally.
- If the odds ratio was less than 1, it would mean that Harry had a lower estimated odds of death than Sally, and thus that Harry had a lower estimated probability of death than Sally.

Remember that the odds ratio is a fraction describing two positive numbers (odds can only be non-negative) so that the smallest possible odds ratio is 0.

13.3.4 Interpreting the rest of the model output from glm

res_modA

Residual Deviance: 89.49

AIC: 93.49

In addition to specifying the logistic regression coefficients, we are also presented with information on degrees of freedom, deviance (null and residual) and AIC.

- The degrees of freedom indicate the sample size.
 - Recall that we had n = 134 subjects in the data. The "Null" model includes only an intercept term (which uses 1 df) and we thus have n 1 (here 133) degrees of freedom available for estimation.
 - In our res_modA model, a logistic regression is fit including a single slope (resection) and an intercept term. Each uses up one degree of freedom to build an estimate, so we have n 2 = 134 2 = 132 residual df remaining.
- The AIC or Akaike Information Criterion (lower values are better) is also provided. This is helpful if we're comparing multiple models for the same outcome.

13.3.5 Deviance and Comparing Our Model to the Null Model

- The deviance (a measure of the model's *lack of fit*) is available for both the null model (the model with only an intercept) and for our model (res_modA) predicting our outcome, mortality.
- The deviance test, though available in R (see below) isn't really a test of whether the model works well. Instead, it assumes the model is true, and then tests to see if the coefficients are detectably different from zero. So it isn't of much practical use.
 - To compare the deviance statistics, we can subtract the residual deviance from the null deviance to describe the impact of our model on fit.
 - Null Deviance Residual Deviance can be compared to a χ^2 distribution with Null DF Residual DF degrees of freedom to obtain a global test of the in-sample predictive power of our model.
 - We can see this comparison more directly by running anova on our model:

anova(res_modA)

```
Analysis of Deviance Table

Model: binomial, link: logit

Response: died

Terms added sequentially (first to last)

Df Deviance Resid. Df Resid. Dev

NULL 133 101.943
resection 1 12.45 132 89.493
```

To complete a deviance test and obtain a p value, we can run the following code to estimate the probability that a chi-square distribution with a single degree of freedom would exhibit an improvement in deviance as large as 12.45.

```
pchisq(12.45, 1, lower.tail = FALSE)
```

[1] 0.0004179918

The p value for the deviance test here is about 0.0004. But, again, this isn't a test of whether the model is any good - it assumes the model is true, and then tests some consequences.

- Specifically, it tests whether (if the model is TRUE) some of the model's coefficients are non-zero.
- That's not so practically useful, so I discourage you from performing global tests of a logistic regression
 model with a deviance test.

13.3.6 Using glance with a logistic regression model

We can use the glance function from the broom package to obtain the null and residual deviance and degrees of freedom. Note that the deviance for our model is related to the log likelihood by -2*logLik.

```
glance(res_modA)
```

```
null.deviance df.null logLik AIC BIC deviance df.residual 101.9431 133 -44.74646 93.49292 99.2886 89.49292 132
```

The glance result also provides the AIC, and the BIC (Bayes Information Criterion), each of which is helpful in understanding comparisons between multiple models for the same outcome (with smaller values of either criterion indicating better models.) The AIC is based on the deviance, but penalizes you for making the model more complicated. The BIC does the same sort of thing but with a different penalty.

Again we see that we have a null deviance of 101.94 on 133 degrees of freedom. Including the **resection** information in the model decreased the deviance to 89.49 points on 132 degrees of freedom, so that's a decrease of 12.45 points while using one degree of freedom, a statistically significant reduction in deviance.

13.4 Interpreting the Model Summary

Let's get a more detailed summary of our res_modA model, including 95% confidence intervals for the coefficients:

```
summary(res_modA)
Call:
glm(formula = died ~ resection, family = "binomial", data = resect)
Deviance Residuals:
   Min
              1Q
                  Median
                                3Q
                                        Max
-1.1844 -0.5435 -0.3823 -0.2663
                                     2.4501
Coefficients:
            Estimate Std. Error z value Pr(>|z|)
                         0.8799
(Intercept) -4.4337
                                -5.039 4.67e-07 ***
resection
              0.7417
                         0.2230
                                  3.327 0.000879 ***
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for binomial family taken to be 1)
   Null deviance: 101.943 on 133 degrees of freedom
Residual deviance: 89.493 on 132 degrees of freedom
```

AIC: 93.493

```
Number of Fisher Scoring iterations: 5
```

```
confint(res_modA, level = 0.95)
```

Waiting for profiling to be done...

```
2.5 % 97.5 % (Intercept) -6.344472 -2.855856 resection 0.322898 1.208311
```

Some elements of this summary are very familiar from our work with linear models.

- We still have a five-number summary of residuals, although these are called *deviance* residuals.
- We have a table of coefficients with standard errors, and hypothesis tests, although these are Wald z-tests, rather than the t tests we saw in linear modeling.
- We have a summary of global fit in the comparison of null deviance and residual deviance, but without a formal p value. And we have the AIC, as discussed above.
- We also have some new items related to a *dispersion* parameter and to the number of Fisher Scoring Iterations.

Let's walk through each of these elements.

13.4.1 Wald Z tests for Coefficients in a Logistic Regression

The coefficients output provides the estimated coefficients, and their standard errors, plus a Wald Z statistic, which is just the estimated coefficient divided by its standard error. This is compared to a standard Normal distribution to obtain the two-tailed p values summarized in the Pr(>|z|) column.

- The interesting result is resection, which has a Wald Z = 3.327, yielding a p value of 0.00088.
- The hypotheses being tested here are H₀: resection does not have an effect on the log odds of died vs. H_A: resection does have such an effect.
- Another way of stating this is that the *p* value assesses whether the estimated coefficient of **resection**, 0.7417, is statistically detectably different from 0. If the coefficient (on the logit scale) for **resection** was truly 0, this would mean that:
 - the log odds of death did not change based on the resection size,
 - the odds of death were unchanged based on the resection size (the odds ratio would be 1), and
 - the probability of death was unchanged based on the resection size.

In our case, we have a statistically detectable change in the log odds of died associated with changes in resection, according to this p value. We conclude that resection size is associated with a positive impact on death rates (death rates are generally higher for people with larger resections.)

13.4.2 Confidence Intervals for the Coefficients

As in linear regression, we can obtain 95% confidence intervals (to get other levels, change the level parameter in confint) for the intercept and slope coefficients.

Here, we see, for example, that the coefficient of resection has a point estimate of 0.7417, and a confidence interval of (0.3229, 1.208). Since this is on the logit scale, it's not that interpretable, but we will often exponentiate the model and its confidence interval to obtain a more interpretable result on the odds ratio scale.

```
exp(coef(res_modA))
```

```
(Intercept) resection 0.01186995 2.09947754
```

exp(confint(res_modA))

Waiting for profiling to be done...

2.5 % 97.5 %

(Intercept) 0.001756429 0.05750655 resection 1.381124459 3.34782604

From this output, we can estimate the odds ratio for death associated with a 1 cm increase in resection size is 2.099, with a 95% CI of (1.38, 3.35). - If the odds ratio was 1, it would indicate that the odds of death did not change based on the change in resection size. - Here, it's clear that the estimated odds of death will be larger (odds > 1) for subjects with larger resection sizes. Larger odds of death also indicate larger probabilities of death. This confidence interval indicates that with 95% confidence, we conclude that increases in resection size are associated with statistically detectable increases in the odds of death. - If the odds ratio was less than 1 (remember that it cannot be less than 0) that would mean that subjects with larger resection sizes were associated with smaller estimated odds of death.

13.4.3 Deviance Residuals

In logistic regression, it's certainly a good idea to check to see how well the model fits the data. However, there are a few different types of residuals. The residuals presented here by default are called deviance residuals. Other types of residuals are available for generalized linear models, such as Pearson residuals, working residuals, and response residuals. Logistic regression model diagnostics often make use of multiple types of residuals.

The deviance residuals for each individual subject sum up to the deviance statistic for the model, and describe the contribution of each point to the model likelihood function.

The deviance residual, d_i , for the ith observation in a model predicting y_i (a binary variable), with the estimate being $\hat{\pi}_i$ is:

$$d_i = s_i \sqrt{-2[y_i log \hat{\pi}_i + (1 - y_i) log (1 - \hat{\pi}_i)]},$$

where s_i is 1 if $y_i = 1$ and $s_i = -1$ if $y_i = 0$.

Again, the sum of the deviance residuals is the deviance.

13.4.4 Dispersion Parameter

The dispersion parameter is taken to be 1 for glm fit using either the binomial or Poisson families. For other sorts of generalized linear models, the dispersion parameter will be of some importance in estimating standard errors sensibly.

13.4.5 Fisher Scoring iterations

The solution of a logistic regression model involves maximizing a likelihood function. Fisher's scoring algorithm in our res_modA needed five iterations to perform the logistic regression fit. All that this tells you is that the model converged, and didn't require a lot of time to do so.

13.5 Plotting a Simple Logistic Regression Model

Let's plot the logistic regression model res_modA for died using the extent of the resection in terms of probabilities. We can use either of two different approaches:

- we can plot the fitted values from our specific model against the original data, using the augment function from the broom package, or
- we can create a smooth function called binomial_smooth that plots a simple logistic model in an analogous way to geom_smooth(method = "lm") for a simple linear regression.

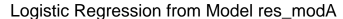
13.5.1 Using augment to capture the fitted probabilities

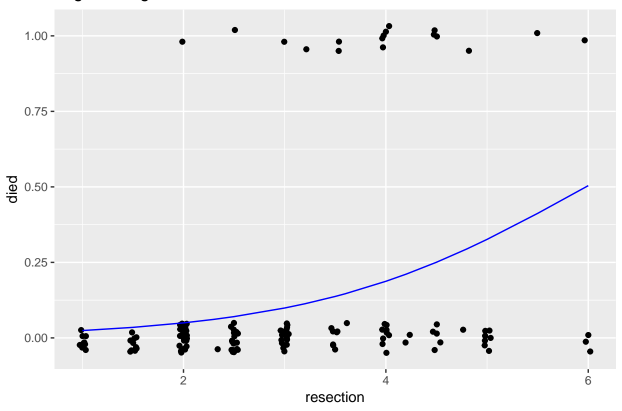
```
res_A_aug <- augment(res_modA, resect,</pre>
                     type.predict = "response")
head(res_A_aug)
  id age prior resection intubated died
                                           .fitted
                                                      .se.fit
                                                                  .resid
     34
                                0
                                     0 0.07046791 0.02562381 -0.3822929
1
  1
            1
                     2.5
2
  2
                                 0
     57
            0
                     5.0
                                     0 0.32622637 0.08605551 -0.8886631
3 3 60
            1
                     4.0
                                1
                                   1 0.18740037 0.04269795 1.8300317
4 4 62
                     4.2
                                0
                                   0 0.21104240 0.04871389 -0.6885386
            1
5
  5
     28
            0
                     6.0
                                 1
                                     1 0.50409637 0.14302982 1.1704596
                                      0 0.09897375 0.02867196 -0.4565542
  6
     52
            0
                     3.0
                                 0
         .hat
                 .sigma
                             .cooksd .std.resid
1 0.010024061 0.8258481 0.0003876961 -0.3842235
2 0.033691765 0.8227475 0.0087350915 -0.9040227
3 0.011972088 0.8107264 0.0265893468 1.8410857
4 0.014252277 0.8243062 0.0019617278 -0.6934983
```

This approach augments the resect data set with fitted, residual and other summaries of each observation's impact on the fit, using the "response" type of prediction, which yields the fitted probabilities in the .fitted column.

13.5.2 Plotting a Logistic Regression Model's Fitted Values

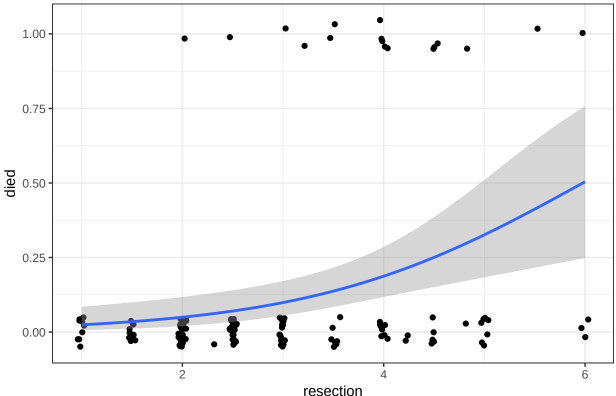
5 0.081835623 0.8196110 0.0477480056 1.2215077 6 0.009218619 0.8255581 0.0005157780 -0.4586733





13.5.3 Plotting a Simple Logistic Model using binomial_smooth





As expected, we see an increase in the model probability of death as the extent of the resection grows larger.

13.6 How well does Model A classify subjects?

A natural question to ask is how well does our model classify patients in terms of likelihood of death.

We could specify a particular rule, for example: if the predicted probability of death is 0.5 or greater, then predict "Died".

```
Predict Alive 114 16
Predict Died 3 1
```

And perhaps build the linked table of row probabilities which tells us, for example, that 87.69% of the patients predicted by the model to be alive actually did survive.

```
round(100*prop.table(
    table(res_A_aug$rule.5, res_A_aug$died), 1), 2)
```

```
0 1
Predict Alive 87.69 12.31
```

```
Predict Died 75.00 25.00
```

Or the table of column probabilities which tell us, for example, that 97.44% of those who actually survived were predicted by the model to be alive.

```
round(100*prop.table(
   table(res_A_aug$rule.5, res_A_aug$died), 2), 2)
```

```
0 1
Predict Alive 97.44 94.12
Predict Died 2.56 5.88
```

We'll discuss various measures of concordance derived from this sort of classification later.

13.7 Receiver Operating Characteristic Curve Analysis

One way to assess the predictive accuracy within the model development sample in a logistic regression is to consider an analyses based on the receiver operating characteristic (ROC) curve. ROC curves are commonly used in assessing diagnoses in medical settings, and in signal detection applications.

The accuracy of a "test" can be evaluated by considering two types of errors: false positives and false negatives.

In our res_modA model, we use resection size to predict whether the patient died. Suppose we established a value R, so that if the resection size was larger than R cm, we would predict that the patient died, and otherwise we would predict that the patient did not die.

A good outcome of our model's "test", then, would be when the resection size is larger than R for a patient who actually died. Another good outcome would be when the resection size is smaller than R for a patient who survived.

But we can make errors, too.

- A false positive error in this setting would occur when the resection size is larger than R (so we predict the patient dies) but in fact the patient does not die.
- A false negative error in this case would occur when the resection size is smaller than R (so we predict the patient survives) but in fact the patient dies.

Formally, the true positive fraction (TPF) for a specific resection cutoff R, is the probability of a positive test (a prediction that the patient will die) among the people who have the outcome died = 1 (those who actually die).

$$TPF(R) = Pr(resection > R|subject died)$$

Similarly, the false positive fraction (FPF) for a specific cutoff R is the probability of a positive test (prediction that the patient will die) among the people with died = 0 (those who don't actually die)

$$FPF(R) = Pr(resection > R|subject did not die)$$

The True Positive Rate is referred to as the sensitivity of a diagnostic test, and the True Negative rate (1 - the False Positive rate) is referred to as the specificity of a diagnostic test.

Since the cutoff R is not fixed in advanced, we can plot the value of TPF (on the y axis) against FPF (on the x axis) for all possible values of R, and this is what the ROC curve is. Others refer to the Sensitivity on the Y axis, and 1-Specificity on the X axis, and this is the same idea.

Before we get too far into the weeds, let me show you some simple situations so you can understand what you might learn from the ROC curve. The web page http://blog.yhat.com/posts/roc-curves.html provides source materials.

13.7.1 Interpreting the Area under the ROC curve

The AUC or Area under the ROC curve is the amount of space underneath the ROC curve. Often referred to as the c statistic, the AUC represents the quality of your TPR and FPR overall in a single number. The C statistic ranges from 0 to 1, with C = 0.5 for a prediction that is no better than random guessing, and C = 1 for a perfect prediction model.

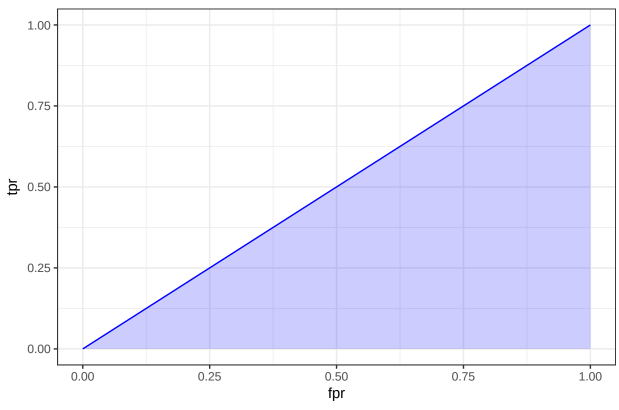
Next, I'll build a simulation to demonstrate several possible ROC curves in the sections that follow.

13.7.1.1 What if we are guessing?

If we're guessing completely at random, then the model should correctly classify a subject (as died or not died) about 50% of the time, so the TPR and FPR will be equal. This yields a diagonal line in the ROC curve, and an area under the curve (C statistic) of 0.5.

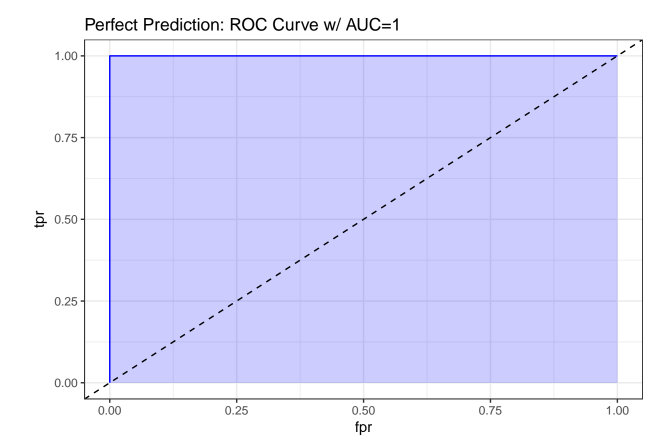
There are several ways to do this on the web, but I'll show this one, which has some bizarre code, but that's a function of using a package called ROCR to do the work. It comes from this link





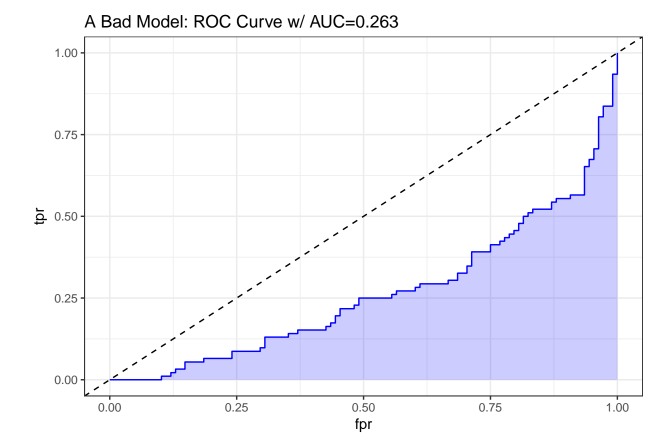
13.7.1.2 What if we classify things perfectly?

If we're classifying subjects perfectly, then we have a TPR of 1 and an FPR of 0. That yields an ROC curve that looks like the upper and left edges of a box. If our model correctly classifies a subject (as died or not died) 100% of the time, the area under the curve (c statistic) will be 1.0. We'll add in the diagonal line here (in a dashed black line) to show how this model compares to random guessing.



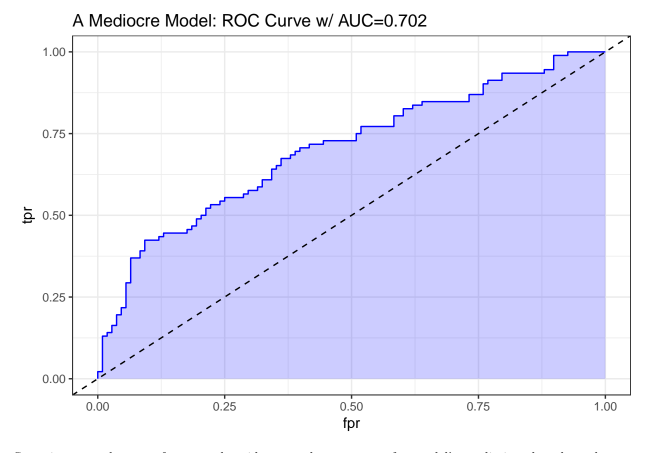
13.7.1.3 What does "worse than guessing" look like?

A bad classifier will appear below and to the right of the diagonal line we'd see if we were completely guessing. Such a model will have a c statistic below 0.5, and will be valueless.



13.7.1.4 What does "better than guessing" look like?

An "OK" classifier will appear above and to the left of the diagonal line we'd see if we were completely guessing. Such a model will have a c statistic above 0.5, and might have some value. The plot below shows a very fairly poor model, but at least it's better than guessing.



Sometimes people grasp for a rough guide as to the accuracy of a model's predictions based on the area under the ROC curve. A common thought is to assess the C statistic much like you would a class grade.

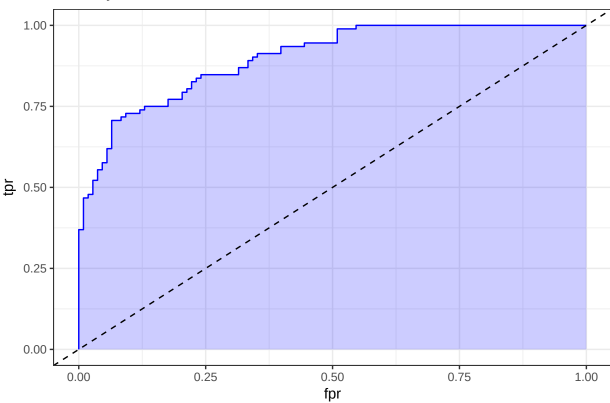
C statistic	Interpretation
0.90 to 1.00	model does an excellent job at discriminating "yes" from "no"
	(A)
0.80 to 0.90	model does a good job (B)
0.70 to 0.80	model does a fair job (C)
0.60 to 0.70	model does a poor job (D)
0.50 to 0.60	model fails (F)
below 0.50	model is worse than random guessing

13.7.1.5 What does "pretty good" look like?

A strong and good classifier will appear above and to the left of the diagonal line we'd see if we were completely guessing, often with a nice curve that is continually increasing and appears to be pulled up towards the top left. Such a model will have a c statistic well above 0.5, but not as large as 1. The plot below shows a stronger model, which appears substantially better than guessing.

```
pred_good <- prediction(sim.temp$p_good, sim.temp$y)
perf_good <- performance(pred_good, measure = "tpr", x.measure = "fpr")
auc_good <- performance(pred_good, measure="auc")
auc_good <- round(auc_good@y.values[[1]],3)
roc_good <- data.frame(fpr=unlist(perf_good@x.values),</pre>
```

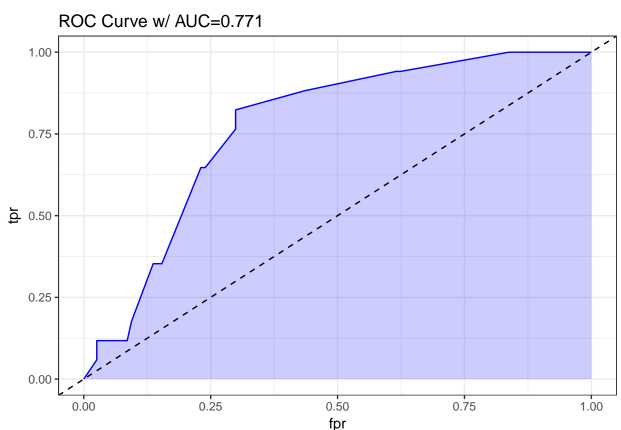
A Pretty Good Model: ROC Curve w/ AUC=0.899



13.8 The ROC Plot for res_modA

Let me show you the ROC curve for our res_modA model.

```
ggplot(roc.data, aes(x=fpr, ymin=0, ymax=tpr)) +
    geom_ribbon(alpha=0.2, fill = "blue") +
    geom_line(aes(y=tpr), col = "blue") +
    geom_abline(intercept = 0, slope = 1, lty = "dashed") +
    labs(title = paste0("ROC Curve w/ AUC=", auc)) +
    theme_bw()
```



Based on the C statistic (AUC = 0.771) this would rank somewhere near the high end of a "fair" predictive model by this standard, not quite to the level of a "good" model.

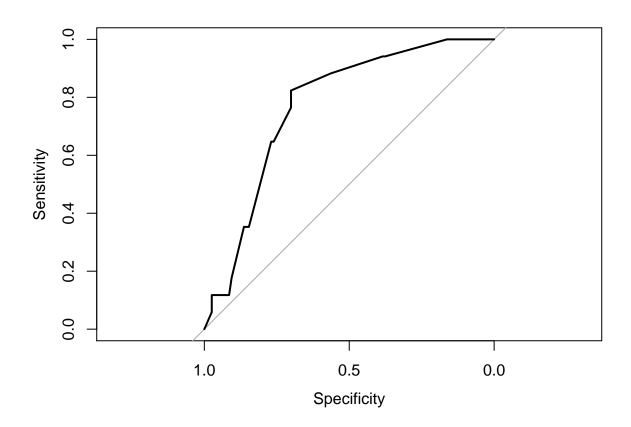
13.8.1 Another way to plot the ROC Curve

If we've loaded the pROC package, we can also use the following (admittedly simpler) approach to plot the ROC curve, without ggplot2, and to obtain the C statistic, and a 95% confidence interval around that C statistic.

Data: predict(res_modA, type = "response") in 117 controls (resect\$died 0) < 17 cases (resect\$died 1).

Area under the curve: 0.7707 95% CI: 0.67-0.8715 (DeLong)

plot(roc.modA)



13.9 Assessing Residual Plots from Model A

Residuals are certainly less informative for logistic regression than they are for linear regression: not only do yes/no outcomes inherently contain less information than continuous ones, but the fact that the adjusted response depends on the fit hampers our ability to use residuals as external checks on the model.

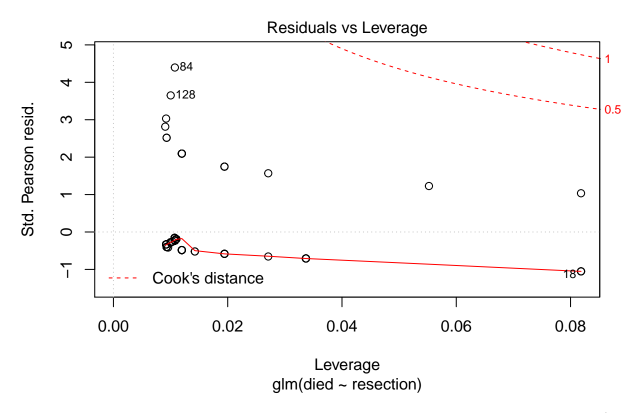
This is mitigated to some extent, however, by the fact that we are also making fewer distributional assumptions in logistic regression, so there is no need to inspect residuals for, say, skewness or heteroskedasticity.

• Patrick Breheny, University of Kentucky, Slides on GLM Residuals and Diagnostics

The usual residual plots are available in R for a logistic regression model, but most of them are irrelevant in the logistic regression setting. The residuals shouldn't follow a standard Normal distribution, and they will not show constant variance over the range of the predictor variables, so plots looking into those issues aren't helpful.

The only plot from the standard set that we'll look at in many settings is plot 5, which helps us assess influence (via Cook's distance contours), and a measure related to leverage (how unusual an observation is in terms of the predictors) and standardized Pearson residuals.

plot(res_modA, which = 5)



In this case, I don't see any highly influential points, as no points fall outside of the Cook's distance (0.5 or 1) contours.

13.10 Model B: A "Kitchen Sink" Logistic Regression Model

Call: glm(formula = died ~ resection + age + prior + intubated, family = binomial,
 data = resect)

Coefficients:

(Intercept) resection age prior intubated -5.152886 0.612211 0.001173 0.814691 2.810797

Degrees of Freedom: 133 Total (i.e. Null); 129 Residual

Null Deviance: 101.9

Residual Deviance: 67.36 AIC: 77.36

13.10.1 Comparing Model A to Model B

```
anova(res_modA, res_modB)
Analysis of Deviance Table
Model 1: died ~ resection
Model 2: died ~ resection + age + prior + intubated
 Resid. Df Resid. Dev Df Deviance
               89.493
1
       132
       129
               67.359 3
2
                          22.134
The addition of age, prior and intubated reduces the lack of fit by 22.134 points, at a cost of 3 degrees of
freedom.
glance(res_modA)
 null.deviance df.null
                                     AIC
                                             BIC deviance df.residual
                          logLik
      101.9431
                   133 -44.74646 93.49292 99.2886 89.49292
                                                                 132
glance(res_modB)
 null.deviance df.null
                        logLik
                                   AIC
                                           BIC deviance df.residual
                   133 -33.6793 77.3586 91.8478 67.3586
By either AIC or BIC, the larger model (res_modB) looks more effective.
13.10.2 Interpreting Model B
summary(res_modB)
glm(formula = died ~ resection + age + prior + intubated, family = binomial,
   data = resect)
Deviance Residuals:
             1Q
                 Median
                              3Q
                                      Max
-1.7831 -0.3741 -0.2386 -0.2014
                                   2.5228
Coefficients:
            Estimate Std. Error z value Pr(>|z|)
(Intercept) -5.152886    1.469453    -3.507    0.000454 ***
resection
          age
           0.814691
                      0.704785 1.156 0.247705
prior
intubated 2.810797
                      0.658395 4.269 1.96e-05 ***
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for binomial family taken to be 1)
   Null deviance: 101.943 on 133 degrees of freedom
Residual deviance: 67.359 on 129 degrees of freedom
AIC: 77.359
```

```
Number of Fisher Scoring iterations: 6
```

It appears that the intubated predictor adds significant value to the model, by the Wald test.

Let's focus on the impact of these variables through odds ratios.

```
2.5 % 97.5 % (Intercept) 0.0002408626 0.0837263 resection 1.0804548590 3.3495636 age 0.9618416869 1.0442885 prior 0.5485116610 9.1679931 intubated 4.7473282453 64.6456919
```

At a 5% significance level, we might conclude that:

- larger sized resections are associated with a meaningful rise (est OR: 1.84, 95% CI 1.08, 3.35) in the odds of death, holding all other predictors constant,
- the need for intubation at the end of surgery is associated with a substantial rise (est OR: 16.6, 95% CI 4.7, 64.7) in the odds of death, holding all other predictors constant, but that
- older age as well as having a prior tracheal surgery appears to be associated with an increase in death risk, but not to an extent that we can declare statistically significant.

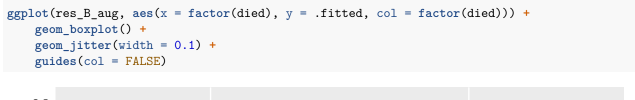
13.11 Plotting Model B

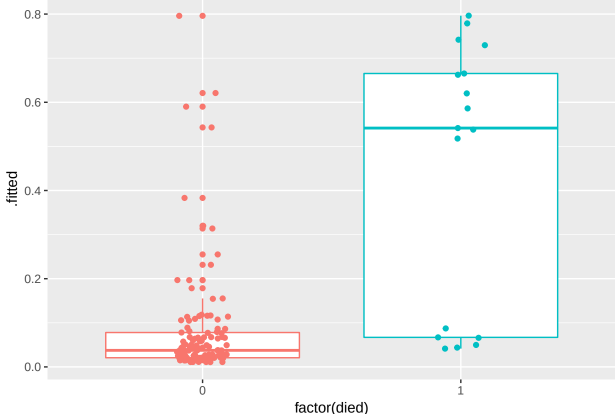
Let's think about plotting the fitted values from our model, in terms of probabilities.

13.11.1 Using augment to capture the fitted probabilities

```
res_B_aug <- augment(res_modB, resect,</pre>
                     type.predict = "response")
head(res_B_aug)
  id age prior resection intubated died
                                             .fitted
                                                        .se.fit
                                                                    .resid
  1
      34
             1
                     2.5
                                  0
                                       0 0.05908963 0.03851118 -0.3490198
1
  2
2
      57
             0
                     5.0
                                  0
                                       0 0.11660492 0.06253774 -0.4979613
3
  3
      60
                     4.0
                                       1 0.72944600 0.15010423 0.7943172
             1
                                  1
4
  4
     62
                     4.2
                                       0 0.15522494 0.09607978 -0.5808354
5
  5
      28
             0
                     6.0
                                  1
                                       1 0.79641141 0.14588554 0.6747435
6
  6
      52
             0
                     3.0
                                  0
                                       0 0.03713809 0.01933270 -0.2751191
                             .cooksd .std.resid
        .hat
                 .sigma
1 0.02667562 0.7247491 0.0003536652 -0.3537702
2 0.03796756 0.7240341 0.0010829917 -0.5076925
3 0.11416656 0.7215778 0.0107925872
4 0.07039819 0.7234665 0.0029937671 -0.6024273
5 0.13126049 0.7225958 0.0088920280 0.7239256
6 0.01045207 0.7250114 0.0000823406 -0.2765683
```

13.11.2 Plotting Model B Fits by Observed Mortality



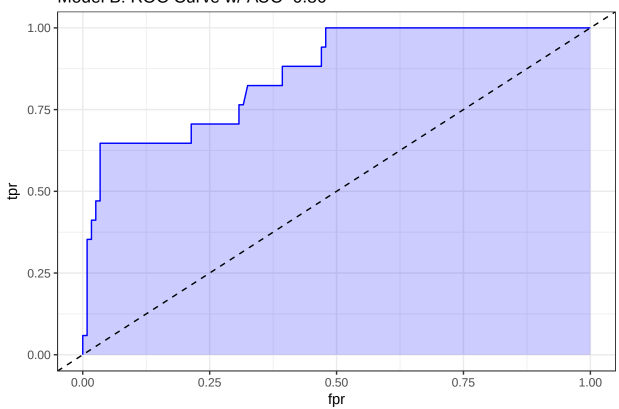


Certainly it appears as though most of our predicted probabilities (of death) for the subjects who actually survived are quite small, but not all of them. We also have at least 6 big "misses" among the 17 subjects who actually died.

13.11.3 The ROC curve for Model B

```
geom_ribbon(alpha=0.2, fill = "blue") +
geom_line(aes(y=tpr), col = "blue") +
geom_abline(intercept = 0, slope = 1, lty = "dashed") +
labs(title = paste0("Model B: ROC Curve w/ AUC=", auc)) +
theme_bw()
```

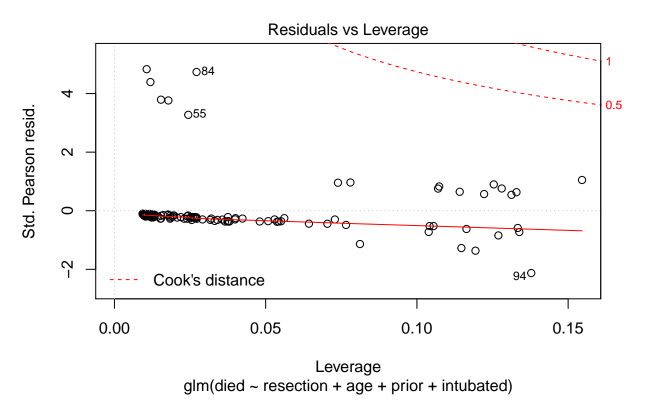




The area under the curve (C-statistic) is 0.86, which certainly looks like a more discriminating fit than model A with resection alone.

13.11.4 Residuals, Leverage and Influence

```
plot(res_modB, which = 5)
```



Again, we see no signs of deeply influential points in this model.

13.12 Logistic Regression using 1rm

To obtain the Nagelkerke R^2 and the C statistic, as well as some other summaries, I'll now demonstrate the use of lrm from the rms package to fit a logistic regression model.

We'll return to the original model, predicting death using resection size alone.

```
dd <- datadist(resect)
options(datadist="dd")

res_modC <- lrm(died ~ resection, data=resect, x=TRUE, y=TRUE)
res_modC</pre>
```

Logistic Regression Model

```
lrm(formula = died ~ resection, data = resect, x = TRUE, y = TRUE)
```

		Model Likelihood		Discrimination		Rank Discrim.	
		Ratio Test		Indexes		Indexes	
0bs	134	LR chi2	12.45	R2	0.167	С	0.771
0	117	d.f.	1	g	1.037	Dxy	0.541
1	17	Pr(> chi2)	0.0004	gr	2.820	gamma	0.582
max deri	v 2e-06			gp	0.110	tau-a	0.121
				Brier	0.103		

```
Coef S.E. Wald Z Pr(>|Z|)
Intercept -4.4337 0.8799 -5.04 <0.0001
resection 0.7417 0.2230 3.33 0.0009
```

This output specifies the following:

- Obs = The number of observations used to fit the model, with 0 = the number of zeros and 1 = the number of ones in our outcome, died. Also specified is the maximum absolute value of the derivative at the point where the maximum likelihood function was estimated. I wouldn't worry about that practically, as all you will care about is whether the iterative function-fitting process converged, and R will warn you in other ways if it doesn't.
- A likelihood ratio test (drop in deviance test) subtracting the residual deviance from the null deviance obtain the Likelihood Ratio χ^2 statistic, subtracting residual df from null df to obtain degrees of freedom, and comparing the resulting test statistic to a χ^2 distribution with the appropriate degrees of freedom to determine a p value.
- A series of discrimination indexes, including the Nagelkerke R², symbolized R2, and several others we'll discuss shortly.
- A series of rank discrimination indexes, including the C statistic (area under the ROC curve) and Somers' D (Dxy), and several others.
- A table of coefficients, standard errors, Wald Z statistics and p values based on those Wald statistics.

The C statistic is estimated to be 0.771, with an associated (Nagelkerke) $R^2 = 0.167$, both indicating at best mediocre performance for this model, as it turns out.

13.12.1 Interpreting Nagelkerke R²

There are many ways to calculate R^2 for logistic regression.

- At the unfortunate URL linked here (unfortunate because the term "pseudo" is misspelled) there is a nice summary of the key issue, which is that there are at least three different ways to think about R^2 in linear regression that are equivalent in that context, but when you move to a categorical outcome, which interpretation you use leads you down a different path for extension to the new type of outcome.
- Paul Allison, for instance, describes several at this link in a post entitled "What's the Best R-Squared for Logistic Regression?"
- Jonathan Bartlett looks at McFadden's pseudo R² in some detail (including some R code) at this link, in a post entitled "R squared in logistic regression"

The Nagelkerke approach that is presented as R2 in the 1rm output is as good as most of the available approaches, and has the positive feature that it does reach 1 if the fitted model shows as much improvement as possible over the null model (which predicts the mean response for all subjects, and has $R^2 = 0$). The greater the improvement, the higher the Nagelkerke R^2 .

For model A, our Nagelkerke $R^2 = 0.167$, which is pretty poor. It doesn't technically mean that 16.7% of any sort of variation has been explained, though.

13.12.2 Interpreting the C statistic and Plotting the ROC Curve

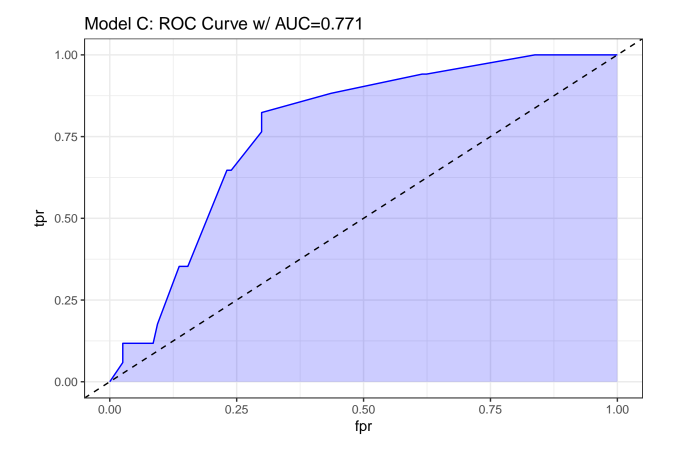
The C statistic is a measure of the area under the receiver operating characteristic curve. This link has some nice material that provides some insight into the C statistic and ROC curve.

- Recall that C ranges from 0 to 1. 0 = BAD, 1 = GOOD.
 - values of C less than 0.5 indicate that your prediction model is not even as good as simple random guessing of "yes" or "no" for your response.
 - C = 0.5 for random guessing

- -C = 1 indicates a perfect classification scheme one that correctly guesses "yes" for all "yes" patients, and for none of the "no" patients.
- The closer C is to 1, the happier we'll be, most of the time.
 - Often we'll call models with 0.5 < C < 0.8 poor or weak in terms of predictive ability by this measure
 - $-0.8 \le C < 0.9$ are moderately strong in terms of predictive power (indicate good discrimination)
 - $-C \ge 0.9$ usually indicates a very strong model in this regard (indicate excellent discrimination)

We've seen the ROC curve for this model before, when we looked at model res_modA fitted using glm in the previous chapter. But, just for completeness, I'll include it.

Note. I change the initial predict call from type = "response" for a glm fit to type = "fitted" in a lrm fit. Otherwise, this is the same approach.



13.12.3 The C statistic and Somers' D

- The C statistic is directly related to **Somers' D statistic**, abbreviated D_{xy} , by the equation C = 0.5 + (D/2).
 - Somers' D and the ROC area only measure how well predicted values from the model can rankorder the responses. For example, predicted probabilities of 0.01 and 0.99 for a pair of subjects are no better than probabilities of 0.2 and 0.8 using rank measures, if the first subject had a lower response value than the second.
 - Thus, the C statistic (or D_{xy}) may not be very sensitive ways to choose between models, even though they provide reasonable summaries of the models individually.
 - This is especially true when the models are strong. The Nagelkerke R² may be more sensitive.
- But as it turns out, we sometimes have to look at the ROC shapes, as the summary statistic alone isn't enough.

In our case, Somers D (Dxy) = .541, so the C statistic is 0.771.

13.12.4 Validating the Logistic Regression Model Summary Statistics

Like other regression-fitting tools in rms, the lrm function has a special validate tool to help perform resampling validation of a model, with or without backwards step-wise variable selection. Here, we'll validate our model's summary statistics using 100 bootstrap replications.

```
set.seed(432001)
validate(res_modC, B = 100)
```

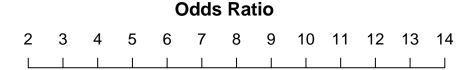
Dxy	0.5415	0.5325	0.5415	-0.0090	0.5505	100
R2	0.1666	0.1664	0.1666	-0.0002	0.1668	100
Intercept	0.0000	0.0000	0.1425	-0.1425	0.1425	100
Slope	1.0000	1.0000	1.0742	-0.0742	1.0742	100
Emax	0.0000	0.0000	0.0416	0.0416	0.0416	100
D	0.0854	0.0872	0.0854	0.0017	0.0837	100
U	-0.0149	-0.0149	-0.0004	-0.0145	-0.0004	100
Q	0.1004	0.1021	0.0859	0.0162	0.0841	100
В	0.1025	0.1032	0.1046	-0.0014	0.1039	100
g	1.0369	1.0247	1.0369	-0.0122	1.0491	100
gp	0.1101	0.1082	0.1101	-0.0019	0.1119	100

Recall that our area under the curve (C statistic) = 0.5 + (Dxy/2), so that we can also use the first row of statistics to validate the C statistic. Accounting for optimism in this manner, our corrected estimates are Dxy = 0.551, so C = 0.776, and Nagelkerke $R^2 = 0.167$.

13.12.5 Plotting the Summary of the 1rm approach

The summary function applied to an 1rm fit shows the effect size comparing the 25th to the 75th percentile of resection.

plot(summary(res_modC))



resection – 4:2

```
summary(res_modC)
```

Effects Response : died

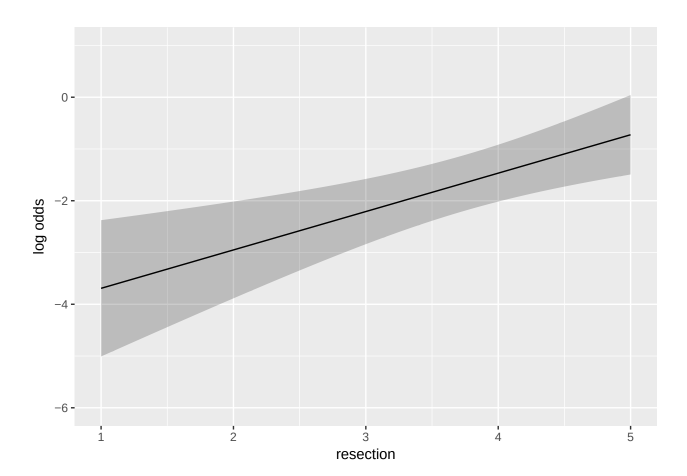
Factor Low High Diff. Effect S.E. Lower 0.95 Upper 0.95 resection 2 4 2 1.4834 0.44591 0.6094 2.3574 Odds Ratio 2 4 2 4.4078 NA 1.8393 10.5630

So, a move from a resection of 2 cm to a resection of 4 cm is associated with an estimated effect on the log odds of death of 1.48 (with standard error 0.45), or with an estimated effect on the odds ratio for death of 4.41, with 95% CI (1.84, 10.56).

13.12.6 Plot In-Sample Predictions for Model C

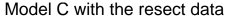
Here we plot the effect of resection (and 95% confidence intervals) across the range of observed values of resection on the log odds of death. Note the linear effect of resection size on the log odds scale.

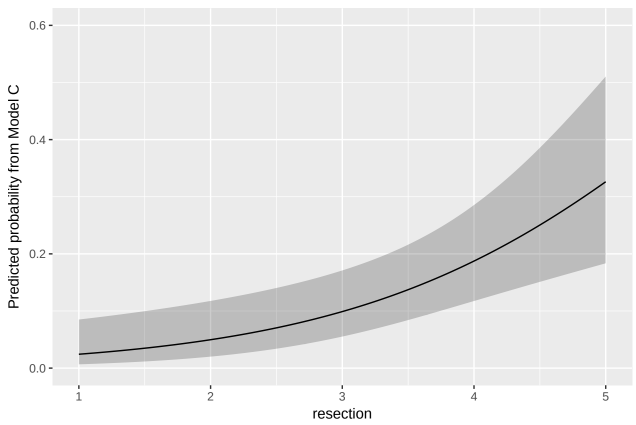
```
ggplot(Predict(res_modC))
```



By applying the plogis function within the Predict command, we can plot the effect of resection on the estimated probability of death. Note the non-linear effect on this probability in this logistic regression model.

```
ggplot(Predict(res_modC, fun = plogis)) +
   labs(y = "Predicted probability from Model C",
        title = "Model C with the resect data")
```





The Predict function itself provides the raw material being captured in this plot.

```
head(Predict(res_modC, fun = plogis))
```

```
resection
                           yhat
                                       lower
                                                 upper .predictor.
resection.1 1.000000 0.02431476 0.006636502 0.08505223
                                                         resection
resection.2 1.020101 0.02467096 0.006789313 0.08559056
                                                          resection
resection.3 1.040201 0.02503224 0.006945549 0.08613277
                                                          resection
resection.4 1.060302 0.02539867 0.007105283 0.08667889
                                                         resection
resection.5 1.080402 0.02577033 0.007268589 0.08722896
                                                         resection
resection.6 1.100503 0.02614728 0.007435542 0.08778304
                                                         resection
```

Response variable (y):

Limits are 0.95 confidence limits

13.12.7 ANOVA from the 1rm approach

```
anova(res_modC)
```

Wald Statistics Response: died

Factor Chi-Square d.f. P resection 11.07 1 9e-04 TOTAL 11.07 1 9e-04

The ANOVA approach applied to a 1rm fit provides a Wald test for the model as a whole. Here, the use of

resection is a significant improvement over a null (intercept-only) model. The p value is 9×10^{-4} .

13.12.8 Are any points particularly influential?

I'll use a cutoff for dfbeta here of 0.3, instead of the default 0.2, because I want to focus on truly influential points. Note that we have to use the data frame version of resect as show.influence isn't tibble-friendly.

```
inf.C <- which.influence(res_modC, cutoff=0.3)
inf.C</pre>
```

```
$Intercept
[1] "84" "128"

$resection
[1] "84"

show.influence(object = inf.C, dframe = data.frame(resect))
```

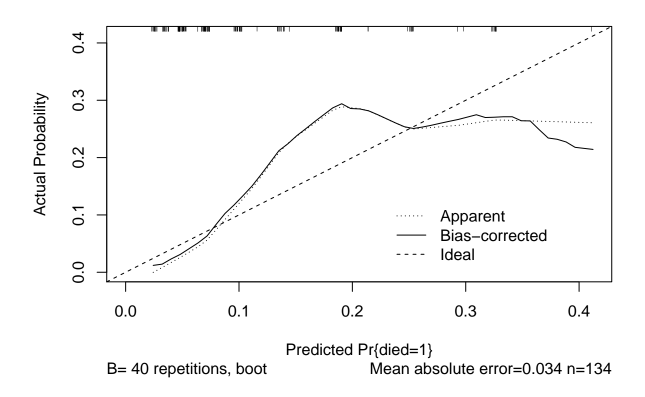
```
Count resection
84 2 *2.0
128 1 2.5
```

It appears that observation 84 may have a meaningful effect on both the intercept and the coefficient for resection.

13.12.9 A plot of the model's calibration curve

The calibrate function applied to a 1rm fit provides an assessment of the impact of overfitting on our model. The function uses bootstrapping (or cross-validation) to get bias-corrected estimates of predicted vs. observed values based on nonparametric smoothers for logistic regressions. In order to obtain this curve, you need to set both x = TRUE and y = TRUE when fitting the model. The errors here refer to the difference between the model predicted values and the corresponding bias-corrected calibrated values.

```
plot(calibrate(res_modC))
```

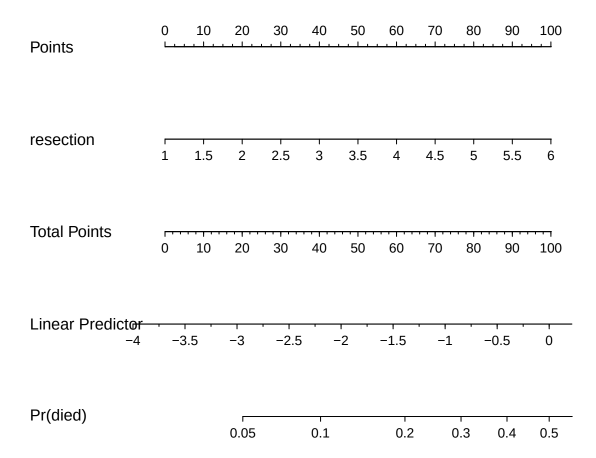


n=134 Mean absolute error=0.034 Mean squared error=0.00239 0.9 Quantile of absolute error=0.104

Here, we see that this model deviates fairly substantially from the "ideal" model, especially as the predicted probabilities increase over 0.1.

13.12.10 A Nomogram for Model C

We use the plogis function within a nomogram call to get R to produce fitted probabilities (of our outcome, died) in this case.



Since there's no non-linearity in the right hand side of our simple logistic regression model, the nomogram is straightforward. We calculate the points based on the resection by traveling up, and then travel down in a straight vertical line from total points through the linear (log odds) predictor straight to a fitted probability. Note that fitted probabilities above 0.5 are not possible within the range of observed resection values in this case.

13.13 Model D: An Augmented Kitchen Sink Model

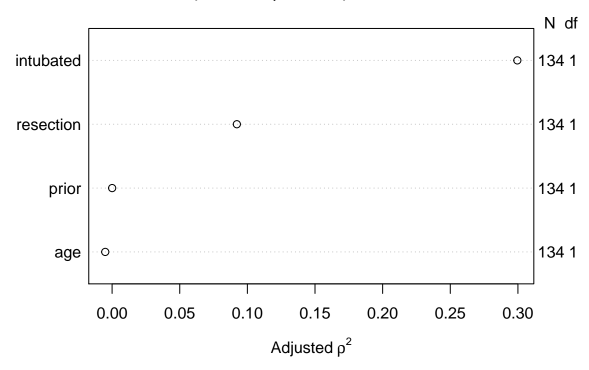
Can we predict survival from the patient's age, whether the patient had prior tracheal surgery or not, the extent of the resection, and whether intubation was required at the end of surgery?

13.13.1 Spearman ρ^2 Plot

Let's start by considering the limited use of non-linear terms for predictors that look important in a Spearman ρ^2 plot.

```
plot(spearman2(died ~ age + prior + resection + intubated, data=resect))
```





The most important variable appears to be whether intubation was required, so I'll include intubated's interaction with the linear effect of the next most (apparently) important variable, resection, and also a cubic spline for resection, with three knots. Since prior and age look less important, I'll simply add them as linear terms.

13.13.2 Fitting Model D using 1rm

Note the use of %ia% here. This insures that only the linear part of the resection term will be used in the interaction with intubated.

13.13.3 Assessing Model D using 1rm's tools

```
res_modD
```

Logistic Regression Model

lrm(formula = died ~ age + prior + rcs(resection, 3) + intubated +
 intubated %ia% resection, data = resect, x = TRUE, y = TRUE)

		Model Likelihood		Discri	Discrimination		Rank Discrim.	
		Ratio	Test	Ind	exes	Indexes		
Obs	134	LR chi2	38.08	R2	0.464	C	0.880	
0	117	d.f.	6	g	2.382	Dxy	0.759	
1	17	Pr(> chi2)	<0.0001	gr	10.825	gamma	0.770	
max deriv	9e-08			gp	0.172	tau-a	0.169	
				Brier	0.067			
		Coef	S.E. Wald	l Z Pr(>	Z)			
Intercept		-11.3636	4.9099 -2.3	0.02	06			
age		0.0000	0.0210 0.0	0.99	88			
prior		0.6269	0.7367 0.8	0.39	47			
resection		3.3799	1.9700 1.7	2 0.08	62			
resection'		-4.2104	2.7035 -1.5	6 0.11	94			
intubated		0.4576	2.7848 0.1	6 0.86	95			
intubated * :	resection	0.6188	0.7306 0.8	5 0.39	70			

- The model likelihood ratio test suggests that at least some of these predictors are helpful.
- The Nagelkerke R² of 0.46, and the C statistic of 0.88 indicate a meaningful improvement in discrimination over our model with resection alone.
- The Wald Z tests see some potential need to prune the model, as none of the elements reaches statistical significance without the others. The product term between intubated and resection, in particular, doesn't appear to have helped much, once we already had the main effects.

13.13.4 ANOVA and Wald Tests for Model D

0.3970 0.1194 0.0003 0.3970 0.3970

anova(res_modD)

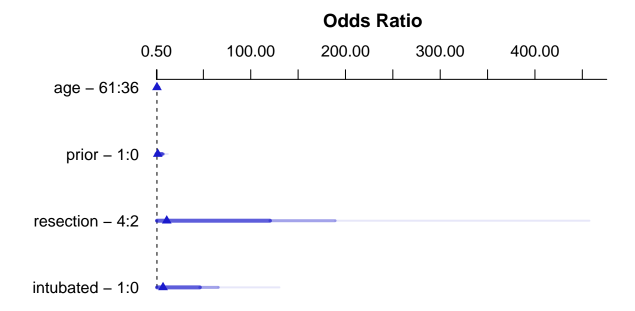
Wald Statistics Response: died Factor Chi-Square d.f. age 0.00 0.72 1 prior resection (Factor+Higher Order Factors) 4.95 All Interactions 0.72 1 Nonlinear 2.43 1 intubated (Factor+Higher Order Factors) 16.45 2 All Interactions 0.72 1 intubated * resection (Factor+Higher Order Factors) 0.72 1 TOTAL NONLINEAR + INTERACTION 2.56 2 TOTAL 23.90 0.9988 0.3947 0.1753

0.2783 0.0005

Neither the interaction term nor the non-linearity from the cubic spline appears to be statistically significant, based on the Wald tests via ANOVA. However it is clear that intubated has a significant impact as a main effect.

13.13.5 Effect Sizes in Model D

```
plot(summary(res_modD))
```



Adjusted to:resection=2.5 intubated=0

Effects	Response	:	died

summary(res_modD)

Factor	Low	High	Diff.	Effect	S.E.	Lower 0.95	Upper 0.95
age	36	61	25	-0.00080933	0.52409	-1.02800	1.0264
Odds Ratio	36	61	25	0.99919000	NA	0.35772	2.7910
prior	0	1	1	0.62693000	0.73665	-0.81688	2.0707
Odds Ratio	0	1	1	1.87190000	NA	0.44181	7.9307
resection	2	4	2	2.42930000	1.43510	-0.38342	5.2419
Odds Ratio	2	4	2	11.35000000	NA	0.68153	189.0400
intubated	0	1	1	2.00470000	1.11220	-0.17513	4.1845
Odds Ratio	0	1	1	7.42380000	NA	0.83934	65.6610

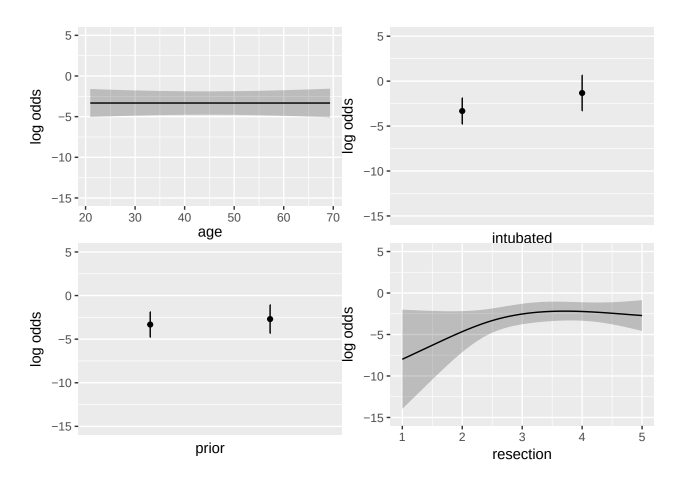
Adjusted to: resection=2.5 intubated=0

The effect sizes are perhaps best described in terms of odds ratios. The odds ratio for death isn't significantly different from 1 for any variable, but the impact of resection and intubated, though not strong enough to be significant, look more substantial (if poorly estimated) than the effects of age and prior.

13.13.6 Plot In-Sample Predictions for Model D

Here are plots of the effects across the range of each predictor (holding the others constant) on the log odds scale. Note the non-linear effect of resection implied by the use of a spline there.

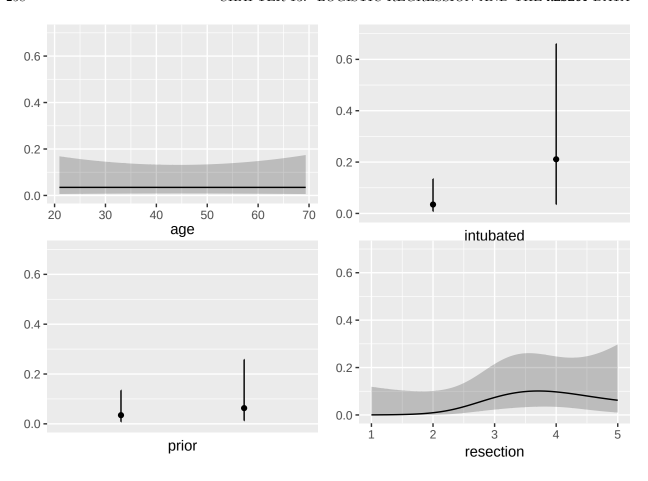
ggplot(Predict(res_modD))



We can also capture and plot these results on the probability scale, as follows¹.

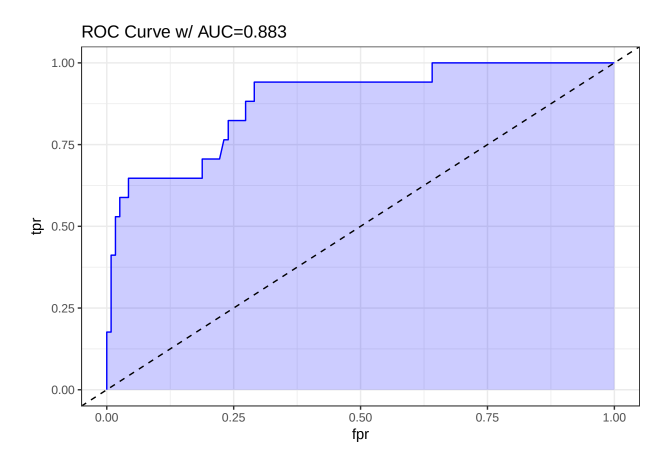
ggplot(Predict(res_modD, fun = plogis))

¹Although I've yet to figure out how to get the y axis relabeled properly without simply dumping the Predict results into a new tibble and starting over with creating the plots.



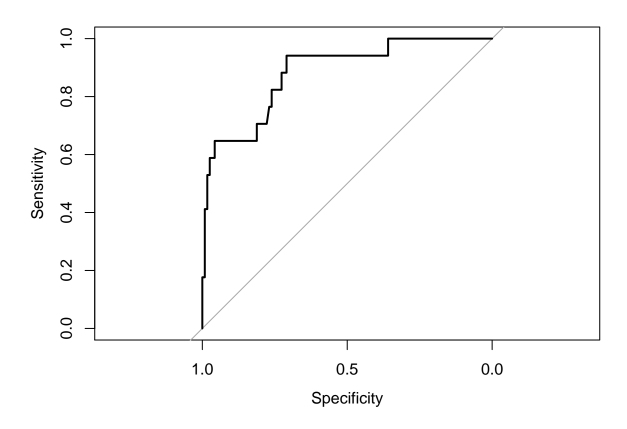
13.13.7 Plotting the ROC curve for Model D

Again, remember to use type = "fitted" with a lrm fit.



The AUC fitted with ROCR (0.883) is slightly different than what 1rm has told us (0.880), and this also happens if we use the pROC approach, demonstrated below.

```
## requires pROC package
roc.modD <-
    roc(resect$died ~ predict(res_modD, type="fitted"),
        ci = TRUE)
roc.modD</pre>
```



13.13.8 Validation of Model D summaries

```
set.seed(432002)
validate(res_modD, B = 100)
```

Divergence or singularity in 6 samples

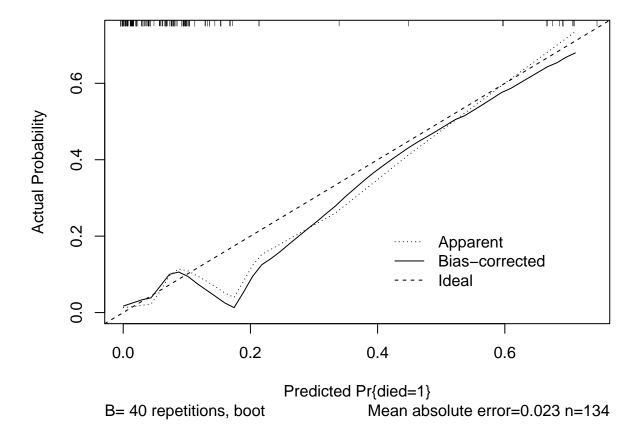
	<pre>index.orig</pre>	training	test	${\tt optimism}$	${\tt index.corrected}$	n
Dxy	0.7652	0.8162	0.7142	0.1020	0.6632	94
R2	0.4643	0.5382	0.3991	0.1391	0.3252	94
Intercept	0.0000	0.0000	-0.3919	0.3919	-0.3919	94
Slope	1.0000	1.0000	0.7201	0.2799	0.7201	94
Emax	0.0000	0.0000	0.1530	0.1530	0.1530	94
D	0.2767	0.3258	0.2322	0.0936	0.1831	94
U	-0.0149	-0.0149	0.1998	-0.2147	0.1998	94
Q	0.2916	0.3407	0.0324	0.3083	-0.0167	94
В	0.0673	0.0595	0.0739	-0.0144	0.0817	94
g	2.3819	4.2214	2.2024	2.0190	0.3629	94
gp	0.1720	0.1777	0.1591	0.0186	0.1534	94

The C statistic indicates fairly strong discrimination, at C=0.88, although after validation, this looks much weaker (based on Dxy = 0.6632, we would have C=0.5+0.6632/2=0.83) and the Nagelkerke R^2 is also reasonably good, at 0.46, although this, too, is overly optimistic, and we bias-correct through our validation study to 0.33.

13.13.9 Calibration Plot for Model D

plot(calibrate(res_modD))

Divergence or singularity in 5 samples



n=134 Mean absolute error=0.023 Mean squared error=0.00151 0.9 Quantile of absolute error=0.062

This larger model is perhaps a bit better calibrated than the simple model on resection alone, but there's still a lot of variation from the ideal (diagonal line.)

13.14 Model E: Fitting a Reduced Model in light of Model D

Can you suggest a reduced model (using a subset of the independent variables in model D) that adequately predicts survival?

Based on the anova for model D and the Spearman rho-squared plot, it appears that a two-predictor model using intubation and resection may be sufficient. Neither of the other potential predictors shows a statistically detectable effect in its Wald test.

Logistic Regression Model

```
lrm(formula = died \sim intubated + resection, data = resect, x = TRUE, y = TRUE)
```

		Model Lik	elihood	Discrin	Discrimination		Rank Discrim.	
		Ratio	Test	Inde	Indexes		Indexes	
0bs	134	LR chi2	33.27	R2	0.413	С	0.867	
0	117	d.f.	2	g	1.397	Dxy	0.734	
1	17	Pr(> chi2)	<0.0001	gr	4.043	gamma	0.757	
max	deriv 5e-10			gp	0.160	tau-a	0.164	
				Brier	0.073			

```
Coef S.E. Wald Z Pr(>|Z|)
Intercept -4.6370 1.0430 -4.45 <0.0001
intubated 2.8640 0.6479 4.42 <0.0001
resection 0.5475 0.2689 2.04 0.0418
```

The model equation is that the log odds of death is -4.637 + 2.864 intubated +0.548 resection.

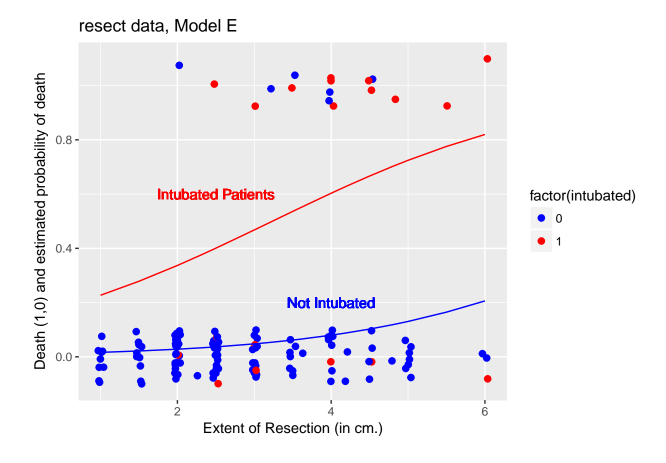
This implies that:

- for intubated patients, the equation is -1.773 + 0.548 resection, while
- for non-intubated patients, the equation is -4.637 + 0.548 resection.

We can use the ilogit function within the faraway package to help plot this.

13.14.1 A Plot comparing the two intubation groups

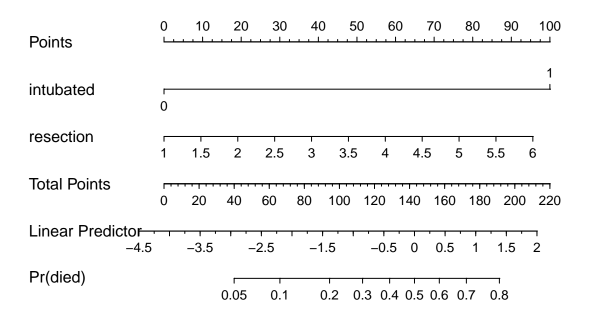
```
ggplot(resect, aes(x = resection, y = died,
                   col = factor(intubated))) +
    scale_color_manual(values = c("blue", "red")) +
   geom_jitter(size = 2, height = 0.1) +
    geom_line(aes(x = resection,
                  y = faraway::ilogit(-4.637 + 0.548*resection)),
              col = "blue") +
    geom_line(aes(x = resection,
                  y = faraway::ilogit(-1.773 + 0.548*resection)),
              col = "red") +
    geom_text(x = 4, y = 0.2, label = "Not Intubated",
              col="blue") +
    geom_text(x = 2.5, y = 0.6, label = "Intubated Patients",
             col="red") +
    labs(x = "Extent of Resection (in cm.)",
         y = "Death (1,0) and estimated probability of death",
         title = "resect data, Model E")
```



The effect of intubation appears to be very large, compared to the resection size effect.

13.14.2 Nomogram for Model E

A nomogram of the model would help, too.



Again, we see that the effect of intubation is enormous, compared to the effect of resection. Another way to see this is to plot the effect sizes directly.

13.14.3 Effect Sizes from Model E

plot(summary(res_modE))



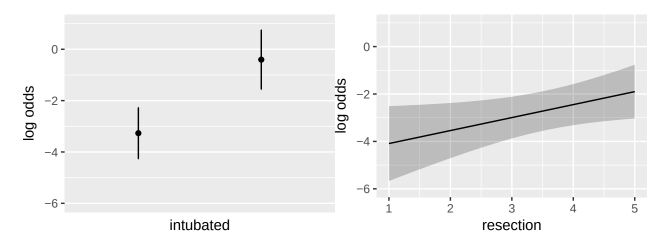
summary(res_modE)

	Effe	ects		Response : died					
Factor	Low	High	Diff.	Effect	S.E.	Lower 0.95	Upper 0.95		
intubated	0	1	1	2.8640	0.64790	1.59410	4.1338		
Odds Ratio	0	1	1	17.5310	NA	4.92390	62.4160		
resection	2	4	2	1.0949	0.53783	0.04082	2.1491		
Odds Ratio	2	4	2	2.9890	NA	1.04170	8.5769		

13.14.4 Plot In-Sample Predictions for Model E

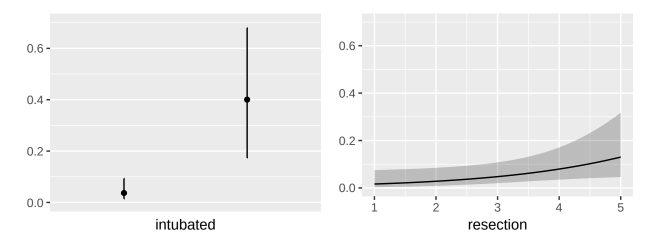
Here are plots of the effects across the range of each predictor (holding the other constant) on the log odds scale.

ggplot(Predict(res_modE))



We can also capture and plot these results on the probability scale, as follows.

```
ggplot(Predict(res_modE, fun = plogis))
```



13.14.5 ANOVA for Model E

anova(res_modE)

Wald	Statistics	Response:	died
waru	DUGUISUICS	nesponse.	$a_{\perp}e_{\alpha}$

Factor	Chi-Square	d.f.	P
intubated	19.54	1	<.0001
resection	4.14	1	0.0418
TOTAL	25.47	2	<.0001

13.14.6 Validation of Model E

validate(res_modE, method="boot", B=40)

	index.orig	training	test	optimism	index.corrected	n
Dxy	0.7340	0.7286	0.7309	-0.0022	0.7363	40
R2	0.4128	0.4222	0.3996	0.0227	0.3901	40
Intercept	0.0000	0.0000	0.0062	-0.0062	0.0062	40
Slope	1.0000	1.0000	0.9604	0.0396	0.9604	40
Emax	0.0000	0.0000	0.0100	0.0100	0.0100	40
D	0.2408	0.2445	0.2319	0.0126	0.2282	40
U	-0.0149	-0.0149	0.0091	-0.0241	0.0091	40
Q	0.2558	0.2594	0.2228	0.0367	0.2191	40

```
B 0.0727 0.0687 0.0776 -0.0089 0.0816 40 g 1.3970 1.4728 1.3697 0.1030 1.2939 40 gp 0.1597 0.1539 0.1561 -0.0023 0.1620 40
```

Our bootstrap validated assessments of discrimination and goodness of fit look somewhat more reasonable now.

13.14.7 Do any points seem particularly influential?

```
As a last step, I'll look at influence, and residuals, associated with model E.
```

```
inf.E <- which.influence(res_modE, cutoff=0.3)
inf.E</pre>
```

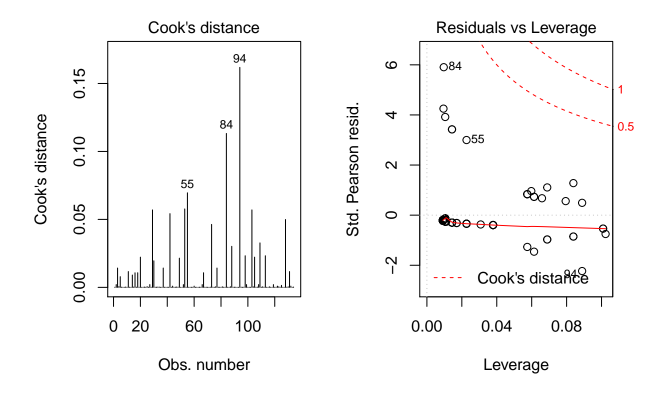
```
$Intercept
[1] "84" "94"

$resection
[1] "84" "94"

show.influence(inf.E, dframe = data.frame(resect))
```

```
Count resection 84 2 *2 94 2 *6
```

13.14.8 Fitting Model E using glm to get plots about influence



Using this glm residuals approach, we again see that points 84 and 94 have the largest influence on our model E.

13.15 Concordance: Comparing Model C, D and E's predictions

To start, we'll gather the predictions fomade by each model (C, D and E) on the probability scale, in one place. Sadly, augment from broom doesn't work well with 1rm fits, so we have to do this on our own.

```
# A tibble: 6 x 9
           age prior resection intubated
                                           died
                                                        С
                                                                       Ε
  <int> <int>
                          <dbl>
                                     <int> <int>
                                                   <dbl>
                                                           <dbl>
                                                                   <dbl>
               <int>
            34
                           2.50
                                         0
                                                0 0.0705 0.0632 0.0367
1
                   1
      2
           57
                   0
2
                           5.00
                                         0
                                                0 0.326
                                                          0.0620 0.130
3
      3
           60
                   1
                           4.00
                                                1 0.187
                                                          0.791
                                         1
4
      4
           62
                   1
                           4.20
                                         0
                                                0 0.211
                                                         0.158
                                                                 0.0881
5
      5
            28
                   0
                           6.00
                                         1
                                                1 0.504 0.711 0.819
6
      6
                   0
                           3.00
                                                0 0.0990 0.0737 0.0477
           52
```

And now, we'll use the gather command to arrange the models and predicted probabilities in a more useful manner for plotting.

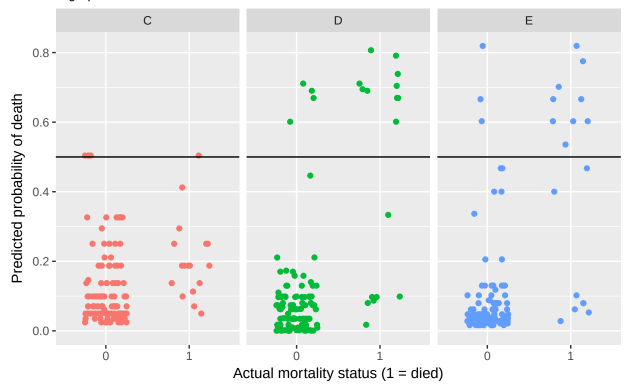
```
res_p <- resect_preds %>%
    gather("model", "prediction", 7:9) %>%
    select(id, died, model, prediction)
head(res_p)
```

```
# A tibble: 6 x 4
   id died model prediction
 <int> <int> <chr> <dbl>
   1 0 C
                0.0705
1
    2
        0 C
2
                 0.326
                0.187
3
    3 1 C
4
   4 0 C
                0.211
5
   5 1 C
                0.504
    6 0 C
6
                  0.0990
```

Here's the resulting plot.

Comparing Predictions for our Three Models

A graphical view of concordance



We could specify a particular rule, for example: if the predicted probability of death is 0.5 or greater, then predict "Died".

C Predict Alive 114 16
Predict Died 3 1
D Predict Alive 113 7
Predict Died 4 10
E Predict Alive 114 8
Predict Died 3 9

And perhaps build the linked table of row probabilities...

```
round(100*prop.table(
   ftable(table(res_p$model, res_p$rule.5, res_p$died))
   ,1),2)
```

```
0 1
C Predict Alive 87.69 12.31
Predict Died 75.00 25.00
D Predict Alive 94.17 5.83
```

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```
Predict Died 28.57 71.43
E Predict Alive 93.44 6.56
Predict Died 25.00 75.00
```

For example, in model E, 93.44% of those predicted to be alive actually survived, and 75% of those predicted to die actually died.

- Model D does a little better in one direction (94.17% of those predicted by Model D to be alive actually survived) but worse in the other (71.43% of those predicted by Model D to die actually died.)
- Model C does worse than each of the others in both predicting those who survive and those who die.

Note that the approaches discussed here would be useful if we had a new sample to predict on, as well. We could then compare the errors for that new data made by this sort of classification scheme either graphically or in a table.

13.16 Conclusions

It appears that intubated status and, to a lesser degree, the extent of the resection both play a meaningful role in predicting death associated with tracheal carina resection surgery. Patients who are intubated are associated with worse outcomes (greater risk of death) and more extensive resections are also associated with worse outcomes.

Chapter 14

Logistic Regression and the smartcle1 data

14.1 The smartcle1 data

Recall that the smartcle1.csv data file available on the Data and Code page of our website describes information on 11 variables for 1036 respondents to the BRFSS 2016, who live in the Cleveland-Elyria, OH, Metropolitan Statistical Area. As we've discussed in previous work, the variables in the smartcle1.csv file are listed below, along with (in some cases) the BRFSS items that generate these responses.

Variable	Description
SEQNO	respondent identification number (all begin with 2016)
physhealth	Now thinking about your physical health, which includes physical illness and injury, for how many days during the past 30 days was your physical health not good?
menthealth	Now thinking about your mental health, which includes stress, depression, and problems with emotions, for how many days during the past 30 days was your mental health not good?
poorhealth	During the past 30 days, for about how many days did poor physical or mental health keep you from doing your usual activities, such as self-care, work, or recreation?
genhealth	Would you say that in general, your health is (five categories: Excellent, Very Good, Good, Fair or Poor)
bmi	Body mass index, in kg/m ²
female	Sex, $1 = \text{female}$, $0 = \text{male}$
internet30	Have you used the internet in the past 30 days? $(1 = yes, 0 = no)$
exerany	During the past month, other than your regular job, did you participate in any physical activities or exercises such as running, calisthenics, golf, gardening, or walking for exercise? $(1 = yes, 0 = no)$
sleephrs	On average, how many hours of sleep do you get in a 24-hour period?
alcdays	How many days during the past 30 days did you have at least one drink of any alcoholic beverage such as beer, wine, a malt beverage or liquor?

In this section, we'll use some of the variables described above to predict the binary outcome: exerany.

14.2 Thinking About Non-Linear Terms

We have enough observations here to consider some non-linearity for our model.

In addition, since the genhealth variable is an ordinal variable and multi-categorical, we should consider how to model it. We have three options:

- 1. include it as a factor in the model (the default approach)
- 2. build a numeric version of that variable, and then restrict our model to treat that numeric variable as ordinal (forcing the categories to affect the exerany probabilities in an ordinal way), rather than as a simple nominal factor (so that if the effect of fair vs. good was to decrease the probability of 'exerany', then the effect of poor vs. good would have to decrease the probability at least as much as fair vs. good did.) Treating the genhealth variable as ordinal could be accomplished with the scored function in the rms package.
- 3. build a numeric version of genhealth and then use the catg function to specify the predictor as nominal and categorical, but this will lead to essentially the same model as choice 1.

Suppose we've decided to treat the genhealth data as categorical, without restricting the effect of its various levels to be ordinal. Suppose also that we've decided to include the following seven variables in our model for exerany:

- physhealth
- menthealth
- genhealth
- bmi
- female
- internet30
- sleephrs

Suppose we have a subject matter understanding that:

- the impact of bmi on exerany is affected by female, so we plan a female x bmi interaction term
- we're using internet30 as a proxy for poverty, and we think that an interaction with self-reported genhealth will be helpful in our model as well.

Note that we do have some missing values in some of these predictors, so we'll have to deal with that soon.

```
n variables: 8
```

Variable type: factor variable missing complete

variable missing complete n n_unique genhealth 3 1033 1036 5 top_counts ordered 2_V: 350, 3_G: 344, 1_E: 173, 4_F: 122 FALSE

Variable type: integer

```
sd p0 p25 median p75 p100
  variable missing complete
                                n mean
   exerany
                 3
                       1033 1036 0.76 0.43
                                            0
                                                 1
                                                         1
                                                             1
    female
                 0
                       1036 1036 0.6 0.49
                                                 0
                                                             1
                                                                  1
                 6
internet30
                       1030 1036 0.81 0.39 0
                                                 1
                                                        1
                                                             1
                                                                  1
menthealth
                       1025 1036 2.72 6.82 0
                                                             2
                                                                 30
                11
                       1019 1036 3.97 8.67 0
physhealth
                17
                                                        0
                                                                 30
```

```
sleephrs 8 1028 1036 7.02 1.53 1 6 7 8 20

Variable type: numeric

variable missing complete n mean sd p0 p25 median p75 p100

bmi 84 952 1036 27.89 6.47 12.71 23.7 26.68 30.53 66.06
```

14.3 A First Model for exerany (Complete Case Analysis)

Suppose we develop a main-effects kitchen sink model (model m1 below) fitted to these predictors without the benefit of any non-linear terms except the two pre-planned interactions. We'll run the model quickly here to ensure that the code runs, in a complete case analysis, without drawing any conclusions, really.

```
Frequencies of Missing Values Due to Each Variable
exerany internet30 genhealth bmi female physhealth
3 6 3 84 0 17
menthealth sleephrs
11 8
```

Logistic Regression Model

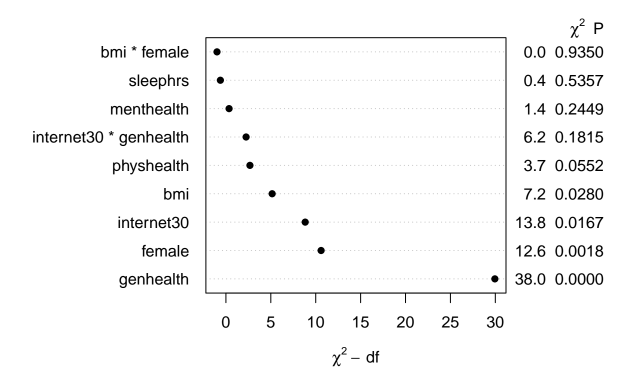
bmi * female

```
lrm(formula = exerany ~ internet30 * genhealth + bmi * female +
    physhealth + menthealth + sleephrs, data = smartcle1)
```

		Model Like Ratio T			iminatio	on	Rank Discrim. Indexes	
Obs	924	LR chi2	134.17	R2	0.20	03	С	0.739
0	219	d.f.	15	g	1.02		Dxy	0.479
1	705	Pr(> chi2)		gr	2.7		gamma	
max deriv	6e-13			gp	0.1	74	tau-a	0.173
				Brier	0.19	53		
			Coef	S.E.	Wald Z	Pr(>	Z)	
Intercept			2.9284	0.9951	2.94	0.003	33	
internet30			0.7234	0.7095	1.02	0.30	79	
genhealth=2	VeryGood		-0.2267	0.7653	-0.30	0.76	71	
genhealth=3	Good		-0.7950	0.7100	-1.12	0.26	29	
genhealth=4	Fair		-1.2818	0.7482	-1.71	0.086	67	
genhealth=5	Poor		-0.5215	0.8590	-0.61	0.543	38	
bmi			-0.0323	0.0220	-1.47	0.142	27	
female			-0.5789	0.7883	-0.73	0.463	27	
physhealth			-0.0208	0.0109	-1.92	0.05	52	
menthealth			-0.0142	0.0122	-1.16	0.24	49	
sleephrs			-0.0338	0.0547	-0.62	0.53	57	
internet30 ×	genheal	th=2_VeryGoo	od 0.1524	0.8295	0.18	0.854	42	
internet30 ×	genheal	th=3_Good	-0.0064	0.7702	-0.01	0.993	34	
internet30 ×	genheal	th=4_Fair	-0.3235	0.8272	-0.39	0.69	57	
internet30 ×	genheal	th=5_Poor	-1.7847	0.9994	-1.79	0.074	41	

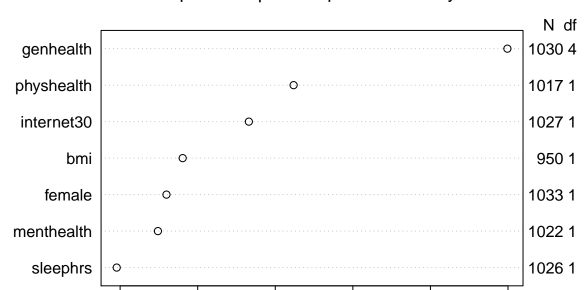
-0.0022 0.0266 -0.08 0.9350

plot(anova(m1))



14.4 Building a Larger Model: Spearman ρ^2 Plot

Before we impute, we might also consider the use of a Spearman ρ^2 plot to decide how best to spend degrees of freedom on non-linear terms in our model for exerany using these predictors. Since we're already planning some interaction terms, I'll keep them in mind as I look at this plot.



Spearman ρ^2 Response: exerany

We see that the best candidate for a non-linear term is the genhealth variable, according to this plot, followed by the physhealth and internet30 predictors, then bmi. I will wind up fitting a model including the following non-linear terms...

Adjusted ρ^2

0.04

0.06

0.08

0.10

• our pre-planned female x bmi and internet30 x genhealth interaction terms,

0.02

• a new genhealth x physhealth interaction term,

0.00

- a restricted cubic spline with 5 knots for physhealth
- a restricted cubic spline with 4 knots for bmi (so the interaction term with female will need to account for this and restrict our interaction to the linear piece of bmi)

14.5 A Second Model for exerany (Complete Cases)

Here's the resulting model fit without worrying about imputation yet. This is just to make sure our code works. Note that I'm inserting the main effects of our interaction terms explicitly before including the interaction terms themselves, and that I need to use %ia% to include the interaction terms where one of the terms is included in the model with a spline. Again, I won't draw any serious conclusions yet.

Frequencies of Missing Values Due to Each Variable exerany bmi physhealth female internet30 genhealth

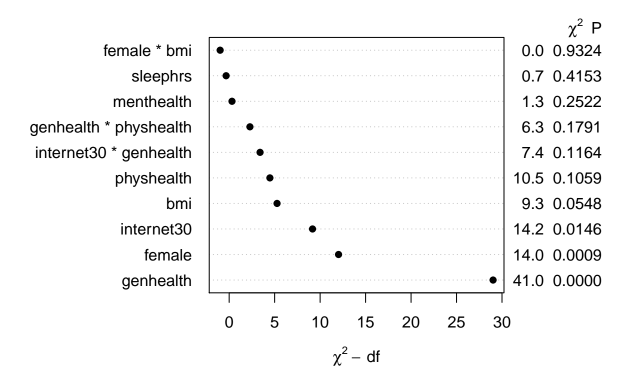
```
3 84 17 0 6 3
menthealth sleephrs
11 8
```

Logistic Regression Model

```
lrm(formula = exerany ~ rcs(bmi, 4) + rcs(physhealth, 5) + female +
   internet30 * genhealth + genhealth %ia% physhealth + female %ia%
   bmi + menthealth + sleephrs, data = smartcle1)
```

		Model Likel			ood	Discrimination			Rank Discrim. Indexes	
			atio '				dexes	. –		
	924	LR ch	112	142	2.51	R2	0.2		C	0.743
	219	d.f.			22	g	1.0		Dxy	0.485
	705	Pr(>	chi2)	<0.0	0001	gr	2.86		gamma	0.485
max deriv 6e	-13					gp	0.1		tau-a	0.176
						Brier	0.1	51		
				Co	ef	S.E.	Wald Z	Pr(>	IZI)	
Intercept				2	2.2998	1.9789	1.16	0.245	52	
bmi				C	.0052	0.0801	0.06	0.948	32	
bmi'				-0	.2988	0.3555	-0.84	0.400	06	
bmi''				C	.8809	0.9615	0.92	0.359	96	
physhealth				C	.0410	0.0868	0.47	0.636	32	
physhealth'				-0	.2944	0.3888	-0.76	0.448	39	
female				-0	.6307	0.8228	-0.77	0.443	34	
internet30				C	.6722	0.7134	0.94	0.346	30	
genhealth=2_Ve	ryGood			-0	.3092	0.7700	-0.40	0.688	30	
genhealth=3_Go	od			-0	.7559	0.7194	-1.05	0.293	34	
genhealth=4_Fa	ir			-1	.1611	0.7646	-1.52	0.128	39	
genhealth=5_Po	or			-2	.3680	1.2846	-1.84	0.065	53	
genhealth=2_Ve	ryGood	* phy	sheal	th C	.0099	0.0824	0.12	0.904	13	
genhealth=3_Go	od * p	hyshea	lth	-0	.0406	0.0772	-0.53	0.599	92	
genhealth=4_Fa	ir * p	hyshea	lth	-0	.0407	0.0773	-0.53	0.598	38	
genhealth=5_Po	or * p	hyshea	lth	C	.0534	0.0868	0.62	0.538	32	
female * bmi				-0	.0023	0.0275	-0.08	0.932	24	
menthealth				-0	.0143	0.0125	-1.15	0.252	22	
sleephrs				-0	.0457	0.0561	-0.81	0.415	53	
internet30 * g	enheal	th=2_V	eryGo	od C	.2174	0.8330	0.26	0.794	11	
internet30 * g	enheal	th=3_G	lood	C	.0375	0.7767	0.05	0.961	15	
internet30 * g	enheal	th=4_F	air'	-0	.3080	0.8371	-0.37	0.712	29	
internet30 * g	enheal	th=5_P	oor	-1	.9411	1.0150	-1.91	0.055	58	

plot(anova(m2))



14.6 Dealing with Missing Data via Simple Imputation

One approach we might take in this problem is to use simple imputation to deal with our missing values. I will proceed as follows:

- 1. Omit all cases where the outcome exerany is missing.
- 2. Determine (and plot) the remaining missingness.
- 3. Use simple imputation for all predictors, and build a new data set with "complete" data.
- 4. Re-fit the proposed models using this new data set.

14.6.1 Omit cases where the outcome is missing

We need to drop the cases where exerany is missing in smartcle1. We'll begin creating an imputed data set, called smartcle_imp0, by filtering on complete data for exerany, as follows.

```
Hmisc::describe(smartcle1$exerany)
smartcle1$exerany
       n missing distinct
                                Info
                                          Sum
                                                             Gmd
                                                  Mean
    1033
                         2
                               0.546
                                          786
                                                0.7609
                                                          0.3642
smartcle_imp0 <- smartcle1 %>%
    filter(complete.cases(exerany)) %>%
    select(SEQNO, exerany, physhealth, menthealth,
           genhealth, bmi, female, internet30, sleephrs)
```

```
Hmisc::describe(smartcle_imp0$exerany)
```

14.6.2 Plot the remaining missingness

We'll look at the missing values (excluding the subject ID: SEQNO) in our new data set. Of course, we can get a count of missing values within each variable with skim or with:

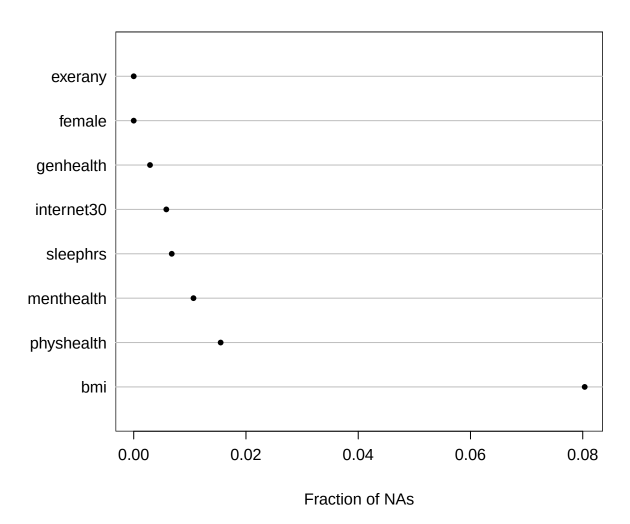
```
colSums(is.na(smartcle_imp0))
```

```
SEQNO exerany physhealth menthealth genhealth bmi 0 0 16 11 3 83 female internet30 sleephrs 0 6 7
```

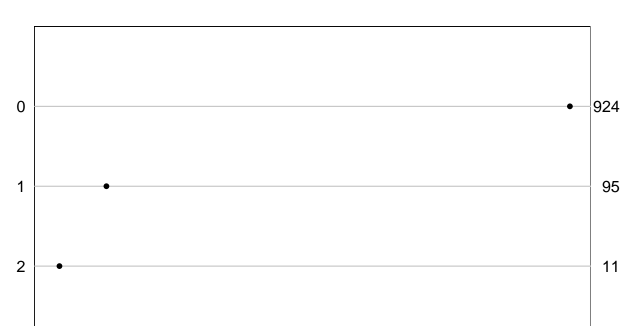
The Hmisc package has a plotting approach which can help identify missingness, too.

```
naplot(naclus(select(smartcle_imp0, -SEQNO)))
```

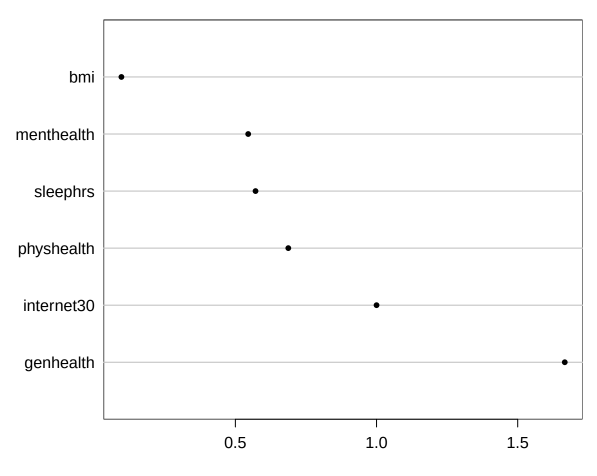
Fraction of NAs in each Variable



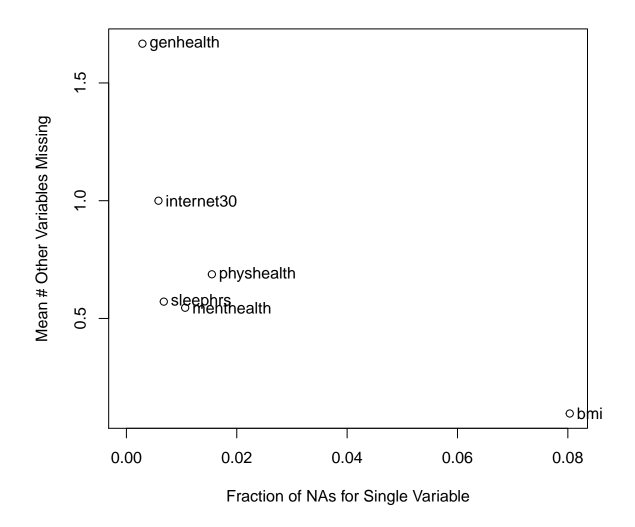
Number of Missing Variables Per Observation



Mean Number of Other Variables Missing for Observations where Indicated Variable is NA



Mean Number of NAs



We can also get a useful accounting of missing data patterns, with the md.pattern function in the mice package.

mice::md.pattern(smartcle_imp0)

	SEQNO	exerany	${\tt female}$	genhealth	${\tt internet30}$	sleephrs	menthealth
924	1	1	1	1	1	1	1
8	1	1	1	1	1	1	1
6	1	1	1	1	1	1	0
76	1	1	1	1	1	1	1
1	1	1	1	1	0	1	1
4	1	1	1	1	1	0	1
3	1	1	1	1	1	1	0
1	1	1	1	0	1	1	0
1	1	1	1	1	1	1	1
4	1	1	1	1	0	1	1
1	1	1	1	1	1	0	1
1	1	1	1	1	1	0	1

```
1
                                           0
                                                          1
                                                                      1
                                                                                    0
                     1
                              1
                                           1
                                                          0
                                                                      1
                                                                                    1
  1
          1
          1
                     1
                              1
                                           0
                                                          1
                                                                      0
                                                                                    1
                     0
                              0
                                           3
                                                          6
                                                                      7
          0
                                                                                   11
     physhealth bmi
924
                      1
                           0
                 1
  8
                 0
                      1
                           1
  6
                 1
                      1
                           1
 76
                 1
                      0
                           1
  1
                 1
                      1
                           1
  4
                 1
                      1
                           1
                           2
                 0
  3
                      1
  1
                 1
                      1
                           2
                      0
                           2
  1
                 0
  4
                      0
                           2
                 1
  1
                 0
                      1
                           2
                      0
                           2
  1
                 1
  1
                 0
                      1
                           3
                 0
                      0
                           3
  1
  1
                 0
                      1
                           3
               16
                    83 126
```

We can also do this with na.pattern in the Hmisc package, but then we have to get the names of the columns, too, so that we can read off the values.

```
na.pattern(smartcle_imp0)
pattern
924
                      76
                1
8
                1
                      1
                            1
    1
                                  1
001110000
names(smartcle_imp0)
[1] "SEQNO"
                  "physhealth" "menthealth" "genhealth"
          "exerany"
[6] "bmi"
          "female"
                  "internet30" "sleephrs"
```

14.6.3 Use simple imputation, build a new data set

The only variables that require no imputation are exerany and female. In this case, we need to impute:

- 83 bmi values (which are quantitative)
- 16 physhealth values (quantitative, must fall between 0 and 30)
- 11 menthealth values (quantitative, must fall between 0 and 30)
- 7 sleephrs values (quantitative, must fall between 0 and 24)
- 6 internet30 values (which are 1/0)
- and 3 genhealth values (which are multi-categorical, so we need to convert them to numbers in order to get the imputation process to work properly)

```
smartcle_imp0 <- smartcle_imp0 %>%
  mutate(genh_n = as.numeric(genhealth))
smartcle_imp0 %>% count(genhealth, genh_n)
```

```
# A tibble: 6 x 3
  genhealth genh_n
                          n
  <fct>
               <dbl> <int>
                1.00
1 1_Excellent
                        172
2 2_VeryGood
                2.00
                        350
                3.00
                        344
3 3 Good
                4.00
4 4_Fair
                        121
5 5_Poor
                5.00
                         43
6 <NA>
               NA
                          3
```

I'll work from the bottom up, using various simputation functions to accomplish the imputations I want. In this case, I'll use predictive mean matching for the categorical data, and linear models or elastic net approaches for the quantitative data. Be sure to set a seed beforehand so you can replicate your work.

```
set.seed(432234)

smartcle_imp1 <- smartcle_imp0 %>%
    impute_pmm(genh_n ~ female) %>%
    impute_pmm(internet30 ~ female + genh_n) %>%
    impute_lm(sleephrs ~ female + genh_n) %>%
    impute_lm(menthealth ~ female + sleephrs) %>%
    impute_en(physhealth ~ female + genh_n + sleephrs) %>%
    impute_en(bmi ~ physhealth + genh_n)
```

After the imputations are complete, I'll back out of the numeric version of genhealth, called genh_n back to my original variable, then check to be sure I now have no missing values.

```
# A tibble: 5 x 3
  genhealth genh_n
                         n
  <fct>
               <dbl> <int>
1 1_Excellent
                1.00
                       172
2 2_VeryGood
                2.00
                       351
3 3_Good
                3.00
                       346
                4.00
4 4_Fair
                       121
                5.00
5 5_Poor
                         43
colSums(is.na(smartcle_imp1))
```

```
SEQNO exerany physhealth menthealth genhealth bmi 0 0 0 0 0 0 0 0 female internet30 sleephrs genh_n 0 0 0 0
```

OK. Looks good. I now have a data frame called smartcle_imp1 with no missingness, which I can use to fit my logistic regression models. Let's do that next, and then return to the problem of accounting for missingness through multiple imputation.

14.7 Refitting Model 1 with simply imputed data

Using the numeric version of the genhealth data, called genh_n, will ease the reviewing of later output, so we'll do that here, making sure R knows that genh_n describes a categorical factor.

Logistic Regression Model

```
lrm(formula = exerany ~ internet30 * catg(genh_n) + bmi * female +
    physhealth + menthealth + sleephrs, data = smartcle_imp1,
    x = TRUE, y = TRUE)
```

		Model Li	kelihood	Discri	${ t mination}$	Rank Di	iscrim.
		Ratio	Test	Inde	exes	Inde	exes
Obs	1033	LR chi2	151.09	R2	0.204	C	0.741
0	247	d.f.	15	g	1.023	Dxy	0.482
1	786	Pr(> chi2	<0.0001	gr	2.780	gamma	0.483
max de:	riv 5e-14			gp	0.175	tau-a	0.176
				Brier	0.154		

```
Coef
                             S.E. Wald Z Pr(>|Z|)
Intercept
                      3.3390 0.9731 3.43 0.0006
internet30
                      0.6487 0.7010 0.93 0.3548
genh_n=2
                     -0.4938 0.7459 -0.66 0.5080
                     -0.9270 0.6972 -1.33 0.1836
genh_n=3
genh_n=4
                     -1.3060 0.7378 -1.77 0.0767
genh_n=5
                     -0.8093 0.8329 -0.97 0.3312
bmi
                     -0.0412 0.0217 -1.90 0.0570
female
                     -0.9473 0.7722 -1.23 0.2199
                     -0.0229 0.0102 -2.25 0.0246
physhealth
menthealth
                     -0.0200 0.0114 -1.76 0.0784
sleephrs
                     -0.0415 0.0492 -0.84 0.3994
internet30 * genh_n=2 0.3701 0.8065 0.46 0.6463
internet30 * genh_n=3 0.1295 0.7543 0.17 0.8637
internet30 * genh n=4 -0.2194 0.8108 -0.27 0.7867
internet30 * genh_n=5 -1.4080 0.9636 -1.46 0.1440
bmi * female
                      0.0118 0.0261 0.45 0.6497
```

All right. We've used 1033 observations, which is correct (after deleting the three with missing exerany. The model shows a Nagelkerke R^2 value of 0.204, and a C statistic of 0.741 after imputation. The likelihood ratio (drop in deviance) test is highly significant.

14.7.1 Validating Summary Statistics

```
set.seed(432099)
validate(m1_a)
```

	<pre>index.orig</pre>	training	test	${\tt optimism}$	${\tt index.corrected}$	n
Dxy	0.4823	0.5012	0.4632	0.0380	0.4444	40
R2	0.2039	0.2218	0.1878	0.0340	0.1699	40
Intercept	0.0000	0.0000	0.0936	-0.0936	0.0936	40
Slope	1.0000	1.0000	0.9001	0.0999	0.9001	40
Emax	0.0000	0.0000	0.0401	0.0401	0.0401	40
D	0.1453	0.1592	0.1329	0.0264	0.1189	40
U	-0.0019	-0.0019	0.0013	-0.0032	0.0013	40
Q	0.1472	0.1612	0.1316	0.0295	0.1177	40
В	0.1539	0.1507	0.1566	-0.0058	0.1597	40
g	1.0226	1.0868	0.9722	0.1146	0.9080	40
gp	0.1751	0.1818	0.1673	0.0145	0.1606	40

It appears that the model's description of summary statistics is a little optimistic for both the C statistic (remember that C = 0.5 + Dxy/2) and the Nagelkerke R^2 . This output suggests that in a new sample of data, our model might be better expected to show a C statistic near ...

$$C = 0.5 + \frac{Dxy}{2} = 0.5 + \frac{0.4444}{2} = 0.7222$$

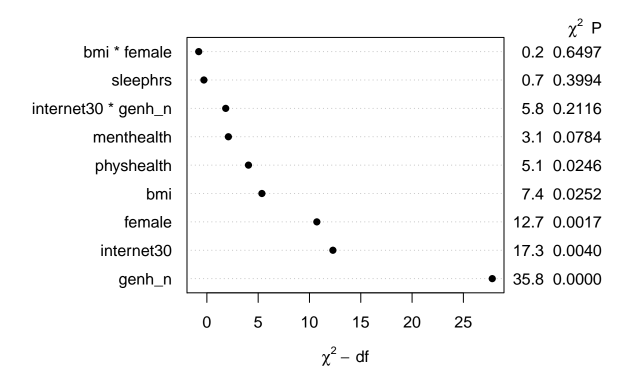
rather than the 0.741 we saw initially, and that the Nagelkerke R^2 in new data will be closer to 0.17, than to the nominal 0.204 we saw above. So, as we walk through some other output for this model, remember that the C statistic wasn't great here (0.72 after validation), so our ability to discriminate exercisers from non-exercisers is still a problem.

14.7.2 ANOVA for the model

anova(m1_a)

Next, let's look at the ANOVA comparisons for this model.

Wald Statistics Respons	e: exerany		
Factor	Chi-Square	d.f.	P
internet30 (Factor+Higher Order Factors)	17.28	5	0.0040
All Interactions	5.84	4	0.2116
<pre>genh_n (Factor+Higher Order Factors)</pre>	35.81	8	<.0001
All Interactions	5.84	4	0.2116
bmi (Factor+Higher Order Factors)	7.36	2	0.0252
All Interactions	0.21	1	0.6497
female (Factor+Higher Order Factors)	12.72	2	0.0017
All Interactions	0.21	1	0.6497
physhealth	5.05	1	0.0246
menthealth	3.10	1	0.0784
sleephrs	0.71	1	0.3994
<pre>internet30 * genh_n (Factor+Higher Order Facto</pre>	rs) 5.84	4	0.2116
<pre>bmi * female (Factor+Higher Order Factors)</pre>	0.21	1	0.6497
TOTAL INTERACTION	6.15	5	0.2917
TOTAL	126.95	15	<.0001
plot(anova(m1_a))			

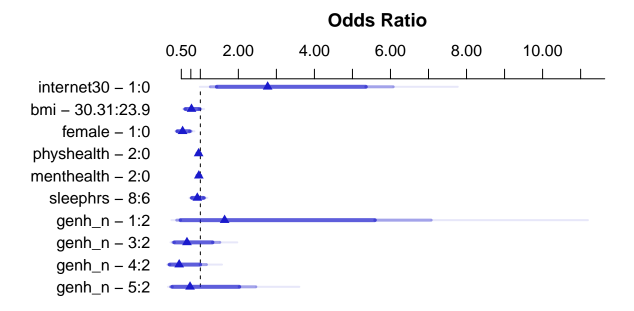


It looks like several of the variables (genhealth, internet30, female, bmi and physhealth) are carrying statistically significant predictive value here.

14.7.3 Summarizing Effect Size

How big are the effects we see?

plot(summary(m1_a))



Adjusted to:internet30=0 genh_n=2 bmi=27.03416 female=0

summary(m1_a)

Effects			Response : exerany				
Factor	Low	High	Diff.	Effect	S.E.	Lower 0.95	Upper 0.95
internet30	0.0	_		1.018800			
Odds Ratio	0.0	1.00	1.00	2.770000	NA	1.263800	6.0710000
bmi	23.9	30.31	6.41	-0.264180	0.138810	-0.536240	0.0078783
Odds Ratio	23.9	30.31	6.41	0.767840	NA	0.584950	1.0079000
female	0.0	1.00	1.00	-0.627270	0.177840	-0.975830	-0.2787200
Odds Ratio	0.0	1.00	1.00	0.534050	NA	0.376880	0.7567500
physhealth	0.0	2.00	2.00	-0.045843	0.020393	-0.085813	-0.0058727
Odds Ratio	0.0	2.00	2.00	0.955190	NA	0.917770	0.9941400
menthealth	0.0	2.00	2.00	-0.040089	0.022778	-0.084733	0.0045549
Odds Ratio	0.0	2.00	2.00	0.960700	NA	0.918760	1.0046000
sleephrs	6.0	8.00	2.00	-0.082947	0.098439	-0.275880	0.1099900
Odds Ratio	6.0	8.00	2.00	0.920400	NA	0.758900	1.1163000
genh_n - 1:2	2.0	1.00	NA	0.493760	0.745930	-0.968240	1.9558000
Odds Ratio	2.0	1.00	NA	1.638500	NA	0.379750	7.0693000
genh_n - 3:2	2.0	3.00	NA	-0.433290	0.431840	-1.279700	0.4131100
Odds Ratio	2.0	3.00	NA	0.648370	NA	0.278120	1.5115000
genh_n - 4:2	2.0	4.00	NA	-0.812200	0.490270	-1.773100	0.1487200
Odds Ratio	2.0	4.00	NA	0.443880	NA	0.169800	1.1603000
genh_n - 5:2	2.0	5.00	NA	-0.315510	0.620110	-1.530900	0.8998700
Odds Ratio	2.0	5.00	NA	0.729420	NA	0.216340	2.4593000

Adjusted to: internet30=0 genh_n=2 bmi=27.03416 female=0

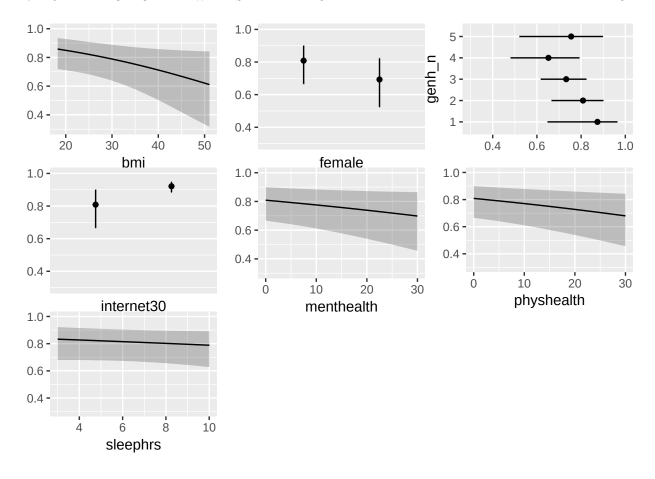
This output is easier to read as a result of using small *numeric* labels in genh_n, rather than the lengthy labels in genhealth. The sensible things to interpret are the odds ratios.

- holding all other predictors constant, the effect of moving from internet30 = 0 to internet30 = 1 is that the odds of exerany increase by a factor of 2.77.
 - Suppose Harry and Steve have the same values of all predictors in this model except that Harry used the internet and Steve did not.
 - So the odds of exercising for Harry (who used the internet) are 2.77 times higher than the odds of exercising for Steve (who didn't use the internet), assuming that all other predictors are the same.
 - We also have a 95% confidence interval for this odds ratio, which is (1.26, 6.07). Since 1 is not in that interval, the data don't seem to be consistent with internet30 having no effect on exerany.
- the odds ratio comparing two subjects with the same predictors except that Harry has a BMI of 30.31 (the 75th percentile of observed BMIs in our sample) and Marty has a BMI of 23.9 (the 25th percentile) is that Harry has 0.767 times the odds of exercising that Marty does. So Harry's probability of exercise will also be lower than Marty's.
 - The 95% confidence interval in this case is (0.58, 1.01), and because 1 is in that interval, we cannot conclude that the effect of bmi meets the standard for statistical significance at the 5% level.
- A similar approach can be used to describe the odds ratios associated with each predictor.
- Note that each of the categories in genh_n is compared to a single baseline category. Here, that's category 2. R will pick the modal category: the one that appears most often in the data. The comparisons of each category against category 2 are not significant in each case, at the 5% level.

14.7.4 Plotting the Model with ggplot and Predict

Let's look at a series of plots describing the model on the probability scale.

ggplot(Predict(m1_a, fun = plogis))



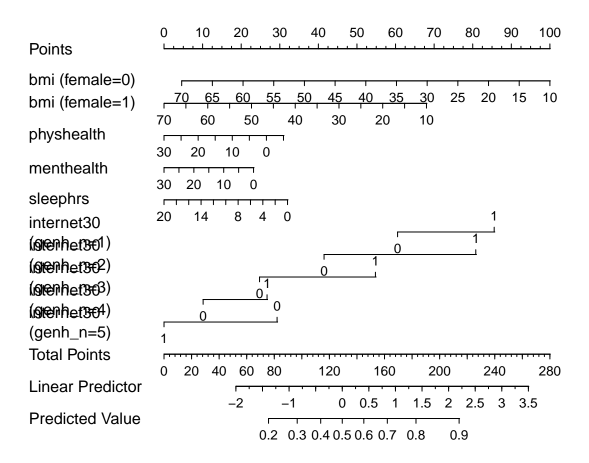
This helps us describe what is happening in terms of direction at least. For example,

- As bmi increases, predicted Pr(exerany) decreases.
- People who accessed the internet in the past 30 days have higher morel probabilities of exercising.

Do any of these plots fail to make sense to you? Is anything moving in a surprising direction?

14.7.5 Plotting the model with a nomogram

```
plot(nomogram(m1_a, fun = plogis))
```

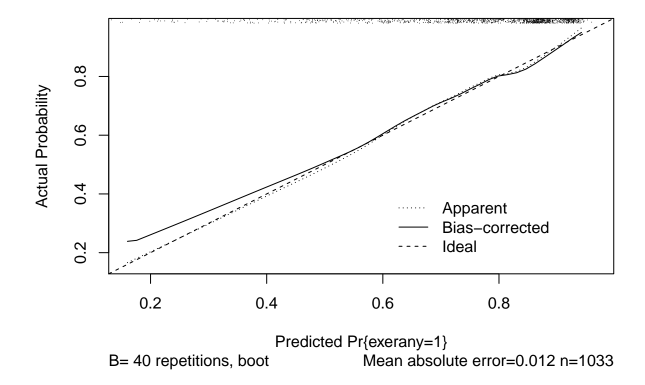


Note the impact of our interaction terms, and how we have two lines for bmi and five lines for internet30 that come out of our product terms. As with any nomogram, to make a prediction we:

- 1. find the values of each of our predictors in the scales, and travel vertically up to the Points line to read off the Points for that predictor.
- 2. sum up the Points across all predictors, and find that location in the Total Points line.
- 3. move vertically down from the total points line to find the estimated "linear predictor" (log odds ratio) and finally the "predicted value" (probability of our outcome exerany = 1.)

14.7.6 Checking the Calibration of our model

plot(calibrate(m1_a))



n=1033 Mean absolute error=0.012 Mean squared error=0.00022 0.9 Quantile of absolute error=0.02

The model appears better calibrated for predicted probabilities above 0.5 or so than it does for lower predicted probabilities. This isn't surprising given the rug plot at the top, which shows we make many more predictions at the high end of the scale.

To test the goodness of fit, we can use the following omnibus test:

Our non-significant p value suggests that we cannot detect anything that's obviously wrong in the model in terms of goodness of fit. That's comforting.

14.8 Refitting Model 2 with simply imputed data

I'll walk through the same tasks for Model m2 that I did above for Model m1. Again, we're running this model after simple imputation of missing values.

Using the numeric version of the genhealth data, called genh_n, will ease the reviewing of later output, so we'll do that here, making sure R knows that genh_n describes a categorical factor.

Logistic Regression Model

```
lrm(formula = exerany ~ rcs(bmi, 4) + rcs(physhealth, 5) + female +
  internet30 * catg(genh_n) + catg(genh_n) %ia% physhealth +
  female %ia% bmi + menthealth + sleephrs, data = smartcle_imp1,
  x = TRUE, y = TRUE)
```

		Model Likelihood		Discrim	Discrimination		Rank Discrim.	
		Ratio	Test	Inde	exes	Inde	exes	
Obs	1033	LR chi2	159.51	R2	0.214	C	0.744	
0	247	d.f.	22	g	1.051	Dxy	0.487	
1	786	Pr(> chi2)	<0.0001	gr	2.860	gamma	0.487	
max	deriv 5e-12			gp	0.179	tau-a	0.177	
				Brier	0 152			

```
Coef
                             S.E.
                                    Wald Z Pr(>|Z|)
Intercept
                      2.7525 1.9074 1.44 0.1490
bmi
                     -0.0072 0.0763 -0.09 0.9250
bmi'
                     -0.2772 0.3288 -0.84 0.3992
bmi''
                     0.9199 0.9873 0.93 0.3515
physhealth
                     0.0287 0.0905 0.32 0.7514
physhealth'
                     -0.1396 0.3555 -0.39 0.6946
female
                     -0.9550 0.8017 -1.19 0.2336
                     0.6099 0.7040 0.87 0.3863
internet30
                     -0.5059 0.7507 -0.67 0.5004
genh_n=2
genh n=3
                     -0.8611 0.7053 -1.22 0.2221
genh_n=4
                     -1.1827 0.7535 -1.57 0.1165
genh_n=5
                     -2.6842 1.1987 -2.24 0.0251
genh_n=2 * physhealth -0.0253 0.0864 -0.29 0.7698
genh_n=3 * physhealth -0.0475 0.0842 -0.56 0.5723
genh_n=4 * physhealth -0.0471 0.0842 -0.56 0.5758
genh_n=5 * physhealth 0.0497 0.0917 0.54 0.5879
female * bmi
                      0.0111 0.0268 0.41 0.6803
menthealth
                     -0.0200 0.0116 -1.72 0.0859
sleephrs
                     -0.0536 0.0503 -1.07 0.2868
internet30 * genh_n=2 0.4193 0.8091 0.52 0.6043
internet30 * genh_n=3 0.1624 0.7592 0.21 0.8306
internet30 * genh_n=4 -0.1802 0.8176 -0.22 0.8255
internet30 * genh_n=5 -1.6615 0.9820 -1.69 0.0907
```

All right. We've again used 1033 observations, which is correct (after deleting the three with missing exerany. The model shows a Nagelkerke R^2 value of 0.214, and a C statistic of 0.744 after imputation. Each of these results are a little better than what we saw with $\mathtt{m1}$ _a but only a little. The likelihood ratio (drop in deviance)

test is still highly significant.

14.8.1 Validating Summary Statistics

set.seed(432009)
validate(m2_a)

	index.orig	training	test	${\tt optimism}$	${\tt index.corrected}$	n
Dxy	0.4870	0.5058	0.4590	0.0468	0.4402	40
R2	0.2145	0.2340	0.1849	0.0491	0.1654	40
Intercept	0.0000	0.0000	0.1729	-0.1729	0.1729	40
Slope	1.0000	1.0000	0.8473	0.1527	0.8473	40
Emax	0.0000	0.0000	0.0692	0.0692	0.0692	40
D	0.1534	0.1695	0.1307	0.0387	0.1147	40
U	-0.0019	-0.0019	0.0041	-0.0060	0.0041	40
Q	0.1554	0.1714	0.1266	0.0448	0.1106	40
В	0.1521	0.1502	0.1567	-0.0065	0.1585	40
g	1.0507	1.1335	0.9626	0.1708	0.8799	40
gp	0.1786	0.1866	0.1642	0.0224	0.1562	40

Again, the model's description of summary statistics is a little optimistic for both the C statistic and the Nagelkerke R². In a new sample of data, model m2_a might be better expected to show a C statistic near ...

$$C = 0.5 + \frac{Dxy}{2} = 0.5 + \frac{0.4402}{2} = 0.7201$$

rather than the 0.744 we saw initially, and that the Nagelkerke R² in new data will be closer to 0.165, than to the nominal 0.215 we saw above. So, after validation, this model actually looks worse than model m1_a.

Model	nominal C	nominal R ²	validated C	validated R ²
m1_a	0.741	0.204	0.722	0.170
$m2_a$	0.744	0.214	0.720	0.165

Again, as we walk through other output for model m2_a, remember that the our ability to discriminate exercisers from non-exercisers is still very much in question using either model.

14.8.2 ANOVA for the model

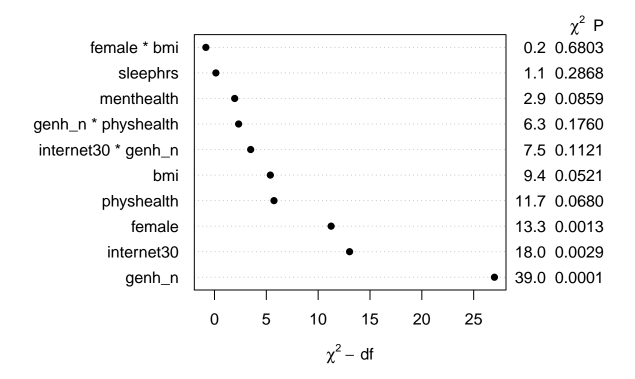
Next, let's look at the ANOVA comparisons for this model.

Mold Ctotictics

anova(m2_a)

Wald	Statistics	Response:	exerany		
Factor			Chi-Square	d.f.	P
bmi (Factor+Higher	Order Factors)		9.39	4	0.0521
All Interactions			0.17	1	0.6803
Nonlinear			1.46	2	0.4831
physhealth (Factor-	Higher Order Factors	3)	11.74	6	0.0680
All Interactions			6.33	4	0.1760
Nonlinear			0.15	1	0.6946
female (Factor+High	ner Order Factors)		13.25	2	0.0013
All Interactions			0.17	1	0.6803

internet30 (Factor+Higher Order Factors)	18.03	5	0.0029	
All Interactions	7.49	4	0.1121	
genh_n (Factor+Higher Order Factors)	39.00	12	0.0001	
All Interactions	11.80	8	0.1604	
<pre>genh_n * physhealth (Factor+Higher Order Factors)</pre>	6.33	4	0.1760	
<pre>female * bmi (Factor+Higher Order Factors)</pre>	0.17	1	0.6803	
menthealth	2.95	1	0.0859	
sleephrs	1.13	1	0.2868	
<pre>internet30 * genh_n (Factor+Higher Order Factors)</pre>	7.49	4	0.1121	
TOTAL NONLINEAR	1.65	3	0.6474	
TOTAL INTERACTION	12.10	9	0.2075	
TOTAL NONLINEAR + INTERACTION	13.40	12	0.3409	
TOTAL	131.03	22	<.0001	
plot(anova(m2_a))				

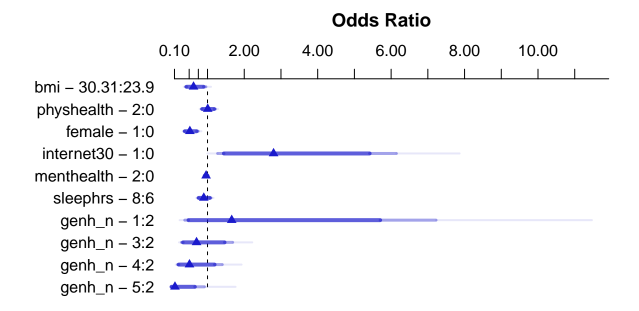


Here, it looks like just three of the variables (genhealth, internet30, and female) are carrying statistically significant predictive value.

14.8.3 Summarizing Effect Size

How big are the effects we see?

plot(summary(m2_a))



Adjusted to:bmi=27.03416 physhealth=0 female=0 internet30=0 genh_n=2

summary(m2_a)

Effects				Response : exerany			
Factor	Low	High	Diff.	Effect	S.E.	Lower 0.95	Upper 0.95
bmi	23.9	30.31	6.41	-0.4797500	0.221460	-0.913810	-0.045691
Odds Ratio	23.9	30.31	6.41	0.6189400	NA	0.400990	0.955340
physhealth	0.0	2.00	2.00	0.0055399	0.095955	-0.182530	0.193610
Odds Ratio	0.0	2.00	2.00	1.0056000	NA	0.833160	1.213600
female	0.0	1.00	1.00	-0.6558700	0.183810	-1.016100	-0.295600
Odds Ratio	0.0	1.00	1.00	0.5189900	NA	0.361990	0.744080
internet30	0.0	1.00	1.00	1.0291000	0.401140	0.242910	1.815300
Odds Ratio	0.0	1.00	1.00	2.7986000	NA	1.275000	6.143200
menthealth	0.0	2.00	2.00	-0.0399460	0.023258	-0.085530	0.005638
Odds Ratio	0.0	2.00	2.00	0.9608400	NA	0.918030	1.005700
sleephrs	6.0	8.00	2.00	-0.1072500	0.100690	-0.304600	0.090096
Odds Ratio	6.0	8.00	2.00	0.8983000	NA	0.737420	1.094300
genh_n - 1:2	2.0	1.00	NA	0.5059000	0.750690	-0.965430	1.977200
Odds Ratio	2.0	1.00	NA	1.6585000	NA	0.380820	7.222700
genh_n - 3:2	2.0	3.00	NA	-0.3552100	0.447130	-1.231600	0.521140
Odds Ratio	2.0	3.00	NA	0.7010200	NA	0.291830	1.684000
genh_n - 4:2	2.0	4.00	NA	-0.6768400	0.517450	-1.691000	0.337350
Odds Ratio	2.0	4.00	NA	0.5082200	NA	0.184330	1.401200
genh_n - 5:2	2.0	5.00	NA	-2.1783000	1.066000	-4.267700	-0.089045
Odds Ratio	2.0	5.00	NA	0.1132300	NA	0.014015	0.914800

Adjusted to: bmi=27.03416 physhealth=0 female=0 internet30=0 genh_n=2

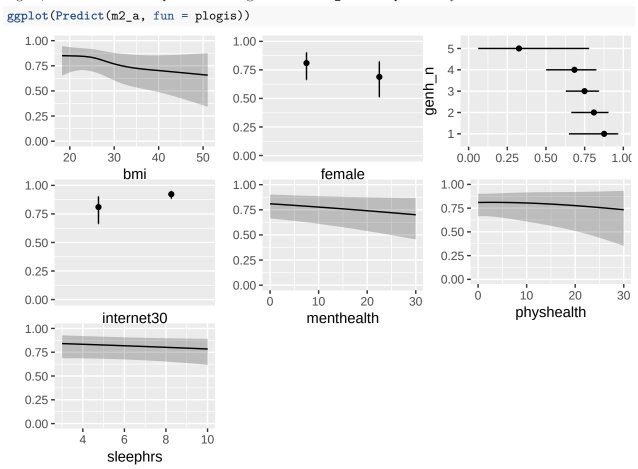
This output is easier to read as a result of using small *numeric* labels in genh_n, rather than the lengthy labels in genhealth. The sensible things to interpret are the odds ratios. For example,

- holding all other predictors constant, the effect of moving from internet30 = 0 to internet30 = 1 is that the odds of exerany increase by a factor of 2.80.
- the odds ratio comparing two subjects with the same predictors except that Harry has a BMI of 30.31 (the 75th percentile of observed BMIs in our sample) and Marty has a BMI of 23.9 (the 25th percentile) is that Harry has 0.619 times the odds of exercising that Marty does. So Harry's probability of exercise will also be lower.

By sex, which group has a larger probability of exerany, holding all other predictors constant, by this model? Females or Males?

14.8.4 Plotting the Model with ggplot and Predict

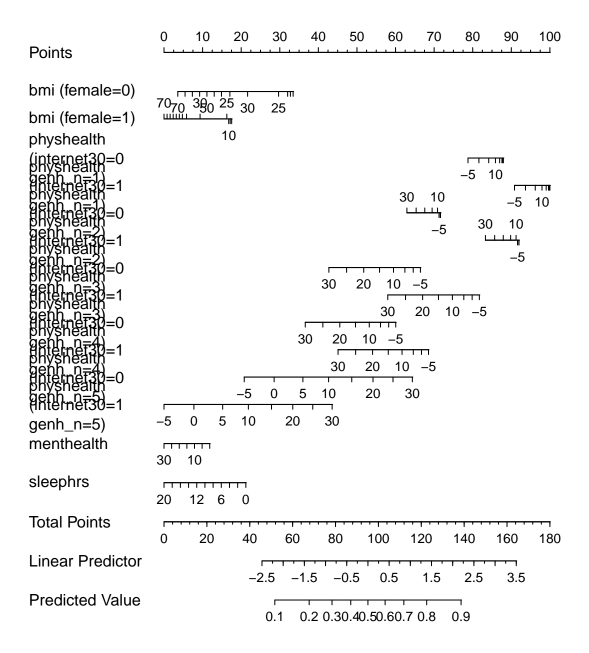
Again, consider a series of plots describing the model m2_a on the probability scale.



Note the small kink in the bmi plot. To what do you attribute this?

14.8.5 Plotting the model with a nomogram

plot(nomogram(m2_a, fun = plogis))



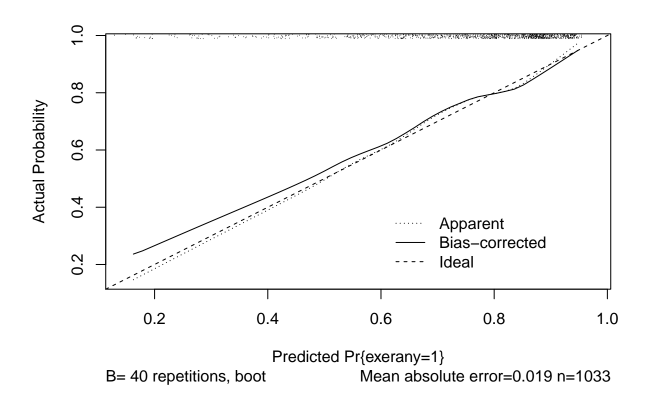
Note the impact of our interaction terms, and the cubic splines in bmi and physhealth. As with any nomogram, to make a prediction we:

- 1. find the values of each of our predictors in the scales, and travel vertically up to the Points line to read off the Points for that predictor.
- 2. sum up the Points across all predictors, and find that location in the Total Points line.

3. move vertically down from the total points line to find the estimated "linear predictor" (log odds ratio) and finally the "predicted value" (probability of our outcome exerany = 1.)

14.8.6 Checking the Calibration of our model

plot(calibrate(m2_a))



n=1033 Mean absolute error=0.019 Mean squared error=0.00048 0.9 Quantile of absolute error=0.027

The model appears to be a bit less well calibrated than m1_a was.

To test the goodness of fit, we can use the following omnibus test:

round(residuals(m2_a, type = "gof"),3)

Sum of squared errors	Expected value HO	SD
157.102	157.011	0.522
Z	P	
0.174	0.862	

Our non-significant p value suggests that we cannot detect anything that's obviously wrong in the model in terms of goodness of fit.

14.9 Comparing Model 2 to Model 1 after simple imputation

We can refit the models with glm and then compare them with anova, aic and bic approaches, if we like.

14.9.1 Comparison by Analysis of Deviance

```
anova(m1_a_glm, m2_a_glm)
Analysis of Deviance Table
Model 1: exerany ~ internet30 * factor(genh_n) + bmi * female + physhealth +
    menthealth + sleephrs
Model 2: exerany ~ rcs(bmi, 4) + rcs(physhealth, 5) + female + internet30 *
    factor(genh_n) + factor(genh_n) %ia% physhealth + female %ia%
    bmi + menthealth + sleephrs
 Resid. Df Resid. Dev Df Deviance
1
       1017
               985.32
       1010
                976.89 7
                             8.4245
To obtain a p value, we can compare this drop in deviance to a \chi^2 distribution with 7 df, as follows:
pchisq(8.4245, 7, lower.tail = FALSE)
[1] 0.2966531
```

So there's no statistically significant advantage apparent from fitting the larger $\mathtt{m2_a}$ model.

14.9.2 Comparing AIC and BIC

```
glance(m1_a_glm)

null.deviance df.null logLik AIC BIC deviance df.residual
1  1136.406 1032 -492.6596 1017.319 1096.363 985.3192 1017

glance(m2_a_glm)

null.deviance df.null logLik AIC BIC deviance df.residual
1  1136.406 1032 -488.4473 1022.895 1136.52 976.8946 1010
```

Model m1_a_glm shows lower AIC and BIC than does m2_a_glm, again suggesting no meaningful advantage for the larger model.

14.10 Dealing with Missing Data via Multiple Imputation

Coming soon.

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