### Classical statistical models

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**BIOF 339** 

### Statistical models

# All models are wrong, but some are useful

G.E.P. Box

#### **Models**

Models are our way of understanding nature, usually using some sort of mathematical expression

Famous mathematical models include Newton's second law of motion, the laws of thermodynamics, the ideal gas law

All probability distributions, like Gaussian, Binomial, Poisson, Gamma, are models

Mendel's laws are models **that result in** particular mathematical models for inheritance and population prevalence

#### **Models**

We use models all the time to describe our understanding of different processes

- Cause-and-effect relationships
- Supply-demand curves
- Financial planning
- Optimizing travel plans (perhaps including traffic like Google Maps)
- Understanding the effects of change
  - Climate change
  - Rule changes via Congress or companies
  - Effect of a drug on disease outcomes
  - Effect of education and behavioral patterns on future earnings

#### **Data-driven models**

Can we use data collected on various aspects of a particular context to understand the relationships between the different aspects?

- How does increased smoking affect your risk of getting lung cancer? (causality/association)
  - o Does genetics matter?
  - Does the kind of smoking matter?
  - Does gender matter?

#### **Data-driven models**

Can we use data collected on various aspects of a particular context to understand the relationships between the different aspects?

- What is your lifetime risk of breast cancer? (prediction)
  - What if you have a sister with breast cancer?
  - What if you had early menarche?
  - What if you are of Ashkenazi Jewish heritage?

The Gail Model from NCI

#### **Association models**

These are more traditional, highly interpretative models that look at **how** different predictors affect outcome.

- Linear regression
- Logistic regression
- Cox proportional hazards regression
- Decision trees

Since these models have a particular known structure determined by the modeler, they can be used on relatively small datasets

You can easily understand which predictors have more "weight" in influencing the outcome

You can literally write down how a prediction would be made

#### **Predictive models**

These are more recent models that primarily look to provide good predictions of an outcome, and the way the predictions are made is left opaque (often called a *black box*)

- Deep Learning (or Neural Networks)
- Random Forests
- Support Vector Machines
- Gradient Boosting Machines

These models require data to both determine the structure of the model as well as make the predictions, so they require lots of data to *train* on

The relative "weight" of predictors in influencing the **predictions** can be obtained

The effect of individual predictors is not easily interpretable, though this is changing

They require a different **philosophic perspective** than traditional association models

#### R for statistical models

We've seen that R is great for data munging and data visualizations

R also can fit a wide variety of statistical models to data.

In fact, most new models first are implemented in R (see CRAN and GitHub)

Today we'll describe some standard popular models. Fitting most models follow the same pattern of code.

#### **Datasets**

We will use the pbc data from the survival package, and the in-built mtcars dataset.

```
library(survival)
str(pbc)
```

```
'data.frame':
               418 obs. of 20 variables:
          : int 1 2 3 4 5 6 7 8 9 10 ...
$ id
$ time
                400 4500 1012 1925 1504 2503 1832 2466 2400 51 ...
          : int
$ status : int 2 0 2 2 1 2 0 2 2 2 ...
$ trt
         : int 111122212...
$ age
      : num
                58.8 56.4 70.1 54.7 38.1 ...
          : Factor w/ 2 levels "m", "f": 2 2 1 2 2 2 2 2 2 2 ...
$ ascites : int 1 0 0 0 0 0 0 0 0 1 ...
$ hepato : int 1 1 0 1 1 1 1 0 0 0 ...
$ spiders : int 1 1 0 1 1 0 0 0 1
$ edema : num
$ bili
          : int 261 302 176 244 279 248 322 280 562 200 ...
$ albumin : num 2.6 4.14 3.48 2.54 3.53 3.98 4.09 4 3.08 2.74 ...
$ copper : int 156 54 210 64 143 50 52 52 79 140 ...
$ alk.phos: num 1718 7395 516 6122 671 ...
$ ast
         : num
               137.9 113.5 96.1 60.6 113.2 ...
$ trig
          : int 172 88 55 92 72 63 213 189 88 143 ...
$ platelet: int 190 221 151 183 136 NA 204 373 251 302 ...
$ protime : num 12.2 10.6 12 10.3 10.9 11 9.7 11 11 11.5 ...
$ stage
          : int 4 3 4 4 3 3 3 3 2 4
```

### The formula interface

# Representing model relationships

In R, there is a particularly convenient way to express models, where you have

- one dependent variable
- one or more independent variables, with possible transformations and interactions

```
y \sim x1 + x2 + x1:x2 + I(x3^2) + x4*x5
```

y depends on ...

- x1 and x2 linearly
- the interaction of x1 and x2 (represented as x1:x2)
- the square of x3 (the I() notation ensures that the ^ symbol is interpreted correctly)
- x4, x5 and their interaction (same as x4 + x5 + x4:x5)

# Representing model relationships

```
y \sim x1 + x2 + x1:x2 + I(x3^2) + x4*x5
```

This interpretation holds for the vast majority of statistical models in R

 For decision trees and random forests and neural networks, don't add interactions or transformations, since the model will try to figure those out on their own

```
myLinearModel <- lm(chol ~ bili + albumin + copper + sex, data = pbc)</pre>
```

Note that everything in R is an **object**, so you can store a model in a variable name.

This statement runs the model and stored the fitted model in myLinearModel

R does not interpret the model, evaluate the adequacy or appropriateness of the model, or comment on whether looking at the relationship between cholesterol and bilirubin makes any kind of sense.

It just fits the model it is given

```
myLinearModel
```

```
Call:
lm(formula = chol ~ bili + albumin + copper + sex, data = pbc)

Coefficients:
(Intercept) bili albumin copper sexf
221.0571 22.7113 28.9076 -0.1888 -9.7605
```

Not very informative, is it?

summary(myLinearModel)

```
Call:
lm(formula = chol ~ bili + albumin + copper + sex, data = pbc)
Residuals:
                                 Max
   Min
            10 Median
                          3Q
-580.83 -90.62 -34.79 37.96 1297.16
Coefficients:
           Estimate Std. Error t value Pr(>|t|)
(Intercept) 221.0571 135.6962 1.629
                                       0.104
bili
        22.7113
                    3.2821 6.920 3.14e-11 ***
albumin 28.9076
                      33.8309
                              0.854
                                       0.394
copper -0.1888
                    0.1743 -1.083
                                       0.280
sexf
            -9.7605
                      40.8253 -0.239
                                       0.811
Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
Residual standard error: 214.3 on 277 degrees of freedom
  (136 observations deleted due to missingness)
Multiple R-squared: 0.1638, Adjusted R-squared: 0.1517
F-statistic: 13.56 on 4 and 277 DF, p-value: 4.147e-10
```

A little better

broom::tidy(myLinearModel)

```
# A tibble: 5 x 5
             estimate std.error statistic p.value
 term
                         <dbl>
 <chr>
               <dbl>
                                  <dbl>
                                          <dbl>
1 (Intercept) 221.
                       136.
                                  1.63 1.04e- 1
2 bili
                      3.28
              22.7
                                  6.92 3.14e-11
              28.9
3 albumin
                                 0.854 3.94e- 1
                       33.8
4 copper
              -0.189
                      0.174
                                 -1.08 2.80e- 1
5 sexf
              -9.76
                        40.8
                                 -0.239 8.11e- 1
```

broom::glance(myLinearModel)

library(gtsummary)
tbl\_regression(myLinearModel)

Characteristic	Beta	95% CI <sup>1</sup>	p-value	
bili	23	16, 29	< 0.001	
albumin	29	-38, 96	0.4	
copper	-0.19	-0.53, 0.15	0.3	
sex				
m	_	_		
f	-9.8	-90, 71	0.8	
<sup>1</sup> CI = Confidence Interval				

library(stargazer)
stargazer(myLinearModel, type='html')

	Dependent variable:		
	chol		
bili	22.711***		
	(3.282)		
albumin	28.908		
	(33.831)		
copper	-0.189		
	(0.174)		
sexf	-9.760		
	(40.825)		
Constant	221.057		

We do need some sense as to how well this model fit the data

```
# install.packages('ggfortify')
library(ggfortify)
autoplot(myLinearModel)
```

Let's see if we have some strangeness going on

```
ggplot(pbc, aes(x = bili))+geom_density()
```

We'd like this to be a bit more "Gaussian" for better behavior

Let's see if we have some strangeness going on

```
ggplot(pbc, aes(x = log(bili)))+geom_density()
```

```
myLinearModel2 <- lm(chol~log(bili) + albumin + copper + sex, data = pbc)
summary(myLinearModel2)</pre>
```

```
Call:
lm(formula = chol ~ log(bili) + albumin + copper + sex, data = pbc)
Residuals:
   Min
            10 Median
                          3Q
                                 Max
-448.77 -96.23 -26.77 40.76 1221.21
Coefficients:
           Estimate Std. Error t value Pr(>|t|)
(Intercept) 128.3685 132.9579 0.965
                                      0.3351
log(bili) 124.2339 14.8852 8.346 3.39e-15 ***
albumin 53.6093
                     33.2245
                             1.614 0.1078
copper -0.3775 0.1743 -2.166
                                     0.0312 *
           19.6595
sexf
                      39.1715 0.502 0.6161
Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
Residual standard error: 207.4 on 277 degrees of freedom
  (136 observations deleted due to missingness)
Multiple R-squared: 0.2163, Adjusted R-squared: 0.205
F-statistic: 19.11 on 4 and 277 DF, p-value: 6.792e-14
```

tbl\_regression(myLinearModel2)

Characteristic	Beta	95% CI <sup>1</sup>	p-value	
log(bili)	124	95, 154	< 0.001	
albumin	54	-12, 119	0.11	
copper	-0.38	-0.72, -0.03	0.031	
sex				
m	_	_		
f	20	-57, 97	0.6	
<sup>1</sup> CI = Confidence Interval				

autoplot(myLinearModel2)

# Just the residual plot, please

autoplot(myLinearModel2, which=1)

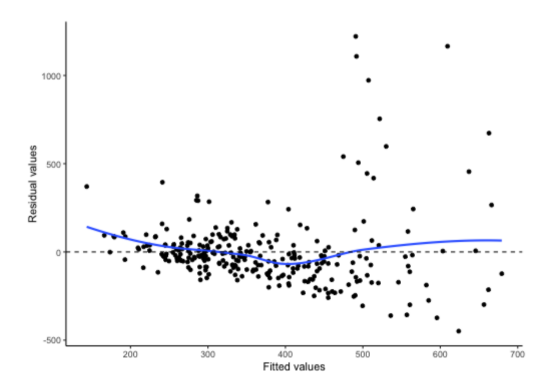
## Just the residual plot, please

```
d <- broom::augment(myLinearModel2, newdata=pbc)
d</pre>
```

```
# A tibble: 418 x 22
     id time status
                                      ascites hepato spiders edema bili chol
                      trt age sex
  <int> <int> <int> <dbl> <fct>
                                        <int> <int> <int> <dbl> <dbl> <int>
                          58.8 f
          400
                                                                   14.5
                                                                          261
         4500
                                                                    1.1
                                                                          302
                        1 56.4 f
         1012
                        1 70.1 m
                                                              0.5
                                                                    1.4
                                                                          176
         1925
                        1 54.7 f
                                                              0.5
                                                                    1.8
                                                                          244
        1504
                        2 38.1 f
                                                                    3.4
                                                                          279
      6
         2503
                                                                         248
                        2 66.3 f
                                                                    0.8
                        2 55.5 f
                                                                          322
        1832
         2466
                        2 53.1 f
                                                                    0.3
                                                                         280
         2400
                                                                    3.2
                                                                          562
                        1 42.5 f
                                                   0
           51
                        2 70.6 f
                                                                   12.6
                                                                          200
 ... with 408 more rows, and 10 more variables: albumin <dbl>, copper <int>,
   alk.phos <dbl>, ast <dbl>, trig <int>, platelet <int>, protime <dbl>,
   stage <int>, .fitted <dbl>, .resid <dbl>
```

# Just the residual plot, please

```
ggplot(d, aes(x = .fitted, y = .resid))+geom_point()+ geom_smooth(se=F)+
  labs(x = 'Fitted values', y = 'Residual values')+
  geom_hline(yintercept=0, linetype=2) +
  theme_classic()
```



### **Predictions**

```
head(predict(myLinearModel2, newdata = pbc))
```

```
1 2 3 4 5 6
560.7384 361.4248 277.4503 333.0571 435.3173 314.7947
```

The newdata has to have the same format and components as the original data the model was trained on

# **Categorical predictors**

```
myLM3 <- lm(chol ~ log(bili) + sex, data = pbc)
broom::tidy(myLM3)</pre>
```

```
# A tibble: 3 x 5
             estimate std.error statistic p.value
  term
 <chr>
                <dbl>
                          <dbl>
                                    <dbl>
                                             <dbl>
 (Intercept)
                283.
                                    7.71 2.14e-13
                           36.6
2 log(bili)
                                    8.22 7.37e-15
                 99.6
                           12.1
3 sexf
                 32.5
                           37.8
                                    0.858 3.92e- 1
```

R has a somewhat unfortunate notation for categorical variables here, as {variable name}{level}

# The logistic transformation

For an outcome which is binary (0/1), what is really modeled is the **probability** that the outcome is 1, usually denoted by p.

However, we know  $0 \le p \le 1$ , so what if the model gives a prediction outside this range!!

The logistic transform takes p to

$$\operatorname{logit}(p) = \operatorname{log}\!\left(rac{p}{1-p}
ight)$$

and we model logit(p), which has a range from  $-\infty$  to  $\infty$ 

Logistic regression is a special case of a **generalized linear model**, so the function we use to run a logistic regression is glm

```
myLR <- glm(spiders ~ albumin + bili + chol, data = pbc, family = binomial)
myLR
```

```
Call: glm(formula = spiders ~ albumin + bili + chol, family = binomial, data = pbc)

Coefficients:
(Intercept) albumin bili chol
2.3326484 -0.9954927 0.0995915 -0.0003176

Degrees of Freedom: 283 Total (i.e. Null); 280 Residual
(134 observations deleted due to missingness)
Null Deviance: 341.4
Residual Deviance: 315.2 AIC: 323.2
```

- We have to add the family = binomial as an argument, since this is a special kind of GLM
- All these models only use complete data; they kick out rows with missing data

broom::tidy(myLR)

```
# A tibble: 4 x 5
              estimate std.error statistic p.value
  term
                 <dbl>
                          <dbl>
                                   <dbl> <dbl>
 <chr>
1 (Intercept) 2.33
                       1.30
                                   1.80 0.0717
2 albumin
             -0.995
                       0.362
                                  -2.75 0.00595
3 bili
                       0.0344
                                 2.89 0.00381
             0.0996
4 chol
             -0.000318 0.000615
                                  -0.517 0.605
```

broom::glance(myLR)

tbl\_regression(myLR)

Characteristic	log(OR) <sup>1</sup>	95% CI <sup>1</sup>	p-value	
albumin	-1.0	-1.7, -0.30	0.006	
bili	0.10	0.04, 0.17	0.004	
chol	0.00	0.00, 0.00	0.6	
<sup>1</sup> OR = Odds Ratio, CI = Confidence Interval				

tbl\_regression(myLR, exponentiate = TRUE)

Characteristic	OR <sup>1</sup>	95% CI <sup>1</sup>	p-value	
albumin	0.37	0.18, 0.74	0.006	
bili	1.10	1.04, 1.19	0.004	
chol	1.00	1.00, 1.00	0.6	
<sup>1</sup> OR = Odds Ratio, CI = Confidence Interval				

# Predictions from logistic regression

```
head(predict(myLR))
```

```
1 2 3 4 5 6
1.10554163 -1.77506554 -1.04814132 -0.09414055 -0.93144911 -1.62851203
```

These are on the "wrong" scale. We would expect probabilities

```
head(predict(myLR, type='response'))
```

```
1 2 3 4 5 6
0.7512970 0.1449135 0.2595822 0.4764822 0.2826308 0.1640343
```

or you can use plogis(predict(myLR)) for the inverse logistic transform

# **Model selection**

### How to get the "best" model

Generally getting to the best model involves

- looking at a lot of graphs
- Fitting lots of models
- Comparing the model fits to see what seems good

Sometimes if you have two models that fit about the same, you take the smaller, less complex model (Occam's Razor)

Generally it is not recommended that you use automated model selection methods. It screws up your error rates and may not be the right end result for your objectives

Model building and selection is an art

### Clues to follow

You can look at the relative weights (size of coefficient and its p-value) of different predictors

• These weights will change once you change the model, so be aware of that

You can trim the number of variables based on collinearities

• If several variables are essentially measuring the same thing, use one of them

You can look at residuals for clues about transformations

You can look at graphs, as well as science, for clues about interactions (synergies and antagonisms)

#### **Automated model selection**

```
# install.packages('leaps')
library(leaps)
mtcars1 <- mtcars %>% mutate(across(c(cyl, vs:carb), as.factor))
all_subsets <- regsubsets(mpg~., data = mtcars1)
all_subsets</pre>
```

```
Subset selection object
Call: regsubsets.formula(mpg ~ ., data = mtcars1)
16 Variables (and intercept)
      Forced in Forced out
          FALSE
cyl6
                    FALSE
cyl8
          FALSE
                     FALSE
          FALSE
disp
                     FALSE
          FALSE
                     FALSE
hp
          FALSE
                     FALSE
drat
          FALSE
                     FALSE
wt
          FALSE
                     FALSE
gsec
          FALSE
                     FALSE
vs1
          FALSE
                     FALSE
am1
          FALSE
                     FALSE
gear4
          FALSE
                     FALSE
gear5
          FALSE
                     FALSE
carb2
          FALSE
                     FALSE
carb3
carb4
          FALSE
                     FALSE
          FALSE
                     FALSE
carb6
          FALSE
carb8
                     FALSE
 subsets of each size up to 8
Selection Algorithm: exhaustive
```

#### **Automated model selection**

Which has the best  $R^2$ ?

```
ind <- which.max(summary(all_subsets)$adjr2)
summary(all_subsets)$which[ind,]</pre>
```

```
(Intercept)
                                cyl8
                                            disp
                                                                      drat
                   cyl6
                                                           hp
       TRUE
                   TRUE
                               FALSE
                                           FALSE
                                                         TRUE
                                                                     FALSE
         wt
                   qsec
                                 vs1
                                             am1
                                                        gear4
                                                                     gear5
       TRUE
                  FALSE
                                TRUE
                                            TRUE
                                                        FALSE
                                                                     FALSE
      carb2
                  carb3
                               carb4
                                            carb6
                                                        carb8
      FALSE
                  FALSE
                               FALSE
                                           FALSE
                                                        FALSE
```