### **Bivariate Data**

2020

In this section we review the relationship between 2 variables.

### Bivariate categorical data

We use the same functions as in the case of one categorical variable.

Let's say we want to evaluate if the students who smoke study less?

And we have the following data for the smoke and amount variables. Where amount is presented as follows:

```
1 – less than 5 hours of studying;
```

- 2 5-10 hours of studying;
- 3 more than 10 hours of studying.

```
> smokes <- c("Yes", "No", "No", "Yes", "No", "Yes", "Yes", "Yes", "Yes", "No", "Yes")
> amount <- c(1, 2, 2, 3, 3, 1, 2, 1, 3, 2)
> table(smokes, amount)
    amount
smokes 1 2 3
No 0 2 2
Yes 3 2 1
```

Let's review the proportions.

To see it clearer we set the numbers to be rounded to the 3 decimal place

```
> options(digits = 3)
```

We can calculate proportions using the prop.table function or the conditional proportions using:

```
1 means that the proportions in the rows will sum to 1
```

2 means that the proportions in the columns will sum to 1

```
    > prop.table(table(smokes, amount))
        amount
smokes 1 2 3
        No 0.0 0.2 0.2
        Yes 0.3 0.2 0.1
    > prop.table(table(smokes, amount), 1)
        amount
smokes 1 2 3
        No 0.000 0.500 0.500
        Yes 0.500 0.333 0.167
    > prop.table(table(smokes, amount), 2)
        amount
smokes 1 2 3
```

No 0.000 0.500 0.667

#### Another example

```
> hair <- c("blond", "blond", "black", "blond", "brown", "brown",
           "brown", "brown", "black", "brown", "black", "brown",
           "black", "black", "brown", "brown", "brown",
+
           "brown", "brown", "black", "brown", "black", "brown",
+
           "blond", "blond", "black", "blond", "brown", "brown",
           "brown", "brown", "black", "brown", "black", "brown", "brown", "brown", "black", "brown", "black", "brown",
+
           "blond", "blond", "black", "blond", "brown", "brown")
+
> eyes <- c("blue", "green", "brown", "blue", "green", "brown",
           "brown", "black", "black", "green", "brown", "brown", "green", "black", "black", "brown", "brown", "black", "green", "black", "brown", "brown", "black", "brown", "blue", "green", "brown", "brown", "black", "brown", "blue", "green", "brown", "brown", "black",
+
+
           "black", "green", "brown", "blue", "green", "brown", "brown", "black", "brown", "blue", "green", "brown",
+
           "blue", "green", "brown", "blue", "green", "brown")
```

### > table(hair, eyes)

eyes

hair black blue brown green black 4 0 6 4 blond 0 5 2 2 brown 7 2 11 5

### > prop.table(table(hair, eyes))

eyes

hair black blue brown green black 0.0833 0.0000 0.1250 0.0833 blond 0.0000 0.1042 0.0417 0.0417 brown 0.1458 0.0417 0.2292 0.1042

### > prop.table(table(hair, eyes), 1)

eyes

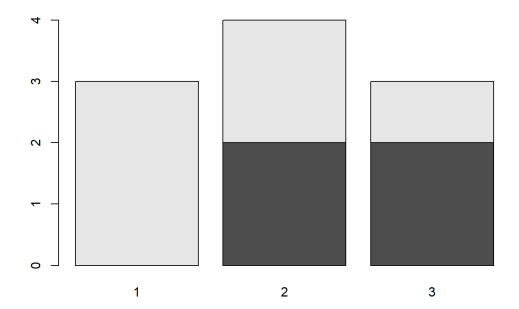
hair black blue brown green black 0.286 0.000 0.429 0.286 blond 0.000 0.556 0.222 0.222 brown 0.280 0.080 0.440 0.200

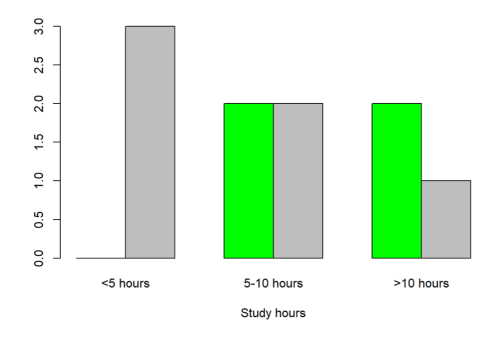
#### > prop.table(table(hair, eyes), 2)

eyes

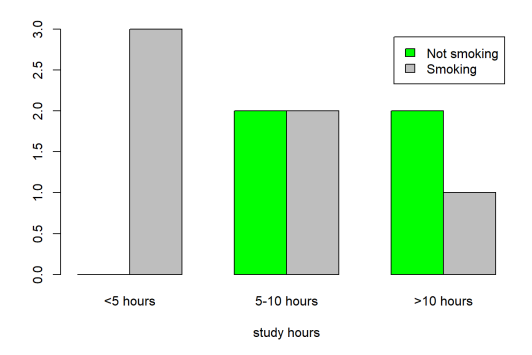
hair black blue brown green black 0.364 0.000 0.316 0.364 blond 0.000 0.714 0.105 0.182 brown 0.636 0.286 0.579 0.455 We can plot the data using barplot and pie functions.

### > barplot(table(smokes, amount))

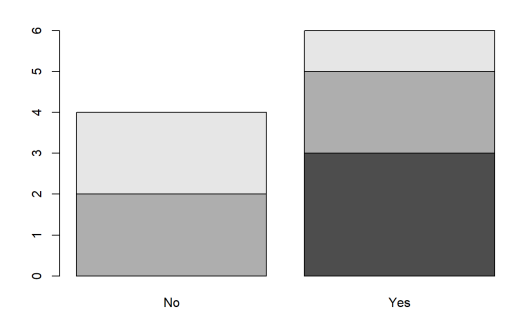




#### 

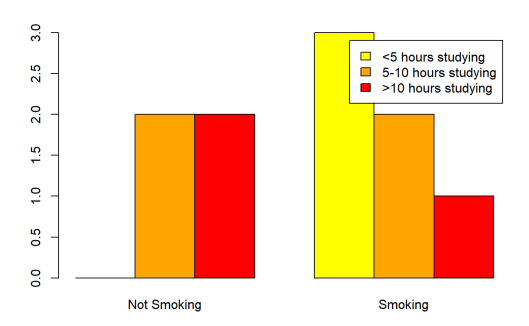


### > barplot(table(amount, smokes))



```
> barplot(table(amount, smokes),
+ names.arg = c("Not Smoking", "Smoking"),
```

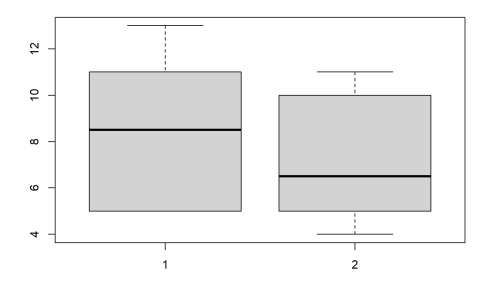
- + legend.text = c("<5 hours studying","5-10 hours studying",">10 hours studying"),
- + beside = TRUE,
- + col = c("Yellow", "Orange", "Red"))



### Categorical vs. numerical data

Simple example might be a drug test, where you have data for an experimental group and for a control group

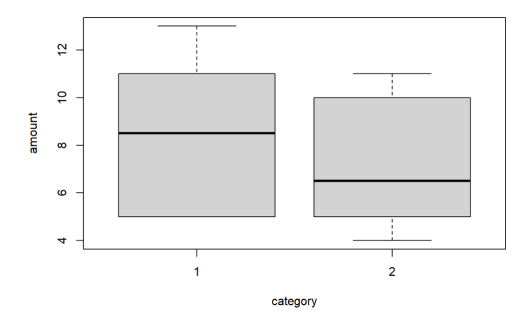
- > group.experimental <- c(5, 5, 5, 13, 7, 11, 11, 9, 8, 9)
- > group.control <- c(11, 8, 4, 5, 9, 5, 10, 5, 4, 10)
- > boxplot(group.experimental, group.control)



From the plot we see that the control group has smaller observations than the experimental group.

Another example will be if we have the same data, but structured differently. Here we can use the response ~ predictor, which will break up the values in response, by the categories in category

```
> amount = c(5, 5, 5, 13, 7, 11, 11, 9, 8, 9, 11, 8, 4, 5, 9, 5, 10, 5, 4, 10) > category = c(1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 2, 2, 2, 2, 2, 2, 2, 2, 2, 2) > boxplot(amount \sim category)
```



### Numerical vs. numerical data

Comparing two numerical variables can be done in different ways. If the two variables are thought to be independent samples you might like to compare their distributions in some manner. However, if you expect a relationship between the variables, you might like to look for that by plotting pairs of points.

#### Comparing two distributions with plot

Let's compare the distributions of the old and the new prices of homes from the home data frame:

> library(UsingR)

### > head(home)

old new

1 64200 257500

2 72100 276800

3 87600 364600

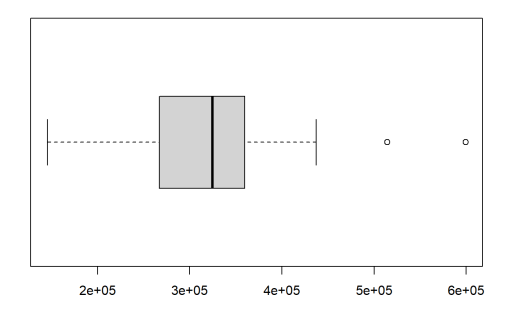
4 59000 160400

5 83200 333500

6 49100 145600

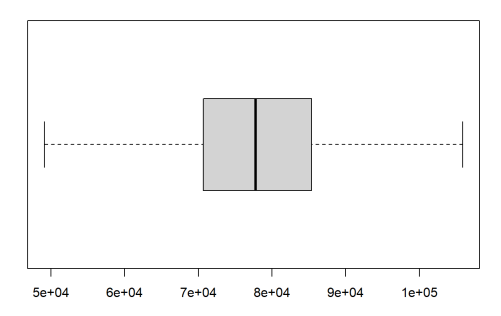
> boxplot(home\$new, horizontal = TRUE, main = "New Home Prices")

### **New Home Prices**



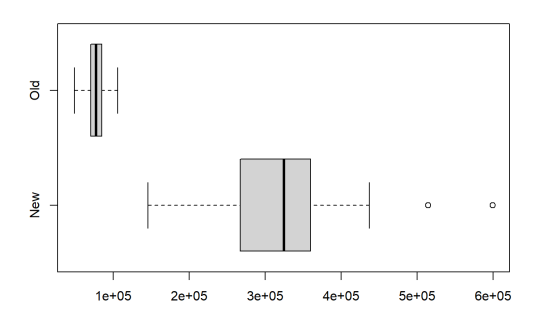
> boxplot(home\$old, horizontal = TRUE, main = "Old Home Prices")

### **Old Home Prices**



### > boxplot(home\$new, home\$old, horizontal = TRUE, names = c("New", "Old"))

scale puts the two data sets on the same scale so they can sensibly be compared



```
> head(home$old)
[1] 64200 72100 87600 59000 83200 49100
> mean(home$old)
[1] 78707
> sd(home$old)
[1] 15590
> head((home$old - mean(home$old)) / sd(home$old))
[1] -0.931 -0.424 0.570 -1.264 0.288 -1.899
> head(scale(home$old))
   [,1]
[1,] -0.931
[2,]-0.424
[3,] 0.570
[4,] -1.264
[5,] 0.288
[6,] -1.899
> head(home$new)
[1] 257500 276800 364600 160400 333500 145600
> mean(home$new)
[1] 329320
> sd(home$new)
[1] 121774
> head((home$new - mean(home$new)) / sd(home$new))
> head(scale(home$new))
    [,1]
[1,] -0.5898
[2,] -0.4313
```

[3,] 0.2897

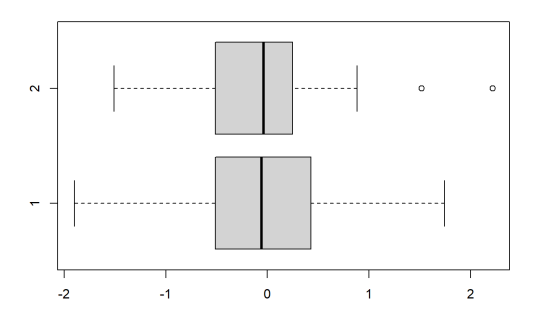
[4,] -1.3872

[5,] 0.0343

[6,] -1.5087

> boxplot(c(scale(home\$old)), c(scale(home\$new)), horizontal = TRUE)

Another example with different data frame on the same house prices



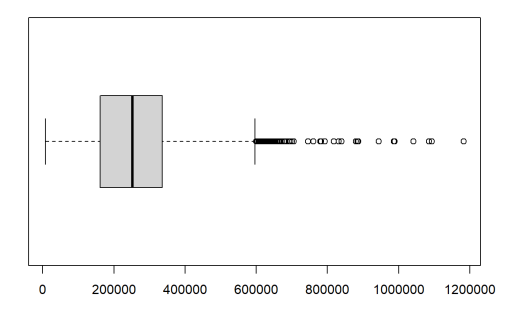
- > library(MASS)
- > head(homedata)

y1970 y2000

- 1 89700 359100
- 2 118400 504500
- 3 116400 477300
- 4 122000 500400
- 5 91500 433900
- 6 102800 464800

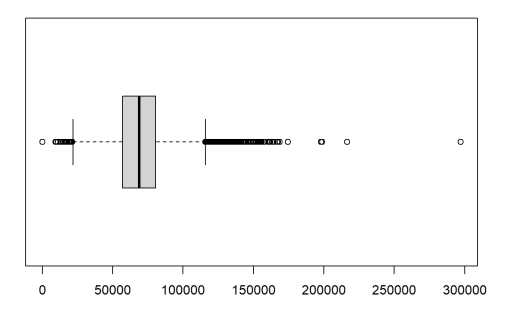
### > boxplot(homedata\$y2000, horizontal = TRUE, main = "Home Prices from 2000")

**Home Prices from 2000** 

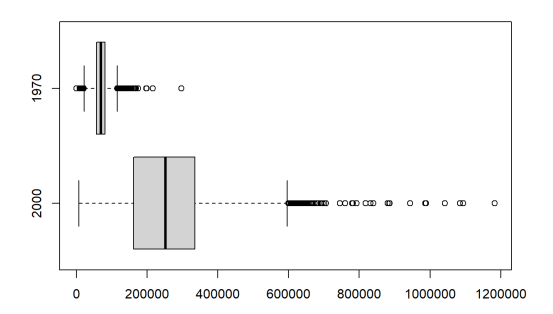


> boxplot(homedata\$y1970, horizontal = TRUE, main = "Home Prices from 1970")

**Home Prices from 1970** 



> boxplot(homedata\$y2000, homedata\$y1970, horizontal = TRUE, names = c("2000", "1970"))



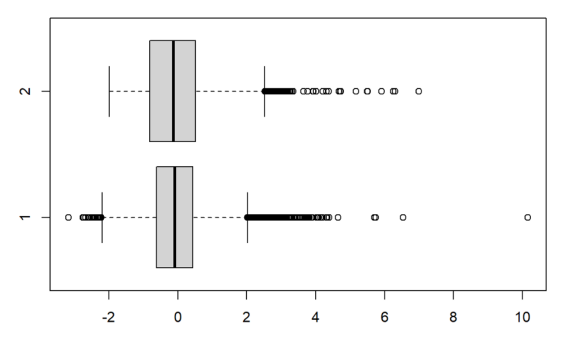
```
> head(homedata$y1970)
[1] 89700 118400 116400 122000 91500 102800
> mean(homedata$y1970)
[1] 70821
> sd(homedata$y1970)
[1] 22284
> head((homedata$y1970 - mean(homedata$y1970)) / sd(homedata$y1970))
[1] 0.847 2.135 2.045 2.297 0.928 1.435
> head(scale(homedata$y1970))
   [,1]
[1,] 0.847
[2,] 2.135
[3,] 2.045
[4,] 2.297
[5,] 0.928
[6,] 1.435
> head(homedata$y2000)
[1] 359100 504500 477300 500400 433900 464800
> mean(homedata$y2000)
[1] 268370
> sd(homedata$y2000)
[1] 130734
> head((homedata$y2000 - mean(homedata$y2000)) / sd(homedata$y2000))
[1] 0.694 1.806 1.598 1.775 1.266 1.503
> head(scale(homedata$y2000))
   [,1]
[1,] 0.694
[2,] 1.806
```

[3,] 1.598

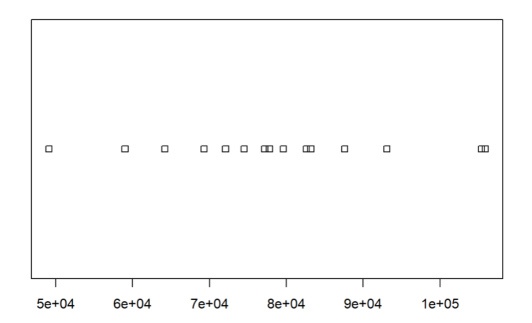
[4,] 1.775 [5,] 1.266

[6,] 1.503

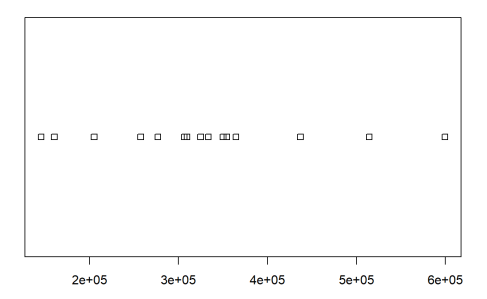
> boxplot(c(scale(homedata\$y1970)), c(scale(homedata\$y2000)), horizontal = TRUE)



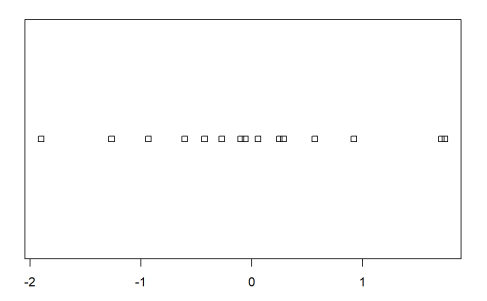
Also we can present the two distributions using stripcharts or dotplots > stripchart(home\$old)



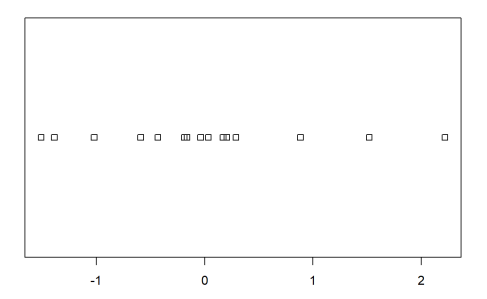
# > stripchart(home\$new)



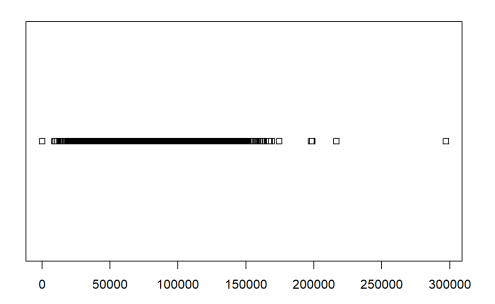
# > stripchart(scale(home\$old))



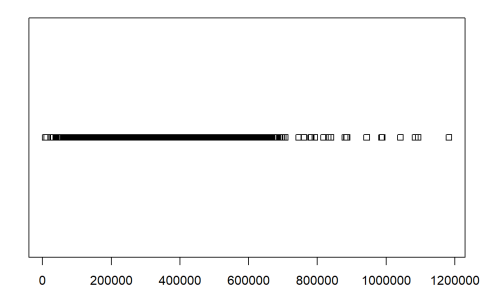
# > stripchart(scale(home\$new))



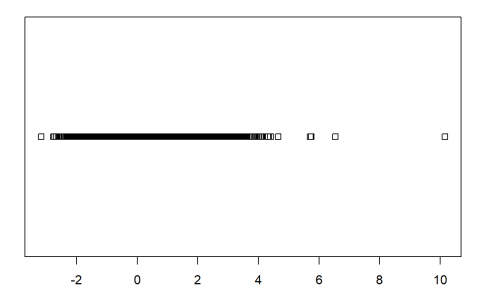
# > stripchart(homedata\$y1970)



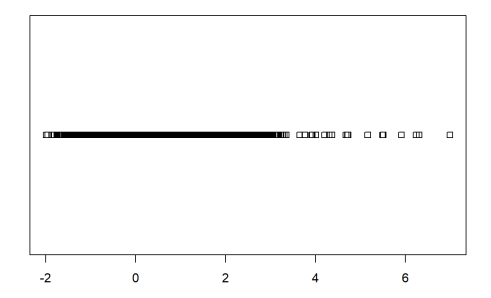
# > stripchart(homedata\$y2000)



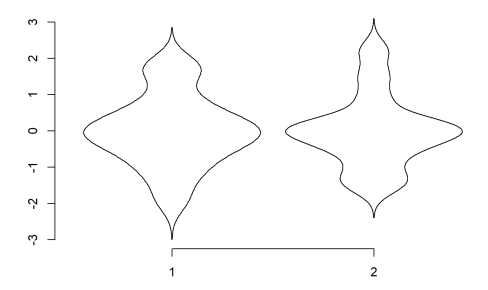
# > stripchart(scale(homedata\$y1970))



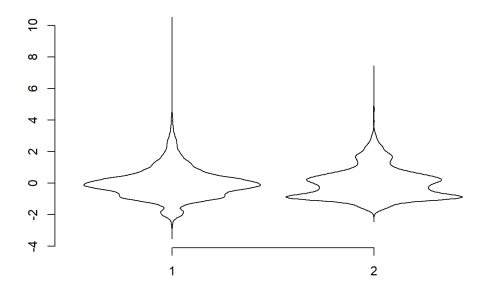
### > stripchart(scale(homedata\$y2000))



We can also compare shapes of distributions using the violin plots > violinplot(scale(home\$old), scale(home\$new)) > simple.violinplot(scale(home\$old),scale(home\$new))

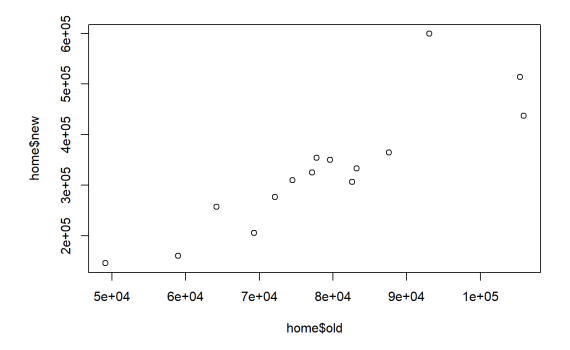


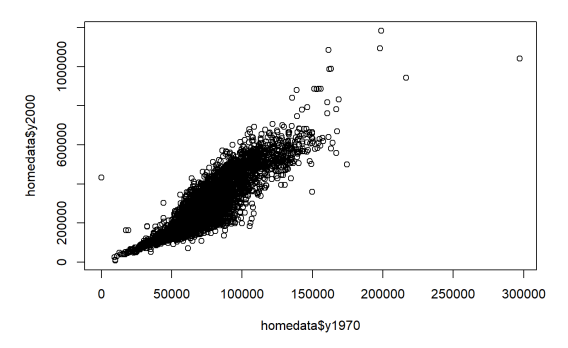
- > violinplot(scale(homedata\$y1970), scale(homedata\$y2000))
- > simple.violinplot(scale(homedata\$y1970),scale(homedata\$y2000))



### **Comparing relationships**

To investigate one numerical variable against another, we can use the scatter plot > plot(home\$old, home\$new)





It looks like we have a strong linear trend. The modeling of such relationships is a common statistical practice. It allows us to make predictions of the y variable based on the value of the x variable.

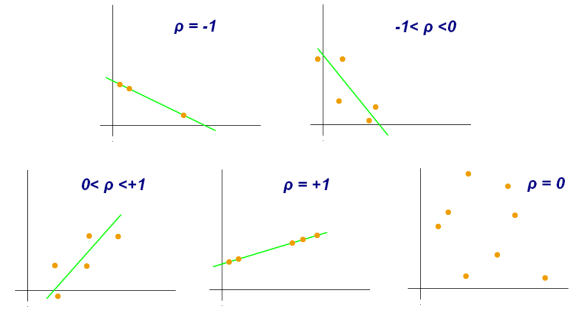
#### Pearson correlation coefficient R

The Pearson correlation coefficient indicates the strength of a linear relationship between two variables.

$$R = \frac{\sum (x_i - \overline{X})(y_i - \overline{Y})}{\sqrt{\sum (x_i - \overline{X})^2 \sum (y_i - \overline{Y})^2}} \in [-1, 1]$$

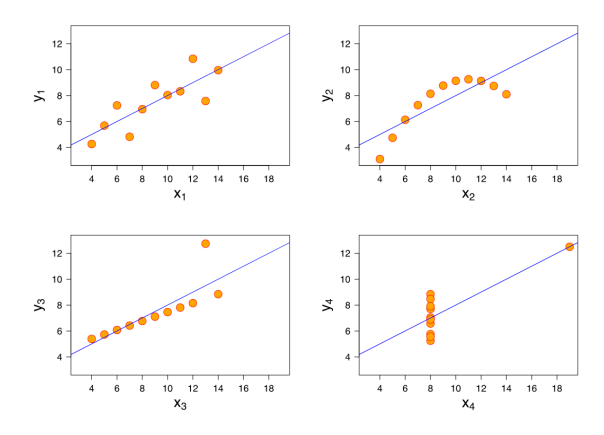
This is a scaled covariance between X and Y. It measures how one variable varies as the other does. A value of

- -1 indicates strong negative linear relationship
- 0 indicates no linear relationship
- +1 indicates strong positive linear relationship



Pearson correlation value generally does not completely characterize their relationship.

The following image displays four data sets called Anscombe's quartet. In each case, the mean and standard deviation of each variable is the same, and the correlation between the two random variables is 0.816. However, the data look very different when graphed.



# Time for some practice. Let's play some funny online games to predict correlations Game 1, Game 2

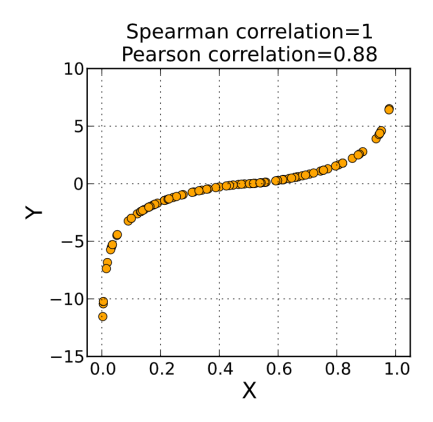
```
> cov(home$old, home$new)
[1] 1.67e+09
> cov(homedata$y1970, homedata$y2000)
[1] 2.61e+09
> cor(home$old, home$new)
[1] 0.881
> cor(homedata$y1970, homedata$y2000)
[1] 0.896
> cor(home$old, home$new)^2
[1] 0.776
> cor(homedata$y1970, homedata$y2000)^2
[1] 0.803
```

### Spearman rank correlation

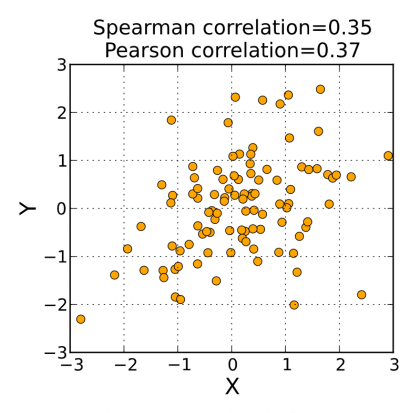
Same thing only applied to the ranks of the data.

```
> cor(rank(home$old), rank(home$new))
[1] 0.925
> cor(rank(homedata$y1970), rank(homedata$y2000))
[1] 0.888
```

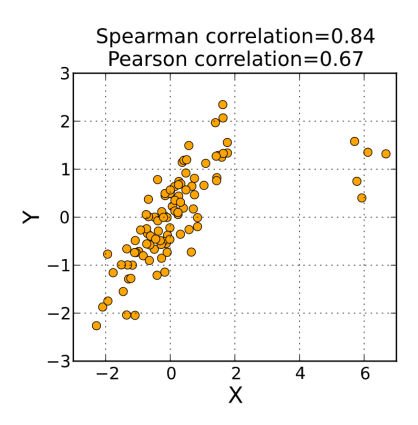
Spearman rank correlation is close to 1 (or -1) if there is a strong increasing (decreasing) trend in the data. (The trend need not be linear.)



When the data are roughly elliptically distributed and there are no prominent outliers, the Spearman correlation and Pearson correlation give similar values.



The Spearman correlation is less sensitive than the Pearson correlation to strong outliers that are in the tails of both samples.



### Simple linear regression

Linear regression is the name of a procedure that fits a straight line to the data. The idea is that the x value is something the experimenter controls, the y value one the experimenter measures. The line is used to predict the value of y for a known value of x. The variable x is the **predictor** variable and y the **response** variable.

Suppose we write the equation of the line as

$$\hat{y} = b_0 + b_1 x$$

Then, for each  $x_i$  the predicted value would be

$$\hat{y} = b_0 + b_1 x_i$$

But the measured value is  $y_i$ , the difference is called the residual and is simply

$$\varepsilon_i = y_i - \hat{y}_i$$

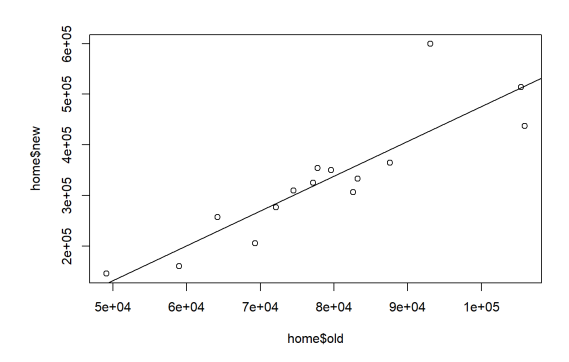
The method of least square is used to choose the values of  $b_0$  and  $b_1$  that minimize the sum or the squares of the residual errors  $\sum_i (y_i - \hat{y}_i)^2$ 

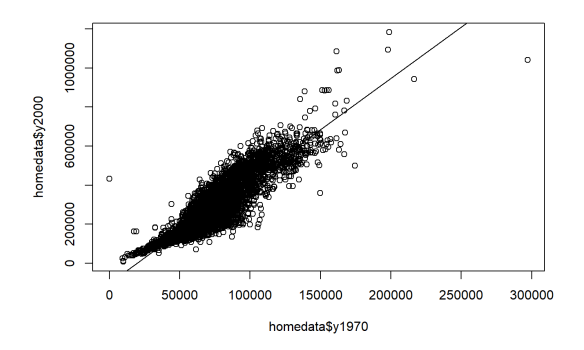
Solving, gives

$$b_0 = \overline{Y} - b_1 \overline{X}, \qquad b_1 = \frac{s_{xy}}{s_x^2} = \frac{\sum (x_i - \overline{X})(y_j - \overline{Y})}{\sum (x_i - \overline{X})^2}$$

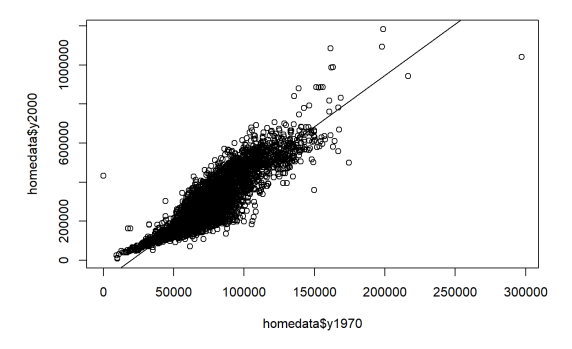
This is, a line with slope given by  $b_1$ .

- > plot(home\$old, home\$new)
- > abline(lm(home\$new ~ home\$old))





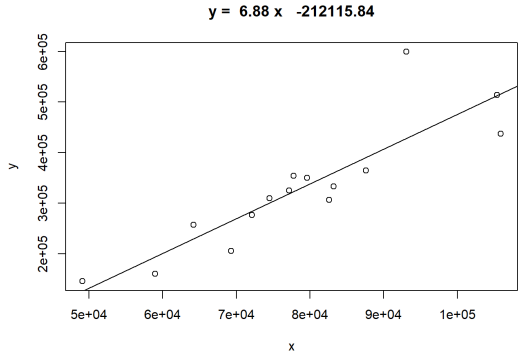
- > plot(homedata\$y2000 ~ homedata\$y1970)
- > abline(lm(homedata\$y2000 ~ homedata\$y1970))



abline function points lines on the current graph window Im function for the linear model

The syntax  $y \sim x$  tells R to model the y variable as a linear function of x simple. Im function will make the same plot and return the regression coefficients

### > simple.lm(home\$old, home\$new)



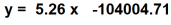
Call:  $Im(formula = y \sim x)$ 

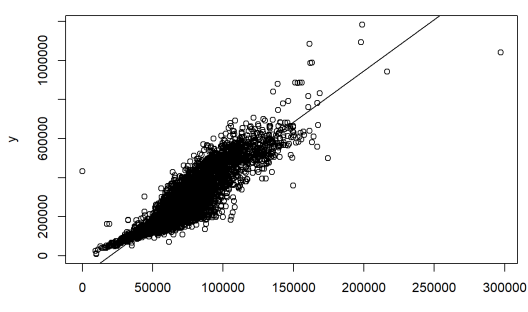
Coefficients:

(Intercept)

-2.12e+05 6.88e+00

> pl <- simple.lm(homedata\$y1970, homedata\$y2000)





X

### > attributes(pl)

\$names

- [1] "coefficients" "residuals" "effects" "rank"
- [5] "fitted.values" "assign" "qr" "df.residual"
- [9] "xlevels" "call" "terms" "model"

\$class

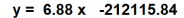
[1] "lm"

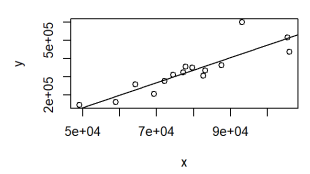
So we can directly access the coefficients, residuals and other parameters

### > pl\$coefficients

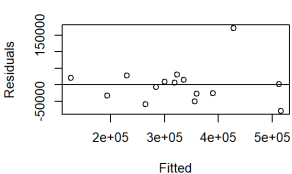
### Residual plot

> simple.lm(home\$old, home\$new, show.residuals = TRUE)

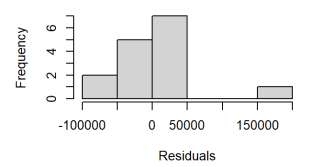




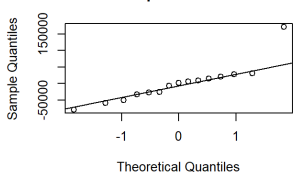
Residuals vs. fitted



hist of residuals



normal plot of residuals



Call:

 $Im(formula = y \sim x)$ 

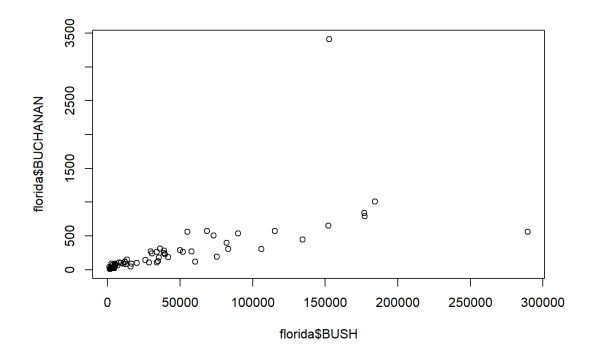
Coefficients:

(Intercept) x -2.12e+05 6.88e+00 The lower left is a histogram of the residuals. If the standard model is applicable, then this should appear "bell" shaped.

### **Locating points**

Consider the florida data frame containing results of year 2000 US presidential election in Florida. Let's review the BUSH and Buchanan votes.

```
> head(florida)
      County
                GORE
                          BUSH BUCHANAN NADER BROWN HAGELIN HARRIS
MCREYNOLDS
1 ALACHUA 47300 34062
                           262 3215 658
                                           42
                                                      658
                                                 4
   BAKER 2392 5610
                         73
                             53
                                 17
                                       3
                                           0
                                                  0
    BAY 18850 38637
                        248 828 171
                                             5
                                                   3
                                        18
4 BRADFORD 3072 5413
                           65
                                    28
                                          2
                                                    0
                              84
                                              0
5 BREVARD 97318 115185
                           570 4470 643
                                            39
                                                 11
                                                        11
6 BROWARD 386518 177279
                             789 7099 1212
                                             128
                                                   49
                                                          35
 MOOREHEAD PHILLIPS Total
     21
           20 86242
1
2
      3
           3 8154
3
     37
           18 58815
4
     3
           2 8669
5
     76
           72 218395
     123
           74 573306
> cor(florida$BUSH, florida$BUCHANAN)
[1] 0.624
> cor(rank(florida$BUSH), rank(florida$BUCHANAN))
[1] 0.944
> plot(florida$BUSH, florida$BUCHANAN)
```



### > simple.lm(florida\$BUSH, florida\$BUCHANAN)

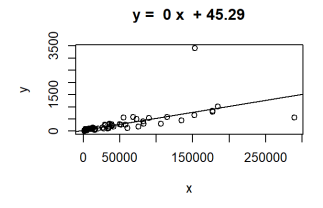
Call:

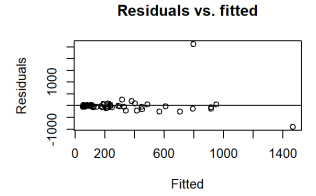
 $Im(formula = y \sim x)$ 

Coefficients:

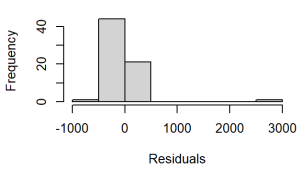
(Intercept) x 45.28986 0.00492

### > simple.lm(florida\$BUSH, florida\$BUCHANAN, show.residuals = TRUE)

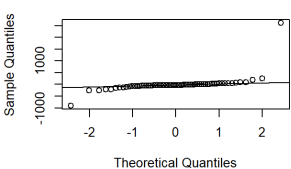




## hist of residuals



### normal plot of residuals



Call:  $Im(formula = y \sim x)$ 

Coefficients:

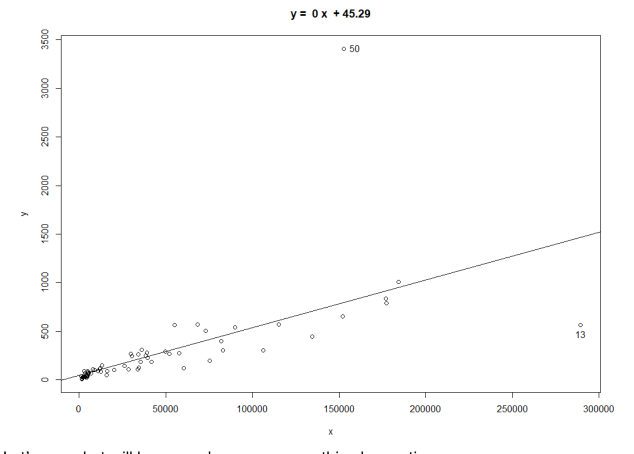
(Intercept) x

45.28986 0.00492

We see a strong linear relationship, except for two "outliers". How can we identify these points? identify find the index of the closest (x, y) coordinates to the mouse click.

- > simple.lm(florida\$BUSH, florida\$BUCHANAN)
- > identify(florida\$BUSH, florida\$BUCHANAN)

Click on the two outliers and then press 'Esc' to finish and print the identified coordinates.



Let's see what will happen when we remove this observations. > florida[c(13,50),]

County GORE BUSH BUCHANAN NADER BROWN HAGELIN HARRIS MCREYNOLDS

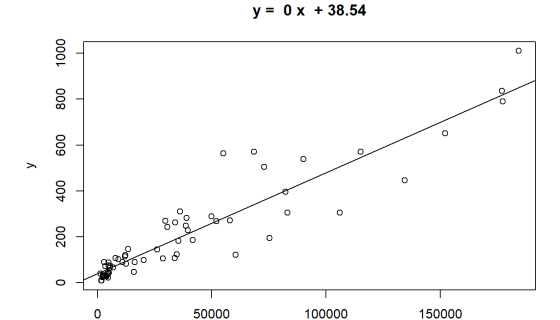
13 DADE 328702 289456 561 5355 759 119 88 36

50 PALM BEACH 268945 152846 3407 5564 743 143 45 302

MOOREHEAD PHILLIPS Total

13 124 69 625269 50 103 188 432286

- > florida.cleaned <- florida[-c(13, 50), ]
- > linearmodel <- simple.lm(florida.cleaned\$BUSH, florida.cleaned\$BUCHANAN)



> linearmodel\$coefficients[1]

(Intercept) 38.5

> linearmodel\$coefficients[2]

X

0.0044

- > bush.palm.beach <- 152846
- > buchanan.palm.beach <- linearmodel\$coefficients[1] + linearmodel\$coefficients[2] \* bush.palm.beach

X

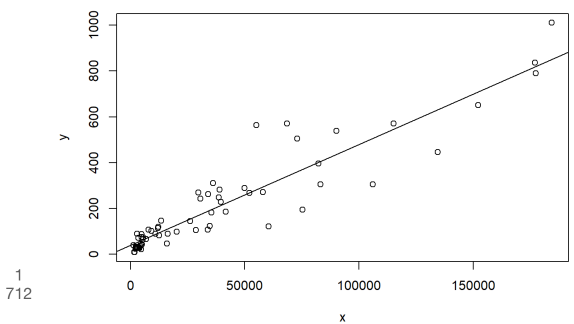
> buchanan.palm.beach

(Intercept)

712

> simple.lm(florida.cleaned\$BUSH, florida.cleaned\$BUCHANAN, pred =
florida\$BUSH[50])

$$y = 0 x + 38.54$$



Call:

 $Im(formula = y \sim x)$ 

Coefficients:

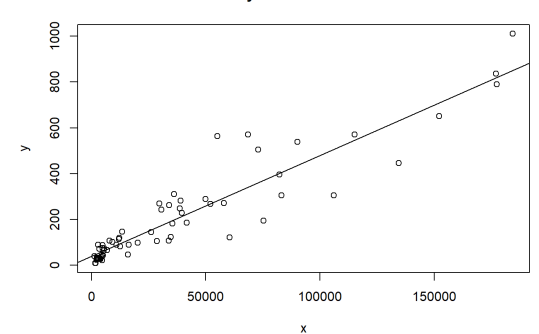
(Intercept) x 38.5363 0.0044

- > bush.dade <- 289456
- > buchanan.dade <- linearmodel\$coefficients[1] + linearmodel\$coefficients[2] \* bush.dade
- > buchanan.dade

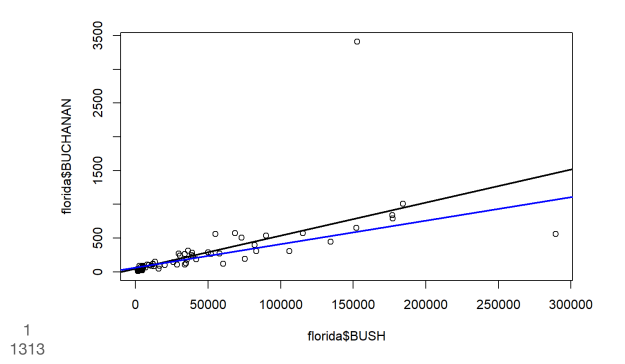
(Intercept)

1313

$$y = 0 x + 38.54$$



# > simple.lm(florida.cleaned\$BUSH, florida.cleaned\$BUCHANAN, pred = florida\$BUSH[13])



Call:

 $Im(formula = y \sim x)$ 

Coefficients:

(Intercept) x 38.5363 0.0044

We expect Buchanan to have received 712 votes in Palm Beach and 1313 in Dade, not the actual received.

- > plot(florida\$BUSH, florida\$BUCHANAN)
- > abline(lm(florida\$BUCHANAN ~ florida\$BUSH), lwd = 2)
- > abline(65.6, 0.00348, col = "Blue", lwd = 2)