# Date Science and Applications with R Statistical models

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# Agenda

- Using data frames for statistical purposes
- Manipulation of data into more convenient forms
- (Re-)Introduction to linear models and the model space

So You've Got A Data Frame. What can we do with it?

- Plot it: examine multiple variables and distributions
- Test it: compare groups of individuals to each other
- Check it: does it conform to what we'd like for our needs?

# Test Case: Birth weight data

## Included in R already:

```
library(tidyverse)
library(lubridate)
library(MASS)
data(birthwt)
summary(birthwt)
```

```
##
         low
                                            lwt
                           age
                                                             race
    Min.
           :0.0000
                      Min.
                              :14.00
                                                               :1.000
##
                                       Min.
                                              : 80.0
                                                        Min.
##
    1st Qu.:0.0000
                      1st Qu.:19.00
                                       1st Qu.:110.0
                                                        1st Qu.:1.000
##
    Median :0.0000
                      Median :23.00
                                       Median :121.0
                                                        Median :1.000
##
    Mean
           :0.3122
                      Mean
                             :23.24
                                       Mean
                                              :129.8
                                                        Mean
                                                               :1.847
    3rd Ou.:1.0000
                      3rd Qu.:26.00
                                       3rd Qu.:140.0
                                                        3rd Ou.:3.000
##
                                              :250.0
##
    Max.
           :1,0000
                      Max.
                             :45.00
                                       Max.
                                                        Max.
                                                               :3.000
##
        smoke
                           ptl
                                              ht
                                                                 ui
                                                                   :0.0000
##
    Min.
           :0.0000
                      Min.
                              :0.0000
                                        Min.
                                               :0.00000
                                                           Min.
                                        1st Qu.:0.00000
                                                           1st Qu.:0.0000
##
    1st Qu.:0.0000
                      1st Qu.:0.0000
    Median :0.0000
                                        Median :0.00000
                                                           Median :0.0000
##
                      Median :0.0000
##
    Mean
           :0.3915
                      Mean
                             :0.1958
                                        Mean
                                               :0.06349
                                                           Mean
                                                                  :0.1481
##
    3rd Qu.:1.0000
                      3rd Qu.:0.0000
                                        3rd Qu.:0.00000
                                                           3rd Qu.:0.0000
                             :3.0000
##
           :1.0000
                                        Max. :1.00000
                                                           Max.
                                                                  :1.0000
    Max.
                      Max.
         ftv
                                                                          3/83
##
                           bwt
```

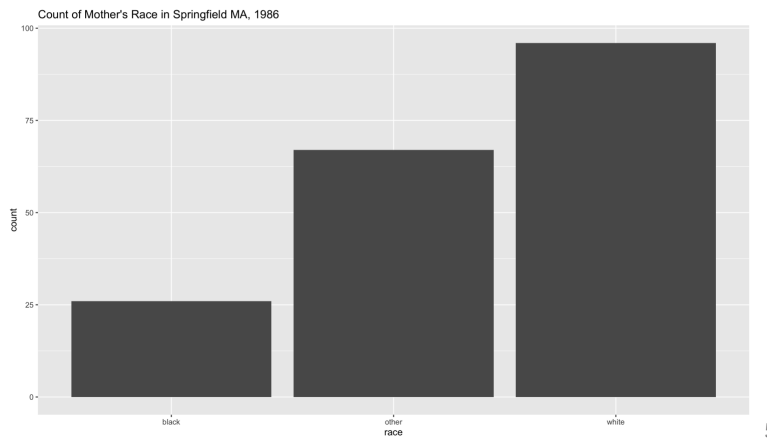
#### Make it readable!

## Let's make all the factors more descriptive.

```
birthwt$race <- factor(c("white", "black", "other")[birthwt$race])
birthwt$mother.smokes <- factor(c("No", "Yes")[birthwt$mother.smokes
birthwt$uterine.irr <- factor(c("No", "Yes")[birthwt$uterine.irr + 1]
birthwt$hypertension <- factor(c("No", "Yes")[birthwt$hypertension +</pre>
```

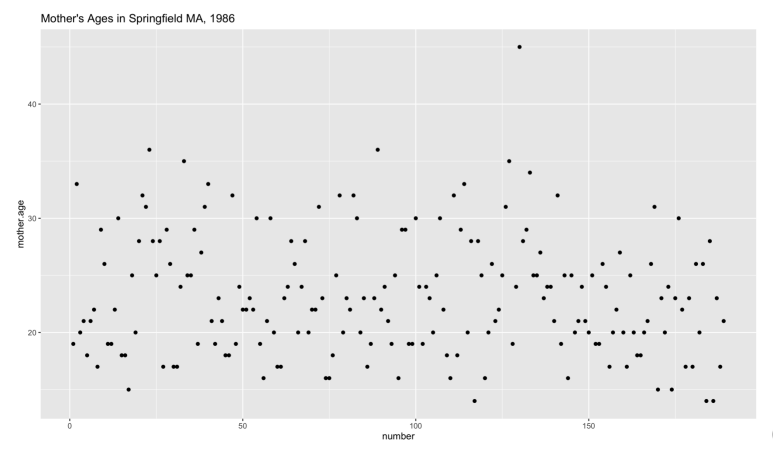
# Bar plot for race

```
birthwt %>% ggplot(aes(x = race))+
  geom_bar()+
  labs(title = "Count of Mother's Race in Springfield MA, 1986")
```



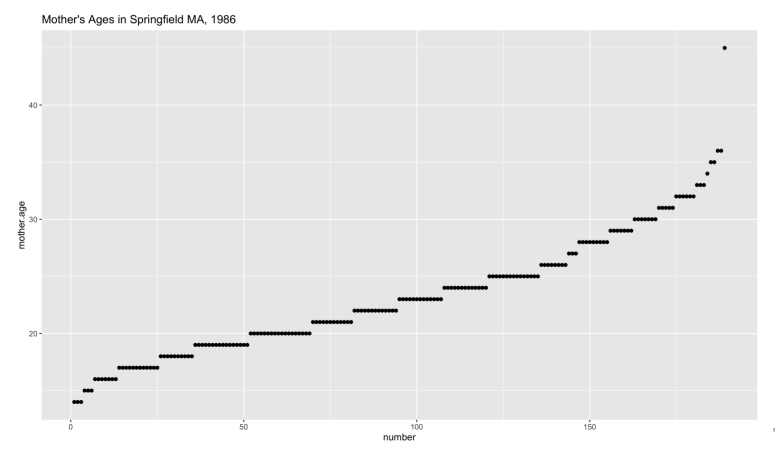
# Scatter plot for mother's ages

```
birthwt %>% ggplot(aes(x = 1:nrow(birthwt), y = mother.age))+
  geom_point()+
  labs(x = 'number', title = "Mother's Ages in Springfield MA, 1986")
```



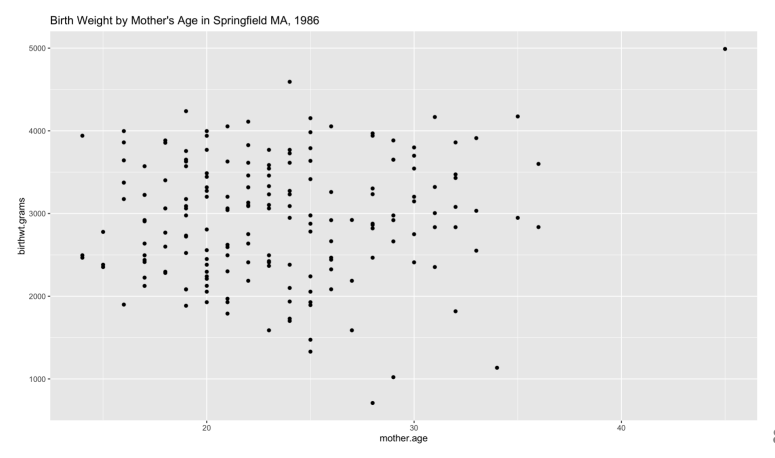
# Sorted mother's ages

```
birthwt %>% arrange(mother.age) %>% ggplot(aes(x = 1:nrow(birthwt), y
geom_point()+
labs(x = 'number', title = "Mother's Ages in Springfield MA, 1986")
```



# Birth weight versus mother's ages

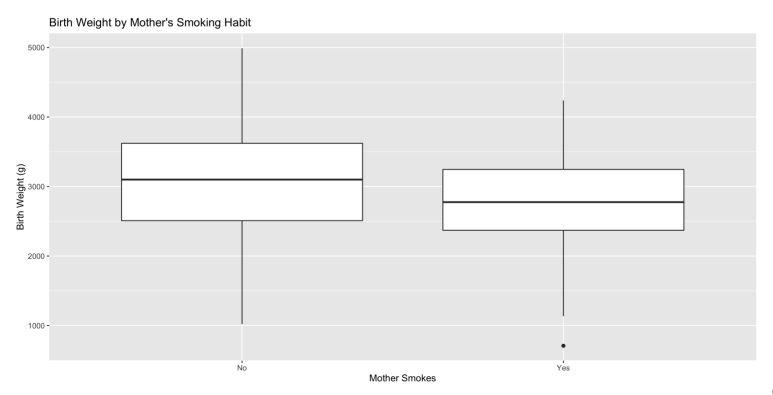
```
birthwt %>% ggplot(aes(x = mother.age, y = birthwt.grams))+
  geom_point()+
  labs(title = "Birth Weight by Mother's Age in Springfield MA, 1986")
```



# **Boxplot**

Let's fit some models to the data pertaining to our outcome(s) of interest.

```
birthwt %>% ggplot(aes(x = mother.smokes, y = birthwt.grams))+
  geom_boxplot()+
  labs(title = "Birth Weight by Mother's Smoking Habit", y = "Birth Weight")
```



# Basic statistical testing

Tough to tell! Simple two-sample t-test:

#### Does this difference match the linear model?

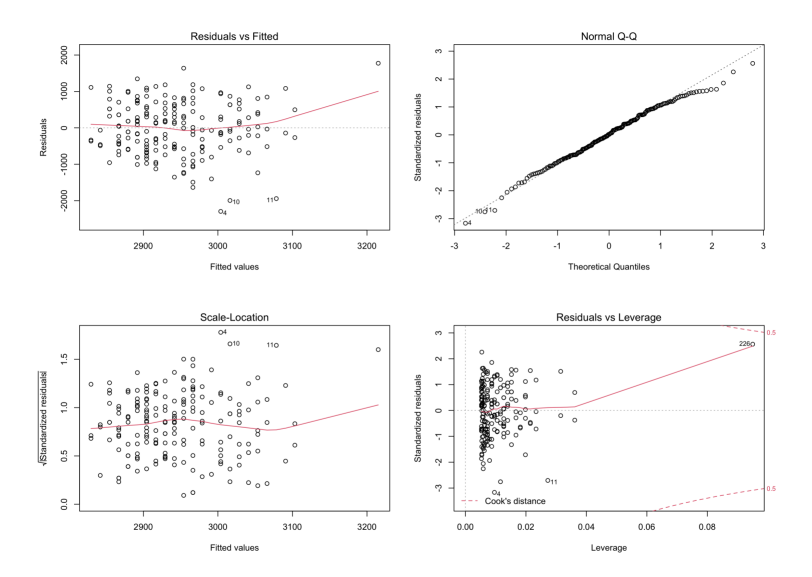
```
linear.model.1 <- lm (birthwt.grams ~ mother.smokes, data=birthwt)</pre>
summary(linear.model.1)
##
## Call:
## lm(formula = birthwt.grams ~ mother.smokes, data = birthwt)
##
## Residuals:
##
      Min 10 Median 30
                                    Max
## -2062.9 -475.9 34.3 545.1 1934.3
##
## Coefficients:
##
                  Estimate Std. Error t value Pr(>|t|)
## (Intercept) 3055.70 66.93 45.653 < 2e-16 ***
## mother.smokesYes -283.78 106.97 -2.653 0.00867 **
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 717.8 on 187 degrees of freedom
## Multiple R-squared: 0.03627, Adjusted R-squared: 0.03112
```

## F-statistic: 7.038 on 1 and 187 DF, p-value: 0.008667

# Basic statistical testing

Does this difference match the linear model?

```
linear.model.2 <- lm (birthwt.grams ~ mother.age, data=birthwt)</pre>
summary(linear.model.2)
##
## Call:
## lm(formula = birthwt.grams ~ mother.age, data = birthwt)
##
## Residuals:
##
       Min
            1Q Median
                                 30
                                         Max
## -2294.78 -517.63 10.51 530.80 1774.92
##
## Coefficients:
              Estimate Std. Error t value Pr(>|t|)
##
## (Intercept) 2655.74 238.86
                                  11.12 <2e-16 ***
## mother.age 12.43 10.02 1.24 0.216
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 728.2 on 187 degrees of freedom
## Multiple R-squared: 0.008157, Adjusted R-squared: 0.002853
## F-statistic: 1.538 on 1 and 187 DF, p-value: 0.2165
```



# **Detecting Outliers**

##

These are the default diagnostic plots for the analysis. Note that our oldest mother and her heaviest child are greatly skewing this analysis as we suspected.

```
birthwt.noout <- birthwt %>% filter(mother.age <= 40)</pre>
linear.model.3 <- lm (birthwt.grams ~ mother.age, data=birthwt.noout)</pre>
summary(linear.model.3)
##
## Call:
## lm(formula = birthwt.grams ~ mother.age, data = birthwt.noout)
##
## Residuals:
                 1Q Median
##
       Min
                                  30
                                          Max
## -2245.89 -511.24 26.45 540.09 1655.48
##
## Coefficients:
##
              Estimate Std. Error t value Pr(>|t|)
                                   11.57 <2e-16 ***
## (Intercept) 2833.273 244.954
## mother.age 4.344 10.349 0.42 0.675
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

## More complex models

## Add in smoking behavior:

```
linear.model.3a <- lm (birthwt.grams ~ + mother.smokes + mother.age,
summary(linear.model.3a)
##
## Call:
## lm(formula = birthwt.grams ~ +mother.smokes + mother.age, data = birthwt.r
##
## Residuals:
      Min
               10 Median
##
                              30
                                     Max
## -2081.22 -459.82 43.56 548.22 1551.51
##
## Coefficients:
##
                Estimate Std. Error t value Pr(>|t|)
## (Intercept) 2954.582 246.280 11.997 <2e-16 ***
## mother.age 3.621 10.208 0.355 0.7232
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 707.1 on 185 degrees of freedom
## Multiple R-squared: 0.03401, Adjusted R-squared: 0.02357
```

## F-statistic: 3.257 on 2 and 185 DF, p-value: 0.04072

# More complex models

## Add in smoking behavior:

```
linear.model.3b <- lm (birthwt.grams ~ mother.age + mother.smokes*rac
summary(linear.model.3b)
##
## Call:
## lm(formula = birthwt.grams ~ mother.age + mother.smokes * race,
##
      data = birthwt.noout)
##
## Residuals:
       Min
                 10 Median
##
                                   30
                                           Max
## -2343.52 -413.66 39.91 480.36
                                      1379.90
##
## Coefficients:
##
                             Estimate Std. Error t value Pr(>|t|)
                             3017.352
  (Intercept)
                                         265.606 11.360 < 2e-16 ***
## mother.age
                               -8.168
                                          10.276 -0.795 0.42772
## mother.smokesYes
                             -316.500
                                         275.896 -1.147 0.25282
## raceother
                              -18.901
                                         193.665 -0.098 0.92236
## racewhite
                              584.042
                                         206.320 2.831 0.00517 **
## mother.smokesYes:raceother 258.999
                                         349.871 0.740 0.46010
## mother.smokesYes:racewhite -271.594
                                         314.268
                                                  -0.864
                                                         0.38862
                                                                   17 / 83
```

# Everything Must Go (In)

Let's do a kitchen sink model on this new data set:

```
linear.model.4 <- lm (birthwt.grams ~ ., data=birthwt.noout)</pre>
summary(linear.model.4)
##
## Call:
## lm(formula = birthwt.grams ~ ., data = birthwt.noout)
##
## Residuals:
          1Q Median
##
      Min
                             30
                                    Max
## -985.04 -274.13 -13.87 262.53 1146.50
##
## Coefficients:
                       Estimate Std. Error t value Pr(>|t|)
##
## (Intercept)
                      3360.5163
                                 215,4112
                                         15.600 < 2e-16 ***
## birthwt.below.2500 -1116.3933 70.8578 -15.755 < 2e-16 ***
## mother.age
                -16.0321 6.4159 -2.499 0.013373 *
## mother.weight
                        1.9317
                                   1.1208 1.723 0.086545 .
## raceother
               68.8145 101.4451 0.678 0.498441
## racewhite
            247.0241 96.4935 2.560 0.011302 *
## mother.smokesYes -157.7041 68.6205 -2.298 0.022719 *
## previous.prem.labor
                        95.9825
                                  65.3329 1.469 0.143573
```

# Everything Must Go (In), Except What Must Not

Whoops! One of those variables was birthwt.below.2500 which is a function of the outcome.

```
linear.model.4a <- lm (birthwt.grams ~ . - birthwt.below.2500, data=k
summary(linear.model.4a)
##
## Call:
## lm(formula = birthwt.grams ~ . - birthwt.below.2500, data = birthwt.noout)
##
## Residuals:
                10 Median
##
       Min
                                 30
                                         Max
## -1761.10 -454.81 46.43 459.78 1394.13
##
## Coefficients:
                     Estimate Std. Error t value Pr(>|t|)
##
## (Intercept)
                     2545,584
                                323.204 7.876 3.21e-13 ***
## mother.age
                                  9.909 -1.222 0.223243
                      -12.111
## mother.weight
                                  1.710 2.801 0.005656 **
                       4.789
## raceother
                      155.605
                                156.564 0.994 0.321634
## racewhite
             494.545
                                147.153 3.361 0.000951 ***
## mother.smokesYes -335.793
                                104.613 -3.210 0.001576 **
## previous.prem.labor
                      -32,922
                                         -0.329 0.742838
                                                                20 / 83
                                100.185
```

## **Generalized Linear Models**

Maybe a linear increase in birth weight is less important than if it's below a threshold like 2500 grams (5.5 pounds). Let's fit a generalized linear model instead:

```
glm.0 <- glm (birthwt.below.2500 ~ . - birthwt.grams, data=birthwt.nc
```

## **Generalized Linear Models**

The default value is a Gaussian model (a standard linear model). Change this:

```
glm.1 <- glm (birthwt.below.2500 ~ . - birthwt.grams, data=birthwt.nc
summary(glm.1)
##
## Call:
## glm(formula = birthwt.below.2500 ~ . - birthwt.grams, family = binomial(li
      data = birthwt.noout)
##
##
## Deviance Residuals:
      Min
               1Q Median
##
                                30
                                       Max
## -1.8938 -0.8222 -0.5363 0.9848
                                    2.2069
##
## Coefficients:
                      Estimate Std. Error z value Pr(>|z|)
##
                     1.721830 1.258897 1.368 0.17140
## (Intercept)
## mother.age
                   -0.027537 0.037718 -0.730 0.46534
## mother.weight -0.015474 0.006919 -2.237 0.02532 *
## raceother
             -0.395505 0.537685 -0.736 0.46199
## racewhite
            -1.269006 0.527180 -2.407 0.01608 *
## mother.smokesYes 0.931733 0.402359 2.316 0.02058 *
## previous.prem.labor
                     0.539549
                                0.345413 1.562
                                                0.11828
                                                               24 / 83
```

# What Do We Do With This, Anyway?

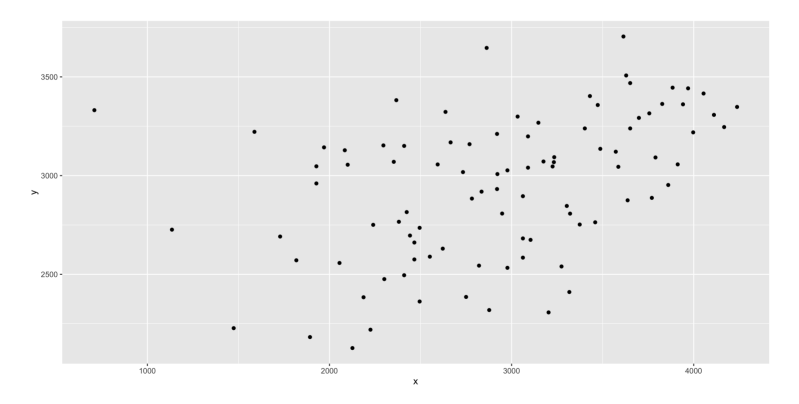
Let's take a subset of this data to do predictions.

```
odds <- seq(1, nrow(birthwt.noout), by=2)
birthwt.in <- birthwt.noout[odds,]</pre>
birthwt.out <- birthwt.noout[-odds,]</pre>
linear.model.half <- lm (birthwt.grams ~ . - birthwt.below.2500, data
summary (linear.model.half)
##
## Call:
## lm(formula = birthwt.grams ~ . - birthwt.below.2500, data = birthwt.in)
##
## Residuals:
       Min
                 10 Median
##
                                  30
                                          Max
## -1705.17 -303.11 26.48 427.18 1261.57
##
## Coefficients:
                      Estimate Std. Error t value Pr(>|t|)
##
## (Intercept)
                      2514.891
                                  450.245 5.586 2.81e-07 ***
## mother.age
                        7.052
                                  14.935 0.472 0.63801
## mother.weight
                    2.683 2.885 0.930 0.35501
## raceother
                       113.948 224.519 0.508 0.61312
## racewhite
                       466.219
                                  204.967 2.275 0.02548 *
                                                                   26 / 83
```

```
birthwt.predict <- predict (linear.model.half)
cor (birthwt.in$birthwt.grams, birthwt.predict)</pre>
```

#### ## [1] 0.508442

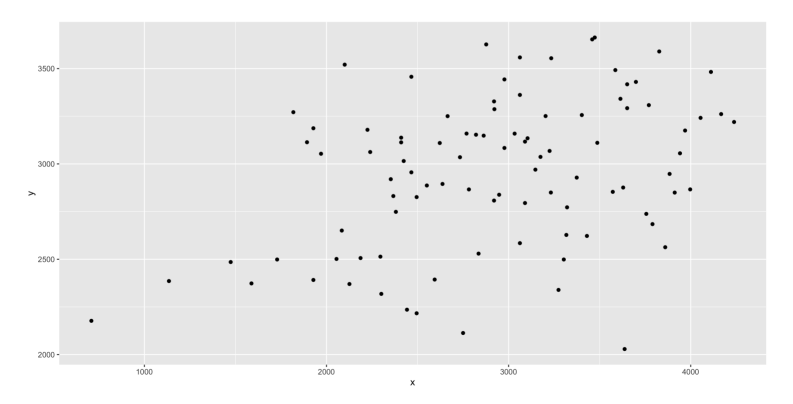
```
tibble(x = birthwt.out$birthwt.grams, y = birthwt.predict) %>%
  ggplot (aes(x = x, y = y)) + geom_point()
```



# What Do We Do With This, Anyway?

birthwt.predict.out <- predict (linear.model.half, birthwt.out)
cor (birthwt.out\$birthwt.grams, birthwt.predict.out)</pre>

## [1] 0.3749431



# Random number generators

- We made reference to random number generation without going under the hood.
- How *does* R get "random" numbers?
- It doesn't, really -- it uses a trick that should be indistinguishable from the real McCoy

Pseudorandom generators produce a deterministic sequence that is indistiguishable from a true random sequence if you don't know how it started.

## Example: runif, where we know where it started

```
runif(1:10)

## [1] 0.9183728 0.7319108 0.8286768 0.9801534 0.4271460 0.6820300 0.0821624
## [8] 0.8520019 0.2604056 0.2969868

set.seed(10)
runif(1:10)
```

[1] 0.50747820 0.30676851 0.42690767 0.69310208 0.08513597 0.2254**36**6/283

# Basic version: Linear Congruential Generator

```
seed <- 10
new.random <- function (a=5, c=12, m=16) {
  out <- (a*seed + c) %% m
  seed <<- out
  return(out)
}
out.length <- 20
variates <- rep (NA, out.length)
for (kk in 1:out.length) variates[kk] <- new.random()
variates</pre>
```

## [1] 14 2 6 10 14 2 6 10 14 2 6 10 14 2 6 10 14 2 6 10

# Try again

### Period 8:

```
variates <- rep (NA, out.length)
for (kk in 1:out.length) variates[kk] <- new.random(a=131, c=7, m=16)
variates
## [1] 5 6 9 2 13 14 1 10 5 6 9 2 13 14 1 10 5 6 9 2</pre>
```

# Try again, again

## Period 16:

```
variates <- rep (NA, out.length)
for (kk in 1:out.length) variates[kk] <- new.random(a=129, c=7, m=16)
variates
## [1] 9 0 7 14 5 12 3 10 1 8 15 6 13 4 11 2 9 0 7 14</pre>
```

# Try again, at last

## Numerical Recipes uses

```
variates <- rep (NA, out.length)
for (kk in 1:out.length) variates[kk] <- new.random(a=1664545, c=1013)
variates

## [1] 1037207853 2090831916 4106096907 768378826 3835752553 1329121000
## [7] 2125006663 2668506502 3581687205 2079234980 2067291011 2197025090
## [13] 3748878561 2913996384 758844863 4029469438 2836748829 1458315036
## [19] 2399149563 2766656186</pre>
```

# How To Distinguish Non-Randomness

- We've covered period: if it's missing some values, it's distinguishable
- Uniformity of distribution in the limitx
- Autocorrelation
- Dimensional distribution -- not a problem for 1-D distributions, but can be for 2+-D

# How does R get everything we need?

## A few distributions of interest:

- Uniform(0,1)
- Bernoulli(p)
- Binomial(n,p)
- Gaussian(0,1)
- Exponential(1)
- Gamma(a)

# In R: everything we need

Suppose we were working with the Exponential distribution.

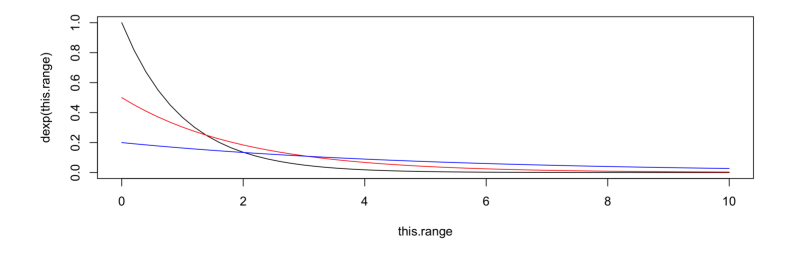
- rexp() generates variates from the distribution.
- dexp() gives the probability density function.
- pexp() gives the cumulative distribution function.
- qexp() gives the quantiles.

#### dexp()

```
dexp(0:5)
```

## [1] 1.000000000 0.367879441 0.135335283 0.049787068 0.018315639 0.00673794

```
this.range <- 0:50/5
plot (this.range, dexp(this.range), ty="l")
lines (this.range, dexp(this.range, rate=0.5), col="red")
lines (this.range, dexp(this.range, rate=0.2), col="blue")</pre>
```

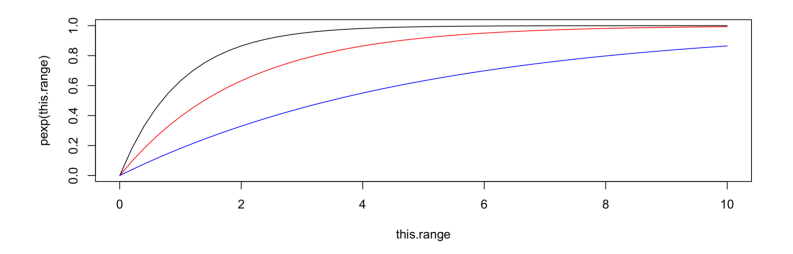


#### pexp()

```
pexp(0:5)
```

## [1] 0.0000000 0.6321206 0.8646647 0.9502129 0.9816844 0.9932621

```
this.range <- 0:50/5
plot (this.range, pexp(this.range), ty="l")
lines (this.range, pexp(this.range, rate=0.5), col="red")
lines (this.range, pexp(this.range, rate=0.2), col="blue")</pre>
```



#### qexp()

```
qexp(0:5)

## Warning in qexp(0:5): 产生了NaNs

## [1] 0 Inf NaN NaN NaN NaN

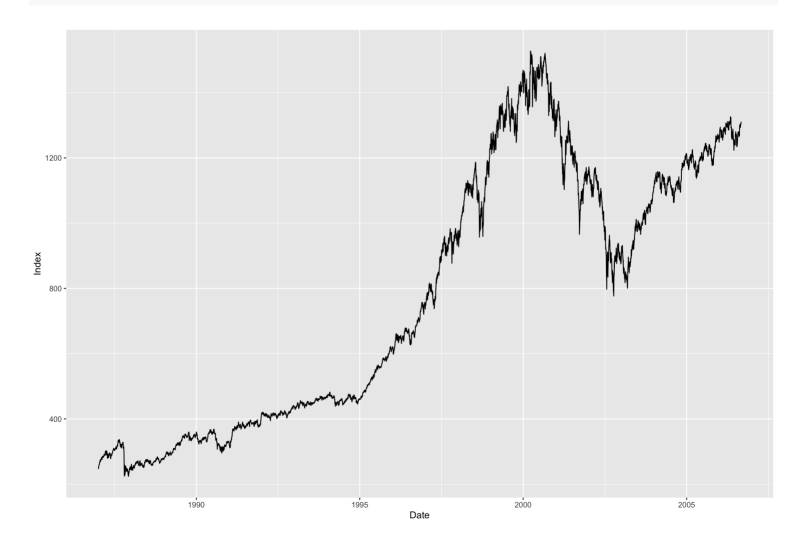
this.range <- seq(0,1,by=0.01)
plot (this.range, qexp(this.range), ylim = c(0, 10), ty="l")
lines (this.range, qexp(this.range, rate=0.5), col="red")
lines (this.range, qexp(this.range, rate=0.2), col="blue")</pre>
```

#### Let's Grab some Data

The Standard and Poor's 500, or simply the S\&P 500, is a stock market index tracking the stock performance of 500 large companies listed on exchanges in the United States. It is one of the most commonly followed equity indices.

```
library(readxl)
SP <- read_excel("data/Stock_Bond.xls") %>% dplyr::select(Date, `S&P_
rename(Index = `S&P_AC`)
```

SP %>% ggplot(aes(x = Date, y = Index)) + geom\_line()



## Let's Transform Some Data

The price  $p_t$  doesn't matter, what matters are the returns  $r_t = \log \left( p_t / p_{t-1} 
ight)$ 

```
returns <- na.omit(as.vector(diff(log(SP$Index))))
summary(returns)

## Min. 1st Qu. Median Mean 3rd Qu. Max.
## -0.2289972 -0.0046537 0.0004976 0.0003368 0.0056195 0.0870888

plot(returns, type="l")</pre>
```

### The Data's Distribution

quantile (x, probs) calculates the quantiles at probs from x

```
quantile(returns,c(0.25,0.5,0.75))

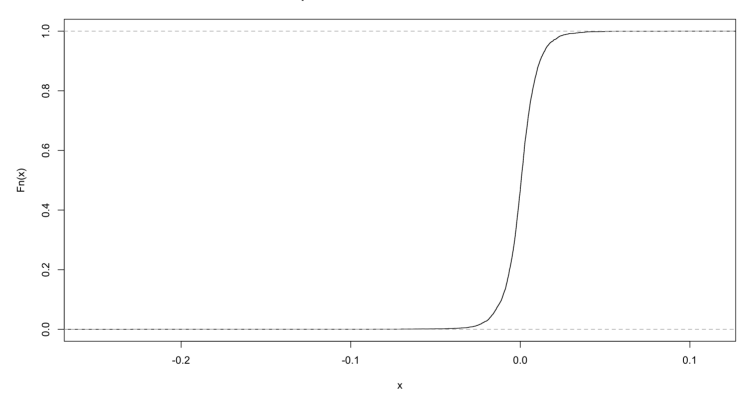
## 25% 50% 75%
## -0.0046537538 0.0004976042 0.0056195438
```

ecdf() - e mpirical c umulative d istribution f unction; no assumptions but also no guess about distribution between the observations

In math, ECDF is often written as  $\widehat{F}$  or  $\widehat{F}_n$ 

plot(ecdf(returns), main="Empirical CDF of S&P 500 index returns")

#### **Empirical CDF of S&P 500 index returns**



Conceptually, quantile and ecdf are inverses to each other

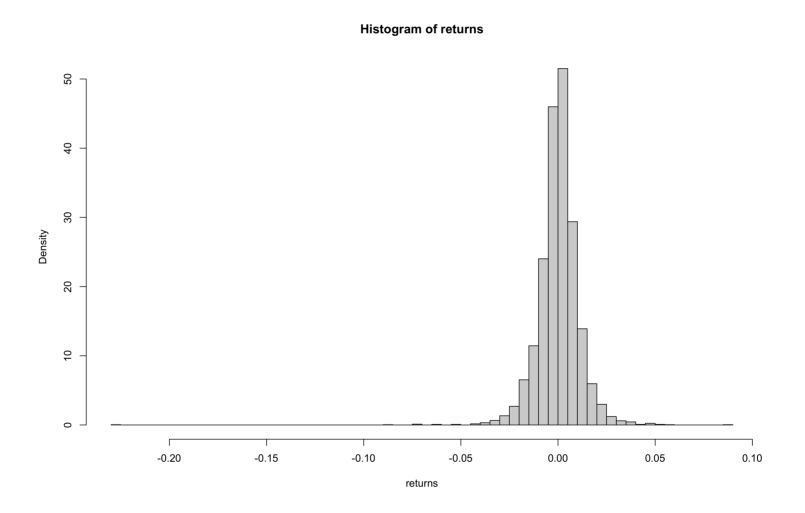
## Getting Probability Densities from Data

hist(x) calculates a histogram from x

- divide the data range up into equal-width bins and *count* how many fall into each bin
- Or divide bin counts by (total count)\*(width of bin), and get an estimate of the probability density function (pdf)

  Produces plot as a default side-effect

### hist(returns,n=101,probability=TRUE)



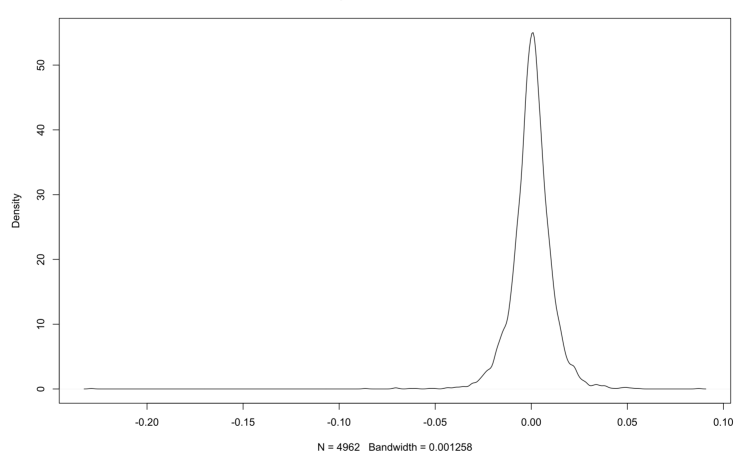
# Probability Densities from Data (cont'd.)

density (x) estimates the density of x by counting how many observations fall in a little window around each point, and then smoothing "Bandwidth"  $\approx$  width of window around each point Technically, a "kernel density estimate"

Remember: density() is an *estimate* of the pdf, not The Truth

density returns a collection of x, y values, suitable for plotting

#### Estimated pdf of S&P 500 index returns



# Probability Densities from Data (cont'd.)

```
hist(returns,n=101,probability=TRUE)
lines(density(returns),lty="dashed")
```

# Getting distributions from data (cont'd.)

table() - tabulate outcomes, most useful for discrete spaces; remember to normalize if you want probabilities

```
plot(table(cats$Sex)/nrow(cats),ylab="probability")
```

### Who Cares About the Distribution of the Data?

- Overly detailed: every single observation recorded as a separate tick
  - Too much information
- The exact set of samples would never repeat if we re-ran things anyway
  - That information is wrong
- Try to *summarize* what will *generalize* to other situations
  - Use a model, remember the model's parameters

### R commands for distributions

- d*foo* = the probability *d* ensity (if continuous) or probability mass function of *foo* (pdf or pmf)
- pfoo = the cumulative *p* robability function (CDF)
- qfoo = the q uantile function (inverse to CDF)
- *rfoo* = draw *r* andom numbers from foo (first argument always the number of draws)

?Distributions to see which distributions are built in

If you write your own, follow the conventions

# **Examples**

```
dnorm(x=c(-1,0,1),mean=1,sd=0.1)
## [1] 5.520948e-87 7.694599e-22 3.989423e+00
pnorm(q=c(2,-2)) # defaults to mean=0,sd=1
## [1] 0.97724987 0.02275013
dbinom(5,size=7,p=0.7,log=TRUE)
## [1] -1.146798
qchisq(p=0.95,df=5)
## [1] 11.0705
rt(n=4,df=2)
## [1] 0.4397369 1.1463390 -0.3783058 -1.3546069
```

# Displaying Probability Distributions

curve is very useful for the d, p, q functions:

```
curve(dgamma(x,shape=45,scale=1.9),from=0,to=200)
```

### How Do We Fit Distributional Models to the Data?

- Match moments (mean, variance, etc.)
- Match other summary statistics
- Maximize the likelihood

#### Method of Moments (MM), Closed Form

- Pick enough moments that they **identify** the parameters
  At least 1 moment per parameter; algebraically independent
- Write equations for the moments in terms of the parameters e.g., for gamma

$$\mu=as\ ,\ \sigma^2=as^2$$

• Do the algebra by hand to solve the equations

$$a=\mu^2/\sigma^2~,~s=\sigma^2/\mu$$

• Code up the formulas (did this in lab 3)

```
gamma.est_MM <- function(x) {
  m <- mean(x); v <- var(x)
  return(c(shape=m^2/v, scale=v/m))
}</pre>
```

## MM, Numerically

- Write functions to get moments from parameters (usually algebra)
- Set up the difference between data and model as another function

```
gamma.mean <- function(shape,scale) { return(shape*scale) }
gamma.var <- function(shape,scale) { return(shape*scale^2) }
gamma.discrepancy <- function(shape,scale,x) {
  return((mean(x)-gamma.mean(shape,scale))^2 + (var(x)-gamma.mean(shape,scale))
}</pre>
```

• Minimize it

## More Generally...

- Nothing magic about moments
- Match other data summaries, say the median
  - Or even more complicated things, like ratios of quantiles
  - You did this in lab
- If you can't solve exactly for parameters from the summaries, set up a discrepancy function and minimize it
  - You are doing this in the HW
- The summaries just have to converge on population values

### Maximum Likeihood

- Usually we think of the parameters as fixed and consider the probability of different outcomes,  $f(x;\theta)$  with  $\theta$  constant and x changing
- **Likelihood** of a parameter value =  $L(\theta)$  = what probability does  $\theta$  give to the data?
  - For continuous variables, use probability density
  - $\circ \ f(x; heta)$  but letting heta change while data constant
  - $\circ$  *Not* the probability of  $\theta$ , if that even makes sense
- **Maximum likelihood** = guess that the parameter is whatever makes the data most likely
- Most likely parameter value = maximum likelihood estimate = MLE

### Likelihood in Code

• With independent data points  $x_1, x_2, x_n$ , likelihood is

$$L( heta) = \prod_{i=1}^n f(x_i; heta)$$

• Multiplying lots of small numbers is numerically bad; take the log:

$$\ell( heta) = \sum_{i=1}^n \log f(x_i; heta)$$

• In pseudo-code:

```
loglike.foo <- function(params, x) {
  sum(dfoo(x=x,params,log=TRUE))
}</pre>
```

### What Do We Do with the Likelihood?

- We maximize it!
- Sometimes we can do the maximization by hand with some calculus
  - For Gaussian, MLE = just match the mean and variance
  - $\circ~$  For Pareto, MLE  $\widehat{a} = 1 + 1/{\log{(x/x_{\min})}}$
- Doing numerical optimization
  - Stick in a minus sign if we're using a minimization function

## Why Use the MLE?

- Usually (but not always) *consistent*: converges on the truth as we get more data
- Usually (but not always) *efficient*: converges on the truth at least as fast as anything else
- There are some parameters where the maximum isn't well-defined (e.g.  $x_{\min}$  for a Pareto)
- Sometimes the data is too aggregated or mangled to use the MLE (as with the income data in lab 5)

### fitdistr

MLE for one-dimensional distributions can be done through fitdistr in the MASS package

It knows about most the standard distributions, but you can also give it arbitrary probability density functions and it will try to maximize them A starting value for the optimization is optional for some distributions, required for others (including user-defined densities)

Returns the parameter estimates and standard errors SEs come from large n approximations so use cautiously

## fitdistr Examples

Fit the gamma distribution to the cats' hearts:

```
require(MASS)
fitdistr(cats$Hwt, densfun="gamma")

## shape rate
## 20.2998092 1.9095724
## ( 2.3729243) ( 0.2259941)
```

Returns: estimates above, standard errors below

Fit the Students t distribution to the returns:

Here parameters are location (m), scale (s) and degrees of freedom (very heavy tails)

# fitdistr Examples (cont'd.)

#### User-defined density:

```
dpareto <- function(x,exponent,xmin,log=FALSE) {</pre>
   f <- (exponent-1)/xmin * (x/xmin)^(-exponent)</pre>
  f <- ifelse(x<xmin,NA,f)</pre>
   if(!log) { return(f) } else (return(log(f)))
# Fit pareto to large absolute returns
   # Parameters given outside the "start" list are fixed
fitdistr(abs(returns)[abs(returns)>0.05], densfun=dpareto,
          start=list(exponent=2.5), xmin=0.05)
## Warning in stats::optim(x = c(0.0529756421537337, 0.228997226565671, 0.051
## use "Brent" or optimize() directly
##
    exponent
##
   3,9960938
    (0.8309668)
##
```

# **Checking Your Estimator**

• simulate, then estimate; estimates should converge as the sample grows

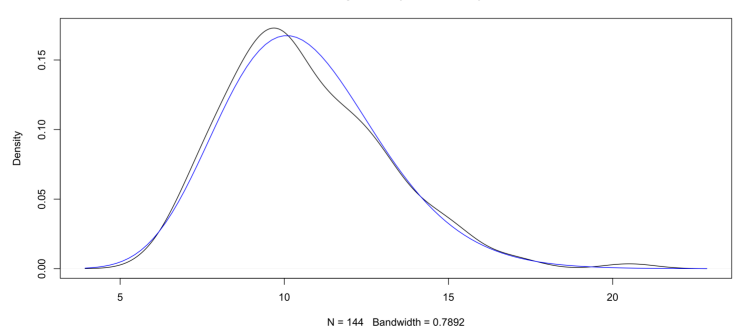
```
gamma.est_MM(rgamma(100, shape=19, scale=45))
     shape scale
##
## 20.72856 41.69062
gamma.est_MM(rgamma(1e5,shape=19,scale=45))
##
     shape scale
## 19.00673 44.88559
gamma.est_MM(rgamma(1e6,shape=19,scale=45))
##
     shape scale
## 18,99762 45,01647
```

# Checking the Fit

Use your eyes: Graphic overlays of theory vs. data

```
plot(density(cats$Hwt))
cats.gamma <- gamma.est_MM(cats$Hwt)
curve(dgamma(x,shape=cats.gamma["shape"],scale=cats.gamma["scale"]),a</pre>
```

#### density.default(x = cats\$Hwt)



# Checking the Fit (cont'd.)

• Calculate summary statistics *not* used in fitting, compare them to the fitted model

# Quantile-Quantile (QQ) Plots

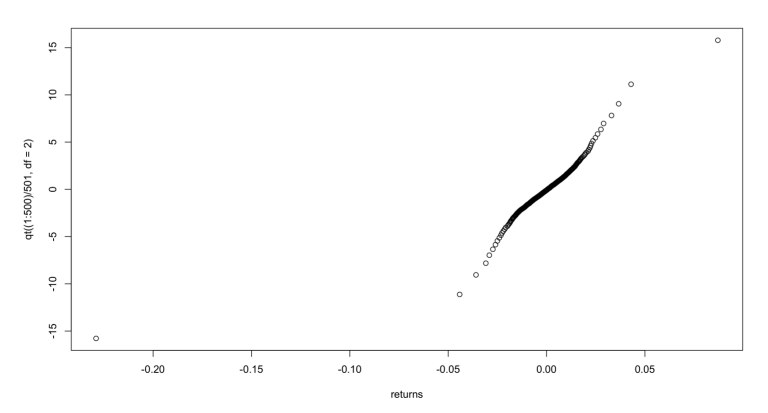
- Plot theoretical vs. actual quantiles
- or plot quantiles of two samples against each other
- Ideally, a straight line when the distributions are the same
- qqnorm, qqline are specialized for checking normality

```
qqnorm(returns); qqline(returns)
```

# QQ Plots (cont'd)

• qqplot(x,y) will do a Q-Q plot of one vector against another

```
qqplot(returns,qt((1:500)/501,df=2))
```



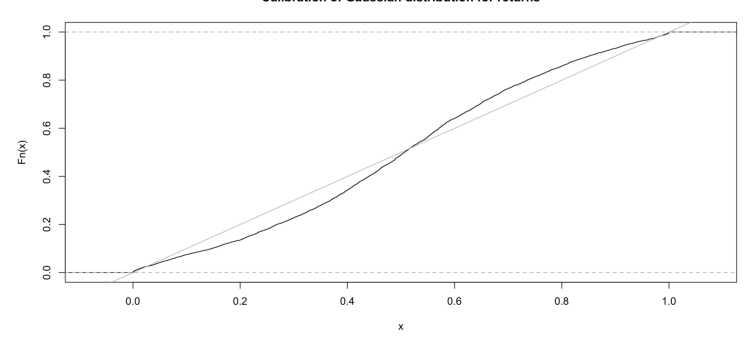
### **Calibration Plots**

- If the distribution is right, 50% of the data should be below the median, 90% should be below the 90th percentile, etc.
- Special case of **calibration** of probabilities: events with probability p% should happen about p% of the time, not more and not less
- We can look at calibraton by calculating the (empirical) CDF of the (theoretical) CDF and plotting
  - Ideal calibration plot is a straight line up the diagonal
  - Systematic deviations are a warning sign

# Making a Calibration Plot

```
plot(ecdf(pnorm(returns, mean=mean(returns), sd=sd(returns))),
    main="Calibration of Gaussian distribution for returns")
abline(0,1,col="grey")
```

#### Calibration of Gaussian distribution for returns



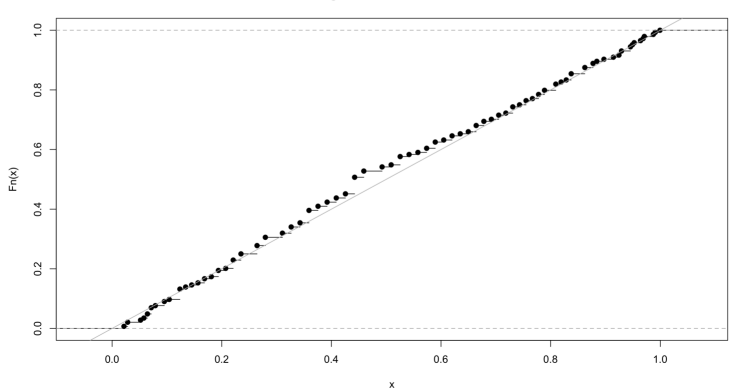
Again, way too many large changes (in either direction)

# Calibration Plots (cont'd.)

```
CXW.t <- coefficients(fitdistr(returns,"t"))
plot(ecdf(pt((returns-CXW.t[1])/CXW.t[2], df=CXW.t[3])),
         main="Calibration of t distribution for returns")
abline(0,1,col="grey")</pre>
```

# Calibration Plots (cont'd.)

#### Calibration of gamma distribution for cats' hearts



# Calibration Plots (cont'd.)

Challenge: Write a general function for making a calibraton plot, taking a data vector, a cumulative probability function, and a parameter vector

#### **Kolmogorov-Smirnov Test**

- How much should the QQ plot or calibration plot wiggle around the diagonal?
- Answer a different question...
- Biggest gap between theoretical and empirical CDF:

$$D_{KS} = \max_{x} \left| F(x) - \widehat{F}(x) 
ight|$$

- Useful because  $D_{KS}$  always has the same distribution if the theoretical CDF is fixed and correct, and K+S calculated this back in the day
- Also works for comparing the empirical CDFs of two samples, to see if they came from the same distribution

## KS Test, Data vs. Theory

```
ks.test(returns,pnorm,mean=0,sd=0.0125)
```

```
##
## One-sample Kolmogorov-Smirnov test
##
## data: returns
## D = 0.10893, p-value < 2.2e-16
## alternative hypothesis: two-sided</pre>
```

- More complicated (and not properly handled by built-in R) if parameters are estimated
  - Estimating parameters makes the fit look better than it really is, so it doesn't help save the model when it gets really rejected (like this one is)

Hack: Estimate using (say) 90% of the data, and then check the fit on the remaining 10%

```
train <- sample(1:length(returns),size=round(0.9*length(returns)))
CWX.t_train <- coefficients(fitdistr(returns[train],"t"))
returns.test_standardized <- (returns[-train]-CWX.t_train[1])/CWX.t_t
ks.test(returns.test_standardized,pt,df=CWX.t_train[3])</pre>
```

```
##
## One-sample Kolmogorov-Smirnov test
##
## data: returns.test_standardized
## D = 0.037817, p-value = 0.4772
## alternative hypothesis: two-sided
```

• Can also test whether two samples come from same distribution

```
n <- length(returns)
half <- round(n/2)
ks.test(returns[1:half], returns[(half+1):n])

##

## Two-sample Kolmogorov-Smirnov test
##

## data: returns[1:half] and returns[(half + 1):n]
## D = 0.099154, p-value = 5.103e-11
## alternative hypothesis: two-sided</pre>
```

## Chi-Squared Test for Discrete Distributions

Compare an actual table of counts to a hypothesized probability distribution e.g., as many up days as down?

```
up_or_down <- ifelse(returns > 0, 1, -1)
# 1936 down days, 1772 up days
chisq.test(table(up_or_down),p=c(1/2,1/2))

##
## Chi-squared test for given probabilities
##
## data: table(up_or_down)
## X-squared = 20.896, df = 1, p-value = 4.85e-06
```

#### **Chi-Squared Test: Degrees of Freedom**

- The p-value calculated by chisq.test assumes that all the probabilities in p were fixed, not estimated from the data used for testing, so df = number of cells in the table -1
- If we estimate q parameters, we need to subtract q degrees of freedom

## Chi-Squared Test for Continuous Distributions

- Divide the range into bins and count the number of observations in each bin; this will be x in chisq.test()
- Use the CDF function p *foo* to calculate the theoretical probability of each bin; this is p
- Plug in to chisq.test
- If parameters are estimated, adjust
- hist() gives us break points and counts:

```
cats.hist <- hist(cats$Hwt,plot=FALSE)
cats.hist$breaks

## [1] 6 8 10 12 14 16 18 20 22

cats.hist$counts

## [1] 20 45 42 23 11 2 0 1</pre>
```

# Chi-Squared for Continuous Data (cont'd.)

Use these for a  $\chi^2$  test:

Don't need to run hist first; can also use cut to discretize (see ?cut)

- This is all a bit old-school
  - Loss of information from discretization
  - $\circ$  Lots of work just to use  $\chi^2$
- Try e.g. ks.test with an independent test set

## Summary

- Visualizing and computing empirical distribution
- Parametric distributions are models
- Methods of fitting: moments, generalized moments, likelihood
- Methods of checking: visual comparisons, other statistics, tests, calibration

### Aside: Some Math for MM and GMM

- Parameter  $\theta$  is a p-dimensional vector, true value =  $\theta^*$
- Introduce  $q \geq p$  functionals  $g_1, \ldots g_q$ , which we can calculate either from the parameter  $\theta$  or from the data  $x_{1:n}$
- ullet Assume that for each  $i,g_i(x_{1:n}) o g_i( heta^*)$
- Define

$$\hat{ heta}_{GMM} = \operatorname{argmin}_{ heta} \sum_{i=1}^q \left( g_i( heta) - g_i(x_{1:n}) 
ight)^2$$

# Math for MM and GMM (cont'd.)

- ullet Shouldn't be hard to believe that  $\hat{ heta}_{GMM} 
  ightarrow heta^*$
- But why give equal attention to every functional?
  - More weight on the more-precisely-measured functionals
  - $\circ$  More weight on the more-sensitive-to heta functionals
  - Less weight on partially-redundant functionals
- Abbreviate  $g(\theta)$  for  $(g_1(\theta), \dots g_q(\theta))$ , and likewise  $g(x_{1:n})$ , so

$$\hat{ heta}_{GMM} = \operatorname{argmin}_{ heta}(g( heta) - g(x_{1:n}))^T (g( heta) - g(x_{1:n}))^T$$

• Generalize by introducing any positive-definite matrix  $\Omega$ :

$$\hat{ heta}_{GMM} = \operatorname{argmin}_{ heta}(g( heta) - g(x_{1:n}))^T \Omega(g( heta) - g(x_{1:n}))$$

- Optimal  $\Omega$  turns out to be the variance matrix of  $g( heta^*)$
- Iterative approximation: start with no weighting, estimate that variance matrix, re-do the estimate with weights, etc.

### Aside: Some Math for the MLE

• More convenient to work with the mean log likelihood:

$$\Lambda( heta) = rac{1}{n} \sum_{i=1}^n \log f(X_i; heta)$$

• This is a sample average so the law of large numbers applies:

$$\Lambda( heta) o \mathbf{E}[\Lambda( heta)] = \lambda( heta)$$

• The true parameter has higher average log-likelihood than anything else: if  $\theta \neq \theta^*$ 

$$heta 
eq heta^* \ \Rightarrow \lambda( heta) < \lambda( heta^*)$$

• Some extra conditions are needed for

$$\hat{ heta}_{MLE} 
ightarrow heta^*$$