

T-test example using RA Fisher's Cat Data

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An example of doing “inference by eye” using R.A. Fisher's cat data. A good summary of this idea is Cummings et al. 2007. Error bars in experimental biology. <http://jcb.rupress.org/content/177/1/7.short>

Load the data

```
library(MASS)
data("cats")
```

Look at the data

```
dim(cats)

## [1] 144  3

summary(cats)

##   Sex           Bwt           Hwt
##   F:47   Min.    :2.000   Min.    : 6.30
##   M:97   1st Qu.:2.300   1st Qu.: 8.95
##           Median :2.700   Median :10.10
##           Mean   :2.724   Mean    :10.63
##           3rd Qu.:3.025   3rd Qu.:12.12
##           Max.    :3.900   Max.    :20.50
```

Summarize the body weight (Bwt) data old-school using summaryBy()

A more modern way would be to use dplyr()

```
library(dplyr)

#get the mean and SD
cat.df1 <- summaryBy(Bwt ~ Sex, data = cats, FUN = c(mean,sd))

#get the sample size using length()
cat.df2 <- summaryBy(Bwt ~ Sex, data = cats, FUN = c(length))

#make a combined dataframe
cat.df3 <- merge(cat.df1,cat.df2)

#calculate the standard error SE by hand

cat.df3$SE <- cat.df3$Bwt.sd/sqrt(cat.df3$Bwt.length)
```

Look at the results

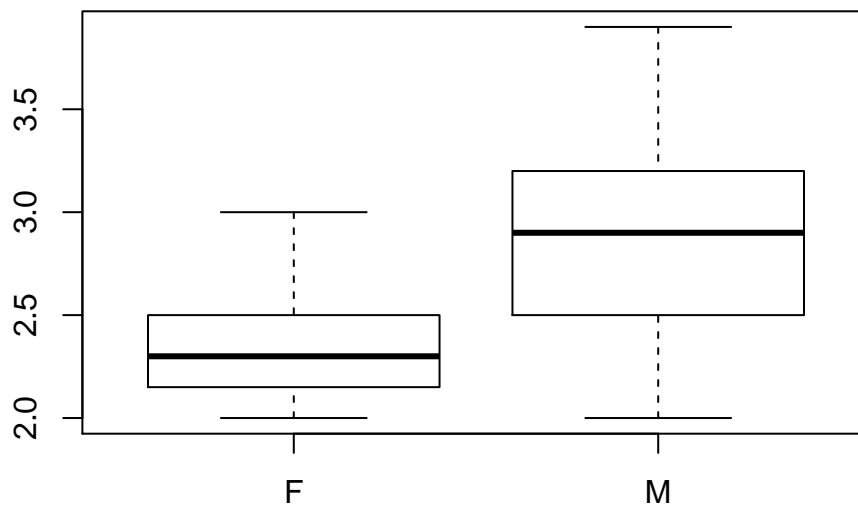
```
cat.df3
```

```
##   Sex Bwt.mean   Bwt.sd Bwt.length      SE
## 1  F 2.359574 0.2739879      47 0.03996525
## 2  M 2.900000 0.4674844      97 0.04746585
```

Plot the data

Visualize the raw data

```
par(mfrow = c(1,1),mar = c(3,3.5,1,1))
boxplot(Bwt ~ Sex, data = cats)
```



Plot the means with error bars

This uses the `errbar()` function. A modern contemporary way would use `ggplot2` and possibly its extension using `ggpubr`.

The real data

This is the actual data. The 95% confidence intervals do not overlap, which indicates that the p-value for the t-test will be less than 0.05.

```
library(Hmisc)
par(mfrow = c(1,2),mar = c(3,3.5,1,1))

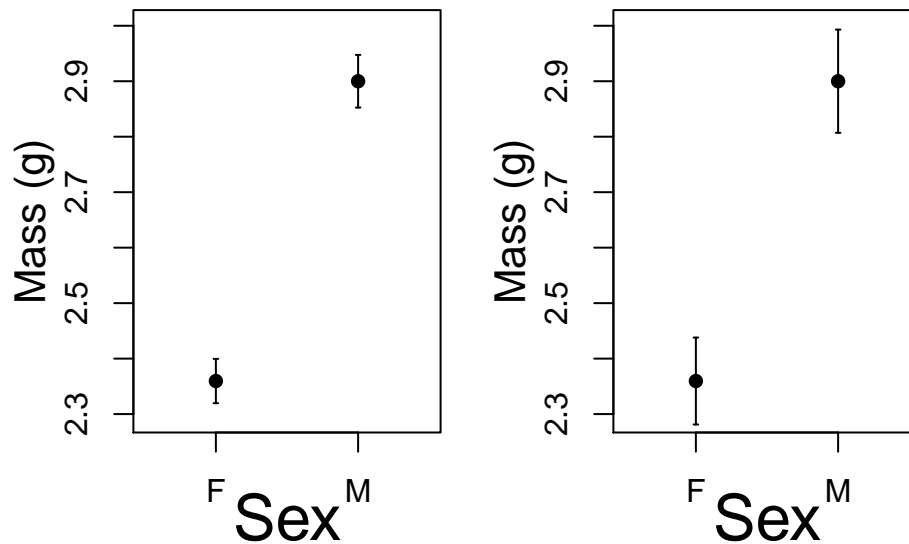
y.lim <- c(2.295,3)
errbar(1:2,
      y = cat.df3$Bwt.mean,
      yplus = cat.df3$Bwt.mean + cat.df3$SE,
      yminus = cat.df3$Bwt.mean - cat.df3$SE,
      xlab = "",
```

```

ylab = "",
xlim=c(0.5,2.5),
ylim = y.lim,
xaxt="n",cex =1)
axis(side=1,at=1:2,labels=cat.df3$Sex)
mtext("Sex", side = 1, line = 2, cex = 2)
mtext("Mass (g)", side = 2, line = 2.1, cex = 1.3)

errbar(1:2,
  y = cat.df3$Bwt.mean,
  yplus =cat.df3$Bwt.mean + 1.96*cat.df3$SE,
  yminus = cat.df3$Bwt.mean-1.96*cat.df3$SE,
  xlab = "",
  ylab = "",
  xlim=c(0.5,2.5),
  ylim = y.lim,
  xaxt="n",cex =1)
axis(side=1,at=1:2,labels=cat.df3$Sex)
mtext("Sex", side = 1, line = 2, cex = 2)
mtext("Mass (g)", side = 2, line = 2.1, cex = 1.3)

```



Modified data with a non-significant different

Make an alternative version of the data where there isn't a difference between the male and female cats

```

cat.df3.mod <- cat.df3
cat.df3.mod$Bwt.mean[1] <- cat.df3$Bwt.mean[2]-cat.df3$Bwt.mean[2]*0.0425

```

The overlap of the error bars here is greater than 1/2 the length of the bar; therefore the p-value for a t-test will be > 0.05 .

```

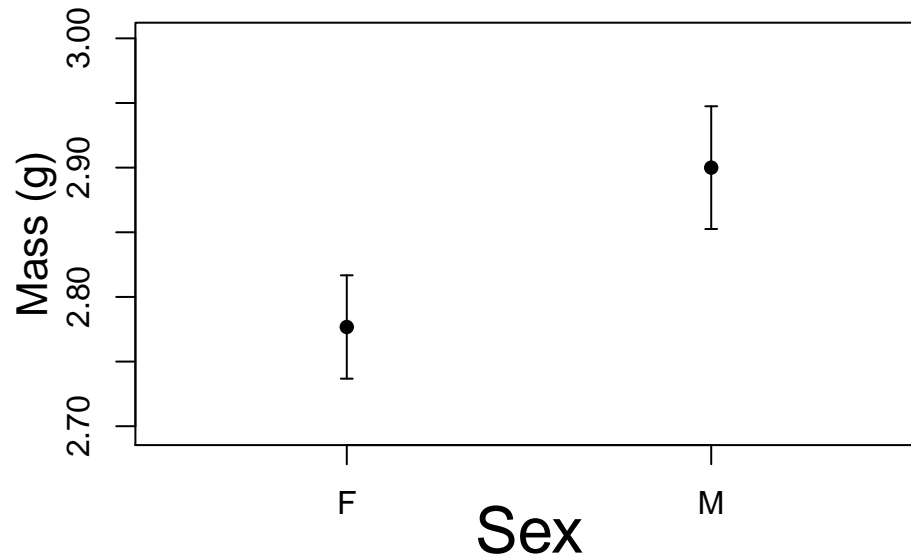
y.lim <- c(2.6975,3)
par(mar = c(3,3.5,1,1))
errbar(1:2,
  y = cat.df3.mod$Bwt.mean,
  yplus =cat.df3.mod$Bwt.mean + cat.df3.mod$SE,

```

```

yminus = cat.df3.mod$Bwt.mean-cat.df3.mod$SE,
xlab = "",
ylab = "",
xlim=c(0.5,2.5),
ylim = y.lim,
xaxt="n",cex =1)
axis(side=1,at=1:2,labels=cat.df3.mod$Sex)
mtext("Sex", side = 1, line = 2, cex = 2)
mtext("Mass (g)", side = 2, line = 2.1, cex = 1.3)

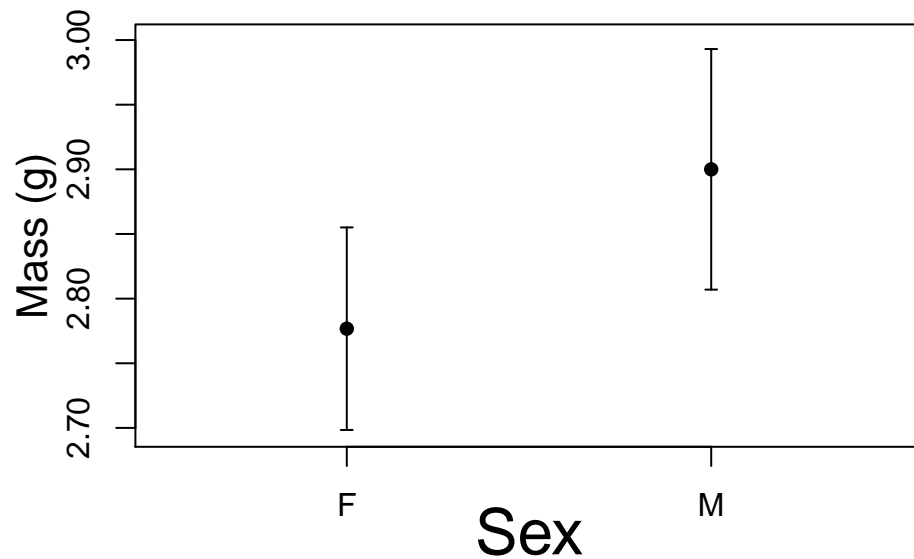
```



```

errbar(1:2,
  y = cat.df3.mod$Bwt.mean,
  yplus =cat.df3.mod$Bwt.mean + 1.96*cat.df3.mod$SE,
  yminus = cat.df3.mod$Bwt.mean-1.96*cat.df3.mod$SE,
  xlab = "",
  ylab = "",
  xlim=c(0.5,2.5),
  ylim = y.lim,
  yaxt="n",cex =1)
axis(side=1,at=1:2,labels=cat.df3.mod$Sex)
mtext("Sex", side = 1, line = 2, cex = 2)
mtext("Mass (g)", side = 2, line = 2.1, cex = 1.3)

```



T-test

A t-test for the difference between female and male cats.

```
t.test(Bwt ~ Sex, data = cats)
```

```
##
## Welch Two Sample t-test
##
## data: Bwt by Sex
## t = -8.7095, df = 136.84, p-value = 8.831e-15
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -0.6631268 -0.4177242
## sample estimates:
## mean in group F mean in group M
##      2.359574      2.900000
```

```
summary(lm(Bwt ~ -1+Sex, data = cats))
```

```
##
## Call:
## lm(formula = Bwt ~ -1 + Sex, data = cats)
##
## Residuals:
##      Min       1Q   Median       3Q      Max
## -0.90000 -0.25957 -0.05957  0.30000  1.00000
##
## Coefficients:
##      Estimate Std. Error t value Pr(>|t|)
## SexF    2.35957    0.06051   39.00  <2e-16 ***
## SexM    2.90000    0.04212   68.86  <2e-16 ***
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
```

```
## Residual standard error: 0.4148 on 142 degrees of freedom
## Multiple R-squared:  0.9778, Adjusted R-squared:  0.9775
## F-statistic: 3131 on 2 and 142 DF,  p-value: < 2.2e-16
```