Agenda

- 1. More ANOVA
- 2. ANOVA lab

ANOVA Statisticians overload the term ANOVA. We use it when we are doing nested F-tests, when we are testing the significance of a categorical variable within a larger linear regression model, and we use it when we are just predicting a quantitative variable with a categorical one.

Just to get some intuition, lets compare the results from a few different approaches to using ANOVA.

```
require(mosaic)
bloodp = read.csv("http://www.math.smith.edu/~bbaumer/mth247/labs/bloodpress.csv")
m1 = lm(BP \sim Weight, data=bloodp)
anova(m1)
## Analysis of Variance Table
##
## Response: BP
          Df Sum Sq Mean Sq F value
           1 505.47 505.47 166.86 1.528e-10 ***
## Weight
## Residuals 18 54.53
                      3.03
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
m2 = lm(BP \sim Age, data=bloodp)
anova(m2)
## Analysis of Variance Table
##
## Response: BP
            Df Sum Sq Mean Sq F value
            1 243.27 243.266 13.825 0.001574 **
## Residuals 18 316.73 17.596
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
m3 = aov(BP~ Weight + Age, data=bloodp)
summary(m3)
              Df Sum Sq Mean Sq F value Pr(>F)
##
## Weight
             1 505.5 505.5 1781.3 < 2e-16 ***
              1 49.7
                        49.7
                                175.2 2.22e-10 ***
## Age
## Residuals
             17
                  4.8
                           0.3
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
m4 = aov(BP~ Age + Weight, data=bloodp)
summary(m4)
##
              Df Sum Sq Mean Sq F value Pr(>F)
              1 243.27 243.27 857.3 5.48e-16 ***
## Age
## Weight
             1 311.91 311.91 1099.2 < 2e-16 ***
## Residuals 17 4.82
                          0.28
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

```
m5 = aov(BP~., data=bloodp)
summary(m5)
##
           Df Sum Sq Mean Sq F value Pr(>F)
           1 243.27 243.27 1466.914 9.38e-15 ***
## Weight
            1 311.91 311.91 1880.844 1.89e-15 ***
            1 1.77 1.77 10.660 0.00615 **
## BSA
            1 0.34 0.34 2.021 0.17871
## Dur
## Pulse
            1 0.12 0.12 0.742 0.40458
## Stress 1 0.44 0.44 2.666 0.12649
## Residuals 13 2.16 0.17
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
m6 = aov(BP~Stress+Pulse+Dur+BSA+Weight+Age, data=bloodp)
summary(m6)
            Df Sum Sq Mean Sq F value Pr(>F)
## Stress
           1 15.04 15.04 90.714 3.16e-07 ***
## Pulse
            1 306.93 306.93 1850.831 2.09e-15 ***
            1 0.72 0.72 4.337 0.0576 .
## Dur
            1 173.87 173.87 1048.444 8.15e-14 ***
## BSA
           1 27.95 27.95 168.533 8.09e-09 ***
## Weight
## Age
            1 33.33 33.33 200.986 2.76e-09 ***
## Residuals 13 2.16 0.17
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

ANOVA lab For your convenience, here is the code from the ANOVA lab

```
require(mosaic)
require(Stat2Data)
data(FruitFlies)
head(FruitFlies)
tally(~Treatment, data=FruitFlies)
# Set the reference level
FruitFlies = FruitFlies %>%
 mutate(Treatment = relevel(Treatment, ref="none"))
d1plot = dotPlot(~Longevity, data=FruitFlies)
d2plot = dotPlot(~Longevity | Treatment, data=FruitFlies, layout=c(1,5))
# arrange the two plots horizontally
print(d1plot, position=c(0, 0, 0.5, 1), more=TRUE)
print(d2plot, position=c(0.5, 0, 1, 1))
b1plot = bwplot(~Longevity, data=FruitFlies)
b2plot = bwplot(~Longevity | Treatment, data=FruitFlies, layout=c(1,5))
# arrange the two plots horizontally
print(b1plot, position=c(0, 0, 0.5, 1), more=TRUE)
print(b2plot, position=c(0.5, 0, 1, 1))
```

```
gstats = favstats(Longevity ~ Treatment, data=FruitFlies)
gstats
fm.null = lm(Longevity ~ 1, data=FruitFlies)
summary(fm.null)
mean(~Longevity, data=FruitFlies)
fitted = fitted.values(fm.null)
ladd(panel.abline(v=coef(fm.null), col="red", lwd=3), plot=d2plot)
fm.aov = aov(Longevity ~ Treatment, data=FruitFlies)
summary(fm.aov)
model.tables(fm.aov)
fm.ref = lm(Longevity ~ Treatment, data=FruitFlies)
summary(fm.ref)
anova(fm.ref)
mean(Longevity~Treatment, data = FruitFlies) - mean(~Longevity, data = FruitFlies)
dotPlot(~Longevity | Treatment, data=FruitFlies, layout=c(1,5), panel=function(x,...){
  panel.dotPlot(x, ...)
 fitted = mean(x)
 panel.abline(v=fitted, col="red")
})
par(mfrow=c(2,2))
plot(fm.aov)
par(mfrow=c(1,1))
sd(Longevity~Treatment, data=FruitFlies)
max(sd(Longevity~Treatment, data=FruitFlies)) / min(sd(Longevity~Treatment, data=FruitFlies))
data(Hawks)
```