BMEG 802 – Advanced Biomedical Experimental Design and Analysis

One Way (Between) Analysis of Variance (ANOVA)

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Recap

- Regression
 - Bivariate
 - Linear (Derivation)
 - Nonlinear
 - Multiple Regression
- Correlation
 - Pearson's r
 - lacksquare Spearman's ho

1 Way (Between) ANOVA

- develop logic & rationale for ANOVA (and formulas) based on a General Linear Model Approach
- any phenomenon is affected by multiple factors
- observed value on dependent variable (DV) = sum of effects of known factors + sum of effects of unknown factors
- similar idea to traditional approach of "accounting for variance" due to various factors
 - equivalent mathematically

General Linear Model (GLM)

Let's develop a model that expresses dependent variable (DV) as a sum of known and unknown factors

$$DV = C + F + R$$

- C = constant factors (known)
- F = factors systematically varied (known)
- R = randomly varying factors (unknown)

Notation looks like this:

$$Y_i = \beta_0 + \beta_1 \cdot X 1_i + \beta_2 \cdot X 2_i + \ldots + \beta_n \cdot X n_i + \epsilon_i$$

Single Group Example

- a little artificial (who ever does experiments using just one group?)
- but it will help us develop the ideas
- imagine we collect scores on some DV for a group of subjects
- we want to compare the group mean to some known population mean
- e.g. IQ scores where by definition, $\mu=100$ and $\sigma=15$

Single Group Example

We know that:

$$H_0: \bar{Y} = \mu$$

$$H_1: \bar{Y} \neq \mu$$

Let's reformulate in terms of a GLM of the effects on DV:

$$H_0: \overline{Y} = \mu + \epsilon$$
; where $\mu = 100$

$$H_1: \overline{Y} = \hat{\mu} + \epsilon$$
; where $\hat{\mu} = Y_i$

Terminology:

 ${\it H}_{\rm 0}$ is the "Restricted Model" — no parameters need to be estimated

 H_0 is the "Full Model" — we need to estimate one parameter (can you see what it is?)

- how well do these two models fit our data?
- let's use the sum of squared deviations of our model from the data, as a measure of goodness of fit

$$H_0: \sum_{i=1}^N (e_i^2) = \sum_{i=1}^N (Y_i - 100)^2$$

$$H_1: \sum_{i=1}^N (e_i^2) = \sum_{i=1}^N (Y_i - \hat{\mu})^2 = \sum_{i=1}^N (Y_i - \hat{Y})^2$$

- SSE about the sample mean is lower than SSE about any other number
- so the error for H_0 will be greater than for H_1
- so the relevant question then is, how much greater must H_0 error be, for us to reject H_0 ?

- Consider the Proportional Increase in Error (PIE)
 - $(E_{restricted} E_{full})/E_{full}$
- PIE gives error increase for H_0 (restricted) compared to H_1 as (full) a % of H_1 error
- We want a model that is both
 - adequate (low error)
 - simple (few parameters to estimate)
- question: why do we want a simpler model?
 - philosophical reasons (Occam's razer)
 - statistical reasons (over-fitting)

- how big is increase in error with H_0 (restricted model), per unit of simplicity?
- let's design a test statistic that takes into account simplicity
- simplicity will be related to the number of parameters we have to estimate
- degrees of freedom (df): # of independent observations in the dataset minus # of independent parameters that need to be estimated
- higher df = a simpler model

Let's normalize model errors (PIE) by model df - This is called the F-statistic!

$$F = \frac{(E_{restricted} - E_{full})/(df_{restricted} - df_{full})}{(E_{full}/df_{full})}$$

We can compute F_{obs} and calculate the probability of obtaining that F-statistic by using the F-distribution!

Two Group Example

Let's look at a more realistic situation

- 2 groups, 10 subjects in each group
 - test mean of group 1 vs mean of group 2
 - do we accept H_0 or H_1 ?
- we will formulate this question as before in terms of 2 linear models
 - full vs restricted model
 - is the error for the restricted model significantly higher than for the full model?
 - is the decrease in error for the full model large enough to justify the need to estimate a greater # parameters?

Two Group Example: Hypotheses and Models

```
H_0: \mu = \mu_1 = \mu_2
restricted model: Y_{ii} = \mu + \epsilon_{ii}
H_1: \mu_1 \neq \mu_2
full model: Y_{ii} = \mu_i + \epsilon_{ii}
i = individual (1, 2, ..., 10)
i = group (1 or 2)
```

Restricted model:

• each score Y_{ii} is the result of a single population mean plus random error ϵ_{ii}

Full model:

• each score Y_{ii} is the result of a different group mean plus random error ϵ_{ii}

Deciding between full and restricted model

how do we decide between these two competing accounts of the data?

key question

- will a restricted model with fewer parameters be a significantly less adequate representation of the data than a full model with a parameter for each group?
- we have a trade-off between simplicity (fewer parameters) and adequacy (ability to accurately represent the data)

Error for the Restricted Model

- Sum of squared deviations of each observation from the estimate of the population mean (given by the grand mean of all of the data)
- Here we need to estimate 1 parameter, $\hat{\mu}$

$$E_{restricted} = \sum_{j=1}^{n_j} \sum_{i=1}^{n_i} (Y_{ij} - \hat{\mu})^2$$

$$\hat{\mu} = \frac{1}{n_i \cdot n_j} \sum_{j=1}^{n_j} \sum_{i=1}^{n_i} (Y_{ij})$$

Error for the Full Model

• Here we need to estimate 2 parameter (one for each group), $\hat{\mu_1}$ and $\hat{\mu_2}$

$$E_{full} = \sum_{j=1}^{n_j} \sum_{i=1}^{n_i} (Y_{ij} - \hat{\mu_j})^2 = \sum_{i=1}^{n_i} (Y_{i1} - \hat{\mu_1})^2 + \sum_{i=1}^{n_i} (Y_{i2} - \hat{\mu_2})^2$$

$$\hat{\mu_1} = \frac{1}{n_1} \sum_{i=1}^{n_i} (Y_{i1})$$

$$\hat{\mu_2} = \frac{1}{n_2} \sum_{i=1}^{n_i} (Y_{i2})$$

Deciding between Full and Restricted Model

Now we formulate our measure of proportional increase in error (PIE) as before using the F-statistic:

$$F = \frac{(E_{restricted} - E_{full})/(df_{restricted} - df_{full})}{(E_{full}/df_{full})}$$

Model Comparison Approach vs Traditional Approach to ANOVA

Traditional formulation of ANOVA asks the same question in a different way:

• is the variability between groups (variance due to differences between groups) greater than expected on the basis of the within-group variability (the variability within a group) observed, and random sampling of group members?

Maxwell, Delaney, Kelley, Chapter 3: Proof that these two approaches are mathematically equivelant

- both use sum of squares
- both use F-statistic

Assumptions of the F test

- 1. the scores on the dependent variable Y are normally distributed in the population (and normally distributed within each group)
- 2. the population variances of scores on Y are equal for all groups
- 3. scores are independent of one another

Violation of Assumption

- ANOVA is somewhat robust to violations of normality
- ANOVA is somewhat robust to homogeneity of variance
- ANOVA is NOT robust to violations of independence

- We can use ANOVA / GLM for 1 sample and 2 sample tests.
 - But, equivalent to using a t-test
- ANOVA are most widely used when there is greater than 2 groups. Acts as the "omnibus" test to decide whether you have "permission" to perform follow-up mean comparisons.
- Lets perform an ANOVA using R for the following simple example. We want to know if the following three groups are different from one another.

Let's determine if there is a main effect of group.

Group 1	Group 2	Group 3
4	7	6
5	4	9
2	6	8
1	3	5
3	5	7
mean = 3	mean = 5	mean = 7

$$H_0: \mu_1 = \mu_2 = \mu_3$$

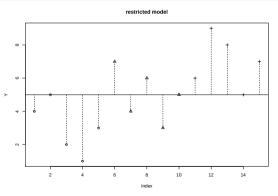
restricted model: $Y_{ij} = \mu + \epsilon_{ij}$

$$H_1: \mu_1 \neq \mu_2 \neq \mu_3$$

full model: $Y_{ij} = \mu_i + \epsilon_{ij}$

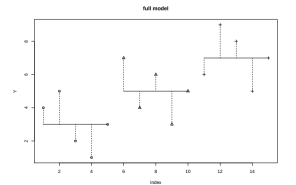
Plot Restricted Model: Single parameter, μ

```
Y <- c(4,5,2,1,3,7,4,6,3,5,6,9,8,5,7)
myFac <- c(1,1,1,1,2,2,2,2,2,3,3,3,3,3)
plot(Y, pch=myFac,main="restricted model")
abline(h=mean(Y))
for (i in 1:length(Y)) {
    lines(c(i,i), c(Y[i], mean(Y)), lty=2)}
```



Plot Full Model: three parameters, μ_1, μ_2, μ_3

```
Y <- c(4,5,2,1,3,7,4,6,3,5,6,9,8,5,7)
myFac <- c(1,1,1,1,1,2,2,2,2,2,3,3,3,3,3)
plot(Y, pch=myFac, main="full model")
for (j in 1:3) {
    w <- which(myFac==j)
    lines(c(min(w),max(w)),c(mean(Y[w]),mean(Y[w])))
    for (i in 1:length(w)) {
        lines(c(w[i],w[i]), c(Y[w[i]], mean(Y[w])), lty=2) }}</pre>
```



```
Using the aov() in R
m1 <- aov(Y ~ factor(mvFac))</pre>
summary(m1)
##
                 Df Sum Sq Mean Sq F value Pr(>F)
## factor(mvFac) 2 40
                              20.0 8 0.0062 **
## Residuals 12 30 2.5
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
Here we find that there is statistically significant main effect of group (p = 0.006)!
```

Alternatively we can build a linear model, using lm(), and then pass it through aov()

```
m2 <- lm(Y - factor(myFac))
summary(m2)</pre>
```

```
##
## Call:
## lm(formula = Y ~ factor(myFac))
##
## Residuals:
     Min
             10 Median
                                Max
      -2
             -1
                     0
## Coefficients:
##
                 Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                   3.0000
                             0.7071
                                      4.243 0.00114 **
## factor(mvFac)2
                  2.0000
                             1.0000
                                      2.000 0.06866 .
## factor(mvFac)3 4.0000
                            1.0000 4.000 0.00176 **
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 1.581 on 12 degrees of freedom
## Multiple R-squared: 0.5714, Adjusted R-squared:
## F-statistic:
                   8 on 2 and 12 DF. p-value: 0.006196
```

In this case the estimate for the first group (called Intercept in the anova output) is 3.0000. The estimate for the mean of group two is equal to the Intercept plus 2.0000, which equals 5.0000. Likewise the estimate for group three is 3.0000 + 4.0000 which equals 7.0000.

anova(m2)

... now running an F-test on our linear model:

```
## Analysis of Variance Table

##
## Response: Y

## Df Sum Sq Mean Sq F value Pr(>F)

## factor(myFac) 2 40 20.0 8 0.006196 **

## Residuals 12 30 2.5

## ---

## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

The F-test of the main effect of the factor is called an omnibus test. A significant test indicates only that the population means are not equal —we would need to perform follow-up tests to find out specifically which groups differ.

Testing Normality

Use Shapiro-Wilk test on EACH group to test for normality (i.e., normal: p > 0.05).

```
shapiro.test(Y[1:5]) # group 1
##
   Shapiro-Wilk normality test
## data: Y[1:5]
## W = 0.98676, p-value = 0.9672
shapiro.test(Y[5:10]) # group 2
##
    Shapiro-Wilk normality test
##
## data: Y[5:10]
## W = 0.91994, p-value = 0.505
shapiro.test(Y[11:15]) # group 3
##
   Shapiro-Wilk normality test
## data: Y[11:15]
## W = 0.98676, p-value = 0.9672
```

Testing Homogeneity of Variances

Bartlett test to test whether variances are similar (i.e., equal homogeneity of variance: p>0.05)

```
##
## Bartlett test of homogeneity of variances
##
## data: Y by factor(myFac)
## Bartlett's K-squared = 0. df = 2. p-value = 1
```

bartlett.test(Y ~ factor(mvFac))

There is no violation of Homogeneity of Variances: p = 1.0

- Data not normal? Can use transforms (e.g., log, square root, etc) OR a nonparametric version of the 1-way ANOVA, known as the Kruskal Wallis.
- Can perform Welch's corrections to F-test if variances are not equal using the oneway.test()
 OR use Kruskal Wallis.

Follow up Mean Comparison's

If ANOVA significant, as well as no violations of either normality or homogeneity of variance, you can perform follow-up mean compomparisons to test for differences between groups. Remember to correct for multiple comparisons!

```
pval_1v2 = t.test(Y[1:5], Y[5:10], alternative = "two.sided")$p.value # G1 vs G2
pval_1v3 = t.test(Y[1:5], Y[11:15], alternative = "two.sided")$p.value # G1 vs G3
pval_2v3 = t.test(Y[5:10], Y[11:15], alternative = "two.sided")$p.value # G2 vs G3
pvals = c(pval_1v2, pval_1v3, pval_2v3)
p.adjust(pvals, method = "holm", n = length(pvals))
```

[1] 0.12147302 0.01184932 0.08119087

Interpretation: There is a significant main of effect of group (p = 0.006), where Group 3 is significantly greater than Group 1 (p = 0.012).

Graphing with GG Plot

Next Week

- Factorial (2-way, 3-way, etc.) ANOVA
- Kruskal Wallis