STA610 Lab01 ANOVA and REANOVA

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- Write down your answers in any blank sheet and submit your work in paper during the lab.
- Your work will not be graded. As long as you submit, you will get a full credit.
- For those who missed the lab today, you can submit it via email to me for half credit.

1. ANOVA as a linear regression

Consider an one-way ANOVA model where an ith unit of a jth group is modeled as

$$y_{ij} = \theta_j + \epsilon_{ij}, \quad i \in [n_j], \quad j \in [m] \quad (n_j = n \quad \forall j)$$

$$\downarrow_{ij} \quad iid \quad N(0, \sigma^2)$$

Another way to look at this model is as follows. First, we reparameterize the group mean parameters θ_i as

$$\theta_j = \begin{cases} \alpha & j = 1\\ \alpha + \beta_{j-1} & j \ge 2 \end{cases}$$

Then you can check that the above model can be written as

$$y = \alpha 1_{mn} + X\beta + \epsilon$$

Q1-1: Write down the form of a design matrix X.

The hypothesis of testing no difference in group means can be written as

$$H_0: \theta_1 = \cdots = \theta_p \quad \text{vs} \quad H_1: \theta_i \neq \theta_i \quad \exists (i,j)$$

Q1-2: Re-express the above H_0 using β .

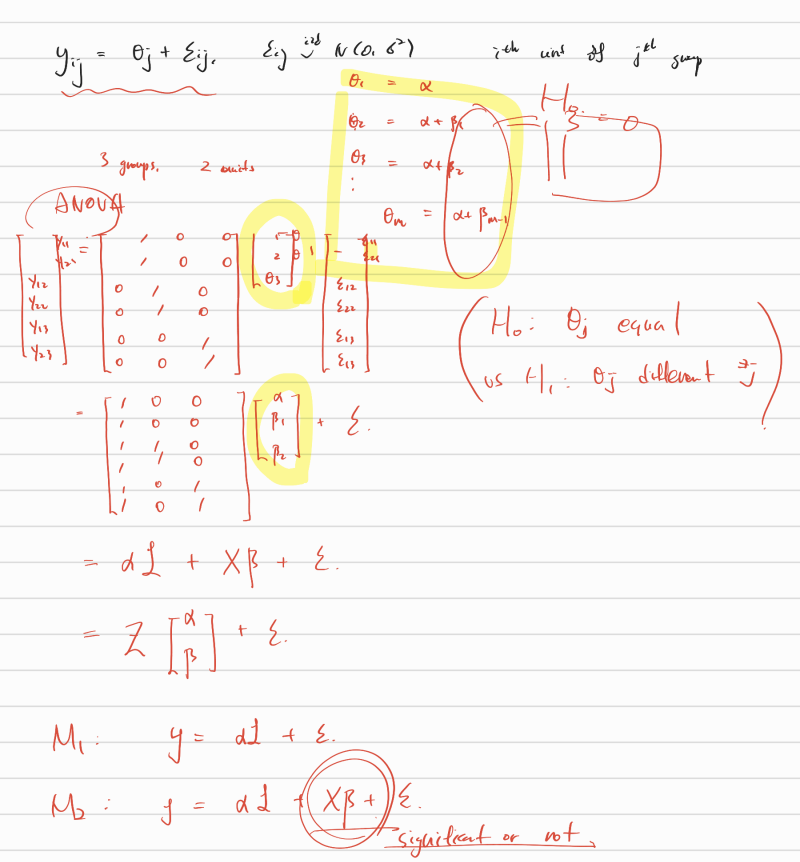
This tells us that ANOVA can be seen as a linear regression, and its hypothesis testing is equivalent to testing a submodel of linear regression. We do not proceed from here, but the general idea is as follows. Note that we wrote ANOVA decomposition as SST = SSA + SSW where

$$SST = \sum_{j=1}^{m} \sum_{i=1}^{n} (y_{ij} - \bar{y})^{2} \qquad \text{leagth dervor using c(Ln)}$$

$$SSA = \sum_{j=1}^{m} \sum_{i=1}^{n} (\bar{y}_{j} - \bar{y})^{2} \qquad \underbrace{SST - SSW}_{\text{by using c(Ln, X)}}_{\text{by using c(Ln, X)}}$$

$$SSW = \sum_{j=1}^{m} \sum_{i=1}^{n} (y_{ij} - \bar{y}_{j})^{2} \qquad \text{leagth dervor using c(Ln, X)}_{\text{by using c(Ln, X)}}$$

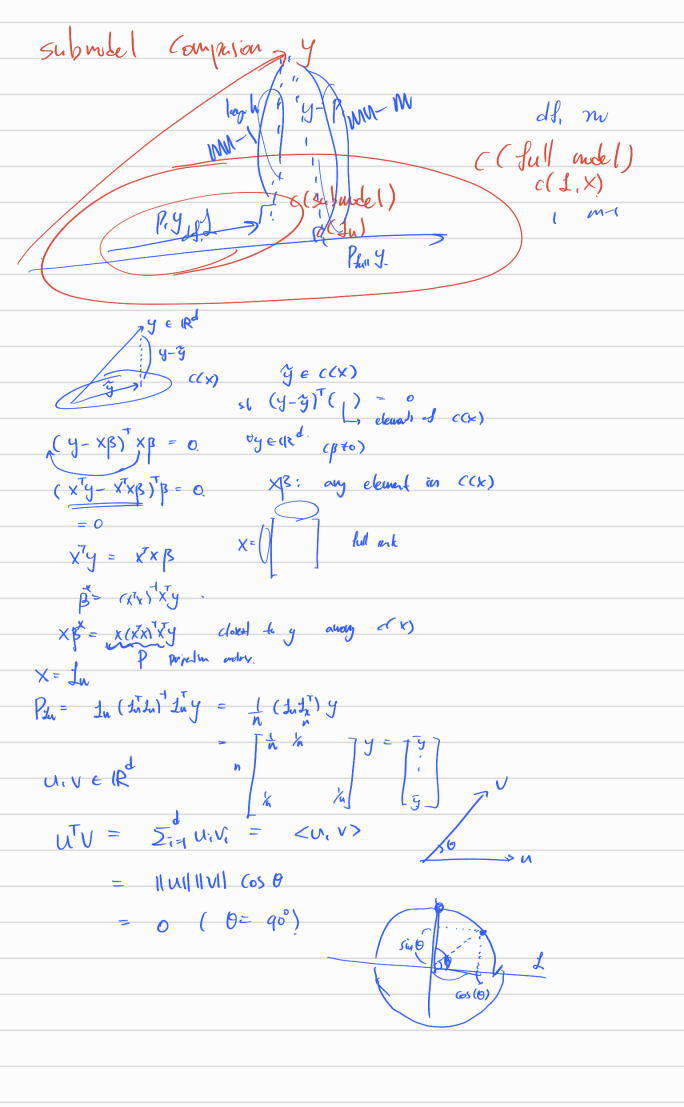
For a matrix X, we write c(X) the vector space its columns expand, i.e., column space. Also, we write P_X an orthogonal projection matrix onto c(X). For convenience, let $Z = cbind(1_{mn}, X)$. The dimension of c(Z)is m.



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$$C(I_n)$$
 vs $C(I_n, XI)$

- i) paroject y onto c(·): y= Py.
- 11) Compare the reduly 11 y- Ry11 s 14- Ry11



((I-p)y) (I-p)y= y (I-p) (I-p)y= y (I-p)7

- 1. One can see that $SST = \|(I P_1)y\|^2 = y^T(I P_1)y$, $SSA = y^T(P_Z P_1)y$ and $SSW = y^T(I P_Z)y$.
- 2. $(I P_1)y$ is a vector y projected onto the column space orthogonal to $c(1_{mn})$, whose dimension is mn 1.
- 3. $(I P_Z)y$ is a vector y projected onto the column space orthogonal to c(Z), whose dimension is mn m = m(n-1).
- 4. $(P_Z P_1)y$ is a vector y projected onto the subspace of c(Z) that are orthogonal to $c(1_{mn})$, whose dimension is m-1

Using these, the F-statistics of testing H_0 is

 $c(Z) \cap c(Zu)$

Note that $SSA = y^T(P_Z - P_1)y = y^T(I - P_1)y - y^T(I - P_Z)y$. A different interpretation of $y^T(I - P_Z)y$ is to see it as a residual sum of squares of a model with a design matrix Z. In this aspect, large SSA means that by including X, we see a large decrease in the residuals, i.e., the full model with Z fits the data better than the intercept only model.

```
library(tidyverse)
URL <- "https://campus.murraystate.edu/academic/faculty/cmecklin/STA565/wheat.txt"
wheat <- read.table(URL,header=TRUE)</pre>
```

str(wheat)

```
## 'data.frame': 30 obs. of 3 variables:
## $ variety : chr "A" "A" "A" "A" ...
## $ location: int 1 2 3 4 5 6 1 2 3 4 ...
## $ yield : num 35.3 31 32.7 36.8 37.2 33.1 33.7 32.2 31.4 32.7 ...
lm1 = lm(yield ~ 1, data = wheat)
lm2 = lm(yield ~ 1 + as.factor(location), data = wheat)
anova(lm1, lm2)
```

Q1-3: Conduct the f-test of treatment effect as learned in the class and check that the test statistics is the same.

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2. REANOVA and covariance structure

Consider a REANOVA model $\epsilon_{ij} \stackrel{iid}{\sim} N(0, \sigma^2), \quad E[a_j \epsilon_{ij}] = 0$

where μ , σ^2 and τ^2 are some unknown fixed parameter.

In the class, we saw that $E(y_{ij}) = \mu$, $V(y_{ij}) = \sigma^2 + \tau^2$ and $Cov(y_{1j}, y_{2j}) = \tau^2$. Also, since y_{ij} is a sum of Gausssian random variables, it itself also follows normal distribution. From this, we can write a joint distribution of $\mathbf{y}_i = [y_{1j}, \cdots, y_{nj}]^T$ for a group j as

Q2-1: Write down the covariance matrix Σ_j as follows:

$$(\Sigma_j)_{kl} = \begin{cases} ? & (k=l) & \text{Cov} (\forall i, \forall j) = \text{U(i)} - \text{C}^2 + \text{C}^2 \\ ? & (k \neq l) & \text{C} \end{cases}$$

Combining all y_j , we can rewrite the above REANOVA model in a matrix-vector form as

(or (dry (1, 1)) $y > N(\mu 1_{mn}, \Sigma)$

Q2-2: What is $Cov(y_{ij}, y_{ij})$? Using this, how can we write Σ ?

In contrast, if we change the model as

 $\begin{aligned} y_{ij} &= \mu + \alpha_j + \epsilon_{ij}, & i \in [n_j], \ j \in [m] \ (n_j = n \ \forall j) \\ \epsilon_{ij} &\stackrel{iid}{\sim} N(0, \sigma^2) & \text{v(4ij)} = \delta^2 \quad \text{Cov(4ij, 4ij)} > 0. \end{aligned}$

where α_j is also some unknown fixed parameter, then we have

 $y_j \sim N((\mu + \alpha_j)1), \sigma^2 I_n$

Q2-3: Write down the joint model for

The key takeaway is that, by adding another source of randomness a_i for group-wise variation, marginally we are modelling the covariance structure of y as Σ .