R09 - Analysis for Experiments with Two Factors Two-way ANOVA and Contrasts

STAT 587 (Engineering) Iowa State University

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Two factors

Consider the question of the affect of variety and density on yield under various experimental designs:

- Balanced, complete design
- Unbalanced, complete
- Incomplete

We will also consider the problem of finding the density that maximizes yield.

Data

An experiment was run on tomato plants to determine the effect of

- 3 different varieties (A,B,C) and
- 4 different planting densities (10,20,30,40)

on yield.

A balanced completely randomized design (CRD) with replication was used.

- complete: each treatment (variety × density) is represented
- balanced: each treatment has the same number of replicates
- randomized: treatment was randomly assigned to the plot
- replication: each treatment is represented more than once

This is also referred to as a full factorial or fully crossed design.

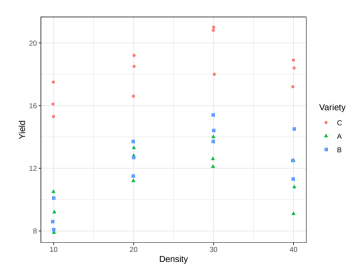
Hypotheses

- How does variety affect mean yield?
 - How is the mean yield for variety A different from B on average?
 - How is the mean yield for variety A different from B at a particular value for density?
- How does density affect mean yield?
 - How is the mean yield for density 10 different from density 20 on average?
 - How is the mean yield for density 10 different from density 20 at a particular value for variety?
- How does density affect yield differently for each variety?

For all of these questions, we want to know

- is there any effect and
- if yes, what is the magnitude and direction of the effect.

Confidence/credible intervals can answer these questions.



Summary statistics

```
# A tibble: 12 x 5
# Groups: Variety [3]
  Variety Density
                     n mean
  <fct>
            <int> <int> <dbl> <dbl>
 1 C
               10
                     3 16.3 1.11
               20
                     3 18.1 1.35
               30
                     3 19.9
                            1.68
               40
                     3 18.2
                            0.874
               10
                     3 9.2 1.30
               20
                     3 12.4 1.10
               30
                     3 12.9 0.985
                     3 10.8 1.7
               40
               10
                     3 8.93 1.04
10 B
               20
                     3 12.6 1.10
11 B
               30
                     3 14.5 0.854
12 B
               40
                     3 12.8 1.62
```

Two-way ANOVA

- Setup: Two categorical explanatory variables with I and J levels respectively
- Model:

$$Y_{ijk} \stackrel{ind}{\sim} N(\mu_{ij}, \sigma^2)$$

where Y_{ijk} is the

- kth observation at the
- ith level of variable 1 (variety) with i = 1, ..., I and the
- jth level of variable 2 (density) with $j=1,\ldots,J$.

Consider the models:

- Additive/Main effects: $\mu_{ij} = \mu + \nu_i + \delta_j$
- Cell-means: $\mu_{ij} = \mu + \nu_i + \delta_j + \gamma_{ij}$

	10	20	30	40
Α	μ_{11}	μ_{12}	μ_{13}	μ_{14}
В	μ_{21}	μ_{22}	μ_{23}	μ_{24}
С	μ_{31}	μ_{32}	μ_{33}	μ_{34}

As a regression model

- 1. Assign a reference level for both variety (C) and density (40).
- 2. Let V_i and D_i be the variety and density for observation i.
- 3. Build indicator variables, e.g. $I(V_i = A)$ and $I(D_i = 10)$.
- 4. The additive/main effects model:

$$\begin{array}{ll} \mu_i = & \beta_0 \\ & +\beta_1 \mathrm{I}(V_i = A) + \beta_2 \mathrm{I}(V_i = B) \\ & +\beta_3 \mathrm{I}(D_i = 10) + \beta_4 \mathrm{I}(D_i = 20) + \beta_5 \mathrm{I}(D_i = 30). \end{array}$$

5. The cell-means model:

$$\begin{split} \mu_i &= & \beta_0 \\ &+ \beta_1 \mathrm{I}(V_i = A) + \beta_2 \mathrm{I}(V_i = B) \\ &+ \beta_3 \mathrm{I}(D_i = 10) + \beta_4 \mathrm{I}(D_i = 20) + \beta_5 \mathrm{I}(D_i = 30) \\ &+ \beta_6 \mathrm{I}(V_i = A) \mathrm{I}(D_i = 10) + \beta_7 \mathrm{I}(V_i = A) \mathrm{I}(D_i = 20) + \beta_8 \mathrm{I}(V_i = A) \mathrm{I}(D_i = 30) \\ &+ \beta_9 \mathrm{I}(V_i = B) \mathrm{I}(D_i = 10) + \beta_{10} \mathrm{I}(V_i = B) \mathrm{I}(D_i = 20) + \beta_{11} \mathrm{I}(V_i = B) \mathrm{I}(D_i = 30) \end{split}$$

ANOVA Table

ANOVA Table - Additive/Main Effects model

Source	SS	df	MS	F	
Factor A	SSA	(I-1)	SSA/(I-1)	MSA/MSE	
Factor B	SSB	(J-1)	SSB/(J-1)	MSB/MSE	
Error	SSE	n-I-J+1	SSE/(n-I-J+1)		
Total	SST	n-1	,		

ANOVA Table - Cell-means model

Source	SS	df	MS	
Factor A	SSA	I-1	SSA/(I-1)	MSA/MSE
Factor B	SSB	J-1	SSB/(J-1)	MSB/MSE
Interaction AB	SSAB	(I-1)(J-1)	SSAB /(I-1)(J-1)	MSAB/MSE
Error	SSE	n-IJ	SSE/(n-IJ)	
Total	SST	n-1		

Two-way ANOVA in R

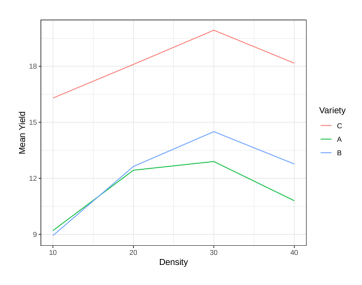
```
tomato$Density = factor(tomato$Density)
m = lm(Yield~Variety+Density, tomato)
drop1(m. test="F")
Single term deletions
Model:
Yield ~ Variety + Density
       Df Sum of Sq RSS AIC F value
                                           Pr(>F)
<none>
                    46.07 20.880
Variety 2 327.60 373.67 92.235 106.659 2.313e-14 ***
Density 3 86.69 132.76 52.980 18.816 4.690e-07 ***
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
m = lm(Yield~Varietv*Density. tomato)
drop1(m, scope = "Variety+Density+Variety:Density, test="F")
Single term deletions
Model:
Yield ~ Variety * Density
               Df Sum of Sa
                            RSS
                                    AIC F value
                                                    Pr(>F)
                            38.040 25.984
<none>
Variety 2 104.749 142.789 69.603 33.0438 1.278e-07 ***
Density
               3
                   19.809 57.849 35.076 4.1660
                                                   0.01648 *
Variety:Density 6
                     8.032 46.072 20.880 0.8445
                                                   0.54836
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

Additive vs cell-means

Opinions differ on whether to use an additive vs a cell-means model when the interaction is not significant. Remember that an insignificant test does not prove that there is no interaction.

	Additive	Cell-means
Interpretation	Direct	More complicated
Estimate of σ^2	Biased	Unbiased

We will continue using the cell-means model to answer the scientific questions of interest.



Two-way ANOVA in R

```
tomato$Density = factor(tomato$Density)
m = lm(Yield~Variety*Density, tomato)
anova(m)
Analysis of Variance Table
Response: Yield
               Df Sum Sq Mean Sq F value
                                             Pr(>F)
                2 327.60 163.799 103.3430 1.608e-12 ***
Variety
Density
                3 86.69 28.896
                                 18.2306 2.212e-06 ***
Variety:Density 6 8.03
                          1.339
                                  0.8445
                                            0.5484
Residuals
               24 38.04
                          1.585
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

Variety comparison

```
library(emmeans)
Warning: package 'emmeans' was built under R version 4.0.2
emmeans(m, pairwise~Variety)
$emmeans
 Variety emmean
                  SE df lower.CL upper.CL
          18.1 0.363 24
                           17.4 18.9
 С
         11.3 0.363 24 10.6
                                 12.1
          12.2 0.363 24 11.5
                                13.0
Results are averaged over the levels of: Density
Confidence level used: 0.95
$contrasts
                    SE df t.ratio p.value
 contrast estimate
 C - A
            6.792 0.514 24 13.214 <.0001
 C - B 5.917 0.514 24 11.512 <.0001
 A - B
           -0.875 0.514 24 -1.702 0.2249
Results are averaged over the levels of: Density
P value adjustment: tukev method for comparing a family of 3 estimates
```

Density comparison

```
emmeans(m, pairwise~Density)
$emmeans
Density emmean SE df lower.CL upper.CL
10
         11.5 0.42 24
                         10.6
                                12.3
        14.4 0.42 24 13.5
                               15.3
20
30
    15.8 0.42 24 14.9
                               16.6
40
        13.9 0.42 24
                        13.0
                               14.8
Results are averaged over the levels of: Variety
Confidence level used: 0.95
$contrasts
                 SE df t.ratio p.value
contrast estimate
10 - 20 -2.911 0.593 24 -4.905 0.0003
10 - 30 -4.300 0.593 24 -7.245 <.0001
10 - 40
          -2.433 0.593 24 -4.100 0.0022
20 - 30
          -1.389 0.593 24 -2.340 0.1169
20 - 40
        0.478 0.593 24 0.805 0.8514
30 - 40
        1.867 0.593 24 3.145 0.0213
Results are averaged over the levels of: Variety
P value adjustment: tukey method for comparing a family of 4 estimates
```

emmeans(m, pairwise~Variety*Density)

\$emmeans

variety	Density	emmean	DE	αı	TOWEL . CL	upper.cr
C	10	16.30	0.727	24	14.80	17.8
A	10	9.20	0.727	24	7.70	10.7
В	10	8.93	0.727	24	7.43	10.4
C	20	18.10	0.727	24	16.60	19.6
A	20	12.43	0.727	24	10.93	13.9
В	20	12.63	0.727	24	11.13	14.1
C	30	19.93	0.727	24	18.43	21.4
A	30	12.90	0.727	24	11.40	14.4
В	30	14.50	0.727	24	13.00	16.0
C	40	18.17	0.727	24	16.67	19.7
A	40	10.80	0.727	24	9.30	12.3
В	40	12.77	0.727	24	11.27	14.3

Confidence level used: 0.95

\$contrasts

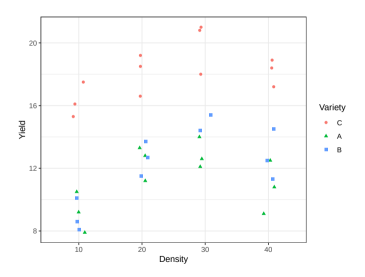
CC	onti	ras	Βt		estimate	SE	aI	t.ratio	p.varue
С	10	-	Α	10	7.1000	1.03	24	6.907	<.0001
С	10	-	В	10	7.3667	1.03	24	7.166	<.0001
С	10	-	С	20	-1.8000	1.03	24	-1.751	0.8276
С	10	-	Α	20	3.8667	1.03	24	3.762	0.0356
С	10	-	В	20	3.6667	1.03	24	3.567	0.0543
С	10	-	С	30	-3.6333	1.03	24	-3.535	0.0582
C	10	_	Α	30	3.4000	1.03	24	3.308	0.0932

Summary

- Use emmeans to answer questions of scientific interest.
- Check model assumptions
- Consider alternative models, e.g. treating density as continuous

Unbalanced design

Suppose for some reason that a variety B, density 30 sample was contaminated. Although you started with a balanced design, the data is now unbalanced. Fortunately, we can still use the tools we have used previously.



Summary statistics

```
# A tibble: 12 x 5
# Groups: Variety [3]
  Variety Density
                     n mean
  <fct>
          <fct> <int> <dbl> <dbl>
 1 C
          10
                     3 16.3 1.11
 2 C
          20
                     3 18.1 1.35
 3 C
          30
                     3 19.9
                            1.68
 4 C
          40
                     3 18.2 0.874
          10
                     3 9.2 1.30
          20
                     3 12.4 1.10
          30
                     3 12.9 0.985
                     3 10.8 1.7
          40
 9 B
          10
                     3 8.93 1.04
10 B
          20
                     3 12.6 1.10
11 B
          30
                     2 14.9 0.707
12 B
          40
                     3 12.8 1.62
```

Two-way ANOVA in R

```
m = lm(Yield~Variety*Density, tomato_unbalanced)
anova(m)
Analysis of Variance Table
Response: Yield
               Df Sum Sq Mean Sq F value
                                           Pr(>F)
Variety
                2 329.99 164.994 102.343 3.552e-12 ***
                3 84.45 28.150 17.461 3.947e-06 ***
Density
Variety:Density 6 8.80
                         1.467
                                  0.910
                                           0.5052
Residuals
               23 37.08
                         1.612
Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
```

Variety comparison

```
emmeans(m, pairwise Variety)
$emmeans
Variety emmean SE df lower.CL upper.CL
         18.1 0.367 23
                          17.4 18.9
        11.3 0.367 23 10.6
                               12.1
        12.3 0.389 23 11.5 13.1
Results are averaged over the levels of: Density
Confidence level used: 0.95
$contrasts
contrast estimate
                 SE df t.ratio p.value
C - A 6.792 0.518 23 13.102 <.0001
C - B 5.817 0.534 23 10.886 <.0001
Δ - B
       -0.975 0.534 23 -1.825 0.1839
Results are averaged over the levels of: Density
P value adjustment: tukey method for comparing a family of 3 estimates
```

Density comparison

```
emmeans(m, pairwise~Density)
$emmeans
Density emmean SE df lower.CL upper.CL
10
         11.5 0.423 23
                          10.6
                                12.4
        14.4 0.423 23
                        13.5
                               15.3
20
30
    15.9 0.457 23
                        15.0 16.9
40
        13.9 0.423 23
                        13.0 14.8
Results are averaged over the levels of: Variety
Confidence level used: 0.95
$contrasts
contrast estimate SE df t.ratio p.value
10 - 20 -2.911 0.599 23 -4.864 0.0004
10 - 30 -4.433 0.623 23 -7.116 <.0001
10 - 40
          -2.433 0.599 23 -4.065 0.0025
20 - 30
          -1.522 0.623 23 -2.443 0.0967
20 - 40
        0.478 0.599 23 0.798 0.8545
30 - 40
          2.000 0.623 23 3.210 0.0189
Results are averaged over the levels of: Variety
P value adjustment: tukey method for comparing a family of 4 estimates
```

Analysis in R

emmeans(m, pairwise~Variety*Density)

\$emmeans

Variety	Density	emmean	SE	df	lower.CL	upper.CL
C	10	16.30	0.733	23	14.78	17.8
A	10	9.20	0.733	23	7.68	10.7
В	10	8.93	0.733	23	7.42	10.4
C	20	18.10	0.733	23	16.58	19.6
A	20	12.43	0.733	23	10.92	13.9
В	20	12.63	0.733	23	11.12	14.1
C	30	19.93	0.733	23	18.42	21.4
A	30	12.90	0.733	23	11.38	14.4
В	30	14.90	0.898	23	13.04	16.8
C	40	18.17	0.733	23	16.65	19.7
A	40	10.80	0.733	23	9.28	12.3
В	40	12.77	0.733	23	11.25	14.3

Confidence level used: 0.95

\$contrasts

```
estimate SE df t.ratio p.value
contrast
C 10 - A 10 7.1000 1.04 23
                            6.849 < .0001
C 10 - B 10
           7.3667 1.04 23
                             7.106 < .0001
C 10 - C 20 -1.8000 1.04 23
                            -1.736 0.8341
```

C 10 - A 20 3.8667 1.04 23 3.730 0.0396 C 10 - B 20 3.6667 1.04 23 3.537 0.0597 C 10 - C 30 -3.6333 1.04 23 -3.505 0.0638

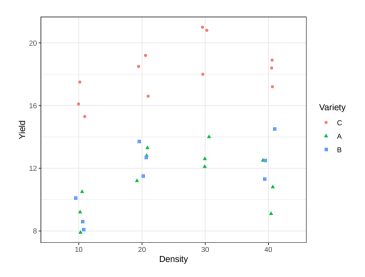
C 10 - A 30 3.4000 1.04 23 3.280 0.1008 C 10 - B 30 1.4000 1.16 23 1.208 0.9828 -1.801 0.8022 C 10 - C 40 -1.8667 1.04 23

Unbalanced Summary

The analysis can be completed just like the balanced design using emmeans to answer scientific questions of interest.

Incomplete design

Suppose none of the samples from variety B, density 30 were obtained. Now the analysis becomes more complicated.



Summary statistics

```
# A tibble: 11 x 5
# Groups: Variety [3]
  Variety Density
                     n mean
  <fct>
          <fct> <int> <dbl> <dbl>
                     3 16.3 1.11
 1 C
          10
 2 C
          20
                     3 18.1 1.35
          30
                     3 19.9
                            1.68
 4 C
          40
                     3 18.2 0.874
          10
                     3 9.2 1.30
          20
                     3 12.4 1.10
          30
                     3 12.9 0.985
                     3 10.8 1.7
          40
 9 B
          10
                     3 8.93 1.04
10 B
          20
                     3 12.6 1.10
11 B
          40
                     3 12.8 1.62
```

Treat as a One-way ANOVA

When the design is incomplete, use a one-way ANOVA combined with contrasts to answer questions of interest. For example, to compare the average difference between B and C, we want to only compare at densities 10, 20, and 40.

	10	20	30	40
Α	μ_{11}	μ_{12}	μ_{13}	μ_{14}
В	μ_{21}	μ_{22}		μ_{24}
C	μ_{31}	μ_{32}	μ_{33}	μ_{34}

Thus, the contrast is

$$\gamma = \frac{1}{3}(\mu_{31} + \mu_{32} + \mu_{34}) - \frac{1}{3}(\mu_{21} + \mu_{22} + \mu_{24})
= \frac{1}{3}(\mu_{31} + \mu_{32} + \mu_{34} - \mu_{21} - \mu_{22} - \mu_{24})$$

The Regression model

The regression model here considers variety-density combination as a single explanatory variable with 11 levels: A10, A20, A30, A40, B10, B20, B40, C10, C20, C30, and C40. Let C40 be the reference level. For observation i, let

- Y_i be the yield
- ullet V_i be the variety
- ullet D_i be the density

The model is then $Y_i \overset{ind}{\sim} N(\mu_i, \sigma^2)$ and

```
\begin{array}{l} \mu_i &= \beta_0 \\ + \beta_1 \mathrm{I}(V_i = A, D_i = 10) + \beta_2 \mathrm{I}(V_i = A, D_i = 20) + \beta_3 \mathrm{I}(V_i = A, D_i = 30) \\ + \beta_5 \mathrm{I}(V_i = B, D_i = 10) + \beta_6 \mathrm{I}(V_i = B, D_i = 20) \\ + \beta_8 \mathrm{I}(V_i = C, D_i = 10) + \beta_9 \mathrm{I}(V_i = C, D_i = 20) + \beta_{10} \mathrm{I}(V_i = C, D_i = 30) \end{array}
```

Two-way ANOVA in R

```
m <- lm(Yield ~ Variety*Density, data=tomato_incomplete)
anova(m)
Analysis of Variance Table
Response: Yield
               Df Sum Sq Mean Sq F value Pr(>F)
Variety
               2 347.38 173.691 104.462 5.868e-12 ***
                3 66.65 22.218 13.362 3.514e-05 ***
Density
Variety:Density 5 7.06
                         1.412
                                  0.849
                                             0.53
Residuals
               22 36.58
                         1.663
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

How can you tell the design is not complete?

One-way ANOVA in R

Contrasts

```
m = lm(Yield ~ VarietyDensity, tomato_incomplete)
em <- emmeans(m. ~ VarietvDensitv)
contrast(em, method = list(
# A10 A20 A30 A40 B10 B20 B40 C10 C20 C30 C40
"C-B" = c( 0, 0, 0, 0, -1, -1, -1, -1, 1, 0, 1)/3, "C-A" = c( -1, -1, -1, -1, 0, 0, 0, 1, 1, 1, 1)/4,
"B-A" = c(-1, -1, 0, -1, 1, 1, 1, 0, 0, 0, 0)/3) %>%
 confint
 contrast estimate SE df lower.CL upper.CL
 C-B
            6.078 0.608 22
                              4.817
                                    7.34
C-A
          6.792 0.526 22 5.700
                                     7.88
           0.633 0.608 22 -0.627
                                     1.89
 B-A
Confidence level used: 0.95
```

Analysis in R

```
m = lm(Yield~Variety:Density, tomato_incomplete)
emmeans (m. pairwise Variety: Density) # We could have used the VarietyDensity model, but this looks nicer
```

14.2

14.3

\$emmeans

```
Variety Density emmean
                          SE df lower.CL upper.CL
                16.30 0.744 22
                                   14.76
        10
                                            17.8
        10
                 9.20 0.744 22
                                   7.66
                                             10.7
        10
                 8.93 0.744 22
                                   7.39
                                             10.5
        20
                18.10 0.744 22
                                  16.56
                                             19.6
        20
                12.43 0.744 22
                                  10.89
                                             14.0
```

12.63 0.744 22 30 19.93 0.744 22 18.39 21.5 30 12.90 0.744 22 11.36 14.4 40 18.17 0.744 22 16.62 19.7 40 10.80 0.744 22 9.26 12.3 12.77 0.744 22

11.09

11.22

5.224 0.0012

Confidence level used: 0.95

C 10 - A 40 5.5000 1.05 22

20

40

\$contrasts

В

```
estimate SE df t.ratio p.value
contrast
C 10 - A 10 7.1000 1.05 22
                             6.744 < .0001
C 10 - B 10 7.3667 1.05 22
                             6.997 < .0001
C 10 - C 20 -1.8000 1.05 22
                            -1.710 0.8157
C 10 - A 20 3.8667 1.05 22
                            3.673 0.0407
C 10 - B 20 3.6667 1.05 22
                            3.483 0.0606
C 10 - C 30 -3.6333 1.05 22
                            -3.451 0.0646
C 10 - A 30 3.4000 1.05 22
                             3.229 0.1007
C 10 - C 40 -1.8667 1.05 22
                           -1.773 0.7829
```

- 10 - 10 - 1000 1 0F 00 - 0F0 0 0F01

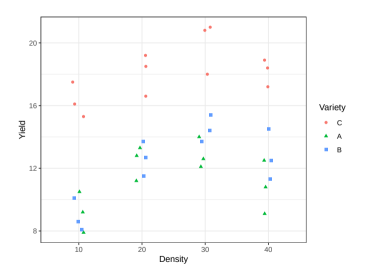
Summary

When dealing with an incomplete design, it is often easier to treat the analysis as a one-way ANOVA and use contrasts to answer scientific questions of interest.

Optimal yield

Now suppose you have the same data set, but your scientific question is different. Specifically, you are interested in choosing a variety-density combination that provides the optimal yield.

You can use the ANOVA analysis to choose from amongst the 3 varieties and one of the 4 densities, but there is no reason to believe that the optimal density will be one of those 4.



Modeling

Considering a single variety, if we assume a linear relationship between Yield (Y_i) and Density (D_i) then the maximum Yield will occur at either $-\infty$ or $+\infty$ which is unreasonable. The easiest way to have a maximum (or minimum) is to assume a quadratic relationship, e.g.

$$E[Y_i] = \mu_i = \beta_0 + \beta_1 D_i + \beta_2 D_i^2$$

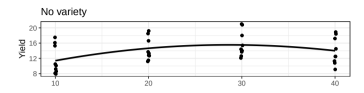
Now we can incorporate Variety (V_i) in many ways. Two options are parallel curves or completely independent curves.

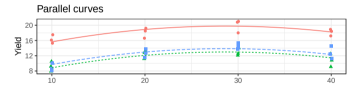
Parallel curves:

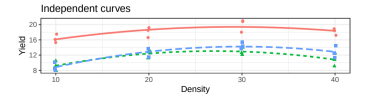
$$\mu_{i} = \beta_{0} + \beta_{1}D_{i} + \beta_{2}D_{i}^{2} + \beta_{3}I(V_{i} = A) + \beta_{4}I(V_{i} = B)$$

Independent curves:

$$\begin{array}{ll} \mu_i = & \beta_0 + \beta_1 D_i + \beta_2 D_i^2 \\ & + \beta_3 \mathrm{I}(V_i = A) + \beta_4 \mathrm{I}(V_i = B) \\ & + \beta_5 \mathrm{I}(V_i = A) D_i + \beta_6 \mathrm{I}(V_i = B) D_i \\ & + \beta_7 \mathrm{I}(V_i = A) D_i^2 + \beta_8 \mathrm{I}(V_i = B) D_i^2 \end{array}$$







Finding the maximum

For a particular variety, there will be an equation like

$$E[Y_i] = \mu_i = \beta_0 + \beta_1 D_i + \beta_2 D_i^2$$

where these β_1 and β_2 need not correspond to any particular β_1 and β_2 we have discussed thus far.

If $\beta_2 < 0$, then the quadratic curve has a maximum and it occurs at $-\beta_1/2\beta_2$.

No variety

```
Call:
lm(formula = Yield ~ Density + I(Density^2), data = tomato)
Residuals:
  Min
          10 Median
                             Max
-4.898 -2.721 -1.320 3.364 6.109
Coefficients:
             Estimate Std. Error t value Pr(>|t|)
(Intercept)
            5.744444 3.128242 1.836 0.0753 .
Density
             0.684111 0.285384
                                2.397 0.0223 *
I(Density^2) -0.011944  0.005618 -2.126
                                         0.0411 *
---
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 3.371 on 33 degrees of freedom
Multiple R-squared: 0.1854, Adjusted R-squared: 0.136
F-statistic: 3.755 on 2 and 33 DF, p-value: 0.03395
```

Parallel curves

```
Call:
lm(formula = Yield ~ Density + I(Density^2) + Variety, data = tomato)
Residuals:
  Min
          10 Median
                              Max
-2.3422 -0.9039 0.1744 0.8082 2.1828
Coefficients:
           Estimate Std. Error t value Pr(>|t|)
(Intercept)
           9.980556
                   1.184193 8.428 1.61e-09 ***
Density
           VarietvA
          -6.791667 0.504942 -13.450 1.76e-14 ***
VarietvB
                    0.504942 -11.718 6.39e-13 ***
          -5.916667
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 1.237 on 31 degrees of freedom
Multiple R-squared: 0.897.Adjusted R-squared: 0.8837
F-statistic: 67.48 on 4 and 31 DF, p-value: 7.469e-15
```

Independent curves

```
Call:
lm(formula = Yield ~ Density * Variety + I(Density^2) * Variety,
   data = tomato)
Residuals:
    Min
              10 Median
                                      Max
-2.04500 -0.82125 -0.01417 0.94000 1.71000
Coefficients:
                     Estimate Std. Error t value Pr(>|t|)
(Intercept)
                    11.808333
                              1.968364 5.999 2.12e-06 ***
Density
                    0.520167
                               0.179570 2.897 0.00739 **
VarietvA
                    -8.458333
                               2.783687 -3.039 0.00523 **
VarietvB
                    -9.733333
                                2.783687 -3.497 0.00165 **
I(Densitv^2)
                    -0.008917
                               0.003535 -2.522 0.01787 *
Density: VarietyA
                  0.199167
                               0.253951 0.784 0.43971
Density: VarietyB
                               0.253951 1.152 0.25924
                     0.292667
VarietvA:I(Densitv^2) -0.004417
                                0.005000
                                         -0.883 0.38482
VarietyB:I(Density^2) -0.004667
                                0.005000
                                         -0.933 0.35889
Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
Residual standard error: 1.225 on 27 degrees of freedom
Multiple R-squared: 0.912.Adjusted R-squared: 0.886
F-statistic: 34.99 on 8 and 27 DF, p-value: 2.678e-12
```

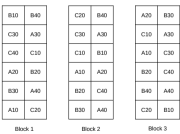
Completely randomized design (CRD)

This semester, we have assumed a completely randomized design. As an example, consider 36 plots and we are randomly assigning our variety-density combinations to the plots such that we have 3 reps of each combination. The result may look something like this

A20	A10	A20	B10	B10	A30
C10	C30	C30	C10	C20	A10
B30	B10	B20	B30	B40	B40
C40	B20	A10	C20	B30	A40
C30	B40	A30	C40	B20	C40
C10	C20	A40	A30	A20	A40

Complete randomized block design (RBD)

A randomized block design is appropriate when there is a nuisance factor that you want to control for. In our example, imagine you had 12 plots at 3 different locations and you expect these locations would have impact on yield. A randomized block design might look like this.



RBD Analysis

Generally, you will want to model a randomized block design using an additive model for the treatment and blocking factor. If you have the replication, you should test for an interaction. Let's compute the degrees of freedom for the ANOVA tables for this current design considering the variety-density combination as the treatment.

V+D+B		T+B		Cell-means	
Factor	df	Factor	df	Factor	df
Variety	2				
Density	3	Treatment	11	Treatment	11
Block	2	Block	2	Block	2
				Treatment x Block	22
Error	28	Error	22	Error	0
Total	35	Total	35	Total	35

The cell-means model does not have enough degrees of freedom to estimate the interaction because there is no replication of the treatment within a block.

Why block?

Consider a simple experiment with 2 blocks each with 3 experimental units and 3 treatments (A, B, C).

Blocked			Unblocked		
	А	O	A	С	
	В	В	В	В	
	A	С	С	А	
	Block 1	Block 2	Block 1	Block 2	

Let's consider 3 possible analyses:

- Blocked experiment using an additive model for treatment and block (RBD)
- Unblocked experiment using only treatment (CRD)
- Unblocked experiment using an additive model for treatment and block

Why block?

Now suppose, the true model is

$$\mu_{ij} = \mu + T_i + B_j$$

where $T_1=T_2=T_3$ and $B_1=0$ and $B_2=\delta$.

In the Blocked experiment using an additive model for treatment and block, the expected treatment differences to all be zero.

In the Unblocked design using only treatment, the expected difference between treatments is

$$\mu_C - \mu_B = \delta$$
 and $\mu_C - \mu_A = \delta/2$.

In the Unblocked design using an additive model for treatment and block, we would have an unbalanced design and it would be impossible to compare B and C.

Summary

Block what you can control; randomize what you cannot.