ANCOVA

Analysis of Covariance

History

Fisher introduced "analysis of covariance" in "Studies in Crop Variation. IV" (Eden and Fisher. 1927).

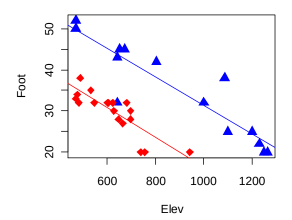
This is a method for factoring out the effects of conditions that are not part of the experimental design but which are there and can be measured.

T. Eden and R. A. Fisher. 1927. Studies in crop variation: IV. The experimental determination of the value of top dressings with cereals. The Journal of Agricultural Science 17: 548-562

Crested ibis nest site: footprint ~ landcover + elevation

Footprint	Elevation	Land cover	Nest site	
22	1230	11	金家村	
32	605	16	七氏山后	
33	471	16	纸坊街5组	
25	1200	11	3组1	
32	602	16	7组	
28	698	16 2组滚沟		
50	471	11	3组2	
52	471	11	蔡河4组	
38	490	16	3组龙泉	
28	648	16	牛河	
20	942	16	代家河	
20	1250	11	草坝4组	
34	477	16	4组云阳	
20	1264	11	华阳中学1号	
32	681	16	4组黄沟	
30	629	16	曹沟	
32	483	16	5组麻洞	
43	643	11	3组分会田	
32	643	11	3组堰岔弯	
30	698	16	后沟	
28	698	16	汤帽	
25	1100	11	草坝5组	
35	533	16	党河电站	
38	1087	11	高峰5组	
45	674	11	7组袁沟	
32	624	16	3组	
20	757	16	沙溪沟	
32	548	16	夏组	
45	653	11 1组石洽		
27	665	16 2组狗家沟		
32	624	16 戴家沟		
20	739	16	池塘岸	
32	1001	11	8组	
42	805	11	2组	

Code	Land cover	类型		
11	Evergreen Needleleaf forest	常绿针叶林		
12	Deciduous Needleleaf forest	落叶针叶林		
13	Evergreen Broadleaf forest	常绿阔叶林		
14	Deciduous Broadleaf forest	落叶阔叶林		
15	Mixed Froest	混交林		
16	Shrub	灌木		
21	Dense Grass	高覆盖度草地		
22	Grass with Moderate Dense	中覆盖度草地		
23	Sparse Grass	低覆盖度草地		
31	Farmlamd	耕地		
41	City and Urban Built-up	城市及建设用地		
51	Harsh Desert	荒漠		
52	Desert	沙漠		
53	Bare Rock	裸露岩石		
61	Wetland	湿地		
62	Ice and Snow	冰川雪被		
63	Waterbody	水体		





ANCOVA

Analysis of Covariance

- Combined use of ANOVA and Regression
 - Adjust for covariate by regressing covariate on the DV, then doing an ANOVA on the adjusted DV.

 $DV = IV \times CV$

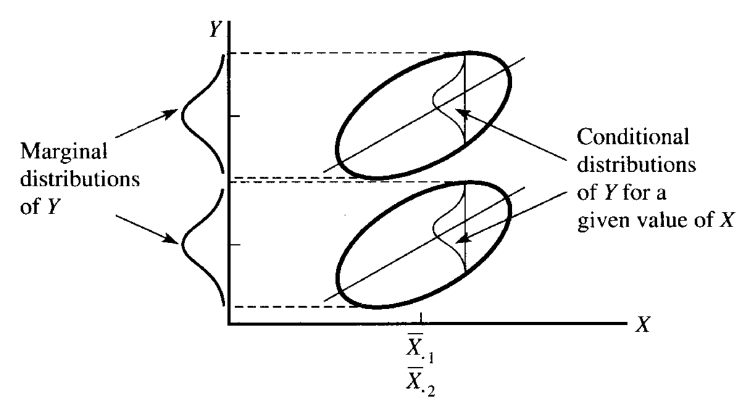
DV: dependent variable (y, continuous)

IV: independent variable (x, categorical)

CV: covariate variable (x, continuous)

Two groups have same X, different Y

$$DV = IV \times CV$$

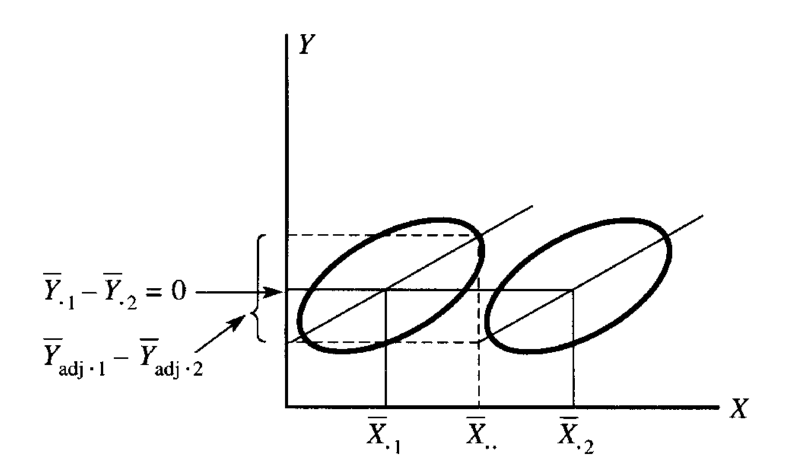


http://en.wikipedia.org/wiki/Marginal_distribution

In probability theory and statistics, the marginal distribution of a subset of a collection of random variables is the probability distribution of the variables contained in the subset. The term marginal variable is used to refer to those variables in the subset of variables being retained. These terms are dubbed "marginal" because they used to be found by summing values in a table along rows or columns, and writing the sum in the margins of the table. The distribution of the marginal variables (the marginal distribution) is obtained by marginalizing over the distribution of the variables being discarded, and the discarded variables are said to have been marginalized out.

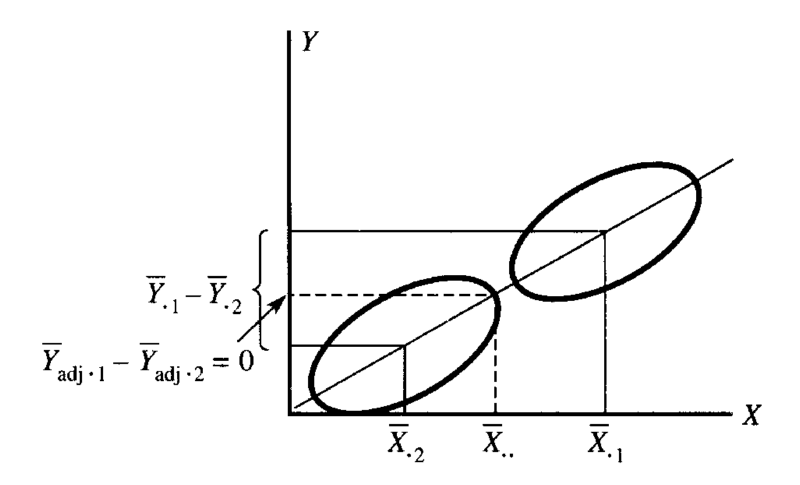
Two groups have different X with same Y, but the adjusted Y are different.

$$DV = IV \times CV$$



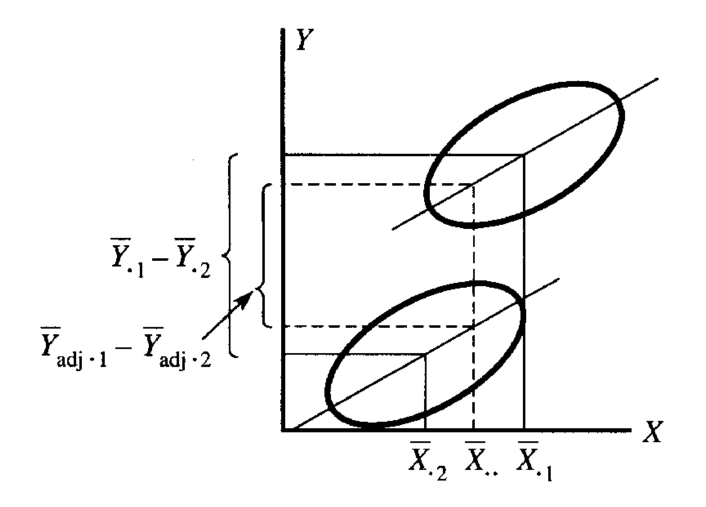
Two groups have different X, the adjusted Y are same

 $DV = IV \times CV$

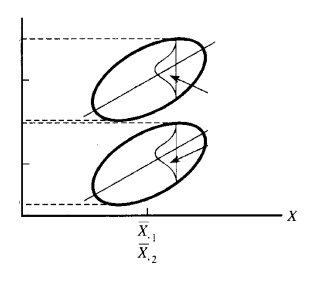


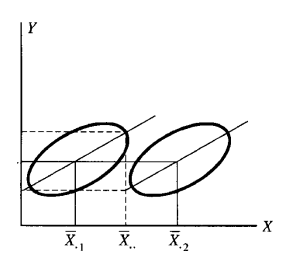
Two groups have different X, the adjusted Y are different

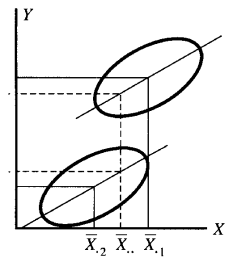
 $DV = IV \times CV$

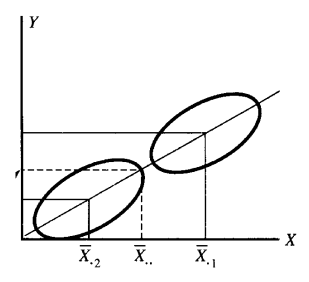


Real effect after adjustment





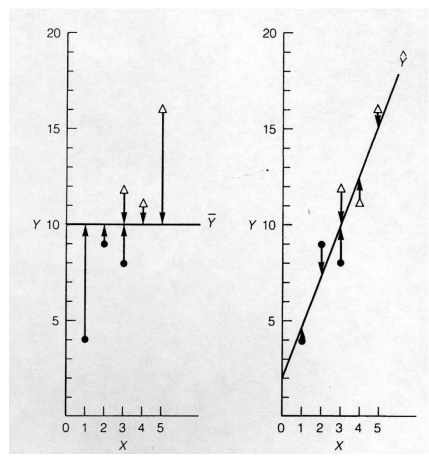




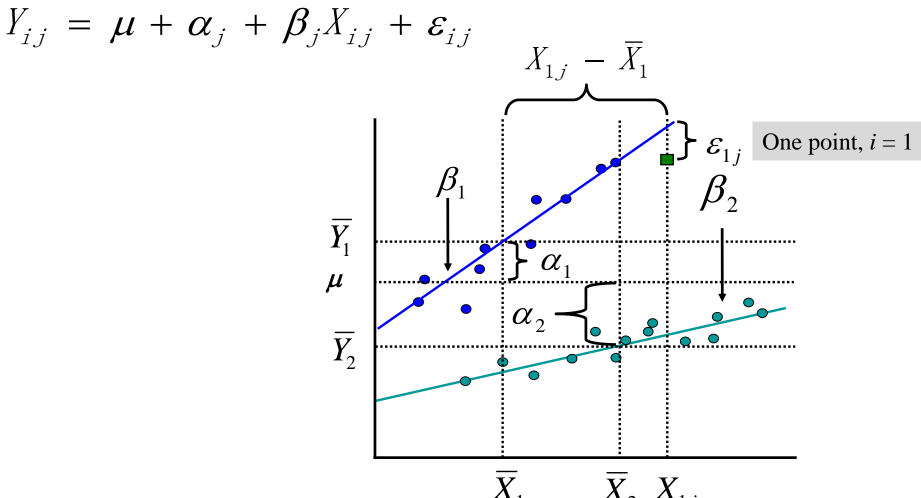
ANCOVA model

• Full model: $Y_{ij} = \mu + \alpha_j + \beta X_{ij} + \varepsilon_{ij}$

- ANCOVA models have a major advantage over ANOVA
 - ANCOVA models have the capability of making a different prediction for each individual, rather than having to make the same prediction for all individuals within a group
 - Predictions are a function of the score of the $\underline{covariate} X_{ii}$



Model effect



Assumptions

Normality of sampling distribution

- Normality on the DV at all of the levels of the IV(s) and the CV(s).
 - This cannot be shown unless you take multiple samples and form sampling distribution.

Homogeneity of Variance

 Equal variances on the DV at all of the levels of the IV(s) and the CV(s).

$$\sigma_1^2 \cong \sigma_2^2 \cong ... \sigma_p^2$$

- This is most important after adjustments have been made, but if you have it before adjustment you are likely to have it afterwards.
- If the assumption of homogeneity of variance fails, a more stringent alpha can be used (e.g. 0.01) or drop the variable from the analysis

Linearity

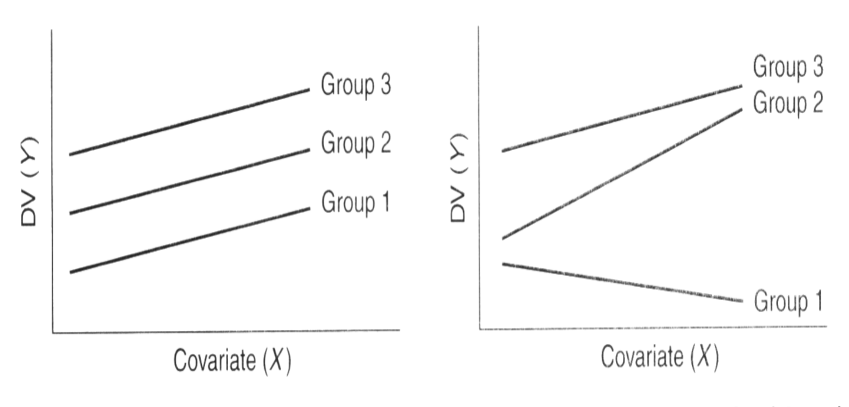
 It is assumed that each DV has a linear relationship with the CV and other CVs.

Check homogeneity of regression

 For many cases, the slopes of the lines predicting the DV from the CV are same for each level of the IV.

- In other words the regression coefficient (B) relating a
 CV to the DV should be the same for each group.
- In still other words, this means no IV by DV interaction.

Homogeneity of Regression



(a) Homogeneity of regression (slopes)

(b) Heterogeneity of regression (slopes)

Dealing with heterogeneous within-group regression slopes

When slopes are clearly heterogeneous

- First, if the slopes themselves are of primary interest, you can contrast slopes across treatment combinations.
- Second, if the treatment (group) effects are the main interest, you can choose certain values of covariate and compare groups at those specific values.
 - e.g. using the mean of X or the value of X for which the distance between regression lines has the most precision.
- Third, use mixed effect models to quantify the effects of IV and CV.

Reliability of Covariates

 It is assumed that each CV is measured without error (this is unrealistic).

 So it is recommended that CVs only be used when they meet a reliability of .8 or more.

Outlier

No outliers

- Test for univariate outliers on the DV and all of the CVs individually
- Test for multivariate outliers in the combined DV and CVs space.

Outlier

- A data point that is distinctly separate from the rest of the data. One definition of outlier is any data point more than 1.5 interquartile ranges (IQRs) below the first quartile or above the third quartile.
- Note: The IQR definition given here is widely used but is not the last word in determining whether a given number is an outlier.

Outlier

Example: For the data 2, 5, 6, 9, 12

minimum = 2

first quartile = 3.5

median = 6

third quartile = 10.5

maximum = 12

$$IQR = 10.5 - 3.5 = 7$$
, so $1.5 \cdot IQR = 10.5$.

To determine if there are outliers we must consider the numbers that are 1.5-IQR or 10.5 beyond the quartiles.

first quartile
$$-1.5 \cdot IQR = 3.5 - 10.5 = -7$$

third quartile $+1.5 \cdot IQR = 10.5 + 10.5 = 21$

Since none of the data are outside the interval from –7 to 21, there are no outliers.

No Multicollinearity/Singularity

 If a CV is highly related to another CV (e.g. at a correlation of .5 or more) then it should not be used to adjust the DV over the other CV.

 One or the other should be removed since they are statistically redundant.

- The problem here is that with unequal samples it is unclear how to calculate the marginal mean.
- Another problem is that the variances then start to overlap one another forcing the within plus between variances to be larger than the total variance.

Biostatistics

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Unaqual cample sizes

data and number of cases					
# Tensile strength in paper manufacturing					
Y < -c(30,35,37,36,34,41,38,42,29,26,33,36,					
28,32,40,41,31,36,42,40,31,30,32,40,					
31,37,41,40,35,40,39,44,32,34,39,45)					
block <- gl(3, 12, 36) # Three blocks					
A <- gl(3, 4, 36) # Three pulp preparation methods					
D 1/4 1 26 // E 1/60					

B <- gl(4, 1, 36) # Four different temperatures Dat <- data.frame(Y, block, A, B)

Dat = Dat[-c(1:3),] # make data unbalanced

summary(aov($Y \sim A*B$, data = Dat)) # type I sum of square Anova(mod <- lm(Y \sim A*B, data = Dat), type = "II") # library(car) Anova(mod <- lm(Y \sim A*B, data = Dat), type = "III")

[7,] [8,] [9,] [10,][11,] [12,] [13,]

[14,]

[15,]

[16,]

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[18,]

[19,]

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[21,]

[22,]

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[24,]

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[27,]

[28,]

[1,]

[2,] [3,]

[4,]

[5,] [6,]

Υ

30

35

37

36 34

41 38

42

29

26

36

42

40

block

31 30 32 2 40 31 3 3 37

3 41 3 40 3 35 40 3

[29,] [30,][31,] 39 3 [32,] 44 3 [33,] 32 3 3 [34,] 34 39 3 [35,] 45 3 [36,]

ANOVA Table (Type I tests)							
	Df	Sum Sq	Mean Sq	F value	Р		
A	2	126	63.02	7.484	0.00352		
В	3	398.9	132.96	15.79	1.33E-05		
A:B	6	81.8	13.63	1.618	0.19147		
Residuals	21	176.8	8.42				
ANOVA Table (Type II tests)							
	Df	Sum Sq	Mean Sq	F value	Р		
A	2	127.5		7.5709	0.003343		
В	3	398.88		15.7898	1.33E-05		
A:B	6	81.76		1.6182	0.191471		
Residuals	21	176.83					
ANOVA Table (Type III tests)							
	Df	Sum Sq	Mean Sq	F value	P		
(Intercept)	1	1740.5		206.6946	2.43E-12		
A	2	20.04		1.19	0.323909		
В	3	156.06		6.1775	0.003539		
A:B	6	81.76		1.6182	0.191471		
Residuals	21	176.83					

- Do not use type 1 sums of squares
- Type 1 sums of squares assumes that the difference in number of subjects is meaningful and gives more weight to the values from larger groups

- Use the type 3 (III) sums of square
- The type 3 sums of square assumes that the data was supposed to be complete, and the difference in the number of subjects is not meaningful
 - Acts like standard multiple regression. Each main effect and interaction is assessed after all other main effects, interactions and covariates are controlled
 - Treats all groups the same small group is weighted equally as a large group (sometimes called the unweighted approach)
 - Are preferable in most cases since they correspond to the variation attributable to an effect after correcting for any other effects in the model.
 They are unaffected by the frequency of observations.

- Number of cases required depends on the number needed to reach appropriate level of power.
- Unbalanced experimental design needs more cases.

Advantages of ANCOVA

- Adjusts for pre-treatment differences between groups.
 - If pre-treatment differences exist because groups were not randomly formed, then ANCOVA will eliminate the bias that may exist with non-random assignment.
- More Power due to decreased variance that must be explained by the IV (smaller error term in the F test).
 - Covariate "accounts for" some of the variance in the DV variance.

 Just like ANOVA the total variance can be separated into within and between groups variance:

$$\sum_{i} \sum_{j} (Y_{ij} - GM_{(y)})^{2} = n \sum_{j} (\overline{Y}_{j} - GM_{(y)})^{2} + \sum_{i} \sum_{j} (Y_{ij} - \overline{Y}_{j})^{2}$$

$$SS_{Total(y)} = SS_{bg(y)} + SS_{wg(y)}$$

 But in ANCOVA you also have a partitioning of the variance in each CV:

$$\sum_{i} \sum_{j} (Z_{ij} - GM_{(z)})^{2} = n \sum_{j} (\bar{Z}_{j} - GM_{(z)})^{2} + \sum_{i} \sum_{j} (Z_{ij} - \bar{Z}_{j})^{2}$$

$$SS_{Total(z)} = SS_{bg(z)} + SS_{wg(z)}$$

 And also a partitioning of the covariation between them (DV and CV):

$$SP_{Total} = SP_{bg} + SP_{wg}$$

 This covariation is used to adjust the between and within groups sums of squares:

$$SS'_{bg(y)} = SS_{bg(y)} - \left[\frac{\left(SP_{bg} + SP_{wg} \right)^{2}}{SS_{bg(z)} + SS_{wg(z)}} - \frac{\left(SP_{wg} \right)^{2}}{SS_{wg(z)}} \right]$$

$$SS'_{wg(y)} = SS_{wg(y)} - \frac{\left(SP_{wg}\right)^2}{SS_{wg(z)}}$$

The adjustment made to the between group scores in last slide can also be conceptualized as:

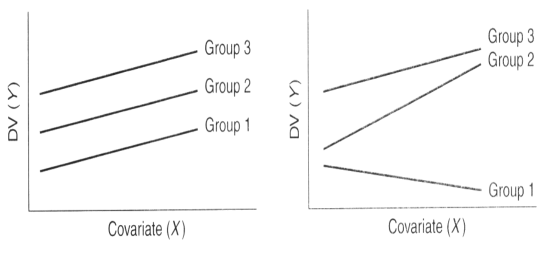
the adjustment is obtained by

- taking each individuals deviation around Y grand mean (before adjustment)
- subtracting from it each person's deviation around the Z grand mean, weighted by the relationship between the two

$$(Y - \overline{Y}_j) = (Y - GM_{(y)}) - \beta_{y \cdot z} (Z - GM_{(z)})$$

Degree of freedom

- Each regression coefficient (slope) we need to estimate (one for every CV) eats up a degree of freedom (if slopes are different).
- This accounts for the smaller error degrees of freedom when compared to regular ANOVA.



(b) Heterogeneity of regression (slopes)

General issues

The basic ANCOVA tests null hypotheses about adjusted factor effects, where the linear relationship between the covariate and the response variable (Y) is taken into account.

These means at all levels of factors are adjusted to the overall mean value for the covariate by the relationship between Y and the covariate.

Xinhai Li

General issues

Homogeneity of within-group regression slopes is tested by including factor by covariate interaction terms in a preliminary model.

In complex models with many categorical explanatory variables, homogeneity of slopes can be checked by combining all factors by the covariate terms into a single interaction term.

Alternatively, homogeneity of slopes may be better tested separately for each component of the analysis.

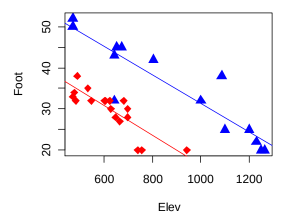
Alternatives to ANCOVA

- When CV and DV have a non-linear relationship
 - Use CV to group similar observations together into blocks. Each block is then used as levels of a BG IV that is crossed with the other BG IV that you are interested in.
 - Blocking may be the best alternative, because it doesn't have the special assumptions of ANCOVA, and it can capture non-linear relationships between CV and DV where ANCOVA only deals with linear relationships.

Example (crested ibis nest site): footprint ~ landcover + elevation

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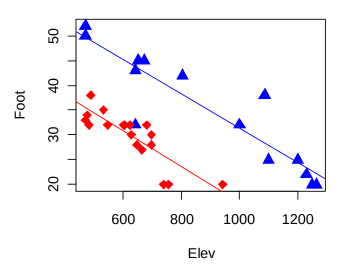


R code – plot data

```
#ANCOVA
#Human footprint index, elevation, and landcover
ibis <- read.csv('d:/ibis.csv',header=T)
ibis$Landcover[ibis$Landcover == 11] <- 1 #Forest
ibis$Landcover[ibis$Landcover == 16] <- 2 #Shrub
ibis$Landcover <- as.factor(ibis$Landcover)

attach(ibis)
plot(Elev, Foot, pch=16+as.numeric(Landcover), col=c('blue', 'red')[as.numeric(Landcover)],cex=1.5)
abline(lm(Foot[Landcover==1]~Elev[Landcover==1]),lty=1, col='blue')
abline(lm(Foot[Landcover==2]~Elev[Landcover==2]),lty=1, col='red')</pre>
```

```
Foot Elev Landcover Nestsite
22 1230 11 金家村
32 605 16 七氏山后
```



Compare means

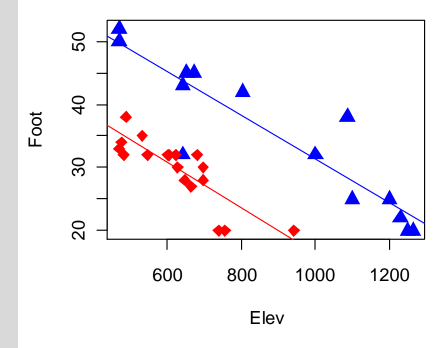
```
options (digits=3)
tapply(Foot, Landcover, mean)
t.test(Foot ~ Landcover)
```

```
> tapply(Foot, Landcover, mean)
    1    2
35.1    29.8
```

> t.test(Foot ~ Landcover)

Welch Two Sample t-test

```
data: Foot by Landcover
t = 1.65, df = 16.5, p-value = 0.118
alternative hypothesis:
true difference in means is not equal to 0
95 percent confidence interval:
-1.5 12.1
sample estimates:
mean in group 1 mean in group 2
35.1
29.8
```



summary(ancova)

anova1 <- Im(Foot~Landcover)
summary(anova1)</pre>

```
Call:
```

Im(formula = Foot ~ Landcover)

Residuals:

Min 1Q Median 3Q Max -15.07 -3.07 2.25 4.00 16.93

Coefficients:

Estimate Std. Error t value Pr(>|t|) (Intercept) 35.07 2.18 16.07 <2e-16 Landcover2 -5.32 2.85 -1.87 0.071 .

Residual standard error: 8.17 on 32 degrees of freedom Multiple R-squared: 0.0985, Adjusted R-squared: 0.0703 F-statistic: 3.5 on 1 and 32 DF, p-value: 0.0707 ancova <- Im(Foot~Landcover*Elev)
summary(ancova)</pre>

Call:

Im(formula = Foot ~ Landcover * Elev)

Residuals:

Min 1Q Median 3Q Max -11.702 -1.475 0.634 1.939 9.670

Coefficients:

Estimate Std. Error t value Pr(>|t|)
(Intercept) 65.96 3.24635 20.32 < 2e-16
Landcover2 -13.41 5.76795 -2.33 0.027
Elev -0.034 0.00346 -10.00 4.6e-11
Landcover2:Elev -0.00153 0.00821 -0.19 0.853

Residual standard error: 3.73 on 30 degrees of freedom Multiple R-squared: 0.824, Adjusted R-squared: 0.806 F-statistic: 46.8 on 3 and 30 DF, p-value: 1.99e-11

ANOVA table

anova(ancova)

Analysis of Varia	Analysis of Variance Table							
Response: Foot	Response: Foot							
	Df	Sum Sq	Mean Sq	F value	Pr(>F)			
Landcover	1	233	233	16.77	0.00029 ***			
Elev	1	1717	1717	123.52	3.7e-12 ***			
Landcover:Elev	1	0	0	0.03	0.85316			
Residuals	30	417	14					

Update model

```
ancova2 = update(ancova, ~. -Landcover:Elev)
anova(ancova, ancova2)
```

```
Analysis of Variance Table

Model 1: Foot ~ Landcover * Elev
Model 2: Foot ~ Landcover + Elev

Res.Df RSS Df Sum of Sq F Pr(>F)
1 30 417
2 31 418 -1 -0.485 0.03 0.85
```

Compare with ANOVA

```
ancova3 = update(ancova2, ~. -Elev)
anova(ancova2, ancova3)
```

```
Analysis of Variance Table

Model 1: Foot ~ Landcover + Elev

Model 2: Foot ~ Landcover

Res.Df RSS Df Sum of Sq F Pr(>F)

1 31 418

2 32 2135 -1 -1717 127 1.6e-12 ***
```

Model selection

```
step(ancova)
                    Start: AIC=93.2
                    Foot ~ Landcover * Elev
                                   Df Sum of Sq RSS AIC
                    - Landcover: Elev 1
                                        0.485
                                                  418 91.3
                                                  417 93.2
                    <none>
                    Step: AIC=91.3
                    Foot ~ Landcover + Elev
                                   Sum of Sq RSS AIC
                               Df
                                               418 91.3
                    <none>
                    - Landcover 1
                                     1229 1646 135.9
                                        1717 2135 144.8
                    - Elev
                    Call:
                    Im(formula = Foot ~ Landcover + Elev)
                    Coefficients:
                    (Intercept) Landcover2
                                              Elev
                      66.2053
                                -14.4522
                                           -0.0349
```

Akaike information criterion (AIC)

The Akaike information criterion is a measure of the relative goodness of fit of a statistical model. It was developed by Hirotsugu Akaike in 1974.

AIC provides a means for comparison among models, a tool for model selection.

$$AIC = 2k - 2\ln(L)$$

where k is the number of parameters in the statistical model, and L is the maximized value of the likelihood function for the estimated model

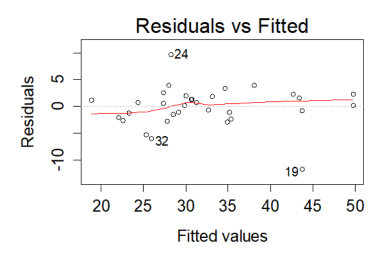
AICc is AIC with a correction for finite sample sizes

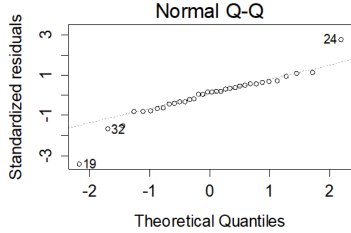
$$AIC_C = AIC + \frac{2k(k+1)}{n-k-1}$$

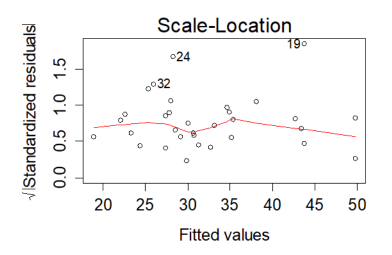
where k denotes the number of model parameters. Thus, AICc is AIC with a greater penalty for extra parameters.

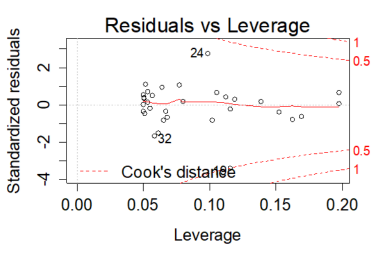
Model check

ancova.final <- step(ancova)
plot(ancova.final)</pre>









56 1 11 17 93

post.score ~ class.type * pre.score * IQ

/* Data for the ANCOVA example (the Trigonometry scores) */

ID	Class type	pre score	post score	IQ
1	1	3	10	122
2	2	24	34	129
3	3	10	21	114
4	1	5	10	121
5	2	18	27	114
6	3	3	18	114

ancova = Im(post.score ~ class.type * pre.score * IQ)

Results

```
scores$class = factor(scores$class)
ancova = Im(post ~ class * pre * IQ,
data = scores)
summary(ancova)
```

Call:

Im(formula = post ~ class * pre * IQ, data = scores)

Residuals:

Min 1Q Median 3Q Max -10.771 -2.900 -0.288 3.055 8.153

Coefficients:

	Estimate	Std. Error	t value	Pr(> t)
(Intercept)	-1.12485	43.56642	-0.03	0.98
class2	-63.85851	54.87780	-1.16	0.25
class3	-29.91929	60.51750	-0.49	0.62
pre	0.50020	4.48630	0.11	0.91
IQ	0.07686	0.35743	0.22	0.83
class2:pre	4.60531	5.57265	0.83	0.41
class3:pre	1.75546	7.85225	0.22	0.82
class2:IQ	0.53693	0.44687	1.20	0.24
class3:IQ	0.34119	0.52100	0.65	0.52
pre:IQ	0.00471	0.03702	0.13	0.90
class2:pre:I0	2 -0.04029	0.04551	-0.89	0.38
class3:pre:I0	Q -0.02008	0.06813	-0.29	0.77

Residual standard error: 4.66 on 44 degrees of freedom Multiple R-squared: 0.484, Adjusted R-squared: 0.355 F-statistic: 3.76 on 11 and 44 DF, p-value: 0.000793

Model selection

```
ancova2 = update(ancova, ~. -class : pre : IQ)
summary(ancova2)
ancova3 = update(ancova2, ~. -class : pre)
summary(ancova3)
ancova4 = update(ancova3, ~. -class : IQ)
summary(ancova4)
ancova5 = update(ancova4, ~. -pre : IQ)
summary(ancova5)
```

```
Call:
Im(formula = post \sim class + pre + IQ, data = scores)
Coefficients:
          Estimate Std. Error t value Pr(>|t|)
(Intercept) -14.8759 8.8927 -1.67
                                   0.1005
class2 -1.4026 1.4889 -0.94
                                   0.3506
                                   0.0024 **
class3 4.9870 1.5609 3.20
                                   1e-05 ***
                  0.1596 4.89
       0.7802
pre
                                   0.0056 **
IQ
          0.2129
                    0.0736 2.89
```

Model selection

step(ancova)

```
Start: AIC=183
post ~ class * pre * IQ
             Df Sum of Sq RSS AIC
- class:pre:IQ 2
                  17.3
                           974 180
                           957 183
<none>
Step: AIC=180
post ~ class + pre + IQ + class:pre + class:IQ + pre:IQ
         Df Sum of Sq RSS AIC
- class:IQ 2
               22.9
                        997 177
- class:pre 2
               37.7
                      1012 178
- pre:IQ
        1
               24.0
                        998 179
<none>
                        974 180
Step: AIC=177
post ~ class + pre + IQ + class:pre + pre:IQ
         Df Sum of Sq RSS AIC
               44.0
- class:pre 2
                    1041 176
                       997 177
<none>
-pre:IQ
        1
              37.9
                      1035 177
```

```
Step: AIC=176
post ~ class + pre + IQ + pre:IQ
       Df Sum of Sq RSS AIC
             30
                    1071 175
- pre:IQ 1
                    1041 176
<none>
                    1396 188
- class 2
             355
Step: AIC=175
post ~ class + pre + IQ
       Df Sum of Sq RSS AIC
                    1071 175
<none>
- IQ
       1 176
                   1247 182
- class 2 334 1405 186
- pre 1
            502
                    1574 195
Call:
Im(formula = post ~ class + pre + IQ, data = scores)
Coefficients:
(Intercept)
            class2
                      class3
                                          IQ
                                 pre
  -14.876
            -1.403
                      4.987
                                0.780
                                         0.213
```

Variance partitioning table

Source	DF	Type I SS	Mean Square	F Value Pr > F
CLASSTYPE	2	115.6381579	57.8190789	2.75 0.0733
PRE	1	493.3922076	493.3922076	23.49 < .0001
IQ	1	175.7215915	175.7215915	8.36 0.0056
Source	DF	Type III SS	Mean Square	F Value Pr > F
CLASSTYPE	2	333.6317170	166.8158585	7.94 0.0010
PRE	1	502.1888091	502.1888091	23.91 <.0001
IQ	1	175.7215915	175.7215915	8.36 0.0056

Childhood sexual abuse

```
# Book: Linear models with R (Faraway 2009)
                                                            15
# Effects of childhood sexual abuse on adult females reported in
# Rodriguez et al. (1997):45 women treated at a clinic,
# who reported childhood sexual abuse (csa), were measured for
                                                            10
                                                                                     9
# post-traumatic stress disorder (ptsd) and
                                                                                   ptsd
# childhood physical abuse (cpa)
                                                            2
library(faraway)
                                                                                     0
data(sexab)
by(sexab, sexab$csa, summary)
                                                                 Abused
                                                                        NotAbused
plot(ptsd ~ csa, sexab)
                                                                      csa
                                                                                                сра
plot(ptsd ~ cpa, pch = as.numeric(sexab$csa),
```

col = as.numeric(sexab\$csa), sexab)

m1 <- Im (ptsd ~ cpa+csa+cpa:csa, sexab) summary (m1) model.matrix (m1)

```
сра
          ptsd
                    csa
2.04786
         9.71365
                   Abused
0.83895
         6.16933
                   Abused
-0.24139 15.15926
                   Abused
-1.11461 11.31277
                   Abused
2.01468 9.95384
                   Abused
6.71131 9.83884
                  Abused
```

m2 <- Im (ptsd ~ cpa+csa, sexab) summary (m2)

Fitted

Residuals

change the reference level

summary (m3)

m3 <- Im (ptsd ~ cpa+csa, sexab)

Childhood sexual abuse

```
call:
lm(formula = ptsd \sim cpa + csa, data = sexab)
Residuals:
   Min
            10 Median
                                   Max
-8.1567 -2.3643 -0.1533
                       2.1466 7.1417
Coefficients:
           Estimate Std. Error t value Pr(>|t|)
                     0.6293
                                6.317
                                       1.87e-08 ***
(Intercept)
             3.9753
                       0.1716 3.209
             0.5506
                                       0.00198
                                                **
cpa
csaAbused
                       0.8219
                                7.632
                                       6.91e-11
             6.2728
                                                ***
                                                            сра
plot(ptsd~cpa, pch=as.numeric(sexab$csa),
    col=as.numeric(sexab$csa), sexab)
 abline (3.9753, 0.5506, col = 'red') # not abused
 abline (10.248, 0.5506) # abused, 10.248 = 3.9753 + 6.2728
plot (fitted (m2), residuals (m2), pch=as.numeric(sexab$csa),
     xlab= "Fitted", ylab="Residuals")
```

```
sexab$csa <- relevel (sexab$csa, ref="NotAbused") # ref="Abused"
```

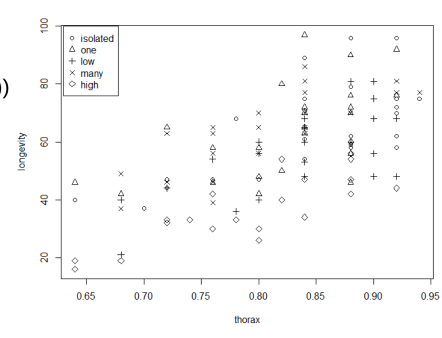
Sexual activity and the life span of male fruitflies

The data for this example come from a study on the sexual activity and the life span of # male fruitflies by Partridge and Farquhar (1981):125 fruitflies were divided randomly # into five groups of 25 each. The response was the longevity of the fruitfly in days. One # group was kept solitary, while another was kept individually with a virgin female each # day. Another group was given eight virgin females per day. As an additional control, the # fourth and fifth groups were kept with one or eight pregnant females per day. Pregnant # fruitflies will not mate. The thorax length of each male was measured as this was known # to affect longevity. The five groups are labeled many, isolated, one, low and high # respectively. The purpose of the analysis is to determine the difference between the five # groups if any.

```
library(faraway)
data (fruitfly)
plot (longevity ~ thorax, fruitfly, pch=unclass (activity))
legend (0.63, 100, levels (fruitfly$activity), pch=1:5)

g <- Im (longevity ~ thorax*activity, fruitfly)
summary (g)
model.matrix(g)
anova(g)

gb <- Im (longevity ~ thorax+activity, fruitfly)
drop1 (gb, test="F") # drop one term, using F test</pre>
```



Sexual activity and the life span of male fruitflies

summary (g)

```
Im(formula = longevity ~ thorax * activity, data = fruitfly)
Coefficients:
                   Estimate
                              Std. Error
                                          t value
                                                 Pr(>|t|)
                                                   0.023 *
                   -50.2420
                              21.8012
                                          -2.305
(Intercept)
thorax
                  136.1268
                            25.9517
                                          5.245
                                                  7.27e-07 ***
activityone
                    6.5172
                              33.8708
                                          0.192
                                                  0.848
activitylow
                    -7.7501
                              33.9690
                                         -0.228
                                                  0.820
                                          -0.035
activitymany
                    -1.1394
                              32.5298
                                                  0.972
                                                                                       drop1 (gb, test="F")
activityhigh
                   -11.0380
                              31.2866
                                          -0.353
                                                  0.725
                                                            Single term deletions
thorax:activityone -4.6771
                            40.6518
                                          -0.115
                                                  0.909
thorax:activitylow
                                                  0.983
                    0.8743 40.4253
                                          0.022
                                                            Model:
                                                   0.868
                              39.3600
                                           0.166
thorax:activitymany
                    6.5478
                                                            longevity ~ thorax + activity
thorax:activityhigh -11.1268
                              38.1200
                                          -0.292
                                                   0.771
                                                                 Df Sum of Sq RSS AIC F value
                                                                                                  Pr(>F)
                                                                            13107 589.92
                                                            <none>
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
                                                                  1 12368.4 25476 670.32 111.348 < 2.2e-16 ***
                                                            thorax
                                                            activity 4 9634.6 22742 650.25 21.684 1.974e-13 ***
Residual standard error: 10.71 on 114 degrees of freedom
Multiple R-squared: 0.6534, Adjusted R-squared: 0.626
                                                            Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
F-statistic: 23.88 on 9 and 114 DF, p-value: < 2.2e-16
```

Missing data

library(faraway) data(chmiss) # Chicago insurance dataset head(chmiss)

	race	fire	theft	age	involact	income
60626	10	6.2	29	60.4	NA	11.744
60640	22.2	9.5	44	76.5	0.1	9.323
60613	19.6	10.5	36	NA	1.2	9.948
60657	17.3	7.7	37	NA	0.5	10.656
60614	24.5	8.6	53	81.4	0.7	9.73
60610	54	34.1	68	52.6	0.3	8.231

model <- Im(involact ~ ., chmiss) **summary**(model)

Coefficients:

Estimate Std. Error t value Pr(>|t|)(Intercept) -1.116483 0.605761 -1.843 0.079475 0.010487 0.003128 3.352 0.003018 ** race 0.043876 0.010319 4.252 0.000356 *** fire theft -0.017220 0.005900 -2.918 0.008215 ** 0.009377 0.003494 2.684 0.013904 * age income 0.068701 0.042156 1.630 0.118077

Residual standard error: 0.3382 on 21 degrees of freedom (20 observations deleted due to missingness) Multiple R-squared: 0.7911, Adjusted R-squared: 0.7414 F-statistic: 15.91 on 5 and 21 DF, p-value: 1.594e-06

Xinhai Li

Replacing missing data with mean

Any case with at least one missing value is omitted from the regression. There are now only 21 degrees of freedom - almost half the data is lost. We can fill in the missing values by their variable means as in:

```
cmeans <- apply (chmiss, 2, mean, na.rm=T)
cmeans</pre>
```

```
race fire theft age involact income 35.60930 11.42444 32.65116 59.96905 0.64773 10.73587
```

```
mchm <- chmiss for (i in c(1, 2, 3, 4, 6)) mchm[is.na (chmiss[,i]), i] <- cmeans[i]
```

```
model <- Im(involact ~ ., mchm) summary(model)
```

Residual standard error: 0.3841 on 38 degrees of freedom (3 observations deleted due to missingness) Multiple R-squared: 0.682, Adjusted R-squared: 0.6401 F-statistic: 16.3 on 5 and 38 DF, p-value: 1.409e-08

Mixed effects ANCOVA

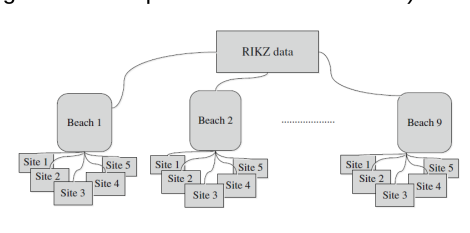


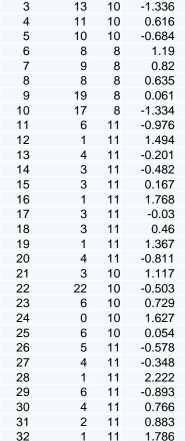
Mixed effects modelling for ANCOVA

Lecture 10. Analysis of covariance

In each inter-tidal area (denoted by 'beach'), five samples were taken, and the macro-fauna and abiotic variables were measured.

Species richness (the number of different species) can be explained by NAP (Normal Amsterdams Peil, the height of a sampling station compared to mean tidal level) and beaches.





11

11

11

10

10

10

10

10

10

10

10

10

10

1.375

-0.06

0.367

1.671

-0.375

-1.005

0.17

2.052

-0.356

0.094

-0.002

2.255

0.865

33

34

35

36

37

38

39

40

42

43

45

10

10

10

Sam Richness

NAP Beach

0.045

-1.036

Biostatistics

Xinhai Li

The random intercept model

$$R_{ij} = \alpha + \beta_1 \times NAP_{ij} + \beta_2 \times Beach_i + \varepsilon_{ij}$$

Beach i

$$\begin{pmatrix} R_{i1} \\ R_{i2} \\ R_{i3} \\ R_{i4} \\ R_{i5} \end{pmatrix} = \begin{pmatrix} 1 & NAP_{i1} \\ 1 & NAP_{i2} \\ 1 & NAP_{i3} \\ 1 & NAP_{i4} \\ 1 & NAP_{i5} \end{pmatrix} \times \begin{pmatrix} \alpha \\ \beta \end{pmatrix} + \begin{pmatrix} 1 \\ 1 \\ 1 \\ 1 \\ 1 \end{pmatrix} \times b_i + \begin{pmatrix} \varepsilon_{i1} \\ \varepsilon_{i2} \\ \varepsilon_{i3} \\ \varepsilon_{i4} \\ \varepsilon_{i5} \end{pmatrix}$$

Assumptions:

The random effects b_i are normally distributed: N(0, d^2).

The errors ε_i are normally distributed.

library(nlme)

RIKZ\$fBeach <- factor(RIKZ\$Beach)

Mlme1 <- lme(Richness ~ NAP, random = ~1 | fBeach, data = RIKZ) summary(Mlme1)

The random intercept model: results

Linear mixed-effects model fit by REML

Data: RIKZ

AIC BIC logLik

247.4802 254.525 -119.7401

Random effects:

Formula: ~1 | fBeach

(Intercept) Residual

StdDev: 2.944065 3.05977

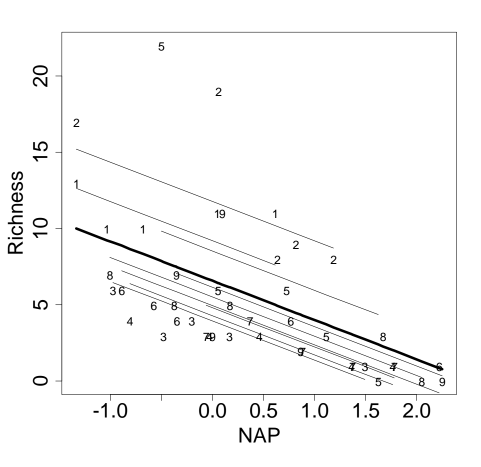
Fixed effects: Richness ~ NAP

Value Std.Error DF t-value p-value (Intercept) 6.581893 1.0957618 35 6.006682 0 NAP -2.568400 0.4947246 35 -5.191574 0

Number of Observations: 45

Number of Groups: 9

The random intercept model: results



The thick line represents the fitted line obtained by the fixed component $6.58 - 2.56 \, NAP_i$, also called the population model.

The other lines are obtained by adding the contribution of \mathbf{b}_i for each beach i to the population fitted curve.

Hence, the random intercept model implies one average curve (the thick line) that is allowed to be shifted up, or down, for each beach by something that is normally distributed with a certain variance d^2 (2.94^2).

The random intercept and slope model

$$\begin{array}{c} \text{Beach i} \\ \begin{pmatrix} R_{i1} \\ R_{i2} \\ R_{i3} \\ R_{i4} \\ R_{i5} \end{pmatrix} = \begin{pmatrix} 1 \text{ MAP}_{i1} \\ 1 \text{ MAP}_{i2} \\ 1 \text{ MAP}_{i3} \\ 1 \text{ MAP}_{i4} \\ 1 \text{ MAP}_{i5} \end{pmatrix} \times \begin{pmatrix} \alpha \\ \beta \end{pmatrix} + \begin{pmatrix} 1 \text{ MAP}_{i1} \\ 1 \text{ MAP}_{i2} \\ 1 \text{ MAP}_{i3} \\ 1 \text{ MAP}_{i4} \\ 1 \text{ MAP}_{i5} \end{pmatrix} \times \begin{pmatrix} b_{i1} \\ b_{i2} \end{pmatrix} + \begin{pmatrix} \varepsilon_{i1} \\ \varepsilon_{i2} \\ \varepsilon_{i3} \\ \varepsilon_{i4} \\ \varepsilon_{i5} \end{pmatrix}$$

Assumptions:
$$\begin{pmatrix} b_{i1} \\ b_{i2} \end{pmatrix} \sim N(0, D)$$
 where $D = \begin{pmatrix} d_{11}^2 & d_{21} \\ d_{12} & d_{22}^2 \end{pmatrix}$

The errors ε_i are normally distributed.

library(nlme)

RIKZ\$fBeach <- factor(RIKZ\$Beach)

Mlme2 <- lme(Richness ~ NAP, random = ~1 + NAP | fBeach, data = RIKZ) summary(Mlme2)

The random intercept and slope model: results

Linear mixed-effects model fit by REML

Data: RIKZ

AIC BIC logLik

244.3839 254.9511 -116.1919

Random effects:

Formula: ~1 + NAP | fBeach

Structure: General positive-definite, Log-Cholesky parametrization

StdDev Corr

(Intercept) 3.549100 (Intr)

NAP 1.715015 -0.99

Residual 2.702785

Fixed effects: Richness ~ NAP

Value Std.Error DF t-value p-value

(Intercept) 6.588729 1.2647708 35 5.209425 0e+00

NAP -2.830029 0.7229514 35 -3.914549 4e-04

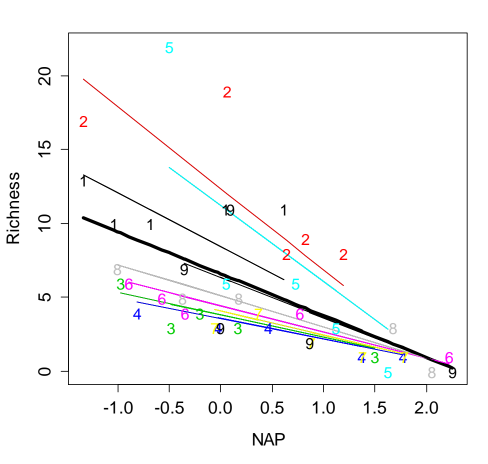
Correlation:

(Intr) NAP -0.819

Number of Observations: 45

Number of Groups: 9

The random intercept and slope model



The variance d_{11}^2 plays the same role as d^2 in the random intercept model; it determines the amount of variation around the population intercept α . The numerical output shows that its estimated value is $3.54^2 = 12.5$.

The model also allows for random variation around the population slope in a similar way as it does for the intercept. The variance d_{22}^2 determines the variation in slopes at the nine beaches. The estimated value of $1.71^2 = 2.92$ shows that there is considerably more variation in intercepts than in slopes at the nine beaches.

Finally, there is a correlation between the random intercepts and slopes. Its value of -0.99 is rather high, but indicates that beaches with a high positive intercept also have a high negative slope.

R code for figures

```
# The Random Intercept and/or slope Model
RIKZ = read.table('D:/softwares/R/library/AED/data/RIKZ.txt',header=T)
library(nlme)
RIKZ$fBeach <- factor(RIKZ$Beach)
Mlme1 <- Ime(Richness ~ NAP, random = ~1 | fBeach, data = RIKZ)
Mlme1 <- Ime(Richness ~ NAP, random = ~1 + NAP | fBeach, data = RIKZ)
summary(MIme1)
# plot regression lines
F0 <- fitted(Mlme1, level = 0) # fitted values obtained by the population model
F1 <- fitted(Mlme1, level = 1) # fitted values obtained by within-beach model
I <- order(RIKZ$NAP); NAPs <- sort(RIKZ$NAP)
plot(NAPs, F0[I], lwd = 4, type = "I",
vlim = c(0, 22), vlab = "Richness", xlab = "NAP")
for (i in 1:9){
  x1 <- RIKZ$NAP[RIKZ$Beach == i]
  y1 <- F1[RIKZ$Beach == i]
  K <- order(x1)
  lines(sort(x1), y1[K])
```

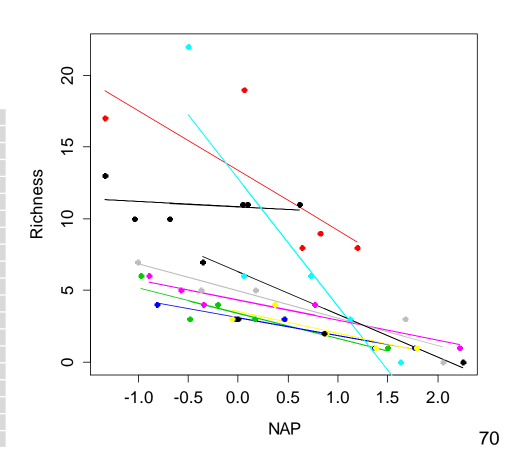
text(RIKZ\$NAP, RIKZ\$Richness, RIKZ\$Beach, cex = 0.9)

model = Im(Richness ~ NAP * fBeach, data = RIKZ)

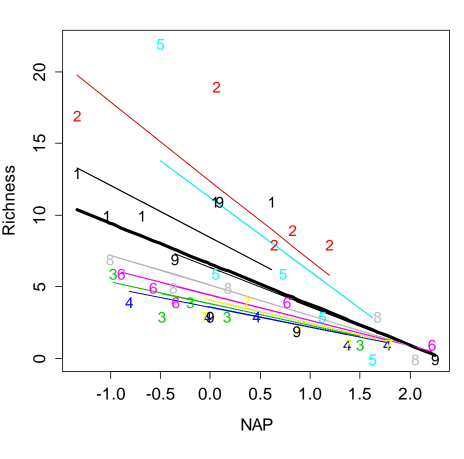
```
model = Im(Richness ~ NAP * fBeach, data = RIKZ) anova(model)
```

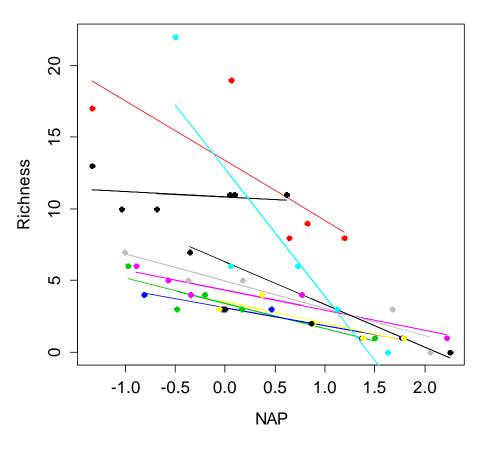
```
pred = predict(model, RIKZ[, c('NAP', 'fBeach')])
plot(Dat$NAP, Dat$Richness, xlab='NAP', ylab='Richness', col='white')
Dat = cbind(RIKZ, pred)
for (i in 1:9) {
    Data = Dat[Dat$Beach==i, ]
    lines(Data$NAP, Data$pred, col=i)
}
```

Coefficients:					
	Estimate	Std.Error	t value	р	
(Intercept)	10.8219	1.3341	8.112	1.03E-08	***
NAP	-0.3718	1.5493	-0.24	0.81214	
fBeach2	2.5238	1.7685	1.427	0.165013	
fBeach3	-7.4212	1.7346	-4.278	0.000211	***
fBeach4	-7.7342	1.8541	-4.171	0.000281	***
fBeach5	1.9609	1.9485	1.006	0.323166	
fBeach6	-6.4973	1.7494	-3.714	0.000938	***
fBeach7	-7.3013	2.2611	-3.229	0.003253	**
fBeach8	-5.8704	1.7981	-3.265	0.002974	**
fBeach9	-4.5268	1.8631	-2.43	0.022036	*
NAP:fBeach2	-3.8034	1.9941	-1.907	0.067169	
NAP:fBeach3	-1.3835	2.0405	-0.678	0.503511	
NAP:fBeach4	-0.8767	1.9528	-0.449	0.657044	
NAP:fBeach5	-8.5283	2.134	-3.996	0.000447	***
NAP:fBeach6	-1.0167	1.829	-0.556	0.582872	
NAP:fBeach7	-1.1458	2.2756	-0.504	0.618683	
NAP:fBeach8	-1.5212	1.8134	-0.839	0.408891	
NAP:fBeach9	-2.5957	1.9537	-1.329	0.1951	



Mixed effect model vs. regular ANCOVA





lme(Richness ~ NAP, random = ~1 + NAP | fBeach,
data = RIKZ)

Im(Richness ~ NAP * fBeach, data = RIKZ)

Rethinking the homogeneity of regression slopes assumption

Always run an ANCOVA model including both the IV, CV and the CVxIV interaction term.

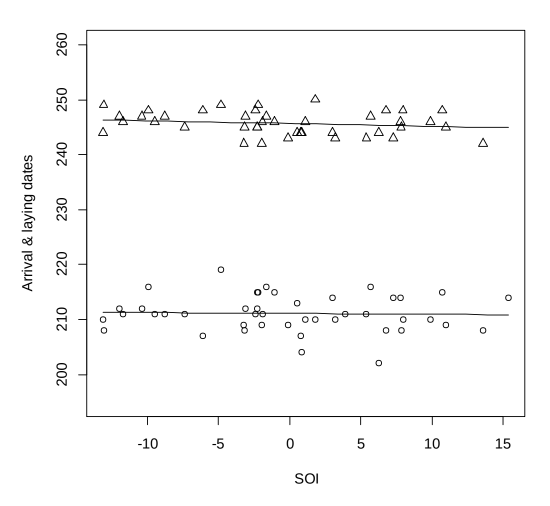
If the CVxIV interaction is significant, ANCOVA should not be performed.

One option is to assess group differences on the DV at particular levels of the CV.

Another option is to use <u>mediation analyses</u> to determine if the CV accounts for the IV's effect on the DV.

ANCOVA with autocorrelation

Year	ArrivalAP	LayingAP	ArrivalCP	LayingCP	ArrivalEP	LayingEP	MSA	SOI
1951	214	NA	NA	NA	NA	NA	0.086	15.38
1952	NA	NA	NA	NA	-10	46	0.127	-0.69
1953	212	245	209	260	NA	NA	0.101	-2.28
1954	NA	NA	NA	NA	NA	NA	0.109	-6.8
1955	NA	NA	NA	NA	NA	NA	0.087	4.08
1956	NA	NA	NA	NA	-9	42	0.086	10.58
1957	215	248	214	259	NA	NA	0.072	10.73
1958	NA	NA	NA	NA	-13	44	0.082	-3.89
1959	209	242	NA	NA	-3	38	0.078	-3.2
1960	NA	NA	NA	NA	NA	NA	0.078	-0.04
1961	NA	NA	NA	NA	NA	NA	0.065	3.83
1962	207	244	210	NA	-12	NA	0.086	0.8
1963	211	243	193	261	-7	41	0.137	5.4
1964	209	242	213	NA	-15	43	0.098	-1.95
1965	202	244	207	262	NA	NA	0.111	6.28
1966	NA	NA	NA	NA	-7	41	0.098	-8.43
1967	NA	NA	NA	NA	-9	38	0.056	-4.24
1968	210	243	212	260	-17	44	0.065	3.2
1969	214	244	209	NA	NA	NA	0.082	3.02
1970	NA	NA	NA	NA	-1	46	0.062	-5.38
1971	211	NA	207	NA	-6	45	0.051	3.93
1971	209	245	201	261	0	43	0.064	10.95
1972	211	245	215	268		50		-7.35
1973	211	243	NA	NA	0 -7	50 44	0.071	
						44	0.063	7.28
1975	210	246	195	259	-7 45		0.106	9.9
1976	208	242 246	198 210	262	-15	44	0.087	13.6
1977	210	246		266	-1	43	0.078	1.11
1978	216	248	213 213	265 265	-5	43	0.064	-9.9
1979	216				-9	43	0.08	-1.65
1980	211	246	206	NA NA	-11	51 45	0.037	-1.91
1981	212	247	213	NA	-5 20		0.054	-3.08
1982	210	250	212		-20	46	0.06	1.8
1983	208	249	216	264	NA 10	NA	0.09	-13.05
1984	NA	NA 040	NA	NA	-10	39	0.088	-8.33
1985	209 204	243 244	208 199	260 263	-4 -10	42 45	0.082	-0.11
1986							0.061	0.86
1987	211	248	210	261	-4	44	0.096	-2.38
1988	210	244	213	263	-4	46	0.048	-13.08
1989	208	245	220	NA	-1	47	0.051	7.82
1990	208	248	219	265	-2	47	0.034	6.77
1991	215	249	215	NA	-3	47	0.054	-2.19
1992	211	247	211	264	-6	44	0.041	-8.78
1993	212	247	209	269	-7	41	0.056	-10.38
1994	211	246	215	262	-7 47	41	0.076	-9.47
1995	212	247	215	263	-17	47	0.065	-11.93
1996	215	245	215	263	-13	46	0.054	-2.27
1997	216	247	214	NA	-6	42	NA	5.69
1998	211	246	217	262	-1	NA 47	NA	-11.67
1999	215	246	220	263	-4	47	NA	-1.08
2000	210	248	208	261	-3	43	NA	7.95
2001	214	246	NA 245	263	-14	47	NA	7.8
2002	213	244	215	NA	-16	44	NA	0.53
2003	207	248	NA 246	264	-9	43	NA	-6.1
2004	208	245	216	265	-11	45	NA	-3.14
2005	219	249	210	266	NA	NA	NA	-4.82



Zuur et al. 2009. *Mixed Effects Models and Extensions in Ecology with R.* Page 343-361 73

ANCOVA with autocorrelated data

```
head(ABirds) # data
```

```
AP <- c(ABirds$ArrivalAP, ABirds$LayingAP)

SOI2 <- c(ABirds$SOI, ABirds$SOI)

Y2 <- c(ABirds$Year, ABirds$Year)

ID <- factor(rep(c("Arrival", "Laying"), each = 55))

library(nlme)

vf2 <- varIdent(form =~ 1 | ID)

M1 <- gls(AP ~ SOI2 + ID + SOI2:ID, weights = vf2, na.action = na.omit)

M2 <- gls(AP ~ SOI2 + ID + SOI2:ID, weights = vf2, na.action = na.omit,

correlation = corAR1(form =~Y2 | ID))

anova(M1, M2)
```

	Model	df	AIC	BIC	logLik	Test	L.Ratio p	-value
М1	1	6	427.8205	442.2608	-207.9102			
M2	2	7	426.0757	442.9228	-206.0379	1 vs 2	3.744727	0.053

Model selection

```
# to compare two models with the same random structure, but with different fixed effect,

# we need to use the maximum likelihood estimation method instead of REML

# (P.356 in Zuur et al. 2009)

M3 <- gls(AP ~ SOI2 + ID + SOI2:ID, weights = vf2, na.action = na.omit, method = "ML",

correlation = corAR1(form =~Y2 | ID))

M4 <- gls(AP ~ SOI2 + ID, weights = vf2, na.action = na.omit, method = "ML",

correlation = corAR1(form =~Y2 | ID))

M5 <- gls(AP ~ ID, weights = vf2, na.action = na.omit, method = "ML",
```

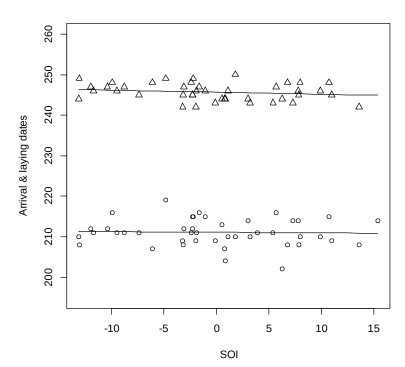
anova(M3, M4, M5)

	Model	df	AIC	BIC	logLik	Test	L.Ratio p-value
M3	1	7	419.0303	436.2107	-202.5152		
M4	2	6	417.1962	431.9222	-202.5981	1 vs 2	0.1658611 0.6838
M5	3	5	416.2205	428.4922	-203.1102	2 vs 3	1.0243171 0.3115

correlation = corAR1(form = ~Y2 | ID))

Plot

```
plot(ABirds$SOI, ABirds$ArrivalAP, ylim = c(195, 260), type = "n",
    ylab = "Arrival & laying dates", xlab='SOI')
points(ABirds$SOI, ABirds$ArrivalAP, pch = 1)
points(ABirds$SOI, ABirds$LayingAP, pch = 2)
MyX <- data.frame(SOI2 = seq(from = min(ABirds$SOI),
        to = max(ABirds\$SOI),
        length = 20), ID = "Arrival")
Pred1 <- predict(M3, newdata = MyX)
lines(MyX$SOI2, Pred1)
MyX <- data.frame(SOI2 = seq(from = min(ABirds$SOI),
        to = max(ABirds\$SOI),
        length = 20), ID = "Laying")
Pred2 <- predict(M3, newdata = MyX)
lines(MyX$SOI2, Pred2)
```



R Model formul	las in R	2020 Lecture 10. Analysis of covariance	Biostatistics Xinhai Li
Regression	y~x	x is a continuous explanat	tory variable
One-way ANOVA	v~sex	sex is a factor (categorica	l variable)

Include main effect and all interaction

As above, don't fit three-way interaction

Common slope but different intercepts

Different slopes and different intercepts

Factor c nested in factor b, within factor a

Factorial experiment with three different

Two continuous explanatory variables

Quadratic. I() indicates as is, so $I(x^2)$ is x

Fit variable and their interactions up to two-

Fit smoothed x an z in a generalized

Transformation specified in model

Includes interaction: x+y+x:y

Quadratic polynomial

additive model

error terms

squared.

way

Regression	y~x	x is a continuous explanatory
One-way ANOVA	y~sex	sex is a factor (categorical var
Two-way ANOVA	y~sex+genotype	sex and genotype are factors

y~n*p*k

y~x+sex

y~x*sex

y~a/b/c

 $V\sim X+Z$

y~x*z

 $y\sim poly(x,2)$

 $y\sim(w+x+z)^2$

 $y \sim s(x) + s(z)$

 $\log(y) \sim I(1/x) + \operatorname{sqrt}(z)$

 $y\sim n*p*k-n:p:k$

y~a/b/c+Error(a/b/c)

 $y \sim x + I(x^2) + z + I(z^2) + x : z$

Factorial ANOVA

Nested ANOVA

Split-plot ANOVA

Multiple regression

Multiple regression

Multiple regression

Multiple regression

Multiple regression

Nonparametric model

explanatory variables

Transformed response &

Three-way ANOVA

Analysis of covariance

Analysis of covariance

Model Operators

Symbol	Explanation
+	indicates inclusion of an explanatory variable, not addition
-	indicates deletion of an explanatory variable, not subtraction
*	indicates inclusion of explanatory variables and all their interactions, not multiplication
<i> </i>	indicates nesting of explanatory variable, not division
	indicates conditioning
•	indicates an interaction, such as x:z
a*b*c	= a + b + c + a:b + a:c + b:c + a:b:c
a/b/c	= a + b%in%a + c%in%b%in%a
(a+b+c)^2	= a + b + c + a:b + a:c + b:c = main effects & up to 2-way interactions
a*b*c-a:b:c	= a + b + c + a:b + a:c + b:c

Statistical Methods

Method	Description
lm	Fits a linear model with normal errors and constant variance: regression, analysis of variance, analysis of covariance
aov	Also fits a linear model with normal errors and constant variance, but oriented towards analysis of variance
glm	Fits generalized linear models by specifying one of a family of error structures (e.g., Poisson for count data) and a particular link function.
gam	Fits generalized additive models by specifying one of a family of error structures (e.g., Poisson for count data) in which continuous explanatory variable can optionally be fitted as arbitrary smoothed functions using non-parametric smoothers rather than a specific parametric functions.
Ime & Imer	Fits linear mixed-effects models including fixed and random effects and allow for the specification of correlation structure amongst the explanatory variables and autocorrelation of the response variable. Imer allows for non-normal errors and non-constant variance with the same error families as glm
nls	Fits non-linear regression models using least squares.
loess	Fits a local regression model with one or more continuous explanatory variables using non-parametric techniques to produce a smoothed model surface.
tree	fits a regression tree model using binary recursive partitioning.

Functions

Functions	Description
summary	Displays the results of a model fit, depending on the method used. Use summary.Im or summary.aov to obtain particular displays.
anova	Displays an analysis of variance table and compares models.
plot	Produces diagnostic plots for model checking.
update	Modifies the last model fit.
coef	Displays the estimated coefficients from a model.
fitted	Displays the fitted values from a model.
resid	Displays the residuals from a model.
predict	Displays the predicted values from a model.
	ancova = $lm(Y\sim X1*X2)$
	coef(ancova)
	fitted(ancova)
	predict(ancova)
	predict(aricova)

resid(ancova)

Assignment

Task:

Develop a ANCOVA experimental design. Generate your own data and FORMALIZE your hypotheses.

Define dependant variable, treatment variable and covariate variable

Check homogeneity of regression, clearly state the model performance

List the model fit statistics