# LA MOLINA NATIONAL AGRARIAN UNIVERSITY

## FACULTY OF ECONOMICS AND PLANNING DEPARTMENT OF STATISTICS AND INFORMATICS



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**PRACTICAL MANUAL** 

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### **PREFACE**

R is a functional programming system exclusive to manage data in statistics and related sciences, such as mathematics, in environments like Windows, Linux and MAC. 'Agricolae' is a package of functions for R applied to agricultural research.

The package 'agricolae' offers a broad functionality in the design of experiments, especially for experiments in agriculture and improvements of plants, which can also be used for other purposes. It contains the following designs: lattice, alpha, cyclic, balanced incomplete block designs, complete randomized blocks, Latin, Graeco-Latin, augmented block designs, divided parcels, divided blocks. It also has several procedures of experimental data analysis, such as the comparisons of treatments of Waller-Duncan, Bonferroni, Duncan, Student-Newman-Keuls, Scheffe, or the classic LSD and Tukey; and non-parametric comparisons, such as Kruskal-Wallis, Friedman, Durbin and Waerden, stability analysis, and other procedures applied in genetics, as well as procedures in biodiversity and descriptive statistics.

For more details on the use of 'agricolae', see the reference manual and the aid system in HTML, which can be found in the menu of R.

#### 1 INSTALLATION OF AGRICOLAE AND USE IN R

### 1.1 INSTALLATION

The main program of R should be already installed in the platform of your computer (Windows, Linux or MAC). If it is not installed yet, you can download it from the R project (<a href="www.r-project.org">www.r-project.org</a>) of a repository CRAN (R Development Core Team, 2013). As it is a free program, no identification is required. The packages can be incorporated through an installation process, directly from the platform of R.

'Agricolae' is a package for R, and as such its installation is just like any other package of R.

For Windows, the R program (version 3.0.0 or higher) is required.

If the R program is already installed in Windows or in another platform, the installation of 'agricolae' can be done directly from the console of R through Internet, that is

install.packages("agricolae)

A repository should be selected and the system is installed automatically.

If there is no Internet connection, it is necessary to copy the file agricolae\_x.x-x. zip for Windows from the page of the R project.

The file agricolae\_x.x-x. zip (De Mendiburu, 2013) can be downloaded from the R repository in the following addresses: <a href="www.r-project.org">www.r-project.org</a> or <a href="http://cran.at.r-project.org/web/packages/agricolae/index.html">http://cran.at.r-project.org/web/packages/agricolae/index.html</a>

The file can be directly incorporated into R installing from the console with the following instruction set if the file is located in the address E: install.packages("E:/agricolae 1.1-4.zip)

It can also be installed from the R menu:

Packages, Install package(s) from local zip files.... Selecting the file zip does not require any unpacking.

For a complete functionality, 'agricolae' requires other packages.

MASS: for the generalized inverse used in the function PBIB.test()

Ime4: for the methods REML and LM in PBIB.test

klaR: for the function triplot() used in the function AMMI()

akima: for the use of the function interpp() used in grid3p() for interpolation

Cluster: for the use of the function consensus()

### 1.2 USE IN R

Since 'agricolae' is a package of functions, these are operational when they are called directly from the console of R and are integrated to all the base functions of R.

The following orders are frequent:

Load the package to the memory: library(agricolae)

Download: detach(package:agricolae)

Once the package is loaded, you can:

List the database: data(package="agricolae") Load the sweet potato data: data(sweetpotato)

See its structure: str(sweetpotato)
Publish its content: fix(sweetpotato)

In order to continue with the command line, do not forget to close the open windows with any R order.

For help: help(sweetpotato); ? sweetpotato To search any functions: apropos("design")

[1] "design.ab"	"design.alpha"	"design.bib"	"design.crd"
[5] "design.cyclic"	"design.dau"	"design.graeco"	"design.lattice"
[9] "design.lsd"	"design.rcbd"	"design.split"	"design.strip"

### 2 DESCRIPTIVE STATISTICS

The package 'agricolae' provides some complementary functions to the R program, specifically for the management of the histogram.

### 2.1 HISTOGRAM

The histogram is constructed with the function graph.freq() and is associated to other functions: polygon.freq, table.freq, stat.freq, intervals.freq, sturges.freq, join.freq, ojiva.freq, and normal.freq.

### Example 1.1 Data generated in R. (students' weight). Figure 2.1

```
c(68, 53, 69.5, 55, 71, 63, 76.5, 65.5, 69, 75, 76, 57, 70.5, 71.5, 56, 81.5, 69, 59, 67.5, 61, 68, 59.5, 56.5, 73, 61, 72.5, 71.5, 59.5, 74.5, 63) -> weight
```

### Load the package 'agricolae':

```
library(agricolae)
par(mfrow=c(2,2))
h1<- graph.freq(weight, col="yellow", frequency =1, axes= FALSE,
main=" Absolute frequency \nh1")
axis(1,h1$breaks); axis(2, las=1)
h2<- graph.freq(weight, frequency =2 , axes= FALSE, main=" polygon
of frequency \nh2")
polygon.freq(h2, col="blue", lwd=2, frequency =2)
TIC<- h2$breaks[2]- h2$breaks[1]
axis(1,c(h2\$mids[1]-TIC, h2\$mids, h2\$mids[6]+TIC),cex=0.6)
axis(2, cex=0.6, las=1)
h3<- graph.freg(weight, col="brown", frequency =3,
                                                           axes=
main="densidad\nh3")
axis(1,h3\$breaks); axis(2,las=1)
h4<- graph.freq(weight, col="blue",
                                          frequency
                                                     =3,
                                                           axes=
                                                                   FALSE,
main="densidad normal\nh4", density=4)
normal.freq(h4, col="red", lty=4,lwd=2, frequency=3)
axis(1,h3$breaks); axis(2,las=1)
```

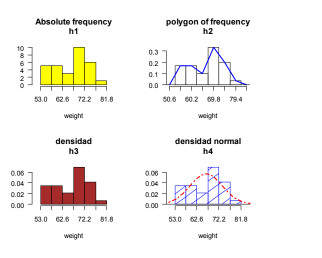


Figure 2.1 Histograms, polygon and density

### 2.2 HANDLING SCALES

It refers to the scale changes in the axes. Figure 2.2

```
par(mfrow=c(2,2))
h5<- graph.freq(weight, axes=FALSE, frequency =1, main=" Absolute
frequency\nh5")
axis(1,h5\$breaks,las=2); axis(2,las=1)
h6<- graph.freg(weight, axes=FALSE, nclass=5, main=" frecuency with
5 classes \nh6")
axis(1,h6\$breaks,las=2); axis(2,las=1)
normal.freq(h6,col="red")
h7<- graph.freq(weight, density=6, col="blue", frequency =3,
axes=FALSE, main=" density \nh7")
lines(density(weight),col="brown",lwd=2)
axis(1,h6\$breaks,las=2); axis(2,las=1)
h8<- graph.freq(weight, border=0, frequency =3, axes=FALSE,
                                                              main="
polygon and density \nh8")
polygon.freq(h8,col="blue", frequency =3)
lines(density(weight),col="brown",lwd=2)
axis(1,h6\$breaks,las=2); axis(2,las=1)
```

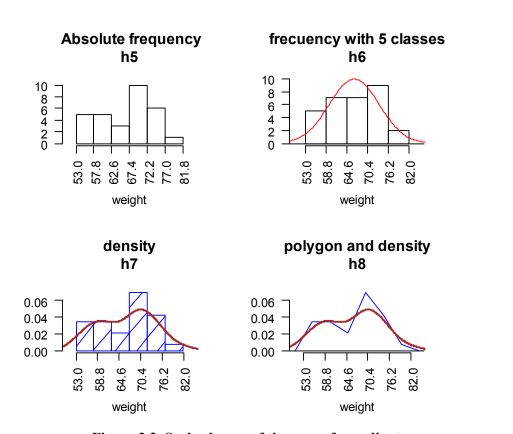


Figure 2.2. Scale change of the axes of coordinates

### 2.3 FREQUENCY TABLES AND STATISTICS

```
Rounded off to two decimals:
                                            stat.freq(h6)
                                            $variance
                                            [1] 50.42133
round(table.freq(h6), 2)
                                            $mean
                                            [1] 66.72667
  Lower Upper Main freq relative CF RCF
                                            $median
   53.0 58.8 55.9
                   5
                           0.17 5 0.17
   58.8 64.6 61.7
                                            [1] 67.08571
                      7
                            0.23 12 0.40
                                            $mode
   64.6 70.4 67.5
                      7
                            0.23 19 0.63
                                                    - ]
                                                             mode
                                            [ -
   70.4
        76.2 73.3
                      9
                            0.30 28 0.93
                                            70.4 76.2
                                                          71.68889
   76.2 82.0 79.1
                      2
                            0.07 30 1.00
```

### 2.4 REPRODUCING HISTOGRAMS AND USE OF hist()

The class of graph.freq() is graph.freq. Figure 2.3

Reproducing the histogram h6 (5 classes)

```
h9<-ojiva.freg(h5,axes=FALSE,type="b",
                                        main="ojiva
                                                           h5\nh9",
                                                      to
col="red")
axis(1, round(h9[,1],1), las=2); axis(2, las=1)
h10<-plot(h6, axes=FALSE, main=" frequency with 5 classes \nh10")
axis(1,h6\$breaks,las=2); axis(2,seq(0,10))
normal.freq(h6,col="red")
summary(h6)
 Lower Upper Main freq relative CF RCF
  53.0 58.8 55.9 5 0.17 5 0.17
                     7
                           0.23 12 0.40
   58.8 64.6 61.7
                    7
   64.6 70.4 67.5
                           0.23 19 0.63
  70.4 76.2 73.3 9
                           0.30 28 0.93
                     2
  76.2 82.0 79.1
                           0.07 30 1.00
```

The class types of the functions hist() and graph.freq() are 'histogram' and 'graph.freq', respectively. However, it is possible to establish compatibility between both functions.

```
hh <- hist(weight,nclass=5, plot=FALSE) # Reports 7 classes
# hist(weight,nclass=4) # Reports 4 classes</pre>
```

In order to show the relative frequencies, you can use graph.freq() with the object hh created by hist(), without modifying the classes.

```
h11<-graph.freq(hh, frequency=2,main="relative\nh11" , col=colors()[467],axes=F) axis(1,h11$breaks, las=2) axis(2,las=1)
```

See the summaries: > summary(hh), summary(h11)

The functions of 'agricolae' for the management of histograms function correctly on the objects created by the function hist() of R.

### 2.5 HISTOGRAM BASED ON GROUPED DATA

If there are grouped data, you can graphic and obtain the histogram summaries with the function graph.freq(), as, for example, in the following table:

```
10-20 20-30 30-40 40-50
              15
                     18
In R we have:
classes \leftarrow c(0, 10, 20, 30, 40, 50)
frec <- c(3, 8, 15, 18, 6)
h12 <- graph.freq(classes, counts=frec, xlab="Classes",
main="Classes")
summary(h12)
  Lower Upper Main freq relative CF RCF
           10
                 5
                       3
                              0.06 3 0.06
                              0.16 11 0.22
     10
           20
                       8
                 15
     20
                 25
                              0.30 26 0.52
           30
                      15
     30
           40
                 35
                      18
                              0.36 44 0.88
                 45
     40
           50
                       6
                              0.12 50 1.00
```

All the functions of 'agricolae' can be applied, including plot().

# plot(h12, xlab="Classes", main="Classes")

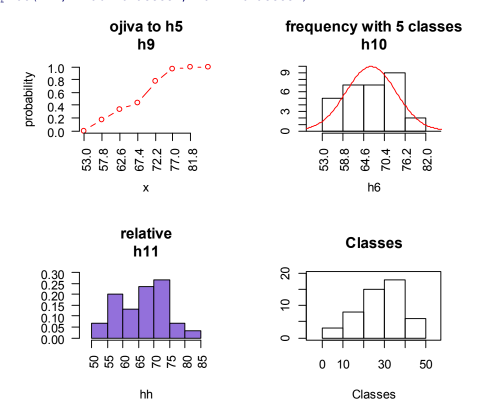


Figure 2.3 New scales for the histograms

### 2.6 JOINING CLASSES

Knowing the students' weight, the original intervals can be changed, joining, for example:

```
intervals<- join.freq(h5$breaks,1:2)</pre>
intervals.freq(h5$breaks)
    lower upper
                          intervals.freq(intervals)
[1,] 53.2 58.0
                               lower upper
[2,] 58.0 62.8
                          [1,] 53.2 62.8
[3,] 62.8 67.6
                          [2,] 62.8 67.6
                          [3,] 67.6 72.4
[4,] 67.6 72.4
[5,] 72.4 77.2
                          [4,] 72.4 77.2
                          [5,] 77.2 82.0
[6, ] 77.2 82.0
                          h13<-graph.freq(weigth,breaks= intervals)</pre>
```

### 3 EXPERIMENT DESIGNS

The package 'agricolae' presents special functions for the creation of the field book for experimental designs. Due to the random generation, this package is quite used in agricultural research.

For this generation, certain parameters are required, as for example the name of each treatment, the number of repetitions, and others, according to the design (Cochran, 1992; Kuehl, 2000; Montgomery, 2002; LeClerg, 1962). There are other parameters of random generation, as the seed to reproduce the same random generation or the generation method (See the reference manual of agriculture http://cran.at.r-project.org/web/packages/agricolae/agricolae.pdf)

### 3.1 COMPLETELY RANDOMIZED DESIGNS

They only require the names of the treatments and the number of their repetitions.

```
trt <- c("A", "B", "C")
repetition \leftarrow c(4, 3, 4)
plan1 <- design.crd(trt,r=repetition)</pre>
  plots trt r
1
      1
         A 1
2
      2
          C 1
3
      3 A 2
4
      4 A 3
5
      5 B 1
      6 C 2
6
7
      7
         C 3
      8 B 2
8
9
      9 A 4
     10 B 3
10
         C 4
11
     11
```

### Excel:

```
write.csv(plan1,"plan1.csv",row.names=FALSE)
```

### 3.2 RANDOMIZED COMPLETE BLOCK DESIGN

They require the names of the treatments and the number of blocks.

```
trt <- c("A", "B", "C")
repetition <- 4
plan2 <- design.rcbd(trt,r=repetition, seed=5, number=101,
first=FALSE)
t(matrix(plan2[ ,3],c(3,4)))
       [,1] [,2] [,3]
[1,] "A" "B" "C"
[2,] "C" "A" "B"
[3,] "C" "A" "B"
[4,] "A" "C" "B"</pre>
```

The plan can be sent to excel as a field book.

### 3.3 LATIN SQUARE DESIGNS

They require the names of the treatments.

```
trt <- c("A", "B", "C", "D")
plan3 <- design.lsd(trt, seed=55, number=101, first=FALSE)
t(matrix(plan3[,4],c(4,4)))
       [,1] [,2] [,3] [,4]
[1,] "A" "B" "D" "C"
[2,] "B" "C" "A" "D"
[3,] "D" "A" "C" "B"
[4,] "C" "D" "B" "A"</pre>
```

### 3.4 GRAECO-LATIN DESIGNS

They require the names of the treatments of each factor of study.

```
T1 <- c("A", "B", "C", "D")

T2 <- 1:4

plan4 <- design.graeco(T1,T2, seed=55, number=101)

t(matrix(paste(plan4[,4],plan4[,5]),c(4,4)))

[,1] [,2] [,3] [,4]

[1,] "A 3" "D 4" "C 1" "B 2"

[2,] "D 1" "A 2" "B 3" "C 4"

[3,] "C 2" "B 1" "A 4" "D 3"

[4,] "B 4" "C 3" "D 2" "A 1"
```

### 3.5 BALANCED INCOMPLETE BLOCK DESIGNS

They require the names of the treatments and the size of the block.

```
trt <- c("A", "B", "C", "D", "Control")
k <- 4
plan5 <- design.bib(trt,k, seed=55, number=101)</pre>
```

According to the produced information, they are five blocks of size 4, being the matrix:

```
t(matrix(plan5[,3],c(4,5)))
    [,1] [,2] [,3]
                               [,4]
[1,] "B"
             "Control" "C"
                               "A"
[2,] "D"
             "A" "C"
                               "B"
[3,] "B"
             "C"
                    "Control" "D"
[4,] "C"
             "D"
                    "A"
                              "Control"
                      "D"
                               "A"
[5,] "Control" "B"
```

It can be observed that the treatments have four repetitions. The parameter lambda has three repetitions, which means that a couple of treatments are together on three occasions. For example, B and E are found in the blocks I, III and V.

### 3.6 CYCLIC DESIGNS

They require the names of the treatments, the size of the block and the number of repetitions. This design is used for 6 to 30 treatments. The repetitions are a multiple of the size of the block; if they are six treatments and the size is 3, then the repetitions can be 6, 9, 12, etc.

```
trt <- c("A", "B", "C", "D", "E", "F")
plan5 <- design.cyclic(trt, k=3, r=6, seed=55, number=101)</pre>
cyclic design
Generator block basic:
1 2 4
1 3 2
Parameters
treatments: 6
Block size: 3
Replication: 6
> plan5$design[[1]]
    [,1] [,2] [,3]
[1,] "F" "A"
              "C"
[2,] "A"
         "D" "B"
[3,] "B"
         "C" "E"
[4,] "D" "F" "C"
[5,] "A" "D" "E"
[6,] "B" "E" "F"
```

```
> plan5$design[[2]]
     [,1] [,2] [,3]
[1,] "D"
          "E"
[2,] "E"
          "F"
                "D"
[3,] "B"
          "C"
                "D"
[4,] "A"
          "F"
                "E"
[5,] "C"
          "B"
                "A"
                "A"
[6,] "B"
```

12 blocks of 4 treatments each have been generated.

### 3.7 LATTICE DESIGNS

They require a number of treatments of a perfect square; for example 9, 16, 25, 36, 49, etc.

They can generate a simple lattice (2 rep.) or a triple lattice (3 rep.)

generating a triple lattice design for 9 treatments 3x3

```
plan6 <- design.lattice(k=3, seed=55, number=101)</pre>
print(plan6)
$square1
      [,1] [,2] [,3]
[1,]
        1 4
[2,]
        2
              9
                   3
[3,]
        5
              7
$square2
       [,1] [,2] [,3]
            1
[1,]
        2
[2,]
              4
        3
[3,]
             8
$square3
       [,1] [,2] [,3]
            4
        2
[1,]
                   6
        3
             1
                   7
[2,]
        9
             8
                   5
[3,]
$plan
  plots sqr block trt
    101
             1
         1
                     1
2
    102
          1
                 1
                     4
    127
         3
                     5
27
```

### 3.8 ALPHA DESIGNS

These designs are generated by the alpha arrangements (Patterson & Williams, 1976). They are similar to the lattice designs, but the tables are rectangular, with s blocks x k treatments. The number of treatments should be equal to s\*k and all the experimental units, r\*s\*k.

```
Genotype<-paste("geno", 1:15, sep = "")</pre>
plan7 <- design.alpha(Genotype, k=3, r=2, seed=55)</pre>
alpha design (0,1) - Serie I
Parameters Alpha design
treatmeans: 15
Block size : 3
Blocks : 5
Replication: 2
Efficiency factor
(E) 0.6363636
<<< Book >>>
plan7$design$rep1
[,1] [,2] [,3]
[1,] "geno8" "geno4" "geno10"
[2,] "geno1" "geno12" "geno14"
[3,] "geno6" "geno2" "geno15"
[4,] "geno7" "geno3" "geno11"
[5,] "geno13" "geno9" "geno5"
plan7$design$rep2
     [,1] [,2]
                        [,3]
[1,] "geno13" "geno7" "geno1"
[2,] "geno8" "geno5" "geno14"
[3,] "geno4" "geno11" "geno15"
[4,] "geno6" "geno3" "geno12"
[5,] "geno10" "geno2"
                         "geno9"
```

### 3.9 AUGMENTED BLOCK DESIGNS

These are designs for two types of treatments: the control treatments (common) and the increased treatments. The common treatments are applied in complete randomized blocks, and the increased treatments, at random. Each treatment should be applied in any block once only. It is understood that the common treatments are of a greater interest; the standard error of the difference is much smaller than when between two increased ones in different blocks. The function design.dau() achieves this purpose.

```
block: 4
[1] "A" "C" "D" "B" "y"
block: 5
[1] "z" "A" "B" "D" "C"
print(plan8)
  plots block trt
   101 1 A
1
2
    102
          1 t
3
    103
          1 B
          1 D
4
   104
   132
         5 D
32
```

For augmented randomized complete block designs, use the function design.crd().

### 3.10 SPLIT-PLOT DESIGNS

These designs have two factors, one is applied in plots and is defined as A in a randomized complete block design; and a second factor, which is applied in the subplots of each plot applied at random. The function design.split() permits to find the experimental plan for this design.

```
t1<-c("A", "B", "C", "D")
t2<-c("a","b","c")
plan9
                  <-design.split(t1,t2,r=3,number=101,seed=45,
first=FALSE)
print(plan9)
  plots block t1 t2
    101 1 A b
          1 A a
2
    101
          1 A c
3
   101
          1 в с
4
    102
36 112
        3 D c
p<-plan9$t1[seq(1,36,3)]</pre>
q<-NULL
for (i in 1:12) q<-c(q,paste(plan9$t2[3*(i-1)+1],plan9$t2[3*(i-1)+1]]
1) +2], plan9$t2[3*(i-1)+3])
In the plots
> print(t(matrix(p,c(4,3))))
    [,1] [,2] [,3] [,4]
[1,] "A" "B"
             "C" "D"
[2,] "B" "D" "C" "A"
[3,] "A" "B" "C" "D"
> print(t(matrix(q,c(4,3))))
    [,1] [,2] [,3]
[1,] "b a c" "c b a" "a b c" "b a c"
[2,] "acb" "acb" "bac" "bac"
[3,] "c b a" "c a b" "a c b" "a b c"
```

### STRIP-PLOT DESIGNS

These designs are used when there are two types of treatments (factors) and are applied separately in large plots, called bands, in a vertical and horizontal direction of the block, obtaining the divided blocks. Each block constitutes a repetition.

```
t1<-c("A", "B", "C")
t2<-c("a", "b", "c", "d")
plan10 <-design.strip(t1,t2,r=3,number=101,seed=45)</pre>
print(plan10)
  plots block t1 t2
1
    101 1 B b
2
    102
            1 в а
    103
3
           1 B d
    104
           1 в с
4
   136 3 A c
36
t3<-paste(plan10$t1,plan10$t2)
B1 < -t (matrix (t3[1:12], c(4,3)))
B2 < -t (matrix(t3[13:24],c(4,3)))
B3 < -t (matrix (t3[25:36], c(4,3)))
> print(B1)
[,1] [,2] [,3] [,4] [1,] "B b" "B a" "B d" "B c"
[2,] "C b" "C a" "C d" "C c"
[3,] "A b" "A a" "A d" "A c"
> print(B2)
[,1] [,2] [,3] [,4] [1,] "A c" "A a" "A d" "A b"
[2,] "C c" "C a" "C d" "C b"
[3,] "B c" "B a" "B d" "B b"
> print(B3)
    [,1] [,2] [,3] [,4]
[1,] "B d" "B a" "B b" "B c"
[2,] "C d" "C a" "C b" "C c"
[3,] "A d" "A a" "A b" "A c"
```

### 4 MULTIPLE COMPARISONS

For the analyses, the following functions of 'agricolae' are used: LSD.test(), HSD.test(), duncan.test(), scheffe.test, waller.test, SNK.test() (Steel, 1996) and durbin.test(), kruskal(), friedman() and waerden.test (Conover, 1999).

For every statistical analysis, the data should be organized in columns. For the demonstration, the 'agricolae' database will be used.

The 'sweetpotato' data correspond to a completely random experiment in field with plots of 50 sweet potato plants, subjected to the virus effect and to a control without virus (See the reference manual of the package).

```
data(sweetpotato)
model<-aov(yield~virus, data=sweetpotato)</pre>
```

```
cv.model(model)
[1] 17.16660
attach(sweetpotato)
mean(yield)
[1] 27,625
```

Model parameters: Degrees of freedom and variance of the error:

```
df<-df.residual(model)
MSerror<-deviance(model)/df</pre>
```

### 4.1 THE LEAST SIGNIFICANT DIFFERENCE (LSD)

It includes the multiple comparison through the method of the minimum significant difference (Least Significant Difference), (Steel, 1997).

```
# comparison <- LSD.test(yield, virus, df, MSerror)</pre>
LSD.test(model, "virus",console=TRUE)
Study:
LSD t Test for yield
Mean Square Error: 22.48917
virus, means and individual (95 %) CI
      yield std.err r
                                     UCL Min. Max.
                             LCL
cc 24.40000 2.084067 3 19.594134 29.20587 21.7 28.5
fc 12.86667 1.246774 3 9.991602 15.74173 10.6 14.9
ff 36.33333 4.233727 3 26.570341 46.09633 28.0 41.8
00 36.90000 2.482606 3 31.175100 42.62490 32.1 40.4
alpha: 0.05; Df Error: 8
Critical Value of t: 2.306004
Least Significant Difference 8.928965
Means with the same letter are not significantly different.
Groups, Treatments and means
        00
                36.9
                 36.33
         ff
а
         CC
                 24.4
b
         fc
                 12.87
C
```

In the function LSD.test(), the multiple comparison was carried out. In order to obtain the probabilities of the comparisons, it should be indicated that groups are not required; thus:

```
# comparison <- LSD.test(yield, virus,df, MSerror, group=F)
comparison <-LSD.test(model, "virus", group=F,console=TRUE)
LSD t Test for yield</pre>
```

Mean Square Error: 22.48917

## virus, means and individual ( 95 %) CI

yield std.err r LCL UCL Min. Max. cc 24.40000 2.084067 3 19.594134 29.20587 21.7 28.5 fc 12.86667 1.246774 3 9.991602 15.74173 10.6 14.9 ff 36.33333 4.233727 3 26.570341 46.09633 28.0 41.8 co 36.90000 2.482606 3 31.175100 42.62490 32.1 40.4

alpha: 0.05; Df Error: 8 Critical Value of t: 2.306004

### Comparison between treatments means

```
Difference
                         pvalue sig
                                          LCL
                                                     UCL
                                    2.604368 20.462299
cc - fc 11.5333333 0.0176377595 *
                                              -3.004368
                                 * -20.862299
cc - ff -11.9333333 0.0150730851
                                * -21.428965 -3.571035
cc - oo -12.5000000 0.0120884239
fc - ff -23.4666667 0.0003023690 *** -32.395632 -14.537701
fc - oo -24.0333333 0.0002574929 *** -32.962299 -15.104368
                                 -9.495632
ff - oo -0.5666667 0.8872673216
                                              8.362299
```

### The significance code "sig" is interpreted as:

```
***: p.valor < 0.001

**: 0.001 <p.valor < 0.01

*: 0.01 < p.valor < 0.05

:: 0.05 < p.valor < 0.10
```

### > comparison

### \$statistics

Mean CV MSerror 27.625 17.1666 22.48917

### \$parameters

Df ntr t.value 8 4 2.306004

### \$means

```
yield std.err r LCL UCL Min. Max. cc 24.40000 2.084067 3 19.594134 29.20587 21.7 28.5 fc 12.86667 1.246774 3 9.991602 15.74173 10.6 14.9 ff 36.33333 4.233727 3 26.570341 46.09633 28.0 41.8 co 36.90000 2.482606 3 31.175100 42.62490 32.1 40.4
```

### \$comparison

```
Difference
                         pvalue sig
                                           LCL
                                                      UCL
cc - fc 11.5333333 0.0176377595 *
                                      2.604368
                                                20.462299
cc - ff -11.9333333 0.0150730851
                                  * -20.862299
                                               -3.004368
cc - oo -12.5000000 0.0120884239
                                 * -21.428965 -3.571035
fc - ff -23.4666667 0.0003023690 *** -32.395632 -14.537701
fc - oo -24.0333333 0.0002574929 *** -32.962299 -15.104368
ff - oo -0.5666667 0.8872673216
                                     -9.495632
                                                 8.362299
```

\$groups NULL

### 4.2 BONFERRONI

With the function LSD.test() we can make adjustments to the probabilities found, as for example the adjustment by Bonferroni.

```
LSD.test(model, "virus", group=F, p.adj= "bon", console=TRUE)
LSD t Test for yield
P value adjustment method: bonferroni
alpha: 0.05; Df Error: 8
Critical Value of t: 3.478879
Comparison between treatments means
         Difference pvalue sig
                                      LCL
cc - fc 11.5333333 0.105827
                                   -1.937064 25.0037305
cc - ff -11.9333333 0.090439
                                . -25.403730
                                                1.5370638
                                . -25.970397
cc - oo -12.5000000 0.072531
                                                0.9703971
cc - oo -12.5000000 0.072531 . -25.970397 0.9703971 fc - ff -23.4666667 0.001814 ** -36.937064 -9.9962695
fc - oo -24.0333333 0.001545 ** -37.503730 -10.5629362
ff - oo -0.5666667 1.000000
                                  -14.037064 12.9037305
```

Other comparison tests can be applied, such as "duncan", "Student-Newman-Keuls", "tukey", and "waller-duncan."

For "duncan", use the function duncan.test(); for "Student-Newman-Keuls", the function SNK.test(); for "tukey", the function HSD.test(); for "scheffe", the function scheffe.test(); and for "waller-duncan", the function waller.test(). The parameters are the same. "Waller" also requires the value of F-calculated of the ANOVA treatments. If the model is used as a parameter, this is no longer necessary.

### 4.3 DUNCAN'S NEW MULTIPLE-RANGE TEST

It corresponds to the Duncan's Test (Steel, 1997).

```
duncan.test(model, "virus", console=TRUE)

Duncan's new multiple range test
for yield

Mean Square Error: 22.48917

virus, means

    yield std.err r Min. Max.
cc 24.40000 2.084067 3 21.7 28.5
fc 12.86667 1.246774 3 10.6 14.9
ff 36.33333 4.233727 3 28.0 41.8
oo 36.90000 2.482606 3 32.1 40.4

alpha: 0.05; Df Error: 8
```

```
Critical Range
8.928965 9.304825 9.514910
Means with the same letter are not significantly different.
Groups, Treatments and means
        00
                36.9
        ff
                36.33
        CC
                24.4
        fc
                12.87
duncan.test(model, "virus", group=FALSE,console=TRUE)
Duncan's new multiple range test
for yield
Mean Square Error: 22.48917
alpha: 0.05; Df Error: 8
Critical Range
      2
               3
8.928965 9.304825 9.514910
Comparison between treatments means
        Difference pvalue sig
                                       LCL
cc - fc 11.5333333 0.017638 *
                                 2.604368 20.462299
                            * -20.862299 -3.004368
cc - ff -11.9333333 0.015073
                             * -21.804825 -3.195175
cc - oo -12.5000000 0.014544
fc - ff -23.4666667 0.000388 *** -32.771492 -14.161842
fc - oo -24.0333333 0.000387 *** -33.548244 -14.518423
ff - oo -0.5666667 0.887267
                                 -9.495632
                                            8.362299
```

### 4.4 STUDENT-NEWMAN-KEULS

Student, Newman and Keuls helped to improve the Newman-Keuls test of 1939, which was known as the Keuls method (Steel, 1997).

```
SNK.test(model, "virus", alpha=0.05,console=TRUE)

Student Newman Keuls Test
for yield

Mean Square Error: 22.48917

virus, means

yield std.err r Min. Max.
cc 24.40000 2.084067 3 21.7 28.5
fc 12.86667 1.246774 3 10.6 14.9
ff 36.33333 4.233727 3 28.0 41.8
oo 36.90000 2.482606 3 32.1 40.4

alpha: 0.05; Df Error: 8
```

```
Critical Range
 8.928965 11.064170 12.399670
Means with the same letter are not significantly different.
Groups, Treatments and means
        00
                36.9
        ff
                36.33
        CC
                24.4
        fc
                12.87
SNK.test(model, "virus", group=FALSE,console=TRUE)
Student Newman Keuls Test
for yield
Mean Square Error: 22.48917
alpha: 0.05; Df Error: 8
Critical Range
                 3
       2
 8.928965 11.064170 12.399670
Comparison between treatments means
      Difference pvalue sig
                                     LCL
cc-fc 11.5333333 0.017638 * 2.604368 20.462299
cc-ff -11.9333333 0.015073 * -20.862299 -3.004368
cc-oo -12.5000000 0.029089 * -23.564170 -1.435830
fc-ff -23.4666667 0.000777 *** -34.530836 -12.402497
fc-oo -24.0333333 0.001162 ** -36.433003 -11.633664
ff-oo -0.5666667 0.887267
                              -9.495632 8.362299
```

### 4.5 TUKEY'S W PROCEDURE (HSD)

This studentized range test, created by Tukey in 1953, is known as the Tukey's HSD (Honestly Significant Differences) Test (Steel, 1997).

```
comparison1 <- HSD.test(model, "virus", console=TRUE)

HSD Test for yield

Mean Square Error: 22.48917

virus, means

        yield std.err r Min. Max.

cc 24.40000 2.084067 3 21.7 28.5

fc 12.86667 1.246774 3 10.6 14.9

ff 36.33333 4.233727 3 28.0 41.8

oo 36.90000 2.482606 3 32.1 40.4

alpha: 0.05; Df Error: 8

Critical Value of Studentized Range: 4.52881</pre>
```

```
Honestly Significant Difference: 12.39967
Means with the same letter are not significantly different.
Groups, Treatments and means
        00
               36.9
ab
        ff
                36.33
bс
        CC
                24.4
С
        fc
                12.87
> comparison1
$statistics
   Mean CV MSerror
                             HSD
 27.625 17.1666 22.48917 12.39967
$parameters
 Df ntr StudentizedRange
                 4.52881
$means
    yield std.err r Min. Max.
cc 24.40000 2.084067 3 21.7 28.5
fc 12.86667 1.246774 3 10.6 14.9
ff 36.33333 4.233727 3 28.0 41.8
00 36.90000 2.482606 3 32.1 40.4
$comparison
NULL
$groups
 trt
        means M
1 oo 36.90000 a
2 ff 36.33333 ab
3 cc 24.40000 bc
4 fc 12.86667 c
```

### 4.6 WALLER-DUNCAN'S BAYESIAN K-RATIO t-TEST

In 1975, Duncan continued the multiple comparison procedures, introducing the criterion of minimizing both experimental errors; for this, he used the Bayes' theorem, obtaining one new test called Waller-Duncan (Steel, 1997).

```
attach (sweetpotato)
then:
waller.test(yield, virus, df, MSerror, Fc= 17.345,
group=F,console=TRUE)
In another case with only invoking the model object:
comparison2 <- waller.test(model, "virus",</pre>
group=FALSE, console=TRUE)
Waller-Duncan K-ratio t Test for yield
This test minimizes the Bayes risk under additive
loss and certain other assumptions.
K ratio
                         100.00000
Error Degrees of Freedom 8.00000
Error Mean Square
                        22.48917
F value
                         17.34478
Critical Value of Waller 2.23600
virus, means
      yield std.err r Min. Max.
cc 24.40000 2.084067 3 21.7 28.5
fc 12.86667 1.246774 3 10.6 14.9
ff 36.33333 4.233727 3 28.0 41.8
00 36.90000 2.482606 3 32.1 40.4
Minimum Significant Difference 8.657906
Comparison between treatments means
        Difference significant
cc - fc 11.5333333
                          TRUE
ff - cc 11.9333333
                         TRUE
oo - cc 12.5000000
                         TRUE
ff - fc 23.466667
                         TRUE
oo - fc 24.0333333
                         TRUE
oo - ff 0.5666667
                         FALSE
```

It is indicated that the virus effect "ff" is not significant to the control "oo."

The found object "compare" has information to make other procedures.

### 4.7 SCHEFFE'S TEST

This method, created by Scheffe in 1959, is very general for all the possible contrasts and their confidence intervals. The confidence intervals for the averages are very broad, resulting in a very conservative test for the comparison between treatment averages (Steel, 1997).

```
# analysis of variance:
model<-aov(yield~virus, data=sweetpotato)</pre>
scheffe.test(model, "virus", group=TRUE, console=TRUE,
main="Yield of sweetpotato\nDealt with different virus")
Study: Yield of sweetpotato
Dealt with different virus
Scheffe Test for yield
Mean Square Error : 22.48917
virus, means
      yield std.err r Min. Max.
cc 24.40000 2.084067 3 21.7 28.5
fc 12.86667 1.246774 3 10.6 14.9
ff 36.33333 4.233727 3 28.0 41.8
00 36.90000 2.482606 3 32.1 40.4
alpha: 0.05; Df Error: 8
Critical Value of F: 4.066181
Minimum Significant Difference: 13.52368
Means with the same letter are not significantly different.
Groups, Treatments and means
            36.9
        00
         ff
                 36.33
ab
        CC
                24.4
        fc
                12.87
```

The minimum significant value is very high.

If you require the approximate probabilities of comparison, you can use the option Group=FALSE.

```
comparison3 <- scheffe.test(model,"virus", group=FALSE,
console=TRUE)

Study:
Scheffe Test for yield
Mean Square Error : 22.48917</pre>
```

```
virus, means
      yield std.err r Min. Max.
cc 24.40000 2.084067 3 21.7 28.5
fc 12.86667 1.246774 3 10.6 14.9
ff 36.33333 4.233727 3 28.0 41.8
00 36.90000 2.482606 3 32.1 40.4
alpha: 0.05; Df Error: 8
Critical Value of F: 4.066181
Comparison between treatments means
         Difference pvalue sig
                                       LCL
                                                    UCL
cc - fc 11.5333333 0.097816 . -1.000348 24.0670149
cc - ff -11.9333333 0.085487
                             . -24.467015 0.6003483
                             . -25.033682 0.0336816
** -36.000348 -10.9329851
cc - oo -12.5000000 0.070607
fc - ff -23.4666667 0.002331
fc - oo -24.0333333 0.001998 ** -36.567015 -11.4996517
                              -13.100348 11.9670149
ff - oo -0.5666667 0.999099
```

### 4.8 MULTIPLE COMPARISON IN FACTORIAL TREATMENTS

In a factorial combined effects of the treatments. Comparetive tests: LSD, HSD, Waller-Duncan, Duncan, Scheffé, SNK can be applied.

```
\# model <-aov (y ~ A * B * C, data)
\# compare <-LSD.test (model, c ("A", "B", "C"),console=TRUE)
```

The comparison is the combination of A:B:C.

Data RCBD design with a factorial clone x nitrogen. The response variable yield:

```
yield < -scan (text =
 "6 7 9 13 16 20 8 8 9
  7 8 8 12 17 18 10 9 12
  9 9 9 14 18 21 11 12 11
 8 10 10 15 16 22 9 9 9 "
block \leftarrowgl (4, 9)
clone <-rep (gl (3, 3, labels = c ("c1", "c2", "c3")), 4)
nitrogen <-rep (gl (3, 1, labels = c ("n1", "n2", "n3")), 12)</pre>
A <-data.frame (block, clone, nitrogen, yield)
head (A)
 block clone nitrogen yield
          с1
                    n1
2
                    n2
                            7
           с1
3
           c1
                    n3
                           9
      1
          c2
                    n1
                           13
5
      1
           c2
                    n2
                           16
           c2
                   n3
                           20
model <-aov (yield ~ block + clone * nitrogen, data = A)</pre>
anova (model)
```

### Analysis of Variance Table Response: yield Df Sum Sq Mean Sq F value block 3 20.75 6.917 5.8246 0.0038746 \*\* 2 497.72 248.861 209.5673 6.370e-16 \*\*\* clone 2 54.06 27.028 22.7602 2.865e-06 \*\*\* nitrogen clone:nitrogen 4 43.28 10.819 9.1111 0.0001265 \*\*\* 24 28.50 Residuals 1.187 Signif. codes: 0 \\*\*\*' 0.001 \\*\*' 0.01 \\*' 0.05 \'.' 0.1 \' 1 out <-LSD.test (model, c("clone", "nitrogen"), main = "Yield ~ block</pre> + nitrogen + clone + clone:nitrogen", console=TRUE) Study: Yield ~ block + nitrogen + clone + clone:nitrogen LSD t Test for yield Mean Square Error: 1.1875 clone:nitrogen, means and individual (95 %) CI

yield std.err r LCL UCL Min. Max. c1:n1 7.50 0.6454972 4 6.167759 8.832241 6 c1:n2 8.50 0.6454972 4 7.167759 9.832241 7 10 c1:n3 9.00 0.4082483 4 8.157417 9.842583 8 10 c2:n1 13.50 0.6454972 4 12.167759 14.832241 12 15 c2:n2 16.75 0.4787136 4 15.761984 17.738016 16 c2:n3 20.25 0.8539126 4 18.487611 22.012389 18 22 c3:n1 9.50 0.6454972 4 8.167759 10.832241 8 11 8 c3:n2 9.50 0.8660254 4 7.712611 11.287389 12 c3:n3 10.25 0.7500000 4 8.702076 11.797924 9 12

alpha: 0.05; Df Error: 24 Critical Value of t: 2.063899

Least Significant Difference 1.590341 Means with the same letter are not significantly different.

```
Groups, Treatments and means
        c2:n3 20.25
а
        c2:n2
               16.75
С
        c2:n1
               13.5
               10.25
        c3:n3
        c3:n1
                9.5
de
        c3:n2
               9.5
de
def
        c1:n3
ef
        c1:n2
               8.5
f
        c1:n1
               7.5
par(mar=c(3,3,2,0))
pic1<-bar.err(out$means, variation="range", ylim=c(5, 25),
bar=FALSE, col=0, las=1)
points(pic1$index,pic1$means,pch=18,cex=1.5,col="blue")
axis(1,pic1$index,labels=FALSE)
title(main="average and range\nclon:nitrogen")
```

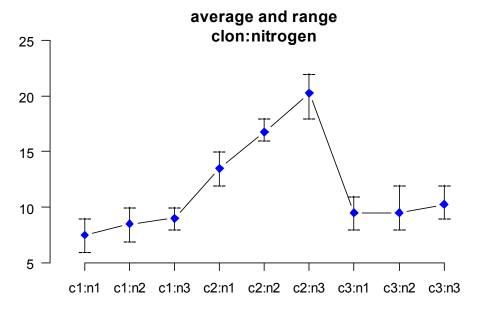


Figure 4.1 Combined clone:nitrogen

### 4.9 GRAPHICS OF THE MULTIPLE COMPARISON

The results of a comparison can be graphically seen with the functions bar.group() and bar.err().

The found object of one comparison is the entry for these functions, Figure 4.2. The objects compare1 and compare2 are used in the following exercise:

comparison1, for the functions bar.group() and bar.err() comparison2, for the function bar.err()

```
par(mfrow=c(2,2))
c1<-colors()[480]; c2=colors()[65]; c3=colors()[15];
c4=colors()[140]
G1<-bar.group(comparison1$groups, ylim=c(0,45),
main="Tukey\nG1",col=c1)
G2<-bar.group(comparison1$groups, horiz=T, xlim=c(0,45),
main="Tukey\nG2",col=c2)
G3<-bar.err(comparison2$means, variation="range",ylim=c(0,45),
col=c3, main="Range\nG3")
G4<-bar.err(comparison2$means, horiz=T, xlim=c(0,45), col=c4,
variation="SE",,main="Standard error \nG4")</pre>
```

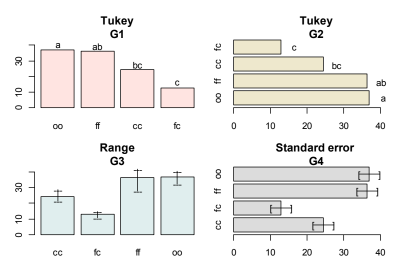


Figure 4.2 Comparison between treatments

### 4.10 ANALYSIS OF BALANCED INCOMPLETE BLOCKS

This analysis can come from balanced or partially balanced designs. The function BIB.test() is for balanced designs, and PBIB.test(), for partially balanced designs. In the following example, the 'agricolae' data will be used.

```
#Example linear estimation and design of experiments. (Joshi, 1987)
# Profesor de Estadistica, Institute of Social Sciences Agra, India
# 6 variedades de trigo en 10 bloques de 3 parcelas cada una.
block < -gl(10,3)
variety<-c(1,2,3,1,2,4,1,3,5,1,4,6,1,5,6,2,3,6,2,4,5,2,5,6,3,4,5,3,
4,6)
y < -c(69, 54, 50, 77, 65, 38, 72, 45, 54, 63, 60, 39, 70, 65, 54, 65, 68, 67, 57, 60, 62,
59, 65, 63, 75, 62, 61, 59, 55, 56)
BIB.test(block, variety, y,console=TRUE)
ANALYSIS BIB: y
Class level information
Block: 1 2 3 4 5 6 7 8 9 10
Trt : 1 2 3 4 5 6
Number of observations:
Analysis of Variance Table
Response: y
            Df
                Sum Sq Mean Sq F value Pr(>F)
block.unadj
             9
                466.97
                       51.885 0.9019 0.54712
trt.adj
             5 1156.44 231.289
                                 4.0206 0.01629 *
Residuals
            15
                862.89 57.526
                0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Signif. codes:
coefficient of variation: 12.6 %
y Means: 60.3
```

```
variety, statistics
     y mean.adj
                     SE r std.err Min. Max.
1 70.2 75.13333 3.728552 5 2.267157 63 77
2 60.0 58.71667 3.728552 5 2.190890
                                    54
                                         65
3 59.4 58.55000 3.728552 5 5.537147
                                   45
                                         75
4 55.0 54.96667 3.728552 5 4.404543
                                         62
                                    38
5 61.4 60.05000 3.728552 5 2.014944
                                    54
                                         65
                                    39 67
6 55.8 54.38333 3.728552 5 4.810405
LSD test
Std.diff : 5.363111
      : 0.05
Alpha
LSD
          : 11.4312
Parameters BIB
Lambda
treatmeans : 6
Block size : 3
Blocks
Replication: 5
Efficiency factor 0.8
<<< Book >>>
Means with the same letter are not significantly different.
Comparison of treatments
Groups, Treatments and means
        1
               75.13
b
        5
               60.05
       2
               58.72
       3
               58.55
b
        4
                54.97
b
        6
                54.38
```

function (block, trt, y, test = c("lsd", "tukey", "duncan", "waller", "snk"), alpha = 0.05, group = TRUE). LSD, Tukey Duncan, Waller-Duncan and SNK, can be used. The probabilities of the comparison can also be obtained. It should only be indicated: group=FALSE, thus:

```
out <-BIB.test(block, trt=variety, y, test="tukey",
group=FALSE, console=TRUE)

ANALYSIS BIB: y
Class level information

Block: 1 2 3 4 5 6 7 8 9 10
Trt : 1 2 3 4 5 6</pre>
Number of observations: 30
```

```
Analysis of Variance Table
Response: y
           Df Sum Sq Mean Sq F value Pr(>F)
block.unadj 9 466.97 51.885 0.9019 0.54712
            5 1156.44 231.289 4.0206 0.01629 *
trt.adj
Residuals
           15 862.89 57.526
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 '' 1
coefficient of variation: 12.6 %
y Means: 60.3
variety, statistics
    y mean.adj
                     SE r std.err Min. Max.
1 70.2 75.13333 3.728552 5 2.267157 63 77
2 60.0 58.71667 3.728552 5 2.190890
                                     54
                                          65
3 59.4 58.55000 3.728552 5 5.537147
                                    45
                                         75
4 55.0 54.96667 3.728552 5 4.404543
                                   38 62
5 61.4 60.05000 3.728552 5 2.014944 54 65
6 55.8 54.38333 3.728552 5 4.810405
                                   39 67
Tukey
Alpha
         : 0.05
Std.err
         : 3.792292
          : 17.42458
Parameters BIB
Lambda
treatmeans : 6
Block size : 3
Blocks
       : 10
Replication: 5
Efficiency factor 0.8
<<< Book >>>
Comparison between treatments means
     Difference pvalue sig
1 - 2 16.4166667 0.070509
1 - 3 16.5833333 0.066649
1 - 4 20.1666667 0.019092
1 - 5 15.0833333 0.109602
1 - 6 20.7500000 0.015510
2 - 3 0.1666667 1.000000
2 - 4 3.7500000 0.979184
2 - 5 -1.3333333 0.999840
2 - 6 4.3333333 0.961588
3 - 4 3.5833333 0.982927
3 - 5 -1.5000000 0.999715
3 - 6 4.1666667 0.967375
4 - 5 -5.0833333 0.927273
4 - 6 0.5833333 0.999997
5 - 6 5.6666667 0.890815
```

The found "outl" object can be used for the functions bar.group() and bar.err() for the bar graphics, in the same way as previously.

```
names(out)
[1] "parameters" "statistics" "comparison" "means" "groups"
bar.group: out$groups
bar.err: out$means
```

### 4.11 PARTIALLY BALANCED INCOMPLETE BLOCKS

The function PBIB.test() (Joshi, 1987) can be used for the lattice and alpha designs.

Consider the following case: Construct the alpha design with 30 treatments, 2 repetitions, and a block size equal to 3.

```
library(agricolae)
library(MASS)
library(lme4)
# alpha design
Genotype<-paste("geno",1:30, sep="")</pre>
r<-2
k < -3
plan<-design.alpha(Genotype,k,r,seed=5)</pre>
alpha design (0,1) - Serie I
Parameters Alpha design
______
treatmeans : 30
Block size : 3
Blocks : 10
Replication: 2
Efficiency factor
(E ) 0.6170213
<<< Book >>>
```

The generated plan is plan\$book.

Suppose that the corresponding observation to each experimental unit is:

```
yield <-c(5,2,7,6,4,9,7,6,7,9,6,2,1,1,3,2,4,6,7,9,8,7,6,4,3,2,2,1,1,2,1,1,2,4,5,6,7,8,6,5,4,3,1,1,2,5,4,2,7,6,6,5,6,4,5,7,6,5,5,4)
```

The data table is constructed for the analysis. In theory, it is presumed that a design is applied and the experiment is carried out; subsequently, the study variables are observed from each experimental unit.

```
data<-data.frame(plan$book, yield)
rm(yield, Genotype)

The analysis:

attach(data)
model <- PBIB.test(block, Genotype, replication, yield, k=3,
group=TRUE, console=TRUE)</pre>
```

### detach (data)

ANALYSIS PBIB: yield

Class level information

block : 20
Genotype : 30

Number of observations: 60

Estimation Method: REML

Parameter Estimates

Variance block:replication 2.834033 replication 0.000000 Residual 2.003098

Fit Statistics

-2 Res Log Likelihood 147.6594 AIC 215.6594 BIC 286.8671

Analysis of Variance Table

Response: yield

Df Sum Sq Mean Sq F value Pr(>F)
Genotype 29 72.006 2.4830 1.2396 0.3668

Residuals 11 22.034 2.0031

coefficient of variation: 31.2 %

yield Means: 4.533333

Parameters PBIB

Genotype 30 block size 3 block/replication 10 replication 2

Efficiency factor 0.6170213

Means with the same letter are not significantly different.

Groups, Treatments and means

7.729 20 13 6.715 ab ab 1 6.505 6.192 abc 8 6.032 24 abcd 5.735 abcd 23 5.473 10 abcd 16 5.455 abcd 21 5.14 abcd abcd 22 5.069 abcd 4 4.874 3 4.794 abcd abcd 14 4.742

```
15
                   4.587
abcd
          27
                   4.563
abcd
abcd
          7
                   4.424
abcd
          5
                   4.286
abcd
          19
                   4.198
abcd
          6
                   4.165
          25
                   3.978
abcd
                   3.943
abcd
          17
bcd
                   3.628
bcd
          28
                   3.495
bcd
          11
                   3.379
bcd
          26
                   3.341
bcd
                   3.052
bcd
          30
bcd
          12
                   2.879
cd
          29
                   2.44
d
          18
                   2.186
```

Comparison between treatments means and its name

<<< to see the objects: means, comparison and groups. >>>

### The adjusted averages can be extracted from the model.

### model\$means

	yield	trt	mean.adj	SE	r	std.err	Min.	Max.
geno1	7.5	1	6.504752	1.313644	2	1.5	6	9
geno10	4.5	2	3.628197	1.313644	2	0.5	4	5
geno11	5.5	3	4.793619	1.310726	2	0.5	5	6
geno12	4.0	4	4.873879	1.313644	2	3.0	1	7
geno8	3.0	29	2.439878	1.310726	2	1.0	2	4
geno9	3.5	30	2.999638	1.310726	2	1.5	2	5

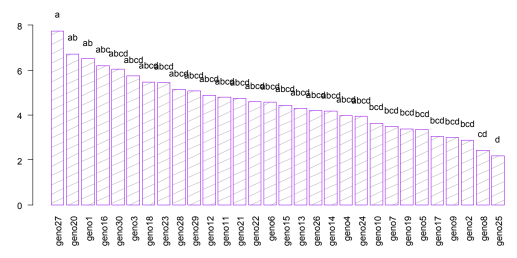


Figure 4.3. Treatment Groups.

### The comparisons:

### model\$comparison

```
Difference
                      stderr
                                pvalue
         2.87655507 1.844369 0.147134
1 - 2
1
   3
         1.71113250 1.576446 0.300944
1
   4
         1.63087260 1.727016 0.365282
1
    5
         2.21879565 1.853044 0.256324
27 - 30
         1.56381175 1.812351 0.406632
    29
         1.05522604 1.850095 0.579894
28 - 30
        0.49546622 1.847786 0.793552
29 - 30 -0.55975982 1.587129 0.730988
```

The data on the adjusted averages and their standard error can be illustrated (figure 4.2), since the created object is very similar to the objects generated by the multiple comparisons.

```
par(mfrow=c(2,2),cex=0.6)
C1<-bar.err(model$means[1:7, ], ylim=c(0,9), col=0, main="C1",
variation="range",border=3)
C2<-bar.err(model$means[8:15,], ylim=c(0,9), col=0, main="C2",
variation="range", border =4)
C3<-bar.err(model$means[16:22,], ylim=c(0,9), col=0, main="C3",
variation="range",border =2)
C4<-bar.err(model$means[23:30,], ylim=c(0,9), col=0, main="C4",
variation="range", border =6)</pre>
```

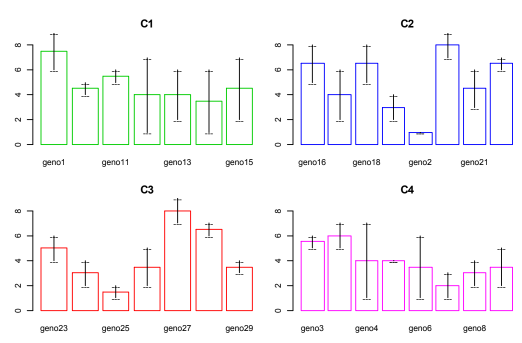


Figure 4.4. Range in each treatment.

Analysis of balanced lattice 3x3, 9 treatments, 4 repetitions.

### Create the data in a text file: latice3x3.txt and read with R:

```
sqr block trt yield
      1
          1 48.76
                         1
                               1
                                   4 14.46
                                                  1
                                                        1
                                                             3 19.68
1
      2
          8 10.83
                               2
                                   6 30.69
                                                        2
                         1
                                                  1
                                                            7 31.00
1
      3
          5 12.54
                                                        3
                         1
                               3
                                   9 42.01
                                                  1
                                                            2 23.00
2
                         2
      4
          5 11.07
                               4
                                  8 22.00
                                                  2
                                                            1 41.00
2
      5
          2 22.00
                         2
                               5
                                   7 42.80
                                                  2
                                                        5
                                                            3 12.90
2
                         2
      6
          9 47.43
                               6
                                   6 28.28
                                                  2
                                                        6
                                                            4 49.95
3
      7
          2 27.67
                         3
                               7
                                   1 50.00
                                                  3
                                                        7
                                                             6 25.00
3
         7 30.00
                         3
                              8
                                   5 24.00
                                                  3
                                                        8
                                                             4 45.57
3
      9
          3 13.78
                         3
                               9
                                  8 24.00
                                                  3
                                                        9
                                                             9 30.00
4
     10
          6 37.00
                         4
                              10
                                   3 15.42
                                                  4
                                                       10
                                                             5 20.00
                         4
                                                  4
4
     11
          4 42.37
                              11
                                   2 30.00
                                                       11
                                                             8 18.00
                                                            1 43.81
4
     12
          9 39.00
                         4
                              12
                                   7 23.80
                                                  4
                                                       12
```

```
library(agricolae)
library (MASS)
library(lme4)
A<-read.table("latice3x3.txt", header=T)
attach(A)
model2<-PBIB.test(block,trt,sqr,yield,k=3,console=TRUE)</pre>
detach(A)
ANALYSIS PBIB: yield
Class level information
block: 12
trt : 9
Number of observations: 36
Estimation Method: REML
Parameter Estimates
          Variance
block:sqr 0.00000
sqr
           0.00000
Residual 56.93724
                      Fit Statistics
-2 Res Log Likelihood
                            198.2320
                            224.2320
ATC.
BIC
                            244.8177
Analysis of Variance Table
Response: yield
          Df Sum Sq Mean Sq F value
                                        Pr(>F)
           8 3749.4 468.68 8.2315 0.0001987 ***
trt
Residuals 16 911.0
                     56.94
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
coefficient of variation: 25.9 %
yield Means: 29.16167
Parameters PBIB
```

```
trt 9
block size 3
block/sqr 3
sqr 4
```

Efficiency factor 0.75

Means with the same letter are not significantly different.

#### Groups, Treatments and means 1 45.89 а ab 9 39.61 ab 4 38.09 7 31.9 bc bс 6 30.24 2 25.67 cd d 8 18.71 d 5 16.9 d 3 15.44

Comparison between treatments means and its name

```
<<< to see the objects: means, comparison and groups. >>>
```

```
par(mar=c(3,3,0,0))
```

bar.group(model2\$group,ylim=c(0,52),density=10)

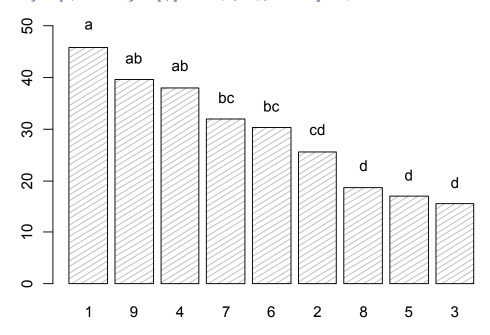


Figure 4.5. Treatment Groups.

### model2\$means

```
      4
      38.0875
      4
      38.0875
      3.772839
      4
      8.027584
      14.46
      49.95

      5
      16.9025
      5
      16.9025
      3.772839
      4
      3.068910
      11.07
      24.00

      6
      30.2425
      6
      30.2425
      3.772839
      4
      2.536390
      25.00
      37.00

      7
      31.9000
      7
      31.9000
      3.772839
      4
      2.906947
      23.80
      42.80

      8
      18.7075
      3.772839
      4
      2.906984
      10.83
      24.00

      9
      39.6100
      9
      39.6100
      3.772839
      4
      3.647335
      30.00
      47.43
```

#### model2\$comparison

```
Difference stderr pvalue
1 - 2 20.2250 5.335599 0.001604
1 - 3 30.4475 5.335599 0.000032
1 - 4 7.8050 5.335599 0.162884
1 - 5 28.9900 5.335599 0.000056
1 - 6 15.6500 5.335599 0.009746
.....
8 - 9 -20.9025 5.335599 0.001228
```

## **4.12 AUGMENTED BLOCKS**

The function DAU.test() can be used for the analysis of the augmented block design.

The data should be organized in a table, containing the blocks, treatments, and the response.

```
block<-c(rep("I",7),rep("II",6),rep("III",7))
trt<-c("A", "B", "C", "D", "q", "k", "1", "A", "B", "C", "D", "e", "i", "A", "B",
"C", "D", "f", "h", "j")
yield<-c(83,77,78,78,70,75,74,79,81,81,91,79,78,92,79,87,81,89,96,
82)
data.frame(block, trt, yield)
   block trt yield
           Α
1
       I
                 83
2
       Ι
           B
                 77
           С
3
       I
                 78
4
       I
           D
                 78
5
                 70
       I
           g
                 75
6
      Ι
           k
7
      I
           1
                 74
8
      ΙI
           Α
                 79
9
      ΙI
           В
                 81
10
     ΙI
           С
                 81
11
     ΙI
                 91
           D
12
     ΙI
                 79
           е
13
     ΙI
                 78
14
     III
           Α
                 92
                 79
15
     III
           В
16
     III
           С
                 87
17
     III
           D
                 81
18
           f
                 89
     III
19
     III
                 96
           h
20
     TTT
           j
                 82
```

The treatments are in each block:

```
by(trt,block,as.character)
block: I
[1] "A" "B" "C" "D" "q" "k" "l"
block: II
[1] "A" "B" "C" "D" "e" "i"
_____
block: III
[1] "A" "B" "C" "D" "f" "h" "i"
With their respective responses:
by(yield,block,as.character)
block: I
[1] 83 77 78 78 70 75 74
block: II
[1] 79 81 81 91 79 78
______
block: III
[1] 92 79 87 81 89 96 82
model<- DAU.test(block,trt,yield,method="lsd",console=TRUE)</pre>
ANALYSIS DAU: yield
Class level information
Block: I II III
Trt: ABCDefghijkl
Number of observations: 20
ANOVA, Treatment Adjusted
Analysis of Variance Table
Response: yield
                           Df Sum Sq Mean Sq F value Pr(>F)
block.unadj
                            2 360.07 180.036
trt.adj 11 285.10 25.918 0.9609 0.5499
Control 3 52.92 17.639 0.6540 0.6092
Control + control.VS.aug. 8 232.18 29.022 1.0760 0.4779
Residuals 6 161.83 26.972
ANOVA, Block Adjusted
Analysis of Variance Table
Response: yield
                     Df Sum Sq Mean Sq F value Pr(>F)
                     11 575.67 52.333
2 69.50 34.750 1.2884 0.3424
trt.unadj
block.adj
                      3 52.92 17.639 0.6540 0.6092
Control
                      7 505.88 72.268 2.6793 0.1253
Augmented
Control vs augmented 1 16.88 16.875 0.6256 0.4591
                      6 161.83 26.972
Residuals
coefficient of variation: 6.4 %
```

yield Means: 81.5

```
Critical Differences (Between) Std Error Diff.
Two Control Treatments
                                            4.240458
Two Augmented Treatments (Same Block)
                                            7.344688
                                         8.211611
Two Augmented Treatments (Different Blocks)
A Augmented Treatment and A Control Treatment 6.360687
Means with the same letter are not significantly different.
Groups, Treatments and means
а
        h
              93.5
        f
                86.5
ab
        Α
               84.67
ab
        D
               83.33
ab
        С
ab
               82
ab
       j
                79.5
       В
ab
                79
ab
       е
                78.25
ab
        k
                78.25
        i
                77.25
ab
ab
        1
                77.25
b
        g
                73.25
Comparison between treatments means
<<< to see the objects: pvalue and means >>>
model$means
    yield std.err r Min. Max. mean.adj
                                             SE block
A 84.66667 11.532563 3 79 92 84.66667 2.998456
B 79.00000 3.464102 3 77 81 79.00000 2.998456
C 82.00000 7.937254 3 78 87 82.00000 2.998456
D 83.33333 11.789826 3 78 91 83.33333 2.998456
e 79.00000
              NA 1 79 79 78.25000 5.193479
                                                   ΙI
f 89.00000
                NA 1 89 89 86.50000 5.193479
                                                  III
g 70.00000
                NA 1 70 70 73.25000 5.193479
                                                   Т
                NA 1 96
                           96 93.50000 5.193479
h 96.00000
                                                  III
i 78.00000
                NA 1 78
                           78 77.25000 5.193479
                                                   ΤT
j 82.00000
                 NA 1 82
                           82 79.50000 5.193479
                                                  III
                 NA 1 75
                            75 78.25000 5.193479
k 75.00000
                                                   I
                 NA 1 74
                          74 77.25000 5.193479
1 74.00000
                                                    Ι
model<- DAU.test(block,trt,yield,method="lsd",group=F,console=TRUE)</pre>
> head(model$comparison,10)
     Difference pvalue sig
A - B 5.666667 0.229886
A - C 2.666667 0.552612
A - D 1.333333 0.763840
A - e 6.416667 0.352008
A - f -1.833333 0.782870
A - g 11.416667 0.122820
A - h -8.833333 0.214268
A - i 7.416667 0.287856
```

A - j 5.166667 0.447652 A - k 6.416667 0.352008

### 4.13 NON-PARAMETRIC COMPARISONS

The functions for non-parametric multiple comparisons included in 'agricolae' are: kruskal(), waerden.test(), friedman() and durbin.test() (Conover, 1999).

The function kruskal() is used for N samples (N>2), populations or data coming from a completely random experiment (populations = treatments).

The function waerden.test(), similar to kruskal-wallis, uses a normal score instead of ranges as kruskal does.

The function friedman() is used for organoleptic evaluations of different products, made by judges (every judge evaluates all the products). It can also be used for the analysis of treatments of the randomized complete block design, where the response cannot be treated through the analysis of variance.

The function durbin.test() for the analysis of balanced incomplete block designs is very used for sampling tests, where the judges only evaluate a part of the treatments.

Montgomery book data (Montgomery, 2002) Included in the 'agricolae' package

```
library(agricolae)
data(corn)
attach(corn)
str(corn)

'data.frame': 34 obs. of 3 variables:
$ method : int 1 1 1 1 1 1 1 1 2 ...
$ observation: int 83 91 94 89 89 96 91 92 90 91 ...
$ rx : num 11 23 28.5 17 17 31.5 23 26 19.5 23 ...
```

For the examples, the 'agricolae' package data will be used.

#### 4.14 KRUSKAL-WALLIS

```
out <- kruskal (observation, method, group=TRUE, main="corn",
console=TRUE)
Study: corn
Kruskal-Wallis test's
Ties or no Ties
Value: 25.62884
degrees of freedom: 3
Pvalue chisq : 1.140573e-05
method, means of the ranks
 observation r
1
   21.83333 9
2
    15.30000 10
3
   29.57143 7
     4.81250 8
```

```
t-Student: 2.042272
      : 0.05
Alpha
LSD
         : 4.9175
Harmonic Mean of Cell Sizes 8.351284
Means with the same letter are not significantly different
Groups, Treatments and mean of the ranks
         3
                 29.57
b
         1
                 21.83
         2
С
                 15.3
                 4.812
```

The object "compares" has the same structure of the comparisons (figure 4.3).

```
par(cex=0.8,mar=c(3,3,1,0))
bar.group(out$groups,ylim=c(0,35),col=colors()[55],las=1)
```

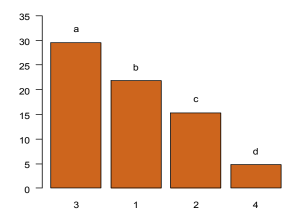


Figure 4.6. Comparison according to Kruskal-Wallis.

## 4.15 FRIEDMAN

```
friedman()
rm(list=ls())
library(agricolae)
data(grass)
attach (grass)
out<-friedman(judge,trt, evaluation,alpha=0.05, group=FALSE,
main="Data of the book of Conover", console=TRUE)
Study: Data of the book of Conover
trt, Sum of the ranks
   evaluation r
t1
         38.0 12
t2
         23.5 12
         24.5 12
t3
         34.0 12
t4
```

```
Friedman's Test
_____
Adjusted for ties
Value: 8.097345
Pvalue chisq : 0.04404214
F value : 3.192198
Pvalue F: 0.03621547
Alpha
       : 0.05
t-Student : 2.034515
Comparison between treatments
Sum of the ranks
       Difference pvalue sig LCL UCL 3.02 25.98
t1 - t2
            13.5 0.022602 * 2.02 24.98
t1 - t3
            4.0 0.483434
t1 - t4
                               -7.48 15.48
t2 - t3
            -1.0 0.860438
                              -12.48 10.48
t2 - t4
            -10.5 0.071736 . -21.98 0.98
                              -20.98 1.98
t3 - t4
            -9.5 0.101742
```

## 4.16 WAERDEN

waerden.test(), with the sweet potato data in the 'agricolae' basis.

```
data(sweetpotato)
attach (sweetpotato)
out <- waerden.test(yield, virus, alpha=0.01, group=TRUE, console=TRUE)
Van der Waerden (Normal Scores) test's
Value: 8.409979
Pvalue: 0.03825667
Degrees of freedom: 3
virus, means of the normal score
        yield std.err r
cc -0.2328353 0.1748697 3
fc -1.0601764 0.2002213 3
ff 0.6885684 0.4396858 3
00 0.6044433 0.2160981 3
t-Student: 3.355387
Alpha
      : 0.01
LSD
        : 1.322487
Means with the same letter are not significantly different
Groups, Treatments and means of the normal score
        ff 0.6886
                0.6044
        00
а
                -0.2328
ab
        CC
        fc
                -1.06
```

The comparison probabilities are obtained with the parameter group=FALSE.

```
> names(out)
[1] "statistics" "parameters" "means" "comparison" "groups"
To see out$comparison
out<-waerden.test(yield, virus, group=F, console=TRUE)</pre>
detach(sweetpotato)
Study:
Van der Waerden (Normal Scores) test's
Value: 8.409979
Pvalue: 0.03825667
Degrees of freedom: 3
virus, means of the normal score
        yield
              std.err r
cc -0.2328353 0.1748697 3
fc -1.0601764 0.2002213 3
ff 0.6885684 0.4396858 3
00 0.6044433 0.2160981 3
Comparison between treatments means
mean of the normal score
       Difference pvalue sig.
                                        LCL
                                                     UCL
cc - fc 0.8273411 0.069032 . -0.08154345 1.73622564
                             * -1.83028827 -0.01251917
cc - ff -0.9214037 0.047582
cc - oo -0.8372786 0.066376
                             . -1.74616316 0.07160593
fc - oo -1.6646197 0.002902 ** -2.57350426 -0.75573516
ff - oo 0.0841251 0.836322
                                -0.82475944 0.99300965
4.17 DURBIN
durbin(); example: Myles Hollander (p. 311) Source: W. Moore and C.I. Bliss. (1942)
days \leftarrow gl(7,3)
chemical<-c("A", "B", "D", "A", "C", "E", "C", "D", "G", "A", "F", "G",
"B", "C", "F", "B", "E", "G", "D", "E", "F")
toxic<-c(0.465,0.343,0.396,0.602,0.873,0.634,0.875,0.325,0.330,
0.423, 0.987, 0.426, 0.652, 1.142, 0.989, 0.536, 0.409, 0.309,
0.609, 0.417, 0.931)
out < - durbin.test(days, chemical, toxic, group=F, console=TRUE,
main="Logarithm of the toxic dose")
Study: Logarithm of the toxic dose
chemical, Sum of ranks
 SIIM
   5
Α
    5
В
```

С

9

```
5
D
Ε
   5
F
   8
G
   5
```

#### Durbin Test \_\_\_\_\_

Value : 7.714286 Df 1 : 6

Df 1 : 6
P-value : 0.2597916
Alpha Alpha : 0.05

Df 2 : 8 t-Student : 2.306004

Least Significant Difference

between the sum of ranks: 5.00689

Parameters BIB Lambda : 1 treatmeans : 7 Block size : 3 Blocks : 7 Replication: 3

Comparison between treatments sum of the ranks

```
Difference pvalue sig
A - B 0 1.000000
A - C
           -4 0.102688
A - D
           0 1.000000
A - E
           0 1.000000
          -3 0.204420
A - F
A - G
           0 1.000000
. . . .
E - F
          -3 0.204420
E - G
           0 1.000000
F - G
           3 0.204420
```

#### 5 **STABILITY ANALYSIS**

In 'agricolae' there are two methods for the study of stability and the AMMI model. These are: a parametric model for a simultaneous selection in yield and stability "SHUKLA'S STABILITY VARIANCE AND KANG'S", and a non-parametric method of Haynes, based on the data range.

## 5.1 PARAMETRIC STABILITY

Use the parametric model, function stability.par().

Prepare a data table where the rows and the columns are the genotypes and the environments, respectively. The data should correspond to yield averages or to another measured variable. Determine the variance of the common error for all the environments and the number of repetitions that was evaluated for every genotype. If the repetitions are different, find a harmonious average that will represent the set. Finally, assign a name to each row that will represent the genotype. We will consider five environments in the following example:

```
\begin{array}{l} v1 <- c(10.2, 8.8, 8.8, 9.3, 9.6, 7.2, 8.4, 9.6, 7.9, 10, 9.3, 8.0, 10.1, 9.4, 10.8, 6.3, 7.4) \\ v2 <- c(7, 7.8, 7.0, 6.9, 7, 8.3, 7.4, 6.5, 6.8, 7.9, 7.3, 6.8, 8.1, 7.1, 7.1, 6.4, 4.1) \\ v3 <- c(5.3, 4.4, 5.3, 4.4, 5.5, 4.6, 6.2, 6.0, 6.5, 5.3, 5.7, 4.4, 4.2, 5.6, 5.8, 3.9, 3.8) \\ v4 <- c(7.8, 5.9, 7.3, 5.9, 7.8, 6.3, 7.9, 7.5, 7.6, 5.4, 5.6, 7.8, 6.5, 8.1, 7.5, 5.0, 5.4) \\ v5 <- c(9, 9.2, 8.8, 10.6, 8.3, 9.3, 9.6, 8.8, 7.9, 9.1, 7.7, 9.5, 9.4, 9.4, 10.3, 8.8, 8.7) \end{array}
```

## For 17 genotypes, the identification is made by letters.

```
study <- data.frame(v1, v2, v3, v4, v5)
rownames(study) <- LETTERS[1:17]</pre>
```

## An error variance of 2 and 4 repetitions is assumed.

```
stability <- stability.par(study, rep=4, MSerror=2)</pre>
```

INTERACTIVE PROGRAM FOR CALCULATING SHUKLA'S STABILITY VARIANCE AND KANG'S

YIELD - STABILITY (YSi) STATISTICS

Environmental index - covariate

Analysis of Variance

	d.f.	Sum o	of Squares	Mean Squares	F	p.value
TOTAL	84		1035.6075			
GENOTYPES	16		120.0875	7.5055	2.65	0.003
ENVIRONMENTS	4		734.2475	183.5619	91.78	<0.001
INTERACTION	64		181.2725	2.8324	1.42	0.033
HETEROGENEITY	16		52.7128	3.2945	1.23	0.281
RESIDUAL	48		128.5597	2.6783	1.34	0.0815
POOLED ERROR	240			2		

Genotype. Stability statistics

```
Mean Sigma-square . s-square . Ecovalence
A 7.86
          1.671833 ns 2.209084 ns
                                     6.567031
           1.822233 ns 1.977299 ns
в 7.22
                                    7.097855
C 7.44
          0.233967 ns 0.134103 ns
                                   1.492208
D 7.42
           4.079567 ns 1.443859 ns 15.064913
E 7.64
           2.037967 ns 2.369090 ns
                                    7.859266
F 7.14
           5.161967 * 6.763106 *
                                   18.885149
G 7.90
           1.759300 ns 1.058092 ns
                                   6.875737
н 7.68
           1.757167 ns 2.028880 ns
                                    6.868208
I 7.34
           5.495300 * 0.423680 ns 20.061619
J 7.54
           4.129967 ns 5.125514 ns 15.242796
K 7.12
           3.848900 ns 4.360772 ns
                                   14.250796
L 7.30
           2.675300 ns 3.610982 ns
                                   10.108678
M 7.66
           3.473167 ns 2.198229 ns 12.924678
N 7.92
                                    3.511972
           0.806233 ns 1.097156 ns
0 8.30
                                    7.553384
           1.951300 ns 1.459578 ns
           3.647833 ns 4.919102 ns 13.541149
P 6.08
Q 5.88
           3.598500 ns 4.353030 ns
                                   13.367031
```

Signif. codes: 0 '\*\*' 0.01 '\*' 0.05 'ns' 1

```
Simultaneous selection for yield and stability (++)
```

```
Yield Rank Adj.rank Adjusted Stab.var Stab.rating YSi ...

A 7.86 14 1 151.671833 0 15 +

B 7.22 5 -1 41.822233 0 4

C 7.44 9 1 100.233967 0 10 +

D 7.42 8 1 94.079567 -2 7

E 7.64 11 1 122.037967 0 12 +

F 7.14 4 -1 35.161967 -4 -1

G 7.90 15 1 161.759300 0 16 +

H 7.68 13 1 141.757167 0 14 +

I 7.34 7 -1 65.495300 -4 2

J 7.54 10 1 11 4.129967 -2 9 +

K 7.12 3 -1 2 3.848900 0 2

L 7.30 6 -1 5 2.675300 0 5

M 7.66 12 1 13 3.473167 0 13 +

N 7.92 16 1 17 0.806233 0 17 +

O 8.30 17 2 19 1.951300 0 19 +

P 6.08 2 -2 0 3.647833 0 0

Q 5.88 1 -3 -2 3.598500 0 -2

Yield Mean: 7.378824
```

```
Yield Mean: 7.378824
YS Mean: 8.352941
LSD (0.05): 0.7384513
------
+ selected genotype
++ Reference: Kang, M. S. 1993. Simultaneous selection for yield and stability: Consequences for growers. Agron. J. 85:754-757.
```

The selected genotypes are: A, C, E, G, H, J, M, N and O. These genotypes have a higher yield and a lower variation. According to the ANOVA, the interaction is significant.

If for example there is an environmental index, it can be added as a covariate. For this case, the altitude of the localities is included.

```
altitude<-c(1200, 1300, 800, 1600, 2400)
stability <- stability.par(study,rep=4,MSerror=2, cova=TRUE,
name.cov= "altitude", file.cov= altitude)</pre>
```

#### 5.2 NON-PARAMETRIC STABILITY

For non-parametric stability, the function in 'agricolae' is stability.nonpar(). The names of the genotypes should be included in the first column, and in the other columns, the response by environments.

```
Ranking...
  v1 v2 v3 v4
A 16.0 8.0 9 14.0 8.0
B 7.5 14.0 5 5.5 10.0
C 7.5 8.0 9 9.0 6.0
D 9.5 6.0 5 5.5 17.0
E 12.5 8.0 11 14.0 3.0
  2.0 17.0 7 7.0 11.0
G 6.0 13.0 16 16.0 15.0
н 12.5 3.0 15 10.5 6.0
I 4.0 4.5 17 12.0 2.0
J 14.0 15.0 9 2.5 9.0 K 9.5 12.0 13 4.0 1.0
L 5.0 4.5 5 14.0 14.0
M 15.0 16.0 3 8.0 12.5
N 11.0 10.5 12 17.0 12.5
0 17.0 10.5 14 10.5 16.0
P 1.0 2.0 2 1.0 6.0
Q 3.0 1.0 1 2.5 4.0
Statistics...
 Mean Rank s1 Z1 s2 Z2
A 7.86 14 5.4 0.02 21.5 0.04
       5 6.2 0.12 25.7 0.02
в 7.22
C 7.44
        9 3.0 2.73 7.5 1.83
D 7.42
        8 7.4 1.20 36.5 1.05
E 7.64
       11 5.6 0.00 21.8 0.03
F 7.14
       4 7.8 1.81 39.2 1.55
G 7.90 15 5.2 0.08 18.7 0.19
H 7.68 13 6.2 0.12 25.3 0.01
        7 8.8 3.87 51.5 5.08
I 7.34
J 7.54
      10 7.2 0.94 34.3 0.71
K 7.12
       3 7.8 1.81 43.0 2.43
L 7.30
        6 7.4 1.20 34.7 0.77
M 7.66 12 7.6 1.49 38.2 1.36
N 7.92
      16 4.2 0.82 14.8 0.57
0 8.30
       17 7.0 0.71 31.7 0.40
P 6.08
       2 6.6 0.35 27.7 0.09
0 5.88 1 7.0 0.71 32.3 0.46
Sum of Z1: 17.97158
Sum of Z2: 16.59462
Test...
The Z-statistics are measures of stability. The test for the
significance
of the sum of Z1 or Z2 are compared to a Chi-Square value of
chi.sum.
individual Z1 or Z2 are compared to a Chi-square value of chi.ind.
                         vs1 vs2 chi.ind chi.sum
             es1 es2
1 7.378824 5.647059 24 2.566667 148.8 8.843605 27.58711
expectation and variance: es1, es2, vs1, vs2
```

#### **5.3** AMMI

The model AMMI uses the biplot constructed through the principal components generated by the interaction environment-genotype. If there is such interaction, the percentage of the two principal components would explain more than the 50% of the total variation; in such case, the biplot would be a good alternative to study the interaction environment-genotype.

The data for AMMI should come from similar experiments conducted in different environments. Homogeneity of variance of the experimental error, produced in the different environments, is required. The analysis is done by combining the experiments.

The data can be organized in columns, thus: environment, genotype, repetition, and variable.

The data can also be the averages of the genotypes in each environment, but it is necessary to consider a harmonious average for the repetitions and a common variance of the error. The data should be organized in columns: environment, genotype, and variable.

When performing AMMI, this generates the BIPLOT graphics; see figure 5.1.

For the application, we consider the data used in the example of parametric stability (study):

```
par(mar=c(4,4,0,0))
rdto <- c(study[,1], study[,2], study[,3], study[,4], study[,5])</pre>
environment \leftarrow gl(5,17)
genotype <- rep(rownames(study),5)</pre>
model<-AMMI(ENV=environment, GEN=genotype, REP=4, Y=rdto, MSE=2,
ylim=c(-2,2), xlim=c(-2,2), number=FALSE)
ANALYSIS AMMI: rdto
Class level information
ENV: 1 2 3 4 5
GEN: A B C D E F G H I J K L M N O P Q
REP: 4
Number of means: 85
Dependent Variable: rdto
Analysis of variance
         Df Sum Sq Mean Sq F value
                                              Pr(>F)
          4 734.2475 183.561882
ENV
REP(ENV) 15
          16 120.0875
                       7.505471 3.752735 3.406054e-06
GEN
ENV:GEN 64 181.2725
                        2.832382 1.416191 3.279630e-02
                       2.000000
Residuals 240 480.0000
Coeff var Mean rdto
19.16584
               7.378824
```

```
Analysis
    percent acum Df
                         Sum.Sq Mean.Sq F.value
                                                       Pr.F
PC1
        38.0 38.0 19 68.96258 3.629609
                                               1.81 0.0225
PC2
        29.8 67.8 17 54.02864 3.178155
                                               1.59 0.0675
PC3
        22.5 90.3 15 40.84756 2.723170
                                               1.36 0.1680
PC4
         9.6 99.9 13 17.43370 1.341054
                                               0.67 0.7915
PC5
         0.0 99.9 11 0.00000 0.000000
                                               0.00 1.0000
require(klaR)
par(mar=c(4,4,0,0))
model <- AMMI (ENV=environment,
                                  GEN=genotype,
                                                    REP=4,
                                                                        MSE=2,
                                                             Y=rdto,
graph="triplot", number=F)
    N
PC2(29.8)
                                                      PC 2 (29.8)
                                                                   PC 3 (22.5)
    0
                               В
                    М
                             Α
    7
    Ŋ
                                             2
        -2
                           O
                 -1
                                                             PC 1 (38)
                       PC 1 (38)
```

Figure 5.1. Biplot and Triplot

In this case, the interaction is significant. The first two components explain 67.8%; then the biplot can provide information about the interaction genotype-environment. With the triplot, 90.3% would be explained.

## **6 SPECIAL FUNCTIONS**

## 6.1 CONSENSUS OF DENDROGRAM

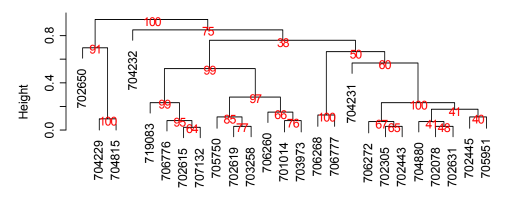
Consensus is the degree or similarity of the vertexes of a tree regarding its branches of the constructed dendrogram. The function to apply is consensus().

The data correspond to a table, with the name of the individuals and the variables in the rows and columns respectively. For the demonstration, we will use the "pamCIP" data of 'agricolae', which correspond to molecular markers of 43 entries of a germplasm bank (rows) and 107 markers (columns).

The program identifies duplicates in the rows and can operate in both cases. The result is a dendrogram, in which the consensus percentage is included, figure 6.1.

```
data(pamCIP)
rownames(pamCIP)<-substr(rownames(pamCIP),1,6)
par(cex=0.8)
output<-consensus(pamCIP,distance="binary", method="complete",
nboot=500)</pre>
```

# **Cluster Dendrogram**



distancia hclust (\*, "complete")

Figure 6.1. Dendrogram, production by consensus()

Duplicates: 18

New data : 25 Records

Consensus hclust

Method distance: binary
Method cluster : complete
rows and cols : 25 107
n-bootstrap : 500

Run time : 20.469 secs

When the dendrogram is complex, it is convenient to extract part of it with the function hcut(), figure 6.2.

hcut(output,h=0.4,group=8,type="t",edgePar = list(lty=1:2, col=colors()[c(42,84)]),main="group 8" ,col.text="blue",cex.text=1,las=1)

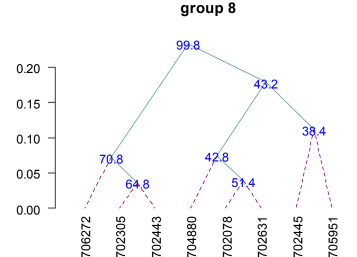


Figure 6.2. Dendrogram, production by hcut()

```
The obtained object "output" contains information about the process:
names (output)
[1] "table.dend" "dendrogram" "duplicates"
```

This means that we can know the duplicates, reconstruct the tree diagram and maintain the interactions.

#### output\$table.dend

```
xaxis
                         height percentage groups
           7.500000 0.02857143
    -6 -24
                                64.0
                                           6-24
       -4 19.500000 0.03571429
                                64.2
                                           3 - 4
   21 23 5.099609 0.93617021 100.0
                                           1-2-3-4-5-6-7-8-9-10-11-
12-13-14-15-16-17-18-19-20-21-22-23-24-25
```

Reproduce the dendrogram:

```
dend<-output$dendrogram
data<-output$table.dend
plot(dend)
text(data[,3],data[,4],data[,5])
Construct a classic dendrogram, figure 6.3
par(mar=c(4,3,1,1))
dend<-as.dendrogram(output$dendrogram)</pre>
plot(dend,type="r",edgePar = list(lty=1:2, col=colors()[c(42,84)])
text(data[,3],data[,4],data[,5],col="blue",cex=1)
```

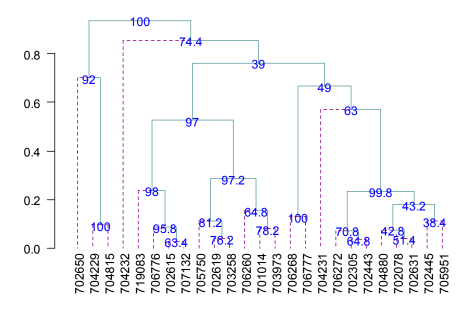


Figure 6.3. Classic dendrogram

## 6.2 MONTECARLO

It is a method for generating random numbers of an unknown distribution. It uses a data set and, through the cumulative behavior of its relative frequency, generates the possible random values that follow the data distribution. These new numbers are used in some simulation process.

The probability density of the original and simulated data can be compared, figure 6.4.

```
data(soil)
set.seed(9473)
simulated <- montecarlo(soil$pH,1000)
par(mar=c(3,0,2,1))
plot(density(soil$pH),axes=F,main="pH density of the soil\ncon
Ralstonia",xlab="",lwd=4)
lines(density(simulated), col="blue", lty=4,lwd=4)
h<-graph.freq(simulated,plot=F)
axis(1,0:12)
legend("topright",c("Original","Simulated"),lty=c(1,4),col=c("black", "blue"), lwd=4)</pre>
```

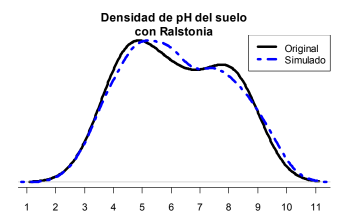


Figure 6.4. Distribution of the simulated and the original data 1000 data have been generated, being the frequency table:

```
round(table.freq(h),2)
  Lower Upper Main freq relative
                                  CF RCF
  2.00 2.79 2.40
                                  12 0.01
                   12
                            0.01
  2.79 3.58 3.19
                     50
                            0.05
                                   62 0.06
  9.11 9.90 9.51
                     49
                            0.05 989 0.99
   9.90 10.69 10.30
                     11
                            0.01 1000 1.00
Some statistics:
summary(soil$pH)
  Min. 1st Qu. Median
                         Mean 3rd Qu.
                                         Max.
        4.700
                 6.100
                         6.154 7.600
                                         8.400
summary(simulated)
  Min. 1st Qu. Median
                         Mean 3rd Qu.
                                         Max.
        4.698
                6.022
                         6.209
                               7.762 10.950
```

### 6.3 RE-SAMPLING IN LINEAR MODEL

It uses the permutation method for the calculation of the probabilities of the sources of variation of ANOVA according to the linear regression model or the design used. The principle is that the Y response does not depend on the averages proposed in the model; hence, the Y values can be permutated and many model estimates can be constructed. On the basis of the patterns of the random variables of the elements under study, the probability is calculated in order to measure the significance.

For a variance analysis, the data should be prepared similarly. The function to use is: resampling.model()

```
data(potato)
potato[,1]<-as.factor(potato[,1])
potato[,2]<-as.factor(potato[,2])
model<-"cutting~variety + date + variety:date"
analysis<-resampling.model(model, potato, k=1000)</pre>
```

The function resampling.model() can be used when the errors have a different distribution from normal.

#### 6.4 SIMULATION IN LINEAR MODEL

Under the assumption of normality, the function generates pseudo experimental errors under the proposed model, and determines the proportion of valid results according to the analysis of variance found.

The function is: simulation.model(). The data are prepared in a table, similarly to an analysis of variance.

Considering the example proposed in the previous procedure:

```
model <- simulation.model(model, potato, k=1000)</pre>
Simulation of experiments
Under the normality assumption
Proposed model: cutting~variety + date + variety:date
Analysis of Variance Table
Response: cutting
              Df Sum Sq Mean Sq F value Pr(>F)
variety
               1 25.087 25.087 7.2580 0.01952 *
date 2 13.892 6.946 2.0096 0.17671
variety:date 2 4.853 2.427 0.7020 0.51484
Residuals 12 41.477 3.456
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 '' 1
Validation of the analysis of variance for the proposed model
Simulations: 1000
                Df F value % Acceptance % Rejection Criterion

      variety
      1 7.2580377
      51.5
      48.5 acceptable

      date
      2 2.0095604
      61.1
      38.9 acceptable

      variety:date
      2 0.7020312
      67.5
      32.5 acceptable
```

The validation is referred to the percentage of decision results equal to the result of the ANOVA decision. Thus, 67.5% of the results simulated on the interaction variety\*date gave the same result of acceptance or rejection obtained in the ANOVA.

#### 6.5 PATH ANALYSIS

It corresponds to the "path analysis" method. The data correspond to correlation matrices of the independent ones with the dependent matrix (XY) and between the independent ones (XX).

It is necessary to assign names to the rows and columns in order to identify the direct and indirect effects.

```
corr.x < -matrix(c(1, 0.5, 0.5, 1), c(2, 2))
corr.y < - rbind(0.6, 0.7)
names<-c("X1","X2")
dimnames(corr.x) <-list(names, names)</pre>
dimnames(corr.y)<-list(names, "Y")</pre>
output<-path.analysis(corr.x,corr.y)</pre>
Direct (Diagonal) and indirect effect path coefficients
______
        X1 X2
X1 0.3333333 0.2666667
X2 0.1666667 0.5333333
Residual Effect^2 = 0.4266667
> output
$Coeff
                 Х2
         X1
X1 0.3333333 0.2666667
X2 0.1666667 0.5333333
$Residual
[1] 0.4266667
```

#### 6.6 LINE X TESTER

It corresponds to a crossbreeding analysis of a genetic design. The data should be organized in a table. Only four columns are required: repetition, females, males, and response. In case it corresponds to progenitors, the females or males field will only be filled with the corresponding one. See the heterosis data.

Example with the heterosis data, locality 2.

```
where <NA> is empty.
If it is a progeny, it comes from a "Female" and a "Male."
If it is a progenitor, it will only be "Female" or "Male."
The following example corresponds to data of the locality 2:
24 progenies
8 females
3 males
3 repetitions
They are 35 treatments (24, 8, 3) applied to three blocks.
data(heterosis)
site2<-subset(heterosis,heterosis[,1]==2)</pre>
site2<-subset(site2[,c(2,5,6,8)],site2[,4]!="Control")
attach(site2)
output1<-lineXtester(Replication, Female, Male, v2)</pre>
detach(site2)
ANALYSIS LINE x TESTER: v2
ANOVA with parents and crosses
_____
                   Df Sum Sq Mean Sq F value Pr(>F)
                   2 0.519190476 0.259595238 9.801 0.0002
Replications
Treatments
                   34 16.101605714 0.473576639 17.879 0.0000
Parents
                  10 7.731490909 0.773149091 29.189 0.0000
Parents vs. Crosses 1 0.005082861 0.005082861 0.192 0.6626
                   23 8.365031944 0.363697041 13.731 0.0000
Crosses
Error
                   68 1.801142857 0.026487395
Total
                  104 18.421939048
ANOVA for line X tester analysis
_____
                   Sum Sq Mean Sq F value Pr(>F)
               7 4.9755431 0.71079187 3.632 0.0191
Lines
Testers
               2 0.6493861 0.32469306 1.659 0.2256
Lines X Testers 14 2.7401028 0.19572163 7.389 0.0000
              68 1.8011429 0.02648739
Error
ANOVA for line X tester analysis including parents
______
                   Df Sum Sq Mean Sq F value Pr(>F)
                   2 0.519190476 0.259595238 9.801 0.0002
Replications
                   34 16.101605714 0.473576639 17.879 0.0000
Treatments
                   10 7.731490909 0.773149091 29.189 0.0000
Parents
                   1 0.005082861 0.005082861
                                              0.192 0.6626
Parents vs. Crosses
                   23 8.365031944 0.363697041 13.731 0.0000
Crosses
                    7 4.975543056 0.710791865
                                              3.632 0.0191
Lines
                   2 0.649386111 0.324693056 1.659 0.2256
Testers
                  14 2.740102778 0.195721627
                                              7.389 0.0000
Lines X Testers 14 2.740102778 0.195721627
Error 68 1.801142857 0.026487395
```

104 18.421939048

Total

```
GCA Effects:
_____
Lines Effects:
Achirana LT-8
               MF-I MF-II Serrana
                                       TPS-2 TPS-25
                                                         TPS-7
                      -0.449
                              0.058 -0.047
0.022 -0.338 0.199
                                                0.414
                                                         0.141
Testers Effects:
TPS-13 TPS-67 TS-15
0.087 0.046 -0.132
SCA Effects:
_____
        Testers
        TPS-13 TPS-67 TS-15
Lines
 Achirana 0.061 0.059 -0.120 LT-8 -0.435 0.519 -0.083
 MF-I
         -0.122 -0.065 0.187
 MF-II -0.194 0.047 0.148
 Serrana 0.032 -0.113 0.081
         0.197 -0.072 -0.124
 TPS-2
 TPS-25
          0.126 -0.200 0.074
 TPS-7
         0.336 -0.173 -0.162
Standard Errors for Combining Ability Effects:
_____
S.E. (gca for line) : 0.05424983
S.E. (gca for tester) : 0.0332211
S.E. (sca effect) : 0.09396346
S.E. (gi - gj)line : 0.07672084
S.E. (gi - gj) tester : 0.04698173
S.E. (sij - skl)tester: 0.1328844
Genetic Components:
============
Cov H.S. (line) : 0.05723003
Cov H.S. (tester): 0.00537381
Cov H.S. (average): 0.003867302
Cov F.S. (average): 0.1279716
F = 0, Adittive genetic variance: 0.01546921
F = 1, Adittive genetic variance: 0.007734604
F = 0, Variance due to Dominance: 0.1128228
F = 1, Variance due to Dominance: 0.05641141
Proportional contribution of lines, testers
and their interactions to total variance
_____
Contributions of lines : 59.48026
Contributions of testers: 7.763104
Contributions of lxt : 32.75663
```

## 6.7 SOIL UNIFORMITY

The Smith index is an indicator of the uniformity, used to determine the parcel size for research purposes. The data correspond to a matrix or table that contains the response per basic unit, a number of n rows x m columns, and a total of n\*m basic units.

For the test, we will use the rice file. The graphic is a result with the adjustment of a model for the parcel size and the coefficient of variation, figure 6.5.

```
data(rice)
table<-index.smith(rice,
main="Interaction between the CV and the parcel size" ,col="red",
type="l",xlab="Size")
uniformity <- data.frame(table$uniformity)</pre>
uniformity
      Size Width Length plots
                                    Vx
                         648 9044.539 13.0
                  1
            1
2
                   2
                         324 7816.068 12.1
            1
                          324 7831.232 12.1
3
      2
                   1
40 162
            Q
                  18
                            4 4009.765 8.6
```

The size is the product of the width x the length of the parcel, and the rectangle size is the product of the width x the length.

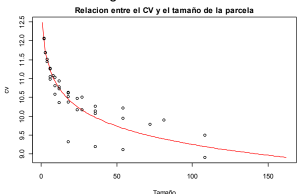


Figure 6.5. Adjustment curve for the optimal size of parcel

### 6.8 CONFIDENCE LIMITS IN BIODIVERSITY INDICES

The biodiversity indices are widely used for measuring the presence of living things in an ecological area. Many programs indicate their value. The function of 'agricolae' is also to show the confidence intervals, which can be used for a statistical comparison. Use the bootstrap procedure. The data are organized in a table; the species are placed in a column; and in another one, the number of individuals. The indices that can be calculated with the function index.bio() of 'agricolae' are: "Margalef", "Simpson.Dom", "Simpson.Div", "Berger.Parker", "McIntosh", and "Shannon."

In the example below, we will use the data obtained in the locality of Paracsho, district of Huasahuasi, province of Tarma in the department of Junín.

The evaluation was carried out in the parcels on 17 November 2005, without insecticide application. The counted specimens were the following:

```
data(paracsho)
species <- paracsho[79:87,4:6]
species</pre>
```

```
Family Number.of.specimens
        Order
      DIPTERA TIPULIDAE
79
               NOCTUIDAE
80 LEPIDOPTERA
                                           1
   NOCTUIDAE PYRALIDAE
81
                                           3
    HEMIPTERA ANTHOCORIDAE
82
                                           1
83
    DIPTERA
              TACHINIDAE
                                          16
      DIPTERA ANTHOCORIDAE
84
                                           3
85
      DIPTERA SCATOPHAGIDAE
                                           5
86
      DIPTERA
                SYRPHIDAE
                                           1
87
      DIPTERA
                 MUSCIDAE
                                           3
```

#### The Shannon index is:

```
output <- index.bio(species[,3],method="Shannon",level=95,nboot=200)
Method: Shannon
Index: 3.52304
95 percent confidence interval:
   3.088775; 4.286088</pre>
```

## 6.9 CORRELATION

The function correlation() of 'agricolae' makes the correlations through the methods of Pearson, Spearman and Kendall for vectors and/or matrices. If they are two vectors, the test is carried out for one or two lines; if it is a matrix one, it determines the probabilities for a difference, whether it is greater or smaller.

For its application, consider the soil data: data(soil)

```
data(soil)
correlation(soil[,2:4],method="pearson")
Correlation Analysis
Method
          : pearson
Alternative: two.sided
$correlation
      pH EC CaCO3
     1.00 0.55 0.73
      0.55 1.00 0.32
CaCO3 0.73 0.32 1.00
$pvalue
               рН
                        EC
                                 CaCO3
     1.000000000 0.0525330 0.004797027
      0.052532997 1.0000000 0.294159813
CaCO3 0.004797027 0.2941598 1.000000000
$n.obs
[1] 13
attach(soil)
correlation(pH, soil[, 3:4], method="pearson")
```

```
Correlation Analysis
Method
         : pearson
Alternative: two.sided
$correlation
    EC CaCO3
pH 0.55 0.73
$pvalue
      EC CaCO3
рН 0.0525 0.0048
$n.obs
[1] 13
correlation(pH, CaCO3, method="pearson")
Pearson's product-moment correlation
data: pH and CaCO3
t = 3.520169 , df = 11 , p-value = 0.004797027
alternative hypothesis: true rho is not equal to 0
sample estimates:
cor
0.7278362
```

### **6.10 OTHER FUNCTIONS**

Desirability functions that facilitate the data management:

tapply.stat() Calculation of statesmen and mathematical operations in columns of a table in relation to grouped factors.

Factor and variable table

Application with 'agricolae' data:

```
data(RioChillon)
attach(RioChillon$babies)
tapply.stat(yield, farmer, function(x) max(x) - min(x))
detach(RioChillon$babies)
           farmer yield
1 AugustoZambrano 7.5
       Caballero 13.4
3
      ChocasAlto 14.1
      FelixAndia 19.4
5
     Huarangal-1 9.8
6
     Huarangal-2 9.1
7
      Huarangal-3 9.4
         Huatocay 19.4
9 IgnacioPolinario 13.1
```

It corresponds to the range of variation in the farmers' yield.

The function "tapply" can be used directly or with function.

If A is a table with columns 1,2 and 3 as category, and 5,6 and 7 as variables, then the following procedures are valid:

```
tapply.stat(A[,5:7], A[,1:3], mean)
tapply.stat(A[,5:7], A[,1:3], function(x) mean(x,na.rm=TRUE))
tapply.stat(A[,c(7,6)], A[,1:2], function(x) sd(x)*100/mean(x))
```

## Coefficient of variation of an experiment

If "model" is the object resulting from an analysis of variance of the function aov() or Im() of R, then the function cv.model() calculates the coefficient of variation.

```
data(sweetpotato)
model <- model<-aov(yield ~ virus, data=sweetpotato)
cv.model(model)
[1] 17.16660</pre>
```

#### Skewness and curtosis

The skewness and curtosis results, obtained by 'agricolae', are equal to the ones obtained by SAS, MiniTab, SPSS, InfoStat, and Excel.

```
If x represents a data set:
> x<-c(3,4,5,2,3,4,5,6,4,NA,7)

skewness is calculated with:
> skewness(x)
[1] 0.3595431

and curtosis with:
> kurtosis(x)
```

#### **Tabular value of Waller-Duncan**

[1]-0.1517996

The function Waller determines the tabular value of Waller-Duncan. For the calculation, value F is necessary, calculated from the analysis of variance of the study factor, with its freedom degrees and the estimate of the variance of the experimental error. Value K, parameter of the function is the ratio between the two types of errors (I and II). To use it, a value associated with the alpha level is assigned. When the alpha level is 0.10, 50 is assigned to K; for 0.05, K=100; and for 0.01, K=500. K can take any value.

Figure 6.6 illustrates the function for different values of K with freedom degrees of 5 for the numerator and 15 for the denominator, and values of calculated F, equal to 2, 4, and 8.

```
q<-5
f<-15
K<-seq(10,1000,100)
n<-length(K)
y<-rep(0,3*n)
dim(y)<-c(n,3)
for(i in 1:n) y[i,1]<-waller(K[i],q,f,Fc=2)
for(i in 1:n) y[i,2]<-waller(K[i],q,f,Fc=4)</pre>
```

```
for(i in 1:n) y[i,3]<-waller(K[i],q,f,Fc=8)
plot(K,y[,1],type="l",col="blue",ylab="waller")
lines(K,y[,2],type="l",col="red",lty=2,lwd=2)
lines(K,y[,3],type="l",col="green",lty=4,lwd=2)
legend("topleft",c("2","4","8"),col=c("blue","red","green"),lty=c(1,8,20),lwd=2,title="Fc")
title(main="Waller in function of K")</pre>
```

#### Waller en función de K

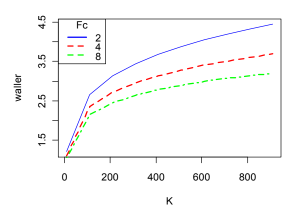


Figure 6.6. Function of Waller to different value of parameters K and Fc

### **AUDPC**

The area under the disease progress curve (AUDPC), (see figure 6.7), calculates the absolute and relative progress of the disease. It is required to measure the disease in percentage terms during several dates, preferably equidistantly.

```
days<-c(7,14,21,28,35,42)
evaluation<-data.frame(E1=10,E2=40,E3=50,E4=70,E5=80,E6=90)
par(mar=c(4,4,1,1))
plot(days, evaluation,type="h",ylim=c(0,100),axes=F,col=
colors()[42],xlab="Days", ylab="Evaluation")
lines(days,evaluation,col= colors()[42])
axis(1,days)
axis(2,seq(0,100,20),las=2)
rect(7,0,42,100)
audpc(evaluation,days)
audpc(evaluation,days,"relative")
text(15,80,"Audpc Absolute = 2030")
text(15,70,"Audpc Relative = 0.58")</pre>
```

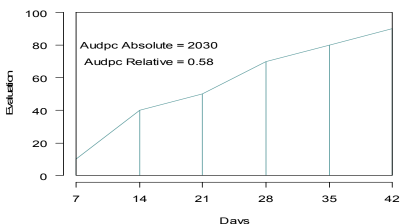


Figure 6.7. AUDPC: Area under the curve

#### **NON-ADDITIVITY**

Tukey's test for non-additivity is used when there are doubts about the additivity veracity of a model. This test confirms such assumption and it is expected to accept the null hypothesis of the non-additive effect of the model.

For this test, all the experimental data used in the estimation of the linear additive model are required.

Use the function nonadditivity() of 'agricolae'. For its demonstration, the experimental data "potato", of the package 'agricolae', will be used. In this case, the model corresponds to the randomized complete block design, where the treatments are the varieties.

```
data(potato)
potato[,1]<-as.factor(potato[,1])</pre>
model<-lm(cutting ~ date + variety,potato)</pre>
df<-df.residual(model)</pre>
MSerror<-deviance(model)/df
attach (potato)
analysis <- nonadditivity (cutting, date, variety, df, MSerror)
detach (potato)
Tukey's test of non-additivity
cutting
P: 15.37166
Q: 77.4444
Analysis of Variance Table
Response: residual
               Df Sum Sq Mean Sq F value Pr(>F)
Non-additivity 1 3.051
                            3.051
                                    0.922 0.3532
Residuals
               14 46.330
                            3.309
```

According to the results, the model is additive because the p.value 0.35 is greater than 0.05.

#### **LATEBLIGHT**

LATEBLIGHT is a mathematical model that simulates the effect of weather, host growth and resistance, and fungicide use on asexual development and growth of Phytophthora infestans on potato foliage.

LATEBLIGHT Version LB2004 was created in October 2004 (Andrade-Piedra et al., 2005a, b and c), based on the C-version written by B.E. Ticknor ('BET 21191 modification of cbm8d29.c'), reported by Doster et al. (1990) and described in detail by Fry et al. (1991) (This version is referred as LB1990 by Andrade-Piedra et al. [2005a]). The first version of LATEBLIGHT was developed by Bruhn and Fry (1981) and described in detail by Bruhn et al. (1980).

```
library(agricolae)
f <- system.file("external/weather.csv", package="agricolae")</pre>
weather <- read.csv(f,header=FALSE)</pre>
f <- system.file("external/severity.csv", package="agricolae")</pre>
severity <- read.csv(f)</pre>
weather[,1]<-as.Date(weather[,1],format = \%m/%d/%Y")
# Parameters dates
dates<-c("2000-03-25","2000-04-09","2000-04-12","2000-04-16","2000-
04 - 22")
dates<-as.Date(dates)</pre>
EmergDate <- as.Date('2000/01/19')</pre>
EndEpidDate <- as.Date("2000-04-22")</pre>
dates<-as.Date(dates)</pre>
NoReadingsH<- 1
RHthreshold <- 90
WS<-weatherSeverity (weather, severity, dates, EmergDate, EndEpidDate,
NoReadingsH, RHthreshold)
# Parameters Lateblight
InocDate<-"2000-03-18"
LGR <- 0.00410
IniSpor <- 0</pre>
SR <- 292000000
IE <- 1.0
LP <- 2.82
InMicCol <- 9
Cultivar <- 'NICOLA'
ApplSys <- "NOFUNGICIDE"
main<-"Cultivar: NICOLA"</pre>
model<-lateblight(WS, Cultivar, ApplSys, InocDate, LGR, IniSpor, SR, IE,</pre>
LP, MatTime='LATESEASON', InMicCol, main=main, type="1", xlim=c(65,95), lw
d=1.5, xlab="Time (days after emergence)", ylab="Severity
(Percentage)")
> head(model$Gfile)
           dates nday MeanSeverity StDevSeverity
                                                     MinObs
                                                                 MaxObs
                               0.1 0.000000 0.100000
Eval1 2000-03-25 66
                                                               0.10000
Eval2 2000-04-09 81
                                         24.722459 -3.922459 45.52246
                               20.8
Eval3 2000-04-12 84
                                        32.710854 24.289146 89.71085
                               57.0
Eval4 2000-04-16 88
                              94.0
                                        7.968689 86.031311 101.96869
4.000000 93.000000 101.00000
                          97.0
Eval5 2000-04-22 94
```

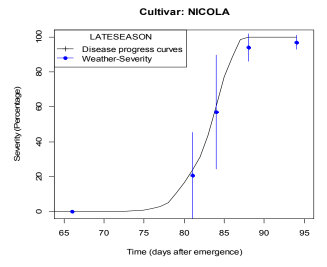


Figure 6.8. lateblight: LATESEASON

```
# reproduce graph
x<- model$Ofile$nday
y<- model $Ofile $SimSeverity
w<- model$Gfile$nday
z<- model$Gfile$MeanSeverity</pre>
Min<-model$Gfile$MinObs
Max<-model$Gfile$MaxObs
plot(x,y,type="1",xlim=c(65,95),lwd=1.5,xlab="Time (days after
emergence)",
ylab="Severity (Percentage)")
points(w,z,col="red",cex=1,pch=19)
npoints <- length(w)</pre>
for ( i in 1:npoints) {
segments(w[i],Min[i],w[i],Max[i],lwd=1.5,col="red")
legend("topleft",c("Disease progress curves","Weather-Severity"),
title="Description",lty=1,pch=c(3,19),col=c("black","red"))
```

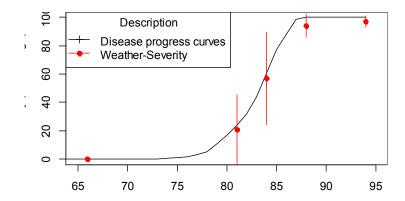


Figure 6.9. lateblight: LATESEASON

Table 6.10. ASCII Character Code Reference for the use of symbols

	ASCII Codes Table used in R								
Code	Symbol	Code	Symbol	Code	Symbol				
92	}	124	ı	64	@				
47	1	60	<	94	۸				
91	[	62	>	35	#				
93	]	61	=	36	\$				
40	(	34	"	37	%				
41	)	126	~	38	&				
123	{	58	:	39	4				
125	}	59	;						

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