



IOWA State University. August, 2007

## **Agricolae – a free statistical library for agricultural research**

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### **Description**

**Agricolae** Version 1.0-3 contains a total 68 statistical routines and 33 data sets.

Agricolae is a statistical library for agricultural research with the goal of supporting developing countries. Focuses on statistical tools used in the breeding program of the International Potato Center for its main commodity crops, potato and sweetpotato. Thus, Agricolae supports a variety of field trial designs, including incomplete block designs techniques, genetic designs, stability analysis, AMMI with biplot and triplot analysis, multiple comparisons of treatments. Other functions include the construction of consensus clusters, optimal size and shape of experimental field plots.

Agricolae was developed using R and is available via the CRAN, repository at <http://www.r-project.org>.

## Planning of field experiment

### Randomize and field book

#### Alpha design, Graeco, latin square, CRD, RCBD , BIB.

The planning of field experiments is one of the main areas of Agricolae.

It supports simple lattice design (`lattice.simple`),  
Factorial a block design (`design.ab`),  
Alpha design (`alpha.design`),  
Balanced Incomplete Block Design (`design.bib`),  
Randomized complete block design (`design.rcbd`),  
Complete randomized design (`design.crd`),  
Graeco-latin square design (`design.graeco`),  
Latin square design (`design.lsd`).

## Planning of field experiment

### Greaco latin

```
args: trt1, trt2, number = 1, seed = 0, kinds = "Super-Duper"
```

```
> T1<-c("a","b","c","d")  
> T2<-c("v","w","x","y")  
> Plan <- design.graeco(T1,T2,number=101)
```

Plots					Treatments			
	[,1]	[,2]	[,3]	[,4]	[,1]	[,2]	[,3]	[,4]
[1,]	101	102	103	104	"d w"	"b v"	"a x"	"c y"
[2,]	105	106	107	108	"b y"	"d x"	"c v"	"a w"
[3,]	109	110	111	112	"a v"	"c w"	"d y"	"b x"
[4,]	113	114	115	116	"c x"	"a y"	"b w"	"d v"

It's not possible to construct: 6,10 and pair >= 14

# Planning of field experiment

## Alpha design

(trt, k, r, number = 1, seed = 0, kinds = "Super-Duper")

```
> Trt <- letters[1:12]
> plan<-design.alpha(trt,k=3, r=2, number=101)
```

```
alpha design (0,1) - Serie I
Parameters Alpha design
=====
treatmeans : 12
Block size : 3
Blocks      : 4
Replication: 2

Efficiency factor
(E ) 0.6470588
<<< Book >>>
```

Field Book					
	plots	cols	block	trt	replication
> plan					
1	101	1	1	j	1
2	102	2	1	h	1
3	103	3	1	c	1
4	104	1	2	d	1
...					
23	123	2	8	h	2
24	124	3	8	e	2

## Comparison of multiple treatments

### Test:

**LSD, HSD, Waller, Durbin, Kruskal Wallis, Friedman, Waerden**

#### Test parametrics:

LSD: Least significant difference and Adjust P-values

HSD: Honestly significant difference Tukey.

Waller: Bayesian t-values for multiple comparisons

#### Test Non parametrics

Kruskal Wallis: Complete randomized design

Friedman: Randomized complete block design

Durbin: Balanced Incomplete Block Design

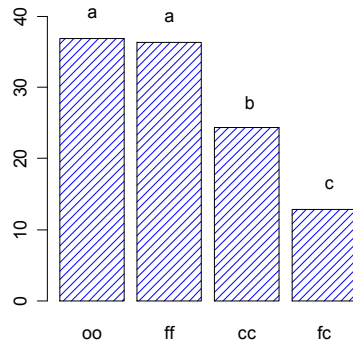
Waerden: The van der Waerden (Normal Scores)

## Comparison of multiple treatments

### Waller-Duncan

(y, trt, DFerror, MSerror, Fc, K = 100, group = TRUE, main = NULL)

```
> attach(sweetpotato)
> model<-aov(yield-virus)
> comparison <- waller.test(yield, virus, DFerror=8, MSerror=22.49, Fc=17.345)
> bar.group(comparison, horiz=FALSE, ylim=c(0,45), density=10, col="blue")
```



Critical Value of Waller 2.236  
 Minimum Significant Difference  
 8.658066  
 Means with the same letter are not significantly different.

#### Groups, Treatments and means

a	oo	36.9
a	ff	36.33333
b	cc	24.4
c	fc	12.86667

## Comparison of multiple treatments

### LSD

(y, trt, DFerror, MSerror, alpha = 0.05, p.adj = c("none", "holm", "hochberg", "bonferroni", "BH", "BY", "fdr"), group = TRUE, main = NULL)

```
> comparison <- LSD.test(yield, virus, DFerror=8, MSerror=22.49,
p.adj="bonferroni")
```

```
group = TRUE

LSD t Test for yield
P value adjustment method: bonferroni
Alpha 0.050000
Error Degrees of Freedom 8.000000
Error Mean Square 22.490000
Critical Value of t 3.478879
```

Least Significant Difference 13.47065  
 Means with the same letter are not significantly different.

#### Groups, Treatments and means

a	oo	36.9
a	ff	36.33333
ab	cc	24.4
b	fc	12.86667

```
group = FALSE

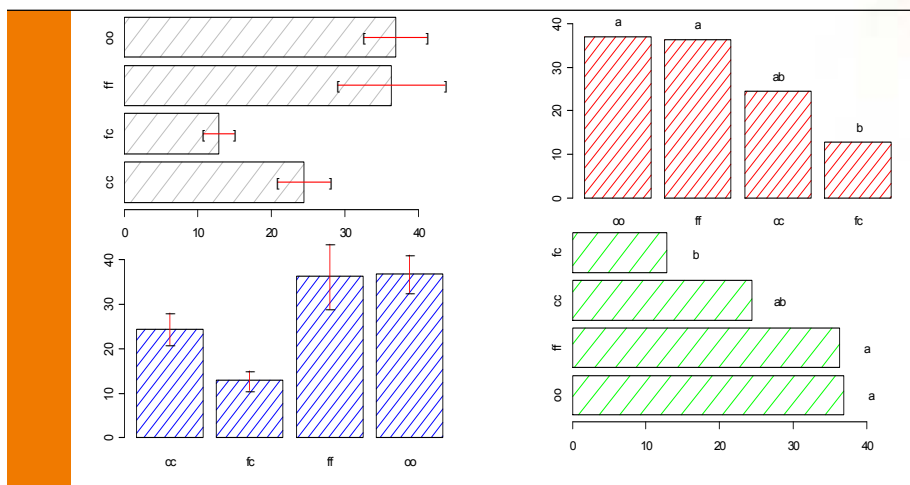
Treatment Means
virus yield std.err replication
1 cc 24.40000 2.084067 3
2 fc 12.86667 1.246774 3
3 ff 36.33333 4.233727 3
4 oo 36.90000 2.482606 3
```

#### Comparison between treatments means

tr.i	tr.j	diff	pvalue
1	2	11.5333333	0.1056
2	1	11.9333333	0.0900
3	1	4.12500000	0.0720
4	2	23.4666667	0.0024
5	2	4.03333333	0.0012
6	3	4.05666667	1.0000

## Comparison of multiple treatments Graphics.

### bar.err & bar.group



## Stability analysis

### AMMI, stability.par, stability.nonpar

**AMMI:** Additive Main Effects and Multiplicative Interaction models are widely used to analyze main effects and genotype by environment (GEN, ENV) interactions in multilocation variety trials. Furthermore, this function generates biplot and triplot graphs as well as principal component analysis.

**stability.par:** SHUKLA'S STABILITY VARIANCE AND KANG'S. This procedure calculates the stability variations as well as the statistics of selection for the yield and the stability

**stability.nonpar:** A method based on the statistical ranges of the study variable per environment for the stability analysis

## Stability analysis

### AMMI

```
(ENV, GEN, REP, Y, MSE=0, number=TRUE, graph="biplot",...)
```

```
> model<- AMMI(ltrv[,2], ltrv[,1], ltrv[,3], ltrv[,5],  
xlim=c(-3,3),ylim=c(-4,4), graph="biplot",number=FALSE)
```

```
ANALYSIS AMMI: ltrv[, 5]  
Class level information
```

```
ENV: Ayac LM-02 SR-02 Hyo-02 LM-03 SR-03  
GEN: 102.18 104.22 121.31 141.28 157.26 163.9 221.19 233.11 235.6  
241.2 255.7 314.12 317.6 319.20 320.16 342.15 346.2 351.26 364.21  
402.7 405.2 406.12 427.7 450.3 506.2 Canchan Desiree Unica  
REP: 1 2 3
```

```
Number of observations: 504
```

```
model Y: ltrv[, 5] ~ ENV + REP%in%ENV + GEN + ENV:GEN  
Random effect REP%in%ENV
```

## Stability analysis

### AMMI

```
(ENV, GEN, REP, Y, MSE=0, number=TRUE, graph="biplot",...)
```

```
Analysis of Variance Table
```

```
Response: Y
```

	Df	Sum Sq	Mean Sq	F value	Pr(>F)
ENV	5	9607.4	1921.5	284.6352	4.957e-12 ***
REP(ENV)	12	81.0	6.8	2.7313	0.00154 **
GEN	27	1367.4	50.6	20.4904	< 2.2e-16 ***
ENV:GEN	135	1764.8	13.1	5.2891	< 2.2e-16 ***
Residuals	324	800.8	2.5		

```
---
```

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

```
Coeff var      Mean ltrv[, 5]  
20.07525      7.831188
```

```
Analysis
```

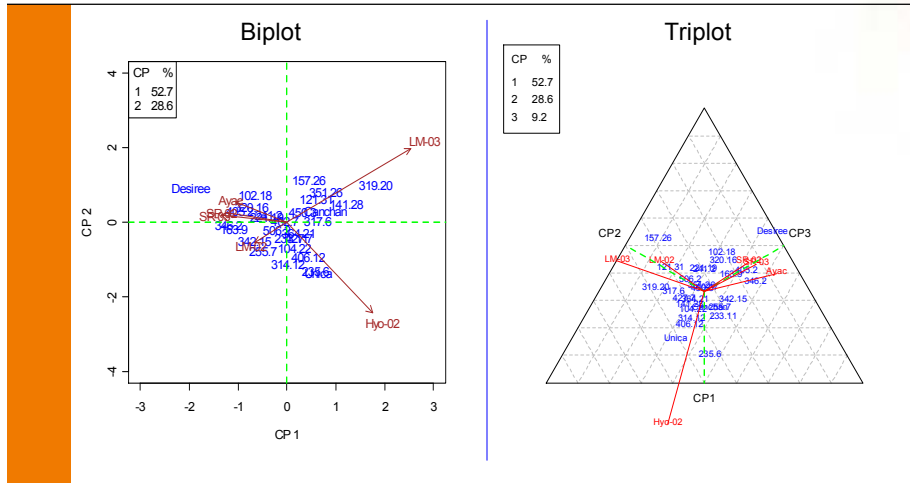
	percent	acum	Df	Sum.Sq	Mean.Sq	F.value	Pr.F
CP1	52.7	52.7	31	929.89935	29.996753	12.14	0.0000
CP2	28.6	81.3	29	503.95903	17.377898	7.03	0.0000

```
... More ...
```

## Stability analysis

### AMMI

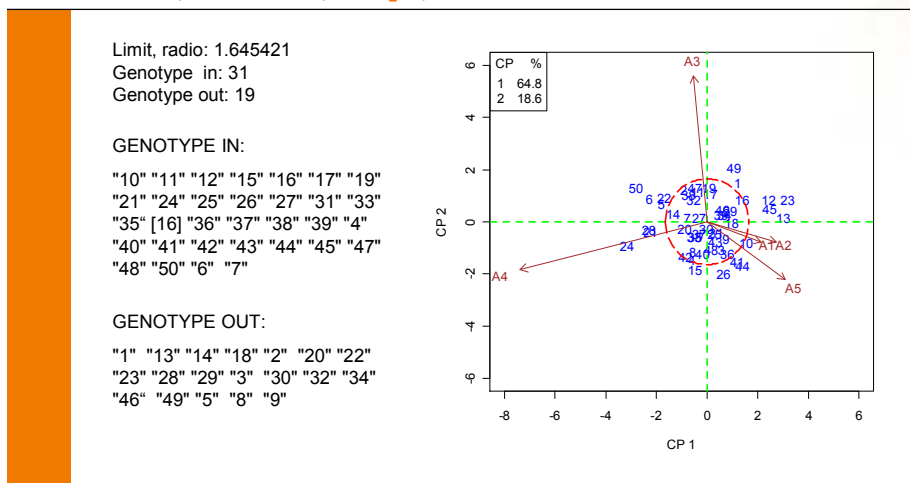
(ENV, GEN, REP, Y, MSE=0, number=TRUE, graph="biplot", ..)



## Stability analysis

### AMMI.contour

(model, distance, shape, ...)



## Stability analysis

### AMMI

```
(ENV,GEN,REP, Y, MSE=0,number=TRUE,graph="biplot",..)
```

Input data:

a) complete or missing value. Experiments in localities under randomized complete block design.

Or

b) Only means and missing value. Estimation variance of error and replication:

MSE = variance error = Mean square error

Rep = constant = Harmonic Mean ( $r_1, r_2, \dots, r_k$ )

## Stability analysis (parametric)

### stability.par

```
(data, rep, MSerror, alpha = 0.1, main = NULL, cova = F,  
name.cov = NULL, file.cov = 0)
```

```
> stability.par(data, rep=4, MSerror=1.8, alpha=0.1, main="Genotype")
```

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

```
YIELD - STABILITY (Ysi) STATISTICS
```

```
Genotype
```

```
Environmental index - covariate
```

```
Analysis of Variance
```

Source	d.f.	Sum of Squares	Mean Squares	F
TOTAL	155	2121.2544		
GENOTYPES	12	101.0877	8.4240	3.31 *
ENVIRONMENTS	11	1684.3067	153.1188	85.07 **
INTERACTION	132	335.8600	2.5444	1.41 **
HETEROGENEITY	12	34.7256	2.8938	1.15 ns
RESIDUAL	120	301.1344	2.5095	1.39 *
POOLED ERROR	432		1.8000	



## Stability analysis (parametric)

### stability.par

```
(data, rep, MSerror, alpha = 0.1, main = NULL, cov =
F, name.cov = NULL, file.cov = 0)
```

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

  Genotype  Yield Rank Adj.rank Adjusted Stab.var Stab.rating YSi ...
1          A 7.383333 11      1      12 2.134311      0 12 +
2          B 6.783333  2     -1      1 1.672824      0  1
3          C 7.250000  9      1     10 0.805606      0 10 +
4          D 6.783333  2     -1      1 2.919766     -2 -1
5          E 7.075000  7     -1      6 1.604036      0  6 +
6          F 6.916667  6     -1      5 3.924945     -2  3
7          G 7.808333 12      2     14 4.043485     -2 12 +
8          H 7.908333 13      2     15 2.899022     -2 13 +
9          I 7.275000 10      1     11 4.251970     -2  9 +
10         J 7.083333  8     -1      7 1.853320      0  7 +
11         K 6.433333  1     -2     -1 2.167039      0 -1
12         L 6.891667  5     -1      4 1.692631      0  4
13         M 6.791667  4     -1      3 3.108168     -2  1

Yield Mean: 7.10641
YS      Mean: 5.846154
LSD (0.05): 0.4514298
-----
+ selected genotype
++ Reference: Kang, M. S. 1993. Simultaneous selection for yield
and stability: Consequences for growers. Agron. J. 85:754-757
```

## Stability analysis (Non-parametric)

Haynes K G, Lambert D H, Christ B J, Weingartner D P, Douches D S, Backlund J E, Fry W and Stevenson W. 1998. Phenotypic stability of resistance to late blight in potato clones evaluated at eight sites in the United States American Journal Potato Research 75, pag 211-217.

```
Stability.nonpar(data, variable=NULL, ranking = FALSE)
```

```
> haynes
  clone FL  MI  ME ...
1  A84118-3 284 1113 1053 ...
2  AO80432-1 254  690 1112 ...
3  AO84275-3 395 1089 1090 ...
4 AWN86514-2 136  296  374 ...
5  B0692-4  87  653  412 ...
6  B0718-3 130  126  329 ...
... ..

stability.nonpar(haynes,"YIELD",ranking=TRUE)

Nonparametric Method for Stability Analysis
-----
Estimation and test of nonparametric measures
Variable: YIELD
Ranking...
      FL MI ME MN ND  NY PA WI
A84118-3  7 11 11 14  8 14.0 12 11
AO80432-1  6  9 13 13 12 12.0 15 14
AO84275-3 10 10 12  8  9  7.0 11 12
AWN86514-2  3  3  3  1  3  3.0  2  1
B0692-4    1  8  4  3  2  2.0  1  3
B0718-3    2  1  2  2  4  4.0  3  4
... ..
```

## Stability analysis Non-parametric

Haynes K G, Lambert D H, Christ B J, Weingartner D P, Douches D S, Backlund J E, Fry W and Stevenson W. 1998. Phenotypic stability of resistance to late blight in potato clones evaluated at eight sites in the United States American Journal Potato Research 75, pag 211-217.

`Stability.nonpar(data, variable=NULL, ranking = FALSE)`

Statistics...

	Mean Rank	s1	Z1	s2	Z2
A84118-3	741.62	13	4.82	0.22	16.70
A080432-1	734.38	12	6.21	0.73	26.57
A084275-3	635.88	9	6.20	0.70	28.53
...					
Sum of Z1:	20.08986				
Sum of Z2:	25.84532				

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.

MEAN	es1	es2	vs1	vs2	chi.ind	chi.sum
561.4609	5.3125	21.25	1.111905	60.75223	8.733011	26.29623

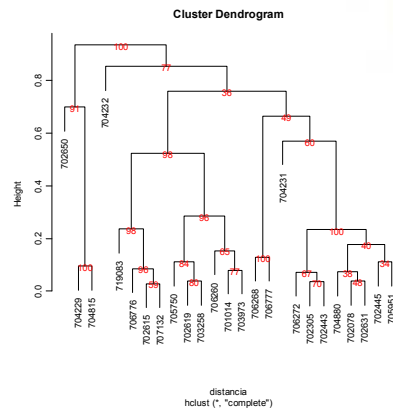
## Consensus cluster

Methods distance and clustering of R, functions `dist()` and `hclust()`.

`(data, distance = c("binary", ..), method = c("complete", ..), nboot = 500, duplicate = TRUE, cex.text = 1, col.text = "red", ...)`

`output<-consensus( pamCIP,distance="binary", method="complete", nboot=500)`

Duplicates: 18  
 New data : 25 Records  
 Consensus hclust  
 Method distance: binary  
 Method cluster : complete  
 rows and cols : 25 107  
 n-boostrap : 500  
 Run time : 16.281 secs





## Consensus cluster

Input: output consensus

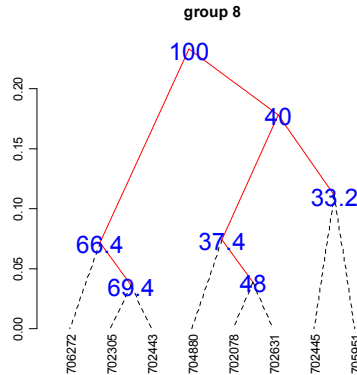
hcut()

```
(consensus, h, group, col.text = "blue", cex.text = 1, ...)
```

```
hcut(output,h=0.4,group=8,type="t",edgePar = list(lty=1:2,
col=2:1),main="group 8" ,col.text="blue",cex.text=2)
```

numbers

```
1 1
2 2
3 1
4 4
5 6
6 2
7 1
8 8
```



## Soil uniformity

Index.smith(data, ...)

```
table<-index.smith(rice, type="l",lty=4, lwd=3,
main="Relationship between CV\n per unit area and plot
size",col="red")
```

Smith's index of soil heterogeneity is used primarily to derive optimum plot size. The index gives a single value as a quantitative measure of soil heterogeneity in an area. The coefficient of variance is used to determine plot size and shape

```
> table
```

```
$model
lm(formula = CV ~
I(log(x)))
Coefficients:
(Intercept)    I(log(x))
 12.4782      -0.7009
```

```
$uniformity
      Size Width Length plots      Vx  CV
[1,]    1     1     1    648 9044.539 13.0
[2,]    2     1     2    324 7816.068 12.1
[3,]    2     2     1    324 7831.232 12.1
[4,]    3     1     3    216 7347.975 11.7
[5,]    3     3     1    216 7355.216 11.7
...
[40,]  162     9    18     4 4009.765  8.6
```

## Soil uniformity

```
Index.smith(data, ...)
```

```
table<-index.smith(rice, type="l",lty=4, lwd=3,
main="Relationship between CV\n per unit area and plot
size",col="red")
```

```
predict(table$model, new=data.frame(x=30))
```

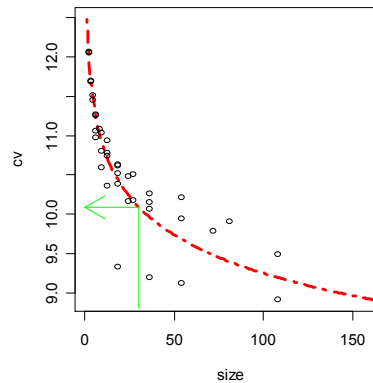
Relationship between CV  
per unit area and plot size

```
[1] 10.09436
```

If plot size = 30 unit <sup>2</sup>  
then CV = 10 %

rice

	V1	V2	V3	V4	V5
1	842	844	808	822	97
2	803	841	870	970	94
3	773	782	860	822	93
4	912	887	815	937	84
5	874	792	803	793	81
6	908	875	899	788	84
7	875	907	921	963	87
8	891	928	871	875	86
9	823	784	754	873	79



## Other functions and data sets

Genetic design: north carolina design, line x tester.  
Biodiversity index and confidence interval.  
Descriptive statistical: cross tabulations, ...  
Model: simulation and resampling.

Data sets main in package 'agricolae':

ComasOxapampa	Data AUDPC Comas - Oxapampa
Glycoalkaloids	Data Glycoalkaloids
RioChillon	Data and analysis Mother and baby trials
clay	Data of Ralstonia population in clay soil
disease	Data evaluation of the disease overtime
huasahuasi	Data of yield in Huasahuasi
melon	Data of yield of melon in a Latin square experiment
natives	Data of native potato
pamCIP	Data Potato Wild
paracsho	Data of Paracsho biodiversity
ralstonia	Data of population bacterial Wilt: AUDPC
soil	Data of soil analysis for 13 localities
sweetpotato	Data of sweetpotato yield
trees	Data of species trees. Pucallpa
wilt	Data of Bacterial Wilt (AUDPC) and soil

## **Agricolae Version 1.0-4**

Please note that there is a new version of the agricolae on the link below

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<http://tarwi.lamolina.edu.pe/~fmendiburu>