Data Analysis with augmentedRCBD

Aravind, J.¹, Mukesh Sankar, S.², Wankhede, D. P.³, and Kaur, V.⁴

2023-05-27

- 1. Division of Germplasm Conservation, ICAR-National Bureau of Plant Genetic Resources, New Delhi.
 - 2. Division of Genetics, ICAR-Indian Agricultural Research Institute, New Delhi.
- 3. Division of Genomic Resources, ICAR-National Bureau of Plant Genetic Resources, New Delhi.
- 4. Division of Germplasm Evaluation, ICAR-National Bureau of Plant Genetic Resources, New Delhi.

Contents

2
2
3
3
3
4
5
5
5
6
6
6
6
7
7
8
0
0
1
1
2
2
5
5
5
7

5 Installation of augmentedRCBD	17
6 Data Format	18
7 Data Analysis for a Single Trait	20
7.1 augmentedRCBD()	21
7.2 print.augmentedRCBD()	
7.3 describe.augmentedRCBD()	47
7.4 freqdist.augmentedRCBD()	48
7.5 gva.augmentedRCBD()	
$7.5~{ t report.augmentedRCBD}$ ()	56
8 Data Analysis for a Multiple Traits	58
8.1 augmentedRCBD.bulk()	58
8.2 print.augmentedRCBD.bulk()	
8.3 report.augmentedRCBD.bulk()	
9 Citing augmentedRCBD	66
10 Session Info	67
References	68

1 Overview

The software augmentedRCBD is built on the R statistical programming language as an add-on (or 'package' in the R lingua franca). It performs the analysis of data generated from experiments in augmented randomised complete block design according to Federer, W.T. (1956a, 1956b, 1961; 1976). It also computes analysis of variance, adjusted means, descriptive statistics, genetic variability statistics etc. and includes options for data visualization and report generation.

This tutorial aims to educate the users in utilising this package for performing such analysis. Utilising ${\tt augmentedRCBD}$ for data analysis requires a basic knowledge of ${\tt R}$ programming language. However, as many of the intended end-users may not be familiar with ${\tt R},$ sections 2 to 4 give a 'gentle' introduction to ${\tt R},$ especially those aspects which are necessary to get ${\tt augmentedRCBD}$ up and running for performing data analysis in a Windows environment. Users already familiar with ${\tt R}$ can feel free to skip to section 5.



2 R software

It is a free software environment for statistical computing and graphics. It is free and open source, platform independent (works on Linux, Windows or MacOS), very flexible, comprehensive with robust interfaces for all the popular programming languages as well as databases. It is strengthened by its diverse library of



add-on packages extending its ability as well as the incredible community support. It is one of the most popular tools being used in academia today (Tippmann, 2015).

3 Getting Started

This section details the steps required to set up the **R** programming environment under a third-party interface called **RStudio** in Windows.

3.1 Installing R

Download and install **R** for Windows from http://cran.r-project.org/bin/windows/base/.

R-3.5.1 for Windows (32/64 bit)

Download R 3.5.1 for Windows (62 megabytes, 32/64 bit)

<u>Installation and other instructions</u> <u>New features in this version</u>

If you want to double-check that the package you have downloaded matches the package distributed by CRAN, you can compare the md5sum of the .exe to the fingerprint on the master server. You will need a version of md5sum for windows: both graphical and command line versions are available.

Frequently asked questions

- Does R run under my version of Windows?
- How do I update packages in my previous version of R?
- Should I run 32-bit or 64-bit R?

Please see the RFAQ for general information about R and the R Windows FAQ for Windows-specific information.

Other builds

- Patches to this release are incorporated in the <u>r-patched snapshot build</u>.
- · A build of the development version (which will eventually become the next major release of R) is available in the r-devel snapshot build
- Previous releases

Note to webmasters: A stable link which will redirect to the current Windows binary release is scranning-nc-nt-windows/base/release.htm.

Last change: 2018-07-02

Fig. 1: The R download location.

3.2 Installing RStudio

The basic command line interface in native **R** is rather limiting. There are several interfaces which enhance it's functionality and ease of use, **RStudio** being one of the most popular among **R** programmers.

 $Download \ and \ install \ \textbf{RStudio} \ for \ Windows \ from \ https://www.rstudio.com/products/rstudio/download/\#download}$

Installers for Supported Platforms

Installers	Size	Date	MD5
RStudio 1.1.456 - Windows Vista/7/8/10	85.8 MB	2018-07-19	24ca3fe0dad8187aabd4bfbb9dc2b5ad
RStudio 1.1.456 - Mac OS X 10.6+ (64-bit)	74.5 MB	2018-07-19	4fc4f4f70845b142bf96dc1a5b1dc556
RStudio 1.1.456 - Ubuntu 12.04-15.10/Debian 8 (32-bit)	89.3 MB	2018-07-19	3493f9d5839e3a3d697f40b7bb1ce961
RStudio 1.1.456 - Ubuntu 12.04-15.10/Debian 8 (64-bit)	97.4 MB	2018-07-19	863ae806120358fa0146e4d14cd75be4
RStudio 1.1.456 - Ubuntu 16.04+/Debian 9+ (64-bit)	64.9 MB	2018-07-19	d96e63548c2add890bac633bdb883f32
RStudio 1.1.456 - Fedora 19+/RedHat 7+/openSUSE 13.1+ (32-bit)	88.1 MB	2018-07-19	1df56c7cd80e2634f8a9fdd11ca1fb2d
RStudio 1.1.456 - Fedora 19+/RedHat 7+/openSUSE 13.1+ (64-bit)	90.6 MB	2018-07-19	5e77094a88fdbddddddb0d35708752462

Zip/Tarballs

Zip/tar archives	Size	Date	MD5
RStudio 1.1.456 - Windows Vista/7/8/10	122.9 MB	2018-07-19	659d6bfe716d8c97acbe501270d89fa3
RStudio 1.1.456 - Ubuntu 12.04-15.10/Debian 8 (32-bit)	90 MB	2018-07-19	63117c159deca4d01221a8069bd45373
RStudio 1.1.456 - Ubuntu 12.04-15.10/Debian 8 (64-bit)	98.3 MB	2018-07-19	c53c32a71a400c6571e36c573f83dfde
RStudio 1.1.456 - Fedora 19+/RedHat 7+/openSUSE 13.1+ (32-bit)	88.8 MB	2018-07-19	f4ba2509fb00e30c91414c6821f1c85f
RStudio 1.1.456 - Fedora 19+/RedHat 7+/openSUSE 13.1+ (64-bit)	91.4 MB	2018-07-19	c60db6467421aa86c772227da0945a13

Source Code

A tarball containing source code for RStudio v1.1.456 can be downloaded from here

Fig. 2: The RStudio download location.

3.3 The RStudio Interface

On opening **RStudio**, the default interface with four panes/windows is visible as follows. Few panes have different tabs.

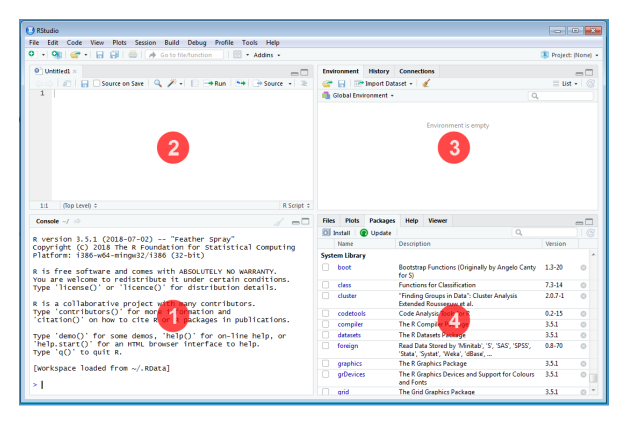


Fig. 3: The default RStudio interface with the four panes.

3.3.1 Console

This is where the action happens. Here any authentic \mathbf{R} code typed after the '>' prompt will be executed after pressing 'Enter' to generate the output.

For example, type **1+1** in the console and press 'Enter'.

1+1

[1] 2

3.3.2 Source

This is where \mathbf{R} Scripts (collection of code) can be created and edited. \mathbf{R} scripts are text files with a . \mathbf{R} extension. \mathbf{R} Code for analysis can be typed and saved in such \mathbf{R} scripts. New scripts can be opened by clicking 'File|New File' and selecting 'R Script'. Code can be selected from \mathbf{R} Scripts and sent to console for evaluation by clicking 'Run' on the 'Source' pane or by pressing 'Ctrl + Enter'.

3.3.3 Environment | History | Connections

The 'Environment' tab shows the list of all the 'objects' (see section 4.3) defined in the current **R** session. It has also some buttons up top to open, save and clear the environment as well as few options for import of data under **Import Dataset**.

The 'History' tab shows a history of all the code that was previously evaluated. This is useful, if you want to go back to some code.

The 'Connections' tab helps to establish and manage connections with different databases and data sources.

3.3.4 Files|Plots|Packages|Help|Viewer

The 'Files' tab shows a sleek file browser to access the file directory in the computer with options to manage the working directory (see section 4.1) under the More button.

The 'Plots' tab shows all the plots generated in \mathbf{R} with buttons to delete unnecessary ones and export useful ones as a pdf file or as an image file.

The 'Packages' tab shows a list of all the \mathbf{R} add-on packages installed. The check box on the left shows whether they are loaded or not. There are also buttons to install and update \mathbf{R} packages.

The 'Viewer' tab shows any web content output generated by an R code.

4 Some Basics

This section describes some basics to enable the users to have a working knowledge in \mathbf{R} in order to use **augmentedRCBD**.

4.1 Working Directory

It is a file path to a folder on the computer which is recognised by **R** as the default location to read files from or write files to. The code **getwd()** shows the current working directory, while **setwd()** can be used to change the existing working directory.

```
# Print current working directory
getwd()
```

[1] "C:/Users/Computer/Documents"

```
# Set new working directory
setwd("C:/Data Analysis/")
getwd()
```

[1] "C:/Data Analysis/"

[1] 3

One key detail is that file paths in \mathbf{R} uses forward slashes (/) as in MacOS or Linux, unlike backward slashes (\(\)) in Windows. This needs to be considered while copying paths from default Windows file explorer.

4.2 Expression and Assignment

Expressions are instructions in the form of code to be entered after the > prompt in the console. Expressions can be a constant, an arithmetic or a condition. A more advanced and most useful expression is a function call (see section 4.3).

```
# Constant
123
[1] 123
# Arithmetic (add two numbers)
1 + 2
```

```
# Condition
34 > 25

[1] TRUE
1 == 2

[1] FALSE
# Function call (mean of a series of numbers)
mean(c(25,56,89,35))

[1] 51.25
```

Information from an expression can be stored as an 'object' (see section 4.3) by assigning a name using the operator '<-'.

```
\# Assign the result of the expression 1 + 2 to an object 'a' a <- 1 + 2 a
```

[1] 3

It is recommended to add comments to explain the code by using the '#' sign. Any code after the '#' sign will be ignored by R.

4.3 Objects and Functions

 ${\tt R}$ is an object-oriented programming language (OOP). Any kind or construct created in ${\tt R}$ is an 'object'. Each object has a 'class' (shown using the ${\tt class}$ () function) and different 'attributes' which defines what operations can be done on that object. There are different types of data structure objects in ${\tt R}$ such as vectors, matrices, factors, data frames, and lists. A 'function' is also an object, which defines a procedure or a sequence of expressions.

4.3.1 Vector

A vector is a collection of elements of a single type (or 'mode'). The common vector modes are 'numeric', 'integer', 'character' and 'logical'. The c() function is used to create vectors. The functions class(), str() and length() show the attributes of vectors.

Vector modes 'numeric' stores real numbers, while 'integer' stores integers, which can be enforced by suffixing elements with 'L'.

```
# A numeric vector
a <- c(1, 2, 3.3)
class(a)

[1] "numeric"
str(a)

num [1:3] 1 2 3.3
length(a)

[1] 3
# An integer vector
b <- c(1L, 2L, 3L)
class(b)</pre>
```

Levels: female male

```
[1] "integer"
str(b)
 int [1:3] 1 2 3
length(b)
[1] 3
The vector mode 'character' store text.
# A character vector
c <- c("one", "two", "three")</pre>
class(c)
[1] "character"
str(c)
 chr [1:3] "one" "two" "three"
length(c)
[1] 3
The vector mode 'logical' stores 'TRUE' OR 'FALSE' logical data.
#logical vector
d <- c(TRUE, TRUE, TRUE, FALSE, TRUE, FALSE)</pre>
class(d)
[1] "logical"
str(d)
 logi [1:6] TRUE TRUE TRUE FALSE TRUE FALSE
length(d)
[1] 6
4.3.2 Factor
A 'factor' in R stores data from categorical data in variables as different levels.
catg <- c("male", "female", "female", "male", "male")</pre>
catg
              "female" "female" "male"
[1] "male"
                                             "male"
is.factor(catg)
[1] FALSE
# Apply the factor function
factor_catg <- factor(catg)</pre>
factor_catg
[1] male
            female female male
                                    male
```

```
is.factor(factor_catg)
[1] TRUE
class(factor_catg)
[1] "factor"
str(factor_catg)
Factor w/ 2 levels "female", "male": 2 1 1 2 2
A character, numeric or integer vector can be transformed to a factor by using the as.factor()
function.
# Conversion of numeric to factor
a \leftarrow c(1, 2, 3.3)
class(a)
[1] "numeric"
str(a)
num [1:3] 1 2 3.3
fac_a <- as.factor(a)</pre>
class(fac_a)
[1] "factor"
str(fac_a)
Factor w/ 3 levels "1", "2", "3.3": 1 2 3
# Conversion of integer to factor
b \leftarrow c(1L, 2L, 3L)
class(b)
[1] "integer"
str(b)
int [1:3] 1 2 3
fac_b <- as.factor(b)</pre>
class(fac_b)
[1] "factor"
str(fac_b)
Factor w/ 3 levels "1", "2", "3": 1 2 3
# Conversion of character to factor
c <- c("one", "two", "three")</pre>
class(c)
[1] "character"
str(c)
 chr [1:3] "one" "two" "three"
```

```
fac_c <- as.factor(c)</pre>
class(fac_c)
[1] "factor"
str(fac_c)
Factor w/ 3 levels "one", "three", ..: 1 3 2
4.3.3 Matrix
A 'matrix' in R is a vector with the attributes 'nrow' and 'ncol'.
# Generate 5 * 4 numeric matrix
m <- matrix(1:20, nrow = 5, ncol = 4)</pre>
     [,1] [,2] [,3] [,4]
[1,]
        1
                   11
              7
                        17
[2,]
        2
                   12
[3,]
        3
              8
                   13
                        18
         4
              9
                  14
                        19
[4,]
[5,]
             10
                   15
class(m)
[1] "matrix" "array"
typeof(m)
[1] "integer"
# Dimensions of m
dim(m)
```

[1] 5 4

4.3.4 List

A 'list' is a container containing different objects. The contents of list need not be of the same type or mode. A list can encompass a mixture of data types such as vectors, matrices, data frames, other lists or any other data structure.

```
w <- list(a, m, d, list(b, c))</pre>
class(w)
[1] "list"
str(w)
List of 4
 $ : num [1:3] 1 2 3.3
 $ : int [1:5, 1:4] 1 2 3 4 5 6 7 8 9 10 ...
 $ : logi [1:6] TRUE TRUE TRUE FALSE TRUE FALSE
 $:List of 2
  ..$ : int [1:3] 1 2 3
  ..$ : chr [1:3] "one" "two" "three"
```

4.3.5 Data Frame

A 'data frame' in \mathbf{R} is a special kind of list with every element having equal length. It is very important for handling tabular data in \mathbf{R} . It is a array like structure with rows and columns. Each column needs to be of a single data type, however data type can vary between columns.

```
L <- LETTERS[1:4]
y < -1:4
z <- c("This", "is", "a", "data frame")</pre>
df \leftarrow data.frame(L, x = 1, y, z)
  Lху
1 A 1 1
               This
2 B 1 2
                 is
3 C 1 3
4 D 1 4 data frame
str(df)
'data.frame':
                 4 obs. of 4 variables:
 $ L: chr "A" "B" "C" "D"
 $ x: num 1 1 1 1
 $ y: int 1 2 3 4
 $ z: chr "This" "is" "a" "data frame"
attributes(df)
$names
[1] "L" "x" "y" "z"
$class
[1] "data.frame"
$row.names
[1] 1 2 3 4
rownames (df)
[1] "1" "2" "3" "4"
colnames (df)
[1] "L" "x" "y" "z"
```

4.3.6 Functions

All of the work in **R** is done by functions. It is an object defining a procedure which takes one or more objects as input (or 'arguments'), performs some action on them and finally gives a new object as output (or 'return'). class(), mean(), getwd(), +, etc. are all functions.

For example the function **mean()** takes a numeric vector as argument and returns the mean as a numeric vector.

```
a <- c(1, 2, 3.3)
mean(a)
```

[1] 2.1

The user can also create custom functions. For example the function **foo** adds two numbers and gives the result.

```
foo <- function(n1, n2) {
  out <- n1 + n2
  return(out)
}
foo(2,3)</pre>
```

[1] 5

4.4 Special Elements

In addition to numbers and text, there are some special elements which can be included in different data objects.

NA (not available) indicates missing data.

```
x <- c(2.5, NA, 8.6)

y <- c(TRUE, FALSE, NA)

z <- c("k", NA, "m", "n", "o")

is.na(x)
```

[1] FALSE TRUE FALSE

is.na(z)

[1] FALSE TRUE FALSE FALSE FALSE

anyNA(x)

[1] TRUE

а

[1] 1.0 2.0 3.3

is.na(a)

[1] FALSE FALSE FALSE

Inf indicates infinity.

1/0

[1] Inf

NaN (Not a Number) indicates any undefined value.

0/0

[1] NaN

4.5 Indexing

The [function is used to extract elements of an object by indexing (numeric or logical). Named elements in lists and data frames can be extracted by using the \$ operator.

Consider a vector a.

```
a \leftarrow c(1, 2, 3.3, 2.8, 6.7)
# Numeric indexing
# Extract first element
a[1]
[1] 1
# Extract elements 2:3
a[2:3]
[1] 2.0 3.3
# Logical indexing
a[a > 3]
[1] 3.3 6.7
Consider a matrix \mathbf{m}.
m <- matrix(1:9, nrow = 3, ncol = 3, byrow = TRUE)</pre>
colnames(m) <- c('a', 'b', 'c')</pre>
     a b c
[1,] 1 2 3
[2,] 4 5 6
[3,] 7 8 9
# Extract elements
m[,2] # 2nd column of matrix
[1] 2 5 8
m[3,] # 3rd row of matrix
abc
7 8 9
m[2:3, 1:3] # rows 2,3 of columns 1,2,3
     abc
[1,] 4 5 6
[2,] 7 8 9
m[2,2] # Element in 2nd column of 2nd row
b
5
m[, 'b'] # Column 'b'
[1] 2 5 8
m[, c('a', 'c')] # Column 'a' and 'c'
     аc
[1,] 1 3
[2,] 4 6
[3,] 7 9
Consider a list {\bf w}.
```

```
w <- list(vec = a, mat = m, data = df, alist = list(b, c))</pre>
# Indexing by number
w[2] # As list structure
$mat
    a b c
[1,] 1 2 3
[2,] 4 5 6
[3,] 7 8 9
w[[2]] # Without list structure
    a b c
[1,] 1 2 3
[2,] 4 5 6
[3,] 7 8 9
# Indexing by name
w$vec
[1] 1.0 2.0 3.3 2.8 6.7
w$data
 Lху
               Z
1 A 1 1
            This
2 B 1 2
              is
3 C 1 3
4 D 1 4 data frame
Consider a data frame df.
df
 Lxy
               Z
            This
1 A 1 1
2 B 1 2
              is
3 C 1 3
4 D 1 4 data frame
# Indexing by number
df[,2] # 2nd column of data frame
[1] 1 1 1 1
df[2] # 2nd column of data frame
 x
1 1
2 1
3 1
4 1
df[3,] # 3rd row of data frame
 Lxyz
3 C 1 3 a
```

```
df[2:3, 1:3] # rows 2,3 of columns 1,2,3

L x y
2 B 1 2
3 C 1 3

df[2,2] # Element in 2nd column of 2nd row

[1] 1
# Indexing by name
df$L

[1] "A" "B" "C" "D"

df$z

[1] "This" "is" "a" "data frame"
```

4.6 Help Documentation

The help documentation regarding any function can be viewed using the ? or help() function. The help documentation shows the default usage of the function including, the arguments that are taken by the function and the type of output object returned ('Value').

```
?ls
help(ls)
?mean
?setwd
```

4.7 Packages

Packages in \mathbf{R} are collections of \mathbf{R} functions, data, and compiled code in a well-defined format. They are add-ons which extend the functionality of \mathbf{R} and at present, there are 19555 packages available for deployment and use at the official repository, the Comprehensive R Archive Network (CRAN).

Valid packages from CRAN can be installed by using the install.packages() command.

```
# Install the package 'readxl' for importing data from excel
install.packages(readxl)
```

Installed packages can be loaded using the function library().

```
# Install the package 'readxl' for importing data from excel
library(readxl)
```

4.8 Importing and Exporting Tabular Data

Tabular data from a spreadsheet can be imported into **R** in different ways. Consider some data such as in Table 1. Copy this data in to a spreadsheet editor such as MS Excel and save it as **augdata.csv**, a comma-separated-value file and **augdata.xlsx**, an Excel file in the working directory (**getwd()**).

Table 1: Example data from an experiment in augmented RCBD design.

trt	y1	y2
1	92	258
2	79	224
3	87	238
4	81	278
7	96	347
11	89	300
12	82	289
1	79	260
2	81	220
3	81	237
4	91	227
5	79	281
9	78	311
1	83	250
2	77	240
3	78	268
4	78	287
8	70	226
6	75	395
10	74	450
	1 2 3 4 7 11 12 1 2 3 4 5 9 1 2 3 4 8 6	1 92 2 79 3 87 4 81 7 96 11 89 12 82 1 79 2 81 3 81 4 91 5 79 9 78 1 83 2 77 3 78 4 78 8 70 6 75

The augdata.csv file can be imported into R using the read.csv() function or the read_csv() function in the readr package.

```
data <- read.csv(file = "augdata.csv")
str(data)

'data.frame': 20 obs. of 4 variables:
$ blk: Factor w/ 3 levels "I","II","III": 1 1 1 1 1 1 1 2 2 2 ...
$ trt: num 1 2 3 4 7 11 12 1 2 3 ...
$ y1 : num 92 79 87 81 96 89 82 79 81 81 ...
$ y2 : num 258 224 238 278 347 300 289 260 220 237 ...</pre>
The argument stringsAsFactors = FALSE reads the text columns as of type character instead of
```

The argument **stringsAsFactors** = **FALSE** reads the text columns as of type **character** instead of the default **factor**.

```
data <- read.csv(file = "augdata.csv", stringsAsFactors = FALSE)
str(data)
'data.frame': 20 obs. of 4 variables:</pre>
```

The augdata.xlsx file can be imported into R using the read_excel() function in the readxl package.

```
library(readxl)
data <- read_excel(path = "augdata.xlsx")

'data.frame': 20 obs. of 4 variables:
$ blk: chr "I" "I" "I" "I" ...</pre>
```

```
$ trt: num 1 2 3 4 7 11 12 1 2 3 ...
$ y1 : num 92 79 87 81 96 89 82 79 81 81 ...
$ y2 : num 258 224 238 278 347 300 289 260 220 237 ...
```

The tabular data can be exported from R to a .csv (comma-separated-value) file by the write.csv() function.

```
write.csv(x = data, file = "augdata.csv")
```

4.9 Additional Resources

To learn more about **R**, there are umpteen number of online tutorials as well as free courses available. Queries about various aspects can be put to the active and vibrant 'R community online.

- Online tutorials
 - http://www.cran.r-project.org/other-docs.html
 - https://bookdown.org/ndphillips/YaRrr/
- Free online courses
 - http://tryr.codeschool.com/
 - https://www.datacamp.com/courses/free-introduction-to-r
- R community support
 - http://stackoverflow.com/
 - **R** help mailing lists: http://www.r-project.org/mail.html

5 Installation of augmentedRCBD

The package augmentedRCBD can be installed using the following functions.

```
# Install from CRAN
install.packages('augmentedRCBD', dependencies=TRUE)

# Install development version from Github
if (!require('devtools')) install.packages('devtools')
library(devtools)
install_github("aravind-j/augmentedRCBD")
```

The stable release is hosted in CRAN (see section 4.7), while the under-development version is hosted as a Github repository. To install from github, you need to use the **install_github()** function from 'devtools package.

Then the package can be loaded using the function

```
library(augmentedRCBD)
```

```
Welcome to augmentedRCBD version 0.1.6
```

```
# To know how to use this package type:
  browseVignettes(package = 'augmentedRCBD')
  for the package vignette.
```

To know whats new in this version type: news(package='augmentedRCBD') for the NEWS file.

- # To cite the methods in the package type: citation(package='augmentedRCBD')
- # To suppress this message use: suppressPackageStartupMessages(library(augmentedRCBD))

The current version of the package is 0.1.6. The previous versions are as follows.

Table 2. Version history of augmentedRCBD R package.

Version	Date
0.1.0	2018-07-10
0.1.1	2019-07-21
0.1.2	2020-03-19
0.1.3	2020-07-27
0.1.4	2021-02-17

To know detailed history of changes use news (package='augmentedRCBD').

6 Data Format

Certain details need to be considered for arranging experimental data for analysis using the augmentedRCBD package.

The data should be in long/vertical form, where each row has the data from one genotype per block. For example, consider the following data (Table 3) recorded for a trait from an experiment laid out in an augmented block design with 3 blocks and 12 genotypes(or treatment) with 6 to 7 genotypes/block. 8 genotypes (Test, G 5 to G 12) are not replicated, while 4 genotypes (Check, G 1 to G 4) are replicated.

Table 3: Data from an experiment in augmented RCBD design.

Block I	G12	G4	G11	$\mathbf{G2}$	G1	G7	$\mathbf{G3}$
	82	81	89	79	92	96	87
Block II	G5	G9	_	$\mathbf{G3}$	$\mathbf{G1}$	G2	G4
	79	78	_	81	79	81	91
Block III	G4	G2	$\mathbf{G1}$	G6	G10	$\mathbf{G3}$	G8
	78	77	83	75	74	78	70

This data needs to be arranged with columns showing block, genotype (or treatment) and the data of the trait for each genotype per block (Table 4).

Table 4: Data from an experiment in augmented RCBD design arranged in long-form.

Block	Treatment	Trait
Block I	G 1	92
Block I	G 2	79
Block I	G 3	87

Block	Treatment	Trait
Block I	G 4	81
Block I	G 7	96
Block I	G 11	89
Block I	G 12	82
Block II	G 1	79
Block II	G 2	81
Block II	G 3	81
Block II	G 4	91
Block II	G 5	79
Block II	G 9	78
Block III	G 1	83
Block III	G 2	77
Block III	G 3	78
Block III	G 4	78
Block III	G 8	70
Block III	G 6	75
Block III	G 10	74

The data for block and genotype (or treatment) can also be depicted as numbers (Table 5).

Table 5: Data from an experiment in augmented RCBD design arranged in long-form (Block and Treatment as numbers).

Block	Treatment	Trait
1	1	92
1	2	79
1	3	87
1	4	81
1	7	96
1	11	89
1	12	82
2	1	79
2	2	81
2	3	81
2	4	91
2	5	79
2	9	78
3	1	83
3	2	77
3	3	78
3	4	78
3	8	70
3	6	75
3	10	74

Multiple traits can be added as additional columns (Table 6).

Table 6: Data from an experiment in augmented RCBD design arranged in long-form (Multiple traits).

Block	Treatment	Trait1	Trait2
Block I	G 1	92	258
Block I	G 2	79	224
Block I	G 3	87	238
Block I	G 4	81	278
Block I	G 7	96	347
Block I	G 11	89	300
Block I	G 12	82	289
Block II	G 1	79	260
Block II	G 2	81	220
Block II	G 3	81	237
Block II	G 4	91	227
Block II	G 5	79	281
Block II	G 9	78	311
Block III	G 1	83	250
Block III	G 2	77	240
Block III	G 3	78	268
Block III	G 4	78	287
Block III	G 8	70	226
Block III	G 6	75	395
Block III	G 10	74	450

Data should preferably be balanced i.e. all the check genotypes should be present in all the blocks. If not, a warning is issued. The number of test genotypes can vary within a block. There should not be any missing values. Rows of genotypes with missing values for one or more traits should be removed.

Such a tabular data should be imported (see section 7.8) into \mathbf{R} as a data frame object (see section 4.3.5). The columns with the block and treatment categorical data should of the type factor (see section 4.3.2), while the column(s) with the trait data should be of the type integer or numeric (see section 4.3.1).

7 Data Analysis for a Single Trait

Analysis of data for a single trait can be performed by using augmentedRCBD function. It generates an object of class augmentedRCBD. Such an object can then be taken as input by the several functions to print the results to console (print.augmentedRCBD), generate descriptive statistics from adjusted means (describe.augmentedRCBD), plot frequency distribution (freqdist.augmentedRCBD) and computed genetic variability statistics (gva.augmentedRCBD). All these outputs can also be exported as a MS Word report using the report.augmentedRCBD function.

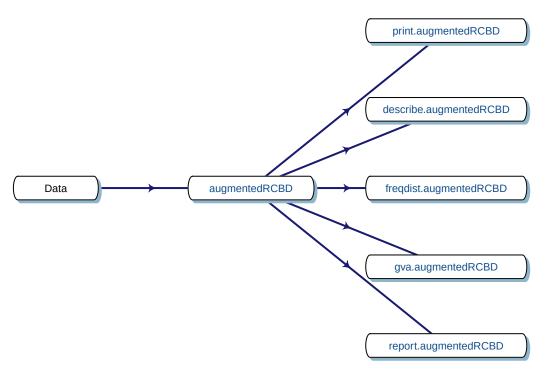


Fig. 4. Workflow for analysis of single traits with augmentedRCBD.

7.1 augmentedRCBD()

Consider the data in Table 1. The data can be imported into ${\bf R}$ as vectors as follows.

```
blk <- c(1, 1, 1, 1, 1, 1, 1, 2, 2, 2, 2, 2, 2, 2, 3, 3, 3, 3, 3, 3, 3, 3)

trt <- c(1, 2, 3, 4, 7, 11, 12, 1, 2, 3, 4, 5, 9, 1, 2, 3, 4, 8, 6, 10)

y1 <- c(92, 79, 87, 81, 96, 89, 82, 79, 81, 81, 91, 79, 78, 83, 77, 78, 78, 70, 75, 74)

y2 <- c(258, 224, 238, 278, 347, 300, 289, 260, 220, 237, 227, 281, 311, 250, 240, 268, 287, 226, 395, 450)
```

The **blk** and **trt** vectors with the block and treatment data need to be converted into factors as follows before analysis.

```
# Convert block and treatment to factors
blk <- as.factor(blk)
trt <- as.factor(trt)</pre>
```

With the data in appropriate format, the analysis can be performed as follows for the trait y1 as follows.

Number of treatments

"12"

Number of check treatments "4"

Number of test treatments "8"

Check treatments "1, 2, 3, 4"

ANOVA, Treatment Adjusted

Df Sum Sq Mean Sq F value Pr(>F)
Block (ignoring Treatments) 2 360.1 180.04 6.675 0.0298 *
Treatment (eliminating Blocks) 11 285.1 25.92 0.961 0.5499
Treatment: Check 3 52.9 17.64 0.654 0.6092
Treatment: Test and Test vs. Check 8 232.2 29.02 1.076 0.4779
Residuals 6 161.8 26.97

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

ANOVA, Block Adjusted

Df Sum Sq Mean Sq F value Pr(>F)
Treatment (ignoring Blocks) 11 575.7 52.33 1.940 0.215
Treatment: Check 3 52.9 17.64 0.654 0.609
Treatment: Test 7 505.9 72.27 2.679 0.125
Treatment: Test vs. Check 1 16.9 16.87 0.626 0.459
Block (eliminating Treatments) 2 69.5 34.75 1.288 0.342
Residuals 6 161.8 26.97

Coefficient of Variation

6.372367

Overall Adjusted Mean

81.0625

Standard Errors

Std. Error of Diff. CD (5%)
Control Treatment Means 4.240458 10.37603
Two Test Treatments (Same Block) 7.344688 17.97180
Two Test Treatments (Different Blocks) 8.211611 20.09309
A Test Treatment and a Control Treatment 6.704752 16.40594

Treatment Means

==========

Treatment	Block	Means	SE	r	Min	Max	Adjusted	Means
1		84.67	3.84	3	79.00	92.00		84.67
10	3	74.00	<na></na>	1	74.00	74.00		77.25
11	1	89.00	<na></na>	1	89.00	89.00		86.50
12	1	82.00	<na></na>	1	82.00	82.00		79.50
2		79.00	1.15	3	77.00	81.00		79.00
3		82.00	2.65	3	78.00	87.00		82.00
4		83.33	3.93	3	78.00	91.00		83.33

5	2	79.00	<na></na>	1	79.00	79.00	78.25
6	3	75.00	<na></na>	1	75.00	75.00	78.25
7	1	96.00	<na></na>	1	96.00	96.00	93.50
8	3	70.00	<na></na>	1	70.00	70.00	73.25
9	2	78.00	<na></na>	1	78.00	78.00	77.25

${\tt Comparisons}$

========

Method : lsd

contrast	estimate	SE	df	t.ratio	p.value	sig
treatment1 - treatment2	5.67	4.24	6	1.336	0.230	
treatment1 - treatment3	2.67	4.24	6	0.629	0.553	
treatment1 - treatment4	1.33	4.24	6	0.314	0.764	
treatment1 - treatment5	6.42	6.36	6	1.009	0.352	
treatment1 - treatment6	6.42	6.36	6	1.009	0.352	
treatment1 - treatment7	-8.83	6.36	6	-1.389	0.214	
treatment1 - treatment8	11.42	6.36	6	1.795	0.123	
treatment1 - treatment9	7.42	6.36	6	1.166	0.288	
treatment1 - treatment10	7.42	6.36	6	1.166	0.288	
treatment1 - treatment11	-1.83	6.36	6	-0.288	0.783	
treatment1 - treatment12	5.17	6.36	6	0.812	0.448	
treatment2 - treatment3	-3.00	4.24	6	-0.707	0.506	
treatment2 - treatment4	-4.33	4.24	6	-1.022	0.346	
treatment2 - treatment5	0.75	6.36	6	0.118	0.910	
treatment2 - treatment6	0.75	6.36	6	0.118	0.910	
treatment2 - treatment7	-14.50	6.36	6	-2.280	0.063	
treatment2 - treatment8	5.75	6.36	6	0.904	0.401	
treatment2 - treatment9	1.75	6.36	6	0.275	0.792	
treatment2 - treatment10	1.75	6.36	6	0.275	0.792	
treatment2 - treatment11	-7.50	6.36	6	-1.179	0.283	
treatment2 - treatment12	-0.50	6.36	6	-0.079	0.940	
treatment3 - treatment4	-1.33	4.24	6	-0.314	0.764	
treatment3 - treatment5	3.75	6.36	6	0.590	0.577	
treatment3 - treatment6	3.75	6.36	6	0.590	0.577	
treatment3 - treatment7	-11.50	6.36	6	-1.808	0.121	
treatment3 - treatment8	8.75	6.36	6	1.376	0.218	
treatment3 - treatment9	4.75	6.36	6	0.747	0.483	
treatment3 - treatment10	4.75	6.36	6	0.747	0.483	
treatment3 - treatment11	-4.50	6.36	6	-0.707	0.506	
treatment3 - treatment12	2.50	6.36	6	0.393	0.708	
treatment4 - treatment5	5.08	6.36	6	0.799	0.455	
treatment4 - treatment6	5.08	6.36	6	0.799	0.455	
treatment4 - treatment7	-10.17	6.36	6	-1.598	0.161	
treatment4 - treatment8	10.08	6.36	6	1.585	0.164	
treatment4 - treatment9	6.08	6.36	6	0.956	0.376	
treatment4 - treatment10	6.08	6.36	6	0.956	0.376	
treatment4 - treatment11	-3.17	6.36	6	-0.498	0.636	
treatment4 - treatment12	3.83	6.36	6	0.603	0.569	
treatment5 - treatment6	0.00	8.21	6	0.000	1.000	

```
treatment5 - treatment7
                        -15.25 8.21 6 -1.857
                                                 0.113
 treatment5 - treatment8
                            5.00 8.21 6 0.609
                                                 0.565
 treatment5 - treatment9
                           1.00 7.34 6
                                         0.136
                                                 0.896
 treatment5 - treatment10
                           1.00 8.21 6 0.122
                                                 0.907
                          -8.25 8.21 6 -1.005
 treatment5 - treatment11
                                                 0.354
 treatment5 - treatment12
                          -1.25 8.21 6 -0.152
                                                 0.884
 treatment6 - treatment7
                          -15.25 8.21 6 -1.857
                                                 0.113
 treatment6 - treatment8 5.00 7.34 6 0.681
                                                 0.521
 treatment6 - treatment9
                          1.00 8.21 6 0.122
                                                 0.907
                           1.00 7.34 6 0.136
 treatment6 - treatment10
                                                 0.896
 treatment6 - treatment11
                          -8.25 8.21 6 -1.005
                                                 0.354
 treatment6 - treatment12 -1.25 8.21 6 -0.152
                                                 0.884
 treatment7 - treatment8
                          20.25 8.21 6 2.466
                                                 0.049
                           16.25 8.21 6 1.979
 treatment7 - treatment9
                                                 0.095
                           16.25 8.21 6 1.979
 treatment7 - treatment10
                                                 0.095
 treatment7 - treatment11
                           7.00 7.34 6 0.953
                                                 0.377
 treatment7 - treatment12
                           14.00 7.34 6 1.906
                                                 0.105
 treatment8 - treatment9
                           -4.00 8.21 6 -0.487
                                                 0.643
 treatment8 - treatment10
                          -4.00 7.34 6 -0.545
                                                 0.606
 treatment8 - treatment11
                          -13.25 8.21 6 -1.614
                                                 0.158
 treatment8 - treatment12
                           -6.25 8.21 6 -0.761
                                                 0.475
                           0.00 8.21 6
 treatment9 - treatment10
                                         0.000
                                                 1.000
 treatment9 - treatment11
                          -9.25 8.21 6 -1.126
                                                 0.303
 treatment9 - treatment12
                          -2.25 8.21 6 -0.274
                                                 0.793
treatment10 - treatment11
                           -9.25 8.21 6 -1.126
                                                 0.303
                           -2.25 8.21 6 -0.274
treatment10 - treatment12
                                                 0.793
                          7.00 7.34 6 0.953
treatment11 - treatment12
                                                 0.377
```

Treatment Groups

Method : 1sd

Treatment	Adjusted	Means	SE	df	lower.CL	upper.CL	Group
8		73.25	5.61	6	59.52	86.98	1
9		77.25	5.61	6	63.52	90.98	12
10		77.25	5.61	6	63.52	90.98	12
5		78.25	5.61	6	64.52	91.98	12
6		78.25	5.61	6	64.52	91.98	12
2		79.00	3.00	6	71.66	86.34	12
12		79.50	5.61	6	65.77	93.23	12
3		82.00	3.00	6	74.66	89.34	12
4		83.33	3.00	6	76.00	90.67	12
1		84.67	3.00	6	77.33	92.00	12
11		86.50	5.61	6	72.77	100.23	12
7		93.50	5.61	6	79.77	107.23	2
class (out1))						

[1] "augmentedRCBD"

Similarly the analysis for the trait y2 can be computed as follows.

```
out2 <- augmentedRCBD(blk, trt, y2, method.comp = "lsd",
              alpha = 0.05, group = TRUE, console = TRUE)
Augmented Design Details
"3"
Number of blocks
Number of treatments
                       "12"
Number of check treatments "4"
Number of test treatments "8"
Check treatments
                       "1, 2, 3, 4"
ANOVA, Treatment Adjusted
Df Sum Sq Mean Sq F value Pr(>F)
Block (ignoring Treatments)
                                2 7019 3510 12.261 0.007597 **
Treatment (eliminating Blocks)
                               11 58965 5360 18.727 0.000920 ***
 Treatment: Check
                                 3
                                   2150
                                           717 2.504 0.156116
 Treatment: Test and Test vs. Check 8 56815
                                            7102 24.810 0.000473 ***
Residuals
                                   1718
                                           286
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
ANOVA, Block Adjusted
_____
                           Df Sum Sq Mean Sq F value Pr(>F)
Treatment (ignoring Blocks) 11 64708 5883 20.550 0.000707 ***
                          3 2150
                                      717 2.504 0.156116
 Treatment: Check
                           7 34863
 Treatment: Test
                                      4980 17.399 0.001366 **
                          1 27694 27694 96.749 6.36e-05 ***
 Treatment: Test vs. Check
Block (eliminating Treatments) 2 1277 639 2.231 0.188645
Residuals
                            6 1717 286
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Coefficient of Variation
_____
6.057617
Overall Adjusted Mean
_____
298.4792
Standard Errors
                                    Std. Error of Diff. CD (5%)
Control Treatment Means
                                            13.81424 33.80224
Two Test Treatments (Same Block)
                                             23.92697 58.54719
Two Test Treatments (Different Blocks)
                                             26.75117 65.45775
A Test Treatment and a Control Treatment
                                             21.84224 53.44603
```

Treatment Means

Treatment	Block	Means	SE	r	Min	Max	Adjusted Means
1		256.00	3.06	3	250.00	260.00	256.00
10	3	450.00	<na></na>	1	450.00	450.00	437.67
11	1	300.00	<na></na>	1	300.00	300.00	299.42
12	1	289.00	<na></na>	1	289.00	289.00	288.42
2		228.00	6.11	3	220.00	240.00	228.00
3		247.67	10.17	3	237.00	268.00	247.67
4		264.00	18.68	3	227.00	287.00	264.00
5	2	281.00	<na></na>	1	281.00	281.00	293.92
6	3	395.00	<na></na>	1	395.00	395.00	382.67
7	1	347.00	<na></na>	1	347.00	347.00	346.42
8	3	226.00	<na></na>	1	226.00	226.00	213.67
9	2	311.00	<na></na>	1	311.00	311.00	323.92

Comparisons

========

Method : 1sd

contrast	${\tt estimate}$	SE	df	t.ratio	p.value	sig
treatment1 - treatment2	28.00	13.81	6	2.027	0.089	
treatment1 - treatment3	8.33	13.81	6	0.603	0.568	
treatment1 - treatment4	-8.00	13.81	6	-0.579	0.584	
treatment1 - treatment5	-37.92	20.72	6	-1.830	0.117	
treatment1 - treatment6	-126.67	20.72	6	-6.113	0.001	***
treatment1 - treatment7	-90.42	20.72	6	-4.363	0.005	**
treatment1 - treatment8	42.33	20.72	6	2.043	0.087	
treatment1 - treatment9	-67.92	20.72	6	-3.278	0.017	*
<pre>treatment1 - treatment10</pre>	-181.67	20.72	6	-8.767	0.000	***
treatment1 - treatment11	-43.42	20.72	6	-2.095	0.081	
treatment1 - treatment12	-32.42	20.72	6	-1.564	0.169	
treatment2 - treatment3	-19.67	13.81	6	-1.424	0.204	
treatment2 - treatment4	-36.00	13.81	6	-2.606	0.040	*
treatment2 - treatment5	-65.92	20.72	6	-3.181	0.019	*
treatment2 - treatment6	-154.67	20.72	6	-7.464	0.000	***
treatment2 - treatment7	-118.42	20.72	6	-5.715	0.001	**
treatment2 - treatment8	14.33	20.72	6	0.692	0.515	
treatment2 - treatment9	-95.92	20.72	6	-4.629	0.004	**
<pre>treatment2 - treatment10</pre>	-209.67	20.72	6	-10.118	0.000	***
<pre>treatment2 - treatment11</pre>	-71.42	20.72	6	-3.447	0.014	*
<pre>treatment2 - treatment12</pre>	-60.42	20.72	6	-2.916	0.027	*
treatment3 - treatment4	-16.33	13.81	6	-1.182	0.282	
treatment3 - treatment5	-46.25	20.72	6	-2.232	0.067	
treatment3 - treatment6	-135.00	20.72	6	-6.515	0.001	***
treatment3 - treatment7	-98.75	20.72	6	-4.766	0.003	**
treatment3 - treatment8	34.00	20.72	6	1.641	0.152	
treatment3 - treatment9	-76.25	20.72	6	-3.680	0.010	*
treatment3 - treatment10	-190.00	20.72	6	-9.169	0.000	***

*	0.047	-2.497	6	-51.75 20.72	treatment3 - treatment11
	0.097	-1.967	6	-40.75 20.72	treatment3 - treatment12
	0.199	-1.444	6	-29.92 20.72	treatment4 - treatment5
**	0.001	-5.727	6	-118.67 20.72	treatment4 - treatment6
**	0.007	-3.977	6	-82.42 20.72	treatment4 - treatment7
	0.051	2.429	6	50.33 20.72	treatment4 - treatment8
*	0.028	-2.892	6	-59.92 20.72	treatment4 - treatment9
***	0.000	-8.381	6	-173.67 20.72	treatment4 - treatment10
	0.138	-1.709	6	-35.42 20.72	treatment4 - treatment11
	0.283	-1.178	6	-24.42 20.72	treatment4 - treatment12
*	0.016	-3.318	6	-88.75 26.75	treatment5 - treatment6
	0.097	-1.963	6	-52.50 26.75	treatment5 - treatment7
*	0.024	3.000	6	80.25 26.75	treatment5 - treatment8
	0.257	-1.254	6	-30.00 23.93	treatment5 - treatment9
**	0.002	-5.374	6	-143.75 26.75	treatment5 - treatment10
	0.844	-0.206	6	-5.50 26.75	treatment5 - treatment11
	0.844	0.206	6	5.50 26.75	treatment5 - treatment12
	0.224	1.355	6	36.25 26.75	treatment6 - treatment7
***	0.000	7.063	6	169.00 23.93	treatment6 - treatment8
	0.070	2.196	6	58.75 26.75	treatment6 - treatment9
	0.061	-2.299	6	-55.00 23.93	treatment6 - treatment10
*	0.021	3.112	6	83.25 26.75	treatment6 - treatment11
*	0.012	3.523	6	94.25 26.75	treatment6 - treatment12
**	0.003	4.962	6	132.75 26.75	treatment7 - treatment8
	0.433	0.841	6	22.50 26.75	treatment7 - treatment9
*	0.014	-3.411	6	-91.25 26.75	treatment7 - treatment10
	0.097	1.964	6	47.00 23.93	treatment7 - treatment11
	0.052	2.424	6	58.00 23.93	treatment7 - treatment12
**	0.006	-4.121	6	-110.25 26.75	treatment8 - treatment9
***	0.000	-9.362	6	-224.00 23.93	treatment8 - treatment10
*	0.018	-3.205	6	-85.75 26.75	treatment8 - treatment11
*	0.031	-2.794	6	-74.75 26.75	treatment8 - treatment12
**	0.005	-4.252	6	-113.75 26.75	treatment9 - treatment10
	0.395	0.916	6	24.50 26.75	treatment9 - treatment11
	0.233	1.327	6	35.50 26.75	treatment9 - treatment12
**	0.002	5.168	6	138.25 26.75	treatment10 - treatment11
**	0.001	5.579	6	149.25 26.75	treatment10 - treatment12
	0.662	0.460	6	11.00 23.93	treatment11 - treatment12

Treatment Groups

Method : lsd

Treatment	Adjusted Means	SE	df	lower.CL	upper.CL	Group
8	213.67	18.27	6	168.95	258.38	12
2	228.00	9.77	6	204.10	251.90	1
3	247.67	9.77	6	223.76	271.57	123
1	256.00	9.77	6	232.10	279.90	1234
4	264.00	9.77	6	240.10	287.90	234
12	288.42	18.27	6	243.70	333.13	345
5	293.92	18.27	6	249.20	338.63	345

```
11
                 299.42 18.27 6 254.70
                                         344.13
                                                    45
                 323.92 18.27 6 279.20
        9
                                         368.63
                                                     56
        7
                 346.42 18.27 6
                                301.70
                                          391.13
                                                     56
                 382.67 18.27 6 337.95
        6
                                         427.38
                                                      67
                 437.67 18.27 6 392.95
       10
                                         482.38
class(out2)
```

[1] "augmentedRCBD"

The data can also be imported as a data frame and then used for analysis. Consider the data frame data imported from Table 1 according to the instructions in section 4.8.

str(data)

Augmented Design Details

```
Number of blocks "3"

Number of treatments "12"

Number of check treatments "4"

Number of test treatments "8"

Check treatments "1, 2, 3, 4"
```

ANOVA, Treatment Adjusted

```
Df Sum Sq Mean Sq F value Pr(>F)
Block (ignoring Treatments)
                                  2 360.1 180.04 6.675 0.0298 *
Treatment (eliminating Blocks)
                                 11 285.1
                                           25.92
                                                    0.961 0.5499
  Treatment: Check
                                  3
                                     52.9
                                           17.64
                                                    0.654 0.6092
  Treatment: Test and Test vs. Check 8 232.2 29.02
                                                    1.076 0.4779
                                   6 161.8
Residuals
                                           26.97
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
ANOVA, Block Adjusted
______
```

Df Sum Sq Mean Sq F value Pr(>F)
Treatment (ignoring Blocks) 11 575.7 52.33 1.940 0.215

Treatment: Check	3	52.9	17.64	0.654	0.609
Treatment: Test	7	505.9	72.27	2.679	0.125
Treatment: Test vs. Check	1	16.9	16.87	0.626	0.459
Block (eliminating Treatments)	2	69.5	34.75	1.288	0.342
Residuals	6	161.8	26.97		

Coefficient of Variation

6.372367

Overall Adjusted Mean

81.0625

Standard Errors

==========

	Std.	Error	of Diff.	CD (5%)
Control Treatment Means			4.240458	10.37603
Two Test Treatments (Same Block)			7.344688	17.97180
Two Test Treatments (Different Blocks)			8.211611	20.09309
A Test Treatment and a Control Treatment			6.704752	16.40594

Treatment Means

Treatment	Block	Means	SE	r	Min	Max	Adjusted	Means
1		84.67	3.84	3	79.00	92.00		84.67
10	3	74.00	<na></na>	1	74.00	74.00		77.25
11	1	89.00	<na></na>	1	89.00	89.00		86.50
12	1	82.00	<na></na>	1	82.00	82.00		79.50
2		79.00	1.15	3	77.00	81.00		79.00
3		82.00	2.65	3	78.00	87.00		82.00
4		83.33	3.93	3	78.00	91.00		83.33
5	2	79.00	<na></na>	1	79.00	79.00		78.25
6	3	75.00	<na></na>	1	75.00	75.00		78.25
7	1	96.00	<na></na>	1	96.00	96.00		93.50
8	3	70.00	<na></na>	1	70.00	70.00		73.25
9	2	78.00	<na></na>	1	78.00	78.00		77.25

${\tt Comparisons}$

========

Method : 1sd

```
contrast estimate SE df t.ratio p.value sig
treatment1 - treatment2 5.67 4.24 6 1.336 0.230
treatment1 - treatment3
                        2.67 4.24 6 0.629
                                             0.553
                       1.33 4.24 6 0.314
                                            0.764
treatment1 - treatment4
                     6.42 6.36 6 1.009
treatment1 - treatment5
                                            0.352
treatment1 - treatment6
                       6.42 6.36 6 1.009 0.352
treatment1 - treatment7 -8.83 6.36 6 -1.389
                                             0.214
                       11.42 6.36 6 1.795
treatment1 - treatment8
                                             0.123
```

```
treatment1 - treatment9
                            7.42 6.36 6
                                          1.166
                                                   0.288
treatment1 - treatment10
                            7.42 6.36 6
                                          1.166
                                                   0.288
                           -1.83 6.36 6 -0.288
treatment1 - treatment11
                                                   0.783
                            5.17 6.36 6
                                          0.812
treatment1 - treatment12
                                                   0.448
treatment2 - treatment3
                           -3.004.246-0.707
                                                   0.506
treatment2 - treatment4
                           -4.33 4.24 6 -1.022
                                                   0.346
treatment2 - treatment5
                            0.75 6.36 6
                                           0.118
                                                   0.910
treatment2 - treatment6
                            0.75 6.36 6
                                           0.118
                                                   0.910
treatment2 - treatment7
                          -14.50 6.36 6 -2.280
                                                   0.063
treatment2 - treatment8
                            5.75 6.36 6
                                          0.904
                                                   0.401
treatment2 - treatment9
                            1.75 6.36 6
                                          0.275
                                                   0.792
treatment2 - treatment10
                            1.75 6.36 6
                                          0.275
                                                   0.792
treatment2 - treatment11
                           -7.50 6.36 6 -1.179
                                                   0.283
treatment2 - treatment12
                           -0.50 6.36 6 -0.079
                                                   0.940
                           -1.33 4.24 6 -0.314
treatment3 - treatment4
                                                   0.764
treatment3 - treatment5
                            3.75 6.36 6
                                         0.590
                                                   0.577
treatment3 - treatment6
                            3.75 6.36 6
                                           0.590
                                                   0.577
treatment3 - treatment7
                          -11.50 6.36 6 -1.808
                                                   0.121
                            8.75 6.36 6
                                          1.376
treatment3 - treatment8
                                                   0.218
treatment3 - treatment9
                            4.75 6.36 6
                                          0.747
                                                   0.483
treatment3 - treatment10
                            4.75 6.36 6
                                         0.747
                                                   0.483
treatment3 - treatment11
                           -4.50 6.36 6 -0.707
                                                   0.506
treatment3 - treatment12
                            2.50 6.36 6
                                         0.393
                                                   0.708
treatment4 - treatment5
                            5.08 6.36 6
                                           0.799
                                                   0.455
                            5.08 6.36 6
                                          0.799
                                                   0.455
treatment4 - treatment6
treatment4 - treatment7
                          -10.17 6.36 6 -1.598
                                                   0.161
                           10.08 6.36 6
treatment4 - treatment8
                                         1.585
                                                   0.164
treatment4 - treatment9
                            6.08 6.36 6
                                          0.956
                                                   0.376
treatment4 - treatment10
                            6.08 6.36 6
                                          0.956
                                                   0.376
                           -3.17 6.36 6 -0.498
treatment4 - treatment11
                                                   0.636
treatment4 - treatment12
                            3.83 6.36 6
                                           0.603
                                                   0.569
treatment5 - treatment6
                            0.00 8.21 6
                                          0.000
                                                   1.000
treatment5 - treatment7
                          -15.25 8.21
                                      6 -1.857
                                                   0.113
treatment5 - treatment8
                            5.00 8.21 6
                                          0.609
                                                   0.565
treatment5 - treatment9
                            1.00 7.34 6
                                          0.136
                                                   0.896
treatment5 - treatment10
                            1.00 8.21 6
                                         0.122
                                                   0.907
                           -8.25 8.21
                                      6 -1.005
treatment5 - treatment11
                                                   0.354
treatment5 - treatment12
                           -1.25 8.21 6 -0.152
                                                   0.884
treatment6 - treatment7
                          -15.25 8.21
                                       6 - 1.857
                                                   0.113
treatment6 - treatment8
                            5.00 7.34 6
                                          0.681
                                                   0.521
treatment6 - treatment9
                            1.00 8.21
                                       6
                                           0.122
                                                   0.907
                            1.00 7.34 6
                                          0.136
                                                   0.896
treatment6 - treatment10
treatment6 - treatment11
                           -8.25 8.21
                                       6 -1.005
                                                   0.354
treatment6 - treatment12
                           -1.25 8.21 6 -0.152
                                                   0.884
treatment7 - treatment8
                           20.25 8.21 6
                                          2.466
                                                   0.049
treatment7 - treatment9
                           16.25 8.21 6
                                         1.979
                                                   0.095
treatment7 - treatment10
                           16.25 8.21 6
                                         1.979
                                                   0.095
treatment7 - treatment11
                            7.00 7.34 6
                                           0.953
                                                   0.377
                                          1.906
                           14.00 7.34 6
treatment7 - treatment12
                                                   0.105
treatment8 - treatment9
                           -4.00 8.21
                                       6 - 0.487
                                                   0.643
                           -4.00 7.34 6 -0.545
treatment8 - treatment10
                                                   0.606
treatment8 - treatment11
                          -13.25 8.21 6 -1.614
                                                   0.158
```

```
treatment8 - treatment12
                          -6.25 8.21 6 -0.761
                                                0.475
treatment9 - treatment10
                          0.00 8.21 6 0.000
                                                1.000
 treatment9 - treatment11
                          -9.25 8.21 6 -1.126
                                                0.303
 treatment9 - treatment12 -2.25 8.21 6 -0.274
                                                0.793
treatment10 - treatment11 -9.25 8.21 6 -1.126
                                                0.303
treatment10 - treatment12
                          -2.25 8.21 6 -0.274
                                                0.793
                          7.00 7.34 6 0.953
                                               0.377
treatment11 - treatment12
```

Treatment Groups

Method : 1sd

Treatment	Adjusted	Means	SE	df	lower.CL	upper.CL	Group
8		73.25	5.61	6	59.52	86.98	1
9		77.25	5.61	6	63.52	90.98	12
10		77.25			63.52	90.98	12
5		78.25	5.61	6	64.52	91.98	12
6		78.25	5.61	6	64.52	91.98	12
2		79.00	3.00	6	71.66	86.34	12
12		79.50	5.61	6	65.77	93.23	12
3		82.00	3.00	6	74.66	89.34	12
4		83.33	3.00	6	76.00	90.67	12
1		84.67	3.00	6	77.33	92.00	12
11		86.50	5.61	6	72.77	100.23	12
7		93.50	5.61	6	79.77	107.23	2

class (out1)

[1] "augmentedRCBD"

Augmented Design Details

```
Number of blocks "3"

Number of treatments "12"

Number of check treatments "4"

Number of test treatments "8"

Check treatments "1, 2, 3, 4"
```

ANOVA, Treatment Adjusted

```
Df Sum Sq Mean Sq F value Pr(>F)

Block (ignoring Treatments) 2 7019 3510 12.261 0.007597 **

Treatment (eliminating Blocks) 11 58965 5360 18.727 0.000920 ***

Treatment: Check 3 2150 717 2.504 0.156116

Treatment: Test and Test vs. Check 8 56815 7102 24.810 0.000473 ***
```

Residuals 6 1718 286

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

ANOVA, Block Adjusted

Df Sum Sq Mean Sq F value Pr(>F)
Treatment (ignoring Blocks) 11 64708 5883 20.550 0.000707 ***
Treatment: Check 3 2150 717 2.504 0.156116
Treatment: Test 7 34863 4980 17.399 0.001366 **
Treatment: Test vs. Check 1 27694 27694 96.749 6.36e-05 ***
Block (eliminating Treatments) 2 1277 639 2.231 0.188645
Residuals 6 1717 286

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

Coefficient of Variation

6.057617

Overall Adjusted Mean

298.4792

Standard Errors

Std. Error of Diff. CD (5%)
Control Treatment Means 13.81424 33.80224
Two Test Treatments (Same Block) 23.92697 58.54719
Two Test Treatments (Different Blocks) 26.75117 65.45775
A Test Treatment and a Control Treatment 21.84224 53.44603

Treatment Means

Treatment	${\tt Block}$	Means	SE	r	Min	Max	Adjusted Means
1		256.00	3.06	3	250.00	260.00	256.00
10	3	450.00	<na></na>	1	450.00	450.00	437.67
11	1	300.00	<na></na>	1	300.00	300.00	299.42
12	1	289.00	<na></na>	1	289.00	289.00	288.42
2		228.00	6.11	3	220.00	240.00	228.00
3		247.67	10.17	3	237.00	268.00	247.67
4		264.00	18.68	3	227.00	287.00	264.00
5	2	281.00	<na></na>	1	281.00	281.00	293.92
6	3	395.00	<na></na>	1	395.00	395.00	382.67
7	1	347.00	<na></na>	1	347.00	347.00	346.42
8	3	226.00	<na></na>	1	226.00	226.00	213.67
9	2	311.00	<na></na>	1	311.00	311.00	323.92

Comparisons

========

Method : 1sd

gontragt	estimate	C E	4£	++i-	p.value	ai a
treatment1 - treatment2		13.81	6	2.027	0.089	sig
treatment1 - treatment3		13.81	6	0.603	0.568	
treatment1 - treatment4		13.81	6	-0.579		
treatment1 - treatment5	-37.92		6	-1.830		
treatment1 - treatment6			6	-6.113		***
treatment1 - treatment7			6	-4.363		**
treatment1 - treatment8		20.72	6	2.043		
treatment1 - treatment9			6	-3.278		*
treatment1 - treatment10			6	-8.767		
treatment1 - treatment11			6	-2.095	0.081	
treatment1 - treatment12			6	-1.564		
treatment2 - treatment3			6	-1.424		
treatment2 - treatment4			6	-2.606		*
treatment2 - treatment5	-65.92		6	-3.181		*
treatment2 - treatment6			6	-7.464		***
treatment2 - treatment7			6	-5.715		**
treatment2 - treatment8		20.72	6	0.692		
treatment2 - treatment9			6	-4.629		**
treatment2 - treatment10				-10.118		
treatment2 - treatment11			6	-3.447		*
treatment2 - treatment12	-60.42		6	-2.916		*
treatment3 - treatment4			6	-1.182		
treatment3 - treatment5	-46.25		6	-2.232		
treatment3 - treatment6			6	-6.515		***
treatment3 - treatment7	-98.75		6	-4.766		**
treatment3 - treatment8		20.72	6	1.641		
treatment3 - treatment9	-76.25		6	-3.680		*
treatment3 - treatment10			6	-9.169	0.010	
treatment3 - treatment11			6	-2.497		*
treatment3 - treatment12	-31.75 -40.75		6	-1.967		•
treatment3 - treatment12	-40.75 -29.92		6	-1.444	0.097	
treatment4 - treatment6			6	-5.727		**
treatment4 - treatment7			6	-3.727		
treatment4 - treatment7 treatment4 - treatment8		20.72	6	2.429	0.007	~ ~
treatment4 - treatment9			6	-2.892		*
treatment4 - treatment10			6		0.028	
				-8.381		^^^
<pre>treatment4 - treatment11 treatment4 - treatment12</pre>	-35.42		6	-1.709	0.138	
treatment4 - treatment12 treatment5 - treatment6				-1.178	0.283	*
treatment5 - treatment7				-3.318		•
treatment5 - treatment7 treatment5 - treatment8				-1.963		*
				3.000		^
treatment5 - treatment9				-1.254		**
treatment5 - treatment10				-5.374		^^
treatment5 - treatment11		26.75	6	-0.206		
treatment5 - treatment12		26.75		0.206		
treatment6 - treatment7		26.75	6	1.355		***
treatment6 - treatment8				7.063		***
treatment6 - treatment9				2.196		
treatment6 - treatment10	-55.00	∠3.93	6	-2.299	0.061	

```
treatment6 - treatment11
                         83.25 26.75 6
                                        3.112
                                               0.021
treatment6 - treatment12
                         94.25 26.75 6
                                        3.523
                                               0.012
 treatment7 - treatment8 132.75 26.75 6
                                        4.962
                                               0.003
                                                     **
                         22.50 26.75 6
 treatment7 - treatment9
                                        0.841
                                               0.433
treatment7 - treatment10 -91.25 26.75 6 -3.411
                                               0.014
treatment7 - treatment11 47.00 23.93 6
                                        1.964
                                               0.097
treatment7 - treatment12
                         58.00 23.93 6
                                        2.424
                                               0.052
 treatment8 - treatment9 -110.25 26.75 6 -4.121
                                               0.006
treatment8 - treatment10 -224.00 23.93 6 -9.362
                                               0.000 ***
treatment8 - treatment11 -85.75 26.75 6 -3.205
                                               0.018
treatment8 - treatment12 -74.75 26.75 6 -2.794
                                               0.031
treatment9 - treatment10 -113.75 26.75 6 -4.252
                                               0.005 **
treatment9 - treatment11 24.50 26.75 6 0.916
                                               0.395
treatment9 - treatment12 35.50 26.75 6 1.327
                                               0.233
treatment10 - treatment11 138.25 26.75 6 5.168
                                               0.002 **
treatment10 - treatment12 149.25 26.75 6 5.579
                                               0.001 **
0.662
```

Treatment Groups

Method : 1sd

Treatment	Adjusted Means	SE	df	lower.CL	upper.CL	Group
8	213.67	18.27	6	168.95	258.38	12
2	228.00	9.77	6	204.10	251.90	1
3	247.67	9.77	6	223.76	271.57	123
1	256.00	9.77	6	232.10	279.90	1234
4	264.00	9.77	6	240.10	287.90	234
12	288.42	18.27	6	243.70	333.13	345
5	293.92	18.27	6	249.20	338.63	345
11	299.42	18.27	6	254.70	344.13	45
9	323.92	18.27	6	279.20	368.63	56
7	346.42	18.27	6	301.70	391.13	56
6	382.67	18.27	6	337.95	427.38	67
10	437.67	18.27	6	392.95	482.38	7

[1] "augmentedRCBD"

class (out2)

Check genotypes are inferred by default on the basis of number of replications. However, if some test genotypes are also replicated, they may also be falsely detected as checks. To avoid this, the checks can be specified by the **checks** argument.

```
Augmented Design Details
```

```
Number of blocks "3"

Number of treatments "12"

Number of check treatments "4"

Number of test treatments "8"

Check treatments "1, 2, 3, 4"
```

ANOVA, Treatment Adjusted

	Df	Sum Sq	Mean Sq	F value	Pr(>F)	
Block (ignoring Treatments)	2	360.1	180.04	6.675	0.0298	*
Treatment (eliminating Blocks)	11	285.1	25.92	0.961	0.5499	
Treatment: Check	3	52.9	17.64	0.654	0.6092	
Treatment: Test and Test vs. Check	8	232.2	29.02	1.076	0.4779	
Residuals	6	161.8	26.97			

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

ANOVA, Block Adjusted

	Df	Sum Sq	Mean Sq F	value	Pr(>F)
Treatment (ignoring Blocks)	11	575.7	52.33	1.940	0.215
Treatment: Check	3	52.9	17.64	0.654	0.609
Treatment: Test	7	505.9	72.27	2.679	0.125
Treatment: Test vs. Check	1	16.9	16.87	0.626	0.459
<pre>Block (eliminating Treatments)</pre>	2	69.5	34.75	1.288	0.342
Residuals	6	161.8	26.97		

Coefficient of Variation

6.372367

Overall Adjusted Mean

81.0625

Standard Errors

	Sta. Error	of Diff. CD (5%)
Control Treatment Means		4.240458 10.37603
Two Test Treatments (Same Block)		7.344688 17.97180
Two Test Treatments (Different Blocks)		8.211611 20.09309
A Test Treatment and a Control Treatment		6.704752 16.40594

Treatment Means

==========

Treatment	Block	Means	SE	r	Min	Max	${\tt Adjusted}$	Means
1		84.67	3.84	3	79.00	92.00		84.67
10	3	74.00	<na></na>	1	74.00	74.00		77.25
11	1	89.00	<na></na>	1	89.00	89.00		86.50
12	1	82.00	<na></na>	1	82.00	82.00		79.50
2		79.00	1.15	3	77.00	81.00		79.00

3		82.00	2.65	3	78.00	87.00	82.00
4		83.33	3.93	3	78.00	91.00	83.33
5	2	79.00	<na></na>	1	79.00	79.00	78.25
6	3	75.00	<na></na>	1	75.00	75.00	78.25
7	1	96.00	<na></na>	1	96.00	96.00	93.50
8	3	70.00	<na></na>	1	70.00	70.00	73.25
9	2	78.00	<na></na>	1	78.00	78.00	77.25

Comparisons

========

Method : lsd

contrast	estimate	SE	df	t.ratio	p.value sig
treatment1 - treatment2	5.67	4.24	6	1.336	0.230
treatment1 - treatment3	2.67	4.24	6	0.629	0.553
treatment1 - treatment4	1.33	4.24	6	0.314	0.764
treatment1 - treatment5	6.42	6.36	6	1.009	0.352
treatment1 - treatment6	6.42	6.36	6	1.009	0.352
treatment1 - treatment7	-8.83	6.36	6	-1.389	0.214
treatment1 - treatment8	11.42	6.36	6	1.795	0.123
treatment1 - treatment9	7.42	6.36	6	1.166	0.288
treatment1 - treatment10	7.42	6.36	6	1.166	0.288
treatment1 - treatment11	-1.83	6.36	6	-0.288	0.783
treatment1 - treatment12	5.17	6.36	6	0.812	0.448
treatment2 - treatment3	-3.00	4.24	6	-0.707	0.506
treatment2 - treatment4	-4.33	4.24	6	-1.022	0.346
treatment2 - treatment5	0.75	6.36	6	0.118	0.910
treatment2 - treatment6	0.75	6.36	6	0.118	0.910
treatment2 - treatment7	-14.50	6.36	6	-2.280	0.063
treatment2 - treatment8	5.75	6.36	6	0.904	0.401
treatment2 - treatment9	1.75	6.36	6	0.275	0.792
treatment2 - treatment10	1.75	6.36	6	0.275	0.792
treatment2 - treatment11	-7.50	6.36	6	-1.179	0.283
treatment2 - treatment12	-0.50	6.36	6	-0.079	0.940
treatment3 - treatment4	-1.33	4.24	6	-0.314	0.764
treatment3 - treatment5	3.75	6.36	6	0.590	0.577
treatment3 - treatment6	3.75	6.36	6	0.590	0.577
treatment3 - treatment7	-11.50	6.36	6	-1.808	0.121
treatment3 - treatment8	8.75	6.36	6	1.376	0.218
treatment3 - treatment9	4.75	6.36	6	0.747	0.483
treatment3 - treatment10	4.75	6.36	6	0.747	0.483
treatment3 - treatment11	-4.50	6.36	6	-0.707	0.506
treatment3 - treatment12	2.50	6.36	6	0.393	0.708
treatment4 - treatment5	5.08	6.36	6	0.799	0.455
treatment4 - treatment6	5.08	6.36	6	0.799	0.455
treatment4 - treatment7	-10.17	6.36	6	-1.598	0.161
treatment4 - treatment8	10.08	6.36	6	1.585	0.164
treatment4 - treatment9	6.08	6.36	6	0.956	0.376
treatment4 - treatment10	6.08	6.36	6	0.956	0.376
treatment4 - treatment11	-3.17	6.36	6	-0.498	0.636

```
treatment4 - treatment12
                           3.83 6.36 6 0.603
                                                 0.569
 treatment5 - treatment6
                           0.00 8.21 6 0.000
                                                 1.000
 treatment5 - treatment7
                         -15.25 8.21 6 -1.857
                                                 0.113
 treatment5 - treatment8
                           5.00 8.21 6 0.609
                                                 0.565
 treatment5 - treatment9
                          1.00 7.34 6 0.136
                                                 0.896
                          1.00 8.21 6 0.122
 treatment5 - treatment10
                                                 0.907
 treatment5 - treatment11
                          -8.25 8.21 6 -1.005
                                                 0.354
 treatment5 - treatment12 -1.25 8.21 6 -0.152
                                                 0.884
 treatment6 - treatment7 -15.25 8.21 6 -1.857
                                                 0.113
                           5.00 7.34 6 0.681
 treatment6 - treatment8
                                                 0.521
 treatment6 - treatment9
                          1.00 8.21 6
                                        0.122
                                                 0.907
 treatment6 - treatment10
                          1.00 7.34 6 0.136
                                                 0.896
 treatment6 - treatment11
                          -8.25 8.21 6 -1.005
                                                 0.354
 treatment6 - treatment12
                          -1.25 8.21 6 -0.152
                                                 0.884
                          20.25 8.21 6 2.466
                                                 0.049
 treatment7 - treatment8
 treatment7 - treatment9
                          16.25 8.21 6 1.979
                                                 0.095
                          16.25 8.21 6 1.979
 treatment7 - treatment10
                                                 0.095
 treatment7 - treatment11
                           7.00 7.34 6 0.953
                                                 0.377
 treatment7 - treatment12
                          14.00 7.34 6 1.906
                                                 0.105
                          -4.00 8.21 6 -0.487
 treatment8 - treatment9
                                                 0.643
 treatment8 - treatment10
                          -4.00 7.34 6 -0.545
                                                 0.606
                          -13.25 8.21 6 -1.614
 treatment8 - treatment11
                                                 0.158
 treatment8 - treatment12
                          -6.25 \ 8.21 \ 6 \ -0.761
                                                 0.475
 treatment9 - treatment10
                          0.00 8.21 6 0.000
                                                 1.000
 treatment9 - treatment11
                          -9.25 8.21 6 -1.126
                                                 0.303
 treatment9 - treatment12
                          -2.25 8.21 6 -0.274
                                                 0.793
treatment10 - treatment11 -9.25 8.21 6 -1.126
                                                 0.303
treatment10 - treatment12
                          -2.25 8.21 6 -0.274
                                                 0.793
treatment11 - treatment12
                          7.00 7.34 6 0.953
                                                 0.377
```

Treatment Groups

Method : 1sd

```
Treatment Adjusted Means
                         SE df lower.CL upper.CL Group
                  73.25 5.61 6
                                  59.52
                                          86.98
       8
       9
                  77.25 5.61 6
                                  63.52
                                          90.98
                                                   12
      10
                 77.25 5.61 6
                                  63.52
                                          90.98
                                                   12
                                                   12
       5
                 78.25 5.61 6
                                  64.52
                                          91.98
       6
                 78.25 5.61 6
                                 64.52
                                          91.98
                                                   12
       2
                 79.00 3.00 6
                               71.66 86.34
                                                   12
                 79.50 5.61 6
                                 65.77 93.23
      12
                                                  12
                                        89.34
       3
                 82.00 3.00 6
                                  74.66
                                                   12
       4
                  83.33 3.00 6
                                 76.00
                                          90.67
                                                  12
                  84.67 3.00 6
       1
                                  77.33
                                          92.00
                                                   12
      11
                  86.50 5.61 6
                                  72.77
                                         100.23
                                                   12
                  93.50 5.61 6
                                  79.77
                                         107.23
```

```
checks = c("1", "2", "3", "4"))
```

Augmented Design Details

Number of blocks "3"
Number of treatments "12"
Number of check treatments "4"
Number of test treatments "8"

Check treatments "1, 2, 3, 4"

ANOVA, Treatment Adjusted

	Df	Sum Sq	Mean Sq	F value	Pr(>F)	
Block (ignoring Treatments)	2	7019	3510	12.261	0.007597	**
Treatment (eliminating Blocks)	11	58965	5360	18.727	0.000920	***
Treatment: Check	3	2150	717	2.504	0.156116	
Treatment: Test and Test vs. Check	8	56815	7102	24.810	0.000473	***
Residuals	6	1718	286			

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

ANOVA, Block Adjusted

	Dİ	Sum Sq	Mean Sq	F value	Pr(>F)	
Treatment (ignoring Blocks)	11	64708	5883	20.550	0.000707	***
Treatment: Check	3	2150	717	2.504	0.156116	
Treatment: Test	7	34863	4980	17.399	0.001366	**
Treatment: Test vs. Check	1	27694	27694	96.749	6.36e-05	***
<pre>Block (eliminating Treatments)</pre>	2	1277	639	2.231	0.188645	
Residuals	6	1717	286			

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

Coefficient of Variation

6.057617

Overall Adjusted Mean

298.4792

Standard Errors

	Std.	Error	of Diff.	CD (5%)
Control Treatment Means			13.81424	33.80224
Two Test Treatments (Same Block)			23.92697	58.54719
Two Test Treatments (Different Blocks)			26.75117	65.45775
A Test Treatment and a Control Treatment			21.84224	53.44603

Treatment Means

Treatment	Block	Means	SE	r	Min	Max	Adjusted Means
1		256.00	3.06	3	250.00	260.00	256.00
10	3	450.00	<na></na>	1	450.00	450.00	437.67
11	1	300.00	<na></na>	1	300.00	300.00	299.42
12	1	289.00	<na></na>	1	289.00	289.00	288.42
2		228.00	6.11	3	220.00	240.00	228.00
3		247.67	10.17	3	237.00	268.00	247.67
4		264.00	18.68	3	227.00	287.00	264.00
5	2	281.00	<na></na>	1	281.00	281.00	293.92
6	3	395.00	<na></na>	1	395.00	395.00	382.67
7	1	347.00	<na></na>	1	347.00	347.00	346.42
8	3	226.00	<na></na>	1	226.00	226.00	213.67
9	2	311.00	<na></na>	1	311.00	311.00	323.92

Comparisons

========

Method : 1sd

contrast	${\tt estimate}$	SE	df	t.ratio	${\tt p.value}$	sig
treatment1 - treatment2	28.00	13.81	6	2.027	0.089	
treatment1 - treatment3	8.33	13.81	6	0.603	0.568	
treatment1 - treatment4	-8.00	13.81	6	-0.579	0.584	
treatment1 - treatment5	-37.92	20.72	6	-1.830	0.117	
treatment1 - treatment6	-126.67	20.72	6	-6.113	0.001	***
treatment1 - treatment7	-90.42	20.72	6	-4.363	0.005	**
treatment1 - treatment8	42.33	20.72	6	2.043	0.087	
treatment1 - treatment9	-67.92	20.72	6	-3.278	0.017	*
<pre>treatment1 - treatment10</pre>	-181.67	20.72	6	-8.767	0.000	***
<pre>treatment1 - treatment11</pre>	-43.42	20.72	6	-2.095	0.081	
<pre>treatment1 - treatment12</pre>	-32.42	20.72	6	-1.564	0.169	
treatment2 - treatment3	-19.67	13.81	6	-1.424	0.204	
treatment2 - treatment4	-36.00	13.81	6	-2.606	0.040	*
treatment2 - treatment5	-65.92	20.72	6	-3.181	0.019	*
treatment2 - treatment6	-154.67	20.72	6	-7.464	0.000	***
treatment2 - treatment7	-118.42	20.72	6	-5.715	0.001	**
treatment2 - treatment8	14.33	20.72	6	0.692	0.515	
treatment2 - treatment9	-95.92	20.72	6	-4.629	0.004	**
treatment2 - treatment10	-209.67	20.72	6	-10.118	0.000	***
<pre>treatment2 - treatment11</pre>	-71.42	20.72	6	-3.447	0.014	*
treatment2 - treatment12	-60.42	20.72	6	-2.916	0.027	*
treatment3 - treatment4	-16.33	13.81	6	-1.182	0.282	
treatment3 - treatment5	-46.25	20.72	6	-2.232	0.067	
treatment3 - treatment6	-135.00	20.72	6	-6.515	0.001	***
treatment3 - treatment7	-98.75	20.72	6	-4.766	0.003	**
treatment3 - treatment8	34.00	20.72	6	1.641	0.152	
treatment3 - treatment9	-76.25	20.72	6	-3.680	0.010	*
treatment3 - treatment10	-190.00	20.72	6	-9.169	0.000	***
<pre>treatment3 - treatment11</pre>	-51.75	20.72	6	-2.497	0.047	*

	0.097	-1.967	6	20.72	-40.75	treatment3 - treatment12
	0.199	-1.444	6	20.72	-29.92	treatment4 - treatment5
**	0.001	-5.727	6	20.72	-118.67	treatment4 - treatment6
**	0.007	-3.977	6	20.72	-82.42	treatment4 - treatment7
	0.051	2.429	6	20.72	50.33	treatment4 - treatment8
*	0.028	-2.892	6	20.72	-59.92	treatment4 - treatment9
***	0.000	-8.381	6	20.72	-173.67	treatment4 - treatment10
	0.138	-1.709	6	20.72	-35.42	treatment4 - treatment11
	0.283	-1.178	6	20.72	-24.42	treatment4 - treatment12
*	0.016	-3.318	6	26.75	-88.75	treatment5 - treatment6
	0.097	-1.963	6	26.75	-52.50	treatment5 - treatment7
*	0.024	3.000	6	26.75	80.25	treatment5 - treatment8
	0.257	-1.254	6	23.93	-30.00	treatment5 - treatment9
**	0.002	-5.374	6	26.75	-143.75	treatment5 - treatment10
	0.844	-0.206	6	26.75	-5.50	treatment5 - treatment11
	0.844	0.206	6	26.75	5.50	treatment5 - treatment12
	0.224	1.355	6	26.75	36.25	treatment6 - treatment7
***	0.000	7.063	6	23.93	169.00	treatment6 - treatment8
	0.070	2.196	6	26.75	58.75	treatment6 - treatment9
	0.061	-2.299	6	23.93	-55.00	treatment6 - treatment10
*	0.021	3.112	6	26.75	83.25	treatment6 - treatment11
*	0.012	3.523	6	26.75	94.25	treatment6 - treatment12
**	0.003	4.962	6	26.75	132.75	treatment7 - treatment8
	0.433	0.841	6	26.75	22.50	treatment7 - treatment9
*	0.014	-3.411	6	26.75	-91.25	treatment7 - treatment10
	0.097	1.964	6	23.93	47.00	treatment7 - treatment11
	0.052	2.424	6	23.93	58.00	treatment7 - treatment12
**	0.006	-4.121	6	26.75	-110.25	treatment8 - treatment9
***	0.000	-9.362	6	23.93	-224.00	treatment8 - treatment10
*	0.018	-3.205	6	26.75	-85.75	treatment8 - treatment11
*	0.031	-2.794	6	26.75	-74.75	treatment8 - treatment12
**	0.005	-4.252	6	26.75	-113.75	treatment9 - treatment10
	0.395	0.916	6	26.75	24.50	treatment9 - treatment11
	0.233	1.327	6	26.75	35.50	treatment9 - treatment12
**	0.002	5.168	6	26.75	138.25	<pre>treatment10 - treatment11</pre>
**	0.001	5.579	6	26.75	149.25	<pre>treatment10 - treatment12</pre>
	0.662	0.460	6	23.93	11.00	<pre>treatment11 - treatment12</pre>

Treatment Groups

Method : 1sd

Treatment Adjusted Means SE df lower.CL upper.CL Group 8 213.67 18.27 6 168.95 258.38 12 2 228.00 9.77 6 204.10 251.90 1 3 247.67 9.77 6 223.76 271.57 123 1 256.00 9.77 6 232.10 279.90 1234 4 264.00 9.77 6 240.10 287.90 234 288.42 18.27 6 243.70 12 333.13 345 5 293.92 18.27 6 249.20 338.63 345 11 299.42 18.27 6 254.70 344.13 45

9	323.92 18	.27 6	279.20	368.63	56
7	346.42 18	.27 6	301.70	391.13	56
6	382.67 18	.27 6	337.95	427.38	67
10	437.67 18	.27 6	392.95	482.38	7

In case the large number of treatments or genotypes, it is advisable to avoid treatment comparisons with the **group = FALSE** argument as it will be memory and processor intensive. Further it is advised to simplify output with **simplify = TRUE** in order to reduce output object size.

If truncate.means = TRUE, then any negative adjusted means will be truncated to zero with a warning.

7.2 print.augmentedRCBD()

The results of analysis in an object of class augmentedRCBD can be printed to the console as follows.

```
# Print results for variable y1
print(out1)
```

Augmented Design Details

```
Number of blocks "3"

Number of treatments "12"

Number of check treatments "4"

Number of test treatments "8"

Check treatments "1, 2, 3, 4"
```

ANOVA, Treatment Adjusted

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

ANOVA, Block Adjusted

```
Df Sum Sq Mean Sq F value Pr(>F)
Treatment (ignoring Blocks) 11 575.7 52.33 1.940 0.215
Treatment: Check 3 52.9 17.64 0.654 0.609
Treatment: Test 7 505.9 72.27 2.679 0.125
Treatment: Test vs. Check 1 16.9 16.87 0.626 0.459
Block (eliminating Treatments) 2 69.5 34.75 1.288 0.342
Residuals 6 161.8 26.97
```

```
Coefficient of Variation
```

6.372367

Overall Adjusted Mean

81.0625

Standard Errors

	Std.	Error	of Diff.	CD (5%)
Control Treatment Means			4.240458	10.37603
Two Test Treatments (Same Block)			7.344688	17.97180
Two Test Treatments (Different Blocks)			8.211611	20.09309
${\tt A}$ Test Treatment and a Control Treatment			6.704752	16.40594

Treatment Means

==========

Treatmen	nt	Block	Means	SE	r	Min	Max	Adjusted	Means
	1		84.67	3.84	3	79.00	92.00		84.67
:	10	3	74.00	<na></na>	1	74.00	74.00		77.25
:	11	1	89.00	<na></na>	1	89.00	89.00		86.50
:	12	1	82.00	<na></na>	1	82.00	82.00		79.50
	2		79.00	1.15	3	77.00	81.00		79.00
	3		82.00	2.65	3	78.00	87.00		82.00
	4		83.33	3.93	3	78.00	91.00		83.33
	5	2	79.00	<na></na>	1	79.00	79.00		78.25
	6	3	75.00	<na></na>	1	75.00	75.00		78.25
	7	1	96.00	<na></na>	1	96.00	96.00		93.50
	8	3	70.00	<na></na>	1	70.00	70.00		73.25
	9	2	78.00	<na></na>	1	78.00	78.00		77.25

Comparisons

Method : lsd

contrast	${\tt estimate}$	SE	df	t.ratio	${\tt p.value}$	sig
treatment1 - treatment2	5.67	4.24	6	1.336	0.230	
treatment1 - treatment3	2.67	4.24	6	0.629	0.553	
treatment1 - treatment4	1.33	4.24	6	0.314	0.764	
treatment1 - treatment5	6.42	6.36	6	1.009	0.352	
treatment1 - treatment6	6.42	6.36	6	1.009	0.352	
treatment1 - treatment7	-8.83	6.36	6	-1.389	0.214	
treatment1 - treatment8	11.42	6.36	6	1.795	0.123	
treatment1 - treatment9	7.42	6.36	6	1.166	0.288	
treatment1 - treatment10	7.42	6.36	6	1.166	0.288	
treatment1 - treatment11	-1.83	6.36	6	-0.288	0.783	
treatment1 - treatment12	5.17	6.36	6	0.812	0.448	
treatment2 - treatment3	-3.00	4.24	6	-0.707	0.506	
treatment2 - treatment4	-4.33	4.24	6	-1.022	0.346	
treatment2 - treatment5	0.75	6.36	6	0.118	0.910	
treatment2 - treatment6	0.75	6.36	6	0.118	0.910	
treatment2 - treatment7	-14.50	6.36	6	-2.280	0.063	

treatment2 - treatment8	5.75 6.36	6	0.904	0.401	
treatment2 - treatment9	1.75 6.36	6	0.275	0.792	
treatment2 - treatment10	1.75 6.36	6	0.275	0.792	
treatment2 - treatment11	-7.50 6.36	6	-1.179	0.283	
treatment2 - treatment12	-0.50 6.36	6	-0.079	0.940	
treatment3 - treatment4	-1.33 4.24	6	-0.314	0.764	
treatment3 - treatment5	3.75 6.36	6	0.590	0.577	
treatment3 - treatment6	3.75 6.36	6	0.590	0.577	
treatment3 - treatment7	-11.50 6.36	6	-1.808	0.121	
treatment3 - treatment8	8.75 6.36	6	1.376	0.218	
treatment3 - treatment9	4.75 6.36	6	0.747	0.483	
treatment3 - treatment10	4.75 6.36	6	0.747	0.483	
treatment3 - treatment11	-4.50 6.36	6	-0.707	0.506	
treatment3 - treatment12	2.50 6.36	6	0.393	0.708	
treatment4 - treatment5	5.08 6.36	6	0.799	0.455	
treatment4 - treatment6	5.08 6.36	6	0.799	0.455	
treatment4 - treatment7	-10.17 6.36	6	-1.598	0.161	
treatment4 - treatment8	10.08 6.36	6	1.585	0.164	
treatment4 - treatment9	6.08 6.36	6	0.956	0.376	
treatment4 - treatment10	6.08 6.36	6	0.956	0.376	
treatment4 - treatment11	-3.17 6.36	6	-0.498	0.636	
treatment4 - treatment12	3.83 6.36	6	0.603	0.569	
treatment5 - treatment6	0.00 8.21	6	0.000	1.000	
treatment5 - treatment7	-15.25 8.21	6	-1.857	0.113	
treatment5 - treatment8	5.00 8.21	6	0.609	0.565	
treatment5 - treatment9	1.00 7.34	6	0.136	0.896	
treatment5 - treatment10	1.00 8.21	6	0.122	0.907	
treatment5 - treatment11	-8.25 8.21	6	-1.005	0.354	
treatment5 - treatment12	-1.25 8.21	6	-0.152	0.884	
treatment6 - treatment7	-15.25 8.21	6	-1.857	0.113	
treatment6 - treatment8	5.00 7.34	6	0.681	0.521	
treatment6 - treatment9	1.00 8.21	6	0.122	0.907	
treatment6 - treatment10	1.00 7.34	6	0.136	0.896	
treatment6 - treatment11	-8.25 8.21	6	-1.005	0.354	
treatment6 - treatment12	-1.25 8.21	6	-0.152	0.884	
treatment7 - treatment8	20.25 8.21	6	2.466	0.049	*
treatment7 - treatment9	16.25 8.21	6	1.979	0.095	
treatment7 - treatment10	16.25 8.21	6	1.979	0.095	
treatment7 - treatment11	7.00 7.34	6	0.953	0.377	
treatment7 - treatment12	14.00 7.34	6	1.906	0.105	
treatment8 - treatment9	-4.00 8.21	6	-0.487	0.643	
treatment8 - treatment10	-4.00 7.34	6	-0.545	0.606	
treatment8 - treatment11	-13.25 8.21	6	-1.614	0.158	
treatment8 - treatment12	-6.25 8.21	6	-0.761	0.475	
treatment9 - treatment10	0.00 8.21	6	0.000	1.000	
treatment9 - treatment11	-9.25 8.21	6	-1.126	0.303	
treatment9 - treatment12	-2.25 8.21	6	-0.274	0.793	
treatment10 - treatment11	-9.25 8.21	6	-1.126	0.303	
treatment10 - treatment12	-2.25 8.21	6	-0.274	0.793	
treatment11 - treatment12	7.00 7.34	6	0.953	0.377	

Treatment Groups

Method : 1sd

Treatment	${\tt Adjusted}$	Means	SE	df	lower.CL	upper.CL	Group
8		73.25	5.61	6	59.52	86.98	1
9		77.25	5.61	6	63.52	90.98	12
10		77.25	5.61	6	63.52	90.98	12
5		78.25	5.61	6	64.52	91.98	12
6		78.25	5.61	6	64.52	91.98	12
2		79.00	3.00	6	71.66	86.34	12
12		79.50	5.61	6	65.77	93.23	12
3		82.00	3.00	6	74.66	89.34	12
4		83.33	3.00	6	76.00	90.67	12
1		84.67	3.00	6	77.33	92.00	12
11		86.50	5.61	6	72.77	100.23	12
7		93.50	5.61	6	79.77	107.23	2

Print results for variable y2 print (out2)

Augmented Design Details

"3" Number of blocks Number of treatments "12" Number of check treatments "4" Number of test treatments "8"

"1, 2, 3, 4" Check treatments

ANOVA, Treatment Adjusted

```
Df Sum Sq Mean Sq F value Pr(>F)
                                              3510 12.261 0.007597 **
Block (ignoring Treatments)
                                  2 7019
Treatment (eliminating Blocks)
                                  11 58965
                                              5360 18.727 0.000920 ***
                                             717 2.504 0.156116
 Treatment: Check
                                  3 2150
 Treatment: Test and Test vs. Check 8 56815 7102 24.810 0.000473 ***
Residuals
                                   6
                                      1718
                                              286
```

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

ANOVA, Block Adjusted

```
Df Sum Sq Mean Sq F value Pr(>F)
Treatment (ignoring Blocks) 11 64708 5883 20.550 0.000707 ***
  Treatment: Check
                            3
                               2150
                                       717
                                            2.504 0.156116
                           7 34863 4980 17.399 0.001366 **
 Treatment: Test
  Treatment: Test vs. Check 1 27694 27694 96.749 6.36e-05 ***
Block (eliminating Treatments) 2 1277
                                            2.231 0.188645
                                       639
                            6 1717
                                        286
Residuals
```

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

Coefficient of Variation

6.057617

Overall Adjusted Mean

298.4792

Standard Errors

	Std.	Error	of Diff.	CD (5%)
Control Treatment Means			13.81424	33.80224
Two Test Treatments (Same Block)			23.92697	58.54719
Two Test Treatments (Different Blocks)			26.75117	65.45775
A Test Treatment and a Control Treatment			21.84224	53.44603

Treatment Means

Treatment	Block	Means	SE	r	Min	Max	Adjusted Means
1		256.00	3.06	3	250.00	260.00	256.00
10	3	450.00	<na></na>	1	450.00	450.00	437.67
11	1	300.00	<na></na>	1	300.00	300.00	299.42
12	1	289.00	<na></na>	1	289.00	289.00	288.42
2		228.00	6.11	3	220.00	240.00	228.00
3		247.67	10.17	3	237.00	268.00	247.67
4		264.00	18.68	3	227.00	287.00	264.00
5	2	281.00	<na></na>	1	281.00	281.00	293.92
6	3	395.00	<na></na>	1	395.00	395.00	382.67
7	1	347.00	<na></na>	1	347.00	347.00	346.42
8	3	226.00	<na></na>	1	226.00	226.00	213.67
9	2	311.00	<na></na>	1	311.00	311.00	323.92

Comparisons

========

Method : 1sd

```
contrast estimate SE df t.ratio p.value sig
treatment1 - treatment2 28.00 13.81 6 2.027 0.089
treatment1 - treatment3
                        8.33 13.81 6 0.603 0.568
treatment1 - treatment4
                        -8.00 13.81 6 -0.579 0.584
treatment1 - treatment5 -37.92 20.72 6 -1.830
                                               0.117
treatment1 - treatment6 -126.67 20.72 6 -6.113 0.001 ***
treatment1 - treatment7 -90.42 20.72 6 -4.363 0.005 **
                       42.33 20.72 6
                                       2.043
treatment1 - treatment8
                                              0.087
treatment1 - treatment9 -67.92 20.72 6 -3.278
                                              0.017
treatment1 - treatment10 -181.67 20.72 6 -8.767
                                              0.000 ***
treatment1 - treatment11 -43.42 20.72 6 -2.095
                                               0.081
treatment1 - treatment12 -32.42 20.72 6 -1.564
                                               0.169
```

```
treatment2 - treatment3
                         -19.67 13.81 6 -1.424
                                                   0.204
treatment2 - treatment4
                         -36.00 13.81 6 -2.606
                                                   0.040
                         -65.92 20.72 6 -3.181
                                                   0.019
treatment2 - treatment5
treatment2 - treatment6 -154.67 20.72 6 -7.464
                                                   0.000 ***
treatment2 - treatment7 -118.42 20.72 6 -5.715
                                                   0.001
treatment2 - treatment8
                          14.33 20.72 6
                                           0.692
                                                   0.515
treatment2 - treatment9
                         -95.92 20.72 6 -4.629
                                                   0.004
                                                   0.000 ***
treatment2 - treatment10 -209.67 20.72 6 -10.118
treatment2 - treatment11
                         -71.42 20.72 6 -3.447
                                                   0.014
                          -60.42 20.72 6 -2.916
treatment2 - treatment12
                                                   0.027
treatment3 - treatment4
                        -16.33 13.81 6 -1.182
                                                   0.282
treatment3 - treatment5 -46.25 20.72 6 -2.232
                                                   0.067
treatment3 - treatment6 -135.00 20.72 6 -6.515
                                                   0.001 ***
treatment3 - treatment7
                          -98.75 20.72 6 -4.766
                                                   0.003
                          34.00 20.72 6
                                           1.641
                                                   0.152
treatment3 - treatment8
treatment3 - treatment9
                         -76.25 20.72 6 -3.680
                                                   0.010
                                                   0.000 ***
treatment3 - treatment10 -190.00 20.72 6 -9.169
treatment3 - treatment11
                          -51.75 20.72 6 -2.497
                                                   0.047
                        -40.75 20.72 6 -1.967
                                                   0.097
treatment3 - treatment12
                        -29.92 20.72 6 -1.444
treatment4 - treatment5
                                                   0.199
treatment4 - treatment6 -118.67 20.72 6 -5.727
                                                   0.001
treatment4 - treatment7
                          -82.42 20.72 6 -3.977
                                                   0.007
treatment4 - treatment8
                           50.33 20.72 6
                                           2.429
                                                   0.051
treatment4 - treatment9
                         -59.92 20.72 6 -2.892
                                                   0.028
treatment4 - treatment10 -173.67 20.72 6 -8.381
                                                   0.000 ***
treatment4 - treatment11
                         -35.42\ 20.72\ 6\ -1.709
                                                   0.138
treatment4 - treatment12 -24.42 20.72 6 -1.178
                                                   0.283
treatment5 - treatment6 -88.75 26.75 6 -3.318
                                                   0.016
treatment5 - treatment7
                          -52.50 26.75 6 -1.963
                                                   0.097
                           80.25 26.75 6
                                           3.000
                                                   0.024
treatment5 - treatment8
treatment5 - treatment9
                         -30.00 23.93 6 -1.254
                                                   0.257
treatment5 - treatment10 -143.75 26.75 6 -5.374
                                                   0.002
treatment5 - treatment11
                          -5.50 26.75 6 -0.206
                                                   0.844
treatment5 - treatment12
                            5.50 26.75 6
                                           0.206
                                                   0.844
treatment6 - treatment7
                          36.25 26.75 6
                                           1.355
                                                   0.224
treatment6 - treatment8
                         169.00 23.93 6
                                           7.063
                                                   0.000 ***
                           58.75 26.75 6
                                                   0.070
treatment6 - treatment9
                                           2.196
treatment6 - treatment10
                          -55.00 23.93 6 -2.299
                                                   0.061
treatment6 - treatment11
                          83.25 26.75 6
                                           3.112
                                                   0.021
treatment6 - treatment12
                           94.25 26.75 6
                                           3.523
                                                   0.012
treatment7 - treatment8
                        132.75 26.75
                                       6
                                           4.962
                                                   0.003
                          22.50 26.75 6
                                           0.841
                                                   0.433
treatment7 - treatment9
treatment7 - treatment10
                         -91.25 26.75 6 -3.411
                                                   0.014
treatment7 - treatment11
                           47.00 23.93 6
                                           1.964
                                                   0.097
treatment7 - treatment12
                           58.00 23.93 6
                                           2.424
                                                   0.052
treatment8 - treatment9 -110.25 26.75 6 -4.121
                                                   0.006
                                                          **
treatment8 - treatment10 -224.00 23.93 6 -9.362
                                                   0.000 ***
treatment8 - treatment11
                          -85.75 26.75 6
                                          -3.205
                                                   0.018
                         -74.75 26.75 6 -2.794
                                                   0.031
treatment8 - treatment12
                                                          **
treatment9 - treatment10 -113.75 26.75 6 -4.252
                                                   0.005
treatment9 - treatment11
                          24.50 26.75 6
                                           0.916
                                                   0.395
treatment9 - treatment12
                           35.50 26.75 6
                                           1.327
                                                   0.233
```

```
treatment10 - treatment11 138.25 26.75 6 5.168 0.002 **
treatment10 - treatment12 149.25 26.75 6 5.579 0.001 **
treatment11 - treatment12 11.00 23.93 6 0.460 0.662
```

Treatment Groups

Method : 1sd

Treatment	Adjusted Means	SE	df	lower.CL	upper.CL	Group
8	213.67	18.27	6	168.95	258.38	12
2	228.00	9.77	6	204.10	251.90	1
3	247.67	9.77	6	223.76	271.57	123
1	256.00	9.77	6	232.10	279.90	1234
4	264.00	9.77	6	240.10	287.90	234
12	288.42	18.27	6	243.70	333.13	345
5	293.92	18.27	6	249.20	338.63	345
11	299.42	18.27	6	254.70	344.13	45
9	323.92	18.27	6	279.20	368.63	56
7	346.42	18.27	6	301.70	391.13	56
6	382.67	18.27	6	337.95	427.38	67
10	437.67	18.27	6	392.95	482.38	7

7.3 describe.augmentedRCBD()

The descriptive statistics such as count, mean, standard error, minimum, maximum, skewness (with p-value from D'Agostino test of skewness (D'Agostino (1970))) and kurtosis (with p-value from Anscombe-Glynn test of kurtosis (Anscombe and Glynn (1983))) for the adjusted means from the results in an object of class augmentedRCBD can be computed as follows.

```
# Descriptive statistics for variable y1
describe.augmentedRCBD(out1)
```

\$Count

[1] 12

\$Mean

[1] 81.0625

\$Std.Error

[1] 1.547002

\$Std.Deviation

[1] 5.358973

\$Min

[1] 73.25

\$Max

[1] 93.5

\$`Skewness(statistic)` skew z

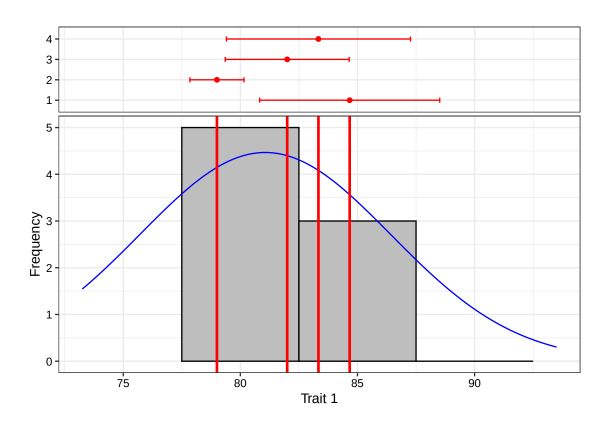
```
0.9250344 1.6745760
$`Skewness(p.value)`
[1] 0.09401746
$`Kurtosis(statistic)`
    kurt
3.522807 1.282305
$`Kurtosis(p.value)`
[1] 0.1997357
# Descriptive statistics for variable y2
describe.augmentedRCBD(out2)
$Count
[1] 12
$Mean
[1] 298.4792
$Std.Error
[1] 18.92257
$Std.Deviation
[1] 65.5497
$Min
[1] 213.6667
$Max
[1] 437.6667
$`Skewness(statistic)`
     skew
0.7449405 1.3680211
$`Skewness(p.value)`
[1] 0.1713055
$`Kurtosis(statistic)`
    kurt
2.787997 0.536812
$`Kurtosis(p.value)`
[1] 0.5913975
```

7.4 freqdist.augmentedRCBD()

The frequency distribution of the adjusted means from the results in an object of class **augmentedRCBD** can be plotted as follows.

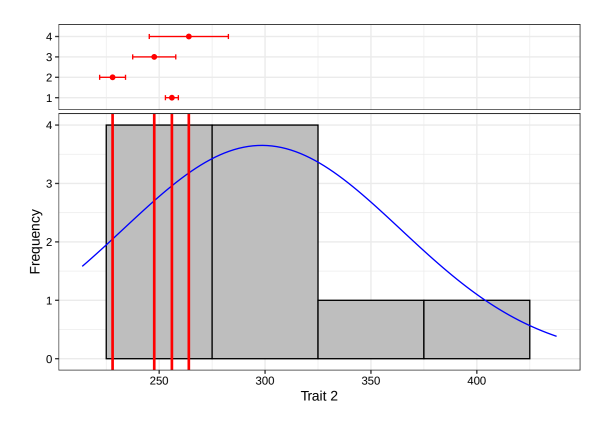
```
# Frequency distribution for variable y1
freq1 <- freqdist.augmentedRCBD(out1, xlab = "Trait 1")</pre>
```

Warning: Removed 2 rows containing missing values (`geom_bar()`).
plot(freq1)



```
# Frequency distribution for variable y2
freq2 <- freqdist.augmentedRCBD(out2, xlab = "Trait 2")</pre>
```

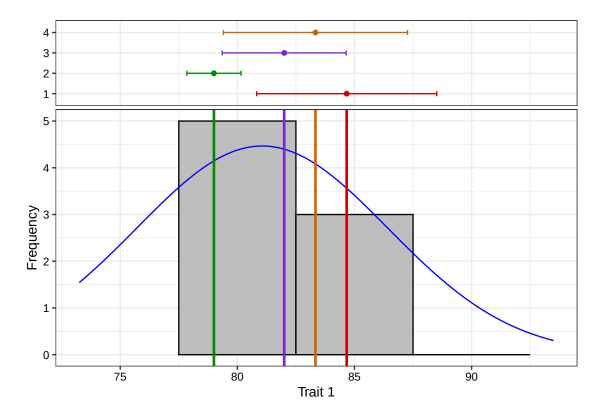
Warning: Removed 2 rows containing missing values (`geom_bar()`).
plot(freq2)



The colours for the check values may be specified using the argument check.col.

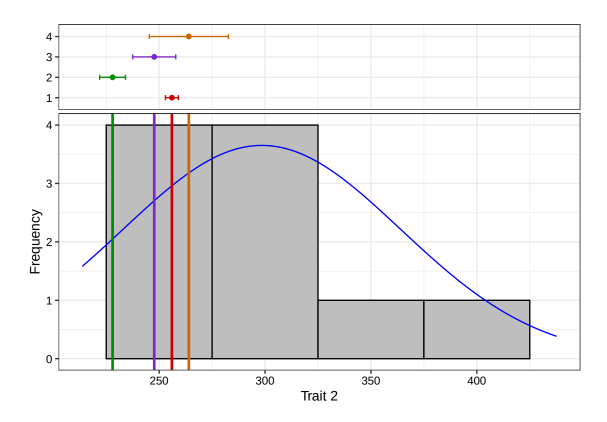
```
colset <- c("red3", "green4", "purple3", "darkorange3")
# Frequency distribution for variable y1
freq1 <- freqdist.augmentedRCBD(out1, xlab = "Trait 1", check.col = colset)</pre>
```

```
Warning: Removed 2 rows containing missing values (`geom_bar()`).
plot(freq1)
```



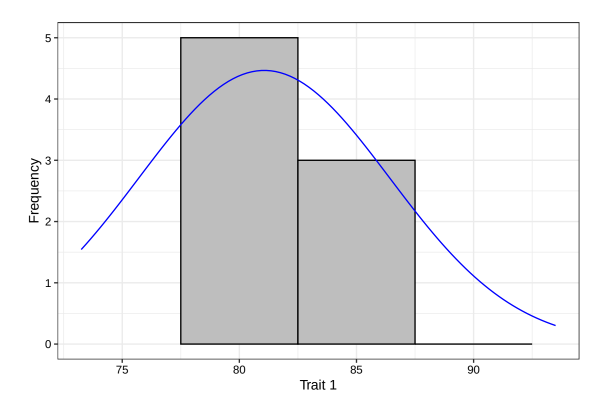
Frequency distribution for variable y2
freq2 <- freqdist.augmentedRCBD(out2, xlab = "Trait 2", check.col = colset)</pre>

```
Warning: Removed 2 rows containing missing values (`geom_bar()`).
plot(freq2)
```

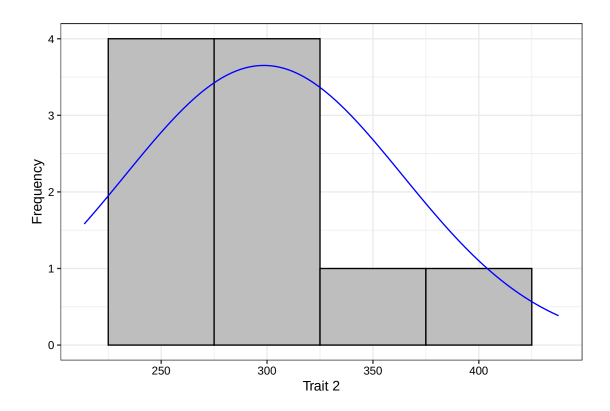


The default the check highlighting can be avoided using the argument highlight.check = FALSE.

```
Warning: Removed 2 rows containing missing values (`geom_bar()`).
plot(freq1)
```



```
Warning: Removed 2 rows containing missing values (`geom_bar()`).
plot(freq2)
```



7.5 gva.augmentedRCBD()

The genetic variability statistics such as mean, phenotypic, genotypic and environmental variation (Federer and Searle (1976)), phenotypic, genotypic and environmental coefficient of variation (Burton (1951), Burton (1952)), category of phenotypic and genotypic coefficient of variation according to Sivasubramaniam and Madhavamenon (1973), broad-sense heritability (H^2) (Lush (1940)), H^2 category according to Robinson (1966), Genetic advance (GA), genetic advance as per cent of mean (GAM) and GAM category according to Johnson et al. (1955) are computed from an object of class **augmentedRCBD** as follows. Genetic variability analysis needs to be performed only if the sum of squares of "Treatment: Test" are significant.

Genetic variability statistics for variable y1 gva.augmentedRCBD(out1)

Warning in gva.augmentedRCBD(out1): P-value for "Treatment: Test" is > 0.05. Genetic variability analysis may not be appropriate for this trait.

\$Mean

[1] 81.0625

\$PV

[1] 72.26786

\$GV

[1] 45.29563

\$EV

```
[1] 26.97222
$GCV
[1] 8.302487
$`GCV category`
[1] "Low"
$PCV
[1] 10.48703
$`PCV category`
[1] "Medium"
$ECV
[1] 6.406759
$hBS
[1] 62.67743
$`hBS category`
[1] "High"
$GA
[1] 10.99216
$GAM
[1] 13.5601
$`GAM category`
[1] "Medium"
# Genetic variability statistics for variable y2
gva.augmentedRCBD(out2)
$Mean
[1] 298.4792
$PV
[1] 4980.411
$GV
[1] 4694.161
$EV
[1] 286.25
$GCV
[1] 22.95435
$`GCV category`
[1] "High"
```

```
$PCV
[1] 23.64387
$`PCV category`
[1] "High"
$ECV
[1] 5.668377
$hBS
[1] 94.25248
$`hBS category`
[1] "High"
$GA
[1] 137.2223
$GAM
[1] 45.97382
$`GAM category`
[1] "High"
```

Negative estimates of variance components if computed are not abnormal. For information on how to deal with these, refer Robinson et al. (1955) and Dudley and Moll (1969).

7.5 report.augmentedRCBD()

The results generated by the analysis can be exported to a MS Word file as follows.

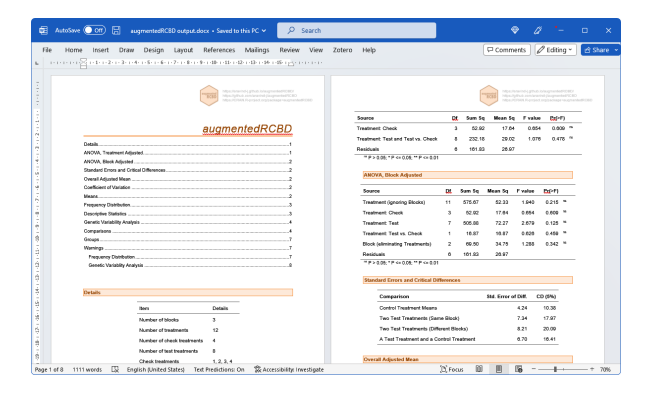


Fig. 6: MS Word report generated with report.agumentedRCBD function.

Alternatively, the analysis results can also be exported to a MS Excel file as follows.

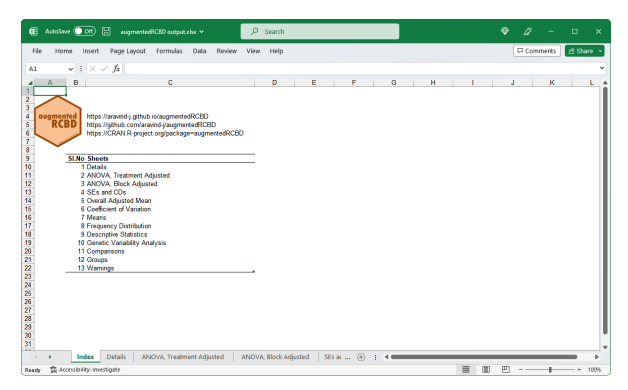


Fig. 7: MS Excel report generated with report.agumentedRCBD function.

8 Data Analysis for a Multiple Traits

Analysis of data for a multiple traits simultaneously can be performed by using augmentedRCBD.bulk function. It generates an object of class augmentedRCBD.bulk. Such an object can then be taken as input by print.augmentedRCBD.bulk to print the results to console. The results can also be exported as a MS Word report using the report.augmentedRCBD.bulk function.

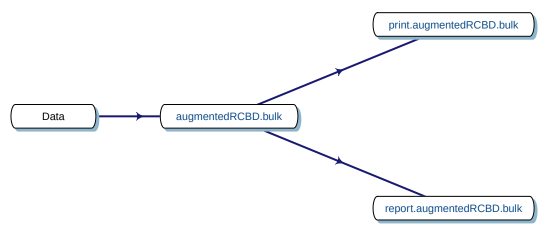


Fig. 8. Workflow for analysis of multiple traits with augmentedRCBD.

8.1 augmentedRCBD.bulk()

Consider the data frame data imported from Table 1 according to the instructions in section 4.8.

str(data)

'data.frame': 20 obs. of 4 variables: \$ blk: Factor w/ 3 levels "1","2","3": 1 1 1 1 1 1 1 2 2 2 ... \$ trt: Factor w/ 12 levels "1","2","3","4",..: 1 2 3 4 7 11 12 1 2 3 ... \$ y1 : num 92 79 87 81 96 89 82 79 81 81 ... \$ y2 : num 258 224 238 278 347 300 289 260 220 237 ... # Convert block and treatment to factors data\$blk <- as.factor(data\$blk) data\$trt <- as.factor(data\$trt)

Rather than performing the analysis individually for each variable/trait separately using augmentedRCBD, the analysis can be performed simultaneously for for both the traits using augmentedRCBD.bulk function. It is a wrapper around the augmentedRCBD core function and its associated helper functions.

However in this case treatment comparisons/grouping by least significant difference or Tukey's honest significant difference method is not computed. Also the output object size is reduced using the **simplify = TRUE** argument in the **augmentedRCBD** function.

The logical arguments **describe**, **freqdist** and **gva** can be used to specify whether to generate the descriptive statistics, frequency distribution plots and genetic variability statistics respectively. If **gva = True**, then plots to compare phenotypic and genotypic coefficient of variation, broad sense heritability and genetic advance over mean between traits are also generated.

```
ANOVA for y1 computed (1/2)
ANOVA for y2 computed (2/2)
Augmented Design Details
_____
Number of blocks
                         "3"
Number of treatments
                         "12"
Number of check treatments "4"
Number of test treatments "8"
Check treatments
                         "1, 2, 3, 4"
                         "2"
Number of traits
Traits
                         "y1, y2"
ANOVA, Treatment Adjusted
```

Mean.Sq

```
Source Df
                                        у1
       Block (ignoring Treatments) 2 180.04 * 3509.67 **
    Treatment (eliminating Blocks) 11 25.92 ns 5360.49 **
                  Treatment: Check 3 17.64 ns 716.75 ns
 Treatment: Test and Test vs. Check 8 29.02 ns 7101.89 **
                         Residuals 6 26.97
                                              286.25
^{n \, s} \, P > 0.05; * P <= 0.05; ** P <= 0.01
ANOVA, Block Adjusted
                                    Mean.Sq
                        Source Df
                                    y1
    Treatment (ignoring Blocks) 11 52.33 ns 5882.50 **
              Treatment: Check 3 17.64 ^{ns} 716.75 ^{ns}
      Treatment: Test vs. Check 1 16.87 ns 27694.41 **
               Treatment: Test 7 72.27 ns 4980.41 **
 Block (eliminating Treatments) 2 34.75 ns
                                          638.58 ns
                    Residuals 6 26.97
                                           286.25
^{n \, s} \, P > 0.05; * P <= 0.05; ** P <= 0.01
Coefficient of Variation
_____
 Trait CV
   y1 6.37
   y2 6.06
Overall Adjusted Mean
_____
 Trait Overall.adjusted.mean
   у1
                    81.06
                    298.48
   y2
Standard Errors
_____
                             Comparison y1
A Test Treatment and a Control Treatment 6.70 21.84
                 Control Treatment Means 4.24 13.81
  Two Test Treatments (Different Blocks) 8.21 26.75
        Two Test Treatments (Same Block) 7.34 23.93
Critical Difference
_____
alpha = 0.05
                             Comparison y1
A Test Treatment and a Control Treatment 16.41 53.45
                 Control Treatment Means 10.38 33.80
   Two Test Treatments (Different Blocks) 20.09 65.46
        Two Test Treatments (Same Block) 17.97 58.55
```

Descriptive Statistics

```
_____
```

Trait Count Mean Std.Error Std.Deviation Min Max Skewness Skewness_sig
y1 12 81.06 1.55 5.36 73.25 93.50 0.93 **
y2 12 298.48 18.92 65.55 213.67 437.67 0.74 **

ns*
**ns*

Kurtosis Kurtosis_sig

3.52

2.79 ns

 $^{n \, s} \, P > 0.05; * P <= 0.05; ** P <= 0.01$

Genetic Variability Analysis

k = 2.063

Trait Mean PV GV EV GCV.category PCV.category ECV y1 + 81.06 72.27 45.30 26.97 8.30 Low 10.49 Medium 6.41 y2 298.48 4980.41 4694.16 286.25 22.95 High 23.64 High 5.67 hBS category GA GAM GAM.category 62.68 High 10.99 13.56 Medium

High

Warning:

94.25

† P-value for "Treatment: Test" is > 0.05. Genetic variability analysis may not be appropriate for

Warning:

‡ Negative GV detected.

GCV, GCV category, hBS, hBS category, GA, GAM and GAM category could not be computed.

High 137.22 45.97

Warning Messages

[Frequency Distribution]

<y1>

Removed 2 rows containing missing values (`geom_bar()`).

<y2>

Removed 2 rows containing missing values (`geom_bar()`).

[GVA]

<y1>

P-value for "Treatment: Test" is > 0.05. Genetic variability analysis may not be appropriate for t

Treatment Means

Treatment Block y1 y2 1 84.67 256.00 10 3 77.25 437.67 11 1 86.50 299.42

```
12
      1 79.50 288.42
        79.00 228.00
2
3
        82.00 247.67
4
        83.33 264.00
5
      2 78.25 293.92
      3 78.25 382.67
6
7
      1 93.50 346.42
      3 73.25 213.67
8
      2 77.25 323.92
```

8.2 print.augmentedRCBD.bulk()

The results of analysis in an object of class augmentedRCBD.bulk can be printed to the console as follows.

```
# Print results
print(bout)
```

Augmented Design Details

```
Number of blocks "3"
Number of treatments "12"
Number of check treatments "4"
Number of test treatments "8"
Check treatments "1, 2, 3, 4"
Number of traits "2"
Traits "y1, y2"
```

ANOVA, Treatment Adjusted

ANOVA, Block Adjusted

Coefficient of Variation

Trait CV

y1 6.37

y2 6.06

Overall Adjusted Mean

=============

Trait Overall.adjusted.mean

y1 81.06

y2 298.48

Standard Errors

Comparison y1 y2

A Test Treatment and a Control Treatment 6.70 21.84

Control Treatment Means 4.24 13.81

Two Test Treatments (Different Blocks) 8.21 26.75

Two Test Treatments (Same Block) 7.34 23.93

Critical Difference

alpha = 0.05

Comparison y1 y2

A Test Treatment and a Control Treatment 16.41 53.45

Control Treatment Means 10.38 33.80

Two Test Treatments (Different Blocks) 20.09 65.46

Two Test Treatments (Same Block) 17.97 58.55

Descriptive Statistics

Trait Count Mean Std.Error Std.Deviation Min Max Skewness Skewness_sig v1 12 81.06 1.55 5.36 73.25 93.50 0.93 ***

y1 12 81.06 1.55 5.36 73.25 93.50 0.93 ^{n s} y2 12 298.48 18.92 65.55 213.67 437.67 0.74

Kurtosis Kurtosis_sig

3.52 ns

2.79 ns

 $^{n \, s} \, P > 0.05; * P <= 0.05; ** P <= 0.01$

Genetic Variability Analysis

k = 2.063

Trait Mean PV GV EV GCV GCV.category PCV PCV.category ECV y1 † 81.06 72.27 45.30 26.97 8.30 Low 10.49 Medium 6.41 y2 298.48 4980.41 4694.16 286.25 22.95 High 23.64 High 5.67

hBS hBS.category GA GAM.category

```
62.68
              High 10.99 13.56
                                     Medium
 94.25
              High 137.22 45.97
                                        High
Warning:
† P-value for "Treatment: Test" is > 0.05. Genetic variability analysis may not be appropriate for
Warning:
‡ Negative GV detected.
GCV, GCV category, hBS, hBS category, GA, GAM and
GAM category could not be computed.
Warning Messages
_____
[Frequency Distribution]
<y1>
Removed 2 rows containing missing values (`geom_bar()`).
<y2>
Removed 2 rows containing missing values (`geom_bar()`).
[GVA]
```

P-value for "Treatment: Test" is > 0.05. Genetic variability analysis may not be appropriate for t

Treatment Means

<y1>

==============

```
Treatment Block y1
      1
             84.67 256.00
      10
            3 77.25 437.67
      11
            1 86.50 299.42
      12
           1 79.50 288.42
              79.00 228.00
       2
       3
              82.00 247.67
       4
              83.33 264.00
            2 78.25 293.92
            3 78.25 382.67
       6
            1 93.50 346.42
       7
            3 73.25 213.67
       8
             2 77.25 323.92
```

8.3 report.augmentedRCBD.bulk()

The results generated by the analysis can be exported to a MS Word file as follows.

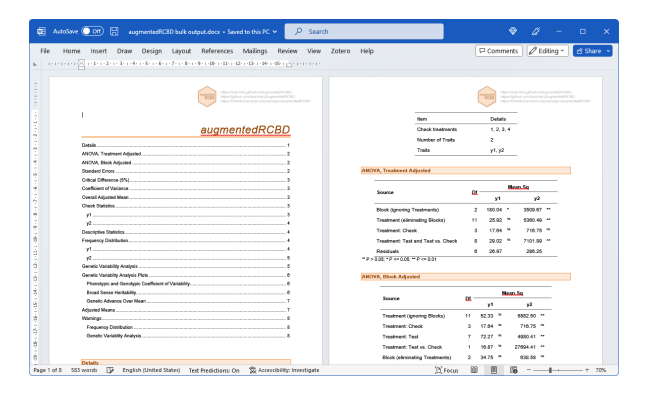


Fig. 9: MS Word report generated with report.agumentedRCBD.bulk function.

Alternatively, the analysis results can also be exported to a MS Excel file as follows.

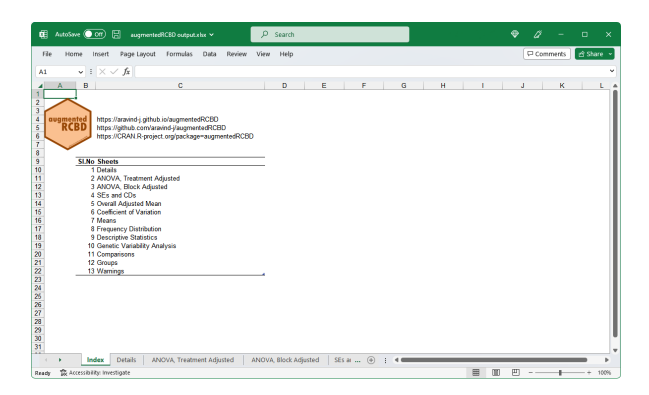


Fig. 10: MS Excel report generated with report.agumentedRCBD.bulk function.

9 Citing augmentedRCBD

citing the package.

```
To cite the R package 'augmentedRCBD' in publications use:

Aravind, J., Mukesh Sankar, S., Wankhede, D. P., and Kaur, V. ().
augmentedRCBD: Analysis of Augmented Randomised Complete Block
Designs. R package version 0.1.6,
https://aravind-j.github.io/augmentedRCBD/https://cran.r-project.org/package=augmentedRCBD.

A BibTeX entry for LaTeX users is

@Manual{,
   title = {augmentedRCBD: Analysis of Augmented Randomised Complete Block Designs},
   author = {J. Aravind and S. {Mukesh Sankar} and Dhammaprakash Pandhari Wankhede and Vikender K
   note = {https://aravind-j.github.io/augmentedRCBD/},
   note = {https://cran.r-project.org/package=augmentedRCBD},
}

This free and open-source software implements academic research by the
```

authors and co-workers. If you use it, please support the project by

10 Session Info

```
sessionInfo()
```

```
R Under development (unstable) (2023-04-28 r84338 ucrt)
Platform: x86_64-w64-mingw32/x64 (64-bit)
Running under: Windows 10 x64 (build 19045)
Matrix products: default
locale:
[1] LC COLLATE=C
                                    LC CTYPE=English India.utf8
[3] LC_MONETARY=English_India.utf8 LC_NUMERIC=C
[5] LC_TIME=English_India.utf8
time zone: Asia/Calcutta
tzcode source: internal
attached base packages:
[1] stats
              graphics grDevices utils
                                             datasets methods
                                                                 base
other attached packages:
[1] diagram_1.6.5
                                             augmentedRCBD_0.1.6
                         shape_1.4.6
loaded via a namespace (and not attached):
 [1] tidyselect_1.2.0
                              farver_2.1.1
                                                      dplyr_1.1.2
                                                      RCurl_1.98-1.12
 [4] bitops_1.0-7
                              fastmap_1.1.1
 [7] TH.data_1.1-2
                             fontquiver_0.2.1
                                                      mathjaxr_1.6-0
[10] promises_1.2.0.1
                             XML_3.99-0.14
                                                      digest_0.6.31
[13] estimability_1.4.1
                             mime_0.12
                                                      lifecycle_1.0.3
[16] multcompView_0.1-9
                              ellipsis_0.3.2
                                                      gfonts_0.2.0
[19] survival_3.5-5
                              magrittr_2.0.3
                                                      compiler_4.4.0
[22] rlang_1.1.0
                              tools_4.4.0
                                                      utf8_1.2.3
[25] yaml_2.3.7
                              data.table_1.14.8
                                                      knitr_1.42
[28] labeling_0.4.2
                              askpass_1.1
                                                      curl_5.0.0
[31] plyr_1.8.8
                              xm12_1.3.4
                                                      multcomp_1.4-23
[34] numform_0.7.0
                             httpcode_0.3.0
                                                      withr_2.5.0
[37] grid_4.4.0
                              fansi_1.0.4
                                                      gdtools_0.3.3
[40] xtable_1.8-4
                              colorspace_2.1-0
                                                      ggplot2_3.4.2
[43] emmeans_1.8.5
                              scales_1.2.1
                                                      MASS_7.3-59
[46] crul_1.3
                              cli_3.6.1
                                                      mvtnorm_1.1-3
[49] rmarkdown 2.21
                              crayon_1.5.2
                                                      ragg_1.2.5
[52] generics_0.1.3
                              rstudioapi_0.14
                                                      httr_1.4.6
[55] reshape2_1.4.4
                              stringr_1.5.0
                                                      splines_4.4.0
[58] vctrs_0.6.2
                             Matrix_1.5-4
                                                      sandwich_3.0-2
[61] jsonlite_1.8.4
                              fontBitstreamVera_0.1.1 systemfonts_1.0.4
[64] glue_1.6.2
                              codetools_0.2-19
                                                      stringi_1.7.12
[67] flextable_0.9.1
                              gtable_0.3.3
                                                      later_1.3.0
                                                      pillar_1.9.0
[70] munsell_0.5.0
                              tibble_3.2.1
[73] htmltools_0.5.5
                              openssl_2.0.6
                                                      R6_2.5.1
[76] textshaping_0.3.6
                              Rdpack_2.4
                                                      evaluate_0.21
```

[79] shiny_1.7.4	lattice_0.21-8	highr_0.10
[82] rbibutils_2.2.13	moments_0.14.1	openxlsx_4.2.5.2
[85] fontLiberation_0.1.0	httpuv_1.6.9	Rcpp_1.0.10
[88] zip_2.3.0	uuid_1.1-0	coda_0.19-4
[91] officer_0.6.2	xfun_0.39	zoo_1.8-12
[94] pkgconfig_2.0.3		

References

- Anscombe, F. J., and Glynn, W. J. (1983). Distribution of the kurtosis statistic b_2 for normal samples. Biometrika 70, 227–234. doi:10.1093/biomet/70.1.227.
- Burton, G. W. (1951). Quantitative inheritance in pearl millet (*Pennisetum glaucum*). Agronomy Journal 43, 409–417. doi:10.2134/agronj1951.00021962004300090001x.
- Burton, G. W. (1952). Qualitative inheritance in grasses. Vol. 1. in *Proceedings of the 6th International Grassland Congress, Pennsylvania State College*, 17–23.
- D'Agostino, R. B. (1970). Transformation to normality of the null distribution of g_1 . Biometrika 57, 679–681. doi:10.1093/biomet/57.3.679.
- Dudley, J. W., and Moll, R. H. (1969). Interpretation and use of estimates of heritability and genetic variances in plant breeding. *Crop Science* 9, 257–262. doi:10.2135/cropsci1969.0011183X000900030001x.
- Federer, W. T. (1956b). Augmented (or Hoonuiaku) Designs. New York: Cornell University Available at: https://ecommons.cornell.edu/handle/1813/32841.
- Federer, W. T. (1956a). Augmented (or Hoonuiaku) designs. The Hawaiian Planters' Record LV(2), 191–208.
- Federer, W. T. (1961). Augmented designs with one-way elimination of heterogeneity. *Biometrics* 17, 447–473. doi:10.2307/2527837.
- Federer, W. T., and Searle, S. R. (1976). Model Considerations and Variance Component Estimation in Augmented Completely Randomized and Randomized Complete Blocks Designs-Preliminary Version. New York: Cornell University Available at: https://hdl.handle.net/1813/32691.
- Johnson, H. W., Robinson, H. F., and Comstock, R. E. (1955). Estimates of genetic and environmental variability in soybeans. *Agronomy journal* 47, 314–318. doi:10.2134/agronj1955.00021962004700070009x.
- Lush, J. L. (1940). Intra-sire correlations or regressions of offspring on dam as a method of estimating heritability of characteristics. *Proceedings of the American Society of Animal Nutrition* 1940, 293–301. doi:10.2527/jas1940.19401293x.
- Robinson, H. F. (1966). Quantitative genetics in relation to breeding on centennial of Mendelism. *Indian Journal of Genetics and Plant Breeding*, 171.
- Robinson, H. F., Comstock, R. E., and Harvey, P. H. (1955). Genetic variances in open pollinated varieties of corn. *Genetics* 40, 45–60. doi:10.1093/genetics/40.1.45.
- Sivasubramaniam, S., and Madhavamenon, P. (1973). Genotypic and phenotypic variability in rice. The Madras Agricultural Journal 60, 1093–1096.
- Tippmann, S. (2015). Programming tools: Adventures with R. Nature News 517, 109. doi:10.1038/517109a.