# Package 'augmentedRCBD'

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```
Title Analysis of Augmented Randomised Complete Block Designs
Version 0.1.1.9000
Description Functions for analysis of data generated from experiments in
     augmented randomised complete block design according to Federer, W.T. (1961)
     <doi:10.2307/2527837>. Computes analysis of variance, adjusted means,
     descriptive statistics, genetic variability statistics etc. Further includes
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```

utils

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```
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```

# **R** topics documented:

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augmented RCBD

Analysis of Augmented Randomised Complete Block Design

# **Description**

augmentedRCBD is a function for analysis of variance of an augmented randomised block design (Federer, 1956; Federer, 1961) and the generation as well as comparison of the adjusted means of the treatments/genotypes.

# Usage

```
augmentedRCBD(
  block,
  treatment,
  y,
  checks = NULL,
  method.comp = c("lsd", "tukey", "none"),
  alpha = 0.05,
  group = TRUE,
  console = TRUE,
  simplify = FALSE,
  truncate.means = TRUE
)
```

# **Arguments**

block Vector of blocks (as a factor).

treatment Vector of treatments/genotypes (as a factor).

y Numeric vector of response variable (Trait).

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checks Character vector of the checks present in treatment levels. If not specified,

checks are inferred from the data on the basis of number of replications of treat-

ments/genotypes.

method.comp Method for comparison of treatments ("1sd" for least significant difference or

"tukey" for Tukey's honest significant difference). If "none", no comparisons will be made, the ANOVA output will be given as a data frame and the adjusted means will be computed directly from treatment and block effects instead of

using emmeans.

alpha Type I error probability (Significance level) to be used for multiple comparisons.

group If TRUE, genotypes will be grouped according to "method.comp". Default is

TRUE.

console If TRUE, output will be printed to console. Default is TRUE. Default is TRUE.

simplify If TRUE, ANOVA output will be given as a data frame instead of a summary.aov

object. Default is TRUE.

truncate.means If TRUE, the negative adjusted means will be truncated to zero. Default is TRUE.

#### **Details**

This function borrows code from DAU. test function of agricolae package (de Mendiburu et al., 2016) as well as from Appendix VIII of Mathur et al., (2008).

#### Value

A list of class augmentedRCBD containing the following components:

Details Details of the augmented design used.

Means A data frame with the "Means", "Block", "SE", "Mix", "Max" and "Adjusted

Means" for each "Treatment".

ANOVA, Treatment Adjusted

An object of class summary. aov for ANOVA table with treatments adjusted.

ANOVA, Block Adjusted

An object of class summary. aov for ANOVA table with block adjusted.

Block effects A vector of block effects.

Treatment effects

A vector of treatment effects.

Std. Errors A data frame of standard error of difference between various combinations along

with critical difference and tukey's honest significant difference (when method.comp

= "tukey") at alpha.

Overall adjusted mean

Overall adjusted mean.

CV Coefficient of variation.

Comparisons A data frame of pairwise comparisons of treatments. This is computed only if

argument group is TRUE

Groups A data frame with compact letter display of pairwise comparisons of treatments.

Means with at least one letter common are not significantly different statistically.

This is computed only if argument group is TRUE

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#### Note

• Data should preferably be balanced i.e. all the check genotypes should be present in all the blocks. If not, a warning is issued.

- There should not be any missing values.
- The number of test genotypes can vary within a block.

In case the large number of treatments or genotypes, it is advisable to avoid 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.

### References

Federer WT (1956). "Augmented (or hoonuiaku) designs." *The Hawaiian Planters' Record*, **LV(2)**, 191–208.

Federer WT (1961). "Augmented designs with one-way elimination of heterogeneity." *Biometrics*, **17**(3), 447–473.

Mathur PN, Muralidharan K, Parthasarathy VA, Batugal P, Bonnot F (2008). *Data Analysis Manual for Coconut Researchers-Bioversity Technical Bulletin No. 14*. Bioversity International. ISBN 978-92-9043-736-9.

de Mendiburu F (2015). agricolae: Statistical Procedures for Agricultural Research. R package version 1.2-8.

#### See Also

```
DAU. test, ea1, emmeans, cld.emmGrid, aug.rcb
```

#### **Examples**

```
# Example data
blk <- c(rep(1,7), rep(2,6), rep(3,7))
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)
data <- data.frame(blk, trt, y1, y2)</pre>
# Convert block and treatment to factors
data$blk <- as.factor(data$blk)</pre>
data$trt <- as.factor(data$trt)</pre>
# Results for variable y1 (checks inferred)
out1 <- augmentedRCBD(data$blk, data$trt, data$y1, method.comp = "lsd",</pre>
                       alpha = 0.05, group = TRUE, console = TRUE)
# Results for variable y2 (checks inferred)
out2 <- augmentedRCBD(data$blk, data$trt, data$y1, method.comp = "lsd",</pre>
                      alpha = 0.05, group = TRUE, console = TRUE)
# Results for variable y1 (checks specified)
out1 <- augmentedRCBD(data$blk, data$trt, data$y1, method.comp = "lsd",</pre>
                       alpha = 0.05, group = TRUE, console = TRUE,
                       checks = c("1", "2", "3", "4"))
# Results for variable y2 (checks specified)
out2 <- augmentedRCBD(data$blk, data$trt, data$y1, method.comp = "lsd",
                       alpha = 0.05, group = TRUE, console = TRUE,
                       checks = c("1", "2", "3", "4"))
```

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```
## Not run:
# Error in case checks not replicated across all blocks
# Check 1 and 4 not replicated in all 3 blocks
trt <- c(1, 2, 3, 14, 7, 11, 12, 1, 2, 3, 4, 5, 9, 13, 2, 3, 4, 8, 6, 10)
data$trt <- as.factor(trt)</pre>
table(data$trt, data$blk)
# Results for variable y1 (checks specified)
out1 <- augmentedRCBD(data$blk, data$trt, data$y1, method.comp = "lsd",</pre>
                       alpha = 0.05, group = TRUE, console = TRUE,
                       checks = c("1", "2", "3", "4"))
## End(Not run)
# Warning in case test treatments are replicated
out1 <- augmentedRCBD(data$blk, data$trt, data$y1, method.comp = "lsd",</pre>
                       alpha = 0.05, group = TRUE, console = TRUE)
out1 <- augmentedRCBD(data$blk, data$trt, data$y1, method.comp = "lsd",</pre>
                       alpha = 0.05, group = TRUE, console = TRUE,
                       checks = c("2", "3"))
```

 $augmented {\tt RCBD.bulk}$ 

Analysis of Augmented Randomised Complete Block Design for Multiple Traits/Characters

# **Description**

augmentedRCBD.bulk is a wrapper around the functions augmentedRCBD, describe.augmentedRCBD, freqdist.augmentedRCBD and gva.augmentedRCBD. It will carry out these analyses for multiple traits/characters from the input data as a data frame object.

#### Usage

```
augmentedRCBD.bulk(
  data,
  block,
  treatment,
  traits,
  checks = NULL,
  alpha = 0.05,
  describe = TRUE,
  freqdist = TRUE,
  gva = TRUE,
  check.col = "red",
  console = TRUE
)
```

# **Arguments**

data

The data as a data frame object. The data frame should possess columns speci-

fying the block, treatment and multiple traits/characters.

block Name of column specifying the blocks in the design as a character string.

treatment Name of column specifying the treatments as a character string.

traits Name of columns specifying the multiple traits/characters as a character vector.

checks Character vector of the checks present in treatment levels. If not specified,

checks are inferred from the data on the basis of number of replications of treat-

ments/genotypes.

alpha Type I error probability (Significance level) to be used for multiple comparisons.

describe If TRUE, descriptive statistics will be computed. Default is TRUE.

frequist If TRUE, frequency distributions be plotted. Default is TRUE.

gva If TRUE, genetic variability analysis will be done. Default is TRUE.

check.col The colour(s) to be used to highlight check values in the plot as a character

vector. Must be valid colour values in R (named colours, hexadecimal represen-

tation, index of colours [1:8] in default R 'palette()' etc.).

console If TRUE, output will be printed to console. Default is TRUE.

#### Value

A list of class augmentedRCBD.bulk containing the following components:

Details Details of the augmented design used and the traits/characters.

ANOVA, Treatment Adjusted

A data frame of mean sum of squares of the specified traits from treatment ad-

justed ANOVA.

ANOVA, Block Adjusted

A data frame of mean sum of squares of the specified traits from block adjusted

**ANOVA** 

Means A data frame of the adjusted means of the treatments for the specified traits.

alpha Type I error probability (Significance level) used.

Std. Errors A data frame of standard error of difference between various combinations for

the specified traits.

CD A data frame of critical difference (at the specified alpha) between various com-

binations for the specified traits.

Overall adjusted mean

A data frame of the overall adjusted mean for the specified traits.

CV A data frame of the coefficient of variance for the specified traits.

Descriptive statistics

A data frame of descriptive statistics for the specified traits.

Frequency distribution

A list of ggplot2 plot grobs of the frequency distribution plots.

Genetic variability analysis

A data frame of genetic variability statistics for the specified traits.

GVA plots A list of three ggplot2 objects with the plots for (a) Phenotypic and Genotypic

CV, (b) Broad sense heritability and (c) Genetic advance over mean

warnings A list of warning messages (if any) captured during model fitting and frequency

distribution plotting.

#### Note

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.

#### See Also

augmentedRCBD, describe.augmentedRCBD, freqdist.augmentedRCBD, gva.augmentedRCBD

# **Examples**

```
# Example data
blk <- c(rep(1,7),rep(2,6),rep(3,7))
trt <- c(1, 2, 3, 4, 7, 11, 12, 1, 2, 3, 4, 5, 9, 1, 2, 3, 4, 8, 6, 10)
y1 \leftarrow 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)
dataf <- data.frame(blk, trt, y1, y2)</pre>
bout <- augmentedRCBD.bulk(data = dataf, block = "blk",</pre>
                            treatment = "trt", traits = c("y1", "y2"),
                            checks = NULL, alpha = 0.05, describe = TRUE,
                            freqdist = TRUE, gva = TRUE,
                            check.col = c("brown", "darkcyan",
                                           "forestgreen", "purple"),
                            console = TRUE)
# Frequency distribution plots
lapply(bout$`Frequency distribution`, plot)
# GVA plots
bout$`GVA plots`
```

describe.augmentedRCBD

Compute Descriptive Statistics from augmentedRCBD Output

# **Description**

 ${\tt describe.augmented RCBD\ computes\ descriptive\ statistics\ from\ the\ adjusted\ means\ in\ an\ object\ of\ class\ augmented RCBD.}$ 

# Usage

```
describe.augmentedRCBD(aug)
```

#### **Arguments**

aug

An object of class augmentedRCBD.

#### **Details**

describe.augmentedRCBD computes the following descriptive statistics from the adjusted means in an object of class augmentedRCBD.

- Count
- Mean
- · Standard deviation
- · Standard error
- Minimum
- Maximum
- Skewness statistic along with p-value from D'Agostino test of skewness (D'Agostino, 1970).
- Kurtosis statistic along with p-value from Anscombe-Glynn test of kurtosis (Anscombe and Glynn, 1983).

#### Value

A list with the following descriptive statistics:

Count The number of treatments/genotypes.

Mean The mean value.
Std.Error The standard error.

Std.Deviation The standard deviation.

Min The minimum value

Max The maximum value

Skewness(statistic)

The skewness estimator.

Skewness(p.value)

The p-value from D'Agostino test of skewness.

Kurtosis(statistic)

The kurtosis estimator.

Kurtosis(p.value)

The p-value from Anscombe-Glynn test of kurtosis.

# References

D'Agostino RB (1970). "Transformation to normality of the null distribution of  $g_1$ ." *Biometrika*, **57**(3), 679–681.

Anscombe FJ, Glynn WJ (1983). "Distribution of the kurtosis statistic  $b_2$  for normal samples." *Biometrika*, **70**(1), 227–234.

#### See Also

augmentedRCBD

#### **Examples**

```
# Example data
blk <- c(rep(1,7),rep(2,6),rep(3,7))
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)
data <- data.frame(blk, trt, y1, y2)</pre>
# Convert block and treatment to factors
data$blk <- as.factor(data$blk)</pre>
data$trt <- as.factor(data$trt)</pre>
# Results for variable y1
out1 <- augmentedRCBD(data$blk, data$trt, data$y1, method.comp = "lsd",</pre>
                       alpha = 0.05, group = TRUE, console = TRUE)
# Results for variable y2
out2 <- augmentedRCBD(data$blk, data$trt, data$y2, method.comp = "lsd",</pre>
                      alpha = 0.05, group = TRUE, console = TRUE)
# Descriptive statistics
describe.augmentedRCBD(out1)
describe.augmentedRCBD(out2)
```

freqdist.augmentedRCBD

Plot Frequency Distribution from augmentedRCBD Output

# Description

freqdist.augmentedRCBD plots frequency distribution from an object of class augmentedRCBD along with the corresponding normal curve and check means with standard errors (if specified by argument highlight.check).

#### Usage

```
freqdist.augmentedRCBD(aug, xlab, highlight.check = TRUE, check.col = "red")
```

# **Arguments**

aug An object of class augmentedRCBD.

xlab The text for x axis label as a character string.

highlight.check

If TRUE, the check means and standard errors are also plotted. Default is TRUE.

check.col The colour(s) to be used to highlight check values in the plot as a character vector. Must be valid colour values in R (named colours, hexadecimal represen-

tation, index of colours [1:8] in default R 'palette()' etc.).

# Value

The frequency distribution plot as a ggplot2 plot grob.

#### See Also

augmented RCBD

#### **Examples**

```
# Example data
blk <- c(rep(1,7),rep(2,6),rep(3,7))
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)
data <- data.frame(blk, trt, y1, y2)</pre>
# Convert block and treatment to factors
data$blk <- as.factor(data$blk)</pre>
data$trt <- as.factor(data$trt)</pre>
# Results for variable y1
out1 <- augmentedRCBD(data$blk, data$trt, data$y1, method.comp = "lsd",</pre>
                       alpha = 0.05, group = TRUE, console = TRUE)
# Results for variable y2
out2 <- augmentedRCBD(data$blk, data$trt, data$y2, method.comp = "lsd",</pre>
                      alpha = 0.05, group = TRUE, console = TRUE)
# Frequency distribution plots
freq1 <- freqdist.augmentedRCBD(out1, xlab = "Trait 1")</pre>
class(freq1)
plot(freq1)
freq2 <- freqdist.augmentedRCBD(out2, xlab = "Trait 2")</pre>
plot(freq2)
# Change check colours
colset <- c("red3", "green4", "purple3", "darkorange3")</pre>
freq1 <- freqdist.augmentedRCBD(out1, xlab = "Trait 1", check.col = colset)</pre>
plot(freq1)
freq2 <- freqdist.augmentedRCBD(out2, xlab = "Trait 2", check.col = colset)</pre>
plot(freq2)
# Without checks highlighted
freq1 <- freqdist.augmentedRCBD(out1, xlab = "Trait 1",</pre>
                                  highlight.check = FALSE)
plot(freq1)
freq2 <- freqdist.augmentedRCBD(out2, xlab = "Trait 2",</pre>
                                  highlight.check = FALSE)
plot(freq2)
```

gva.augmentedRCBD

Perform Genetic Variability Analysis on augmentedRCBD Output

#### **Description**

gva.augmentedRCBD performs genetic variability analysis on an object of class augmentedRCBD.

#### Usage

```
gva.augmentedRCBD(aug, k = 2.063)
```

### **Arguments**

aug An object of class augmentedRCBD.

k The standardized selection differential or selection intensity. Default is 2.063

for 5% selection proportion (see **Details**).

#### **Details**

gva.augmentedRCBD performs genetic variability analysis from the ANOVA results in an object of class augmentedRCBD and computes several variability estimates.

The phenotypic, genotypic and environmental variance ( $\sigma_p^2$ ,  $\sigma_g^2$  and  $\sigma_e^2$ ) are obtained from the ANOVA tables according to the expected value of mean square described by Federer and Searle (1976) as follows:

 $\sigma_p^2$  = Mean sum of squares of test treatments

 $\sigma_e^2 =$  Mean sum of squares of residuals

$$\sigma_q^2 = \sigma_p^2 - \sigma_e^2$$

Phenotypic and genotypic coefficients of variation (PCV and GCV) are estimated according to Burton (1951, 1952) as follows:

$$PCV = \frac{\sigma_p^2}{\sqrt{\overline{x}}} \times 100$$

$$GCV = \frac{\sigma_g^2}{\sqrt{\overline{x}}} \times 100$$

Where  $\overline{x}$  is the mean.

The estimates of PCV and GCV are categorised according to Sivasubramanian and Madhavamenon (1978) as follows:

CV (%)	Category
x < 10	Low
$10 \le x < 20$	Medium
$\geq 20$	High

The broad-sense heritability  $(H^2)$  is calculated according to method of Lush (1940) as follows:

$$H^2 = \frac{\sigma_g^2}{\sigma_p^2}$$

The estimates of broad-sense heritability  $(H^2)$  are categorised according to Robinson (1966) as follows:

$H^2$	Category
x < 30	Low
$30 \le x < 60$	Medium
$\geq 60$	High

Genetic advance (GA) and genetic advance as per cent of mean (GAM) are estimated and cate-

gorised according to Johnson et al., (1955) as follows:

$$GA = k \times \sigma_g \times \frac{H^2}{100}$$

Where the constant k is the standardized selection differential or selection intensity. The value of k at 5% proportion selected is 2.063. Values of k at other selected proportions are available in Appendix Table A of Falconer and Mackay (1996).

$$GAM = \frac{GA}{\overline{x}} \times 100$$

GAM	Category
x < 10	Low
$10 \le x < 20$	Medium
$\geq 20$	High

# Value

A list with the following descriptive statistics:

Mean	The mean value.
PV	Phenotyic variance.
GV	Genotyipc variance.
EV	Environmental variance.
GCV	Genotypic coefficient of variation
GCV category	The GCV category according to Sivasubramaniam and Madhavamenon (1973).
PCV	Phenotypic coefficient of variation
PCV category	The PCV category according to Sivasubramaniam and Madhavamenon (1973).
ECV	Environmental coefficient of variation
hBS	The broad-sense heritability $(H^2)$ (Lush 1940).
hBS category	The $H^2$ category according to Robinson (1966).
GA	Genetic advance (Johnson et al. 1955).
GAM	Genetic advance as per cent of mean (Johnson et al. 1955).
GAM category	The GAM category according to Johnson et al. (1955).

# Note

Genetic variability analysis needs to be performed only if the sum of squares of "Treatment: Test" are significant.

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

# References

Lush JL (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**(1), 293–301.

Burton GW (1951). "Quantitative inheritance in pearl millet (*Pennisetum glaucum*)." Agronomy Journal, **43**(9), 409–417.

Burton GW (1952). "Qualitative inheritance in grasses. Vol. 1." In *Proceedings of the 6th International Grassland Congress, Pennsylvania State College*, 17–23.

Johnson HW, Robinson HF, Comstock RE (1955). "Estimates of genetic and environmental variability in soybeans." *Agronomy journal*, **47**(7), 314–318.

Robinson HF (1966). "Quantitative genetics in relation to breeding on centennial of Mendelism." *Indian Journal of Genetics and Plant Breeding*, 171.

Dudley JW, Moll RH (1969). "Interpretation and use of estimates of heritability and genetic variances in plant breeding." *Crop Science*, **9**(3), 257–262.

Sivasubramaniam S, Madhavamenon P (1973). "Genotypic and phenotypic variability in rice." *The Madras Agricultural Journal*, **60**(9-13), 1093–1096.

Federer WT, Searle SR (1976). "Model Considerations and Variance Component Estimation in Augmented Completely Randomized and Randomized Complete Blocks Designs-Preliminary Version." Technical Report BU-592-M, Cornell University, New York.

Falconer DS, Mackay TFC (1996). *Introduction to quantitative genetics*. Pearson/Prenctice Hall, New York, NY.

#### See Also

augmentedRCBD

# **Examples**

```
# Example data
blk <- c(rep(1,7), rep(2,6), rep(3,7))
trt <- c(1, 2, 3, 4, 7, 11, 12, 1, 2, 3, 4, 5, 9, 1, 2, 3, 4, 8, 6, 10)
y1 \leftarrow 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)
data <- data.frame(blk, trt, y1, y2)</pre>
# Convert block and treatment to factors
data$blk <- as.factor(data$blk)</pre>
data$trt <- as.factor(data$trt)</pre>
# Results for variable y1
out1 <- augmentedRCBD(data$blk, data$trt, data$y1, method.comp = "lsd",
                       alpha = 0.05, group = TRUE, console = TRUE)
# Results for variable v2
out2 <- augmentedRCBD(data$blk, data$trt, data$y2, method.comp = "lsd",
                      alpha = 0.05, group = TRUE, console = TRUE)
# Genetic variability analysis
gva.augmentedRCBD(out1)
gva.augmentedRCBD(out2)
```

print.augmentedRCBD

Prints summary of augmentedRCBD object

# Description

print.augmentedRCBD prints to console the summary of an object of class augmentedRCBD including the augmented design details, ANOVA (Treatment adjusted), ANOVA (Block adjusted), treatment means, coefficient of variation, overall adjusted mean, critical differences and standard errors. The treatment/genotype groups along with the grouping method are also printed if they were computed.

# Usage

```
## S3 method for class 'augmentedRCBD'
print(x, ...)
```

# **Arguments**

x An object of class augmentedRCBD.
... Unused

# See Also

augmentedRCBD

print.augmentedRCBD.bulk

Prints summary of augmentedRCBD.bulk object

# Description

print.augmentedRCBD.bulk prints to console the summary of an object of class augmentedRCBD.bulk including the augmented design details, trait-wise mean sum of squares from ANOVA (Treatment adjusted) and ANOVA (Block adjusted), adjusted means, coefficient of variation, overall adjusted means critical differences, standard errors, descriptive statistics, frequency distribution plots, genetic variability statistics and plots of genetic variability parameters.

### Usage

```
## S3 method for class 'augmentedRCBD.bulk' print(x, ...)
```

# Arguments

x An object of class augmentedRCBD.bulk.... Unused

#### See Also

augmentedRCBD.bulk

report.augmentedRCBD Generate MS Word Report from augmentedRCBD Output

#### **Description**

report.augmentedRCBD generates a tidy report from an object of class augmentedRCBD as docx MS word file using the officer package.

# Usage

```
report.augmentedRCBD(aug, target)
```

#### **Arguments**

aug An object of class augmentedRCBD.
target The path to the docx file to be created.

#### See Also

```
officer, flextable
```

# **Examples**

 $report.augmented RCBD (out, \ file.path (tempdir(), \ "augmented RCBD \ output.docx"))$ 

```
report.augmentedRCBD.bulk
```

Generate MS Word Report from augmentedRCBD.bulk Output

# **Description**

report.augmentedRCBD.bulk generates a tidy report from an object of class augmentedRCBD.bulk as docx MS word file using the officer package.

#### Usage

```
report.augmentedRCBD.bulk(aug.bulk, target)
```

# **Arguments**

aug.bulk An object of class augmentedRCBD.bulk.target The path to the docx file to be created.

# See Also

```
officer, flextable augmentedRCBD.bulk
```

# **Examples**

```
# Example data
blk <- c(rep(1,7), rep(2,6), rep(3,7))
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)
dataf <- data.frame(blk, trt, y1, y2)</pre>
bout <- augmentedRCBD.bulk(data = dataf, block = "blk",</pre>
                           treatment = "trt", traits = c("y1", "y2"),
                           checks = NULL, alpha = 0.05, describe = TRUE,
                           freqdist = TRUE, gva = TRUE,
                           check.col = c("brown", "darkcyan",
                                          "forestgreen", "purple"),
                           console = FALSE)
report.augmentedRCBD.bulk(bout, file.path(tempdir(),
                           "augmentedRCBD bulk output.docx"))
```

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