

Package ‘aridagri’

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Type Package

Title Comprehensive Statistical Tools for Agricultural Research

Version 2.0.4

Description A comprehensive suite of statistical and analytical tools for agricultural research. Includes complete analysis of variance (ANOVA) functions for all experimental designs: Completely Randomized Design (CRD), Randomized Block Design (RBD), Pooled RBD, Split Plot with all variations, Split-Split Plot, Strip Plot, Latin Square, Factorial, Augmented, and Alpha Lattice, with proper error terms and comprehensive Standard Error (SE) and Critical Difference (CD) calculations. Features multiple post-hoc tests: Least Significant Difference (LSD), Duncan Multiple Range Test (DMRT), Tukey Honestly Significant Difference (HSD), Student-Newman-Keuls (SNK), Scheffe, Bonferroni, and Dunnett, along with assumption checking and publication-ready output. Advanced methods include stability analysis using Eberhart-Russell regression, Additive Main Effects and Multiplicative Interaction (AMMI), Finlay-Wilkinson regression, Shukla stability variance, Wricke ecovalence, Coefficient of Variation (CV), and Cultivar Superiority Index as described in Eberhart and Russell (1966) [doi:10.2135/cropsci1966.0011183X000600010011x](https://doi.org/10.2135/cropsci1966.0011183X000600010011x). Thermal indices include Growing Degree Days (GDD), Heliothermal Units (HTU), Photothermal Units (PTU), and Heat Use Efficiency (HUE). Crop growth analysis covers Crop Growth Rate (CGR), Relative Growth Rate (RGR), Net Assimilation Rate (NAR), and Leaf Area Index (LAI). Also provides harvest index, yield gap analysis, economic efficiency indices (Benefit-Cost ratio), nutrient use efficiency calculations, correlation matrix, Principal Component Analysis (PCA), path analysis, and Structural Equation Modeling (SEM). Statistical methods follow Gomez and Gomez (1984, ISBN:0471870927) and Panse and Sukhatme (1985, ISBN:8170271169).

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aridagri-package	<i>aridagri: Comprehensive Statistical Tools for Agricultural Research</i>
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Description

The aridagri package provides comprehensive statistical and analytical tools for agricultural research. It includes complete ANOVA functions for all experimental designs, multiple post-hoc tests, stability analysis methods, thermal indices, crop growth analysis, and advanced statistical methods.

Experimental Design ANOVA Functions

- [anova_crd](#): Completely Randomized Design
- [anova_rbd](#): Randomized Block Design
- [anova_rbd_pooled](#): Pooled RBD (Multi-Environment)
- [anova_latin](#): Latin Square Design
- [anova_factorial](#): Two-Factor Factorial
- [anova_factorial_3way](#): Three-Factor Factorial
- [anova_spd](#): Split Plot Design
- [anova_sspd](#): Split-Split Plot Design
- [anova_strip](#): Strip Plot Design
- [anova_augmented](#): Augmented Block Design
- [anova_alpha_lattice](#): Alpha Lattice Design

Post-Hoc Tests

- [perform_posthoc](#): Multiple comparison tests (LSD, Duncan, Tukey, SNK, Scheffe, Bonferroni, Dunnett)
- [check_assumptions](#): ANOVA assumption checking

Agronomic Analysis Functions

- [stability_analysis](#): Multi-method stability analysis (Eberhart-Russell, AMMI, Finlay-Wilkinson, Shukla, Wricke, CV, Superiority)
- [thermal_indices](#): GDD, HTU, PTU, Heat Use Efficiency
- [crop_growth_analysis](#): CGR, RGR, NAR, LAI
- [harvest_index](#): Harvest index and partitioning
- [yield_gap_analysis](#): Yield gap calculations
- [economic_indices](#): B:C ratio, net returns

Statistical Functions

- [correlation_analysis](#): Correlation matrix with significance
- [pca_analysis](#): Principal component analysis
- [path_analysis](#): Path coefficient analysis
- [sem_analysis](#): Structural equation modeling

Nutrient Analysis Functions

- [nue_calculate](#): Nutrient use efficiency calculations
- [nutrient_response](#): Response curve analysis
- [economic_analysis](#): Economic viability assessment

Author(s)

Lalit Kumar Rolaniya, Ram Lal Jat, Monika Punia, Raja Ram Choudhary

See Also

Useful links:

- <https://github.com/lalitrolaniya/aridagri>
- Report bugs at <https://github.com/lalitrolaniya/aridagri/issues>

anova_alpha_lattice *Alpha Lattice Design ANOVA*

Description

Performs ANOVA for Alpha Lattice (Resolvable Incomplete Block) Design.

Usage

```
anova_alpha_lattice(  
  data,  
  response,  
  genotype,  
  replication,  
  block,  
  verbose = TRUE  
)
```

Arguments

data	Data frame containing the data
response	Name of response variable
genotype	Name of genotype column
replication	Name of replication column
block	Name of incomplete block column (nested within replication)
verbose	Logical. If TRUE (default), prints formatted output to console.

Value

List containing ANOVA results

Author(s)

Lalit Kumar Rolaniya, Ram Lal Jat, Monika Punia, Raja Ram Choudhary

Examples

```
data <- expand.grid(rep = 1:2, block = 1:5, entry = 1:4)  
data$genotype <- paste0("G", 1:nrow(data))  
data$yield <- rnorm(nrow(data), 1200, 150)
```

anova_augmented	<i>Augmented Block Design ANOVA</i>
-----------------	-------------------------------------

Description

Performs ANOVA for Augmented Randomized Block Design where checks are replicated and test entries appear once.

Usage

```
anova_augmented(data, response, genotype, block, check_names, verbose = TRUE)
```

Arguments

data	Data frame containing the data
response	Name of response variable
genotype	Name of genotype/entry column
block	Name of block column
check_names	Vector of check variety names
verbose	Logical. If TRUE (default), prints formatted output to console.

Value

List containing ANOVA results and adjusted means

Author(s)

Lalit Kumar Rolaniya, Ram Lal Jat, Monika Punia, Raja Ram Choudhary

Examples

```
data <- data.frame(
  block = rep(1:5, each = 7),
  genotype = c(rep(c("C1","C2","C3"), 5), paste0("T", 1:20)),
  yield = rnorm(35, 1200, 150)
)
# Note: This is simplified example
```

anova_crd

=====

COMPREHENSIVE EXPERIMENTAL DESIGN ANALYSIS FUNCTIONS Package: aridagri Author: Lalit Kumar Rolaniya ICAR-Indian Institute of Pulses Research, Regional Centre, Bikaner

=====

Completely Randomized Design (CRD) ANOVA

Description

Performs complete ANOVA for Completely Randomized Design with post-hoc tests, assumptions checking, and publication-ready output.

Usage

```
anova_crd(
  data,
  response,
  treatment,
  posthoc = "lsd",
```

```

    alpha = 0.05,
    verbose = TRUE
  )

```

Arguments

data	Data frame containing the experimental data
response	Name of the response variable (as string)
treatment	Name of treatment factor
posthoc	Post-hoc test: "lsd", "duncan", "tukey", "snk", "scheffe", or "all"
alpha	Significance level (default 0.05)
verbose	Logical. If TRUE (default), prints formatted output to console.

Value

List containing ANOVA table, means, post-hoc results, and diagnostics

Author(s)

Lalit Kumar Rolaniya, ICAR-IIPR, Bikaner

Examples

```

data <- data.frame(
  treatment = rep(c("T1", "T2", "T3", "T4"), each = 5),
  yield = c(rnorm(5, 1200, 50), rnorm(5, 1350, 60),
            rnorm(5, 1100, 55), rnorm(5, 1450, 65))
)
anova_crd(data, response = "yield", treatment = "treatment", posthoc = "all")

```

anova_factorial	<i>Factorial ANOVA (Two-Factor)</i>
-----------------	-------------------------------------

Description

Performs two-factor factorial ANOVA with interaction analysis.

Usage

```

anova_factorial(
  data,
  response,
  factor1,
  factor2,
  replication = NULL,
  posthoc = "lsd",

```

```

    alpha = 0.05,
    verbose = TRUE
  )

```

Arguments

data	Data frame containing the data
response	Name of the response variable
factor1	Name of first factor (A)
factor2	Name of second factor (B)
replication	Name of replication factor (optional)
posthoc	Post-hoc test method
alpha	Significance level
verbose	Logical. If TRUE (default), prints formatted output to console.

Value

List containing ANOVA results

Author(s)

Lalit Kumar Rolaniya, ICAR-IIPR, Bikaner

Examples

```

data <- expand.grid(
  rep = 1:4,
  nitrogen = c("N0", "N40", "N80", "N120"),
  phosphorus = c("P0", "P30", "P60")
)
data$yield <- rnorm(nrow(data), 1200, 150)

anova_factorial(data, response = "yield",
  factor1 = "nitrogen", factor2 = "phosphorus",
  replication = "rep")

```

anova_factorial_3way *Three-Factor Factorial ANOVA*

Description

Performs three-factor factorial ANOVA with all interactions.

Usage

```
anova_factorial_3way(  
  data,  
  response,  
  factor1,  
  factor2,  
  factor3,  
  replication = NULL,  
  alpha = 0.05,  
  verbose = TRUE  
)
```

Arguments

data	Data frame containing the data
response	Name of the response variable
factor1	Name of first factor (A)
factor2	Name of second factor (B)
factor3	Name of third factor (C)
replication	Name of replication factor (optional)
alpha	Significance level
verbose	Logical. If TRUE (default), prints formatted output to console.

Value

List containing ANOVA results

Author(s)

Lalit Kumar Rolaniya, Ram Lal Jat, Monika Punia, Raja Ram Choudhary

Examples

```
data <- expand.grid(rep = 1:3, A = c("A1", "A2"), B = c("B1", "B2"), C = c("C1", "C2"))  
data$yield <- rnorm(nrow(data), 1200, 150)  
anova_factorial_3way(data, "yield", "A", "B", "C", "rep")
```

anova_latin

*Latin Square Design ANOVA***Description**

Performs ANOVA for Latin Square Design with row and column blocking.

Usage

```
anova_latin(
  data,
  response,
  treatment,
  row,
  column,
  posthoc = "lsd",
  alpha = 0.05,
  verbose = TRUE
)
```

Arguments

data	Data frame containing the data
response	Name of the response variable
treatment	Name of treatment factor
row	Name of row factor
column	Name of column factor
posthoc	Post-hoc test method
alpha	Significance level
verbose	Logical. If TRUE (default), prints formatted output to console.

Value

List containing ANOVA results

Author(s)

Lalit Kumar Rolaniya, ICAR-IIPR, Bikaner

Examples

```
# 5x5 Latin Square
data <- data.frame(
  row = rep(1:5, each = 5),
  col = rep(1:5, 5),
  treatment = c("A", "B", "C", "D", "E", "B", "C", "D", "E", "A",
```

```

      "C","D","E","A","B", "D","E","A","B","C",
      "E","A","B","C","D"),
  yield = rnorm(25, 1200, 100)
)
anova_latin(data, response = "yield", treatment = "treatment",
            row = "row", column = "col")

```

anova_rbd

*Randomized Block Design (RBD) ANOVA***Description**

Performs complete ANOVA for Randomized Block Design (RCBD) with post-hoc tests, assumptions checking, and publication-ready output.

Usage

```

anova_rbd(
  data,
  response,
  treatment,
  block,
  posthoc = "lsd",
  alpha = 0.05,
  verbose = TRUE
)

```

Arguments

data	Data frame containing the experimental data
response	Name of the response variable
treatment	Name of treatment factor
block	Name of block/replication factor
posthoc	Post-hoc test: "lsd", "duncan", "tukey", "snk", "scheffe", or "all"
alpha	Significance level (default 0.05)
verbose	Logical. If TRUE (default), prints formatted output to console.

Value

List containing ANOVA table, means, post-hoc results

Author(s)

Lalit Kumar Rolaniya, ICAR-IIPR, Bikaner

Examples

```
data <- data.frame(
  rep = rep(1:4, each = 5),
  treatment = rep(c("T1", "T2", "T3", "T4", "T5"), 4),
  yield = c(rnorm(5, 1200, 50), rnorm(5, 1250, 55),
            rnorm(5, 1180, 45), rnorm(5, 1270, 60))
)
anova_rbd(data, response = "yield", treatment = "treatment",
          block = "rep", posthoc = "duncan")
```

anova_rbd_pooled

*Pooled Analysis of RBD Experiments (Multi-Environment/Year)***Description**

Performs pooled ANOVA for RBD experiments conducted across multiple environments, years, or locations. Tests homogeneity of error variances using Bartlett's test before pooling.

Usage

```
anova_rbd_pooled(
  data,
  response,
  treatment,
  environment,
  block,
  posthoc = "lsd",
  alpha = 0.05,
  verbose = TRUE
)
```

Arguments

data	Data frame containing combined data from all environments
response	Name of the response variable
treatment	Name of treatment factor
environment	Name of environment/year/location factor
block	Name of block factor (nested within environment)
posthoc	Post-hoc test method
alpha	Significance level
verbose	Logical. If TRUE (default), prints formatted output to console.

Value

List containing pooled ANOVA, individual ANOVAs, and interaction analysis

Author(s)

Lalit Kumar Rolaniya, ICAR-IIPR, Bikaner

Examples

```
# Data from 3 years
data <- data.frame(
  year = rep(c("Y1", "Y2", "Y3"), each = 20),
  rep = rep(rep(1:4, each = 5), 3),
  treatment = rep(c("T1", "T2", "T3", "T4", "T5"), 12),
  yield = rnorm(60, 1200, 150)
)
anova_rbd_pooled(data, response = "yield", treatment = "treatment",
  environment = "year", block = "rep")
```

anova_spd

```
=====
SPLIT PLOT DESIGN ANALYSIS FUNCTIONS (All Varia-
tions) Package: aridagri Author: Lalit Kumar Rolaniya ICAR-
Indian Institute of Pulses Research, Regional Centre, Bikaner
=====
Split Plot Design ANOVA (Standard)
```

Description

Performs complete ANOVA for Split Plot Design with proper error terms for main plot and sub-plot factors. Includes all standard post-hoc comparisons.

Usage

```
anova_spd(
  data,
  response,
  main_plot,
  sub_plot,
  replication,
  posthoc = "lsd",
  alpha = 0.05,
  verbose = TRUE
)
```

Arguments

data	Data frame containing the experimental data
response	Name of the response variable
main_plot	Name of main plot factor

sub_plot	Name of sub-plot factor
replication	Name of replication/block factor
posthoc	Post-hoc test method
alpha	Significance level
verbose	Logical. If TRUE (default), prints formatted output to console.

Value

List containing ANOVA table, means, and post-hoc results

Author(s)

Lalit Kumar Rolaniya, ICAR-IIPR, Bikaner

Examples

```
data <- data.frame(
  rep = rep(1:3, each = 12),
  irrigation = rep(rep(c("I1", "I2", "I3"), each = 4), 3),
  variety = rep(c("V1", "V2", "V3", "V4"), 9),
  yield = rnorm(36, 1200, 150)
)
anova_spd(data, response = "yield", main_plot = "irrigation",
           sub_plot = "variety", replication = "rep")
```

anova_spd_ab_cd	<i>Split Plot Design with (AB) Main and (CD) Sub</i>
-----------------	--

Description

Performs ANOVA for Split Plot Design where main plot contains factorial combination of AB and sub-plot contains factorial combination of CD. Complex design for multi-factor experiments.

Usage

```
anova_spd_ab_cd(
  data,
  response,
  main_factor1,
  main_factor2,
  sub_factor1,
  sub_factor2,
  replication,
  verbose = TRUE
)
```

Arguments

data	Data frame containing the experimental data
response	Name of the response variable
main_factor1	First factor in main plot (A)
main_factor2	Second factor in main plot (B)
sub_factor1	First factor in sub-plot (C)
sub_factor2	Second factor in sub-plot (D)
replication	Name of replication factor
verbose	Logical. If TRUE (default), prints formatted output to console.

Value

List containing ANOVA table and means

Author(s)

Lalit Kumar Rolaniya, ICAR-IIPR, Bikaner

Examples

```
# Example: Irrigation Tillage (main), Variety Nitrogen (sub)
data <- expand.grid(
  rep = 1:3,
  irrigation = c("I1", "I2"),
  tillage = c("CT", "ZT"),
  variety = c("V1", "V2"),
  nitrogen = c("N1", "N2", "N3")
)
data$yield <- rnorm(nrow(data), 1200, 150)

anova_spd_ab_cd(data, response = "yield",
  main_factor1 = "irrigation", main_factor2 = "tillage",
  sub_factor1 = "variety", sub_factor2 = "nitrogen",
  replication = "rep")
```

anova_spd_ab_main

Split Plot Design with AB in Main Plot

Description

Performs ANOVA for Split Plot Design where main plot contains factorial combination of two factors (AB) and sub-plot contains factor C. Common in irrigation variety as main plot and nitrogen as sub-plot.

Usage

```
anova_spd_ab_main(  
  data,  
  response,  
  main_factor1,  
  main_factor2,  
  sub_plot,  
  replication,  
  posthoc = "lsd",  
  alpha = 0.05,  
  verbose = TRUE  
)
```

Arguments

data	Data frame containing the experimental data
response	Name of the response variable
main_factor1	First factor in main plot (A)
main_factor2	Second factor in main plot (B)
sub_plot	Sub-plot factor (C)
replication	Name of replication factor
posthoc	Post-hoc test method
alpha	Significance level
verbose	Logical. If TRUE (default), prints formatted output to console.

Details

Design structure:

- Main Plot: A B factorial
- Sub-Plot: C
- Error (a): For testing A, B, and AB
- Error (b): For testing C and all interactions with C

Value

List containing ANOVA table and means

Author(s)

Lalit Kumar Rolaniya, ICAR-IIPR, Bikaner

Examples

```
data <- data.frame(
  rep = rep(1:3, each = 24),
  irrigation = rep(rep(c("I1", "I2"), each = 12), 3),
  variety = rep(rep(c("V1", "V2", "V3"), each = 4), 6),
  nitrogen = rep(c("N0", "N1", "N2", "N3"), 18),
  yield = rnorm(72, 1200, 150)
)
anova_spd_ab_main(data, response = "yield",
  main_factor1 = "irrigation", main_factor2 = "variety",
  sub_plot = "nitrogen", replication = "rep")
```

anova_spd_c_main_ab_sub

Split Plot Design with C in Main Plot, AB in Sub-Plot

Description

Performs ANOVA for Split Plot Design where main plot contains single factor C and sub-plot contains factorial combination of AB.

Usage

```
anova_spd_c_main_ab_sub(
  data,
  response,
  main_plot,
  sub_factor1,
  sub_factor2,
  replication,
  posthoc = "lsd",
  alpha = 0.05,
  verbose = TRUE
)
```

Arguments

data	Data frame containing the experimental data
response	Name of the response variable
main_plot	Main plot factor (C)
sub_factor1	First factor in sub-plot (A)
sub_factor2	Second factor in sub-plot (B)
replication	Name of replication factor
posthoc	Post-hoc test method
alpha	Significance level
verbose	Logical. If TRUE (default), prints formatted output to console.

Value

List containing ANOVA table and means

Author(s)

Lalit Kumar Rolaniya, ICAR-IIPR, Bikaner

Examples

```
data <- data.frame(
  rep = rep(1:3, each = 24),
  irrigation = rep(rep(c("I1", "I2", "I3"), each = 8), 3),
  variety = rep(rep(c("V1", "V2"), each = 4), 9),
  nitrogen = rep(c("N1", "N2", "N3", "N4"), 18),
  yield = rnorm(72, 1200, 150)
)
anova_spd_c_main_ab_sub(data, response = "yield",
  main_plot = "irrigation",
  sub_factor1 = "variety", sub_factor2 = "nitrogen",
  replication = "rep")
```

anova_spd_pooled

Pooled Split Plot Design Analysis

Description

Performs pooled analysis of Split Plot Design experiments conducted across multiple environments/years/locations.

Usage

```
anova_spd_pooled(
  data,
  response,
  main_plot,
  sub_plot,
  environment,
  replication,
  verbose = TRUE
)
```

Arguments

data	Data frame containing combined data
response	Name of the response variable
main_plot	Name of main plot factor

sub_plot	Name of sub-plot factor
environment	Name of environment factor
replication	Name of replication factor (nested within environment)
verbose	Logical. If TRUE (default), prints formatted output to console.

Value

List containing pooled ANOVA and component analyses

Author(s)

Lalit Kumar Rolaniya, ICAR-IIPR, Bikaner

Examples

```
data <- expand.grid(
  year = c("Y1", "Y2", "Y3"),
  rep = 1:3,
  irrigation = c("I1", "I2", "I3"),
  variety = c("V1", "V2", "V3", "V4")
)
data$yield <- rnorm(nrow(data), 1200, 180)

anova_sspd_pooled(data, response = "yield", main_plot = "irrigation",
  sub_plot = "variety", environment = "year", replication = "rep")
```

anova_sspd	<i>Split-Split Plot Design ANOVA</i>
------------	--------------------------------------

Description

Performs complete ANOVA for Split-Split Plot Design with proper error terms for main plot, sub-plot, and sub-sub-plot factors. Generates publication-ready ANOVA table with significance levels.

Usage

```
anova_sspd(
  data,
  response,
  main_plot,
  sub_plot,
  sub_sub_plot,
  replication,
  verbose = TRUE
)
```

Arguments

data	Data frame containing the experimental data
response	Name of the response variable (as string)
main_plot	Name of main plot factor
sub_plot	Name of sub-plot factor
sub_sub_plot	Name of sub-sub-plot factor
replication	Name of replication/block factor
verbose	Logical. If TRUE (default), prints formatted output to console.

Value

List containing ANOVA table, means, and significance tests

Author(s)

Lalit Kumar Rolaniya, ICAR-IIPR, Bikaner

Examples

```
# Example with sample data
data <- expand.grid(rep=1:3, A=c('A1','A2'), B=c('B1','B2'), C=c('C1','C2'))
data$yield <- rnorm(24, 1200, 150)
anova_sspd(data, response='yield', main_plot='A', sub_plot='B',
            sub_sub_plot='C', replication='rep')
```

anova_sspd_pooled

Pooled Split-Split Plot Design ANOVA

Description

Performs pooled ANOVA for SSPD experiments across multiple environments.

Usage

```
anova_sspd_pooled(
  data,
  response,
  main_plot,
  sub_plot,
  sub_sub_plot,
  environment,
  replication,
  verbose = TRUE
)
```

Arguments

data	Data frame containing combined data
response	Name of the response variable
main_plot	Name of main plot factor
sub_plot	Name of sub-plot factor
sub_sub_plot	Name of sub-sub-plot factor
environment	Name of environment factor
replication	Name of replication factor
verbose	Logical. If TRUE (default), prints formatted output to console.

Value

List containing pooled ANOVA results

Author(s)

Lalit Kumar Rolaniya, Ram Lal Jat, Monika Punia, Raja Ram Choudhary

Examples

```
data <- expand.grid(env = c("E1", "E2"), rep = 1:3, A = c("A1", "A2"),
                   B = c("B1", "B2"), C = c("C1", "C2"))
data$yield <- rnorm(nrow(data), 1200, 150)
anova_sspd_pooled(data, "yield", "A", "B", "C", "env", "rep")
```

anova_strip	=====
	ADDITIONAL EXPERIMENTAL DESIGN FUNCTIONS
	Package: aridagri Authors: Lalit Kumar Rolaniya,
	Ram Lal Jat, Monika Punia, Raja Ram Choudhary
	=====
	Strip Plot Design ANOVA

Description

Performs ANOVA for Strip Plot (Strip-Split) Design where two factors are applied in horizontal and vertical strips.

Usage

```
anova_strip(  
  data,  
  response,  
  horizontal_factor,  
  vertical_factor,  
  replication,  
  alpha = 0.05,  
  verbose = TRUE  
)
```

Arguments

data	Data frame containing the experimental data
response	Name of the response variable
horizontal_factor	Factor applied in horizontal strips (A)
vertical_factor	Factor applied in vertical strips (B)
replication	Name of replication factor
alpha	Significance level
verbose	Logical. If TRUE (default), prints formatted output to console.

Value

List containing ANOVA results

Author(s)

Lalit Kumar Rolaniya, Ram Lal Jat, Monika Punia, Raja Ram Choudhary

Examples

```
data <- expand.grid(  
  rep = 1:4,  
  irrigation = c("I1", "I2", "I3"),  
  tillage = c("CT", "MT", "ZT")  
)  
data$yield <- rnorm(nrow(data), 1200, 150)  
anova_strip(data, response = "yield", horizontal_factor = "irrigation",  
            vertical_factor = "tillage", replication = "rep")
```

arid_plot

Visualization Functions for aridagri

Description

Generate publication-quality plots for aridagri analyses.

Usage

```
arid_plot(x, type = "bar", ...)
```

Arguments

x	An object from aridagri analysis functions
type	Plot type: "bar", "line", "interaction", "boxplot"
...	Additional arguments passed to plotting functions

Value

No return value; called for its side effect of drawing a plot on the current graphics device. The plot produced depends on the class of x: a correlation heatmap for `aridagri_correlation` objects, a bar chart of treatment means for `aridagri_anova` objects, and a bar chart of the integrated stability ranking for `aridagri_stability` objects. Uses only base graphics, so no additional packages are required.

Author(s)

Lalit Kumar Rolaniya, Ram Lal Jat, Monika Punia, Raja Ram Choudhary

Examples

```
df <- data.frame(  
  yield = c(1200, 1350, 1100, 1450, 1280),  
  wue = c(4.2, 4.8, 3.9, 5.1, 4.5),  
  protein = c(22.1, 23.5, 21.8, 24.2, 22.9)  
)  
result <- correlation_analysis(df, plot = FALSE)  
arid_plot(result)
```

check_assumptions	<i>Check ANOVA Assumptions</i>
-------------------	--------------------------------

Description

Tests assumptions of ANOVA: normality and homogeneity of variances.

Usage

```
check_assumptions(model, verbose = TRUE)
```

Arguments

model	ANOVA model object
verbose	Logical. If TRUE (default), prints formatted output to console.

Value

List containing diagnostic test results

correlation_analysis	<i>Correlation Analysis with Significance</i>
----------------------	---

Description

Computes correlation matrix with significance levels and generates publication-ready correlation table and plot.

Usage

```
correlation_analysis(  
  data,  
  method = "pearson",  
  plot = TRUE,  
  digits = 3,  
  verbose = TRUE  
)
```

Arguments

data	Data frame with numeric variables
method	Correlation method: "pearson", "spearman", or "kendall"
plot	Logical, whether to generate correlation plot
digits	Number of decimal places
verbose	Logical. If TRUE (default), prints formatted output to console.

Value

List with correlation matrix and significance matrix

Author(s)

Lalit Kumar Rolaniya, ICAR-IIPR, Bikaner

Examples

```
data <- data.frame(  
  yield = c(1200, 1350, 1100, 1450, 1280),  
  wue = c(4.2, 4.8, 3.9, 5.1, 4.5),  
  protein = c(22.1, 23.5, 21.8, 24.2, 22.9)  
)  
correlation_analysis(data)
```

crop_growth_analysis *Crop Growth Analysis (CGR, RGR, NAR, LAI)*

Description

Calculates crop growth parameters from sequential harvest data.

Usage

```
crop_growth_analysis(dry_weight, leaf_area, days, verbose = TRUE)
```

Arguments

dry_weight	Vector of dry matter at different stages
leaf_area	Vector of leaf area at different stages
days	Vector of days after sowing
verbose	Logical. If TRUE (default), prints formatted output to console.

Value

Data frame with growth parameters

Author(s)

Lalit Kumar Rolaniya, Ram Lal Jat, Monika Punia, Raja Ram Choudhary

Examples

```
dry_weight <- c(0.5, 2.1, 8.5, 25, 45, 62, 75)
leaf_area <- c(15, 85, 350, 800, 950, 850, 600)
days <- c(15, 30, 45, 60, 75, 90, 105)
crop_growth_analysis(dry_weight, leaf_area, days)
```

economic_analysis	<i>Economic Analysis for Arid Agriculture</i>
-------------------	---

Description

Calculates economic indicators including Cost of Cultivation, Gross Returns, Net Returns, B:C Ratio, and profitability indices for arid farming systems.

Usage

```
economic_analysis(
  yield,
  price,
  cost_fixed,
  cost_variable,
  byproduct_yield = 0,
  byproduct_price = 0,
  verbose = TRUE
)
```

Arguments

yield	Crop yield (kg/ha)
price	Market price (Rs/kg)
cost_fixed	Fixed costs (Rs/ha)
cost_variable	Variable costs (Rs/ha)
byproduct_yield	Byproduct yield (kg/ha), optional
byproduct_price	Byproduct price (Rs/kg), optional
verbose	Logical. If TRUE (default), prints formatted output to console.

Value

Data frame with economic analysis

Author(s)

Lalit Kumar Rolaniya, ICAR-IIPR, Bikaner

Examples

```
economic_analysis(yield = 1200, price = 65, cost_fixed = 15000, cost_variable = 12000)

# With byproduct
economic_analysis(yield = 1200, price = 65, cost_fixed = 15000,
                  cost_variable = 12000, byproduct_yield = 1800, byproduct_price = 5)
```

economic_indices	<i>Economic Efficiency Indices (B:C Ratio)</i>
------------------	--

Description

Calculates B:C ratio and economic efficiency metrics.

Usage

```
economic_indices(gross_return, total_cost, verbose = TRUE)
```

Arguments

- gross_return Gross returns (Rs/ha)
- total_cost Total cost (Rs/ha)
- verbose Logical. If TRUE (default), prints formatted output to console.

Value

Data frame with economic indices

Author(s)

Lalit Kumar Rolaniya, Ram Lal Jat, Monika Punia, Raja Ram Choudhary

Examples

```
economic_indices(gross_return = c(75000, 82000), total_cost = c(35000, 38000))
```

export_results	<i>Export Results to Publication Format</i>
----------------	---

Description

Exports aridagri analysis results to Excel format for publication.

Usage

```
export_results(x, file, format = "xlsx", digits = 3, verbose = TRUE)
```

Arguments

x	An object from aridagri analysis functions
file	Output file path
format	Output format: "xlsx" (default) or "csv"
digits	Number of decimal places
verbose	Logical. If TRUE (default), prints formatted output to console.

Value

Invisibly returns the file path

Author(s)

Lalit Kumar Rolaniya, Ram Lal Jat, Monika Punia, Raja Ram Choudhary

Examples

```
df <- data.frame(  
  yield = c(1200, 1350, 1100, 1450, 1280),  
  wue = c(4.2, 4.8, 3.9, 5.1, 4.5)  
)  
result <- correlation_analysis(df, plot = FALSE)  
export_results(result, tempfile(fileext = ".xlsx"))
```

harvest_index	<i>Harvest Index Calculation</i>
---------------	----------------------------------

Description

Calculates Harvest Index and related partitioning indices.

Usage

```
harvest_index(grain_yield, straw_yield, verbose = TRUE)
```

Arguments

grain_yield	Grain/economic yield
straw_yield	Straw/stover yield
verbose	Logical. If TRUE (default), prints formatted output to console.

Value

Data frame with harvest indices

Author(s)

Lalit Kumar Rolaniya, Ram Lal Jat, Monika Punia, Raja Ram Choudhary

Examples

```
harvest_index(grain_yield = c(1200, 1350, 1100), straw_yield = c(2400, 2500, 2300))
```

nue_calculate	<i>Nutrient Use Efficiency Calculations</i>
---------------	---

Description

Comprehensive nutrient use efficiency calculations including Agronomic Efficiency (AE), Physiological Efficiency (PE), Apparent Recovery Efficiency (ARE), and Partial Factor Productivity (PFP). Essential for INM research in arid regions.

Usage

```
nue_calculate(
  yield_fertilized,
  yield_control,
  nutrient_applied,
  nutrient_uptake_fert = NULL,
  nutrient_uptake_ctrl = NULL,
  biomass_fert = NULL,
  biomass_ctrl = NULL,
  verbose = TRUE
)
```

Arguments

yield_fertilized	Yield with fertilizer application (kg/ha)
yield_control	Yield in control/unfertilized plot (kg/ha)
nutrient_applied	Amount of nutrient applied (kg/ha)
nutrient_uptake_fert	Nutrient uptake in fertilized plot (kg/ha), optional
nutrient_uptake_ctrl	Nutrient uptake in control plot (kg/ha), optional
biomass_fert	Total biomass in fertilized plot (kg/ha), optional
biomass_ctrl	Total biomass in control plot (kg/ha), optional
verbose	Logical. If TRUE (default), prints formatted output to console.

Details

Efficiency calculations:

- AE (Agronomic Efficiency) = $(\text{Yield_fert} - \text{Yield_ctrl}) / \text{Nutrient_applied}$
- PFP (Partial Factor Productivity) = $\text{Yield_fert} / \text{Nutrient_applied}$
- ARE (Apparent Recovery Efficiency) = $(\text{Uptake_fert} - \text{Uptake_ctrl}) / \text{Nutrient_applied} \times 100$
- PE (Physiological Efficiency) = $(\text{Yield_fert} - \text{Yield_ctrl}) / (\text{Uptake_fert} - \text{Uptake_ctrl})$

Value

Data frame with efficiency indices

Author(s)

Lalit Kumar Rolaniya, ICAR-IIPR, Bikaner

Examples

```
# Basic NUE calculation
nue_calculate(yield_fertilized = 1850, yield_control = 1200, nutrient_applied = 40)

# Complete NUE with uptake data
nue_calculate(
  yield_fertilized = 1850,
  yield_control = 1200,
  nutrient_applied = 40,
  nutrient_uptake_fert = 65,
  nutrient_uptake_ctrl = 35
)
```

nutrient_response	<i>Nutrient Response Curve Analysis</i>
-------------------	---

Description

Fits nutrient response curves using quadratic, linear-plateau, or Mitscherlich models to determine economic optimum dose.

Usage

```
nutrient_response(
  dose,
  yield,
  model = "quadratic",
  price_output = 50,
  price_nutrient = 15,
  verbose = TRUE
)
```

Arguments

dose	Numeric vector of nutrient doses (kg/ha)
yield	Numeric vector of corresponding yields (kg/ha)
model	Model type: "quadratic", "linear_plateau", or "mitscherlich"
price_output	Price of output (Rs/kg)
price_nutrient	Price of nutrient (Rs/kg)
verbose	Logical. If TRUE (default), prints formatted output to console.

Value

List with model parameters and economic optimum

Author(s)

Lalit Kumar Rolaniya, ICAR-IIPR, Bikaner

Examples

```
dose <- c(0, 20, 40, 60, 80, 100)
yield <- c(1100, 1350, 1520, 1610, 1650, 1660)
nutrient_response(dose, yield, model = "quadratic",
                  price_output = 60, price_nutrient = 15)
```

path_analysis	<i>Path Coefficient Analysis</i>
---------------	----------------------------------

Description

Performs path analysis to determine direct and indirect effects of independent variables on a dependent variable. Essential for understanding yield contributing factors.

Usage

```
path_analysis(data, dependent, independent, digits = 4, verbose = TRUE)
```

Arguments

data	Data frame with numeric variables
dependent	Name of dependent variable (e.g., "yield")
independent	Character vector of independent variable names
digits	Number of decimal places
verbose	Logical. If TRUE (default), prints formatted output to console.

Value

List with direct effects, indirect effects, and correlation breakdown

Author(s)

Lalit Kumar Rolaniya, ICAR-IIPR, Bikaner

Examples

```
data <- data.frame(
  yield = c(1200, 1350, 1100, 1450, 1280, 1380, 1220, 1400),
  pods = c(45, 52, 42, 58, 48, 54, 46, 56),
  seeds = c(8.2, 9.1, 7.8, 9.5, 8.5, 9.0, 8.3, 9.3),
  weight = c(32, 35, 30, 38, 33, 36, 31, 37)
)
path_analysis(data, dependent = "yield",
  independent = c("pods", "seeds", "weight"))
```

pca_analysis

*Principal Component Analysis***Description**

Performs PCA with visualization suitable for agricultural research data. Includes scree plot, biplot, and variable contributions.

Usage

```
pca_analysis(data, scale = TRUE, ncp = 5, plot = TRUE, verbose = TRUE)
```

Arguments

data	Data frame with numeric variables
scale	Logical, whether to scale variables (default TRUE)
ncp	Number of components to retain (default 5)
plot	Logical, whether to generate plots
verbose	Logical. If TRUE (default), prints formatted output to console.

Value

PCA results with eigenvalues, loadings, and scores

Author(s)

Lalit Kumar Rolaniya, ICAR-IIPR, Bikaner

Examples

```
data <- data.frame(
  yield = rnorm(30, 1200, 200),
  wue = rnorm(30, 4.5, 0.5),
  protein = rnorm(30, 22, 2),
  biomass = rnorm(30, 3500, 500)
)
pca_analysis(data)
```

perform_posthoc	<i>Perform Post-Hoc Tests</i>
-----------------	-------------------------------

Description

Performs multiple comparison tests after ANOVA.

Usage

```
perform_posthoc(  
  model,  
  data,  
  response,  
  treatment,  
  mse,  
  df_error,  
  posthoc = "lsd",  
  alpha = 0.05,  
  verbose = TRUE  
)
```

Arguments

model	ANOVA model object
data	Data frame
response	Response variable name
treatment	Treatment factor name
mse	Mean square error
df_error	Error degrees of freedom
posthoc	Test type: "lsd", "duncan", "tukey", "snk", "scheffe", or "all"
alpha	Significance level
verbose	Logical. If TRUE (default), prints formatted output to console.

Value

List containing post-hoc test results

print.aridagri_pca	<i>Print Method for PCA Results</i>
--------------------	-------------------------------------

Description

Prints a formatted summary of Principal Component Analysis (PCA) results.

Usage

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

Arguments

x	An object of class 'aridagri_pca' from pca_analysis
...	Additional arguments (currently unused)

Value

No return value, called for side effects. Prints the number of components retained by Kaiser criterion and cumulative variance explained to the console. The input object is returned invisibly.

sem_analysis	<i>Structural Equation Modeling for Field Experiments</i>
--------------	---

Description

Performs SEM analysis for agricultural field experiments. Allows testing of hypothesized causal relationships among variables.

Usage

```
sem_analysis(data, model, plot = TRUE, verbose = TRUE)
```

Arguments

data	Data frame with variables
model	Model specification in lavaan syntax
plot	Logical, whether to generate path diagram
verbose	Logical. If TRUE (default), prints formatted output to console.

Value

SEM results including fit indices and parameter estimates

Author(s)

Lalit Kumar Rolaniya, ICAR-IIPR, Bikaner

Examples

```
if (requireNamespace("lavaan", quietly = TRUE)) {
  set.seed(42)
  n <- 100
  nitrogen <- rnorm(n, 60, 10)
  phosphorus <- rnorm(n, 30, 5)
  yield <- 0.5 * nitrogen + 0.3 * phosphorus + rnorm(n, 0, 5)
  df <- data.frame(yield = yield, nitrogen = nitrogen, phosphorus = phosphorus)
  model <- 'yield ~ nitrogen + phosphorus'
  result <- sem_analysis(df, model, plot = FALSE)
}
```

stability_analysis

=====

*ADVANCED AGRONOMIC STATISTICAL METHODS Pack-
age: aridagri Authors: Lalit Kumar Rolaniya, Ram Lal Jat,
Monika Punia, Raja Ram Choudhary ICAR-Indian Institute of
Pulses Research, Regional Centre, Bikaner ICAR-Indian In-
stitute of Groundnut Research, Regional Research Station, Bikaner*

=====

Stability Analysis for Agronomic Traits (Multiple Methods)

Description

Performs comprehensive stability analysis using multiple established methods for evaluating geno-
type/treatment performance across environments.

Methods included:

- Eberhart & Russell (1966): Regression approach
- AMMI: Additive Main effects and Multiplicative Interaction
- Finlay & Wilkinson (1963): Linear regression on environmental mean
- Shukla (1972): Stability variance
- Wricke (1962): Ecovalence
- Coefficient of Variation: CV-based ranking
- Superiority Index: Lin & Binns (1988)

Usage

```
stability_analysis(  
  data,  
  genotype,  
  environment,  
  replication,  
  trait,  
  method = "all",  
  verbose = TRUE  
)
```

Arguments

data	Data frame with genotype/treatment, environment, replication, and trait
genotype	Name of genotype/treatment column
environment	Name of environment/location/year column
replication	Name of replication column
trait	Name of trait/response variable
method	Method: "eberhart", "ammi", "finlay", "shukla", "wricke", "cv", "superiority", or "all"
verbose	Logical. If TRUE (default), prints formatted output to console.

Value

List containing stability parameters and rankings

Author(s)

Lalit Kumar Rolaniya, Ram Lal Jat, Monika Punia, Raja Ram Choudhary

References

Eberhart, S.A. and Russell, W.A. (1966). Crop Science, 6: 36-40.

Examples

```
data <- expand.grid(  
  variety = paste0("V", 1:10),  
  location = paste0("L", 1:5),  
  rep = 1:3  
)  
data$yield <- rnorm(nrow(data), 1200, 200)  
  
stability_analysis(data, genotype = "variety", environment = "location",  
  replication = "rep", trait = "yield", method = "all")
```

thermal_indices	<i>Growing Degree Days (GDD) and Thermal Indices</i>
-----------------	--

Description

Calculates Growing Degree Days, Helio-thermal Units, Photo-thermal Units, and Heat Use Efficiency from temperature data.

Usage

```
thermal_indices(
  tmax,
  tmin,
  base_temp = 10,
  sunshine_hours = NULL,
  day_length = NULL,
  crop_yield = NULL,
  biomass = NULL,
  verbose = TRUE
)
```

Arguments

tmax	Vector of daily maximum temperatures (C)
tmin	Vector of daily minimum temperatures (C)
base_temp	Base temperature (C)
sunshine_hours	Vector of daily sunshine hours (optional)
day_length	Vector of day length in hours (optional)
crop_yield	Crop yield for HUE calculation (optional)
biomass	Biomass for HUE calculation (optional)
verbose	Logical. If TRUE (default), prints formatted output to console.

Value

List with thermal indices

Author(s)

Lalit Kumar Rolaniya, Ram Lal Jat, Monika Punia, Raja Ram Choudhary

Examples

```
tmax <- runif(90, 30, 42)
tmin <- runif(90, 18, 28)
thermal_indices(tmax, tmin, base_temp = 10)
```

yield_gap_analysis	<i>Yield Gap Analysis</i>
--------------------	---------------------------

Description

Calculates yield gaps comparing actual with potential yields.

Usage

```
yield_gap_analysis(  
  actual_yield,  
  potential_yield,  
  attainable_yield = NULL,  
  verbose = TRUE  
)
```

Arguments

actual_yield	Actual yield (kg/ha)
potential_yield	Potential yield (kg/ha)
attainable_yield	Attainable yield (kg/ha, optional)
verbose	Logical. If TRUE (default), prints formatted output to console.

Value

Data frame with yield gap analysis

Author(s)

Lalit Kumar Rolaniya, Ram Lal Jat, Monika Punia, Raja Ram Choudhary

Examples

```
yield_gap_analysis(actual_yield = c(800, 950, 720), potential_yield = 1500)
```

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