

Package ‘fastrda’

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Type Package

Title Fast Redundancy Analysis (RDA) with High-Performance 'C++' Backend

Version 0.1.2

Description Provides a high-performance implementation of redundancy analysis (RDA) in 'C++' using 'Armadillo' and 'OpenMP'. Supports standard and partial RDA, centering, scaling, overall and axis-wise permutation tests, biplot visualization, score extraction, and prediction. Designed for large ecological, genomic, and other multivariate data sets where computational speed and memory efficiency are important.

Depends R (>= 4.0.0)

Imports Rcpp (>= 1.0.0), parallel, ggplot2 (>= 3.3.0)

Suggests RhpcBLASctl, ggrepel (>= 0.9.0), RColorBrewer (>= 1.1-2), knitr, rmarkdown, vegan

LinkingTo Rcpp, RcppArmadillo (>= 0.12.0)

VignetteBuilder knitr

License GPL (>= 3)

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biplotrda

Biplot and Biplot Scores for fastrda Objects

Description

Computes vegan-compatible species and environmental biplot scores from a fitted `fastrda` object, and generates publication-ready ordination biplots using **ggplot2**.

Usage

```
biplot_scores(
  object,
  type = c("species", "environment", "both"),
  scaling = NULL,
  X = NULL,
  ...
)

biplotrda(
  x,
  axes = 1:2,
  type = c("all", "sites", "species", "environment",
           "sites_species", "sites_environment"),
  scaling = 2,
  X = NULL,
  show_ids = FALSE,
  colpal = NULL,
  site_col = NULL,
  species_col = NULL,
  env_col = NULL,
  site_size = 2.5,
  arrow_size = 0.8,
  label_size = 3.5,
  arrow_scale = 1.5,
  alpha = 0.8,
  max_labels = 50,
  title = "fastrda Biplot",
  subtitle = NULL,
  theme = ggplot2::theme_minimal()
)

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

## S3 method for class 'fastrda'
biplot(x, ...)
```

```
## S3 method for class 'fastrda'
plot(x, ...)
```

Arguments

object, x	A fitted fastrda object (or a fastrda_biplot object for print methods).
type	Character string specifying what to compute or display. For biplot_scores: "species", "environment", or "both". For biplotrda: "all", "sites", "species", "environment", "sites_species", or "sites_environment".
scaling	Integer (0-3) specifying the scaling type for scores. If NULL, uses the scaling from the original analysis model.
X	Optional matrix or data frame of environmental variables if not stored in the object.
axes	Integer vector of length 2 specifying the constrained axes to plot. Default is 1:2.
show_ids	Logical; if TRUE, displays site, species, or environmental variable labels. Default is FALSE.
colpal	Character string specifying a color palette name from RColorBrewer . Default is "Set1".
site_col	Custom color for site points (overrides palette).
species_col	Custom color for species arrows (overrides palette).
env_col	Custom color for environmental arrows (overrides palette).
site_size	Numeric; size of site points. Default is 2.5.
arrow_size	Numeric; thickness of arrows. Default is 0.8.
label_size	Numeric; size of labels. Default is 3.5.
arrow_scale	Numeric; scaling multiplier factor for biplot arrows. Default is 1.5.
alpha	Numeric; transparency value for points. Default is 0.8.
max_labels	Integer; maximum number of labels to display when show_ids = TRUE. Default is 50.
title	Character string for the plot title. Default is "fastrda Biplot".
subtitle	Character string for the plot subtitle. Default is NULL.
theme	ggplot2 theme object. Default is ggplot2::theme_minimal().
...	Additional arguments passed to lower-level plot or score functions.

Details

The function `biplot_scores` extracts or dynamically calculates site-weighted canonical coefficients and species scores conforming to the standard **vegan** package definitions across various scaling configurations (0 to 3). It seamlessly supports both matrix and list scaling structure formats in `fastrda v1.4.4`.

The function `biplotrda` (and its S3 generics `biplot.fastrda` and `plot.fastrda`) builds a customizable **ggplot2** graphic supporting point layers for sites, vector arrows for species and environmental variables, automatic axis proportion variance labeling, and intelligent label overlapping control via **ggrepel** (if installed).

Value

For `biplot_scores`, returns a numeric matrix (if type is species or environment) or a list with species and environment components assigned to class `fastrda_biplot`.

For `biplotrda`, `biplot.fastrda`, and `plot.fastrda`, returns a **ggplot2** object representing the ordination biplot.

For `print.fastrda_biplot`, invisibly returns the input object after printing summary text to the console.

See Also

[fastrda](#), [ordiplot](#)

Examples

```
## Not run:
set.seed(42)

n <- 30

env <- data.frame(
  Temperature = seq(10, 30, length.out = n),
  Moisture = rev(seq(20, 80, length.out = n)),
  pH = rnorm(n, 6.8, 0.4)
)

Y <- cbind(
  Sp1 = 1.2 * scale(env$Temperature) + rnorm(n, 0, 0.30),
  Sp2 = 0.9 * scale(env$Temperature) + rnorm(n, 0, 0.35),
  Sp3 = 1.1 * scale(env$Moisture) + rnorm(n, 0, 0.30),
  Sp4 = -0.9 * scale(env$Moisture) + rnorm(n, 0, 0.30),
  Sp5 = 1.0 * scale(env$pH) + rnorm(n, 0, 0.30),
  Sp6 = -0.8 * scale(env$pH) + rnorm(n, 0, 0.30)
)

fit <- fastrda(
  genotype = Y,
  environment = env,
  axes = 2,
  scaling = 2,
  keep_workspace = "none",
  verbose = FALSE
)

bp_scores <- biplot_scores(fit, type = "both")

bp_scores$environment
bp_scores$species

biplotrda(
  fit,
  axes = 1:2,
```

```

    scaling = 2,
    type = "all",
    show_ids = TRUE,
    title = "Simulated RDA Example"
  )

  ## End(Not run)

```

fastrda

Fast Redundancy Analysis and Associated Methods

Description

A high-performance implementation of Redundancy Analysis (RDA) and partial RDA using efficient QR decomposition, complete with integrated S3 methods for prediction, score extraction, permutation tests, summary reporting, and biplot visualization.

Usage

```

fastrda(
  genotype,
  environment,
  covariates = NULL,
  scale = TRUE,
  center = TRUE,
  axes = 3,
  scaling = 2,
  keep_workspace = "minimal",
  store_fitted = FALSE,
  threads = min(2, parallel::detectCores()),
  verbose = TRUE
)

## S3 method for class 'fastrda'
predict(
  object,
  newdata = NULL,
  type = c("lc", "response"),
  rank = NULL,
  scaling = NULL,
  ...
)

## S3 method for class 'fastrda'
scores(
  x,
  display = c("both", "sites", "species", "wa", "lc"),

```

```

    choices = 1:2,
    scaling = NULL,
    ...
)

anova_fastrda(
  object,
  by = c("overall", "axis"),
  permutations = 999,
  threads = min(2, parallel::detectCores())
)

## S3 method for class 'fastrda'
anova(object, ...)

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

## S3 method for class 'fastrda'
summary(object, ...)

## S3 method for class 'anova.fastrda'
print(x, ...)

## S3 method for class 'anova.fastrda.axis'
print(x, ...)

get_site_scores(object, scaling = 2)

```

Arguments

genotype	A numeric matrix or data frame of response variables (e.g., species abundances, genetic variants). Rows represent samples, columns represent variables.
environment	A numeric matrix or data frame of predictor variables. Rows must match the number of rows in genotype.
covariates	An optional numeric matrix or data frame of conditioning variables to be partialled out (partial RDA). Default is NULL.
scale	Logical; if TRUE, variables are scaled to unit variance before analysis. Default is TRUE.
center	Logical; if TRUE, variables are centered to zero mean before analysis. Default is TRUE.
axes	Integer; number of constrained axes to retain. Must be ≥ 1 . Default is 3.
scaling	Integer (0-3); type of scaling for ordination scores: 0 No scaling. 1 Sites scaled by eigenvalue sqrt, species unscaled. 2 Sites unscaled, species scaled by eigenvalue sqrt (vegan default).

	3 Symmetric scaling (both sites and species scaled by eigenvalue fourth root).
keep_workspace	Logical or character specifying workspace retention: "minimal" Store only Q and QtY (default). "compact" Store Q, QtY and Y_res. "full" Store all workspace elements. "none" Store no workspace.
store_fitted	Logical; if TRUE, stores fitted values in the output. Default is FALSE.
threads	Integer; number of threads for parallel computation. Defaults to number of available cores.
verbose	Logical; if TRUE, prints progress and summary messages. Default is TRUE.
object, x	A fitted fastrda object.
newdata	An optional matrix or data frame of new predictor variables for prediction.
type	Character string specifying the prediction type: "lc" Linear combination scores for new data. "response" Reconstructed response variables.
rank	Integer specifying the number of constrained axes to use for prediction or low-rank reconstruction.
choices	Integer vector specifying the constrained axes to extract.
by	Character string specifying the permutation test type: "overall" Global permutation test for the model. "axis" Marginal permutation test for each constrained axis.
permutations	Integer; number of permutations to perform. Default is 999.
display	Character string specifying what scores to extract: "both" Both sites and species scores. "sites" Site scores only. "species" Species scores only. "wa" Alias for site scores (vegan compatibility). "lc" Linear combination scores.
...	Additional arguments passed to other methods.

Details

Redundancy Analysis (RDA) is a multivariate constrained ordination method that finds linear combinations of environmental variables that best explain variation in response variables. It extends PCA by including explanatory variables.

fastrda implements RDA using a reduced-rank formulation based on efficient QR decomposition. For partial RDA models, conditioning variables are handled via Z-residualization.

Key Features:

High Performance C++ core with OpenMP parallelization for large matrices.

Scaled Scores Site and species scores are pre-computed for fast access.

Partial RDA Support for conditioning variables.

Permutation Tests Overall and axis-wise permutation tests.

vegan Compatibility S3 methods mimic vegan interface.

Scaling Types: The scaling parameter controls how site and species scores are scaled:

scaling = 0 No scaling (raw scores).

scaling = 1 Site scores scaled by eigenvalue sqrt, species unscaled.

scaling = 2 Site scores unscaled, species scaled by eigenvalue sqrt (vegan default).

scaling = 3 Symmetric scaling (both scaled by eigenvalue fourth root).

Note on Adjusted R-squared: The semi-partial adjusted R^2 formulation computed in this version differs from the default implementation found in the **vegan** package.

Value

For `fastrda`, an object of class `fastrda` containing a list of model results:

site_scores Pre-computed scaled site scores for the fitted scaling (access via `fit$site_scores`).

raw_site_scores Unscaled site scores for flexible re-scaling.

species_scores Pre-computed scaled species scores for the fitted scaling.

loadings Species canonical coefficients/loadings.

coefficients Environmental coefficients for the full-rank model.

lc_coefficients_raw Raw linear combination coefficients.

eigenvalues Eigenvalues of the constrained axes.

prop_total Proportion of total inertia explained per axis.

prop_constrained Proportion of constrained inertia explained per axis.

n_axes Number of retained constrained axes.

n_samples Number of samples (rows).

df_model Model degrees of freedom.

df_resid Residual degrees of freedom.

df_cond Conditioning (partial) degrees of freedom.

total_inertia Total inertia of the response matrix.

constrained_inertia Inertia explained by constraints.

unconstrained_inertia Residual inertia.

conditioned_inertia Inertia removed by conditioning variables (partial RDA).

R2 Coefficient of determination.

adj_R2 Adjusted R-squared.

pseudo_F Pseudo-F model statistic.

scaling_constants Pre-computed scaling multipliers for all scaling types.

workspace Workspace used for permutation tests (if retained).

call Matched function call.

env_names Column names of the environmental matrix.

species_names Column names of the response matrix.

site_names Row names of the response matrix.

For S3 methods and helper functions:

predict.fastrda Returns either linear combination scores (type = "lc") or reconstructed response matrix (type = "response").

scores.fastrda Returns a matrix of site scores or a list of site/species scores.

anova_fastrda Returns an object of class `anova.fastrda` (overall test) or `anova.fastrda.axis` (axis-wise test).

get_site_scores Returns site scores with optional re-scaling.

summary.fastrda Invisibly returns the `fastrda` object after printing high-level model summaries.

See Also

[biplotrda](#), [biplot_scores](#), and [rda](#) for the reference implementation from the **vegan** package.

Examples

```
set.seed(123)
n <- 100
p <- 200
k <- 3
Y <- matrix(rnorm(n * p), n, p)
X <- matrix(rnorm(n * k), n, k)
Z <- matrix(rnorm(n * 2), n, 2)
colnames(Y) <- paste0("Sp", 1:p)
colnames(X) <- paste0("Env", 1:k)

# Fit standard RDA
fit <- fastrda(genotype = Y, environment = X, axes = 3, verbose = FALSE)

# Print model summary
summary(fit)

# Access pre-computed scores
head(fit$site_scores)
head(fit$species_scores)

# Fit partial RDA with covariates
fit_prda <- fastrda(genotype = Y, environment = X, covariates = Z, axes = 3, verbose = FALSE)

# Extract scores using S3 method
sc <- scores(fit, display = "both", choices = 1:2)

# Predict low-rank linear combination scores (rank = 1 path)
pred_lc <- predict(fit, type = "lc", rank = 1)
```

```
# Predict response matrix for new data
pred_resp <- predict(fit, newdata = X, type = "response")

# Get site scores with different scaling (e.g., scaling = 1)
site_sc <- get_site_scores(fit, scaling = 1)
head(site_sc)

# Permutation tests (isolated to respect CRAN execution time limits)
anv_overall <- anova(fit, permutations = 99)
print(anv_overall)

anv_axis <- anova_fastrda(fit, by = "axis", permutations = 99)
print(anv_axis)
```

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