Package 'PLNmodels'

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Title Poisson Lognormal Models

Version 1.0.0

Description The Poisson-lognormal model and variants (Chiquet, Mariadassou and Robin, 2021 <doi:10.3389/fevo.2021.588292>) can be used for a variety of multivariate problems when count data are at play, including principal component analysis for count data, discriminant analysis, model-based clustering and network inference. Implements variational algorithms to fit such models accompanied with a set of functions for visualization and diagnostic.

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BugReports https://github.com/pln-team/PLNmodels/issues

License GPL (>= 3)

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Barents fish data set

Description

This data set gives the abundance of 30 fish species observed in 89 sites in the Barents sea. For each site, 4 additional covariates are known. Subsample of the original datasets studied by Fossheim et al, 2006.

Usage

barents

barents

Format

A data frame with 6 variables:

- Abundance: A 30 fish species by 89 sites count matrix
- Offset: A 30 fish species by 116 samples offset matrix, measuring the sampling effort in each site
- 4 covariates for latitude, longitude, depth (in meters), temperature (in Celsius degrees).

Source

Data from M. Fossheim and coauthors.

References

Fossheim, Maria, Einar M. Nilssen, and Michaela Aschan. "Fish assemblages in the Barents Sea." Marine Biology Research 2.4 (2006). doi:10.1080/17451000600815698

Examples

data(barents)

coef.PLNfit

Extract model coefficients

Description

Extracts model coefficients from objects returned by PLN() and its variants

Usage

```
## S3 method for class 'PLNfit'
coef(object, type = c("main", "covariance"), ...)
```

Arguments

object	an R6 object with class PLNfit	
type	type of parameter that should be extracted. Either "main" (default) for	
	В	
	or "covariance" for	
	Σ	
	additional parameters for S3 compatibility. Not used	

Value

A matrix of coefficients extracted from the PLNfit model.

coef.PLNLDAfit

See Also

sigma.PLNfit(), vcov.PLNfit(), standard_error.PLNfit()

Examples

```
data(trichoptera)
trichoptera <- prepare_data(trichoptera$Abundance, trichoptera$Covariate)
myPLN <- PLN(Abundance ~ 1 + offset(log(Offset)), data = trichoptera)
coef(myPLN) ## B
coef(myPLN, type = "covariance") ## Sigma</pre>
```

coef.PLNLDAfit Extracts model coefficients from objects returned by PLNLDA()

Description

The method for objects returned by PLNLDA() only returns coefficients associated to the

Θ

part of the model (see the PLNLDA vignette for mathematical details).

Usage

S3 method for class 'PLNLDAfit'
coef(object, ...)

Arguments

object	an R6 object with class PLNLDAfit
	additional parameters for S3 compatibility. Not used

Value

Either NULL or a matrix of coefficients extracted from the PLNLDAfit model.

Examples

```
data(trichoptera)
trichoptera <- prepare_data(trichoptera$Abundance, trichoptera$Covariate)
myPLNLDA <- PLNLDA(Abundance ~ Wind, grouping = Group, data = trichoptera)
coef(myPLNLDA)</pre>
```

coef.PLNmixturefit *Extract model coefficients*

Description

Extracts model coefficients from objects returned by PLN() and its variants

Usage

```
## S3 method for class 'PLNmixturefit'
coef(object, type = c("main", "means", "covariance", "mixture"), ...)
```

Arguments

object type	an R6 object with class PLNm type of parameter that should	ixturefit I be extracted. Either "main" (default) for	
		Θ	
	, "means" for		
		μ	
	, "mixture" for		
		π	
	or "covariance" for		
		Σ	
	additional parameters for S3	compatibility. Not used	

Value

A matrix of coefficients extracted from the PLNfit model.

See Also

sigma.PLNmixturefit()

Examples

coefficient_path Extract the regularization path of a PLNnetwork fit

Description

Extract the regularization path of a PLNnetwork fit

Usage

```
coefficient_path(Robject, precision = TRUE, corr = TRUE)
```

Arguments

Robject	an object with class PLNnetworkfamily, i.e. an output from PLNnetwork()
precision	a logical, should the coefficients of the precision matrix Omega or the covariance matrix Sigma be sent back. Default is TRUE.
corr	a logical, should the correlation (partial in case precision = TRUE) be sent back. Default is TRUE.

Value

Sends back a tibble/data.frame.

Examples

```
data(trichoptera)
trichoptera <- prepare_data(trichoptera$Abundance, trichoptera$Covariate)
fits <- PLNnetwork(Abundance ~ 1, data = trichoptera)
head(coefficient_path(fits))</pre>
```

compute_offset	Compute offsets from a count data using one of several normalization
	schemes

Description

Computes offsets from the count table using one of several normalization schemes (TSS, CSS, RLE, GMPR, etc) described in the literature.

Usage

```
compute_offset(
  counts,
  offset = c("TSS", "GMPR", "RLE", "CSS", "Wrench", "none"),
  ...
)
```

Arguments

counts	Required. An abundance count table, preferably with dimensions names and species as columns.
offset	Optional. Normalization scheme used to compute scaling factors used as offset during PLN inference. Available schemes are "TSS" (Total Sum Scaling, default), "CSS" (Cumulative Sum Scaling, used in metagenomeSeq), "RLE" (Relative Log Expression, used in DESeq2), "GMPR" (Geometric Mean of Pairwise Ratio, introduced in Chen et al., 2018), Wrench (introduced in Kumar et al., 2018) or "none". Alternatively the user can supply its own vector or matrix of offsets (see note for specification of the user-supplied offsets).
	Additional parameters passed on to specific methods (for now CSS and RLE)

Details

RLE has additional pseudocounts and type arguments to add pseudocounts to the observed counts (defaults to 0L) and to compute offsets using only positive counts (if type == "poscounts"). This mimics the behavior of DESeq2::DESeq() when using sfType == "poscounts". CSS has an additional reference argument to choose the location function used to compute the reference quantiles (defaults to median as in the Nature publication but can be set to mean to reproduce behavior of functions cumNormStat* from metagenomeSeq). Wrench has two additional parameters: groups to specify sample groups and type to either reproduce exactly the default Wrench::wrench() behavior (type = "wrench", default) or to use simpler heuristics (type = "simple"). Note that (i) CSS normalization fails when the median absolute deviation around quantiles does not become instable for high quantiles (limited count variations both within and across samples) and/or one sample has less than two positive counts, (ii) RLE fails when there are no common species across all samples (unless type == "poscounts" has been specified) and (iii) GMPR fails if a sample does not share any species with all other samples.

Value

If offset = "none", NULL else a vector of length nrow(counts) with one offset per sample.

References

Chen, L., Reeve, J., Zhang, L., Huang, S., Wang, X. and Chen, J. (2018) GMPR: A robust normalization method for zero-inflated count data with application to microbiome sequencing data. PeerJ, 6, e4600 doi:10.7717/peerj.4600

Paulson, J. N., Colin Stine, O., Bravo, H. C. and Pop, M. (2013) Differential abundance analysis for microbial marker-gene surveys. Nature Methods, 10, 1200-1202 doi:10.1038/nmeth.2658

Anders, S. and Huber, W. (2010) Differential expression analysis for sequence count data. Genome Biology, 11, R106 doi:10.1186/gb20101110r106

Kumar, M., Slud, E., Okrah, K. et al. (2018) Analysis and correction of compositional bias in sparse sequencing count data. BMC Genomics 19, 799 doi:10.1186/s1286401851605

Examples

data(trichoptera)

extract_probs

```
counts <- trichoptera$Abundance
compute_offset(counts)
## Other normalization schemes
compute_offset(counts, offset = "RLE", pseudocounts = 1)
compute_offset(counts, offset = "Wrench", groups = trichoptera$Covariate$Group)
compute_offset(counts, offset = "GMPR")
## User supplied offsets
my_offset <- setNames(rep(1, nrow(counts)), rownames(counts))
compute_offset(counts, offset = my_offset)
```

extract_probs

Extract edge selection frequency in bootstrap subsamples

Description

Extracts edge selection frequency in networks reconstructed from bootstrap subsamples during the stars stability selection procedure, as either a matrix or a named vector. In the latter case, edge names follow igraph naming convention.

Usage

```
extract_probs(
  Robject,
  penalty = NULL,
  index = NULL,
  crit = c("StARS", "BIC", "EBIC"),
  format = c("matrix", "vector"),
  tol = 1e-05
)
```

Arguments

Robject	an object with class PLNnetworkfamily, i.e. an output from PLNnetwork()
penalty	penalty used for the bootstrap subsamples
index	Integer index of the model to be returned. Only the first value is taken into account.
crit	a character for the criterion used to performed the selection. Either "BIC", "ICL", "EBIC", "StARS", "R_squared". Default is ICL for PLNPCA, and BIC for PLNnetwork. If StARS (Stability Approach to Regularization Selection) is chosen and stability selection was not yet performed, the function will call the method stability_selection() with default argument.
format	output format. Either a matrix (default) or a named vector.
tol	tolerance for rounding error when comparing penalties.

Value

Either a matrix or named vector of edge-wise probabilities. In the latter case, edge names follow igraph convention.

Examples

```
data(trichoptera)
trichoptera <- prepare_data(trichoptera$Abundance, trichoptera$Covariate)</pre>
nets <- PLNnetwork(Abundance ~ 1 + offset(log(Offset)), data = trichoptera)</pre>
## Not run:
stability_selection(nets)
probs <- extract_probs(nets, crit = "StARS", format = "vector")</pre>
probs
## End(Not run)
## Not run:
## Add edge attributes to graph using igraph
net_stars <- getBestModel(nets, "StARS")</pre>
g <- plot(net_stars, type = "partial_cor", plot=F)</pre>
library(igraph)
E(g)$prob <- probs[as_ids(E(g))]</pre>
g
## End(Not run)
```

fitted.PLNfit	Extracts model fitted values from objects returned by PLN() and its
	variants

Description

Extracts model fitted values from objects returned by PLN() and its variants

Usage

```
## S3 method for class 'PLNfit'
fitted(object, ...)
```

Arguments

object	an R6 object with class PLNfit
	additional parameters for S3 compatibility. Not used

Value

A matrix of Fitted values extracted from the object object.

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fitted.PLNmixturefit *Extracts model fitted values from objects returned by* PLNmixture() *and its variants*

Description

Extracts model fitted values from objects returned by PLNmixture() and its variants

Usage

```
## S3 method for class 'PLNmixturefit'
fitted(object, ...)
```

Arguments

object	an R6 object with class PLNmixturefit
	additional parameters for S3 compatibility. Not used

Value

A matrix of Fitted values extracted from the object object.

```
getBestModel.PLNPCAfamily
```

Best model extraction from a collection of models

Description

Best model extraction from a collection of models

Usage

```
## S3 method for class 'PLNPCAfamily'
getBestModel(Robject, crit = c("ICL", "BIC"), ...)
getBestModel(Robject, crit, ...)
## S3 method for class 'PLNmixturefamily'
getBestModel(Robject, crit = c("ICL", "BIC"), ...)
## S3 method for class 'PLNnetworkfamily'
getBestModel(Robject, crit = c("BIC", "EBIC", "StARS"), ...)
```

Arguments

Robject	an object with class PLNPCA familly ot PLN network family
crit	a character for the criterion used to performed the selection. Either "BIC", "ICL", "EBIC", "StARS", "R_squared". Default is ICL for PLNPCA, and BIC for PLNnetwork. If StARS (Stability Approach to Regularization Selection) is chosen and stability selection was not yet performed, the function will call the method stability_selection() with default argument.
	additional parameters for StARS criterion (only for PLNnetwork). stability, a scalar indicating the target stability (= $1 - 2$ beta) at which the network is selected. Default is 0.9 .

Value

Send back an object with class PLNPCAfit or PLNnetworkfit

Methods (by class)

- getBestModel(PLNPCAfamily): Model extraction for PLNPCAfamily
- getBestModel(PLNmixturefamily): Model extraction for PLNmixturefamily
- getBestModel(PLNnetworkfamily): Model extraction for PLNnetworkfamily

Examples

```
## Not run:
data(trichoptera)
trichoptera <- prepare_data(trichoptera$Abundance, trichoptera$Covariate)
myPCA <- PLNPCA(Abundance ~ 1 + offset(log(Offset)), data = trichoptera, ranks = 1:4)
myModel <- getBestModel(myPCA)</pre>
```

End(Not run)

getModel.PLNPCAfamily Model extraction from a collection of models

Description

Model extraction from a collection of models

Usage

```
## S3 method for class 'PLNPCAfamily'
getModel(Robject, var, index = NULL)
getModel(Robject, var, index)
## S3 method for class 'PLNmixturefamily'
```

mollusk

```
getModel(Robject, var, index = NULL)
## S3 method for class 'PLNnetworkfamily'
getModel(Robject, var, index = NULL)
```

Arguments

Robject	an R6 object with class PLNPCAfamily or PLNnetworkfamily
var	value of the parameter (rank for PLNPCA, sparsity for PLNnetwork) that identifies the model to be extracted from the collection. If no exact match is found, the model with closest parameter value is returned with a warning.
index	Integer index of the model to be returned. Only the first value is taken into account.

Value

Sends back an object with class PLNPCAfit or PLNnetworkfit.

Methods (by class)

- getModel(PLNPCAfamily): Model extraction for PLNPCAfamily
- getModel(PLNmixturefamily): Model extraction for PLNmixturefamily
- getModel(PLNnetworkfamily): Model extraction for PLNnetworkfamily

Examples

```
## Not run:
data(trichoptera)
trichoptera <- prepare_data(trichoptera$Abundance, trichoptera$Covariate)
myPCA <- PLNPCA(Abundance ~ 1 + offset(log(Offset)), data = trichoptera, ranks = 1:5)
myModel <- getModel(myPCA, 2)</pre>
```

End(Not run)

mollusk

Mollusk data set

Description

This data set gives the abundance of 32 mollusk species in 163 samples. For each sample, 4 additional covariates are known.

Usage

mollusk

Format

A list with 2 two data frames:

Abundance a 163 x 32 data frame of abundancies/counts (163 samples and 32 mollusk species)

Covariate a 163 x 4 data frame of covariates:

site a factor with 8 levels indicating the sampling site

season a factor with 4 levels indicating the season

method a factor with 2 levels for the method of sampling - wood or string

duration a numeric with 3 levels for the time of exposure in week

In order to prepare the data for using formula in multivariate analysis (multiple outputs and inputs), use prepare_data(). Original data set has been extracted from ade4.

Source

Data from Richardot-Coulet, Chessel and Bournaud.

References

Richardot-Coulet, M., Chessel D. and Bournaud M. (1986) Typological value of the benthos of old beds of a large river. Methodological approach. Archiv für Hydrobiologie, 107, 363–383.

See Also

prepare_data()

Examples

```
data(mollusk)
mollusc <- prepare_data(mollusk$Abundance, mollusk$Covariate)</pre>
```

oaks

Oaks amplicon data set

Description

This data set gives the abundance of 114 taxa (66 bacterial OTU, 48 fungal OTUs) in 116 samples. For each sample, 11 additional covariates are known.

Usage

oaks

oaks

Format

A data frame with 13 variables:

- Abundance: A 114 taxa by 116 samples count matrix
- Offset: A 114 taxa by 116 samples offset matrix
- Sample: Unique sample id
- tree: Tree status with respect to the pathogen (susceptible, intermediate or resistant)
- branch: Unique branch id in each tree (4 branches were sampled in each tree, with 10 leaves per branch)
- leafNO: Unique leaf id in each tree (40 leaves were sampled in each tree)
- distTObase: Distance of the sampled leaf to the base of the branch
- distTOtrunk: Distance of the sampled leaf to the base of the tree trunk
- · distTOground: Distance of the sampled leaf to the base of the ground
- pmInfection: Powdery mildew infection, proportion of the upper leaf area displaying mildew symptoms
- orientation: Orientation of the branch (South-West SW or North-East NE)
- readsTOTfun: Total number of ITS1 reads for that leaf
- readsTOTbac: Total number of 16S reads for that leaf

Source

Data from B. Jakuschkin and coauthors.

References

Jakuschkin, B., Fievet, V., Schwaller, L. et al. Deciphering the Pathobiome: Intra- and Interkingdom Interactions Involving the Pathogen Erysiphe alphitoides . Microb Ecol 72, 870–880 (2016). doi:10.1007/s002480160777x

See Also

prepare_data()

Examples

```
data(oaks)
## Not run:
oaks_networks <- PLNnetwork(formula = Abundance ~ 1 + offset(log(0ffset)), data = oaks)
## End(Not run)</pre>
```

Description

Fit the multivariate Poisson lognormal model with a variational algorithm. Use the (g)lm syntax for model specification (covariates, offsets, weights).

Usage

```
PLN(formula, data, subset, weights, control = PLN_param())
```

Arguments

formula	an object of class "formula": a symbolic description of the model to be fitted.
data	an optional data frame, list or environment (or object coercible by as.data.frame to a data frame) containing the variables in the model. If not found in data, the variables are taken from environment(formula), typically the environment from which PLN is called.
subset	an optional vector specifying a subset of observations to be used in the fitting process.
weights	an optional vector of observation weights to be used in the fitting process.
control	a list-like structure for controlling the optimization, with default generated by PLN_param(). See the associated documentation for details.

Value

an R6 object with class ${\sf PLNfit}$

See Also

The class PLNfit and the configuration function PLN_param()

Examples

```
data(trichoptera)
trichoptera <- prepare_data(trichoptera$Abundance, trichoptera$Covariate)
myPLN <- PLN(Abundance ~ 1, data = trichoptera)</pre>
```

PLN

PLNfamily

Description

super class for PLNPCAfamily and PLNnetworkfamily.

Public fields

responses the matrix of responses common to every models

covariates the matrix of covariates common to every models

offsets the matrix of offsets common to every models

weights the vector of observation weights

inception a PLNfit object, obtained when no sparsifying penalty is applied.

models a list of PLNfit object, one per penalty.

Active bindings

criteria a data frame with the values of some criteria (approximated log-likelihood, BIC, ICL, etc.) for the collection of models / fits BIC and ICL are defined so that they are on the same scale as the model log-likelihood, i.e. with the form, loglik - 0.5 penalty

convergence sends back a data frame with some convergence diagnostics associated with the optimization process (method, optimal value, etc)

Methods

Public methods:

- PLNfamily\$new()
- PLNfamily\$postTreatment()
- PLNfamily\$getModel()
- PLNfamily\$plot()
- PLNfamily\$show()
- PLNfamily\$print()
- PLNfamily\$clone()

Method new(): Create a new PLNfamily object.

Usage:

PLNfamily\$new(responses, covariates, offsets, weights, control)

Arguments:

responses the matrix of responses common to every models covariates the matrix of covariates common to every models offsets the matrix of offsets common to every models weights the vector of observation weights control list controlling the optimization and the model *Returns:* A new PLNfamily object

Method postTreatment(): Update fields after optimization

Usage:

PLNfamily\$postTreatment(config)

Arguments:

config a list for controlling the post-treatment.

Method getModel(): Extract a model from a collection of models

Usage:

PLNfamily\$getModel(var, index = NULL)

Arguments:

var value of the parameter (rank for PLNPCA, sparsity for PLNnetwork) that identifies the model to be extracted from the collection. If no exact match is found, the model with closest parameter value is returned with a warning.

index Integer index of the model to be returned. Only the first value is taken into account.

Returns: A PLNfit object

Method plot(): Lineplot of selected criteria for all models in the collection

Usage:

PLNfamily\$plot(criteria, reverse)

Arguments:

- criteria A valid model selection criteria for the collection of models. Includes loglik, BIC (all), ICL (PLNPCA) and pen_loglik, EBIC (PLNnetwork)
- reverse A logical indicating whether to plot the value of the criteria in the "natural" direction (loglik penalty) or in the "reverse" direction (-2 loglik + penalty). Default to FALSE, i.e use the natural direction, on the same scale as the log-likelihood.

Returns: A ggplot2 object

Method show(): User friendly print method

Usage:
PLNfamily\$show()

Method print(): User friendly print method

Usage:

PLNfamily\$print()

Method clone(): The objects of this class are cloneable with this method.

Usage:

PLNfamily\$clone(deep = FALSE)

Arguments:

deep Whether to make a deep clone.

See Also

getModel()

PLNfit

Description

The function PLN() fit a model which is an instance of a object with class PLNfit. Objects produced by the functions PLNnetwork(), PLNPCA(), PLNmixture() and PLNLDA() also enjoy the methods of PLNfit() by inheritance.

This class comes with a set of R6 methods, some of them being useful for the user and exported as S3 methods. See the documentation for coef(), sigma(), predict(), vcov() and standard_error().

Fields are accessed via active binding and cannot be changed by the user.

Active bindings

- n number of samples
- q number of dimensions of the latent space
- p number of species
- d number of covariates
- nb_param number of parameters in the current PLN model
- model_par a list with the matrices of the model parameters: B (covariates), Sigma (covariance), Omega (precision matrix), plus some others depending on the variant)

var_par a list with the matrices of the variational parameters: M (means) and S2 (variances)

- optim_par a list with parameters useful for monitoring the optimization
- latent a matrix: values of the latent vector (Z in the model)
- latent_pos a matrix: values of the latent position vector (Z) without covariates effects or offset
- fitted a matrix: fitted values of the observations (A in the model)
- vcov_coef matrix of sandwich estimator of the variance-covariance of B (need fixed -ie known-covariance at the moment)
- vcov_model character: the model used for the residual covariance
- weights observational weights
- loglik (weighted) variational lower bound of the loglikelihood
- loglik_vec element-wise variational lower bound of the loglikelihood
- BIC variational lower bound of the BIC
- entropy Entropy of the variational distribution
- ICL variational lower bound of the ICL
- R_squared approximated goodness-of-fit criterion
- criteria a vector with loglik, BIC, ICL and number of parameters

Methods

Public methods:

- PLNfit\$new()
- PLNfit\$update()
- PLNfit\$optimize()
- PLNfit\$optimize_vestep()
- PLNfit\$postTreatment()
- PLNfit\$predict()
- PLNfit\$predict_cond()
- PLNfit\$show()
- PLNfit\$print()
- PLNfit\$clone()

Method new(): Initialize a PLNfit model

Usage:

PLNfit\$new(responses, covariates, offsets, weights, formula, control)

Arguments:

- responses the matrix of responses (called Y in the model). Will usually be extracted from the corresponding field in PLNfamily-class
- covariates design matrix (called X in the model). Will usually be extracted from the corresponding field in PLNfamily-class
- offsets offset matrix (called O in the model). Will usually be extracted from the corresponding field in PLNfamily-class
- weights an optional vector of observation weights to be used in the fitting process.
- formula model formula used for fitting, extracted from the formula in the upper-level call control a list-like structure for controlling the fit, see PLN_param().

Method update(): Update a PLNfit object

```
Usage:

PLNfit$update(

B = NA,

Sigma = NA,

Omega = NA,

M = NA,

S = NA,

Ji = NA,

R2 = NA,

Z = NA,

A = NA,

monitoring = NA

)
```

Arguments:

B matrix of regression matrix

Sigma variance-covariance matrix of the latent variables

Omega precision matrix of the latent variables. Inverse of Sigma.

M matrix of variational parameters for the mean

S matrix of variational parameters for the variance

Ji vector of variational lower bounds of the log-likelihoods (one value per sample)

R2 approximate R^2 goodness-of-fit criterion

Z matrix of latent vectors (includes covariates and offset effects)

A matrix of fitted values

monitoring a list with optimization monitoring quantities

Returns: Update the current PLNfit object

Method optimize(): Call to the NLopt or TORCH optimizer and update of the relevant fields

Usage:

PLNfit\$optimize(responses, covariates, offsets, weights, config)

Arguments:

- responses the matrix of responses (called Y in the model). Will usually be extracted from the corresponding field in PLNfamily-class
- covariates design matrix (called X in the model). Will usually be extracted from the corresponding field in PLNfamily-class
- offsets offset matrix (called O in the model). Will usually be extracted from the corresponding field in PLNfamily-class

weights an optional vector of observation weights to be used in the fitting process.

config part of the control argument which configures the optimizer

Method optimize_vestep(): Result of one call to the VE step of the optimization procedure: optimal variational parameters (M, S) and corresponding log likelihood values for fixed model parameters (Sigma, B). Intended to position new data in the latent space.

```
Usage:
PLNfit$optimize_vestep(
  covariates,
  offsets,
  responses,
  weights,
  B = self$model_par$B,
  Omega = self$model_par$Omega,
   control = PLN_param(backend = "nlopt")
)
```

Arguments:

- covariates design matrix (called X in the model). Will usually be extracted from the corresponding field in PLNfamily-class
- offsets offset matrix (called O in the model). Will usually be extracted from the corresponding field in PLNfamily-class
- responses the matrix of responses (called Y in the model). Will usually be extracted from the corresponding field in PLNfamily-class
- weights an optional vector of observation weights to be used in the fitting process.

B Optional fixed value of the regression parameters

Omega precision matrix of the latent variables. Inverse of Sigma.

control a list-like structure for controlling the fit, see PLN_param().

Sigma variance-covariance matrix of the latent variables

Returns: A list with three components:

- the matrix M of variational means,
- the matrix S2 of variational variances
- the vector log.lik of (variational) log-likelihood of each new observation

Method postTreatment(): Update R2, fisher and std_err fields after optimization

```
Usage:
```

```
PLNfit$postTreatment(
   responses,
   covariates,
   offsets,
   weights = rep(1, nrow(responses)),
   config,
   nullModel = NULL
)
```

Arguments:

- responses the matrix of responses (called Y in the model). Will usually be extracted from the corresponding field in PLNfamily-class
- covariates design matrix (called X in the model). Will usually be extracted from the corresponding field in PLNfamily-class
- offsets offset matrix (called O in the model). Will usually be extracted from the corresponding field in PLNfamily-class
- weights an optional vector of observation weights to be used in the fitting process.
- config part of the control argument which configures the optimizer
- nullModel null model used for approximate R2 computations. Defaults to a GLM model with same design matrix but not latent variable.
- jackknife Boolean indicating whether jackknife estimation of bias and variance should be computed for the model parameters. Default is FALSE

Method predict(): Predict position, scores or observations of new data.

Usage:

PLNfit\$predict(newdata, type = c("link", "response"), envir = parent.frame())
Arguments:

- newdata A data frame in which to look for variables with which to predict. If omitted, the fitted values are used.
- type Scale used for the prediction. Either link (default, predicted positions in the latent space) or response (predicted counts).
- envir Environment in which the prediction is evaluated

Returns: A matrix with predictions scores or counts.

PLNfit

Method predict_cond(): Predict position, scores or observations of new data, conditionally on the observation of a (set of) variables

```
Usage:
PLNfit$predict_cond(
    newdata,
    cond_responses,
    type = c("link", "response"),
    var_par = FALSE,
    envir = parent.frame()
)
```

Arguments:

newdata a data frame containing the covariates of the sites where to predict

- cond_responses a data frame containing the count of the observed variables (matching the names of the provided as data in the PLN function)
- type Scale used for the prediction. Either link (default, predicted positions in the latent space) or response (predicted counts).
- var_par Boolean. Should new estimations of the variational parameters of mean and variance be sent back, as attributes of the matrix of predictions. Default to FALSE.

envir Environment in which the prediction is evaluated

Returns: A matrix with predictions scores or counts.

Method show(): User friendly print method

```
Usage:
PLNfit$show(
   model = paste("A multivariate Poisson Lognormal fit with", self$vcov_model,
        "covariance model.\n")
)
```

Arguments: model First line of the print output

Method print(): User friendly print method

Usage:
PLNfit\$print()

Method clone(): The objects of this class are cloneable with this method.

Usage: PLNfit\$clone(deep = FALSE) Arguments:

deep Whether to make a deep clone.

Examples

```
## Not run:
data(trichoptera)
trichoptera <- prepare_data(trichoptera$Abundance, trichoptera$Covariate)</pre>
```

```
myPLN <- PLN(Abundance ~ 1, data = trichoptera)
class(myPLN)
print(myPLN)
## End(Not run)</pre>
```

PLNfit_diagonal

An R6 Class to represent a PLNfit in a standard, general framework, with diagonal residual covariance

Description

The function PLNLDA() produces an instance of an object with class PLNLDAfit.

This class comes with a set of methods, some of them being useful for the user: See the documentation for the methods inherited by PLNfit(), the plot() method for LDA visualization and predict() method for prediction

Super class

PLNmodels::PLNfit -> PLNfit_diagonal

Active bindings

nb_param number of parameters in the current PLN model

vcov_model character: the model used for the residual covariance

Methods

Public methods:

- PLNfit_diagonal\$new()
- PLNfit_diagonal\$clone()

Method new(): Initialize a PLNfit model

Usage:

PLNfit_diagonal\$new(responses, covariates, offsets, weights, formula, control)

Arguments:

- responses the matrix of responses (called Y in the model). Will usually be extracted from the corresponding field in PLNfamily-class
- covariates design matrix (called X in the model). Will usually be extracted from the corresponding field in PLNfamily-class
- offsets offset matrix (called O in the model). Will usually be extracted from the corresponding field in PLNfamily-class
- weights an optional vector of observation weights to be used in the fitting process.
- formula model formula used for fitting, extracted from the formula in the upper-level call control a list for controlling the optimization. See details.

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Method clone(): The objects of this class are cloneable with this method.
Usage:
PLNfit_diagonal\$clone(deep = FALSE)
Arguments:
deep Whether to make a deep clone.

Super classes

PLNmodels::PLNfit -> PLNmodels::PLNLDAfit -> PLNLDAfit_spherical

Active bindings

vcov_model character: the model used for the residual covariance

Methods

Public methods:

- PLNLDAfit_spherical\$new()
- PLNLDAfit_spherical\$clone()

Method new(): Initialize a PLNfit model

```
Usage:
PLNLDAfit_spherical$new(
  grouping,
  responses,
  covariates,
  offsets,
  weights,
  formula,
  control
)
```

Arguments:

grouping a factor specifying the class of each observation used for discriminant analysis.

- responses the matrix of responses (called Y in the model). Will usually be extracted from the corresponding field in PLNfamily-class
- covariates design matrix (called X in the model). Will usually be extracted from the corresponding field in PLNfamily-class
- offsets offset matrix (called O in the model). Will usually be extracted from the corresponding field in PLNfamily-class
- weights an optional vector of observation weights to be used in the fitting process.

formula model formula used for fitting, extracted from the formula in the upper-level call control a list for controlling the optimization. See details.

Method clone(): The objects of this class are cloneable with this method.

Usage: PLNLDAfit_spherical\$clone(deep = FALSE) Arguments: deep Whether to make a deep clone.

Examples

```
## Not run:
data(trichoptera)
trichoptera <- prepare_data(trichoptera$Abundance, trichoptera$Covariate)
myPLN <- PLN(Abundance ~ 1, data = trichoptera)
class(myPLN)
print(myPLN)
## End(Not run)
## Not run:
data(trichoptera)
trichoptera <- prepare_data(trichoptera$Abundance, trichoptera$Covariate)
myPLNLDA <- PLNLDA(Abundance ~ 1, data = trichoptera, control = PLN_param(covariance = "spherical"))
class(myPLNLDA)
print(myPLNLDA)
## End(Not run)
```

PLNfit_fixedcov	An R6 Class to represent a PLNfit in a standard, general framework,
	with fixed (inverse) residual covariance

Description

An R6 Class to represent a PLNfit in a standard, general framework, with fixed (inverse) residual covariance

An R6 Class to represent a PLNfit in a standard, general framework, with fixed (inverse) residual covariance

Super class

PLNmodels::PLNfit -> PLNfit_fixedcov

Active bindings

nb_param number of parameters in the current PLN model

vcov_model character: the model used for the residual covariance

vcov_coef matrix of sandwich estimator of the variance-covariance of B (needs known covariance at the moment)

Methods

Public methods:

- PLNfit_fixedcov\$new()
- PLNfit_fixedcov\$optimize()
- PLNfit_fixedcov\$postTreatment()

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PLNfit_fixedcov

```
    PLNfit_fixedcov$clone()
```

Method new(): Initialize a PLNfit model

Usage:

```
PLNfit_fixedcov$new(responses, covariates, offsets, weights, formula, control)
```

Arguments:

- responses the matrix of responses (called Y in the model). Will usually be extracted from the corresponding field in PLNfamily-class
- covariates design matrix (called X in the model). Will usually be extracted from the corresponding field in PLNfamily-class
- offsets offset matrix (called O in the model). Will usually be extracted from the corresponding field in PLNfamily-class
- weights an optional vector of observation weights to be used in the fitting process.
- formula model formula used for fitting, extracted from the formula in the upper-level call control a list for controlling the optimization. See details.

Method optimize(): Call to the NLopt or TORCH optimizer and update of the relevant fields *Usage*:

PLNfit_fixedcov\$optimize(responses, covariates, offsets, weights, config)

- Arguments:
- responses the matrix of responses (called Y in the model). Will usually be extracted from the corresponding field in PLNfamily-class
- covariates design matrix (called X in the model). Will usually be extracted from the corresponding field in PLNfamily-class
- offsets offset matrix (called O in the model). Will usually be extracted from the corresponding field in PLNfamily-class

weights an optional vector of observation weights to be used in the fitting process. config part of the control argument which configures the optimizer

Method postTreatment(): Update R2, fisher and std_err fields after optimization

```
Usage:
PLNfit_fixedcov$postTreatment(
  responses,
  covariates,
  offsets,
  weights = rep(1, nrow(responses)),
  config,
  nullModel = NULL
)
```

Arguments:

responses the matrix of responses (called Y in the model). Will usually be extracted from the corresponding field in PLNfamily-class

covariates design matrix (called X in the model). Will usually be extracted from the corresponding field in PLNfamily-class

- offsets offset matrix (called O in the model). Will usually be extracted from the corresponding field in PLNfamily-class
- weights an optional vector of observation weights to be used in the fitting process.
- config part of the control argument which configures the optimizer
- nullModel null model used for approximate R2 computations. Defaults to a GLM model with same design matrix but not latent variable.
- jackknife Boolean indicating whether jackknife estimation of bias and variance should be computed for the model parameters. Default is FALSE

Method clone(): The objects of this class are cloneable with this method.

```
Usage:
PLNfit_fixedcov$clone(deep = FALSE)
Arguments:
deep Whether to make a deep clone.
```

Examples

```
## Not run:
data(trichoptera)
trichoptera <- prepare_data(trichoptera$Abundance, trichoptera$Covariate)
myPLN <- PLN(Abundance ~ 1, data = trichoptera)
class(myPLN)
print(myPLN)
```

End(Not run)

PLNfit_spherical	An R6 Class to represent a PLNfit in a standard, general framework,
	with spherical residual covariance

Description

An R6 Class to represent a PLNfit in a standard, general framework, with spherical residual covariance

An R6 Class to represent a PLNfit in a standard, general framework, with spherical residual covariance

Super class

PLNmodels::PLNfit -> PLNfit_spherical

Active bindings

nb_param number of parameters in the current PLN model vcov_model character: the model used for the residual covariance

PLNLDA

Methods

Public methods:

- PLNfit_spherical\$new()
- PLNfit_spherical\$clone()

Method new(): Initialize a PLNfit model

Usage:

```
PLNfit_spherical$new(responses, covariates, offsets, weights, formula, control)
```

Arguments:

- responses the matrix of responses (called Y in the model). Will usually be extracted from the corresponding field in PLNfamily-class
- covariates design matrix (called X in the model). Will usually be extracted from the corresponding field in PLNfamily-class
- offsets offset matrix (called O in the model). Will usually be extracted from the corresponding field in PLNfamily-class
- weights an optional vector of observation weights to be used in the fitting process.

formula model formula used for fitting, extracted from the formula in the upper-level call control a list for controlling the optimization. See details.

Method clone(): The objects of this class are cloneable with this method.

Usage:

PLNfit_spherical\$clone(deep = FALSE)

Arguments:

deep Whether to make a deep clone.

Examples

```
## Not run:
data(trichoptera)
trichoptera <- prepare_data(trichoptera$Abundance, trichoptera$Covariate)
myPLN <- PLN(Abundance ~ 1, data = trichoptera)
class(myPLN)
print(myPLN)
```

End(Not run)

PLNLDA

Poisson lognormal model towards Linear Discriminant Analysis

Description

Fit the Poisson lognormal for LDA with a variational algorithm. Use the (g)lm syntax for model specification (covariates, offsets).

Usage

```
PLNLDA(formula, data, subset, weights, grouping, control = PLN_param())
```

Arguments

formula	an object of class "formula": a symbolic description of the model to be fitted.
data	an optional data frame, list or environment (or object coercible by as.data.frame to a data frame) containing the variables in the model. If not found in data, the variables are taken from environment(formula), typically the environment from which lm is called.
subset	an optional vector specifying a subset of observations to be used in the fitting process.
weights	an optional vector of observation weights to be used in the fitting process.
grouping	a factor specifying the class of each observation used for discriminant analysis.
control	a list-like structure for controlling the optimization, with default generated by PLN_param(). See the associated documentation

Details

The parameter control is a list controlling the optimization with the following entries:

- "covariance" character setting the model for the covariance matrix. Either "full" or "spherical". Default is "full".
- "trace" integer for verbosity.
- "inception" Set up the initialization. By default, the model is initialized with a multivariate linear model applied on log-transformed data. However, the user can provide a PLNfit (typically obtained from a previous fit), which often speed up the inference.
- "ftol_rel" stop when an optimization step changes the objective function by less than ftol multiplied by the absolute value of the parameter. Default is 1e-8
- "ftol_abs" stop when an optimization step changes the objective function by less than ftol multiplied by the absolute value of the parameter. Default is 0
- "xtol_rel" stop when an optimization step changes every parameters by less than xtol multiplied by the absolute value of the parameter. Default is 1e-6
- "xtol_abs" stop when an optimization step changes every parameters by less than xtol multiplied by the absolute value of the parameter. Default is 0
- "maxeval" stop when the number of iteration exceeds maxeval. Default is 10000
- "maxtime" stop when the optimization time (in seconds) exceeds maxtime. Default is -1 (no restriction)
- "algorithm" the optimization method used by NLOPT among LD type, i.e. "CCSAQ", "MMA", "LBFGS", "VAR1", "VAR2". See NLOPT documentation for further details. Default is "CC-SAQ".

Value

an R6 object with class PLNLDAfit()

PLNLDAfit

See Also

The class **PLNLDAfit**

Examples

```
data(trichoptera)
trichoptera <- prepare_data(trichoptera$Abundance, trichoptera$Covariate)
myPLNLDA <- PLNLDA(Abundance ~ 1, grouping = Group, data = trichoptera)</pre>
```

PLNLDAfit

An R6 Class to represent a PLNfit in a LDA framework

Description

The function PLNLDA() produces an instance of an object with class PLNLDAfit.

This class comes with a set of methods, some of them being useful for the user: See the documentation for the methods inherited by PLNfit(), the plot() method for LDA visualization and predict() method for prediction

Super class

PLNmodels::PLNfit -> PLNLDAfit

Active bindings

rank the dimension of the current model

nb_param number of parameters in the current PLN model

model_par a list with the matrices associated with the estimated parameters of the PLN model: B (covariates), Sigma (latent covariance), C (latent loadings), P (latent position) and Mu (group means)

percent_var the percent of variance explained by each axis

corr_map a matrix of correlations to plot the correlation circles

scores a matrix of scores to plot the individual factor maps

group_means a matrix of group mean vectors in the latent space.

Methods

Public methods:

- PLNLDAfit\$new()
- PLNLDAfit\$optimize()
- PLNLDAfit\$postTreatment()
- PLNLDAfit\$setVisualization()
- PLNLDAfit\$plot_individual_map()
- PLNLDAfit\$plot_correlation_map()

- PLNLDAfit\$plot_LDA()
- PLNLDAfit\$predict()
- PLNLDAfit\$show()
- PLNLDAfit\$clone()

Method new(): Initialize a PLNLDAfit object

```
Usage:
PLNLDAfit$new(
  grouping,
  responses,
  covariates,
  offsets,
  weights,
  formula,
  control
)
```

Arguments:

grouping a factor specifying the class of each observation used for discriminant analysis.

responses the matrix of responses (called Y in the model). Will usually be extracted from the corresponding field in PLNfamily-class

covariates design matrix (called X in the model). Will usually be extracted from the corresponding field in PLNfamily-class

- offsets offset matrix (called O in the model). Will usually be extracted from the corresponding field in PLNfamily-class
- weights an optional vector of observation weights to be used in the fitting process.
- formula model formula used for fitting, extracted from the formula in the upper-level call control list controlling the optimization and the model

Method optimize(): Compute group means and axis of the LDA (noted B in the model) in the latent space, update corresponding fields

Usage:

PLNLDAfit\$optimize(grouping, responses, covariates, offsets, weights, config)

Arguments:

grouping a factor specifying the class of each observation used for discriminant analysis.

- responses the matrix of responses (called Y in the model). Will usually be extracted from the corresponding field in PLNfamily-class
- covariates design matrix. Automatically built from the covariates and the formula from the call
- offsets offset matrix (called O in the model). Will usually be extracted from the corresponding field in PLNfamily-class

weights an optional vector of observation weights to be used in the fitting process.

- config list controlling the optimization
- X Abundance matrix.

Method postTreatment(): Update R2, fisher and std_err fields and visualization

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Usage:

PLNLDAfit\$postTreatment(grouping, responses, covariates, offsets, config)

Arguments:

grouping a factor specifying the class of each observation used for discriminant analysis.

- responses the matrix of responses (called Y in the model). Will usually be extracted from the corresponding field in PLNfamily-class
- covariates design matrix (called X in the model). Will usually be extracted from the corresponding field in PLNfamily-class
- offsets offset matrix (called O in the model). Will usually be extracted from the corresponding field in PLNfamily-class

config list controlling the post-treatment

Method setVisualization(): Compute LDA scores in the latent space and update corresponding fields.

Usage:

PLNLDAfit\$setVisualization(scale.unit = FALSE)

Arguments:

scale.unit Logical. Should LDA scores be rescaled to have unit variance

Method plot_individual_map(): Plot the factorial map of the LDA

```
Usage:
PLNLDAfit$plot_individual_map(
  axes = 1:min(2, self$rank),
  main = "Individual Factor Map",
  plot = TRUE
)
```

Arguments:

- axes numeric, the axes to use for the plot when map = "individual" or "variable". Default it c(1,min(rank))
- main character. A title for the single plot (individual or variable factor map). If NULL (the default), an hopefully appropriate title will be used.

plot logical. Should the plot be displayed or sent back as ggplot object

```
Returns: a ggplot graphic
```

Method plot_correlation_map(): Plot the correlation circle of a specified axis for a PLNLDAfit object

```
Usage:
PLNLDAfit$plot_correlation_map(
  axes = 1:min(2, self$rank),
  main = "Variable Factor Map",
  cols = "default",
  plot = TRUE
)
```

Arguments:

- axes numeric, the axes to use for the plot when map = "individual" or "variable". Default it c(1,min(rank))
- main character. A title for the single plot (individual or variable factor map). If NULL (the default), an hopefully appropriate title will be used.
- cols a character, factor or numeric to define the color associated with the variables. By default, all variables receive the default color of the current palette.
- plot logical. Should the plot be displayed or sent back as ggplot object

Returns: a ggplot graphic

Method plot_LDA(): Plot a summary of the PLNLDAfit object

```
Usage:
PLNLDAfit$plot_LDA(
   nb_axes = min(3, self$rank),
   var_cols = "default",
   plot = TRUE
)
```

Arguments:

- nb_axes scalar: the number of axes to be considered when map = "both". The default is min(3,rank).
- var_cols a character, factor or numeric to define the color associated with the variables. By default, all variables receive the default color of the current palette.

plot logical. Should the plot be displayed or sent back as ggplot object

```
Returns: a grob object
```

Method predict(): Predict group of new samples

```
Usage:
PLNLDAfit$predict(
    newdata,
    type = c("posterior", "response", "scores"),
    scale = c("log", "prob"),
    prior = NULL,
    control = PLN_param(backend = "nlopt"),
    envir = parent.frame()
)
```

Arguments:

newdata A data frame in which to look for variables, offsets and counts with which to predict.

- type The type of prediction required. The default are posterior probabilities for each group (in either unnormalized log-scale or natural probabilities, see "scale" for details), "response" is the group with maximal posterior probability and "scores" is the average score along each separation axis in the latent space, with weights equal to the posterior probabilities.
- scale The scale used for the posterior probability. Either log-scale ("log", default) or natural probabilities summing up to 1 ("prob").
- prior User-specified prior group probabilities in the new data. If NULL (default), prior probabilities are computed from the learning set.
- control a list for controlling the optimization. See PLN() for details.

envir Environment in which the prediction is evaluated

Method show(): User friendly print method

Usage:
PLNLDAfit\$show()

Method clone(): The objects of this class are cloneable with this method.

Usage: PLNLDAfit\$clone(deep = FALSE) Arguments: deep Whether to make a deep clone.

See Also

The function PLNLDA.

Examples

```
## Not run:
data(trichoptera)
trichoptera <- prepare_data(trichoptera$Abundance, trichoptera$Covariate)
myPLNLDA <- PLNLDA(Abundance ~ 1, grouping = Group, data = trichoptera)
class(myPLNLDA)
print(myPLNLDA)
```

End(Not run)

PLNLDAfit_diagonal An R6 Class to represent a PLNfit in a LDA framework with diagonal covariance

Description

The function PLNLDA() produces an instance of an object with class PLNLDAfit.

This class comes with a set of methods, some of them being useful for the user: See the documentation for the methods inherited by PLNfit(), the plot() method for LDA visualization and predict() method for prediction

Super classes

PLNmodels::PLNfit -> PLNmodels::PLNLDAfit -> PLNLDAfit_diagonal

Active bindings

vcov_model character: the model used for the residual covariance

Methods

Public methods:

- PLNLDAfit_diagonal\$new()
- PLNLDAfit_diagonal\$clone()

Method new(): Initialize a PLNfit model

```
Usage:
PLNLDAfit_diagonal$new(
  grouping,
  responses,
  covariates,
  offsets,
  weights,
  formula,
  control
)
```

Arguments:

grouping a factor specifying the class of each observation used for discriminant analysis.

responses the matrix of responses (called Y in the model). Will usually be extracted from the corresponding field in PLNfamily-class

- covariates design matrix (called X in the model). Will usually be extracted from the corresponding field in PLNfamily-class
- offsets offset matrix (called O in the model). Will usually be extracted from the corresponding field in PLNfamily-class
- weights an optional vector of observation weights to be used in the fitting process.
- formula model formula used for fitting, extracted from the formula in the upper-level call control a list for controlling the optimization. See details.

Method clone(): The objects of this class are cloneable with this method.

Usage:

PLNLDAfit_diagonal\$clone(deep = FALSE)

Arguments:

deep Whether to make a deep clone.

Examples

```
## Not run:
data(trichoptera)
trichoptera <- prepare_data(trichoptera$Abundance, trichoptera$Covariate)
myPLNLDA <- PLNLDA(Abundance ~ 1, data = trichoptera, control = PLN_param(covariance = "diagonal"))
class(myPLNLDA)
print(myPLNLDA)
```

End(Not run)

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Description

Helper to define list of parameters to control the PLNLDA fit. All arguments have defaults.

Usage

```
PLNLDA_param(
    backend = c("nlopt", "torch"),
    trace = 1,
    covariance = c("full", "diagonal", "spherical"),
    config_post = list(),
    config_optim = list(),
    inception = NULL
)
```

Arguments

backend	optimization back used, either "nlopt" or "torch". Default is "nlopt"
trace	a integer for verbosity.
covariance	character setting the model for the covariance matrix. Either "full", "diagonal" or "spherical". Default is "full".
config_post	a list for controlling the post-treatments (optional bootstrap, jackknife, R2, etc.). See details
config_optim	a list for controlling the optimizer (either "nlopt" or "torch" backend). See de- tails
inception	Set up the parameters initialization: by default, the model is initialized with a multivariate linear model applied on log-transformed data, and with the same formula as the one provided by the user. However, the user can provide a PLNfit (typically obtained from a previous fit), which sometimes speeds up the inference.

Details

The list of parameters config_optim controls the optimizers. When "nlopt" is chosen the following entries are relevant

- "algorithm" the optimization method used by NLOPT among LD type, e.g. "CCSAQ", "MMA", "LBFGS". See NLOPT documentation for further details. Default is "CCSAQ".
- "maxeval" stop when the number of iteration exceeds maxeval. Default is 10000
- "ftol_rel" stop when an optimization step changes the objective function by less than ftol multiplied by the absolute value of the parameter. Default is 1e-8

- "xtol_rel" stop when an optimization step changes every parameters by less than xtol multiplied by the absolute value of the parameter. Default is 1e-6
- "ftol_abs" stop when an optimization step changes the objective function by less than ftol_abs. Default is 0.0 (disabled)
- "xtol_abs" stop when an optimization step changes every parameters by less than xtol_abs. Default is 0.0 (disabled)
- "maxtime" stop when the optimization time (in seconds) exceeds maxtime. Default is -1 (disabled)

When "torch" backend is used, with the following entries are relevant:

- "maxeval" stop when the number of iteration exceeds maxeval. Default is 10000
- "ftol_rel" stop when an optimization step changes the objective function by less than ftol multiplied by the absolute value of the parameter. Default is 1e-8
- "xtol_rel" stop when an optimization step changes every parameters by less than xtol multiplied by the absolute value of the parameter. Default is 1e-6

The list of parameters config_post controls the post-treatment processing, with the following entries:

- jackknife boolean indicating whether jackknife should be performed to evaluate bias and variance of the model parameters. Default is FALSE.
- bootstrap integer indicating the number of bootstrap resamples generated to evaluate the variance of the model parameters. Default is 0 (inactivated).
- variational_var boolean indicating whether variational Fisher information matrix should be computed to estimate the variance of the model parameters (highly underestimated). Default is TRUE.
- rsquared boolean indicating whether approximation of R2 based on deviance should be computed. Default is TRUE

Value

list of parameters configuring the fit.

PLNmixture Poisson lognormal mixture model

Description

Fit the mixture variants of the Poisson lognormal with a variational algorithm. Use the (g)lm syntax for model specification (covariates, offsets).

Usage

```
PLNmixture(formula, data, subset, clusters = 1:5, control = PLNmixture_param())
```

PLNmixturefamily

Arguments

formula	an object of class "formula": a symbolic description of the model to be fitted.
data	an optional data frame, list or environment (or object coercible by as.data.frame to a data frame) containing the variables in the model. If not found in data, the variables are taken from environment(formula), typically the environment from which lm is called.
subset	an optional vector specifying a subset of observations to be used in the fitting process.
clusters	a vector of integer containing the successive number of clusters (or components) to be considered
control	a list-like structure for controlling the optimization, with default generated by PLNmixture_param(). See the associated documentation for details.

Value

an R6 object with class ${\tt PLNmixturefamily},$ which contains a collection of models with class ${\tt PLNmixturefit}$

See Also

The classes PLNmixturefamily, PLNmixturefit and PLNmixture_param()

Examples

PLNmixturefamily An R6 Class to represent a collection of PLNmixturefit

Description

The function PLNmixture() produces an instance of this class.

This class comes with a set of methods, some of them being useful for the user: See the documentation for getBestModel(), getModel() and plot().

Super class

PLNmodels::PLNfamily -> PLNmixturefamily

Active bindings

clusters vector indicating the number of clusters considered is the successively fitted models

Methods

Public methods:

- PLNmixturefamily\$new()
- PLNmixturefamily\$optimize()
- PLNmixturefamily\$smooth()
- PLNmixturefamily\$plot()
- PLNmixturefamily\$plot_objective()
- PLNmixturefamily\$getBestModel()
- PLNmixturefamily\$show()
- PLNmixturefamily\$print()
- PLNmixturefamily\$clone()

Method new(): Initialize all models in the collection.

```
Usage:
PLNmixturefamily$new(
   clusters,
   responses,
   covariates,
   offsets,
   formula,
   control
)
```

Arguments:

clusters the dimensions of the successively fitted models responses the matrix of responses common to every models covariates the matrix of covariates common to every models offsets the matrix of offsets common to every models formula model formula used for fitting, extracted from the formula in the upper-level call control a list for controlling the optimization. See details. control a list for controlling the optimization. See details. Method optimize(): Call to the optimizer on all models of the collection

Usage:

PLNmixturefamily\$optimize(config)

Arguments:

config a list for controlling the optimization

Method smooth(): function to restart clustering to avoid local minima by smoothing the loglikelihood values as a function of the number of clusters

Usage:
PLNmixturefamily\$smooth(control)

Arguments:

control a list to control the smoothing process

Method plot(): Lineplot of selected criteria for all models in the collection

Usage:

PLNmixturefamily\$plot(criteria = c("loglik", "BIC", "ICL"), reverse = FALSE)

Arguments:

- criteria A valid model selection criteria for the collection of models. Any of "loglik", "BIC" or "ICL" (all).
- reverse A logical indicating whether to plot the value of the criteria in the "natural" direction (loglik 0.5 penalty) or in the "reverse" direction (-2 loglik + penalty). Default to FALSE, i.e use the natural direction, on the same scale as the log-likelihood..

Returns: A ggplot2 object

Method plot_objective(): Plot objective value of the optimization problem along the penalty path

Usage:
PLNmixturefamily\$plot_objective()

Returns: a ggplot graph

Method getBestModel(): Extract best model in the collection

Usage:

PLNmixturefamily\$getBestModel(crit = c("BIC", "ICL", "loglik"))

Arguments:

crit a character for the criterion used to performed the selection. Either "BIC", "ICL" or "loglik". Default is ICL

Returns: a PLNmixturefit object

Method show(): User friendly print method

Usage:

PLNmixturefamily\$show()

Method print(): User friendly print method

Usage:
PLNmixturefamily\$print()

Method clone(): The objects of this class are cloneable with this method.

Usage: PLNmixturefamily\$clone(deep = FALSE) Arguments: deep Whether to make a deep clone.

See Also

The function PLNmixture, the class PLNmixturefit

PLNmixturefit An R6 Class to represent a PLNfit in a mixture framework

Description

The function PLNmixture produces a collection of models which are instances of object with class PLNmixturefit. A PLNmixturefit (say, with k components) is itself a collection of k PLNfit.

This class comes with a set of methods, some of them being useful for the user: See the documentation for ...

Active bindings

- n number of samples
- p number of dimensions of the latent space
- k number of components
- d number of covariates

components components of the mixture (PLNfits)

latent a matrix: values of the latent vector (Z in the model)

latent_pos a matrix: values of the latent position vector (Z) without covariates effects or offset

posteriorProb matrix of posterior probability for cluster belonging

memberships vector for cluster index

mixtureParam vector of cluster proportions

optim_par a list with parameters useful for monitoring the optimization

nb_param number of parameters in the current PLN model

entropy_clustering Entropy of the variational distribution of the cluster (multinomial)

entropy_latent Entropy of the variational distribution of the latent vector (Gaussian)

entropy Full entropy of the variational distribution (latent vector + clustering)

loglik variational lower bound of the loglikelihood

PLNmixturefit

- loglik_vec element-wise variational lower bound of the loglikelihood
- BIC variational lower bound of the BIC
- ICL variational lower bound of the ICL (include entropy of both the clustering and latent distributions)

R_squared approximated goodness-of-fit criterion

criteria a vector with loglik, BIC, ICL, and number of parameters

model_par a list with the matrices of parameters found in the model (Theta, Sigma, Mu and Pi)

vcov_model character: the model used for the covariance (either "spherical", "diagonal" or "full")

fitted a matrix: fitted values of the observations (A in the model)

group_means a matrix of group mean vectors in the latent space.

Methods

Public methods:

- PLNmixturefit\$new()
- PLNmixturefit\$optimize()
- PLNmixturefit\$predict()
- PLNmixturefit\$plot_clustering_data()
- PLNmixturefit\$plot_clustering_pca()
- PLNmixturefit\$postTreatment()
- PLNmixturefit\$show()
- PLNmixturefit\$print()
- PLNmixturefit\$clone()

Method new(): Optimize a the

Initialize a PLNmixturefit model

```
Usage:
PLNmixturefit$new(
  responses,
  covariates,
  offsets,
  posteriorProb,
  formula,
  control
)
```

Arguments:

responses the matrix of responses common to every models covariates the matrix of covariates common to every models offsets the matrix of offsets common to every models posteriorProb matrix ofposterior probability for cluster belonging formula model formula used for fitting, extracted from the formula in the upper-level call control a list for controlling the optimization. **Method** optimize(): Optimize a PLNmixturefit model

Usage:

```
PLNmixturefit$optimize(responses, covariates, offsets, config)
```

Arguments:

responses the matrix of responses common to every models covariates the matrix of covariates common to every models offsets the matrix of offsets common to every models config a list for controlling the optimization

Method predict(): Predict group of new samples

```
Usage:
PLNmixturefit$predict(
    newdata,
    type = c("posterior", "response", "position"),
    prior = matrix(rep(1/self$k, self$k), nrow(newdata), self$k, byrow = TRUE),
    control = PLNmixture_param(),
    envir = parent.frame()
)
```

Arguments:

newdata A data frame in which to look for variables, offsets and counts with which to predict.

- type The type of prediction required. The default posterior are posterior probabilities for each group, response is the group with maximal posterior probability and latent is the averaged latent coordinate (without offset and nor covariate effects), with weights equal to the posterior probabilities.
- prior User-specified prior group probabilities in the new data. The default uses a uniform prior.

control a list-like structure for controlling the fit. See PLNmixture_param() for details. envir Environment in which the prediction is evaluated

Method plot_clustering_data(): Plot the matrix of expected mean counts (without offsets, without covariate effects) reordered according the inferred clustering

Usage:

```
PLNmixturefit$plot_clustering_data(
  main = "Expected counts reorder by clustering",
  plot = TRUE,
  log_scale = TRUE
)
```

Arguments:

main character. A title for the plot. An hopefully appropriate title will be used by default.

plot logical. Should the plot be displayed or sent back as ggplot object

log_scale logical. Should the color scale values be log-transform before plotting? Default is TRUE.

Returns: a ggplot graphic

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Method plot_clustering_pca(): Plot the individual map of a PCA performed on the latent coordinates, where individuals are colored according to the memberships

```
Usage:
PLNmixturefit$plot_clustering_pca(
  main = "Clustering labels in Individual Factor Map",
  plot = TRUE
)
```

Arguments:

main character. A title for the plot. An hopefully appropriate title will be used by default. plot logical. Should the plot be displayed or sent back as ggplot object

Returns: a ggplot graphic

Method postTreatment(): Update fields after optimization

```
Usage:
PLNmixturefit$postTreatment(
  responses,
  covariates,
  offsets,
  weights,
  config,
  nullModel
)
```

Arguments:

responses the matrix of responses common to every models

covariates the matrix of covariates common to every models

offsets the matrix of offsets common to every models

weights an optional vector of observation weights to be used in the fitting process.

```
config a list for controlling the post-treatment
```

nullModel null model used for approximate R2 computations. Defaults to a GLM model with same design matrix but not latent variable.

Method show(): User friendly print method

Usage:
PLNmixturefit\$show()

Method print(): User friendly print method

Usage:
PLNmixturefit\$print()

Method clone(): The objects of this class are cloneable with this method.

Usage:
PLNmixturefit\$clone(deep = FALSE)

Arguments:

deep Whether to make a deep clone.

The function PLNmixture, the class PLNmixturefamily

PLNmixture_param Control of a PLNmixture fit

Description

Helper to define list of parameters to control the PLNmixture fit. All arguments have defaults.

Usage

```
PLNmixture_param(
   backend = "nlopt",
   trace = 1,
   covariance = "spherical",
   init_cl = "kmeans",
   smoothing = "both",
   config_post = list(),
   config_optim = list(),
   inception = NULL
)
```

Arguments

backend	optimization back used, either "nlopt" or "torch". Default is "nlopt"
trace	a integer for verbosity.
covariance	character setting the model for the covariance matrices of the mixture compo- nents. Either "full", "diagonal" or "spherical". Default is "spherical".
init_cl	The initial clustering to apply. Either, 'kmeans', CAH' or a user defined clus- tering given as a list of clusterings, the size of which is equal to the number of clusters considered. Default is 'kmeans'.
smoothing	The smoothing to apply. Either, 'none', forward', 'backward' or 'both'. Default is 'both'.
config_post	a list for controlling the post-treatments (optional bootstrap, jackknife, R2, etc.). See details
config_optim	a list for controlling the optimizer (either "nlopt" or "torch" backend). See de- tails
inception	Set up the parameters initialization: by default, the model is initialized with a multivariate linear model applied on log-transformed data, and with the same formula as the one provided by the user. However, the user can provide a PLNfit (typically obtained from a previous fit), which sometimes speeds up the inference.

Details

The list of parameters config_optim controls the optimizers. When "nlopt" is chosen the following entries are relevant

- "it_smooth" number of forward/backward iteration of smoothing. Default is 2.
- "algorithm" the optimization method used by NLOPT among LD type, e.g. "CCSAQ", "MMA", "LBFGS". See NLOPT documentation for further details. Default is "CCSAQ".
- "maxeval" stop when the number of iteration exceeds maxeval. Default is 10000
- "ftol_rel" stop when an optimization step changes the objective function by less than ftol multiplied by the absolute value of the parameter. Default is 1e-8
- "xtol_rel" stop when an optimization step changes every parameters by less than xtol multiplied by the absolute value of the parameter. Default is 1e-6
- "ftol_out" outer solver stops when an optimization step changes the objective function by less than xtol multiply by the absolute value of the parameter. Default is 1e-6
- "maxit_out" outer solver stops when the number of iteration exceeds out.maxit. Default is 50
- "ftol_abs" stop when an optimization step changes the objective function by less than ftol_abs. Default is 0.0 (disabled)
- "xtol_abs" stop when an optimization step changes every parameters by less than xtol_abs. Default is 0.0 (disabled)
- "maxtime" stop when the optimization time (in seconds) exceeds maxtime. Default is -1 (disabled)

When "torch" backend is used, with the following entries are relevant:

- "maxeval" stop when the number of iteration exceeds maxeval. Default is 10000
- "ftol_rel" stop when an optimization step changes the objective function by less than ftol multiplied by the absolute value of the parameter. Default is 1e-8
- "xtol_rel" stop when an optimization step changes every parameters by less than xtol multiplied by the absolute value of the parameter. Default is 1e-6
- "ftol_out" outer solver stops when an optimization step changes the objective function by less than xtol multiply by the absolute value of the parameter. Default is 1e-6
- "maxit_out" outer solver stops when the number of iteration exceeds out.maxit. Default is 50

The list of parameters config_post controls the post-treatment processing, with the following entries:

- jackknife boolean indicating whether jackknife should be performed to evaluate bias and variance of the model parameters. Default is FALSE.
- bootstrap integer indicating the number of bootstrap resamples generated to evaluate the variance of the model parameters. Default is 0 (inactivated).
- variational_var boolean indicating whether variational Fisher information matrix should be computed to estimate the variance of the model parameters (highly underestimated). Default is FALSE.
- rsquared boolean indicating whether approximation of R2 based on deviance should be computed. Default is FALSE

Value

list of parameters configuring the fit.

```
PLNmodels
```

PLNmodels

Description

The Poisson lognormal model and variants can be used for a variety of multivariate problems when count data are at play (including PCA or LDA for count data, network inference). This package implements efficient variational algorithms to fit such models accompanied with a set of functions for visualization and diagnostic.

Multivariate Poisson lognormal model (aka PLN)

See the main function PLN() and the associated methods for manipulation.

Also try vignette("PLN_trichoptera", package="PLNmodels") for an overview.

Rank Constrained Poisson lognormal for Poisson Principal Component Analysis (aka PLNPCA)

See the main function PLNPCA() and the associated methods for manipulation.

The Poisson PCA and the associated variational inference is fully explained in Chiquet el al (2018), see reference below.

Also try vignette("PLNPCA_trichoptera", package="PLNmodels") for an overview.

Sparse Poisson lognormal model for sparse covariance inference for counts (aka PLNnetwork)

See the main function PLNnetwork() and the associated methods for manipulation. Also try vignette("PLNnetwork_trichoptera", package="PLNmodels") for an overview.

Poisson lognormal discriminant analysis (aka PLNLDA)

See the main function PLNLDA() and the associated methods for manipulation. Also try vignette("PLNLDA_trichoptera", package="PLNmodels") for an overview.

Mixtures of Poisson lognormal models for model-based clustering (aka PLNmixture)

See the main function PLNmixture() and the associated methods for manipulation. Also try vignette("PLNmixture_trichoptera", package="PLNmodels") for an overview.

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PLNnetwork

Description

Fit the sparse inverse covariance variant of the Poisson lognormal with a variational algorithm. Use the (g)lm syntax for model specification (covariates, offsets).

Usage

```
PLNnetwork(
   formula,
   data,
   subset,
   weights,
   penalties = NULL,
   control = PLNnetwork_param()
)
```

Arguments

formula	an object of class "formula": a symbolic description of the model to be fitted.
data	an optional data frame, list or environment (or object coercible by as.data.frame to a data frame) containing the variables in the model. If not found in data, the variables are taken from environment(formula), typically the environment from which lm is called.
subset	an optional vector specifying a subset of observations to be used in the fitting process.
weights	an optional vector of observation weights to be used in the fitting process.
penalties	an optional vector of positive real number controlling the level of sparsity of the underlying network. if NULL (the default), will be set internally. See PLNnetwork_param() for additional tuning of the penalty.
control	a list-like structure for controlling the optimization, with default generated by PLNnetwork_param(). See the corresponding documentation for details;

Value

an R6 object with class ${\sf PLNnetworkfamily},$ which contains a collection of models with class ${\sf PLNnetworkfit}$

See Also

The classes PLNnetworkfamily and PLNnetworkfit, and the and the configuration function PLNnetwork_param().

Examples

```
data(trichoptera)
trichoptera <- prepare_data(trichoptera$Abundance, trichoptera$Covariate)
fits <- PLNnetwork(Abundance ~ 1, data = trichoptera)</pre>
```

PLNnetworkfamily An R6 Class to represent a collection of PLNnetworkfit

Description

The function PLNnetwork() produces an instance of this class.

This class comes with a set of methods, some of them being useful for the user: See the documentation for getBestModel(), getModel() and plot()

Super class

PLNmodels::PLNfamily -> PLNnetworkfamily

Active bindings

penalties the sparsity level of the network in the successively fitted models

- stability_path the stability path of each edge as returned by the stars procedure
- stability mean edge stability along the penalty path
- criteria a data frame with the values of some criteria (approximated log-likelihood, (E)BIC, ICL and R2, stability) for the collection of models / fits BIC, ICL and EBIC are defined so that they are on the same scale as the model log-likelihood, i.e. with the form, loglik 0.5 penalty

Methods

Public methods:

- PLNnetworkfamily\$new()
- PLNnetworkfamily\$optimize()
- PLNnetworkfamily\$stability_selection()
- PLNnetworkfamily\$coefficient_path()
- PLNnetworkfamily\$getBestModel()
- PLNnetworkfamily\$plot()
- PLNnetworkfamily\$plot_stars()
- PLNnetworkfamily\$plot_objective()
- PLNnetworkfamily\$show()
- PLNnetworkfamily\$clone()

Method new(): Initialize all models in the collection

Usage:

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```
PLNnetworkfamily$new(
    penalties,
    responses,
    covariates,
    offsets,
    weights,
    formula,
    control
}
```

```
)
```

Arguments:

penalties a vector of positive real number controlling the level of sparsity of the underlying network.

responses the matrix of responses common to every models

covariates the matrix of covariates common to every models

offsets the matrix of offsets common to every models

weights the vector of observation weights

formula model formula used for fitting, extracted from the formula in the upper-level call control a list for controlling the optimization.

Returns: Update current PLNnetworkfit with smart starting values

Method optimize(): Call to the C++ optimizer on all models of the collection

Usage:

PLNnetworkfamily\$optimize(config)

Arguments:

config a list for controlling the optimization.

Method stability_selection(): Compute the stability path by stability selection

```
Usage:
```

```
PLNnetworkfamily$stability_selection(
   subsamples = NULL,
   control = PLNnetwork_param()
)
```

Arguments:

- subsamples a list of vectors describing the subsamples. The number of vectors (or list length) determines the number of subsamples used in the stability selection. Automatically set to 20 subsamples with size $10 \times \text{sqrt}(n)$ if $n \ge 144$ and $0.8 \times n$ otherwise following Liu et al. (2010) recommendations.
- control a list controlling the main optimization process in each call to PLNnetwork. See PLNnetwork() for details.

Method coefficient_path(): Extract the regularization path of a PLNnetworkfamily

Usage:

PLNnetworkfamily\$coefficient_path(precision = TRUE, corr = TRUE)

Arguments:

- precision Logical. Should the regularization path be extracted from the precision matrix Omega (TRUE, default) or from the variance matrix Sigma (FALSE)
- corr Logical. Should the matrix be transformed to (partial) correlation matrix before extraction? Defaults to TRUE

Method getBestModel(): Extract the best network in the family according to some criteria

```
Usage:
PLNnetworkfamily$getBestModel(
  crit = c("BIC", "EBIC", "StARS"),
  stability = 0.9
)
```

Arguments:

- crit character. Criterion used to perform the selection. Is "StARS" is chosen but \$stability field is empty, will compute stability path.
- stability Only used for "StARS" criterion. A scalar indicating the target stability (= 1 2 beta) at which the network is selected. Default is 0.9.

Method plot(): Display various outputs (goodness-of-fit criteria, robustness, diagnostic) associated with a collection of PLNnetwork fits (a PLNnetworkfamily)

```
Usage:
PLNnetworkfamily$plot(
  criteria = c("loglik", "pen_loglik", "BIC", "EBIC"),
  reverse = FALSE,
  log.x = TRUE
)
```

Arguments:

- criteria vector of characters. The criteria to plot in c("loglik", "pen_loglik", "BIC", "EBIC"). Defaults to all of them.
- reverse A logical indicating whether to plot the value of the criteria in the "natural" direction (loglik 0.5 penalty) or in the "reverse" direction (-2 loglik + penalty). Default to FALSE, i.e use the natural direction, on the same scale as the log-likelihood..

log.x logical: should the x-axis be represented in log-scale? Default is TRUE.

Returns: a ggplot graph

Method plot_stars(): Plot stability path

Usage:

PLNnetworkfamily\$plot_stars(stability = 0.9, log.x = TRUE)

Arguments:

stability scalar: the targeted level of stability in stability plot. Default is 0.9. log.x logical: should the x-axis be represented in log-scale? Default is TRUE.

Returns: a ggplot graph

Method plot_objective(): Plot objective value of the optimization problem along the penalty path

PLNnetworkfit

Usage:
PLNnetworkfamily\$plot_objective()
Returns: a ggplot graph

Method show(): User friendly print method

Usage:
PLNnetworkfamily\$show()

Method clone(): The objects of this class are cloneable with this method.

Usage:
PLNnetworkfamily\$clone(deep = FALSE)

Arguments:

deep Whether to make a deep clone.

See Also

The function PLNnetwork(), the class PLNnetworkfit

Examples

```
data(trichoptera)
trichoptera <- prepare_data(trichoptera$Abundance, trichoptera$Covariate)
fits <- PLNnetwork(Abundance ~ 1, data = trichoptera)
class(fits)</pre>
```

PLNnetworkfit

An R6 Class to represent a PLNfit in a sparse inverse covariance framework

Description

The function PLNnetwork() produces a collection of models which are instances of object with class PLNnetworkfit. This class comes with a set of methods, some of them being useful for the user: See the documentation for plot() and methods inherited from PLNfit.

Super classes

```
PLNmodels::PLNfit -> PLNmodels::PLNfit_fixedcov -> PLNnetworkfit
```

Active bindings

vcov_model character: the model used for the residual covariance

penalty the global level of sparsity in the current model

penalty_weights a matrix of weights controlling the amount of penalty element-wise.

n_edges number of edges if the network (non null coefficient of the sparse precision matrix)

nb_param number of parameters in the current PLN model

pen_loglik variational lower bound of the l1-penalized loglikelihood

EBIC variational lower bound of the EBIC

density proportion of non-null edges in the network

criteria a vector with loglik, penalized loglik, BIC, EBIC, ICL, R_squared, number of parameters, number of edges, and graph density

Methods

Public methods:

- PLNnetworkfit\$new()
- PLNnetworkfit\$update()
- PLNnetworkfit\$optimize()
- PLNnetworkfit\$latent_network()
- PLNnetworkfit\$plot_network()
- PLNnetworkfit\$show()
- PLNnetworkfit\$clone()

Method new(): Initialize a PLNnetworkfit object

```
Usage:
PLNnetworkfit$new(
    penalty,
    penalty_weights,
    responses,
    covariates,
    offsets,
    weights,
    formula,
    control
)
```

Arguments:

penalty a positive real number controlling the level of sparsity of the underlying network.

- penalty_weights either a single or a list of p x p matrix of weights (default filled with 1) to adapt the amount of shrinkage to each pairs of node. Must be symmetric with positive values.
- responses the matrix of responses (called Y in the model). Will usually be extracted from the corresponding field in PLNfamily-class
- covariates design matrix (called X in the model). Will usually be extracted from the corresponding field in PLNfamily-class

- offsets offset matrix (called O in the model). Will usually be extracted from the corresponding field in PLNfamily-class
- weights an optional vector of observation weights to be used in the fitting process.
- formula model formula used for fitting, extracted from the formula in the upper-level call control a list for controlling the optimization.

Method update(): Update fields of a PLNnetworkfit object

```
Usage:
PLNnetworkfit$update(
    penalty = NA,
    B = NA,
    Sigma = NA,
    Omega = NA,
    M = NA,
    S = NA,
    Z = NA,
    A = NA,
    Ji = NA,
    R2 = NA,
    monitoring = NA
```

```
)
```

Arguments:

penalty a positive real number controlling the level of sparsity of the underlying network.

B matrix of regression matrix

Sigma variance-covariance matrix of the latent variables

Omega precision matrix of the latent variables. Inverse of Sigma.

- M matrix of mean vectors for the variational approximation
- S matrix of variance vectors for the variational approximation
- Z matrix of latent vectors (includes covariates and offset effects)
- A matrix of fitted values

Ji vector of variational lower bounds of the log-likelihoods (one value per sample)

R2 approximate R^2 goodness-of-fit criterion

monitoring a list with optimization monitoring quantities

Method optimize(): Call to the C++ optimizer and update of the relevant fields

Usage:

PLNnetworkfit\$optimize(responses, covariates, offsets, weights, config)

Arguments:

- responses the matrix of responses (called Y in the model). Will usually be extracted from the corresponding field in PLNfamily-class
- covariates design matrix (called X in the model). Will usually be extracted from the corresponding field in PLNfamily-class
- offsets offset matrix (called O in the model). Will usually be extracted from the corresponding field in PLNfamily-class

weights an optional vector of observation weights to be used in the fitting process. config a list for controlling the optimization

Method latent_network(): Extract interaction network in the latent space

Usage:

```
PLNnetworkfit$latent_network(type = c("partial_cor", "support", "precision"))
```

Arguments:

type edge value in the network. Can be "support" (binary edges), "precision" (coefficient of the precision matrix) or "partial_cor" (partial correlation between species)

Returns: a square matrix of size PLNnetworkfit\$n

Method plot_network(): plot the latent network.

```
Usage:
PLNnetworkfit$plot_network(
  type = c("partial_cor", "support"),
  output = c("igraph", "corrplot"),
  edge.color = c("#F8766D", "#00BFC4"),
  remove.isolated = FALSE,
  node.labels = NULL,
  layout = layout_in_circle,
  plot = TRUE
)
```

Arguments:

type edge value in the network. Either "precision" (coefficient of the precision matrix) or "partial_cor" (partial correlation between species).

output Output type. Either igraph (for the network) or corrplot (for the adjacency matrix)

- edge.color Length 2 color vector. Color for positive/negative edges. Default is c("#F8766D", "#00BFC4"). Only relevant for igraph output.
- remove.isolated if TRUE, isolated node are remove before plotting. Only relevant for igraph output.
- node.labels vector of character. The labels of the nodes. The default will use the column names of the response matrix.
- layout an optional igraph layout. Only relevant for igraph output.
- plot logical. Should the final network be displayed or only sent back to the user. Default is TRUE.

Method show(): User friendly print method

Usage:

PLNnetworkfit\$show()

Method clone(): The objects of this class are cloneable with this method.

Usage:

PLNnetworkfit\$clone(deep = FALSE)

Arguments:

deep Whether to make a deep clone.

PLNnetwork_param

See Also

The function PLNnetwork(), the class PLNnetworkfamily

Examples

```
## Not run:
data(trichoptera)
trichoptera <- prepare_data(trichoptera$Abundance, trichoptera$Covariate)</pre>
nets <- PLNnetwork(Abundance ~ 1, data = trichoptera)</pre>
myPLNnet <- getBestModel(nets)</pre>
class(myPLNnet)
print(myPLNnet)
## End(Not run)
```

Control of PLNnetwork fit

Description

PLNnetwork_param

Helper to define list of parameters to control the PLN fit. All arguments have defaults.

Usage

```
PLNnetwork_param(
  backend = "nlopt",
  trace = 1,
  n_penalties = 30,
 min_ratio = 0.1,
  penalize_diagonal = TRUE,
  penalty_weights = NULL,
  config_post = list(),
  config_optim = list(),
  inception = NULL
)
```

Arguments

optimization back used, either "nlopt" or "torch". Default is "nlopt"	
a integer for verbosity.	
an integer that specifies the number of values for the penalty grid when internally generated. Ignored when penalties is non NULL	
the penalty grid ranges from the minimal value that produces a sparse to this value multiplied by min_ratio. Default is 0.1.	
penalize_diagonal	
boolean: should the diagonal terms be penalized in the graphical-Lasso? Default is \ensuremath{TRUE}	

penalty_weights	
	either a single or a list of p x p matrix of weights (default filled with 1) to adapt the amount of shrinkage to each pairs of node. Must be symmetric with positive values.
config_post	a list for controlling the post-treatments (optional bootstrap, jackknife, R2, etc.). See details
config_optim	a list for controlling the optimizer (either "nlopt" or "torch" backend). See de- tails
inception	Set up the parameters initialization: by default, the model is initialized with a multivariate linear model applied on log-transformed data, and with the same formula as the one provided by the user. However, the user can provide a PLNfit (typically obtained from a previous fit), which sometimes speeds up the inference.

Details

The list of parameters config_optim controls the optimizers. When "nlopt" is chosen the following entries are relevant

- "algorithm" the optimization method used by NLOPT among LD type, e.g. "CCSAQ", "MMA", "LBFGS". See NLOPT documentation for further details. Default is "CCSAQ".
- "maxeval" stop when the number of iteration exceeds maxeval. Default is 10000
- "ftol_rel" stop when an optimization step changes the objective function by less than ftol multiplied by the absolute value of the parameter. Default is 1e-8
- "xtol_rel" stop when an optimization step changes every parameters by less than xtol multiplied by the absolute value of the parameter. Default is 1e-6
- "ftol_out" outer solver stops when an optimization step changes the objective function by less than xtol multiply by the absolute value of the parameter. Default is 1e-6
- "maxit_out" outer solver stops when the number of iteration exceeds out.maxit. Default is 50
- "ftol_abs" stop when an optimization step changes the objective function by less than ftol_abs. Default is 0.0 (disabled)
- "xtol_abs" stop when an optimization step changes every parameters by less than xtol_abs. Default is 0.0 (disabled)
- "maxtime" stop when the optimization time (in seconds) exceeds maxtime. Default is -1 (disabled)

When "torch" backend is used, with the following entries are relevant:

- "maxeval" stop when the number of iteration exceeds maxeval. Default is 10000
- "ftol_out" outer solver stops when an optimization step changes the objective function by less than xtol multiply by the absolute value of the parameter. Default is 1e-6
- "maxit_out" outer solver stops when the number of iteration exceeds out.maxit. Default is 50
- "ftol_rel" stop when an optimization step changes the objective function by less than ftol multiplied by the absolute value of the parameter. Default is 1e-8
- "xtol_rel" stop when an optimization step changes every parameters by less than xtol multiplied by the absolute value of the parameter. Default is 1e-6

PLNPCA

The list of parameters config_post controls the post-treatment processing, with the following entries:

- jackknife boolean indicating whether jackknife should be performed to evaluate bias and variance of the model parameters. Default is FALSE.
- bootstrap integer indicating the number of bootstrap resamples generated to evaluate the variance of the model parameters. Default is 0 (inactivated).
- variational_var boolean indicating whether variational Fisher information matrix should be computed to estimate the variance of the model parameters (highly underestimated). Default is FALSE.
- rsquared boolean indicating whether approximation of R2 based on deviance should be computed. Default is FALSE

Value

list of parameters configuring the fit.

Description

Fit the PCA variants of the Poisson lognormal with a variational algorithm. Use the (g)lm syntax for model specification (covariates, offsets).

Usage

```
PLNPCA(formula, data, subset, weights, ranks = 1:5, control = PLNPCA_param())
```

Arguments

formula	an object of class "formula": a symbolic description of the model to be fitted.
data	an optional data frame, list or environment (or object coercible by as.data.frame to a data frame) containing the variables in the model. If not found in data, the variables are taken from environment(formula), typically the environment from which Im is called.
subset	an optional vector specifying a subset of observations to be used in the fitting process.
weights	an optional vector of observation weights to be used in the fitting process.
ranks	a vector of integer containing the successive ranks (or number of axes to be considered)
control	a list-like structure for controlling the optimization, with default generated by PLNPCA_param(). See the associated documentation. for details.

Value

an R6 object with class PLNPCAfamily, which contains a collection of models with class PLNPCAfit

See Also

The classes PLNPCAfamily and PLNPCAfit, and the configuration function PLNPCA_param().

Examples

```
#' ## Use future to dispatch the computations on 2 workers
## Not run:
future::plan("multisession", workers = 2)
## End(Not run)
data(trichoptera)
trichoptera <- prepare_data(trichoptera$Abundance, trichoptera$Covariate)
myPCA <- PLNPCA(Abundance ~ 1 + offset(log(Offset)), data = trichoptera, ranks = 1:5)
# Shut down parallel workers
## Not run:
future::plan("sequential")
## End(Not run)
```

PLNPCAfamily An R6 Class to represent a collection of PLNPCAfit

Description

The function PLNPCA() produces an instance of this class.

This class comes with a set of methods, some of them being useful for the user: See the documentation for getBestModel(), getModel() and plot().

Super class

PLNmodels::PLNfamily -> PLNPCAfamily

Active bindings

ranks the dimensions of the successively fitted models

Methods

Public methods:

- PLNPCAfamily\$new()
- PLNPCAfamily\$optimize()
- PLNPCAfamily\$getModel()

- PLNPCAfamily\$getBestModel()
- PLNPCAfamily\$plot()
- PLNPCAfamily\$show()
- PLNPCAfamily\$clone()

Method new(): Initialize all models in the collection.

```
Usage:

PLNPCAfamily$new(

ranks,

responses,

covariates,

offsets,

weights,

formula,

control

)

Arguments:
```

ranks the dimensions of the successively fitted models responses the matrix of responses common to every models covariates the matrix of covariates common to every models offsets the matrix of offsets common to every models weights the vector of observation weights formula model formula used for fitting, extracted from the formula in the upper-level call control list controlling the optimization and the model

Method optimize(): Call to the C++ optimizer on all models of the collection

```
Usage:
PLNPCAfamily$optimize(config)
Arguments:
```

config list controlling the optimization.

Method getModel(): Extract model from collection and add "PCA" class for compatibility with factoextra::fviz()

```
Usage:
```

PLNPCAfamily\$getModel(var, index = NULL)

Arguments:

var value of the parameter (rank for PLNPCA, sparsity for PLNnetwork) that identifies the model to be extracted from the collection. If no exact match is found, the model with closest parameter value is returned with a warning.

index Integer index of the model to be returned. Only the first value is taken into account.

Returns: a PLNPCAfit object

Method getBestModel(): Extract best model in the collection

Usage:

```
PLNPCAfamily$getBestModel(crit = c("ICL", "BIC"))
```

Arguments:

crit a character for the criterion used to performed the selection. Either "ICL", "BIC". Default is ICL

Returns: a PLNPCAfit object

Method plot(): Lineplot of selected criteria for all models in the collection

Usage:

```
PLNPCAfamily$plot(criteria = c("loglik", "BIC", "ICL"), reverse = FALSE)
```

Arguments:

- criteria A valid model selection criteria for the collection of models. Any of "loglik", "BIC" or "ICL" (all).
- reverse A logical indicating whether to plot the value of the criteria in the "natural" direction (loglik penalty) or in the "reverse" direction (-2 loglik + penalty). Default to FALSE, i.e use the natural direction, on the same scale as the log-likelihood.

Returns: A ggplot2 object

Method show(): User friendly print method

Usage:
PLNPCAfamily\$show()

Method clone(): The objects of this class are cloneable with this method.

```
Usage:
PLNPCAfamily$clone(deep = FALSE)
```

Arguments:

deep Whether to make a deep clone.

See Also

The function PLNPCA(), the class PLNPCAfit()

Examples

```
data(trichoptera)
trichoptera <- prepare_data(trichoptera$Abundance, trichoptera$Covariate)
myPCAs <- PLNPCA(Abundance ~ 1 + offset(log(Offset)), data = trichoptera, ranks = 1:5)
class(myPCAs)</pre>
```

PLNPCAfit

Description

The function PLNPCA() produces a collection of models which are instances of object with class PLNPCAfit. This class comes with a set of methods, some of them being useful for the user: See the documentation for the methods inherited by PLNfit and the plot() methods for PCA visualization

Super class

PLNmodels::PLNfit -> PLNPCAfit

Active bindings

rank the dimension of the current model

vcov_model character: the model used for the residual covariance

nb_param number of parameters in the current PLN model

entropy entropy of the variational distribution

- latent_pos a matrix: values of the latent position vector (Z) without covariates effects or offset
- model_par a list with the matrices associated with the estimated parameters of the pPCA model: B (covariates), Sigma (covariance), Omega (precision) and C (loadings)

percent_var the percent of variance explained by each axis

corr_circle a matrix of correlations to plot the correlation circles

scores a matrix of scores to plot the individual factor maps (a.k.a. principal components)

- rotation a matrix of rotation of the latent space
- eig description of the eigenvalues, similar to percent_var but for use with external methods
- var a list of data frames with PCA results for the variables: coord (coordinates of the variables), cor (correlation between variables and dimensions), cos2 (Cosine of the variables) and contrib (contributions of the variable to the axes)
- ind a list of data frames with PCA results for the individuals: coord (coordinates of the individuals), cos2 (Cosine of the individuals), contrib (contributions of individuals to an axis inertia) and dist (distance of individuals to the origin).
- call Hacky binding for compatibility with factoextra functions

Methods

Public methods:

- PLNPCAfit\$new()
- PLNPCAfit\$update()
- PLNPCAfit\$optimize()
- PLNPCAfit\$optimize_vestep()

- PLNPCAfit\$project()
- PLNPCAfit\$setVisualization()
- PLNPCAfit\$postTreatment()
- PLNPCAfit\$plot_individual_map()
- PLNPCAfit\$plot_correlation_circle()
- PLNPCAfit\$plot_PCA()
- PLNPCAfit\$show()
- PLNPCAfit\$clone()

Method new(): Initialize a PLNPCAfit object

Usage:

PLNPCAfit\$new(rank, responses, covariates, offsets, weights, formula, control)

Arguments:

rank rank of the PCA (or equivalently, dimension of the latent space)

- responses the matrix of responses (called Y in the model). Will usually be extracted from the corresponding field in PLNfamily
- covariates design matrix (called X in the model). Will usually be extracted from the corresponding field in PLNfamily
- offsets offset matrix (called O in the model). Will usually be extracted from the corresponding field in PLNfamily
- weights an optional vector of observation weights to be used in the fitting process.
- formula model formula used for fitting, extracted from the formula in the upper-level call control a list for controlling the optimization. See details.

Method update(): Update a PLNPCAfit object

```
Usage:

PLNPCAfit$update(

B = NA,

Sigma = NA,

Omega = NA,

C = NA,

M = NA,

S = NA,

Z = NA,

A = NA,

Ji = NA,

R2 = NA,

monitoring = NA

)
```

Arguments:

B matrix of regression matrix

Sigma variance-covariance matrix of the latent variables

Omega precision matrix of the latent variables. Inverse of Sigma.

C matrix of PCA loadings (in the latent space)

- M matrix of mean vectors for the variational approximation
- S matrix of variance vectors for the variational approximation
- Z matrix of latent vectors (includes covariates and offset effects)
- A matrix of fitted values
- Ji vector of variational lower bounds of the log-likelihoods (one value per sample)
- R2 approximate R^2 goodness-of-fit criterion
- monitoring a list with optimization monitoring quantities

Returns: Update the current PLNPCAfit object

Method optimize(): Call to the C++ optimizer and update of the relevant fields

Usage:

PLNPCAfit\$optimize(responses, covariates, offsets, weights, config)

Arguments:

- responses the matrix of responses (called Y in the model). Will usually be extracted from the corresponding field in PLNfamily
- covariates design matrix (called X in the model). Will usually be extracted from the corresponding field in PLNfamily
- offsets offset matrix (called O in the model). Will usually be extracted from the corresponding field in PLNfamily

weights an optional vector of observation weights to be used in the fitting process.

config part of the control argument which configures the optimizer

Method optimize_vestep(): Result of one call to the VE step of the optimization procedure: optimal variational parameters (M, S) and corresponding log likelihood values for fixed model parameters (C, B). Intended to position new data in the latent space for further use with PCA.

```
Usage:
PLNPCAfit$optimize_vestep(
  covariates,
  offsets,
  responses,
  weights = rep(1, self$n),
  control = PLNPCA_param(backend = "nlopt")
)
```

Arguments:

- covariates design matrix (called X in the model). Will usually be extracted from the corresponding field in PLNfamily
- offsets offset matrix (called O in the model). Will usually be extracted from the corresponding field in PLNfamily
- responses the matrix of responses (called Y in the model). Will usually be extracted from the corresponding field in PLNfamily

weights an optional vector of observation weights to be used in the fitting process.

control a list for controlling the optimization. See details.

Returns: A list with three components:

• the matrix M of variational means,

- the matrix S2 of variational variances
- the vector log.lik of (variational) log-likelihood of each new observation

Method project(): Project new samples into the PCA space using one VE step

Usage:

PLNPCAfit\$project(newdata, control = PLNPCA_param(), envir = parent.frame())

Arguments:

newdata A data frame in which to look for variables, offsets and counts with which to predict. control a list for controlling the optimization. See PLN() for details. envir Environment in which the projection is evaluated

Returns:

• the named matrix of scores for the newdata, expressed in the same coordinate system as self\$scores

Method setVisualization(): Compute PCA scores in the latent space and update corresponding fields.

Usage:
PLNPCAfit\$setVisualization(scale.unit = FALSE)

Arguments:

scale.unit Logical. Should PCA scores be rescaled to have unit variance

Method postTreatment(): Update R2, fisher, std_err fields and set up visualization after optimization

```
Usage:
PLNPCAfit$postTreatment(
  responses,
  covariates,
  offsets,
  weights,
  control,
  nullModel
)
```

Arguments:

responses the matrix of responses (called Y in the model). Will usually be extracted from the corresponding field in PLNfamily

- covariates design matrix (called X in the model). Will usually be extracted from the corresponding field in PLNfamily
- offsets offset matrix (called O in the model). Will usually be extracted from the corresponding field in PLNfamily

weights an optional vector of observation weights to be used in the fitting process.

- control a list for controlling the optimization. See details.
- nullModel null model used for approximate R2 computations. Defaults to a GLM model with same design matrix but not latent variable.

Method plot_individual_map(): Plot the factorial map of the PCA

```
Usage:
PLNPCAfit$plot_individual_map(
  axes = 1:min(2, self$rank),
  main = "Individual Factor Map",
  plot = TRUE,
  cols = "default"
)
```

Arguments:

- axes numeric, the axes to use for the plot when map = "individual" or "variable". Default it c(1,min(rank))
- main character. A title for the single plot (individual or variable factor map). If NULL (the default), an hopefully appropriate title will be used.
- plot logical. Should the plot be displayed or sent back as ggplot object
- cols a character, factor or numeric to define the color associated with the individuals. By default, all individuals receive the default color of the current palette.

```
Returns: a ggplot graphic
```

Method plot_correlation_circle(): Plot the correlation circle of a specified axis for a PLNLDAfit object

Usage:

```
PLNPCAfit$plot_correlation_circle(
   axes = 1:min(2, self$rank),
   main = "Variable Factor Map",
   cols = "default",
   plot = TRUE
)
```

Arguments:

- axes numeric, the axes to use for the plot when map = "individual" or "variable". Default it c(1,min(rank))
- main character. A title for the single plot (individual or variable factor map). If NULL (the default), an hopefully appropriate title will be used.
- cols a character, factor or numeric to define the color associated with the variables. By default, all variables receive the default color of the current palette.

plot logical. Should the plot be displayed or sent back as ggplot object

Returns: a ggplot graphic

Method plot_PCA(): Plot a summary of the PLNPCAfit object

```
Usage:
PLNPCAfit$plot_PCA(
    nb_axes = min(3, self$rank),
    ind_cols = "ind_cols",
    var_cols = "var_cols",
    plot = TRUE
)
```

Arguments:

- nb_axes scalar: the number of axes to be considered when map = "both". The default is min(3,rank).
- ind_cols a character, factor or numeric to define the color associated with the individuals. By default, all variables receive the default color of the current palette.
- var_cols a character, factor or numeric to define the color associated with the variables. By default, all variables receive the default color of the current palette.

plot logical. Should the plot be displayed or sent back as ggplot object

Returns: a grob object

Method show(): User friendly print method

Usage:
PLNPCAfit\$show()

Method clone(): The objects of this class are cloneable with this method.

Usage: PLNPCAfit\$clone(deep = FALSE) Arguments: deep Whether to make a deep clone.

See Also

The function PLNPCA, the class PLNPCAfamily

Examples

```
data(trichoptera)
trichoptera <- prepare_data(trichoptera$Abundance, trichoptera$Covariate)
myPCAs <- PLNPCA(Abundance ~ 1 + offset(log(Offset)), data = trichoptera, ranks = 1:5)
myPCA <- getBestModel(myPCAs)
class(myPCA)
print(myPCA)</pre>
```

PLNPCA_param Control of PLNPCA fit

Description

Helper to define list of parameters to control the PLNPCA fit. All arguments have defaults.

Usage

```
PLNPCA_param(
   backend = "nlopt",
   trace = 1,
   config_post = list(),
   config_optim = list(),
   inception = NULL
)
```

PLNPCA_param

Arguments

backend	optimization back used, either "nlopt" or "torch". Default is "nlopt"
trace	a integer for verbosity.
config_post	a list for controlling the post-treatments (optional bootstrap, jackknife, R2, etc.). See details
config_optim	a list for controlling the optimizer (either "nlopt" or "torch" backend). See details
inception	Set up the parameters initialization: by default, the model is initialized with a multivariate linear model applied on log-transformed data, and with the same formula as the one provided by the user. However, the user can provide a PLNfit (typically obtained from a previous fit), which sometimes speeds up the inference.

Details

The list of parameters config_optim controls the optimizers. When "nlopt" is chosen the following entries are relevant

- "algorithm" the optimization method used by NLOPT among LD type, e.g. "CCSAQ", "MMA", "LBFGS". See NLOPT documentation for further details. Default is "CCSAQ".
- "maxeval" stop when the number of iteration exceeds maxeval. Default is 10000
- "ftol_rel" stop when an optimization step changes the objective function by less than ftol multiplied by the absolute value of the parameter. Default is 1e-8
- "xtol_rel" stop when an optimization step changes every parameters by less than xtol multiplied by the absolute value of the parameter. Default is 1e-6
- "ftol_abs" stop when an optimization step changes the objective function by less than ftol_abs. Default is 0.0 (disabled)
- "xtol_abs" stop when an optimization step changes every parameters by less than xtol_abs. Default is 0.0 (disabled)
- "maxtime" stop when the optimization time (in seconds) exceeds maxtime. Default is -1 (disabled)

When "torch" backend is used, with the following entries are relevant:

- "maxeval" stop when the number of iteration exceeds maxeval. Default is 10000
- "ftol_rel" stop when an optimization step changes the objective function by less than ftol multiplied by the absolute value of the parameter. Default is 1e-8
- "xtol_rel" stop when an optimization step changes every parameters by less than xtol multiplied by the absolute value of the parameter. Default is 1e-6

The list of parameters config_post controls the post-treatment processing, with the following entries:

• jackknife boolean indicating whether jackknife should be performed to evaluate bias and variance of the model parameters. Default is FALSE.

- bootstrap integer indicating the number of bootstrap resamples generated to evaluate the variance of the model parameters. Default is 0 (inactivated).
- variational_var boolean indicating whether variational Fisher information matrix should be computed to estimate the variance of the model parameters (highly underestimated). Default is FALSE.
- rsquared boolean indicating whether approximation of R2 based on deviance should be computed. Default is TRUE

Value

list of parameters configuring the fit.

PLN_param Control of a PLN fit

Description

Helper to define list of parameters to control the PLN fit. All arguments have defaults.

Usage

```
PLN_param(
 backend = c("nlopt", "torch"),
 trace = 1,
 covariance = c("full", "diagonal", "spherical", "fixed"),
 Omega = NULL,
 config_post = list(),
 config_optim = list(),
 inception = NULL
)
```

Arguments

backend	optimization back used, either "nlopt" or "torch". Default is "nlopt"
trace	a integer for verbosity.
covariance	character setting the model for the covariance matrix. Either "full", "diagonal", "spherical" or "fixed". Default is "full".
Omega	precision matrix of the latent variables. Inverse of Sigma. Must be specified if covariance is "fixed"
config_post	a list for controlling the post-treatments (optional bootstrap, jackknife, R2, etc.). See details
config_optim	a list for controlling the optimizer (either "nlopt" or "torch" backend). See details

inception Set up the parameters initialization: by default, the model is initialized with a multivariate linear model applied on log-transformed data, and with the same formula as the one provided by the user. However, the user can provide a PLNfit (typically obtained from a previous fit), which sometimes speeds up the inference.

Details

The list of parameters config_optim controls the optimizers. When "nlopt" is chosen the following entries are relevant

- "algorithm" the optimization method used by NLOPT among LD type, e.g. "CCSAQ", "MMA", "LBFGS". See NLOPT documentation for further details. Default is "CCSAQ".
- "maxeval" stop when the number of iteration exceeds maxeval. Default is 10000
- "ftol_rel" stop when an optimization step changes the objective function by less than ftol multiplied by the absolute value of the parameter. Default is 1e-8
- "xtol_rel" stop when an optimization step changes every parameters by less than xtol multiplied by the absolute value of the parameter. Default is 1e-6
- "ftol_abs" stop when an optimization step changes the objective function by less than ftol_abs. Default is 0.0 (disabled)
- "xtol_abs" stop when an optimization step changes every parameters by less than xtol_abs. Default is 0.0 (disabled)
- "maxtime" stop when the optimization time (in seconds) exceeds maxtime. Default is -1 (disabled)

When "torch" backend is used, with the following entries are relevant:

- "maxeval" stop when the number of iteration exceeds maxeval. Default is 10000
- "ftol_rel" stop when an optimization step changes the objective function by less than ftol multiplied by the absolute value of the parameter. Default is 1e-8
- "xtol_rel" stop when an optimization step changes every parameters by less than xtol multiplied by the absolute value of the parameter. Default is 1e-6

The list of parameters config_post controls the post-treatment processing, with the following entries:

- jackknife boolean indicating whether jackknife should be performed to evaluate bias and variance of the model parameters. Default is FALSE.
- bootstrap integer indicating the number of bootstrap resamples generated to evaluate the variance of the model parameters. Default is 0 (inactivated).
- variational_var boolean indicating whether variational Fisher information matrix should be computed to estimate the variance of the model parameters (highly underestimated). Default is FALSE.
- rsquared boolean indicating whether approximation of R2 based on deviance should be computed. Default is TRUE

Value

list of parameters configuring the fit.

plot.PLNfamily

Description

Display the criteria associated with a collection of PLN fits (a PLNfamily)

Usage

```
## S3 method for class 'PLNfamily'
plot(x, criteria = c("loglik", "BIC", "ICL"), reverse = FALSE, ...)
```

Arguments

Х	an R6 object with class PLNfamily
criteria	vector of characters. The criteria to plot in c("loglik", "BIC", "ICL"). Default is c("loglik", "BIC", "ICL").
reverse	A logical indicating whether to plot the value of the criteria in the "natural" direction (loglik - 0.5 penalty) or in the "reverse" direction (-2 loglik + penalty). Default to FALSE, i.e use the natural direction, on the same scale as the log-likelihood.
	additional parameters for S3 compatibility. Not used

Details

The BIC and ICL criteria have the form 'loglik - 1/2 * penalty' so that they are on the same scale as the model log-likelihood. You can change this direction and use the alternate form '-2*loglik + penalty', as some authors do, by setting reverse = TRUE.

Value

Produces a plot representing the evolution of the criteria of the different models considered, highlighting the best model in terms of BIC and ICL (see details).

See Also

plot.PLNPCAfamily() and plot.PLNnetworkfamily()

Examples

```
data(trichoptera)
trichoptera <- prepare_data(trichoptera$Abundance, trichoptera$Covariate)
myPCAs <- PLNPCA(Abundance ~ 1 + offset(log(Offset)), data = trichoptera, ranks = 1:5)
## Not run:
plot(myPCAs)</pre>
```

End(Not run)
plot.PLNLDAfit

Description

LDA visualization (individual and/or variable factor map(s)) for a PLNPCAfit object

Usage

```
## S3 method for class 'PLNLDAfit'
plot(
    x,
    map = c("both", "individual", "variable"),
    nb_axes = min(3, x$rank),
    axes = seq.int(min(2, x$rank)),
    var_cols = "var_colors",
    plot = TRUE,
    main = NULL,
    ...
)
```

Arguments

х	an R6 object with class PLNPCAfit
map	the type of output for the PCA visualization: either "individual", "variable" or "both". Default is "both".
nb_axes	scalar: the number of axes to be considered when map = "both". The default is min(3,rank).
axes	numeric, the axes to use for the plot when map = "individual" or "variable". Default it c(1,min(rank))
var_cols	a character or factor to define the color associated with the variables. By default, all variables receive the default color of the current palette.
plot	logical. Should the plot be displayed or sent back as ggplot2 object
main	character. A title for the single plot (individual or variable factor map). If NULL (the default), an hopefully appropriate title will be used.
	Not used (S3 compatibility).

Value

displays an individual and/or variable factor maps for the corresponding axes, and/or sends back a ggplot2 or gtable object

Examples

```
data(trichoptera)
trichoptera <- prepare_data(trichoptera$Abundance, trichoptera$Covariate)
myPLNLDA <- PLNLDA(Abundance ~ 1, grouping = Group, data = trichoptera)
## Not run:
plot(myPLNLDA, map = "individual", nb_axes = 2)
## End(Not run)</pre>
```

plot.PLNmixturefamily Display the criteria associated with a collection of PLNmixture fits (a PLNmixturefamily)

Description

Display the criteria associated with a collection of PLNmixture fits (a PLNmixturefamily)

Usage

```
## S3 method for class 'PLNmixturefamily'
plot(
    x,
    type = c("criteria", "diagnostic"),
    criteria = c("loglik", "BIC", "ICL"),
    reverse = FALSE,
    ...
)
```

Arguments

х	an R6 object with class PLNmixturefamily
type	a character, either "criteria" or "diagnostic" for the type of plot.
criteria	vector of characters. The criteria to plot in c("loglik", "BIC", "ICL"). Default is c("loglik", "BIC", "ICL").
reverse	A logical indicating whether to plot the value of the criteria in the "natural" direction (loglik - 0.5 penalty) or in the "reverse" direction (-2 loglik + penalty). Default to FALSE, i.e use the natural direction, on the same scale as the log-likelihood.
	additional parameters for S3 compatibility. Not used

Details

The BIC and ICL criteria have the form 'loglik - 1/2 * penalty' so that they are on the same scale as the model log-likelihood. You can change this direction and use the alternate form '-2*loglik + penalty', as some authors do, by setting reverse = TRUE.

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plot.PLNmixturefit

Value

Produces either a diagnostic plot (with type = 'diagnostic') or the evolution of the criteria of the different models considered (with type = 'criteria', the default).

Examples

plot.PLNmixturefit *Mixture visualization of a* PLNmixturefit *object*

Description

Represent the result of the clustering either by coloring the individual in a two-dimension PCA factor map, or by representing the expected matrix of count reorder according to the clustering.

Usage

```
## S3 method for class 'PLNmixturefit'
plot(x, type = c("pca", "matrix"), main = NULL, plot = TRUE, ...)
```

Arguments

Х	an R6 object with class PLNmixturefit
type	character for the type of plot, either "pca", for or "matrix". Default is "pca".
main	character. A title for the plot. If NULL (the default), an hopefully appropriate title will be used.
plot	logical. Should the plot be displayed or sent back as ggplot object
	Not used (S3 compatibility).

Value

a ggplot graphic

Examples

plot.PLNnetworkfamily Display various outputs (goodness-of-fit criteria, robustness, diagnostic) associated with a collection of PLNnetwork fits (a PLNnetworkfamily)

Description

Display various outputs (goodness-of-fit criteria, robustness, diagnostic) associated with a collection of PLNnetwork fits (a PLNnetworkfamily)

Usage

```
## S3 method for class 'PLNnetworkfamily'
plot(
    x,
    type = c("criteria", "stability", "diagnostic"),
    criteria = c("loglik", "pen_loglik", "BIC", "EBIC"),
    reverse = FALSE,
    log.x = TRUE,
    stability = 0.9,
    ...
)
```

Arguments

х	an R6 object with class PLNnetworkfamily
type	a character, either "criteria", "stability" or "diagnostic" for the type of plot.
criteria	vector of characters. The criteria to plot in c("loglik", "BIC", "ICL", "R_squared", "EBIC", "pen_loglik"). Default is c("loglik", "pen_loglik", "BIC", "EBIC"). Only relevant when type = "criteria".
reverse	A logical indicating whether to plot the value of the criteria in the "natural" direction (loglik - 0.5 penalty) or in the "reverse" direction (-2 loglik + penalty). Default to FALSE, i.e use the natural direction, on the same scale as the log-likelihood.
log.x	logical: should the x-axis be represented in log-scale? Default is TRUE.
stability	scalar: the targeted level of stability in stability plot. Default is .9.
	additional parameters for S3 compatibility. Not used

Details

The BIC and ICL criteria have the form 'loglik - 1/2 * penalty' so that they are on the same scale as the model log-likelihood. You can change this direction and use the alternate form '-2*loglik + penalty', as some authors do, by setting reverse = TRUE.

Value

Produces either a diagnostic plot (with type = 'diagnostic'), a stability plot (with type = 'stability') or the evolution of the criteria of the different models considered (with type = 'criteria', the default).

Examples

```
data(trichoptera)
trichoptera <- prepare_data(trichoptera$Abundance, trichoptera$Covariate)
fits <- PLNnetwork(Abundance ~ 1, data = trichoptera)
## Not run:
plot(fits)
## End(Not run)</pre>
```

plot.PLNnetworkfit	Extract and plot the network (partial correlation, support or inverse
	<i>covariance</i>) <i>from a</i> PLNnetworkfit <i>object</i>

Description

Extract and plot the network (partial correlation, support or inverse covariance) from a PLNnetworkfit object

Usage

```
## S3 method for class 'PLNnetworkfit'
plot(
    x,
    type = c("partial_cor", "support"),
    output = c("igraph", "corrplot"),
    edge.color = c("#F8766D", "#00BFC4"),
    remove.isolated = FALSE,
    node.labels = NULL,
    layout = layout_in_circle,
    plot = TRUE,
    ...
)
```

Arguments

х	an R6 object with class PLNnetworkfit
type	character. Value of the weight of the edges in the network, either "partial_cor" (partial correlation) or "support" (binary). Default is "partial_cor".
output	the type of output used: either 'igraph' or 'corrplot'. Default is 'igraph'.
edge.color	Length 2 color vector. Color for positive/negative edges. Default is c("#F8766D", "#00BFC4"). Only relevant for igraph output.

remove.isolated	
	if TRUE, isolated node are remove before plotting. Only relevant for igraph out-
	put.
node.labels	vector of character. The labels of the nodes. The default will use the column names ot the response matrix.
layout	an optional igraph layout. Only relevant for igraph output.
plot	logical. Should the final network be displayed or only sent back to the user. Default is \ensuremath{TRUE} .
	Not used (S3 compatibility).

Value

Send back an invisible object (igraph or Matrix, depending on the output chosen) and optionally displays a graph (via igraph or corrplot for large ones)

Examples

```
data(trichoptera)
trichoptera <- prepare_data(trichoptera$Abundance, trichoptera$Covariate)
fits <- PLNnetwork(Abundance ~ 1, data = trichoptera)
myNet <- getBestModel(fits)
## Not run:
plot(myNet)</pre>
```

End(Not run)

plot.PLNPCAfamily	Display the criteria associated with a collection of PLNPCA fits (a
	PLNPCAfamily)

Description

Display the criteria associated with a collection of PLNPCA fits (a PLNPCAfamily)

Usage

```
## S3 method for class 'PLNPCAfamily'
plot(x, criteria = c("loglik", "BIC", "ICL"), reverse = FALSE, ...)
```

Arguments

х	an R6 object with class PLNPCAfamily
criteria	vector of characters. The criteria to plot in c("loglik", "BIC", "ICL"). Default is c("loglik", "BIC", "ICL").
reverse	A logical indicating whether to plot the value of the criteria in the "natural" direction (loglik - 0.5 penalty) or in the "reverse" direction (-2 loglik + penalty). Default to FALSE, i.e use the natural direction, on the same scale as the log-likelihood.
	additional parameters for S3 compatibility. Not used

Details

The BIC and ICL criteria have the form 'loglik - 1/2 * penalty' so that they are on the same scale as the model log-likelihood. You can change this direction and use the alternate form '-2*loglik + penalty', as some authors do, by setting reverse = TRUE.

Value

Produces a plot representing the evolution of the criteria of the different models considered, highlighting the best model in terms of BIC and ICL (see details).

Examples

```
data(trichoptera)
trichoptera <- prepare_data(trichoptera$Abundance, trichoptera$Covariate)
myPCAs <- PLNPCA(Abundance ~ 1 + offset(log(Offset)), data = trichoptera, ranks = 1:5)
## Not run:
plot(myPCAs)</pre>
```

End(Not run)

plot.PLNPCAfit	PCA visualization (individual and/or variable factor map(s)) for a
	PLNPCAfit object

Description

PCA visualization (individual and/or variable factor map(s)) for a PLNPCAfit object

Usage

```
## S3 method for class 'PLNPCAfit'
plot(
    x,
    map = c("both", "individual", "variable"),
    nb_axes = min(3, x$rank),
    axes = seq.int(min(2, x$rank)),
    ind_cols = "ind_colors",
    var_cols = "var_colors",
    plot = TRUE,
    main = NULL,
    ....
)
```

Arguments

х	an R6 object with class PLNPCAfit
map	the type of output for the PCA visualization: either "individual", "variable" or "both". Default is "both".
nb_axes	<pre>scalar: the number of axes to be considered when map = "both". The default is min(3,rank).</pre>
axes	<pre>numeric, the axes to use for the plot when map = "individual" or map = "variable". Default it c(1,min(rank))</pre>
ind_cols	a character, factor or numeric to define the color associated with the individuals. By default, all variables receive the default color of the current palette.
var_cols	a character, factor or numeric to define the color associated with the variables. By default, all variables receive the default color of the current palette.
plot	logical. Should the plot be displayed or sent back as ggplot object
main	character. A title for the single plot (individual or variable factor map). If NULL (the default), an hopefully appropriate title will be used.
	Not used (S3 compatibility).

Value

displays an individual and/or variable factor maps for the corresponding axes, and/or sends back a ggplot or gtable object

Examples

```
data(trichoptera)
trichoptera <- prepare_data(trichoptera$Abundance, trichoptera$Covariate)
myPCAs <- PLNPCA(Abundance ~ 1 + offset(log(Offset)), data = trichoptera, ranks = 1:5)
myPCA <- getBestModel(myPCAs)
## Not run:
plot(myPCA, map = "individual", nb_axes=2, ind_cols = trichoptera$Group)
plot(myPCA, map = "variable", nb_axes=2)
plot(myPCA, map = "both", nb_axes=2, ind_cols = trichoptera$Group)
## End(Not run)</pre>
```

predict.PLNfit *Predict counts of a new sample*

Description

Predict counts of a new sample

Usage

```
## S3 method for class 'PLNfit'
predict(object, newdata, type = c("link", "response"), ...)
```

Arguments

object	an R6 object with class PLNfit
newdata	A data frame in which to look for variables and offsets with which to predict
type	The type of prediction required. The default is on the scale of the linear predic- tors (i.e. log average count)
	additional parameters for S3 compatibility. Not used

Value

A matrix of predicted log-counts (if type = "link") or predicted counts (if type = "response").

predict.PLNLDAfit *Predict group of new samples*

Description

Predict group of new samples

Usage

```
## S3 method for class 'PLNLDAfit'
predict(
   object,
   newdata,
   type = c("posterior", "response", "scores"),
   scale = c("log", "prob"),
   prior = NULL,
   control = PLN_param(backend = "nlopt"),
   ...
)
```

Arguments

object	an R6 object with class PLNLDAfit
newdata	A data frame in which to look for variables, offsets and counts with which to predict.
type	The type of prediction required. The default are posterior probabilities for each group (in either unnormalized log-scale or natural probabilities, see "scale" for details), "response" is the group with maximal posterior probability and "scores" is the average score along each separation axis in the latent space, with weights equal to the posterior probabilities.
scale	The scale used for the posterior probability. Either log-scale ("log", default) or natural probabilities summing up to 1 ("prob").
prior	User-specified prior group probabilities in the new data. If NULL (default), prior probabilities are computed from the learning set.
control	a list for controlling the optimization. See PLN() for details.
	additional parameters for S3 compatibility. Not used

Value

A matrix of posterior probabilities for each group (if type = "posterior"), a matrix of (average) scores in the latent space (if type = "scores") or a vector of predicted groups (if type = "response").

Examples

predict.PLNmixturefit Prediction for a PLNmixturefit object

Description

Predict either posterior probabilities for each group or latent positions based on new samples

Usage

```
## S3 method for class 'PLNmixturefit'
predict(
    object,
    newdata,
    type = c("posterior", "response", "position"),
    prior = matrix(rep(1/object$k, object$k), nrow(newdata), object$k, byrow = TRUE),
    control = PLNmixture_param(),
    ...
)
```

Arguments

object	an R6 object with class PLNmixturefit
newdata	A data frame in which to look for variables, offsets and counts with which to predict.
type	The type of prediction required. The default posterior are posterior probabil- ities for each group, response is the group with maximal posterior probability and latent is the averaged latent in the latent space, with weights equal to the posterior probabilities.

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predict_cond

prior	User-specified prior group probabilities in the new data. The default uses a uniform prior.
control	a list-like structure for controlling the fit. See ${\tt PLNmixture_param}$ () for details.
	additional parameters for S3 compatibility. Not used

Value

A matrix of posterior probabilities for each group (if type = "posterior"), a matrix of (average) position in the latent space (if type = "position") or a vector of predicted groups (if type = "response").

Examples

predict_cond Predict counts conditionally

Description

Predict counts of a new sample conditionally on a (set of) observed variables

Usage

```
predict_cond(
    object,
    newdata,
    cond_responses,
    type = c("link", "response"),
    var_par = FALSE
)
## S3 method for class 'PLNfit'
predict_cond(
    object,
    newdata,
    cond_responses,
    type = c("link", "response"),
    var_par = FALSE
)
```

Arguments

object	an R6 object with class PLNfit
newdata	A data frame in which to look for variables and offsets with which to predict
cond_responses	a data frame containing the counts of the observed variables (matching the names provided as data in the PLN function) $% \left(\frac{1}{2}\right) =0$
type	The type of prediction required. The default is on the scale of the linear predictors (i.e. log average count)
var_par	Boolean. Should new estimations of the variational parameters of mean and variance be sent back, as attributes of the matrix of predictions. Default to FALSE.

Value

A list containing:

pred	A matrix of predicted log-counts (if type = "link") or predicted counts (if type = "response")
М	A matrix containing E(Z_uncond Y_c) for each given site.
S	A matrix containing Var(Z_uncond Y_c) for each given site (sites are the third dimension of the array)

Methods (by class)

• predict_cond(PLNfit): Predict counts of a new sample conditionally on a (set of) observed variables for a PLNfit

Examples

```
data(trichoptera)
trichoptera_prep <- prepare_data(trichoptera$Abundance, trichoptera$Covariate)
myPLN <- PLN(Abundance ~ Temperature + Wind, trichoptera_prep)
#Condition on the set of the first two species in the dataset (Hym, Hys) at the ten first sites
Yc <- trichoptera$Abundance[1:10, c(1, 2), drop=FALSE]
newX <- cbind(1, trichoptera$Covariate[1:10, c("Temperature", "Wind")])
pred <- predict_cond(myPLN, newX, Yc, type = "response")</pre>
```

prepare_data

Prepare data for use in PLN models

Description

Prepare data in proper format for use in PLN model and its variants. The function (i) merges a count table and a covariate data frame in the most comprehensive way and (ii) computes offsets from the count table using one of several normalization schemes (TSS, CSS, RLE, GMPR, Wrench, etc). The function fails with informative messages when the heuristics used for sample matching fail.

prepare_data

Usage

prepare_data(counts, covariates, offset = "TSS", ...)

Arguments

counts	Required. An abundance count table, preferably with dimensions names and species as columns.
covariates	Required. A covariates data frame, preferably with row names.
offset	Optional. Normalization scheme used to compute scaling factors used as offset during PLN inference. Available schemes are "TSS" (Total Sum Scaling, de- fault), "CSS" (Cumulative Sum Scaling, used in metagenomeSeq), "RLE" (Rel- ative Log Expression, used in DESeq2), "GMPR" (Geometric Mean of Pairwise Ratio, introduced in Chen et al., 2018), Wrench (introduced in Kumar et al., 2018) or "none". Alternatively the user can supply its own vector or matrix of offsets (see note for specification of the user-supplied offsets).
	Additional parameters passed on to compute_offset()

Value

A data.frame suited for use in PLN() and its variants with two specials components: an abundance count matrix (in component "Abundance") and an offset vector/matrix (in component "Offset", only if offset is not set to "none")

Note

User supplied offsets should be either vectors/column-matrices or have the same number of column as the original count matrix and either (i) dimension names or (ii) the same dimensions as the count matrix. Samples are trimmed in exactly the same way to remove empty samples.

References

Chen, L., Reeve, J., Zhang, L., Huang, S., Wang, X. and Chen, J. (2018) GMPR: A robust normalization method for zero-inflated count data with application to microbiome sequencing data. PeerJ, 6, e4600 doi:10.7717/peerj.4600

Paulson, J. N., Colin Stine, O., Bravo, H. C. and Pop, M. (2013) Differential abundance analysis for microbial marker-gene surveys. Nature Methods, 10, 1200-1202 doi:10.1038/nmeth.2658

Anders, S. and Huber, W. (2010) Differential expression analysis for sequence count data. Genome Biology, 11, R106 doi:10.1186/gb20101110r106

Kumar, M., Slud, E., Okrah, K. et al. (2018) Analysis and correction of compositional bias in sparse sequencing count data. BMC Genomics 19, 799 doi:10.1186/s1286401851605

See Also

compute_offset() for details on the different normalization schemes

Examples

```
data(trichoptera)
proper_data <- prepare_data(
  counts = trichoptera$Abundance,
  covariates = trichoptera$Covariate,
  offset = "TSS"
)
proper_data$Abundance
proper_data$Offset</pre>
```

rPLN

PLN RNG

Description

Random generation for the PLN model with latent mean equal to mu, latent covariance matrix equal to Sigma and average depths (sum of counts in a sample) equal to depths

Usage

```
rPLN(
    n = 10,
    mu = rep(0, ncol(Sigma)),
    Sigma = diag(1, 5, 5),
    depths = rep(10000, n)
)
```

Arguments

n	the sample size
mu	vectors of means of the latent variable
Sigma	covariance matrix of the latent variable
depths	Numeric vector of target depths. The first is recycled if there are not n values

Details

The default value for mu and Sigma assume equal abundances and no correlation between the different species.

Value

a n * p count matrix, with row-sums close to depths

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sigma.PLNfit

Examples

```
## 10 samples of 5 species with equal abundances, no covariance and target depths of 10,000
rPLN()
## 2 samples of 10 highly correlated species with target depths 1,000 and 100,000
## very different abundances
mu <- rep(c(1, -1), each = 5)
Sigma <- matrix(0.8, 10, 10); diag(Sigma) <- 1
rPLN(n=2, mu = mu, Sigma = Sigma, depths = c(1e3, 1e5))</pre>
```

sigma.PLNfit Extract variance-covariance of residuals 'Sigma'

Description

Extract the variance-covariance matrix of the residuals, usually noted

Σ

in PLN models. This captures the correlation between the species in the latent space.

Usage

S3 method for class 'PLNfit'
sigma(object, ...)

Arguments

object	an R6 object with class PLNfit
	additional parameters for S3 compatibility. Not used

Value

A semi definite positive matrix of size p, assuming there are p species in the model.

See Also

coef.PLNfit(), standard_error.PLNfit() and vcov.PLNfit() for other ways to access

Examples

.

```
data(trichoptera)
trichoptera <- prepare_data(trichoptera$Abundance, trichoptera$Covariate)
myPLN <- PLN(Abundance ~ 1 + offset(log(Offset)), data = trichoptera)
sigma(myPLN) ## Sigma</pre>
```

sigma.PLNmixturefit Extract variance-covariance of residuals 'Sigma'

Description

Extract the variance-covariance matrix of the residuals, usually noted

Σ

in PLN models. This captures the correlation between the species in the latent space. or PLNmixture, it is a weighted mean of the variance-covariance matrices of each component.

Usage

```
## S3 method for class 'PLNmixturefit'
sigma(object, ...)
```

Arguments

object	an R6 object with class PLNmixturefit
	additional parameters for S3 compatibility. Not used

Value

A semi definite positive matrix of size p, assuming there are p species in the model.

See Also

coef.PLNmixturefit() for other ways to access

Σ

Examples

•

stability_selection Compute the stability path by stability selection

Description

This function computes the StARS stability criteria over a path of penalties. If a path has already been computed, the functions stops with a message unless force = TRUE has been specified.

Usage

```
stability_selection(
   Robject,
   subsamples = NULL,
   control = PLNnetwork_param(),
   force = FALSE
)
```

Arguments

subsamples a list of vectors describing the subsamples. The number of vectors (or list len determines th number of subsamples used in the stability selection. Autom cally set to 20 subsamples with size 10*sqrt(n) if n >= 144 and 0.8*n of)
wise following Liu et al. (2010) recommendations.	igth) nati- ther-
control a list controlling the main optimization process in each call to PLNnetwork. PLNnetwork() for details.	See
force force computation of the stability path, even if a previous one has been detected	cted.

Value

the list of subsamples. The estimated probabilities of selection of the edges are stored in the fields stability_path of the initial Robject with class PLNnetworkfamily

Examples

```
data(trichoptera)
trichoptera <- prepare_data(trichoptera$Abundance, trichoptera$Covariate)
fits <- PLNnetwork(Abundance ~ 1, data = trichoptera)
## Not run:
n <- nrow(trichoptera)
subs <- replicate(10, sample.int(n, size = n/2), simplify = FALSE)
stability_selection(nets, subsamples = subs)</pre>
```

End(Not run)

```
standard_error.PLNPCAfit
```

Component-wise standard errors of B

Description

Extracts univariate standard errors for the estimated coefficient of B. Standard errors are computed from the (approximate) Fisher information matrix.

Usage

```
## S3 method for class 'PLNPCAfit'
standard_error(
  object,
  type = c("variational", "jackknife", "sandwich"),
  parameter = c("B", "Omega")
)
standard_error(
  object,
  type = c("variational", "jackknife", "sandwich"),
  parameter = c("B", "Omega")
)
## S3 method for class 'PLNfit'
standard_error(
 object,
  type = c("variational", "jackknife", "bootstrap", "sandwich"),
  parameter = c("B", "Omega")
)
## S3 method for class 'PLNfit_fixedcov'
standard_error(
 object,
  type = c("variational", "jackknife", "bootstrap", "sandwich"),
  parameter = c("B", "Omega")
)
```

Arguments

object	an R6 object with class PLNfit
type	string describing the type of variance approximation: "variational", "jackknife", "sandwich" (only for fixed covariance). Default is "variational".
parameter	string describing the target parameter: either B (regression coefficients) or Omega (inverse residual covariance)

trichoptera

Value

A p * d positive matrix (same size as B) with standard errors for the coefficients of B

Methods (by class)

- standard_error(PLNPCAfit): Component-wise standard errors of B in PLNPCAfit (not implemented yet)
- standard_error(PLNfit): Component-wise standard errors of B in PLNfit
- standard_error(PLNfit_fixedcov): Component-wise standard errors of B in PLNfit_fixedcov

See Also

vcov.PLNfit() for the complete variance covariance estimation of the coefficient

Examples

trichoptera

Trichoptera data set

Description

Data gathered between 1959 and 1960 during 49 insect trapping nights. For each trapping night, the abundance of 17 Trichoptera species is recorded as well as 6 meteorological variables which may influence the abundance of each species. Finally, the observations (that is to say, the trapping nights), have been classified into 12 groups corresponding to contiguous nights between summer 1959 and summer 1960.

Usage

trichoptera

Format

A list with 2 two data frames:

Abundancea 49 x 17 matrix of abundancies/counts (49 trapping nights and 17 trichoptera species)

Covariate a 49 x 7 data frame of covariates:

Temperature Evening Temperature in Celsius Wind Wind in m/s Pressure Pressure in mm Hg Humidity relative to evening humidity in percentCloudiness proportion of sky coverage at 9pmPrecipitation Nighttime precipitation in mmGroup a factor of 12 levels for the definition of the consecutive night groups

In order to prepare the data for using formula in multivariate analysis (multiple outputs and inputs), use prepare_data(). We only kept a subset of the original meteorological covariates for illustration purposes.

Source

Data from P. Usseglio-Polatera.

References

Usseglio-Polatera, P. and Auda, Y. (1987) Influence des facteurs météorologiques sur les résultats de piégeage lumineux. Annales de Limnologie, 23, 65–79. (code des espèces p. 76) See a data description at http://pbil.univ-lyon1.fr/R/pdf/pps034.pdf (in French)

See Also

prepare_data()

Examples

```
data(trichoptera)
trichoptera <- prepare_data(trichoptera$Abundance, trichoptera$Covariate)</pre>
```

vcov.PLNfit

Calculate Variance-Covariance Matrix for a fitted PLN() model object

Description

Returns the variance-covariance matrix of the main parameters of a fitted PLN() model object. The main parameters of the model correspond to

B

, as returned by coef.PLNfit(). The function can also be used to return the variance-covariance matrix of the residuals. The latter matrix can also be accessed via sigma.PLNfit()

Usage

```
## S3 method for class 'PLNfit'
vcov(object, type = c("main", "covariance"), ...)
```

vcov.PLNfit

Arguments

object	an R6 object with class PLNfit	
type	type of parameter that should be extracted. Either "main" (default)	for
	В	
	or "covariance" for	
	Σ	
	additional parameters for S3 compatibility. Not used	

Value

A matrix of variance/covariance extracted from the PLNfit model. If type="main" and B is a matrix of size d * p, the result is a block-diagonal matrix with p (number of species) blocks of size d (number of covariates). if type="main", it is a symmetric matrix of size p. .

See Also

sigma.PLNfit(), coef.PLNfit(), standard_error.PLNfit()

Examples

```
data(trichoptera)
trichoptera <- prepare_data(trichoptera$Abundance, trichoptera$Covariate)
myPLN <- PLN(Abundance ~ 1 + offset(log(Offset)), data = trichoptera)
vcov(myPLN) ## variance-covariance of B
vcov(myPLN, type = "covariance") ## Sigma</pre>
```

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