

Package ‘ALDEx3’

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Title Linear Models for Sequence Count Data

Version 1.2.0

Description Provides scalable generalized linear and mixed effects models tailored for sequence count data analysis (e.g., analysis of 16S or RNA-seq data). Uses Dirichlet-multinomial sampling to quantify uncertainty in relative abundance or relative expression conditioned on observed count data.
Implements scale models as a generalization of normalizations which account for uncertainty in scale (e.g., total abundances) as described in Nixon et al. (2025) <[doi:10.1186/s13059-025-03609-3](https://doi.org/10.1186/s13059-025-03609-3)> and McGovern et al. (2025) <[doi:10.1101/2025.08.05.668734](https://doi.org/10.1101/2025.08.05.668734)>.

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aldex	<i>ALDEx3 Linear Models</i>
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Description

ALDEx3 Linear Models

Usage

```
aldex(
  Y,
  X,
  data = NULL,
  method = "lm",
  nsample = 2000,
  scale = NULL,
  streamsize = 8000,
  n.cores = detectCores() - 1,
  return.pars = c("X", "estimate", "std.error", "p.val", "p.val.adj", "logComp",
    "logScale"),
  p.adjust.method = "BH",
  test = "t.HC3",
```

```

onesided = FALSE,
...
)

```

Arguments

Y	an (D x N) matrix of sequence count, N is number of samples, D is number of taxa or genes
X	either a formula (in which case DATA must be non-null) or a model matrix of dimension P x N (P is number of linear model covariates). If a formula is passed it should not include the target Y, e.g., should simply be "~condition-1" (note the lack of the left hand side). If using lme4, this should be the formula including random effects.
data	a data frame for use with formula, must have N rows
method	(default lm) The regression method; "lm": linear regression, "lme4": linear mixed effects regression with lme4; "nlme": linear mixed effects models with nlme, REQUIRES "random" argument representing random effects be passed into aldex function, can also pass "correlation" as argument (see nlme documentation for how to use "correlation" argument); or "blmm": an ALDEx3-specific approximate mixed-effects engine that uses one batched profiled mixed-model anchor fit per feature, draw-specific local covariance updates, and exact conditional fixed-effect solves. The approximation is only in the variance-component optimisation step. See the mixed-effects vignette for details, validation guidance, and the runtime comparison with exact lme4.
nsample	number of monte carlo replicates
scale	the scale model, can be a function or an N x nsample matrix (samples should be given on log2-scale; e.g., samples should be of log of system scale). The API for writing your own scale models is documented below in examples.
streamsize	(default 8000) memory footprint (approximate) at which to use streaming. This should be thought of as the number of Mb for each streaming chunk. If $DN/nsample * 8/1000000$ is less than streamsize then no streaming will be performed. Note, to conserve memory, samples from the Dirichlet and scale models will not be returned if streaming is used. Streaming can be turned off by setting streamsize=Inf.
n.cores	(default detectCores()-1) If method is 'lme4' or 'blmm', use this many cores for feature-level parallelism across taxa/features. For method = "blmm", parallelism is across features, not across Monte Carlo draws within a feature.
return.pars	what results should be returned, see return section below.
p.adjust.method	(default BH) The method for multiple hypothesis test correction. See p.adjust for all available methods.
test	(default t.HC3), "t", t test is performed for each covariate (fast); "t.HC0" Heteroskedasticity-Robust Standard Errors used (HC0; White's; slower); "t.HC3" (default) Heteroskedasticity-Robust Standard Errors used (HC3; unlike HC0, this includes a leverage adjustment and is better for small sample sizes or when there are data with high leverage; slowest). To learn more about these, look at Long and Ervin (2000) Using Heteroscedasticity Consistent Standard Errors in the Linear Regression Model, The American Statistician.

onesided (default: FALSE) if sided return p-values for two-sided test. Otherwise if "lower" or "upper" return one-sided test corresponding to test that estimate is negative or positive respectively.

... parameters to be passed to the scale model (if a function is provided), may also be random or correlation arguments for nlme.

Value

a list with elements controlled by parameter `reeturn.pars`. Options include: - `X`: $P \times N$ covariate matrix - `estimate`: $(P \times D \times \text{nsample})$ array of linear model estimates - `std.error`: $(P \times D \times \text{nsample})$ array of standard error for the estimates - `p.val`: $(P \times D)$ matrix, unadjusted p-value for two-sided t-test - `p.val.adj`: $(P \times D)$ matrix, p-value for two-sided t-test adjusted according to `p.adj.method` - `logScale`: $(N \times S)$ matrix, samples of the log scale from the scale model - `logComp`: $(D \times N \times S)$ array, samples of the log composition from the multinomial-Dirichlet - `streaming`: boolean, detnote if streaming was used. - `random.effects` ($Pr \times N \times S$): if using mixed effects models, return all Pr random-effects covariance terms and the residual variance. For `method = "blmm"`, correlated random-effect covariance terms are returned when present, matching the `lme4` naming convention as closely as possible. Note, `logScale` and `logComp` are not returned if streaming is active.

Author(s)

Justin Silverman, Kyle McGovern

Examples

```
Y <- matrix(1:110, 10, 11)
condition <- c(rep(0, 5), rep(1, 6))
data <- data.frame(condition=condition)
## demonstrate formula interface and passing optional argument (gamma) to
## the scale model (clr)
res <- aldex(Y, ~condition, data, nsample=2000, scale=clr.sm, gamma=0.5)

## demonstrating how to write a custom scale model, I will write a model
## that generalizes total sum scaling (where we assume no change between
## conditions)
## Functions can include parameters X (model matrix), Y, and logComp.
## If included in the function definition, those parameters will be passed
## dynamically when aldex is running. Other optional parameters (gamma)
## can be passed as additional arguments to the aldex function
tss <- function(X, logComp, gamma=0.5) {
  P <- nrow(X)
  nsample <- dim(logComp)[3]
  LambdaScale <- matrix(rnorm(P*nsample,0,gamma), P, nsample)
  logScale <- t(X)%*% LambdaScale
  return(logScale)
}
res <- aldex(Y, ~condition, data, nsample=2000, scale=tss, gamma=0.75)
```

aldex.effect *ALDEx3 Effect Diagnostics*

Description

ALDEx2-inspired effect diagnostics for an ALDEx3 binary contrast

Usage

```
aldex.effect(object, contrast)
```

Arguments

object	An object returned by aldex.
contrast	The exact model coefficient or unambiguous binary data column to summarize.

Details

aldex.effect is a helper for effect-size diagnostics that Greg Gloor and ALDEx2 users commonly inspect when evaluating pairwise effects. It is not a replacement for summary.aldex, which reports model estimates, standard errors and adjusted p-values. This function summarizes a single binary ALDEx3 model contrast using Cohen's-d-based Monte Carlo diagnostics.

For a selected contrast row k , feature d , Monte Carlo draw s , and binary group indicators $x = \text{object}\$X[k,]$, let $B[d, s]$ be the fitted ALDEx3 coefficient stored in $\text{object}\$estimate[k, d, s]$. Let $\log W[d, i, s] = \text{object}\$logComp[d, i, s] + \text{object}\$logScale[i, s]$ be the reconstructed log abundance for sample i . aldex.effect requires logComp and logScale, so it is unavailable when aldex streams large jobs and omits those arrays. For each feature and Monte Carlo draw, the pooled within-group variance is $((n_0 - 1) * \text{var}(\log W[d, x == 0, s]) + (n_1 - 1) * \text{var}(\log W[d, x == 1, s])) / (n_0 + n_1 - 2)$, where n_0 and n_1 are the group sizes.

The returned columns are calculated as follows:

- estimate: the mean of $B[d, s]$ over Monte Carlo draws.
- pooled.SD: the mean of the pooled within-group standard deviation, $\sqrt{\text{pooled variance}}$, over Monte Carlo draws.
- cohens.d: the mean of $B[d, s] / \text{pooled.SD}[d, s]$ over Monte Carlo draws. This is Cohen's-d-based and is not the same as the historical robust ALDEx2 effect statistic based on between/within dispersion.
- overlap: an ALDEx2-style directional uncertainty diagnostic. It counts the Monte Carlo Cohen's d draws below and above zero, smooths those two counts with a 0.5 pseudocount, converts them with the ALDEx/Aitchison mean, and returns the smaller smoothed sign probability. Values near 0 indicate that nearly all Monte Carlo effect draws have the same sign. Values near 0.5 indicate uncertainty about direction. overlap is not a p-value, a confidence interval probability, or a literal area of overlap between two density curves.

Value

A data.frame with columns parameter, entity, estimate, pooled.SD, cohens.d, and overlap.

Author(s)

Greg Gloor, Justin Silverman

aldex.lm.sim.clr	<i>Simulation function solely for testing and exploring ALDEx3, Truth is in CLR Coordinates.</i>
------------------	--

Description

Not designed to create realistic data. Does not add any noise to linear regression! True W is in CLR Coordinates

Usage

```
aldex.lm.sim.clr(D = 10, N = 11, P = 2, depth = 10000)
```

Arguments

D	number of taxa/genes
N	number of samples
P	number of covariates
depth	sum of counts for each multinomial draw

Value

a list with elements Y, X, W, and Lambda

Author(s)

Justin Silverman

aldex.mem.sim	<i>Simulation for testing mixed effects models.</i>
---------------	---

Description

Includes random intercept, option to include random slope, second random intercept, and time-correlation.

Usage

```
aldex.mem.sim(
  D,
  days,
  subjects,
  depth = 10000,
  location = 1,
  random_slope = FALSE,
  corr = 0,
  rho_ar1 = 0,
  sd_resid = 0.1
)
```

Arguments

D	number of taxa/genes
days	num days (i.e., repeated measurements) for each subject
subjects	num of subjects to simulate
depth	sum of counts for each multinomial draw
location	second random intercept, if 0 ignore, else simulate num of locations
random_slope	If true, simulate a random slope for each subject.
corr	The correlation between slope/random intercept
rho_ar1	The ar1 time-correlation, if 0, don't simulate
sd_resid	The residual error time-correlation.

Author(s)

Kyle McGovern

aldex.plot

Plot Method for ALDEx3 objects with pairwise comparisons

Description

Simple plots for an ALDEx3 object

Usage

```
aldex.plot(
  object,
  plot = c("volcano", "effect", "MA", "water"),
  contrast = NULL,
  threshold = 0.05,
  min.diff = 0.5,
```

```

    cohen = 0.5,
    sig.col = rgb(1, 0, 0, 0.5),
    water.show = 5,
    water.col = c("red", "blue"),
    water.names = TRUE,
    ...
  )

```

Arguments

<code>object</code>	An object of class <code>aldex</code>
<code>plot</code>	type of plot (default='volcano')
<code>contrast</code>	the name of the comparison to plot, must be provided
<code>threshold</code>	FDR significance threshold (default=0.05)
<code>min.diff</code>	(default=0.5) used for MA (display), and waterfall (cutoff) plots
<code>cohen</code>	Cohen's d threshold for effect plot (slope)
<code>sig.col</code>	color for significant features
<code>water.show</code>	number features to show in waterfall plot
<code>water.col</code>	vector of colors for waterfall plot
<code>water.names</code>	logical, show names of features for waterfall plot
<code>...</code>	additional plot parameters (cex)

Details

Provides volcano, effect, MA and waterfall plots from an ALDEx3 object.

This method plots combinations of adjusted p-values from `object$p.val.adj`, posterior estimates, standard errors and log abundance values averaged across Monte Carlo samples. The result is returned as a single plot of the desired type. Effect plots use the ALDEx2-inspired diagnostics returned by `aldex.effect`. The contrast must identify exactly one binary model coefficient. Effect diagnostics and MA plots require `logComp` and `logScale`, which are not returned when `aldex` streams large jobs.

Value

the desired plot

Author(s)

Greg Gloor

aldex.pvals	<i>Calculate p-values adjusting for changes in sign as described by Nixon et al. (2024) in Beyond Normalization: Incorporating Scale Uncertainty in Microbiome and Gene Expression Analysis (internal only)</i>
-------------	---

Description

Calculate p-values adjusting for changes in sign as described by Nixon et al. (2024) in Beyond Normalization: Incorporating Scale Uncertainty in Microbiome and Gene Expression Analysis (internal only)

Usage

```
aldex.pvals(p.lower, p.upper, p.adjust.method, onesided = FALSE)
```

Arguments

p.lower	A P x D x S matrix for P covariates, D taxa/genes, and S monte carlo samples representing the lower tail p.values
p.upper	A P x D x S matrix for P covariates, D taxa/genes, and S monte carlo samples representing the upper tail p.values
p.adjust.method	An adjutment method for p.adjust
onesided	(default: FALSE) if sided return p-values for two-sided test. Otherwise if "lower" or "upper" return one-sided test corresponding to test that estimate is negative or positive respectively.

Value

A list with P x D matrices with the non-adjusted and adjusted p-values.

Author(s)

Justin Silverman, Kyle McGovern

References

Nixon G, Gloor GB, Silverman JD (2025). "Incorporating scale uncertainty in microbiome and gene expression analysis as an extension of normalization". Genome Biology. doi:10.1186/s13059-025036093

blmm

*Fast approximate mixed-effects engine for ALDEx3.***Description**

Fast approximate mixed-effects engine for ALDEx3.

Usage

```
blmm(logW, formula, data, n.cores = 1L)
```

Arguments

logW	numeric array (N x D x S)
formula	lme4 mixed-effects formula
data	data.frame for formula evaluation
n.cores	number of worker processes for feature-level parallelism. If n.cores > 1, features are distributed across workers; the per-feature draw loop remains serial.

Details

Internal engine called by [aldex](#) when method = "blmm".

Uses an lme4-style profiled Gaussian LMM formulation. One batched anchor REML problem is fit per feature across all S MC draws; draw-specific local covariance updates are obtained via a single Newton step; exact conditional GLS fixed-effect solves are then run per draw. The approximation is only in the variance-component step — fixed effects and standard errors are exact conditional on the updated covariance parameters.

Features for which the approximate path fails are silently re-fit with the exact lme4 engine and a consolidated warning is issued naming the affected features and their error messages.

Value

named list of arrays (P x D x S): estimate, std.error, df, p.lower, p.upper, and (Pr x D x S) random.eff — same layout as `sr.mem`.

 clr.sm

Default CLR-based scale model (with optional scale uncertainty)

Description

Implements the default scale model described in Nixon et al. (Beyond Normalizations / scale-uncertainty framework). This model generalizes the centered log-ratio (CLR) normalization by treating the (log) scale as a latent random variable and allowing additive uncertainty around the CLR-implied scale differences via a Gaussian term with standard deviation γ .

Usage

```
clr.sm(X, logComp, gamma = 0.5)
```

Arguments

<code>X</code>	A numeric design matrix used to model scale variation across samples. This is the covariate/design matrix passed internally by <code>aldex()</code> to the scale model. Rows correspond to regression coefficients (e.g., intercept and covariates after contrasts/encoding) and columns correspond to samples. If the analysis includes only an intercept (no covariates), <code>X</code> is typically a 1 x N matrix of ones. (This parameter is automatically passed by <code>aldex</code>)
<code>logComp</code>	A numeric array of Monte Carlo log-compositions with dimensions features x samples x nsample. This is produced internally by ALDEx3 from Dirichlet-multinomial Monte Carlo sampling and log-ratio representation. <code>##'</code> (This parameter is automatically passed by <code>aldex</code>)
<code>gamma</code>	Non-negative scalar. Standard deviation of the Gaussian perturbation that relaxes the CLR assumption about scale. $\gamma = 0$ yields the pure CLR assumption; recommended default values in the scale-uncertainty literature are often around 0.5, but appropriate values depend on how strongly you trust the CLR scale assumption in the current study.

Details

In the limit $\gamma = 0$, this reduces to the CLR assumption (no scale uncertainty beyond the CLR-implied scale). Larger γ values represent increasing uncertainty about the CLR-implied scale differences.

Value

A numeric matrix of dimension $N \times \text{nsample}$ giving Monte Carlo samples of the log-scale for each sample (rows) and each Monte Carlo draw (columns).

Author(s)

Justin Silverman

References

Nixon G, Gloor GB, Silverman JD (2025). "Incorporating scale uncertainty in microbiome and gene expression analysis as an extension of normalization". *Genome Biology*. doi:10.1186/s13059-025036093

coefficient.sm	<i>Coefficient-based scale model with user-specified prior on fixed effects</i>
----------------	---

Description

Draws Monte Carlo samples of the log2 scale by sampling fixed-effect coefficients from a multivariate normal distribution and mapping them through the design matrix X . This scale model is useful when you want to encode prior information about how covariates (e.g., treatment, batch, time) affect scale, rather than specifying scale moments directly per sample.

Usage

```
coefficient.sm(X, logComp, c.mu = NULL, c.cor = NULL)
```

Arguments

X	A numeric design matrix passed internally by <code>aldex()</code> to the scale model. Rows correspond to fixed-effect coefficients/covariates ($P = \text{nrow}(X)$) and columns correspond to samples ($N = \text{ncol}(X)$). (Automatically supplied by <code>aldex()</code> .)
<code>logComp</code>	A numeric array of Monte Carlo log-compositions with dimensions <code>features x samples x nsample</code> . This scale model uses <code>nsample</code> (the number of Monte Carlo draws) but does not otherwise use <code>logComp</code> . (Automatically supplied by <code>aldex()</code> .)
<code>c.mu</code>	Numeric vector of length P giving the mean of the fixed effect coefficients in log2-scale space. Must not be <code>NULL</code> .
<code>c.cor</code>	Numeric $P \times P$ covariance matrix for the fixed effect coefficients in log2-scale space. Must not be <code>NULL</code> .

Details

Specifically, for each Monte Carlo draw $b^{(m)} \sim N(c.mu, c.cor)$, the per-sample log2 scale is computed as $b^{(m)T} X$, producing an $N \times \text{nsample}$ matrix of log2-scale draws.

For example, with an intercept and a treatment indicator where treatment is expected to increase log2 scale by ~ 1 on average, one might use `c.mu = c(0, 1)` and `c.cor = diag(c(0.25, 0.25))` (i.e., SD 0.5 for each coefficient, independent).

Value

A numeric matrix of dimension $N \times \text{nsample}$ giving Monte Carlo draws of the log2 scale for each sample (rows) across `nsample` draws (columns).

Author(s)

Kyle McGovern

cohensd*Cohen's D*

Description

Function to compute cohensd on the results provided by the aldex function

Usage

```
cohensd(m, var)
```

Arguments

m	the output of a call to aldex
var	if aldex was called with X being a pre-computed model matrix, this var should be an integer corresponding to a binary covariate indicating the desired effect size to calculate (an effect size between two groups indicated by the binary covariate). For example, if the third covariate in the model is an indicator denoting health (0) and disease (1) then set var=3. In contrast, if X was a formula (in which case the data argument should have been specified) then var can be set to the unquoted or quoted name of the binary condition variable (e.g., var=condition or var="condition").

Details

WARNING: this function is experimental and requires users read the documentation fully.

Value

A (D x nsample)-matrix of Cohen's d statistics for the variable of interest.

Author(s)

Justin Silverman

combine.streams	<i>Combine output from aldex streams (internal only)</i>
-----------------	--

Description

Takes a list l, where every element is itself a list of 3D arrays. Each array should have matching first and second dimintions. This function combines those arrays along the third dimension.

Usage

```
combine.streams(l)
```

Arguments

l	a list, where every element is itself a list of 3D arrays.
---	--

Value

a list of 3D arrays

Author(s)

Justin Silverman

fflm	<i>Freaking Fask Linear Models</i>
------	------------------------------------

Description

Tailored for ALDEx3 where covariates are shared between massive numbers of linear regressions where only Y is changing.

Usage

```
fflm(Y, X, test = "t.HC3")
```

Arguments

Y	a numeric array (N x D X S) where D is the number of taxa/genes, N is the number of samples, and S is the number of posterior samples
X	a numeric matrix (N x P) where P is number of covariates
test	(default t.HC3), "t", t test is performed for each covariate (fast); "t.HC0" Heteroskedasticity-Robust Standard Errors used (HC0; White's; slower); "t.HC3" (default) Heteroskedasticity-Robust Standard Errors used (HC3; unlike HC0, this includes a leverage adjustment and is better for small sample sizes or when there are data with high leverage; slowest). To learn more about these, loko at Long and Ervin (2000) Using Heteroscedasticity Consistent Standard Errors in the Linear Regression Model, The American Statistician.

Value

A list of (P x D x S)-arrays with the OLS point estimates, the standard errors, and the two-sided p-values for each coefficient (P), of each model fit to each taxa (D) and each posterior sample (S)

Author(s)

Justin Silverman

gut_crohns_data	<i>Gut Crohn's microbiome dataset (list)</i>
-----------------	--

Description

This dataset contains a matrix and a data frame: genus-level microbiome profiles and corresponding sample metadata from a Crohn's disease case-control cohort. The dataset is used in examples and vignettes throughout the package.

Usage

```
gut_crohns_data
```

Format

A named list of one matrix and a dataframe:

counts A matrix with read counts with 195 rows and 95 columns)

metadata A data frame with 95 rows and 7 columns containing subject-level covariates.

Details

The counts matrix has one row per sample and one column per genus. The metadata data frame has one row per sample with metadata, critically Health.status either CD or Control, and Average cell count per gram frozen feces.

Source

Vandeputte D, Falony G, Vieira-Silva S, Wang J, Sailer M, Theis S, Raes J (2017). "Quantitative microbiome profiling links gut community variation to microbial load". *Nature*, 551, 507–511.
[doi:10.1038/nature24460](https://doi.org/10.1038/nature24460)

miniclo	<i>Closure operation</i>
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Description

Closure operation

Usage

```
miniclo(X)
```

Arguments

`X` should be a D x N matrix, closes along the D dimension

Value

D x N matrix with columns summing to 1

Author(s)

Justin Silverman

oral_mouthwash_data	<i>Oral microbiome perturbation dataset (list)</i>
---------------------	--

Description

This dataset contains a matrix and a data frame: genus-level microbiome profiles and corresponding sample metadata from an oral microbiome perturbation study. 28 participants' oral microbiomes were measured before, 15 minutes after, and 2 hours after perturbation with either a water control, antiseptic mouthwash, alcohol-free mouthwash, or soda. The dataset is used in examples and vignettes throughout the package.

Usage

```
oral_mouthwash_data
```

Format

A named list of one matrix and a dataframe:

`counts` A matrix with read counts with 116 rows and 81 columns)

`metadata` A data frame with 81 rows and 38 columns containing sample-level covariates.

Details

The counts matrix has one row per sample and one column per genus. The metadata data frame has one row per sample with metadata, critically the participant ID, flow cytometry average (and replicate) cells, time_c (time points), and treat (the perturbations)

Source

Marotz C, Morton JT, Navarro P, Coker J, Belda-Ferre P, Knight R (2021). "Quantifying live microbial load in human saliva samples over time reveals stable composition and dynamic load". *mSystems*, 6(3), e01182-21. doi:[10.1128/mSystems.0118220](https://doi.org/10.1128/mSystems.0118220)

req	<i>Test object contains elements</i>
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Description

Test object contains elements

Usage

```
req(obj, names)
```

Arguments

obj	object (list type)
names	names of elements required to be in the list

Author(s)

Justin Silverman

rLogDirichlet	<i>Function for sampling Dirichlet random variables (base-2 normalized via log-sum-exp for stability)</i>
---------------	---

Description

Function for sampling Dirichlet random variables (base-2 normalized via log-sum-exp for stability)

Usage

```
rLogDirichlet(n, alpha)
```

Arguments

n	number of samples
alpha	D-vector of Dirichlet parameters

Value

a D x n matrix of samples

Author(s)

Justin Silverman

sample.sm	<i>Sample-specific scale model with user-specified mean and variance/covariance</i>
-----------	---

Description

Draws Monte Carlo samples of the log2 scale for each sample using user-supplied moments. This scale model is useful when external measurements (e.g., qPCR, flow cytometry, spike-ins) provide information about absolute scale, or when you want to encode prior information about scale on a per-sample basis.

Usage

```
sample.sm(X, logComp, s.mu = NULL, s.var = NULL, s.cor = NULL)
```

Arguments

X	A numeric design matrix passed internally by aldex() to the scale model. Columns correspond to samples ($N = \text{ncol}(X)$). This scale model does not use X directly, but N is inferred from it. (Automatically supplied by aldex().)
logComp	A numeric array of Monte Carlo log-compositions with dimensions features x samples x nsample. This scale model uses nsample to determine the number of Monte Carlo draws, but does not otherwise use logComp. (Automatically supplied by aldex().)
s.mu	Numeric vector of length N giving the mean of the log2 scale for each sample. Must not be NULL.
s.var	Numeric vector of length N giving the marginal variance of the log2 scale for each sample. Use this when assuming samples' log2 scales are independent. Must be NULL if s.cor is provided.
s.cor	Numeric N x N covariance matrix for the log2 scale across samples. Use this when encoding correlations between samples' log2 scales. Must be NULL if s.var is provided.

Details

Exactly one of `s.var` or `s.cor` must be provided:

- `s.var`: independent per-sample log2-scale variance (diagonal covariance)
- `s.cor`: full $N \times N$ log2-scale covariance matrix

The returned matrix has N rows (samples) and `nsample` columns (Monte Carlo draws), consistent with the ALDEx3 scale-model interface.

Value

A numeric matrix of dimension $N \times \text{nsample}$ giving Monte Carlo draws of the log2 scale for each sample (rows) across `nsample` draws (columns).

Author(s)

Kyle McGovern

`sr.mem`

Implementation of SR-MEM: scale-reliant mixed effects models.

Description

Implementation of SR-MEM: scale-reliant mixed effects models.

Usage

```
sr.mem(logW, formula, data, n.cores, method, mem.args)
```

Arguments

<code>logW</code>	a numeric array ($N \times D \times S$) where D is the number of taxa, N is the number of samples, and S is the number of posterior samples
<code>formula</code>	an lme4 formula with fixed and random effects for lmer
<code>data</code>	A data.frame with the random and fixed effects items in formula
<code>n.cores</code>	The number of cores for parallelization. If <code>n.cores=1</code> , no parallelization is used
<code>method</code>	The mem method to use: either <code>nlme</code> or <code>lme4</code>
<code>mem.args</code>	Additional arguments including random or correlation

Value

A list of $(P \times D \times S)$ -arrays with the fixed effect point estimates, the standard errors, degrees of freedom and the lower and upper p-values for each coefficient (P), of each model fit to each taxa (D) and each posterior sample (S) and a $(Pr \times D \times S)$ -array with (Pr) random effects.

Author(s)

Kyle McGovern

summary.aldex	<i>Summary Method for ALDEx3 Objects</i>
---------------	--

Description

Summarize an ALDEx3 result object

Usage

```
## S3 method for class 'aldex'
summary(object, ignore.intercept = TRUE, ...)
```

Arguments

object	An object of class aldex
ignore.intercept	(default=TRUE), ignore intercept when creating summary table
...	Additional arguments (currently ignored).

Details

Provides a summary of the adjusted p-values, estimates, and standard errors from an ALDEx3 result object.

This method extracts adjusted p-values from `object$p.val.adj`, along with posterior estimates and standard errors averaged across Monte Carlo samples. The result is returned as a long-format `data.frame` suitable for downstream analysis or visualization.

Value

A `data.frame` with columns `parameter`, `entity`, `p.val.adjusted`, `estimate`, and `std.error`.

Author(s)

Justin Silverman

tss.sm	<i>TSS-centered scale model (with optional scale uncertainty)</i>
--------	---

Description

Implements a total-sum-scaling (TSS)-centered variant of the default scale-uncertainty model described in Nixon et al. (Beyond Normalizations / scale-uncertainty framework). Unlike [clr.sm](#), which is centered on the CLR-implied scale, this model is centered on the TSS assumption that there is no systematic change in scale across.

Usage

```
tss.sm(X, logComp, gamma = 0.5)
```

Arguments

<code>X</code>	A numeric design matrix passed internally by <code>aldex()</code> to the scale model. Rows correspond to fixed-effect coefficients/covariates ($P = \text{nrow}(X)$) and columns correspond to samples ($N = \text{ncol}(X)$). (Automatically supplied by <code>aldex()</code> .)
<code>logComp</code>	A numeric array of Monte Carlo log-compositions with dimensions features $\times$ samples $\times$ <code>nsample</code> . This scale model uses <code>nsample</code> (the number of Monte Carlo draws) but does not otherwise use <code>logComp</code> . (Automatically supplied by <code>aldex()</code> .)
<code>gamma</code>	Non-negative scalar. Standard deviation of the Gaussian perturbation applied to the scale-model coefficients (in log2 space). <code>gamma = 0</code> implies no scale uncertainty (all draws are centered at zero effect); larger values allow greater departures from the TSS-centered assumption.

Details

Scale uncertainty is introduced via an additive Gaussian perturbation on the (log2) fixed effects. For each Monte Carlo draw, a coefficient vector is sampled as $b^{(m)} \sim N(0, \gamma^2 I)$, and the per-sample log2 scale is computed as $b^{(m)T} X$. Larger values of `gamma` correspond to weaker confidence in the TSS-centered assumption (more allowed scale variation); `gamma = 0` yields no scale variation beyond the model center.

Note: `logComp` is included to match the ALDEx3 scale-model interface and to determine `nsample`, but it is not otherwise used by this model.

Value

A numeric matrix of dimension $N \times \text{nsample}$ giving Monte Carlo draws of the log2 scale for each sample (rows) across `nsample` draws (columns).

Author(s)

Justin Silverman

References

Nixon G, Gloor GB, Silverman JD (2025). "Incorporating scale uncertainty in microbiome and gene expression analysis as an extension of normalization". *Genome Biology*. doi:10.1186/s13059-025036093

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