

# Package ‘LRDE’

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**Type** Package

**Title** Differential Expression Analysis with Long Read RNA-Seq Data

**Version** 0.99.6

**Description** Provides hurdle negative binomial models for differential expression analysis with long-read RNA-Seq data.

**Depends** R (>= 4.6.0)

**Imports** stats, SummarizedExperiment

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**biocViews** Software, DifferentialExpression, Sequencing, RNASeq, LongRead, GeneExpression, Regression

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**BugReports** <https://github.com/ziyang773/LRDE/issues>

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LRDE-package	<i>LRDE: Differential Expression Analysis for Long-Read RNA-Seq Data</i>
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## Description

The **LRDE** package provides statistical methods for differential expression analysis of long-read RNA sequencing (RNA-Seq) data using a hurdle negative binomial generalized linear model (hurdle-NB GLM).

It implements procedures for:

- Estimation of sample-specific size factors for normalization
- Modeling zero inflation via group-specific expression probabilities
- Gene-wise (tag-wise) dispersion estimation
- Statistical testing for differential expression

These methods are designed to address key challenges in long-read RNA-Seq data, including limited sample sizes and excess zero counts (dropout events).

## Details

The main functions in this package include:

- **prepareDGE**: Prepare count data for analysis. Converts supported input types (`matrix`, `data.frame`, `DGEList`, `DESeqDataSet`, and `SummarizedExperiment`) into a standardized format.
- **sizeFactorsEst**: Estimate sample-specific size factors for normalization.
- **tagwiseEst**: Estimate gene-specific (tag-wise) dispersion parameters for a hurdle negative binomial model using prior information from bin-level estimates.
- **hurdle\_LRT**: Perform gene-wise likelihood ratio tests (LRT) for differential expression.
- **hurdle\_Wald\_Test**: Perform gene-wise Wald tests for differential expression.

## Typical workflow:

1. Prepare data using `prepareDGE`
2. Normalize counts with `sizeFactorsEst`
3. Estimate tag-wise dispersions using `tagwiseEst`
4. Perform differential expression testing with `hurdle_LRT` or `hurdle_Wald_Test`

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**See Also**

[prepareDGE](#), [sizeFactorsEst](#), [tagwiseEst](#), [hurdle\\_LRT](#), [hurdle\\_Wald\\_Test](#)

**Examples**

```
# Load the package
library(LRDE)

# Simulate count data
set.seed(123)
mat <- matrix(rnbinom(300, size = 5, mu = 5), nrow = 50)
grp <- factor(c("A", "A", "A", "B", "B", "B"))

# Prepare data
y <- prepareDGE(mat, grp)

# Normalize counts
y <- sizeFactorsEst(y)

# Estimate dispersions
y <- tagwiseEst(y)

# Differential expression testing
y <- hurdle_Wald_Test(y)
y <- hurdle_LRT(y)

# Access results
head(y$lrt_stats)
head(y$p.values)
```

---

hurdle\_LRT

*Likelihood Ratio Test for Hurdle Negative Binomial Model*

---

**Description**

Performs gene-wise likelihood ratio tests (LRT) for differential expression using a hurdle negative binomial model with fixed zero probabilities and tag-wise dispersions.

**Usage**

```
hurdle_LRT(y)
```

**Arguments**

**y** A list-like object returned from `tagwiseEst()` containing:

- counts** Numeric matrix of gene expression counts (genes x samples).
- samples** Data frame with columns `group` (factor) and `size.factor` (numeric).
- tagwise.disp** Numeric vector of estimated tag-wise dispersions.
- zero\_prob\_matrix** Matrix of zero probabilities per group per gene.

**Details**

For each gene:

- The null model assumes a single shared mean across groups.
- The alternative model estimates group-specific means.
- Zero probabilities and dispersions are fixed from prior estimates.
- When one group has all zero counts, a one-sided Z-test is applied instead.

**Value**

The input object `y` with two additional elements:

**lrt\_stats** Numeric vector of LRT statistics for each gene.

**p.values** Numeric vector of corresponding p-values.

**Examples**

```
set.seed(123)
mat <- matrix(rnbinom(30, size = 5, mu = 5), nrow = 5)
grp <- c("A", "A", "A", "B", "B", "B")
y <- prepareDGE(mat, grp)
y <- sizeFactorsEst(y)
y <- tagwiseEst(y)
y <- hurdle_LRT(y)
```

---

hurdle\_Wald\_Test

*Wald Test for Hurdle Negative Binomial Model*

---

**Description**

Performs gene-wise Wald tests for differential expression using a hurdle negative binomial model with fixed zero probabilities and tag-wise dispersions.

**Usage**

```
hurdle_Wald_Test(y)
```

**Arguments**

**y** A list-like object returned from `tagwiseEst()` containing:

- counts** Numeric matrix of gene expression counts (genes x samples).
- samples** Data frame with columns `group` (factor) and `size.factor` (numeric).
- tagwise.disp** Numeric vector of estimated tag-wise dispersions.
- zero\_prob\_matrix** Matrix of zero probabilities per group per gene.

**Details**

For each gene:

- Zero probabilities and dispersions are fixed from prior estimates.
- The model estimates group-specific mean parameters.
- When one group has all zero counts, a one-sided Z-test is applied instead.
- Otherwise, a standard two-sided Wald test is applied on the log-difference of group means.

**Value**

The input object `y` with two additional elements:

**wald\_stats** Numeric vector of Wald statistics for each gene.

**p.values** Numeric vector of corresponding p-values.

**Examples**

```
set.seed(123)
mat <- matrix(rnbinom(30, size = 5, mu = 5), nrow = 5)
grp <- c("A", "A", "A", "B", "B", "B")
y <- prepareDGE(mat, grp)
y <- sizeFactorsEst(y)
y <- tagwiseEst(y)
y <- hurdle_Wald_Test(y)
```

---

```
prepareDGE
```

---

*Prepare Count Data for Differential Expression Analysis*

---

**Description**

Converts various supported input types to a standardized list format for downstream differential expression analysis. Supports `matrix`, `data.frame`, `DGEList`, `DESeqDataSet`, and `SummarizedExperiment` objects.

**Usage**

```
prepareDGE(data, group)
```

**Arguments**

data	A numeric matrix, data.frame, or supported object containing counts.
group	A vector of group labels for the columns/samples of data. Must be the same length as the number of columns in data.

**Details**

This function performs input validation.

- Checks for non-negative numeric values and absence of NA.
- Ensures group labels match the number of samples.
- Automatically assigns column names if missing.
- Returns a list suitable for use with hurdle model-based DE functions.

**Value**

A list with two elements:

**counts** An integer matrix of counts.

**samples** A data.frame containing sample-level metadata: group, lib.size, and size.factor.

**Examples**

```
# Example with a matrix
set.seed(123)
mat <- matrix(rnbinom(30, size = 5, mu = 5), nrow = 5)
grp <- c("A", "A", "A", "B", "B", "B")
y <- prepareDGE(mat, grp)

# Example with a SummarizedExperiment
if (requireNamespace("SummarizedExperiment", quietly = TRUE)) {
  se <- SummarizedExperiment::SummarizedExperiment(assays = list(counts = mat))
  y_se <- prepareDGE(se, grp)
  y
}
```

---

priorEst

*Estimate Bin-wise Priors for Hurdle Model Parameters (Internal)*

---

**Description**

Internal function to group genes by similar mean expression levels and perform bin-wise estimation of prior parameters for the hurdle negative binomial model. Within each bin, genes are pooled and a global model is fitted to estimate group-specific non-zero probabilities and log dispersion.

**Usage**

```
priorEst(y, n_bins = NULL)
```

**Arguments**

**y** A list object produced by `sizeFactorsEst()`, containing counts, samples, and `baseMean`.

**n\_bins** Optional integer specifying the number of bins. If `NULL`, the number of bins is determined automatically.

**Details**

Genes are partitioned into bins based on  $\log(\text{baseMean})$ . For each bin, a hurdle negative binomial model is fitted using `nll_hurdle()` on pooled counts to obtain bin-level estimates of:

- Group-specific non-zero probabilities
- Log-dispersion parameter ( $\log(\phi)$ )

The estimated parameters are mapped back to individual genes to provide stabilized priors for downstream modeling.

**Value**

The input object `y` with added components:

**prob\_matrix** Matrix of group-specific non-zero probabilities.

**prior\_log\_phi\_gene** Gene-wise  $\log(\phi)$  estimates.

**prior\_bins** `Data.frame` of bin-level parameter estimates.

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sizeFactorsEst                      *Estimate Size Factors for Normalization*

---

**Description**

Computes sample-specific size factors for long-read RNA-Seq data, used to normalize counts for differential expression analysis.

**Usage**

```
sizeFactorsEst(y, type = c("poscounts", "ratio"), locfunc = stats::median)
```

## Arguments

y	A count matrix ( <code>matrix</code> or <code>data.frame</code> ) or the output of <code>prepareDGE()</code> . If a <code>matrix/data.frame</code> is provided, the function assumes two equal-sized groups.
type	Character string specifying the method for estimating size factors: <b>"poscounts"</b> Geometric mean-based method. <b>"ratio"</b> Simple ratio method using the mean of log counts. Default is "poscounts".
locfunc	Function to summarize log-ratios across genes. Defaults to <code>median</code> .

## Details

This function implements two methods for size factor estimation:

- **poscounts**: Computes a geometric mean of positive counts per gene, then calculates ratios for each sample. Normalizes so that the geometric mean of size factors equals 1.
- **ratio**: Uses the mean of log-counts per gene across samples to compute ratios.

The function automatically normalizes counts using the estimated size factors and stores gene-level normalized means in `baseMean`.

## Value

A list (same structure as `prepareDGE()` output) with:

**counts** Original count matrix (integer).

**samples** Data frame with sample information, updated `size.factor`.

**baseMean** Normalized mean of counts per gene.

## Examples

```
# Using a count matrix
#' set.seed(123)
mat <- matrix(rnbinom(30, size = 5, mu = 5), nrow = 5)
grp <- c("A", "A", "A", "B", "B", "B")
y <- prepareDGE(mat, grp)
y <- sizeFactorsEst(y, type = "poscounts")
```

---

`tagwiseEst`*Tag-wise Dispersion Estimation for Hurdle Negative Binomial Model*

---

### Description

Estimate gene-specific (tag-wise) dispersion parameters for a hurdle negative binomial model using prior information derived from bin-level estimates.

### Usage

```
tagwiseEst(y)
```

### Arguments

`y` A list object created by `prepareDGE` with size factors estimated, containing counts and sample information.

### Details

This function performs the following steps:

1. Retrieves bin-level prior estimates of zero probabilities and log-dispersion for each gene via `priorEst`.
2. Fixes the zero probabilities and optimizes only the mean parameters and dispersion for each gene individually.
3. Uses the internal function `nll_hurdle_fixed_P` to compute the negative log-likelihood with fixed zero probabilities.

The resulting `tagwise.disp` will be used for downstream differential expression analysis.

### Value

The input `y` object augmented with:

**tagwise.disp** Numeric vector of estimated gene-wise dispersions.

**zero\_prob\_matrix** Numeric matrix of fixed zero probabilities for each gene and group.

### Examples

```
set.seed(123)
mat <- matrix(rnbinom(30, size = 5, mu = 5), nrow = 5)
grp <- c("A", "A", "A", "B", "B", "B")
y <- prepareDGE(mat, grp)
y <- sizeFactorsEst(y)
y <- tagwiseEst(y)
head(y$tagwise.disp)
head(y$zero_prob_matrix)
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