# Package 'LDAcoop'

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Type Package

**Title** Analysis of Data from Limiting Dilution Assay (LDA) with or without Cellular Cooperation

Version 0.1.2

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URL https://github.com/ZytoHMGU/LDAcoop

BugReports https://github.com/ZytoHMGU/LDAcoop/issues

Description Cellular cooperation compromises the established method of calculating clonogenic activity from limiting dilution assay (LDA) data. This tool provides functions that enable robust analysis in presence or absence of cellular cooperation. The implemented method incorporates the same cooperativity module to model the non-linearity associated with cellular cooperation as known from the colony formation assay (Brix et al. (2021) <doi:10.1038/s41596-021-00615-0>: ``Analysis of clonogenic growth in vitro." Nature protocols).

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**Encoding UTF-8** 

LazyData true

RoxygenNote 7.2.3

**Depends** R (>= 3.5.0)

Imports Hmisc

**Suggests** knitr, rmarkdown, testthat (>= 3.0.0)

VignetteBuilder knitr

Config/testthat/edition 3

NeedsCompilation no

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# Repository CRAN

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# Description

LDA data from 11 cell lines, up to 4 biological replicates and up to 6 treatments.

## Usage

```
data(LDAdata)
```

## **Format**

```
data.frame with columns: "name", "replicate", "Group", "S-value", "# Tested", "# Clonal growth"
```

```
data(LDAdata)
```

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LDA\_activity LDA\_activity

#### Description

calculation of clonogenic activities from data collected in a limiting dilution assay (LDA) experiment (i.e. cells, wells, positive wells, group).

#### Usage

```
LDA_activity(x, name = "LDA cells")
```

#### **Arguments**

x numeric data.frame or matrix with three columns (cells, wells, positive wells,

group (optional))

name optional: experiment name (e.g. name of cell line)

#### Value

list object with LDA-activities as returned by LDA\_activity\_single

#### **Examples**

#### **Description**

calculation of clonogenic activity from data collected by a limiting dilution assay (LDA) experiment (i.e. numbers of: cells seeded, wells, positive wells).

## Usage

```
LDA_activity_single(x, name = "cell line a", treat = "no")
```

# Arguments

x numeric data.frame or matrix with three columns (cells, wells, positive wells)

name optional: experiment name (e.g. name of cell line) treat optional: treatment (e.g. irradiation dose in Gy)

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#### Value

list object with estimated activity, 95 84 matrix, fit-object and p-value for cooperativity-test

## **Examples**

LDA\_plot

LDA\_plot

### **Description**

plot clonogenic activity and survival (at more than one treatment group) for data from limiting dilution assay (LDA) experiments.

### Usage

```
LDA_plot(LDA_tab, uncertainty = "act", xlim = NULL, uncertainty.band = FALSE)
```

#### **Arguments**

```
LDA_tab LDA data.frame ("cells", "wells", "positive", "group", "replicate")
uncertainty method for uncertainty calculation ("act", "ep")
xlim setting xlim of clonogenic activity plot
uncertainty.band
plotting of uncertainty bands TRUE/FALSE
```

#### Value

none

```
data(LDAdata)
Z1 <- subset.data.frame(LDAdata,subset = name == unique(LDAdata$name)[1])
LDA_plot(Z1[,c("S-value","# Tested","# Clonal growth","Group","replicate")])</pre>
```

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```
LDA_plot_activity LDA_plot_activity
```

## **Description**

generate clonogenic activity estimation plot (frequency of negative wells over the number of cells seeded) for data of limiting dilution assay (LDA) experiments. Input is an data object as returned by the preprocessing function LDA\_prepare\_plot().

#### Usage

```
LDA_plot_activity(LDA_obj, xlim = NULL, uncertainty.band = FALSE)
```

## Arguments

```
LDA_obj list returned from LDA_prepare_plot

xlim manually setting the xlim

uncertainty.band

plotting uncertainty bands TRUE/FALSE
```

## Value

none

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LDA\_plot\_SF

LDA\_plot\_SF

## **Description**

generate clonogenic survival plot (estimated clonogenic survival over treatment) for data from limiting dilution assay (LDA). Input is an data object as returned by the preprocessing function LDA\_prepare\_plot().

## Usage

```
LDA_plot_SF(LDA_obj)
```

#### **Arguments**

LDA\_obj

list returned from LDA\_prepare\_plot

#### Value

none

## **Examples**

LDA\_prepare\_plot

LDA\_prepare\_plot

#### **Description**

analyze limiting dilution assay (LDA) data and collect information for plotting.

#### Usage

```
LDA_prepare_plot(LDA_tab, uncertainty = "act")
```

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## Arguments

LDA\_tab LDA data.frame ("cells", "wells", "positive", "group", "replicate")

uncertainty method for approximation of uncertainties of survival fractions (SF): activity

based ("act") or by error propagation ("ep")

#### Value

none

#### **Examples**

LDA\_survival

LDA\_survival

## Description

calculation of clonogenic survival in a table of data from a limiting dilution assay (LDA) experiment (i.e. cells, wells, positive wells, group).

#### Usage

```
LDA_survival(x, name = "cell line a")
```

# **Arguments**

x numeric data.frame or matrix with three columns (cells, wells, positive wells, group)

name optional: experiment name (e.g. name of cell line)

#### Value

list object with LDA-activities as returned by LDA\_activity\_single

#### **Examples**

```
LDA_survival_single LDA_survival_single
```

## **Description**

calculate clonogenic survival fraction from LDA\_activity objects.

#### Usage

```
LDA_survival_single(act.0, act.x)
```

## Arguments

```
act.0 reference activity
act.x activity after treatment
```

#### Value

list object with survival fraction, estimated confidence intervals (by error propagation through first order Taylor series approximation and by combination of 84

LDA\_table 9

LDA_table LD
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#### **Description**

show table with activities and clonogenic survival from limiting dilution assay (LDA) data.

## Usage

```
LDA_table(x, ref_class = "unknown", uncertainty = "act")
```

#### **Arguments**

ref\_class

x numeric data.frame or matrix with at least three columns (cells, wells, positive wells, group (optional))

name of reference class for calculation of SF values

uncertainty method for calculating the uncertainty bands of survival fractions ("act" (default)

for combining the activity confidence intervals; "ep" for error propagation via

first order Taylor series expansion.)

#### Value

table

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