# Package 'qPLEXdata'

October 17, 2019

Type Package

Title Data accompanying qPLEXanalyzer package
Version 1.2.0
<b>Date</b> 2018-11-14
<b>Description</b> qPLEX-RIME and Full proteome TMT mass spectrometry datasets.
<b>Depends</b> R (>= 3.5), qPLEXanalyzer
Imports utils, knitr, MSnbase, dplyr
VignetteBuilder knitr
License GPL-2
biocViews ExperimentData, MassSpectrometryData, Proteome
NeedsCompilation no
git_url https://git.bioconductor.org/packages/qPLEXdata
git_branch RELEASE_3_9
git_last_commit c3fe91f
git_last_commit_date 2019-05-02
Date/Publication 2019-10-17
Author Kamal Kishore Developer [aut, cre]
Maintainer Kamal Kishore Developer <kamal.fartiyal84@gmail.com></kamal.fartiyal84@gmail.com>
R topics documented:
exp1_specificity exp2_Xlink exp3_OHT_ESR1 exp4_OHT_FP exp5_PDX exp6_ER exp7_NCOA3 exp8_CBP exp9_PolII qPLEXdata
Index

2 exp2\_Xlink

exp1\_specificity

exp1\_specificity dataset

# **Description**

In this experiment we have used the qPLEX-RIME approach to identify ER specific interactors. We performed replicate ER RIME pull-downs in five independent biological replicates and an equal number of matched IgG mock samples was included.

# Usage

```
data(exp1_specificity)
```

#### **Format**

An object of class list related to peptides quantification. It consists of qPLEX-RIME data of 10 samples divided into two conditions (ER and IgG).

#### Value

An object of class list related to peptides quantification.

exp2\_Xlink

exp2\_Xlink dataset

# **Description**

An ER qPLEX-RIME experiment was performed to compare two different ways of cell crosslinking. MCF7 cells were double crosslinked with DSG/formaldehyde (double) or with formaldehyde alone (single). Four biological replicates were obtained for each condition along with two IgG pooled samples from each replicate.

# Usage

```
data(exp2_Xlink)
```

# **Format**

An object of class list related to peptides quantification. It consists of qPLEX-RIME data of 10 samples divided into three conditions (FA, DSG.FA and IgG).

#### Value

exp3\_OHT\_ESR1 3

exp3\_OHT\_ESR1

exp3\_OHT\_ESR1 dataset

#### **Description**

Three ER qPLEX-RIME (10plex) experiments were performed to investigate the dynamics of the ER complex assembly upon 4-hydrotamoxifen (OHT) treatment at 2h, 6h and 24h or at 24h post-treatment with the drug-vehicle alone (ethanol). Two biological replicates of each condition were included in each experiment to finally consider a total of six replicates per time point. Additionally, MCF7 cells were treated with OHT or ethanol and cross-linked at 24h post-treatment in each experiment to be used for mock IgG pull-downs and to enable discrimination of non-specific binding in the same experiment. This is a timecourse experiment to study the effect of tamoxifen in ER interactome using qPLEX-RIME method.

# Usage

```
data(exp3_OHT_ESR1)
```

#### **Format**

An object of class list related to peptides quantification. It consists of qPLEX-RIME data from three experimental runs. Each run contains 10 samples divided into five conditions (IgG, vehicle, tam.2h, tam.6h and tam.24h).

#### Value

An object of class list related to peptides quantification.

exp4\_OHT\_FP

exp4\_OHT\_FP dataset

# **Description**

We performed two 10plex-TMT time-course experiments to study the effect of 4-hydrotamoxifen (OHT) on total protein levels. MCF7 cells were treated with OHT for 2h, 6h, 24h or for 24h with the drug-vehicle alone (ethanol) and a total number of four biological replicates were obtained. This is a timecourse experiment to study the effect of tamoxifen on full proteome.

#### Usage

```
data(exp4_OHT_FP)
```

#### **Format**

An object of class list related to peptides quantification. It consists of total proteome data from two experimental runs. Each run contains 10 samples divided into four conditions (vehicle, tam.2h, tam.6h and tam.24h).

#### Value

4 exp6\_ER

exp5\_PDX

exp5\_PDX dataset

#### **Description**

An ER qPLEX-RIME experiment was performed using three independent ER+ human Patient Derived Xenograft (PDX) tumour material. Cryosections of each tumour were double-crosslinked and each tumour was split in two parts that were used for ER and IgG RIME pull-down assays. One of the tumours was split in three different parts to be used as ER or IgG qPLEX-RIME in order to assess technical variability.

#### Usage

data(exp5\_PDX)

#### **Format**

An object of class list related to peptides quantification. It consists of qPLEX-RIME data of 10 samples divided into two conditions (PDX and IgG).

#### Value

An object of class list related to peptides quantification.

exp6\_ER

exp6 ER dataset

# **Description**

An ER qPLEX-RIME experiment was performed using five independent ER-positive human breast cancer tumours. Cryosections of each tumour were double-crosslinked and each tumour was split in two parts that were used for ER and IgG RIME pull-down assays.

## Usage

data(exp6\_ER)

#### **Format**

An object of class list related to peptides quantification. It consists of qPLEX-RIME data of 10 samples divided into two conditions (ER and IgG).

#### Value

exp7\_NCOA3 5

exp7\_NCOA3

exp7\_NCOA3 dataset

# **Description**

In this experiment we have used the qPLEX-RIME method to identify and characterize NCOA3 (SRC-3) associated proteins. We performed NCOA3 RIME pull-downs in five independent biological replicates and in five matched IgG mock samples.

# Usage

```
data(exp7_NCOA3)
```

#### **Format**

An object of class list related to peptides quantification. It consists of qPLEX-RIME data of 10 samples divided into two conditions (NCOA3 and IgG).

#### Value

An object of class list related to peptides quantification.

exp8\_CBP

exp8\_CBP dataset

#### **Description**

A qPLEX-RIME experiment was designed for the characterization of the CBP (CREB-binding protein) interactome. Five independent biological replicates of CBP RIME pull-downs and five IgG RIME pull-downs were prepared for this experiment.

# Usage

```
data(exp8_CBP)
```

# **Format**

An object of class list related to peptides quantification. It consists of qPLEX-RIME data of 10 samples divided into two conditions (CBP and IgG).

# Value

6 qPLEXdata

exp9\_PolII

exp9\_PolII dataset

# **Description**

The qPLEX-RIME method was applied for the characterization of the largest and catalytic component of RNA polymerase II (RPB1). Particularly, the phosphorylated form at Serine 5 in the Cterminal domain (CTD) was used as the bait protein. Five biological replicates of RNA polymerase II RIME pull-downs and five IgG pull-downs were included for the identification and characterization of RNA polymerase II-associated proteins.

#### Usage

```
data(exp9_PolII)
```

# **Format**

An object of class list related to peptides quantification. It consists of qPLEX-RIME data of 10 samples divided into two conditions (PolII and IgG).

#### Value

An object of class list related to peptides quantification.

qPLEXdata

Available datasets in the qPLEXdata package

# Description

This function lists the datasets available in qPLEXdata package

# Usage

```
qPLEXdata()
```

#### Value

A list of datasets

# **Examples**

```
qPLEXdata()
```

# **Index**

```
*Topic datasets
    exp1\_specificity, 2
    exp2_Xlink, 2
    exp3_OHT_ESR1, 3
    exp4_OHT_FP, 3
    exp5_PDX, 4
    exp6_ER, 4
    exp7_NCOA3, 5
    exp8_CBP, 5
    exp9_PolII, 6
*Topic data
    exp1_specificity, 2
    exp2_Xlink, 2
    exp3_OHT_ESR1, 3
    exp4_OHT_FP, 3
    exp5_PDX, 4
    exp6_ER, 4
    exp7_NCOA3, 5
    exp8_CBP, 5
    exp9_PolII, 6
exp1_specificity, 2
exp2_Xlink, 2
exp3_OHT_ESR1, 3
exp4_OHT_FP, 3
exp5_PDX, 4
exp6_ER, 4
exp7_NCOA3, 5
exp8_CBP, 5
exp9_PolII, 6
list, 2-6
qPLEXdata, 6
```