Package 'Xeva'

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```
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Description The Xeva package provides efficient and powerful functions for patient-
      drived xenograft (PDX) based pharmacogenomic data analysis.
     This package contains a set of functions to perform analysis of patient-
      derived xenograft data. This package was developed by the BHKLab, for further informa-
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ABC

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Description

area between curves Computes the area between two time-volume curves.

Usage

```
ABC(
  contr.time = NULL,
  contr.volume = NULL,
  treat.time = NULL,
  treat.volume = NULL)
```

Arguments

```
contr.time Time vector for control.

contr.volume Volume vector for control.

treat.time Time vector for treatment.

treat.volume Volume vector for treatment.
```

Value

Returns batch response object.

Examples

addExperimentalDesign Add a new experimental design

Description

Add a new experimental design in the expDesign slot.

```
addExperimentalDesign(
  object,
  treatment = NULL,
  control = NULL,
  batch.id = NULL,
  replace = FALSE
)

## S4 method for signature 'XevaSet'
```

4 angle

```
addExperimentalDesign(
  object,
  treatment = NULL,
  control = NULL,
  batch.id = NULL,
  replace = FALSE
)
```

Arguments

object The Xeva dataset.

treatment The model.id of treatment.

control The model.id of control.

batch.id The batch.id for a new batch.

replace If TRUE, replace an old batch with new values.

Value

Returns Xeva dataset with new experimental design added.

Examples

angle

compute angle Computes the angle between two time-volume curves.

Description

compute angle Computes the angle between two time-volume curves.

Usage

```
angle(
  contr.time = NULL,
  contr.volume = NULL,
  treat.time = NULL,
  treat.volume = NULL,
  degree = TRUE
)
```

Arguments

degree Default TRUE will give angle in degrees and FALSE will return in radians.

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Value

Returns batch response object.

Examples

```
contr.time <- treat.time <- c(0, 3, 7, 11, 18, 22, 26, 30, 32, 35)
contr.volume<- contr.time * tan(60*pi/180)
treat.volume<- treat.time * tan(15*pi/180)
ang <- angle(contr.time, contr.volume, treat.time, treat.volume)
print(ang)
par(pty="s")
xylimit <- range(c(contr.time, contr.volume, treat.time, treat.volume))
plot(contr.time, contr.volume, type = "b", xlim = xylimit, ylim = xylimit)
lines(treat.time, treat.volume, type = "b")
abline(lm(contr.volume~contr.time))
abline(lm(treat.volume~treat.time))</pre>
```

AUC

area under the curve AUC Returns area under the curve

Description

area under the curve AUC Returns area under the curve

Usage

```
AUC(time, volume)
```

Arguments

time A vector of time points recorded for the experiment.

volume First vector of volume.

Value

Returns angle and slope object.

```
time <- c(0, 3, 7, 11, 18, 22, 26, 30, 32, 35)
volume1<- time * tan(30*pi/180)
volume2<- time * tan(45*pi/180)
auc1 <- AUC(time, volume1)
auc2 <- AUC(time, volume2)
par(pty="s")
xylimit <- range(c(time, volume1, volume2))
plot(time, volume1, type = "b", xlim = xylimit, ylim = xylimit)
lines(time, volume2, type = "b")
abline(lm(volume1~time))
abline(lm(volume2~time))</pre>
```

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batchInfo

Get batch information

Description

Get batch information from a Xeva dataset.

Usage

```
batchInfo(
  object,
  batch = NULL,
  model.id = NULL,
  model.id.type = c("any", "control", "treatment")
)

## S4 method for signature 'XevaSet'
batchInfo(
  object,
  batch = NULL,
  model.id = NULL,
  model.id.type = c("any", "control", "treatment")
)
```

Arguments

object The Xeva object from which batch information is obtained.

batch Name of the batch. Default NULL.

model.id Model ID for which need to be searched in the batches. Default NULL.

model.id.type Type of the model ID in a batch. See the Details section below.

Details

By default this function will return the names of all the batches present in the dataset. If a batch specified, it will return the experiment design (control and treatment model IDs) of that particular batch. If model.id is specified, it will return the names of all the batches where this particular model.id is present. If both batch and model.id are specified, batch will take precedent.

For model.id.type, the default value 'any' will return all the batch IDs where the given model ID is present in any arm (ie. control or treatment) of the batch. It can also be restricted to look only for treatment (or control) arm by specifying the type.

Value

A Vector with batch names.

```
data(brca)
##to get all the batch names
batch.name <- batchInfo(brca)</pre>
```

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```
##to get a specific batch
batch.design <- batchInfo(brca, batch=batch.name[1])
##to get all the batches where a model.id is present
batchInfo(brca, model.id="X.6047.uned")</pre>
```

brca

PDXE breast cancer dataset

Description

A Xeva object containing only breast cancer PDXs from the PDXE dataset For details about PDX-MI, see: Gao et al. High-throughput screening using patient-derived tumor xenografts to predict clinical trial drug response. Nature medicine, 21(11):1318, 2015.

Usage

```
data(brca)
```

Format

An object of class XevaSet of length 1.

Source

```
https://www.nature.com/articles/nm.3954?draft=journal
```

createXevaSet

XevaSet constructor

Description

A constructor to create XevaSet. Only objects returned by this constructor are expected to work with the XevaSet methods.

```
createXevaSet(
  name,
  model = data.frame(),
  drug = data.frame(),
  experiment = data.frame(),
  expDesign = list(),
  modelSensitivity = data.frame(),
  batchSensitivity = data.frame(),
  molecularProfiles = list(),
  modToBiobaseMap = data.frame()
)
```

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Arguments

name A character string detailing the name of the dataset.

model A data. frame containing the annotations for all the models used in the experi-

ment.

drug A data. frame containing the annotations for all the drugs profiled in the dataset,

across all data types.

experiment A data. frame containing all experiment information.

expDesign A list containing name of the batch, control and treatment model.id

modelSensitivity

A data. frame containing sensitivity for each model

batchSensitivity

A data. frame containing sensitivity for each batch

molecularProfiles

A list of ExpressionSet objects containing different molecular profiles.

modToBiobaseMap

A data. frame containing model.id corresponding Biobase object id and name of the molecular Profiles

Details

This function creates a XevaSet object. It takes different model infromation and genomic data as input. For detailed discription of all varaibles please see Xeva vignette section "Creating new Xeva object"

Value

Returns Xeva object

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dosePlot

plot dose data

Description

plot data for dose in model.id

Usage

```
dosePlot(
  object,
  model.id,
  max.time = NULL,
  treatment.only = FALSE,
  vol.normal = FALSE,
  concurrent.time = FALSE,
  point.shape = 21,
  point.size = 3,
  line.size = 4,
  point.color = "#878787",
  line.color = "#bababa",
  fill.col = c("#f5f5f5", "#E55100"),
  modify.x.axis = FALSE
)
```

Arguments

object Xeva object. model.id one or multiple model.id Maximum time point of the plot. Default NULL will plot complete data max.time treatment.only Default FALSE. Given full data treatment.only=TRUE will plot data only during treatment vol.normal Default FALSE. If TRUE, volume will be normalized concurrent.time Default FALSE. If TRUE, cut the batch data such that control and treatment will end at the same time point point.shape shape of the point point.size size of the point line.size size of the line point.color color for point line.color color for line fill.col a vector with color to fill modify.x.axis Default FALSE

Value

A ggplot2 plot

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Examples

```
data(brca)
dosePlot(brca, model.id=c("X.6047.LJ16","X.6047.LJ16.trab"), fill.col=c("#f5f5f5", "#993404"))
```

downloadXevaSet

Download a XevaSet object or table of available XevaSet objects

Description

This function allows you to see the available XevaSet object and download them for use with this package. The XevaSet have been extensively curated and organised within a XevaSet class, enabling use with all the analysis tools provided in Xeva.

Usage

```
downloadXevaSet(
  name = NULL,
  saveDir = file.path(".", "XevaSet"),
  XevaSetFileName = NULL,
  verbose = TRUE
)
```

Arguments

name Character string, the name of the XevaSet to download.

saveDir Character string with the folder path where the XevaSet should be saved. De-

faults to './XevaSet/'. Will create directory if it does not exist.

XevaSetFileName

character string, the file name to save the dataset under

verbose bool Should status messages be printed during download. Defaults to TRUE.

Value

A data frame if name is NULL, showing all the available XevaSet objects. If name is specified, it will download the dataset from our server

drugInform

Get drug information Get the drug information slot from a XevaSet object.

Description

Get drug information Get the drug information slot from a XevaSet object.

```
drugInform(object)
## S4 method for signature 'XevaSet'
drugInform(object)
```

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Arguments

object The XevaSet to retrieve drug information from.

Value

A data. frame with the drug annotations.

Examples

```
data(brca)
head(drugInform(brca))
```

drugSensitivitySig

get drug sensitivity values

Description

Given a Xeva object and drug name, this function will return sensitivity values for all the genes/features.

Usage

```
drugSensitivitySig(
  object,
  drug,
  mDataType = NULL,
  molData = NULL,
  features = NULL,
  model.ids = NULL,
  model2bidMap = NULL,
  sensitivity.measure = "slope",
  fit = c("lm", "CI", "pearson", "spearman", NA),
  standardize = c("SD", "rescale", "none"),
  nthread = 1,
  tissue = NULL,
  verbose = TRUE
)
```

Arguments

object The Xeva dataset.
drug Name of the drug.
mDataType Molecular data type.

molData External data matrix. Rows as features and columns as samples.

features Set which molecular data features to use. Default NULL will use all features.

model.ids Set which model.id to use from the dataset. Default NULL will use all model.ids.

model2bidMap A data.frame with model.id and biobase.id. Default NULL will use internal

mapping.

sensitivity.measure

Name of the sensitivity measure.

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fit Association method to use, can be 'lm', 'CI', 'pearson' or 'spearman'. If 'NA'

only the data will be return. Default 1m.

standardize Default SD. Name of the method to use for data standardization before fitting.

nthread number of threads

tissue tissue type. Default NULL uses 'tissue' from object.

verbose Default TRUE will show information

Details

Method to compute association can be specified by fit. It can be one of the:

• "lm" for linear models

• "CI" for concordance index

• "pearson" for Pearson correlation

• "spearman" for Spearman correlation

If fit is set to NA, processed data (an ExpressionSet) will be returned.

A matrix of values can be directly passed to molData. In case where a model.id maps to multiple biobase.ids, the first biobase.id in the data.frame will be used.

Value

A data. frame with features and values.

Examples

getExperiment

Get PDX experiment data

Description

For a given model.id, getExperiment will

getExperiment 13

Usage

```
getExperiment(
  object,
  model.id = NULL,
  batch = NULL,
  patient.id = NULL,
  drug = NULL,
  control.name = NULL,
  treatment.only = FALSE,
  max.time = NULL,
  vol.normal = FALSE,
  log.volume = FALSE,
  return.list = FALSE,
  impute.value = FALSE,
  concurrent.time = FALSE
)
## S4 method for signature 'XevaSet'
getExperiment(
  object,
  model.id = NULL,
  batch = NULL,
  patient.id = NULL,
  drug = NULL,
  control.name = NULL,
  treatment.only = FALSE,
  max.time = NULL,
  vol.normal = FALSE,
  log.volume = FALSE,
  return.list = FALSE,
  impute.value = FALSE,
  concurrent.time = FALSE
)
```

Arguments

impute.value

object The XevaSet object. model.id The model.id for which data is required, multiple IDs are allowed. Batch name from the XevaSet or experiment design. batch Patient id from the XevaSet. Default NULL. patient.id Name of the drug. drug Name of drug used as control. Default NULL. control.name Default FALSE. If TRUE, give data for non-zero dose periods only (if dose data treatment.only are available). max.time Maximum time for data. vol.normal If TRUE it will normalize the volume. Default FALSE. If TRUE log of the volume will be used. Default FALSE. log.volume return.list Default FALSE will return a data.frame.

Default FALSE. If TRUE, impute the missing values.

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```
concurrent.time
```

Default FALSE. If TRUE, cut the batch data such that control and treatment will end at same time point.

Value

a data. fram will all the the values stored in experiment slot

Examples

getMolecularProfiles Get molecular profiles from a XevaSet object

Description

This function serves to get molecular profiles from a XevaSet object.

Usage

```
getMolecularProfiles(object, data.type)
```

Arguments

object The XevaSet.

data.type character, where one of the molecular data types is needed.

Value

An ExpressionSet where sample names are the biobase.id of the model.

```
data(brca)
brca.RNA <- getMolecularProfiles(brca, data.type="RNASeq")</pre>
```

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1mm

linear mixed model

Description

Comput the linear mixed model (lmm) statistics for a PDX batch

Usage

```
1mm(data)
```

Arguments

data

a data.frame containg a batch data

Details

The input data.frame (data) must contain these columns: model.id, volume, time, exp.type

Value

Returns a fit object

Examples

```
data(repdx)
data <- getExperiment(repdx, batch = "P1")$model
lmm(data)</pre>
```

modelInfo

modelInfo Generic Generic for modelInfo method

Description

modelInfo Generic Generic for modelInfo method

Usage

```
modelInfo(object, mDataType = NULL)
## S4 method for signature 'XevaSet'
modelInfo(object, mDataType = NULL)
```

Arguments

object Xeva object

mDataType Molecular data type.

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Value

A data. frame with the model annotations.

Examples

```
data(brca)
mid <- modelInfo(brca)
head(mid)</pre>
```

mRECIST

Computes the mRECIST

Description

 $\ensuremath{\mathsf{mRECIST}}$ Returns the mRECIST for given volume response.

Usage

```
mRECIST(time, volume, min.time = 10, return.detail = FALSE)
```

Arguments

time Value of best response.

volume Value of best average response.

min.time Minimum time after which tumor volume will be considered.

return.detail Default FALSE. If TRUE, return all intermediate values.

Value

Returns the mRECIST.

```
time <- c(0, 3, 7, 11, 18, 22, 26, 30, 32, 35)
volume<- c(250.8, 320.4, 402.3, 382.6, 384, 445.9, 460.2, 546.8, 554.3, 617.9)
mRECIST(time, volume, min.time=10, return.detail=FALSE)
```

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PDXMI

PDX-MI data

Description

A dataset containing PDX models minimal information (PDX-MI) standard and corresponding Xeva variable.

Usage

```
data(PDXMI)
```

Format

An object of class data. frame with 45 rows and 4 columns.

Details

For details about PDX-MI, see:

Meehan, Terrence F., et al. "PDX-MI: minimal information for patient-derived tumor xenograft models." Cancer research 77.21 (2017): e62-e66.

Source

```
http://cancerres.aacrjournals.org/lookup/doi/10.1158/0008-5472.CAN-17-0582
```

plotmRECIST

To plot mRECIST values

Description

plotmRECIST plots the mRECIST matrix obtained from summarizeResponse.

```
plotmRECIST(
  mat,
  control.name = NA,
  control.col = "#238b45",
  drug.col = "black",
  colPalette = NULL,
  name = "Drug & Models",
  sort = TRUE,
  row_fontsize = 12,
  col_fontsize = 12,
  draw_plot = TRUE
)
```

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Arguments

mat The mRECIST matrix where rows are drugs and columns are patients.

control.name Name of the control.

control.col Color of the control.

drug.col Color of the drug names.

colPalette Color palette for mRECIST values.

name Title of the plot.

sort If matrix should be sorted before plotting.

row_fontsize Size of the row name font.
col_fontsize Size of the column name font.

draw_plot Default TRUE will plot the figure. If FALSE, return an object.

Value

mRECIST plot.

Examples

```
data(brca)
brca.mr <- summarizeResponse(brca, response.measure = "mRECIST", group.by="patient.id")
plotmRECIST(as.matrix(brca.mr), control.name = "untreated")</pre>
```

plotPDX

Plot batch data

Description

Plot data for a batch.id, experiment design or model.id

```
plotPDX(
  object,
  batch = NULL,
  patient.id = NULL,
  drug = NULL,
  model.id = NULL,
  model.color = NULL,
  control.name = NULL,
  max.time = NULL,
  treatment.only = FALSE,
  vol.normal = FALSE,
  impute.value = TRUE,
  concurrent.time = FALSE,
  control.col = "#e41a1c",
  treatment.col = "#377eb8",
  title = "",
  xlab = "Time",
```

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```
ylab = "Volume",
  log.y = FALSE,
  SE.plot = c("all", "none", "errorbar", "ribbon"),
  aspect.ratio = c(1, NULL),
  minor.line.size = 0.5,
  major.line.size = 0.7
plotBatch(
  object,
  batch = NULL,
  patient.id = NULL,
  drug = NULL,
  control.name = NULL,
  max.time = NULL,
  treatment.only = FALSE,
  vol.normal = FALSE,
  impute.value = TRUE,
  concurrent.time = FALSE,
  control.col = "#6baed6",
  treatment.col = "#fc8d59",
  title = "",
  xlab = "Time"
  ylab = "Volume",
  log.y = FALSE,
  SE.plot = c("all", "none", "errorbar", "ribbon"),
  aspect.ratio = c(1, NULL),
  minor.line.size = 0.5,
  major.line.size = 0.7
)
```

Xeva object.

Arguments

object

batch Batch name or experiment design list. patient.id Patient id from the XevaSet. Default NULL. drug Name of the drug. Default NULL. model.id One or multiple model.id. Default NULL. model.color Color for model.id. Default NULL. control.name Name of the control sample. Maximum time point of the plot. Default NULL will plot complete data. max.time treatment.only Default FALSE. Given full data treatment.only=TRUE will plot data only during treatment. vol.normal Default FALSE. If TRUE, volume will be normalized. impute.value Default TRUE will impute values if missing. concurrent.time Default FALSE. If TRUE, cut the batch data such that control and treatment will end at the same time point. Color for control plots. control.col

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treatment.col	Color for treatment plots.	
title	Title of the plot.	
xlab	Title of the x-axis.	
ylab	Title of the y-axis.	
log.y	Default FALSE. If TRUE, y-axis will be log-transformed.	
SE.plot	Plot type. Default "all" will plot all plots and average curves. Possible values are "all", "none", "errorbar", and "ribbon".	
aspect.ratio	Default 1 will create a plot of equal width and height.	
minor.line.size		
	Line size for minor lines. Default 0.5.	
major.line.size		
	Line size for major lines. Default 0.7.	

Value

A ggplot2 plot with control and treatment batch data.

Examples

print.batchResponse

Print the batch response

Description

Print the batch response

Usage

```
## S3 method for class 'batchResponse' print(x, ...)
```

Arguments

x batchResponse object... Other arguments

Value

prints the batchResponse

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print.modelResponse

Print the model response

Description

Print the model response

Usage

```
## S3 method for class 'modelResponse' print(x, ...)
```

Arguments

x modelResponse object

... Other arguments

Value

prints the modelResponse

print.pdxBatch

Print the pdx batch

Description

Print the pdx batch

Usage

```
## S3 method for class 'pdxBatch' print(x, ...)
```

Arguments

x pdxBatch object... Other arguments

Value

prints pdxBatch

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repdx

Example PDX dataset

Description

A Xeva object containing anonymous PDX data with replicates. Each batch has 5 replicates.

Usage

```
data(repdx)
```

Format

An object of class XevaSet of length 1.

response

compute PDX response

Description

response Computes the drug response of an individual PDX model or batch.

Usage

```
response(
  object,
  model.id = NULL,
  batch = NULL,
  res.measure = c("mRECIST", "slope", "AUC", "angle", "abc", "TGI", "lmm"),
  treatment.only = FALSE,
  max.time = NULL,
  impute.value = TRUE,
  min.time = 10,
  concurrent.time = TRUE,
  vol.normal = FALSE,
  log.volume = FALSE,
  verbose = TRUE
)
```

Arguments

object Xeva object.

model.id model.id for which the durg response is to be computed.

batch. id or experiment design for which the drug response is to be computed.

res.measure Drug response measure. See Details below

treatment.only Default FALSE. If TRUE, give data for non-zero dose periods only (if dose data

are available).

max.time Maximum time for data.

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impute.value Default FALSE. If TRUE, impute the missing values.

min.time Default 10 days. Used for mRECIST computation.

concurrent.time Default FALSE. If TRUE, cut the batch data such that control and treatment will end at same time point.

vol.normal If TRUE it will normalize the volume. Default FALSE.

log.volume If TRUE log of the volume will be used for response calculation. Default FALSE verbose Default TRUE will print information.

Details

At present the following response measures are implemented

- · mRECIST Computes mRECIST for individual PDX models
- slope Computes slope of the fitted individual PDX curves
- AUC Computes area under a PDX curve for individual PDX models
- angle Computes angle between treatment and control PDX curves
- abc Computes area between the treatment and control PDX curves
- TGI Computes tumor growth inhibition using treatment and control PDX curves
- lmm Computes linear mixed model (lmm) statistics for a PDX batch

Value

Returns model or batch drug response object.

Examples

selectModelIds

To select model IDs based on drug name and/or tissue type.

Description

To select model IDs based on drug name and/or tissue type.

```
selectModelIds(object, drug = NULL, drug.match.exact = TRUE, tissue = NULL)
## S4 method for signature 'XevaSet'
selectModelIds(object, drug = NULL, drug.match.exact = TRUE, tissue = NULL)
```

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Arguments

object The XevaSet.
drug Name of the drug.
drug.match.exact

Default TRUE.

tissue Tumor type. Default NULL.

Value

A vector with the matched model.ids.

Examples

```
data(brca)
df = selectModelIds(brca, drug="trastuzumab", drug.match.exact=TRUE, tissue="BRCA")
head(df)
df2 = selectModelIds(brca, drug="trastuzumab", drug.match.exact=FALSE)
head(df2)
```

sensitivity

Get sensitivity for an Xeva object

Description

Given a Xeva object, it will return a data. frame detailing sensitivity information.

Usage

```
sensitivity(object, type = c("model", "batch"), sensitivity.measure = NULL)
```

Arguments

object The Xeva dataset.

type Sensitivity type (either model or batch).

sensitivity.measure

Name of the sensitivity.measure. Default NULL will return all sensitivity

measures.

Value

A data. frame with model or batch ID and sensitivity values.

```
data(brca)
head(sensitivity(brca, type="batch"))
head(sensitivity(brca, type="model"))
```

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setResponse	set PDX response
-------------	------------------

Description

setResponse sets response of all PDXs in an Xeva object.

Usage

```
setResponse(
  object,
  res.measure = c("mRECIST", "slope", "AUC", "angle", "abc", "TGI", "lmm"),
  min.time = 10,
  treatment.only = FALSE,
  max.time = NULL,
  vol.normal = FALSE,
  impute.value = TRUE,
  concurrent.time = TRUE,
  log.volume = FALSE,
  verbose = TRUE
)
```

Arguments

	,	
	object	Xeva object.
	res.measure	Response measure, multiple measures are allowed. See Details below
	min.time	Minimum number of days for mRECIST computation. Default 10 days.
	treatment.only	Default FALSE. If TRUE, give data for non-zero dose periods only (if dose data are available). $$
	max.time	Maximum number of days to consider for analysis. Data by ond this will be discarded. Default $\mbox{\scriptsize NULL}$ takes full data.
	vol.normal	If TRUE it will will normalize the volume. Default FALSE
	impute.value	Default FALSE. If TRUE, impute the missing volume values.
concurrent.time		
		Default FALSE. If TRUE, cut the batch data such that control and treatment will end at same time point.
	log.volume	If TRUE log of the volume will be used for response calculation. Default ${\sf FALSE}$
	verbose	Default TRUE will print information.

Details

At present fellowing response measure are implemented

- mRECIST Computes mRECIST for indivial PDX model
- slope Computes slope of the fitted indivial PDX curve
- AUC Computes area under a PDX curve for indivial PDX model
- angle Computes angle between treatment and control PDX curves
- abc Computes area between the treatment and control PDX curves
- TGI Computes tumor growth inhibition using treatment and control PDX curves
- lmm Computes linear mixed model (lmm) statistics for a PDX batch

26 slope

Value

Returns updated Xeva object.

Examples

```
data(brca)
brca <- setResponse(brca, res.measure = c("mRECIST"), verbose=FALSE)</pre>
```

slope

Computes slope

Description

slope returns the slope for given time and volume data.

Usage

```
slope(time, volume, degree = TRUE)
```

Arguments

 $\mbox{time} \qquad \qquad \mbox{A vector of time}.$

volume A vector of volume.

degree Default TRUE will give angle in degrees and FALSE will return in radians.

Value

Returns the slope and a fit object.

```
time <- c(0, 3, 7, 11, 18, 22, 26, 30, 32, 35)
volume<- c(250.8, 320.4, 402.3, 382.6, 384, 445.9, 460.2, 546.8, 554.3, 617.9)
sl <- slope(time, volume)
par(pty="s")
xylimit <- range(c(time, volume))
plot(time, volume, type = "b", xlim = xylimit, ylim = xylimit)
abline(lm(volume~time))</pre>
```

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subsetXeva

Subset Xeva object.

Description

Subset Xeva object.

Usage

```
subsetXeva(object, ids, id.name, keep.batch = TRUE)
```

Arguments

object The XevaSet object.

ids IDs to be selected for.

id.name Names of the IDs.

keep.batch Default TRUE. If FALSE, remove all other model.ids from the experiemt design

that do not belong to selection.

Value

New Xeva object.

Examples

```
data(brca)
print(brca)
df <- subsetXeva(brca, ids = c("X-1004", "X-1008", "X-1286"), id.name="patient.id", keep.batch=TRUE)
print(df)</pre>
```

 $\verb|summarizeMolecularProfiles| \\$

Summarize molecular profiles

Description

This function serves to get molecular profiles from a XevaSet object.

```
summarizeMolecularProfiles(
  object,
  drug,
  mDataType,
  tissue = NULL,
  sensitivity.measure = NULL,
  unique.model = TRUE,
  batch = NULL
)
```

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Arguments

object The XevaSet.
drug Name of the drug.

mDataType character, where one of the molecular data types is needed.

tissue Default NULL will return all tissue types.

sensitivity.measure

Default NULL will return all sensitivity measures.

unique.model Default TRUE will return only one sequncing ID, in the case where one model ID

maps to several sequencing IDs.

batch Name of the batch. Default NULL.

Details

If a sequencing sample belongs to multiple models, summarizeMolecularProfiles will create a separate column for each model.

• All models without molecular data will be removed from the output ExpressionSet.

Value

An ExpressionSet where sample names are model.id and sensitivity measures will be presented in pData.

Examples

summarizeResponse

Summarize Response of PDXs

Description

This function summarizes the drug response information of PDXs.

```
summarizeResponse(
  object,
  response.measure = "mRECIST",
  model.id = NULL,
  batch.id = NULL,
  group.by = "patient.id",
  summary.stat = c(";", "mean", "median"),
  tissue = NULL
)
```

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Arguments

object	The XevaSet object.		
response.measure			
	character indicating which response measure to use. Use the responseMeasures function to find out what measures are available for each XevaSet.		
model.id	The model.id for which data is required.		
batch.id	A vector of batch names. Default NULL will return all batches.		
group.by	Default patient.id. Dictates how the models should be grouped together. See details below.		
summary.stat	Dictates which summary method to use if multiple IDs are found.		
tissue	Name of the tissue. Default NULL		

Details

There can be two types of drug response measure.

- Per model response: One response value for each Model, eg. mRECIST_recomputed for each model.
- Per batch response: One response value for each Batch, eg. angle between treatment and control groups.

For the per model response output, columns will be model.id (or group.by). For the per batch response output, the group.by value can be "batch.name".

Value

A matrix with rows as drug names, column as group.by. Each cell contains response.measure for the pair.

Examples

```
data(brca)
brca.mR <- summarizeResponse(brca, response.measure = "mRECIST", group.by="patient.id")</pre>
```

TGI	tumor growth inhibition (TGI) Computes the tumor growth inhibition
	(TGI) between two time-volume curves

Description

tumor growth inhibition (TGI) Computes the tumor growth inhibition (TGI) between two time-volume curves

```
TGI(contr.volume, treat.volume)
```

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Arguments

Value

Returns batch response object

Examples

```
contr.volume <- c(1.35, 6.57, 13.94, 20.39, 32.2, 39.26, 46.9, 53.91)
treat.volume <- c(0.4, 1.26, 2.59, 3.62, 5.77, 6.67, 7.47, 8.98, 9.29, 9.44)
TGI(contr.volume, treat.volume)
```

waterfall

waterfall plot Creates waterfall plot for a given drug.

Description

waterfall plot Creates waterfall plot for a given drug.

Usage

```
waterfall(
  object,
  res.measure,
  drug = NULL,
  group.by = NULL,
  summary.stat = c(";", "mean", "median"),
  tissue = NULL,
  model.id = NULL,
  model.type = NULL,
  type.color = "#cc4c02",
  legend.name = NULL,
  yname = NULL,
  title = NULL,
  sort = TRUE
)
```

Arguments

object The XevaSet object

res.measure PDX model drug response measure

drug Name of the drug

group.by Group drug response data

summary.stat How to summarize multiple values

tissue Tissue type

model.id Indicates which model.id to plot. Default NULL will plot all models

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model.type Type of model, such as mutated or wild type type.color A list with colors used for each type in the legend

legend.name Name of the legend yname Name for the y-axis title Title of the plot

sort Default TRUE will sort the data

Value

waterfall plot in ggplot2

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