# Package 'DGEobj'

October 12, 2022

Type Package

Title Differential Gene Expression (DGE) Analysis Results Data Object

Version 1.1.2

Description Provides a flexible container to manage and annotate Differential Gene Expression (DGE) analysis results (Smythe et. al (2015) <doi:10.1093/nar/gkv007>). The DGEobj has data slots for row (gene), col (samples), assays (matrix n-rows by m-samples dimensions) and metadata (not keyed to row, col, or assays). A set of accessory functions to deposit, query and retrieve subsets of a data workflow has been provided. Attributes are used to capture metadata such as species and gene model, including reproducibility information such that a 3rd party can access a DGEobj history to see how each data object was created or modified. Since the DGEobj is customizable and extensible it is not limited to RNA-seq analysis types of workflows -- it can accommodate nearly any data analysis workflow that starts from a matrix of assays (rows) by samples (columns).

**Depends** R (>= 3.5.0)

License GPL-3

Language en-US

**Encoding UTF-8** 

Imports assertthat, magrittr, stringr, utils

**Suggests** biomaRt, conflicted, dplyr, edgeR, GenomicRanges, glue, knitr, rmarkdown, testthat

biocViews

RoxygenNote 7.1.2

VignetteBuilder knitr

NeedsCompilation no

**Author** John Thompson [aut],

Connie Brett [aut, cre],

Isaac Neuhaus [aut],

Ryan Thompson [aut]

Maintainer Connie Brett < connie@aggregate-genius.com>

**Repository** CRAN

**Date/Publication** 2022-05-16 07:10:12 UTC

2 DGEobj-package

# **R** topics documented:

	DGEobj-package	2
	addItem	3
	addItems	4
	annotateDGEobj	5
	as.list.DGEobj	$\epsilon$
	baseType	
	baseTypes	
	dim.DGEobj	
	dimnames.DGEobj	
	getAttributes	
	getBaseType	
	getItem	10
	getItems	11
	getType	
	initDGEobj	
	initDGEobjDef	
	inventory	
	newType	
	print.DGEobj	
	resetDGEobj	
	rmItem	
	setAttributes	
	showAttributes	
	showMeta	
	showTypes	
	subset.DGEobj	
	[.DGEobj	
	<u>.                                    </u>	
dex		23

DGEobj-package

DGEobj Package Overview

## **Description**

DGEobj is an S3 data class that provides a flexible container for Differential Gene Expression (DGE) analysis results. The DGEobj class is designed to be extensible allowing definition of new data types as needed. A set of accessory functions to deposit, query and retrieve subsets of a data workflow has been provided. Attributes are used to capture metadata such as species and gene model, including reproducibility information such that a 3rd party can access a DGEobj history to see how each data object was created or modified.

addItem 3

#### **Details**

Operationally, the DGEobj is influenced by the RangedSummarizedExperiment (RSE). The DGEobj has data slots for row (gene), col (samples), assays (anything with n-rows by m-samples dimensions) and metadata (anything that can't be keyed to row, col or assay). The key motivation for creating the DGEobj data structure is that the RSE only allows one data item each in the row and col slots and thus is unsuitable for capturing the plethora of data objects created during a typical DGE workflow. The DGEobj data structure can hold any number of row and col data objects and thus is suitable for capturing the multiple steps of a downstream analysis.

Certain object types, primarily the count matrix and associated row and column info, are defined as unique which means only one instance of that type may be added to the DGEobj.

When multiple objects of one type are included in a DGEobj (e.g. two different fits), the concept of parent attributes is used to associate downstream data objects (e.g. contrasts) with the appropriate data object they are derived from.

#### **More Information**

```
browseVignettes(package = 'DGEobj')
```

addItem

Add a data item

#### **Description**

Add a data item

#### Usage

```
addItem(
  dgeObj,
  item,
  itemName,
  itemType,
  funArgs = match.call(),
  itemAttr,
  parent = "",
  overwrite = FALSE,
  init = FALSE
)
```

#### **Arguments**

dgeObj A class DGEobj created by function initDGEobj()
item The data item to be deposited in the DGEobj
itemName The user-assigned name for this data item
itemType The type attribute. See showTypes() to see the

The type attribute. See showTypes() to see the predefined types – types are

extensible with the newType() function.

4 addItems

(optional) A text field to annotate how the data object was created. If the result funArgs

of match.call() is passed as this argument, the name and arguments used in the

current function are captured

itemAttr (optional) A named list of attributes to add directly to the item

parent (optional) itemName of the parent of this item

overwrite Whether to overwrite a matching data object stored in the itemName slot (default

= FALSE)

init Internal Use (default = FALSE)

#### Value

A DGEobj

## **Examples**

```
## Not run:
  myFunArgs <- match.call() # Capture calling function and arguments
  myDGEobj <- addItem(myDGEobj, item</pre>
                                           = MyCounts,
                                  itemName = "counts",
                                  itemType = "counts",
                                  funArgs = myFunArgs)
## End(Not run)
```

addItems

Add multiple data items

## Description

Add multiple data items

## Usage

```
addItems(dgeObj, itemList, itemTypes, parents, itemAttr, overwrite = FALSE)
```

## **Arguments**

dgeObj	A class DGEobj created by function initDGEobj()
itemList	A named list of data items to add to DGEobj
itemTypes	A list of type values for each item on itemList
parents	(optional) A list of parent values for each item on it

n itemList (optional, but highly

recommended)

(optional) An named list of attributes to add to each item. These attributes will itemAttr

be attached to all items in the call.

Whether to overwrite a matching data object stored in the itemName slot (default overwrite

= FALSE)

annotateDGEobj 5

#### Value

```
A DGEobj
```

#### **Examples**

annotateDGEobj

Add annotations

### Description

Reads an annotation file containing key/value pairs or a named list and attaches them attributes to a DGEobj. If a file is used, it should be a text file containing key/value pairs separated by an equals sign. The keys argument specifies which keys we want to capture as attributes on the DGEobj.

## Usage

```
annotateDGEobj(dgeObj, annotations, keys = NULL)
```

#### **Arguments**

dge0bj A object of class DGEobj created by function initDGEobj()

annotations Either a character string path to a file with annotations given as key/value pairs

separated by an equal sign, or a named list of key/value pairs

keys By default (value = NULL), all keys are read in and applied as DGEobj at-

tributes. Use the keys argument to specify a specific list of keys to read from the

file.

#### Value

A DGEobj

as.list.DGEobj

#### **Examples**

as.list.DGEobj

Cast as a simple list

## **Description**

Cast as a simple list

## Usage

```
## S3 method for class 'DGEobj'
as.list(x, ...)
```

#### **Arguments**

x A DGEobj

... Additional parameters

## Value

A named list

```
# example DGEobj
exObj <- readRDS(system.file("miniObj.RDS", package = "DGEobj"))
mylist <- as.list(exObj)</pre>
```

baseType 7

 ${\sf baseType}$ 

Get the baseType of an internal data item

## Description

Get the baseType of an internal data item

## Usage

```
baseType(dgeObj, type)
```

## Arguments

dge0bj A class DGEobj created by function initDGEobj()
type An item type for which you want the baseType

#### Value

character string

## **Examples**

```
# example DGEobj
exObj <- readRDS(system.file("miniObj.RDS", package = "DGEobj"))
baseType(exObj, type = "DGEList")</pre>
```

baseTypes

Get a list of the available baseTypes

## Description

Get a list of the available baseTypes

## Usage

```
baseTypes(dgeObj)
```

## Arguments

dge0bj (optional) A class DGEobj created by function initDGEobj()

## Value

A character vector of baseTypes

8 dimnames.DGEobj

## **Examples**

```
# Global definition of baseTypes
baseTypes()

# example DGEobj
exObj <- readRDS(system.file("miniObj.RDS", package = "DGEobj"))

# Basetypes from a specific DGEobj
baseTypes(exObj)</pre>
```

dim.DGEobj

Get the "assay" dimensions (row/genes by col/samples)

## **Description**

Returns the dimensions of the assay data (baseType)

#### Usage

```
## S3 method for class 'DGEobj'
dim(x)
```

## **Arguments**

Х

A class DGEobj created by function initDGEobj()

#### Value

An integer vector [r,c] with a length of 2.

dimnames.DGEobj

Get the "assay" names (row/genes by col/samples)

## Description

Returns a list of length 2 containing the the assay data names (baseType)

## Usage

```
## S3 method for class 'DGEobj'
dimnames(x)
```

#### **Arguments**

Х

A class DGEobj created by function initDGEobj()

getAttributes 9

## Value

A list of length 2 containing rownames and colnames of the DGEobj

getAttributes

Get all attributes

## **Description**

Get all user-defined attributes from a DGEobj except for any listed in the excludeList argument.

## Usage

```
getAttributes(
  dgeObj,
  excludeList = list("dim", "dimnames", "names", "row.names", "class")
)
```

## Arguments

dgeObj A DGEobj

excludeList A list of attribute names to exclude from the output (default = list("dim", "dim-

names", "names", "row.names"))

## Value

A named list

```
# example DGEobj
exObj <- readRDS(system.file("miniObj.RDS", package = "DGEobj"))
getAttributes(exObj)
# Get the formula attribute from the design (if set)
attr(exObj$design, "formula")</pre>
```

10 getItem

getBaseType

Retrieve data items by baseType

## Description

Retrieve data items by baseType

## Usage

```
getBaseType(dgeObj, baseType)
```

## Arguments

dgeObj A class DGEobj created by function initDGEobj()
baseType One or more of: ["row", "col", "assay", "meta"]

#### Value

A list of data items

## **Examples**

```
# example DGEobj
exObj <- readRDS(system.file("miniObj.RDS", package = "DGEobj"))
Assays <- getBaseType(exObj, baseType = "assay")
AssaysAndMeta <- getBaseType(exObj, c("assay", "meta"))</pre>
```

getItem

Retrieve a data item by name

## Description

Retrieve a data item by name

## Usage

```
getItem(dgeObj, itemName)
```

## **Arguments**

dge0bj A class DGEobj created by function initDGEobj()

itemName Name of item to retrieve

getItems 11

## Value

The requested data item

## **Examples**

```
# example DGEobj
exObj <- readRDS(system.file("miniObj.RDS", package = "DGEobj"))
MyCounts <- getItem(exObj, "counts")</pre>
```

getItems

Retrieve multiple data items by name

## Description

Retrieve multiple data items by name

## Usage

```
getItems(dgeObj, itemNames)
```

## Arguments

dge0bj A class DGEobj created by function initDGEobj()

itemNames A character string, character vector, or list names to retrieve

#### Value

A list

```
# example DGEobj
exObj <- readRDS(system.file("miniObj.RDS", package = "DGEobj"))
myList <- getItems(exObj, list("counts", "geneData"))
names(myList)</pre>
```

12 initDGEobj

	_
get	ype
geti	ype

Retrieve data items by type

## **Description**

Retrieve data items by type

## Usage

```
getType(dgeObj, type, parent)
```

## **Arguments**

dge0bj A class DGEobj created by function initDGEobj()

type A type or list of types to retrieve

parent (optional) Filter return list for common parent (e.g. useful to select one set of

contrast results when multiple fits have been performed)

## Value

A list of data items

## **Examples**

```
# example DGEobj
exObj <- readRDS(system.file("miniObj.RDS", package = "DGEobj"))

MyRawData <- getType(exObj, type = list("counts", "design", "geneData"))</pre>
```

 $\verb"initDGE" obj$ 

Initialize with base data (primaryAssayData, row annotations, col annotations)

## **Description**

Initialize with base data (primary Assay Data, row annotations, col annotations)

initDGEobj 13

#### Usage

```
initDGEobj(
  primaryAssayData,
  rowData,
  colData,
  level,
  customAttr,
  allowShortSampleIDs = FALSE,
  DGEobjDef = initDGEobjDef()
)
```

#### **Arguments**

primaryAssayData

A numeric matrix or dataframe with row and colnames. Each column represents a sample. Each row represents and assay. This is typically the counts matrix in

a DGE RNA-Seq experiment.

rowData Gene, exon, isoform or protein level annotation. Rownames must match the

rownames in primaryAssayData

colData A dataframe describing the experiment design. Rownames much match the col-

names(primaryAssayData)

level One of "gene", "exon", "isoform" or "protein"

customAttr (optional) Named list of attributes

 $\verb|allowShortSampleIDs||$ 

Using sequential integer rownames (even if typed as character) is discouraged and by default will abort the DGEobj creation. If you have a legitimate need to have short sample names composed of numeric characters, you can set this

argument to TRUE (default = FALSE)

DGEobjDef An object definition. Defaults to the global DGEobj definition (initDGEob-

jDef()) and you usually shouldn't change this unless you're customizing the

object for new data types.

#### Value

A DGEobj

14 initDGEobjDef

GeneModel = "Ensembl.R89"))

initDGEobjDef

Instantiate a class DGEobjDef object.

## **Description**

Instantiate a class DGEobjDef object.

## Usage

```
initDGEobjDef(levels, primaryAssayNames, types, uniqueTypes)
```

## Arguments

levels A character string or vector providing names for new levels

primaryAssayNames

A character string or vector, must be the same length as levels This argument

supplies the primary Assay Names for the corresponding levels.

types A named character vector of new types where the values indicate the basetype

for each named type (optional)

uniqueTypes A name or vector of names to add to the uniqueType list (optional)

#### Value

A class DGEobjDef object suitable for use with initDGEobj

inventory 15

inventory	Retrieve the object inventory
Invented y	Retrieve the object thivehior,

## Description

Retrieve the object inventory

#### Usage

```
inventory(dgeObj, verbose = FALSE)
```

## Arguments

dgeObj A class DGEobj created by function initDGEobj()

verbose Include funArgs column in the output (default = FALSE)

## Value

A data.frame summarizing the data contained in the DGEobj

## **Examples**

```
# example DGEobj
exObj <- readRDS(system.file("miniObj.RDS", package = "DGEobj"))
inventory(exObj)</pre>
```

newType

Add a new type definition to a DGEobj

#### **Description**

Add a new type definition to a DGEobj

#### Usage

```
newType(dgeObj, itemType, baseType, uniqueItem = FALSE)
```

## Arguments

dge0bj A class DGEobj created by function initDGEobj()

itemType The name of the new type to create

baseType The baseType of the new item. One of [row, col, assay, meta]

uniqueItem If set to TRUE, only one instance of the new type is allowed in a DGEobj

16 print.DGEobj

## Value

A DGEobj

## **Examples**

print.DGEobj

Print the Inventory

## Description

Print the Inventory

## Usage

```
## S3 method for class 'DGEobj'
print(x, ..., verbose = FALSE)
```

## Arguments

x A class DGEobj created by function initDGEobj()

... Additional parameters

verbose Include funArgs column in the output (default = FALSE)

```
# example DGEobj
exObj <- readRDS(system.file("miniObj.RDS", package = "DGEobj"))
print(exObj)</pre>
```

resetDGEobj 17

resetDGEobj

Reset to original data

## **Description**

During a workflow, a DGEobj typically gets filtered down to remove samples that fail QC or non-expressed genes. The resetDGEobj() function produces a new DGEobj with the original unfiltered data. Resetting an object does not restore changes to attributes or class, but does revert changes made with addItem() and rmItem(). Reset requires that \*\_orig data is still in the DGEobj.

### Usage

```
resetDGEobj(dgeObj)
```

### **Arguments**

dgeObj

A class DGEobj created by function initDGEobj()

## Value

A DGEobj

## **Examples**

```
#example object
exObj <- readRDS(system.file("miniObj.RDS", package = "DGEobj"))
# subset to first 10 rows to show reset functionality
exObj <- exObj[c(1:10), ]

exObj <- resetDGEobj(exObj)
dim(exObj)</pre>
```

rmItem

Removes a named data item

## **Description**

Removes a named data item

## Usage

```
rmItem(dgeObj, itemName)
```

18 setAttributes

## **Arguments**

dge0bj A class DGEobj created by function initDGEobj()

itemName Name of the item to remove

## Value

A DGEobj

## **Examples**

```
# example DGEobj
exObj <- readRDS(system.file("miniObj.RDS", package = "DGEobj"))
exObj <- rmItem(exObj, "design")</pre>
```

setAttributes

Set attributes

## Description

Set one or more attributes on a DGEobj or on a specific item within a DGEobj.

## Usage

```
setAttributes(dgeObj, attribs)
```

## **Arguments**

dgeObj A DGEobj

attribs A named list of attribute/value pairs

#### **Details**

This function adds attributes without deleting the attributes that are already present. Any named attribute that already exists in the object will be updated. To remove an attribute from an object pass NULL as the attribute value.

#### Value

A DGEobj

showAttributes 19

#### **Examples**

```
# example DGEobj
exObj <- readRDS(system.file("miniObj.RDS", package = "DGEobj"))</pre>
# Assign attributes to a DGEobj
MyAttributes <- list(Platform</pre>
                                    = "RNA-Seq",
                     Instrument = "HiSeq",
                     Vendor
                                   = "Unknown",
                     readType
                                   = "PE",
                     readLength = 75,
                     strandSpecific = TRUE)
exObj <- setAttributes(exObj, MyAttributes)</pre>
# Set attributes on an item inside a DGEobj
MyAttributes <- list(normalized = FALSE,</pre>
                     LowIntFilter = "FPK >5 in >= 1 group")
exObj[["counts"]] <- setAttributes(exObj[["counts"]], MyAttributes)</pre>
```

showAttributes

Print attributes

## Description

This function prints all attributes regardless of the class of the attribute value.

#### Usage

```
showAttributes(
  dgeObj,
  skipList = c("dim", "dimnames", "rownames", "colnames", "listData", "objDef")
)
```

## Arguments

dgeObj A DGEobj

skipList A character vector of attributes to skip. Use this to avoid printing certain lengthy

attributes like rownames. Defaults to c("dim", "dimnames", "rownames", "col-

names", "listData", "objDef")

#### **Details**

\*Note\* Use showMeta() to only retrieve attributes that are key/value pairs.

20 showTypes

## **Examples**

```
# example DGEobj
exObj <- readRDS(system.file("miniObj.RDS", package = "DGEobj"))
showAttributes(exObj)</pre>
```

showMeta

Retrieve the Key/Value metadata attributes that have a character value and length of 1

## **Description**

Retrieve the Key/Value metadata attributes that have a character value and length of 1

## Usage

```
showMeta(dgeObj)
```

## Arguments

dgeObj

A DGEobj with attributes

## Value

A data.frame with "Attribute" and "Value" columns

## **Examples**

```
# example DGEobj
exObj <- readRDS(system.file("miniObj.RDS", package = "DGEobj"))
showMeta(exObj)</pre>
```

showTypes

Returns and prints the list of all defined types

## **Description**

Returns and prints the list of all defined types

## Usage

```
showTypes(dgeObj)
```

subset.DGEobj 21

## **Arguments**

dge0bj

A class DGEobj created by function initDGEobj()

#### Value

data.frame

## **Examples**

```
# example DGEobj
exObj <- readRDS(system.file("miniObj.RDS", package = "DGEobj"))</pre>
showTypes(ex0bj)
```

subset.DGEobj

Subset internal row or column data

#### **Description**

Subset internal row or column data

#### Usage

```
## S3 method for class 'DGEobj'
subset(x, ..., row, col, drop = FALSE, debug = FALSE)
```

## **Arguments**

A class DGEobj created by function initDGEobj() Х Additional parameters Row index for the subset row Col index for the subset col Included for compatibility only drop debug

(default = FALSE) Set to TRUE to get additional information on the console if

subsetting a DGEobj fails with a dimension error.

#### Value

A DGEobj

```
# example DGEobj
exObj <- readRDS(system.file("miniObj.RDS", package = "DGEobj"))</pre>
exObj <- subset(exObj, 1:10, 5:50)</pre>
```

22 [.DGEobj

[.DGEobj

Subset with square brackets

# Description

Subset with square brackets

## Usage

```
## S3 method for class 'DGEobj' x[...]
```

## Arguments

x A DGEobj

... Additional parameters

## Value

A DGEobj

# **Index**

```
[.DGEobj, 22
addItem, 3
addItems, 4
annotateDGEobj, 5
as.list.DGEobj, 6
baseType, 7
baseTypes, 7
DGEobj-package, 2
dim.DGEobj, 8
dimnames.DGEobj, 8
getAttributes, 9
{\tt getBaseType},\, {\color{red} 10}
getItem, 10
getItems, 11
getType, 12
initDGEobj, 12
initDGEobjDef, 14
inventory, 15
newType, 15
print.DGEobj, 16
resetDGEobj, 17
rmItem, 17
setAttributes, 18
showAttributes, 19
showMeta, 20
showTypes, 20
{\tt subset.DGEobj}, {\tt 21}
```