# Package 'SeuratObject'

October 24, 2023

```
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     'zzz.R'
     'generics.R'
     'keymixin.R'
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     'default.R'
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     'centroids.R'
     'command.R'
     'compliance.R'
     'data.R'
     'jackstraw.R'
     'dimreduc.R'
     'segmentation.R'
     'molecules.R'
     'spatial.R'
     'fov.R'
     'neighbor.R'
     'seurat.R'
     'sparse.R'
     'utils.R'
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SeuratObject-package SeuratObject: Data Structures for Single Cell Data

#### **Description**

Defines S4 classes for single-cell genomic data and associated information, such as dimensionality reduction embeddings, nearest-neighbor graphs, and spatially-resolved coordinates. Provides data access methods and R-native hooks to ensure the Seurat object is familiar to other R users. See Satija R, Farrell J, Gennert D, et al (2015) doi:10.1038/nbt.3192, Macosko E, Basu A, Satija R, et al (2015) doi:10.1016/j.cell.2015.05.002, and Stuart T, Butler A, et al (2019) doi:10.1016/j.cell.2019.05.031, Hao Y, Hao S, et al (2021) doi:10.1016/j.cell.2021.04.048 and Hao Y, et al (2023) doi:10.1101/2022.02.24.481684 for more details.

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#### See Also

# Useful links:

- https://mojaveazure.github.io/seurat-object/
- https://github.com/mojaveazure/seurat-object
- Report bugs at https://github.com/mojaveazure/seurat-object/issues

.DollarNames.SeuratCommand

Dollar-sign Autocompletion

#### **Description**

Autocompletion for \$ access on a SeuratCommand object

### Usage

```
## S3 method for class 'SeuratCommand'
.DollarNames(x, pattern = "")
```

#### **Arguments**

x A SeuratCommand object

pattern A regular expression. Only matching names are returned.

#### Value

The parameter name matches for pattern

#### See Also

Command log object and interaction methods \$.SeuratCommand(), LogSeuratCommand(), SeuratCommand-class, [.SeuratCommand(), as.list.SeuratCommand()

AddMetaData

Add in metadata associated with either cells or features.

### **Description**

Adds additional data to the object. Can be any piece of information associated with a cell (examples include read depth, alignment rate, experimental batch, or subpopulation identity) or feature (ENSG name, variance). To add cell level information, add to the Seurat object. If adding feature-level metadata, add to the Assay object (e.g. object[["RNA"]])

```
AddMetaData(object, metadata, col.name = NULL)
## S3 method for class 'Assay'
AddMetaData(object, metadata, col.name = NULL)
## S3 method for class 'Assay5'
```

as.Centroids

```
AddMetaData(object, metadata, col.name = NULL)
## S3 method for class 'Seurat'
AddMetaData(object, metadata, col.name = NULL)
```

# Arguments

object An object

metadata A vector, list, or data.frame with metadata to add

col. name A name for meta data if not a named list or data.frame

#### Value

object with metadata added

#### **Examples**

```
cluster_letters <- LETTERS[Idents(object = pbmc_small)]
names(cluster_letters) <- colnames(x = pbmc_small)
pbmc_small <- AddMetaData(
  object = pbmc_small,
  metadata = cluster_letters,
  col.name = 'letter.idents'
)
head(x = pbmc_small[[]])</pre>
```

as.Centroids

Convert Segmentation Layers

### **Description**

Convert Segmentation Layers

```
as.Centroids(x, nsides = NULL, radius = NULL, theta = NULL, ...)
as.Segmentation(x, ...)
## S3 method for class 'Segmentation'
as.Centroids(x, nsides = NULL, radius = NULL, theta = NULL, ...)
## S3 method for class 'Centroids'
as.Segmentation(x, ...)
```

as.Graph 9

### **Arguments**

x An object
nsides The number of sides to represent cells/spots; pass Inf to plot as circles
radius Radius of shapes when plotting
theta Angle to adjust shapes when plotting
... Arguments passed to other methods

#### Value

```
as.Centroids: A Centroids object
as.Segmentation: A Segmentation object
```

as.Graph

Coerce to a Graph Object

### Description

Convert a matrix (or Matrix) to a Graph object

# Usage

```
as.Graph(x, ...)
## S3 method for class 'Matrix'
as.Graph(x, ...)
## S3 method for class 'matrix'
as.Graph(x, ...)
## S3 method for class 'Neighbor'
as.Graph(x, weighted = TRUE, ...)
```

### **Arguments**

x The matrix to convert

... Ignored

weighted If TRUE, fill entries in Graph matrix with value from the nn.dist slot of the

Neighbor object

#### Value

A Graph object

### See Also

Other graph: Graph-class

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#### **Examples**

```
# converting sparse matrix
mat <- Matrix::rsparsematrix(nrow = 10, ncol = 10, density = 0.1)
rownames(x = mat) <- paste0("feature_", 1:10)
colnames(x = mat) <- paste0("cell_", 1:10)
g <- as.Graph(x = mat)

# converting dense matrix
mat <- matrix(data = 1:16, nrow = 4)
rownames(x = mat) <- paste0("feature_", 1:4)
colnames(x = mat) <- paste0("cell_", 1:4)
g <- as.Graph(x = mat)</pre>
```

as.list.SeuratCommand Coerce a SeuratCommand to a list

### Description

Coerce a SeuratCommand to a list

# Usage

```
## S3 method for class 'SeuratCommand'
as.list(x, complete = FALSE, ...)
```

#### **Arguments**

```
    x A SeuratCommand object
    complete Include slots besides just parameters (eg. call string, name, timestamp)
    Ignored
```

#### Value

A list with the parameters and, if complete = TRUE, the call string, name, and timestamp

#### See Also

Command log object and interaction methods \$.SeuratCommand(), .DollarNames.SeuratCommand(), LogSeuratCommand(), SeuratCommand-class, [.SeuratCommand()

### **Examples**

```
cmd <- pbmc_small[["NormalizeData.RNA"]]
as.list(cmd)
as.list(cmd, complete = TRUE)</pre>
```

as.matrix.LogMap

as.matrix.LogMap

Coerce Logical Maps to Matrices

# Description

Coerce a logical map to a matrix; this removes all logical map class capabilities from the object and returns a base-R matrix object

### Usage

```
## S3 method for class 'LogMap'
as.matrix(x, ...)
```

# Arguments

x A LogMap object
... Ignored

### Value

A base-R matrix created from x

### See Also

Logical map objects, validity, and interaction methods: LogMap-validity, LogMap, droplevels.LogMap(), intersect.LogMap(), labels.LogMap()

### **Examples**

```
map <- LogMap(letters[1:10])
map[['obs']] <- c(1, 3, 7)
mat <- as.matrix(map)
mat
class(mat)</pre>
```

as.Neighbor

Coerce to a Neighbor Object

### **Description**

Convert objects to Neighbor objects

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### Usage

```
as.Neighbor(x, ...)
## S3 method for class 'Graph'
as.Neighbor(x, ...)
```

# Arguments

x An object to convert to Neighbor

... Arguments passed to other methods

### Value

A Neighbor object

as.Seurat

Coerce to a Seurat Object

# Description

Convert objects to Seurat objects

# Usage

```
as.Seurat(x, ...)
```

# Arguments

x An object to convert to class Seurat

... Arguments passed to other methods

### Value

A Seurat object generated from x

as.sparse 13

as.sparse

Cast to Sparse

### **Description**

Convert dense objects to sparse representations

# Usage

```
as.sparse(x, ...)
## S3 method for class 'data.frame'
as.sparse(x, row.names = NULL, ...)
## S3 method for class 'Matrix'
as.sparse(x, ...)
## S3 method for class 'matrix'
as.sparse(x, ...)
## S3 method for class 'ngCMatrix'
as.sparse(x, ...)
```

#### **Arguments**

x An object

... Arguments passed to other methods

row.names NULL or a character vector giving the row names for the data; missing values are

not allowed

### Value

A sparse representation of the input data

Assay-class

The Assay Class

### **Description**

The Assay object is the basic unit of Seurat; each Assay stores raw, normalized, and scaled data as well as cluster information, variable features, and any other assay-specific metadata. Assays should contain single cell expression data such as RNA-seq, protein, or imputed expression data.

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### **Slots**

counts Unnormalized data such as raw counts or TPMs
data Normalized expression data
scale.data Scaled expression data
assay.orig Original assay that this assay is based off of. Used to track assay provenance
var.features Vector of features exhibiting high variance across single cells
meta.features Feature-level metadata

misc A named list of unstructured miscellaneous data

key A one-length character vector with the object's key; keys must be one or more alphanumeric characters followed by an underscore "\_" (regex pattern "^[a-zA-Z][a-zA-Z0-9]\*\_\$")

#### See Also

```
v3 Assay object, validity, and interaction methods: $.Assay(), Assay-validity, CreateAssayObject(), [.Assay(), [[.Assay(), dim.Assay(), dimnames.Assay(), merge.Assay(), split.Assay(), subset.Assay()
```

Assay-validity

V3 Assay Validity

#### **Description**

Validation of Assay objects is handled by validObject

#### data Validation

blah

counts Validation

blah

scale.data Validation

blah

#### Feature-Level Meta Data Validation

blah

#### Variable Feature Validation

blah

Assay5-class 15

#### **Key Validation**

Keys must be a one-length character vector; a key must be composed of one of the following:

- An empty string (eg. "''") where nzchar() == 0
- An string composed of one or more alphanumeric values (both lower- and upper-case) that ends with an underscore ("\_"); the first character must be a letter

Keys that are not empty strings are validated with the regex "^[a-zA-Z][a-zA-Z0-9]\*\_\$"

#### See Also

```
validObject
v3 Assay object, validity, and interaction methods: $.Assay(), Assay-class, CreateAssayObject(),
[.Assay(), [[.Assay(), dim.Assay(), dimnames.Assay(), merge.Assay(), split.Assay(),
subset.Assay()
```

#### **Examples**

```
rna <- pbmc_small[["RNA"]]
validObject(rna)</pre>
```

Assay5-class

The v5 Assay Object

### **Description**

The v5 Assay is the typical Assay class used in **Seurat** v5; ...

#### **Slots**

- layers A named list containing expression matrices; each matrix should be a two-dimensional object containing some subset of cells and features defined in the cells and features slots. Cell and feature membership is recorded in the cells and features slots, respectively
- cells A logical mapping of cell names and layer membership; this map contains all the possible cells that this assay can contain. New layers must have some subset of cells present in this map
- features A <u>logical mapping</u> of feature names and layer membership; this map contains all the possible features that this assay can contain. New layers must have some subset of features present in this map
- default A one-length integer with the end index of the default layer; the default layer be all layers up to and including the layer at index default
- assay.orig Original assay that this assay is based off of; used to track assay provenance
- meta.data A data frame with feature-level meta data; should have the same number of rows as features
- misc A named list of unstructured miscellaneous data
- key A one-length character vector with the object's key; keys must be one or more alphanumeric characters followed by an underscore "\_" (regex pattern "^[a-zA-Z][a-zA-Z0-9]\*\_\$")

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#### See Also

```
v5 Assay object, validity, and interaction methods: $.Assay5(), Assay5-validity, [.Assay5(), [[.Assay5(), dim.Assay5(), dimnames.Assay5(), merge.Assay5(), split.Assay5(), subset.Assay5()
```

Assay5-validity

V5 Assay Validity

#### **Description**

Validation of Assay5 objects is handled by validObject

### **Layer Validation**

blah

#### **Key Validation**

Keys must be a one-length character vector; a key must be composed of one of the following:

- An empty string (eg. "''") where nzchar() == 0
- An string composed of one or more alphanumeric values (both lower- and upper-case) that ends with an underscore ("\_"); the first character must be a letter

Keys that are not empty strings are validated with the regex "^[a-zA-Z][a-zA-Z0-9]\*\_\$"

#### See Also

```
validObject
```

```
v5 Assay object, validity, and interaction methods: $.Assay5(), Assay5-class, [.Assay5(), [[.Assay5(), dim.Assay5(), dimnames.Assay5(), merge.Assay5(), split.Assay5(), subset.Assay5()
```

AssayData

Get and Set Assay Data

### **Description**

General accessor and setter functions for Assay objects. GetAssayData can be used to pull information from any of the expression matrices (eg. "counts", "data", or "scale.data"). SetAssayData can be used to replace one of these expression matrices

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### Usage

```
GetAssayData(object, ...)
SetAssayData(object, layer, new.data, slot = deprecated(), ...)
## S3 method for class 'Seurat'
GetAssayData(object, assay = NULL, layer = NULL, slot = deprecated(), ...)
## S3 method for class 'Seurat'
SetAssayData(
  object,
 layer = "data",
 new.data,
  slot = deprecated(),
  assay = NULL,
)
## S3 method for class 'Assay'
GetAssayData(
  object,
  layer = c("data", "scale.data", "counts"),
  slot = deprecated(),
)
## S3 method for class 'Assay'
SetAssayData(
 object,
 layer = c("data", "scale.data", "counts"),
 new.data,
  slot = deprecated(),
)
```

# Arguments

object	An object
	Arguments passed to other methods
layer	Name of layer to get or set
new.data	New assay data to add
slot	[Deprecated] Specific assay data to get or set
assay	Specific assay to get data from or set data for; defaults to the default assay

### Value

GetAssayData: returns the specified assay data

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SetAssayData: object with the assay data set

#### Lifecycle

#### [Superseded]

GetAssayData and SetAssayData have been superseded. To fetch expression matrices, use LayerData; to set expression data, use LayerData<-

### **Examples**

```
# Get assay data from the default assay in a Seurat object
GetAssayData(object = pbmc_small, slot = "data")[1:5,1:5]
# Set an Assay layer through the Seurat object
count.data <- GetAssayData(object = pbmc_small[["RNA"]], layer = "counts")</pre>
count.data <- as.matrix(x = count.data + 1)</pre>
new.seurat.object <- SetAssayData(</pre>
    object = pbmc_small,
    layer = "counts",
    new.data = count.data,
    assay = "RNA"
)
# Get the data directly from an Assay object
GetAssayData(pbmc_small[["RNA"]], layer = "data")[1:5,1:5]
# Set an Assay layer directly
count.data <- GetAssayData(pbmc_small[["RNA"]], layer = "counts")</pre>
count.data \leftarrow as.matrix(x = count.data + 1)
new.assay <- SetAssayData(pbmc_small[["RNA"]], layer = "counts", new.data = count.data)</pre>
```

Assays

Query Specific Object Types

#### **Description**

List the names of Assay, DimReduc, Graph, Neighbor objects

```
Assays(object, ...)

Graphs(object, slot = NULL)

Neighbors(object, slot = NULL)

Reductions(object, slot = NULL)
```

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```
## S3 method for class 'Seurat'
Assays(object, slot = deprecated(), ...)
```

### **Arguments**

object A Seurat object
... Ignored

slot Name of component object to return

#### Value

If slot is NULL, the names of all component objects in this Seurat object. Otherwise, the specific object specified

### **Examples**

```
Assays(pbmc_small)
Graphs(pbmc_small)
Reductions(object = pbmc_small)
```

AttachDeps

Attach Required Packages

### Description

Helper function to attach required packages. Detects if a package is already attached and if so, skips it. Should be called in .onAttach

# Usage

```
AttachDeps(deps)
```

### **Arguments**

deps

A character vector of packages to attach

#### Value

Invisibly returns NULL

### Lifecycle

### [Superseded]

AttachDeps has been superseded as of **SeuratObject** v5.0.0; as an alternative, list dependencies in the Depends section of DESCRIPTION

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#### **Examples**

```
# Use in your .onAttach hook
if (FALSE) {
   .onAttach <- function(libname, pkgname) {
     AttachDeps(c("SeuratObject", "rlang"))
   }
}</pre>
```

Boundaries

Get, Set, and Query Segmentation Boundaries

# Description

Get, Set, and Query Segmentation Boundaries

### Usage

```
Boundaries(object, ...)

DefaultBoundary(object)

DefaultBoundary(object, ...) <- value

Molecules(object, ...)

## S3 method for class 'FOV'
Boundaries(object, ...)

## S3 method for class 'FOV'
DefaultBoundary(object)

## S3 replacement method for class 'FOV'
DefaultBoundary(object, ...) <- value

## S3 method for class 'FOV'
Molecules(object, ...)
```

# Arguments

object An object

... Arguments passed to other methods

value The name of a segmentation boundary to set as default

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### Value

Boundaries: The names of all segmentation boundaries present within object

DefaultBoundary: The name of the default segmentation boundary

DefaultBoundary<-: object with the default segmentation boundary set to value

Molecules: The names of all molecule sets present within object

CastAssay

Cast Assay Layers

# Description

Cast layers in v5 assays to other classes

# Usage

```
CastAssay(object, to, ...)
## S3 method for class 'Assay5'
CastAssay(object, to, layers = NA, verbose = TRUE, ...)
```

### Arguments

object	An object
to	Either a class name or a function that takes a layer and returns the same layer as a new class
• • •	If to is a function, arguments passed to to
layers	A vector of layers to cast; defaults to all layers
verbose	Show progress updates

### Value

object with the layers cast to class specified by to

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Cells

Cell and Feature Names

#### **Description**

Get the cell and feature names of an object

#### Usage

```
Cells(x, ...)
Features(x, ...)
## Default S3 method:
Cells(x, ...)
## S3 method for class 'Assay5'
Cells(x, layer = NULL, simplify = TRUE, ...)
## S3 method for class 'Assay5'
Features(x, layer = NULL, simplify = TRUE, ...)
## S3 method for class 'DimReduc'
Cells(x, ...)
## S3 method for class 'Neighbor'
Cells(x, ...)
```

### **Arguments**

x An object

... Arguments passed to other methods

layer Layer to pull cells/features for; defaults to default layer; if NA, returns all cells

for the assay

simplify Simplify the cell/feature names into a single vector; if FALSE, separates each

cell/feature names by layer

### Value

Cell: A vector of cell names

Features: A vector of feature names

#### See Also

```
dimnames.Assay5(), dimnames.Assay(), dimnames.Seurat()
```

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#### **Examples**

```
Cells(x = pbmc_small)
```

CellsByIdentities

Get cell names grouped by identity class

#### **Description**

Get cell names grouped by identity class

### Usage

```
CellsByIdentities(object, idents = NULL, cells = NULL, return.null = FALSE)
```

### **Arguments**

object A Seurat object

idents A vector of identity class levels to limit resulting list to; defaults to all identity

class levels

cells A vector of cells to grouping to

return.null If no cells are requested, return a NULL; by default, throws an error

#### Value

A named list where names are identity classes and values are vectors of cells belonging to that class

# **Examples**

```
CellsByIdentities(object = pbmc_small)
```

 ${\tt CellsByImage}$ 

Get a vector of cell names associated with an image (or set of images)

# Description

Get a vector of cell names associated with an image (or set of images)

```
CellsByImage(object, images = NULL, unlist = FALSE)
```

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### **Arguments**

object Seurat object

images Vector of image names

unlist Return as a single vector of cell names as opposed to a list, named by image

name.

#### Value

A vector of cell names

#### **Examples**

```
## Not run:
CellsByImage(object = object, images = "slice1")
## End(Not run)
```

Centroids-class

The Centroids Class

# Description

The Centroids Class

### **Slots**

```
cells (character [n]) A vector of cell names; there should be as many cell names as there are points and no duplicate names
```

nsides (integer [1L]) The number of sides to draw when plotting centroids; must be either 0L for circles or greater than 3

radius (numeric [1L]) The radius of the shape when plotting the centroids

theta (numeric [1L]) The angle in degrees to adjust the shape when plotting the centroids

#### See Also

Centroids methods: Centroids-methods

Segmentation layer classes: Centroids-methods, Molecules-class, Molecules-methods, Segmentation-class, Segmentation-methods

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Centroids-methods

Centroids Methods

### **Description**

Methods for Centroids objects

```
## S3 method for class 'Centroids'
Cells(x, ...)
## S3 method for class 'Centroids'
GetTissueCoordinates(object, full = TRUE, ...)
## S3 method for class 'Centroids'
Radius(object)
## S3 method for class 'Centroids'
RenameCells(object, new.names = NULL, ...)
## S3 method for class 'Centroids'
Theta(object)
## S3 method for class 'Centroids'
is.finite(x)
## S3 method for class 'Centroids'
is.infinite(...)
## S3 method for class 'Centroids'
length(x)
## S3 method for class 'Centroids'
lengths(x, use.names = TRUE)
## S3 method for class 'Centroids'
subset(x, cells = NULL, ...)
## S4 method for signature 'Centroids, character, ANY, ANY'
x[i, j, ..., drop = TRUE]
## S4 method for signature 'Centroids, numeric, ANY, ANY'
x[i, j, ..., drop = TRUE]
## S4 method for signature 'Centroids'
show(object)
```

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#### **Arguments**

x, object A Centroids object

... Arguments passed to other methods

full Expand the coordinates to the full polygon

new.names vector of new cell names

use.names Ignored

i, cells A vector of cells to keep; if NULL, defaults to all cells

j, drop Ignored

#### **Details**

GetTissueCoordinates: Get cell spatial coordinates

Radius: Get the centroid radius RenameCells: Update cell names

Theta: Get the offset angle

is.finite, is.infinite: Test to see if the centroids are circular or polygonal

length: Get the number of sides for the polygonal centroid lengths: Generate a run-length encoding of the cells present

subset, [: Subset a Centroids object to certain cells

show: Display an object summary to stdout

#### Value

GetTissueCoordinates: A data frame with three columns:

• "x": the x-coordinate

• "y": the y-coordinate

• "cell": the cell name

If full is TRUE, then each coordinate will indicate a vertex for the cell polygon; otherwise, each coordinate will indicate a centroid for the cell

Radius The radius of the centroids

RenameCells: object with the cells renamed to new.names

Theta: The offset angle in degrees

is.finite: TRUE if the centroids are polygonal, FALSE if circular

is.infinite: The opposite of is.finite

length: 0 if the centroids are circular, otherwise the number of sides of the polygonal centroid

lengths: An rle object for the cells

subset, [: x subsetted to the cells specified by cells/i

show: Invisibly returns NULL

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### See Also

#### Centroids-class

Segmentation layer classes: Centroids-class, Molecules-class, Molecules-methods, Segmentation-class, Segmentation-methods

CheckGC

Conditional Garbage Collection

# Description

Call gc only when desired

### Usage

```
CheckGC(option = "SeuratObject.memsafe")
```

# Arguments

```
option ..
```

#### Value

Invisibly returns NULL

CheckLayersName

Check layers names for the input list

### **Description**

Check layers names for the input list

### Usage

```
CheckLayersName(matrix.list, layers.type = c("counts", "data"))
```

### Arguments

```
matrix.list A list of matrices
```

layers.type layers type, such as counts or data

28 CreateAssay5Object

Command

Get SeuratCommands

### **Description**

Pull information on previously run commands in the Seurat object.

### Usage

```
Command(object, ...)
## S3 method for class 'Seurat'
Command(object, command = NULL, value = NULL, ...)
```

### **Arguments**

object An object

... Arguments passed to other methods

command Name of the command to pull, pass NULL to get the names of all commands run

value Name of the parameter to pull the value for

#### Value

Either a SeuratCommand object or the requested parameter value

CreateAssay50bject

Create a v5 Assay object

#### **Description**

Create an Assay5 object from a feature expression matrix; the expected format of the matrix is features x cells

```
CreateAssay50bject(
  counts = NULL,
  data = NULL,
  min.cells = 0,
  min.features = 0,
  csum = NULL,
  fsum = NULL,
  ...
)
```

CreateAssayObject 29

#### **Arguments**

counts A two-dimensional expression matrix
data Optional prenormalized data matrix

min.cells Include features detected in at least this many cells; will subset the counts matrix

as well. To reintroduce excluded features, create a new object with a lower cutoff

fsum Function for calculating cell sums
Function for calculating feature sums
... Arguments passed to other methods

#### Value

An Assay5 object

Create Assay Object Create an Assay object

#### **Description**

Create an Assay object from a feature (e.g. gene) expression matrix. The expected format of the input matrix is features x cells.

#### Usage

```
CreateAssayObject(
  counts,
  data,
  min.cells = 0,
  min.features = 0,
  key = NULL,
  check.matrix = FALSE,
  ...
)
```

#### **Arguments**

counts Unnormalized data such as raw counts or TPMs
data Prenormalized data; if provided, do not pass counts

min.cells Include features detected in at least this many cells. Will subset the counts

matrix as well. To reintroduce excluded features, create a new object with a

lower cutoff

min. features Include cells where at least this many features are detected

key Optional key to initialize assay with

check.matrix Check counts matrix for NA, NaN, Inf, and non-integer values

... Arguments passed to as. sparse

30 CreateCentroids

#### **Details**

Non-unique cell or feature names are not allowed. Please make unique before calling this function.

#### Value

```
A Assay object
```

#### See Also

```
v3 Assay object, validity, and interaction methods: $.Assay(), Assay-class, Assay-validity, [.Assay(), [[.Assay(), dim.Assay(), dimnames.Assay(), merge.Assay(), split.Assay(), subset.Assay()
```

# **Examples**

```
## Not run:
pbmc_raw <- read.table(
   file = system.file('extdata', 'pbmc_raw.txt', package = 'Seurat'),
   as.is = TRUE
)
pbmc_rna <- CreateAssayObject(counts = pbmc_raw)
pbmc_rna
## End(Not run)</pre>
```

CreateCentroids

Create a Centroids Objects

# Description

Create a Centroids Objects

### Usage

```
CreateCentroids(coords, nsides, radius, theta)
```

### **Arguments**

coords	The coordinates of cell/spot centroids
nsides	The number of sides to represent cells/spots; pass Inf to plot as circles
radius	Radius of shapes when plotting
theta	Angle to adjust shapes when plotting

#### Value

A Centroids object

CreateDimReducObject Create a DimReduc object

# Description

Create a DimReduc object

### Usage

```
CreateDimReducObject(
  embeddings = new(Class = "matrix"),
  loadings = new(Class = "matrix"),
  projected = new(Class = "matrix"),
  assay = NULL,
  stdev = numeric(),
  key = NULL,
  global = FALSE,
  jackstraw = NULL,
  misc = list()
)
```

# Arguments

embeddings	A matrix with the cell embeddings
loadings	A matrix with the feature loadings
projected	A matrix with the projected feature loadings
assay	Assay used to calculate this dimensional reduction
stdev	Standard deviation (if applicable) for the dimensional reduction
key	A character string to facilitate looking up features from a specific DimReduc
global	Specify this as a global reduction (useful for visualizations)
jackstraw	Results from the JackStraw function
misc	list for the user to store any additional information associated with the dimensional reduction

#### Value

A DimReduc object

#### See Also

```
Dimensional reduction object, validity, and interaction methods DimReduc-class, DimReduc-validity, [.DimReduc(), [[.DimReduc(), dim.DimReduc(), merge.DimReduc(), print.DimReduc(), subset.DimReduc()
```

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### **Examples**

```
data <- GetAssayData(pbmc_small[["RNA"]], slot = "scale.data")
pcs <- prcomp(x = data)
pca.dr <- CreateDimReducObject(
  embeddings = pcs$rotation,
  loadings = pcs$x,
  stdev = pcs$sdev,
  key = "PC",
  assay = "RNA"
)</pre>
```

CreateF0V

Create Spatial Coordinates

# Description

Create Spatial Coordinates

```
CreateFOV(coords, ...)
## S3 method for class 'Centroids'
CreateFOV(
  coords,
 molecules = NULL,
  assay = "Spatial",
 key = NULL,
 name = NULL,
  . . .
)
## S3 method for class 'data.frame'
CreateFOV(
  coords,
  type = c("segmentation", "centroids"),
  nsides = Inf,
  radius = NULL,
  theta = 0L,
 molecules = NULL,
  assay = "Spatial",
  key = NULL,
  name = NULL,
)
```

CreateMolecules 33

```
## S3 method for class 'list'
CreateFOV(coords, molecules = NULL, assay = "Spatial", key = NULL, ...)
## S3 method for class 'Segmentation'
CreateFOV(
   coords,
   molecules = NULL,
   assay = "Spatial",
   key = NULL,
   name = NULL,
   ...
)
```

### **Arguments**

coords

type

... Arguments passed to other methods

Molecules

A data.frame with spatially-resolved molecule information or a Molecules object

Assay

Name of associated assay

Key

Key for these spatial coordinates

Name

When coords is a data.frame, Centroids, or Segmentation, name to store coordinates as

When providing a data.frame, specify if the coordinates represent a cell seg-

mentation or voxel centroids

nsides The number of sides to represent cells/spots; pass Inf to plot as circles

radius Radius of shapes when plotting
theta Angle to adjust shapes when plotting

Spatial coordinates

### Value

A FOV object

#### See Also

FOV-class

Create Molecules Create a Molecules Object

### Description

Create a Molecules Object

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#### Usage

```
CreateMolecules(coords, ...)
## S3 method for class 'data.frame'
CreateMolecules(coords, key = "", ...)
## S3 method for class 'Molecules'
CreateMolecules(coords, ...)
## S3 method for class 'NULL''
CreateMolecules(coords, ...)
```

### **Arguments**

coords

Spatial coordinates for molecules; should be a data frame with three columns:

- "x": x-coordinates for each molecule
- "y": y-coordinates for each molecule
- "gene": gene name for each molecule

... Arguments passed to other methods

key A key to set for the molecules

#### Value

A Molecules object

CreateSegmentation

Create a Segmentation Objects

### Description

Create a Segmentation Objects

### Usage

```
CreateSegmentation(coords)
## S3 method for class 'data.frame'
CreateSegmentation(coords)
## S3 method for class 'Segmentation'
CreateSegmentation(coords)
```

#### **Arguments**

coords

The coordinates of cell segmentations

CreateSeuratObject 35

### Value

A Segmentation object

CreateSeuratObject

Create a Seurat object

### **Description**

Create a Seurat object from raw data

```
CreateSeuratObject(
  counts,
  assay = "RNA",
 names.field = 1,
 names.delim = "_",
 meta.data = NULL,
 project = "CreateSeuratObject",
)
## Default S3 method:
CreateSeuratObject(
  counts,
  assay = "RNA",
 names.field = 1L,
 names.delim = "_",
 meta.data = NULL,
 project = "SeuratProject",
 min.cells = 0,
 min.features = 0,
)
## S3 method for class 'Assay'
CreateSeuratObject(
  counts,
  assay = "RNA",
  names.field = 1L,
 names.delim = "_",
 meta.data = NULL,
 project = "SeuratProject",
)
```

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```
## S3 method for class 'Assay5'
CreateSeuratObject(
  counts,
  assay = "RNA",
  names.field = 1L,
  names.delim = "_",
  meta.data = NULL,
  project = "SeuratProject",
  ...
)
```

#### **Arguments**

counts

features as rows or an Assay-derived object Name of the initial assay assay names.field For the initial identity class for each cell, choose this field from the cell's name. E.g. If your cells are named as BARCODE\_CLUSTER\_CELLTYPE in the input matrix, set names.field to 3 to set the initial identities to CELLTYPE. For the initial identity class for each cell, choose this delimiter from the cell's names.delim column name. E.g. If your cells are named as BARCODE-CLUSTER-CELLTYPE, set this to "-" to separate the cell name into its component parts for picking the relevant field. Additional cell-level metadata to add to the Seurat object. Should be a data.frame meta.data where the rows are cell names and the columns are additional metadata fields. Row names in the metadata need to match the column names of the counts matrix. project Project name for the Seurat object Arguments passed to other methods Include features detected in at least this many cells. Will subset the counts min.cells

Either a matrix-like object with unnormalized data with cells as columns and

matrix as well. To reintroduce excluded features, create a new object with a

#### Value

A Seurat object

min.features

### Note

In previous versions (<3.0), this function also accepted a parameter to set the expression threshold for a 'detected' feature (gene). This functionality has been removed to simplify the initialization process/assumptions. If you would still like to impose this threshold for your particular dataset, simply filter the input expression matrix before calling this function.

Include cells where at least this many features are detected

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## **Examples**

```
## Not run:
pbmc_raw <- read.table(
   file = system.file('extdata', 'pbmc_raw.txt', package = 'Seurat'),
   as.is = TRUE
)
pbmc_small <- CreateSeuratObject(counts = pbmc_raw)
pbmc_small
## End(Not run)</pre>
```

Crop

Crop Coordinates

# Description

**Crop Coordinates** 

## Usage

```
Crop(object, x = NULL, y = NULL, coords = c("plot", "tissue"), ...)
## S3 method for class 'FOV'
Crop(object, x = NULL, y = NULL, coords = c("plot", "tissue"), ...)
```

# Arguments

object	An object
x, y	Range to crop x/y limits to; if NULL, uses full range of x/y
coords	Coordinate system to execute crop; choose from:
	• "plot": Coordinates as shown when plotting
	• "tissue": Coordinates from GetTissueCoordinates
	Arguments passed to other methods

#### Value

```
object cropped to the region specified by x and y
```

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DefaultAssay

Default Assay

#### **Description**

Get and set the default assay

### Usage

```
DefaultAssay(object, ...)
DefaultAssay(object, ...) <- value</pre>
## S3 method for class 'Graph'
DefaultAssay(object, ...)
## S3 replacement method for class 'Graph'
DefaultAssay(object, ...) <- value</pre>
## S3 method for class 'Assay'
DefaultAssay(object, ...)
## S3 replacement method for class 'Assay'
DefaultAssay(object, ...) <- value</pre>
## S3 method for class 'Assay5'
DefaultAssay(object, ...)
## S3 replacement method for class 'Assay5'
DefaultAssay(object, ...) <- value</pre>
## S3 method for class 'SeuratCommand'
DefaultAssay(object, ...)
## S3 method for class 'DimReduc'
DefaultAssay(object, ...)
## S3 replacement method for class 'DimReduc'
DefaultAssay(object, ...) <- value</pre>
## S3 method for class 'Seurat'
DefaultAssay(object, ...)
## S3 replacement method for class 'Seurat'
DefaultAssay(object, ...) <- value</pre>
```

DefaultDimReduc 39

#### **Arguments**

object An object

... Arguments passed to other methods value Name of assay to set as default

#### Value

 $\label{eq:default} \textbf{DefaultAssay: The name of the default assay}$ 

DefaultAssay<-: An object with the default assay updated

## **Examples**

```
# Get current default assay
DefaultAssay(object = pbmc_small)

# Create dummy new assay to demo switching default assays
new.assay <- pbmc_small[["RNA"]]
Key(object = new.assay) <- "RNA2_"
pbmc_small[["RNA2"]] <- new.assay
# switch default assay to RNA2
DefaultAssay(object = pbmc_small) <- "RNA2"
DefaultAssay(object = pbmc_small)</pre>
```

DefaultDimReduc

Find the default DimReduc

#### **Description**

Searches for DimReducs matching "umap", "tsne", or "pca", case-insensitive, and in that order. Priority given to DimReducs matching the DefaultAssay or assay specified (eg. "pca" for the default assay weights higher than "umap" for a non-default assay)

## Usage

```
DefaultDimReduc(object, assay = NULL)
```

## **Arguments**

object A Seurat object

assay Name of assay to use; defaults to the default assay of the object

#### Value

The default DimReduc, if possible

DefaultFOV

# **Examples**

```
DefaultDimReduc(pbmc_small)
```

DefaultF0V

Get and Set the Default FOV

# Description

Get and Set the Default FOV

# Usage

```
DefaultFOV(object, ...)
DefaultFOV(object, ...) <- value

## S3 method for class 'Seurat'
DefaultFOV(object, assay = NULL, ...)

## S3 replacement method for class 'Seurat'
DefaultFOV(object, assay = NA, ...) <- value</pre>
```

# Arguments

object	A Seurat Object
	Arguments passed to other methods
value	The name of the FOV to set as the default
assay	Name of assay to get or set default FOV for; pass NA to get or set the global default ${\sf FOV}$

## Value

```
DefaultFOV: The name of the default FOV

DefaultFOV<-: object with the default FOV set to value
```

DefaultLayer 41

DefaultLayer

Default Layer

## **Description**

Get and set the default layer

## Usage

```
DefaultLayer(object, ...)
DefaultLayer(object, ...) <- value

## S3 method for class 'Assay'
DefaultLayer(object, ...)

## S3 method for class 'Assay5'
DefaultLayer(object, ...)

## S3 replacement method for class 'Assay5'
DefaultLayer(object, ...) <- value</pre>
```

## **Arguments**

object An object

... Arguments passed to other methods

value Name of layer to set as default

#### Value

DefaultLayer: The name of the default layer

DefaultLayer<-: An object with the default layer updated

dim.Assay

Feature and Cell Numbers

## **Description**

Feature and Cell Numbers

#### Usage

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

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### **Arguments**

Х

An Assay object

## Value

A two-length numeric vector with the total number of features and cells in x

#### See Also

```
v3 Assay object, validity, and interaction methods: $.Assay(), Assay-class, Assay-validity,
CreateAssayObject(), [.Assay(), [[.Assay(), dimnames.Assay(), merge.Assay(), split.Assay(),
subset.Assay()
```

# **Examples**

```
rna <- pbmc_small[["RNA"]]
dim(rna)</pre>
```

dim.Assay5

Feature and Cell Numbers

# Description

Feature and Cell Numbers

# Usage

```
## S3 method for class 'Assay5'
dim(x)
```

## Arguments

Х

An Assay5 object

### Value

A two-length numeric vector with the total number of features and cells in x

#### See Also

```
v5 Assay object, validity, and interaction methods: $.Assay5(), Assay5-class, Assay5-validity, [.Assay5(), [[.Assay5(), dimnames.Assay5(), merge.Assay5(), split.Assay5(), subset.Assay5()
```

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dim.DimReduc

Dimensional Reduction Meta-Information

## **Description**

Pull meta-information about cells and dimensions for a given dimensional reduction; cell meta-information is stored as row meta-information (eg. nrow, rownames) and dimension meta-information is stored as column meta-information (eg. ncol, colnames)

#### Usage

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

## S3 method for class 'DimReduc'
dimnames(x)

## S3 method for class 'DimReduc'
length(x)

## S3 method for class 'DimReduc'
names(x)
```

## Arguments

Х

A DimReduc object

#### Value

```
dim: The number of cells (nrow) and dimensions (ncol) dimnames: The cell (row) and dimension (column) names length: The number of dimensions names: The dimension identifiers
```

#### See Also

Cells

```
Dimensional reduction object, validity, and interaction methods CreateDimReducObject(), DimReduc-class, DimReduc-validity, [.DimReduc(), [[.DimReduc(), merge.DimReduc(), print.DimReduc(), subset.DimReduc()
```

```
pca <- pbmc_small[["pca"]]
pca
dim(pca)</pre>
```

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```
# nrow is number of cells
nrow(pca)

# rownames pulls cell names
head(rownames(pca))

# ncol and length are number of dimensions
ncol(pca)
length(pca)

# colnames and names pull dimension identifiers
head(colnames(pca))
head(names(pca))
```

dim.Seurat

Feature and Cell Numbers

## **Description**

Feature and Cell Numbers

## Usage

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

### **Arguments**

Χ

A Seurat object

#### Value

A two-length numeric vector with the total number of features and cells in x

#### See Also

```
Seurat object, validity, and interaction methods $.Seurat(), Seurat-class, Seurat-validity,
[[.Seurat(), [[<-,Seurat, NULL, [[<-,Seurat, dimnames.Seurat(), merge.Seurat(), names.Seurat(),
subset.Seurat()</pre>
```

```
# Get the number of features in an object
nrow(pbmc_small)
# Get the number of cells in an object
ncol(pbmc_small)
```

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dimnames.Assay

Assay-Level Feature and Cell Names

#### **Description**

Get and set feature and cell names in v5 Assays

#### Usage

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

## S3 replacement method for class 'Assay'
dimnames(x) <- value</pre>
```

### **Arguments**

x An Assay object

value A two-length list where the first entry is the existing feature names for x and the

second entry is the *updated* cell names for x

#### Value

dimnames: A two-length list with the following values:

- A character vector will all features in x
- A character vector will all cells in x

dimnames<-: x with the cell names updated to those in value[[2L]]

#### See Also

```
v3 Assay object, validity, and interaction methods: $.Assay(), Assay-class, Assay-validity,
CreateAssayObject(), [.Assay(), [[.Assay(), dim.Assay(), merge.Assay(), split.Assay(),
subset.Assay()
Cells(), dimnames.Assay5(), dimnames.Seurat()
```

```
rna <- pbmc_small[["RNA"]]
# Feature and cell names can be acquired with `rownames` and `colnames`
head(rownames(rna))
head(colnames(rna))
# Cell names can be updated with `colnames<-`
colnames(rna)[1] <- "newcell"
head(colnames(rna))</pre>
```

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dimnames.Assay5

Assay-Level Feature and Cell Names

### **Description**

Get and set feature and cell names in v5 Assays

#### Usage

```
## S3 method for class 'Assay5'
dimnames(x)
## S3 replacement method for class 'Assay5'
dimnames(x) <- value</pre>
```

#### **Arguments**

x An Assay5 object

value A two-length list with updated feature and/or cells names

#### Value

dimnames: A two-length list with the following values:

- A character vector with all features in x
- A character vector with all cells in x

dimnames<-: x with the feature and/or cell names updated to value

#### See Also

```
v5 Assay object, validity, and interaction methods: $.Assay5(), Assay5-class, Assay5-validity, [.Assay5(), [[.Assay5(), dim.Assay5(), merge.Assay5(), split.Assay5(), subset.Assay5() Cells(), dimnames.Assay(), dimnames.Seurat()
```

dimnames.Seurat

Feature and Cell Names

### **Description**

Get and set feature and cell inames in Seurat objects

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#### Usage

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

## S3 replacement method for class 'Seurat'
dimnames(x) <- value</pre>
```

### **Arguments**

x A Seurat object

value A two-length list with updated feature and/or cells names

#### Value

dimnames: A two-length list with the following values:

- A character vector with all features in the default assay
- A character vector with all cells in x

dimnames<-: x with the feature and/or cell names updated to value

#### See Also

```
Seurat object, validity, and interaction methods $.Seurat(), Seurat-class, Seurat-validity,
[[.Seurat(), [[<-, Seurat, NULL, [[<-, Seurat, dim.Seurat(), merge.Seurat(), names.Seurat(),
subset.Seurat()
Cells(), dimnames.Assay5(), dimnames.Assay()</pre>
```

# **Examples**

```
# Get the feature names of an object
head(rownames(pbmc_small))

# Get the cell names of an object
head(colnames(pbmc_small))

colnames(pbmc_small)[1] <- "newcell"
head(colnames(pbmc_small))</pre>
```

DimReduc-class

The Dimensional Reduction Class

#### **Description**

The DimReduc object stores a dimensionality reduction taken out in Seurat; each DimReduc consists of a cell embeddings matrix, a feature loadings matrix, and a projected feature loadings matrix.

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#### **Slots**

```
cell.embeddings Cell embeddings matrix (required)

feature.loadings Feature loadings matrix (optional)

feature.loadings.projected Projected feature loadings matrix (optional)

assay.used Name of assay used to generate DimReduc object

global Is this DimReduc global/persistent? If so, it will not be removed when removing its associated assay

stdev A vector of standard deviations

jackstraw A JackStrawData-class object associated with this DimReduc

misc A named list of unstructured miscellaneous data

key A one-length character vector with the object's key; keys must be one or more alphanumeric
```

characters followed by an underscore "\_" (regex pattern "^[a-zA-Z][a-zA-Z0-9]\*\_\$")

#### See Also

Dimensional reduction object, validity, and interaction methods CreateDimReducObject(), DimReduc-validity, [.DimReduc(), [[.DimReduc(), dim.DimReduc(), merge.DimReduc(), print.DimReduc(), subset.DimReduc()

DimReduc-validity

Dimensional Reduction Validity

## Description

Validation of DimReduc objects is handled by validObject

#### **Cell Embeddings Validation**

The cell embeddings matrix must be a numeric matrix of dimensions  $n_{cells}$  by  $d_{dimensions}$ ; row names must be the cell names and column names must be the dimension identifier. The dimension identifier must be "key\_dimension" (eg. "PC\_1"). Dimension identifiers must be in order and cannot be skipped

# **Feature and Projected Feature Loadings Validation**

blah

#### **Standard Deviations Validation**

blah

Distances 49

#### **Key Validation**

Keys must be a one-length character vector; a key must be composed of one of the following:

- An empty string (eg. "''") where nzchar() == 0
- An string composed of one or more alphanumeric values (both lower- and upper-case) that ends with an underscore ("\_"); the first character must be a letter

Keys that are not empty strings are validated with the regex "^[a-zA-Z][a-zA-Z0-9]\*\_\$"

#### See Also

```
Dimensional reduction object, validity, and interaction methods CreateDimReducObject(), DimReduc-class, [.DimReduc(), [[.DimReduc(), dim.DimReduc(), merge.DimReduc(), print.DimReduc(), subset.DimReduc()
```

Distances

Get the Neighbor nearest neighbors distance matrix

# Description

Get the Neighbor nearest neighbors distance matrix

## Usage

```
Distances(object, ...)
## S3 method for class 'Neighbor'
Distances(object, ...)
```

#### **Arguments**

object An object

.. Arguments passed to other methods

#### Value

The distance matrix

50 droplevels.LogMap

droplevels.LogMap

Drop Unused Logical Map Values

# Description

Remove any unused values from a logical map

# Usage

```
## S3 method for class 'LogMap'
droplevels(x, ...)
```

## **Arguments**

```
x A LogMap object
... Ignored
```

### Value

x with values not present in any observation removed

### See Also

```
Logical\ map\ objects,\ validity,\ and\ interaction\ methods:\ LogMap-validity,\ LogMap,\ as.\ matrix.\ LogMap(),\ intersect.\ LogMap(),\ labels.\ LogMap()
```

```
map <- LogMap(letters[1:10])
map[['obs']] <- c(1, 3, 7)
map[['entry']] <- c(2, 7, 10)

# Remove unused values
map <- droplevels(map)
map
map[[]]</pre>
```

Embeddings 51

Embeddings

Get Cell Embeddings

# Description

Get Cell Embeddings

# Usage

```
Embeddings(object, ...)
## S3 method for class 'DimReduc'
Embeddings(object, ...)
## S3 method for class 'Seurat'
Embeddings(object, reduction = "pca", ...)
```

## **Arguments**

object An object

... Arguments passed to other methods

reduction Name of reduction to pull cell embeddings for

### Value

The embeddings matrix

## **Examples**

```
# Get the embeddings directly from a DimReduc object
Embeddings(object = pbmc_small[["pca"]])[1:5, 1:5]
# Get the embeddings from a specific DimReduc in a Seurat object
Embeddings(object = pbmc_small, reduction = "pca")[1:5, 1:5]
```

FetchData

Access cellular data

## **Description**

Retrieves data (feature expression, PCA scores, metrics, etc.) for a set of cells in a Seurat object

52 FetchData

## Usage

```
FetchData(object, ...)
## S3 method for class 'DimReduc'
FetchData(object, vars, cells = NULL, ...)
## S3 method for class 'Seurat'
FetchData(
   object,
   vars,
   cells = NULL,
   layer = NULL,
   clean = TRUE,
   slot = deprecated(),
   ...
)
```

# Arguments

object	An object
	Arguments passed to other methods
vars	List of all variables to fetch, use keyword "ident" to pull identity classes
cells	Cells to collect data for (default is all cells)
layer	Layer to pull feature data for
clean	Remove cells that are missing data; choose from:
	• "all": consider all columns for cleaning
	<ul> <li>"ident": consider all columns except the identity class for cleaning</li> </ul>
	• "project": consider all columns except the identity class for cleaning; fill missing identity values with the object's project
	• "none": do not clean
	Passing TRUE is a shortcut for "ident"; passing FALSE is a shortcut for "none"
slot	Deprecated in favor of layer

#### Value

A data frame with cells as rows and cellular data as columns

```
pc1 <- FetchData(object = pbmc_small, vars = 'PC_1')
head(x = pc1)
head(x = FetchData(object = pbmc_small, vars = c('groups', 'ident')))</pre>
```

FilterObjects 53

FilterObjects

Find Sub-objects of a Certain Class

#### **Description**

Get the names of objects within a Seurat object that are of a certain class

### Usage

```
FilterObjects(object, classes.keep = c("Assay", "StdAssay", "DimReduc"))
```

### **Arguments**

object A Seurat object

classes.keep A vector of names of classes to get

#### Value

A vector with the names of objects within the Seurat object that are of class classes. keep

### Lifecycle

### [Deprecated]

FilterObjects was deprecated in version 5.0.0; use .FilterObjects instead

#### **Examples**

FilterObjects(pbmc\_small)

FOV-class

The Field of View Object

#### **Description**

A modern container for storing coordinates of spatially-resolved single cells. Capable of storing multiple cell segmentation boundary masks. Supports coordinates for spatially-resolved molecule (FISH) data. Compatible with SpatialImage

#### **Slots**

molecules A named list of Molecules objects defining spatially-resolved molecular coordinates boundaries A named list of Segmentation and Centroids objects defining spatially-resolved boundaries

assay A character naming the associated assay of the spatial coordinates

key A one-length character vector with the object's key; keys must be one or more alphanumeric characters followed by an underscore "\_" (regex pattern "^[a-zA-Z][a-zA-Z0-9]\*\_\$")

54 FOV-methods

#### See Also

FOV-methods

FOV-methods

FOV Methods

## **Description**

Methods for FOV objects

## Usage

```
## S3 method for class 'FOV'
Cells(x, boundary = NULL, ...)
## S3 method for class 'FOV'
Features(x, set = NULL, ...)
## S3 method for class 'FOV'
FetchData(object, vars, cells = NULL, simplify = TRUE, ...)
## S3 method for class 'FOV'
GetTissueCoordinates(object, which = NULL, ...)
## S3 method for class 'FOV'
Keys(object, ...)
## S3 method for class 'FOV'
RenameCells(object, new.names = NULL, ...)
## S3 method for class 'FOV'
x$i, ...
## S3 method for class 'FOV'
x[i, j, ...]
## S3 method for class 'FOV'
x[[i, ...]]
## S3 method for class 'FOV'
length(x)
## S3 method for class 'FOV'
names(x)
## S3 method for class 'FOV'
subset(x, cells = NULL, features = NULL, ...)
```

FOV-methods 55

```
## S4 replacement method for signature 'FOV, character, missing, Centroids'
x[[i, j, ...]] <- value

## S4 replacement method for signature 'FOV, character, missing, Molecules'
x[[i, j, ...]] <- value

## S4 replacement method for signature 'FOV, character, missing, `NULL`'
x[[i, j, ...]] <- value

## S4 replacement method for signature 'FOV, character, missing, Segmentation'
x[[i, j, ...]] <- value

## S4 method for signature 'FOV'
show(object)</pre>
```

## **Arguments**

x, object	A FOV object
boundary, set	Name of segmentation boundary or molecule set to extract cell or feature names for; pass NA to return all cells or feature names
• • •	Arguments passed to other methods
vars	A vector of variables to fetch; can be the name of a segmentation boundary, to get tissue coordinates, or molecule names, to get molecule coordinates
simplify	If only returning either boundary or molecule coordinates, return a single data frame instead of a list
which	Name of segmentation boundary or molecule set
new.names	vector of new cell names
i, cells	For [[ and [[<-, the name of a segmentation or "molecules"; for FetchData, subset. and [, a vector of cells to keep
j, features	For subset and [, a vector of features to keep; for [[<-, not used
value	For [[<-, a replacement Molecules, Centroids, or Segmentation object; otherwise NULL to remove the boundary stored at i

#### Details

The following methods are defined for interacting with a FOV object:

Cells: Get cell names

Features: Get spatially-resolved molecule names

FetchData: Fetch boundary and/or molecule coordinates from a FOV object

GetTissueCoordinates: Get boundary or molecule coordinates from a FOV object

Keys: Get the keys of molecule sets contained within a FOV object

RenameCells: Update cell names

\$, [[: Extract a segmentation boundary

56 FOV-methods

length: Get the number of segmentation layers in a FOV object

names: Get the names of segmentation layers and molecule sets

subset, [: Subset a FOV object

[[<-: Add or remove segmentation layers and molecule information to/from a FOV object

show: Display an object summary to stdout

#### Value

Cells: A vector of cell names

Features: A vector of spatially-resolved molecule names; if no molecular information present, returns NULL

FetchData: If both molecule and boundary coordinates are requested, then a two-length list:

- "molecules": A data frame with the molecule coordinates requested. If molecules requested are keyed, the keys are preserved in the data frame
- "coordinates": A data frame with coordinates from the segmentation boundaries requested

If simplify is TRUE and only one data frame is generated, then only the data frame is returned. Otherwise, a one-length list is returned with the single data frame generated

GetTissueCoordinates: ...

Keys: A named vector of molecule set keys; names are the names of the molecule sets and values are the keys for the respective molecule set

RenameCells: object with the cells renamed to new.names

\$, [[: The segmentation boundary or spatially-resolved molecule information stored at i

length: The number of segmentation layers (Segmentation or Centroids objects)

names: A vector of segmentation boundary and molecule set names

subset: x with just the cells and features specified

[[<-: Varies depending on the class of value:

- If value is NULL, returns x with the boundary i removed; also allows removing molecules; does not allow removing the default segmentation
- If value is a Molecules, returns x with value stored in molecules; requires that i is "molecules"
- Otherwise, stores value as a segmentation boundary named i

show: Invisibly returns NULL

### See Also

FOV-class

FOV-validity 57

FOV-validity

FOV Validity

### **Description**

Validation of FOV objects is handled by validObject

# **Boundary Validation**

blah

## **Molecule Validation**

blah

## See Also

validObject

GetImage

Get image data

## **Description**

Get image data

## Usage

```
GetImage(object, mode = c("grob", "raster", "plotly", "raw"), ...)
## S3 method for class 'Seurat'
GetImage(
  object,
  mode = c("grob", "raster", "plotly", "raw"),
  image = NULL,
  ...
)
```

# Arguments

```
object An object

mode How to return the image; should accept one of "grob", "raster", "plotly", or "raw"

... Arguments passed to other methods

image Name of SpatialImage object to pull image data for; if NULL, will attempt to select an image automatically
```

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## Value

```
Image data, varying depending on the value of mode:
```

```
"grob" An object representing image data inheriting from grob objects (eg. rastergrob)
```

"raster" An object of class raster

"plotly" A list with image data suitable for Plotly rendering, see plotly::layout for more details

"raw" The raw image data as stored in the object

## See Also

layout

GetTissueCoordinates Get tissue coordinates

# Description

Get tissue coordinates

### Usage

```
GetTissueCoordinates(object, ...)
## S3 method for class 'Seurat'
GetTissueCoordinates(object, image = NULL, ...)
```

# Arguments

object An object

... Arguments passed to other methods

image Name of SpatialImage object to get coordinates for; if NULL, will attempt to

select an image automatically

#### Value

A data frame with tissue coordinates

Graph-class 59

Graph-class

The Graph Class

### Description

The Graph class inherits from dgCMatrix. We do this to enable future expandability of graphs.

#### **Slots**

assay.used Optional name of assay used to generate Graph object

#### See Also

```
dgCMatrix-class
Other graph: as.Graph()
```

**HVFInfo** 

Highly Variable Features

## **Description**

Get and set variable feature information for an Assay object. HVFInfo and VariableFeatures utilize generally variable features, while SVFInfo and SpatiallyVariableFeatures are restricted to spatially variable features

#### Usage

```
HVFInfo(object, method, status = FALSE, ...)
VariableFeatures(object, method = NULL, ...)
VariableFeatures(object, ...) <- value
SVFInfo(object, method, status, ...)
SpatiallyVariableFeatures(object, method, ...)
## S3 method for class 'Seurat'
HVFInfo(
   object,
   method = NULL,
   status = FALSE,
   assay = NULL,
   selection.method = deprecated(),</pre>
```

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```
)
## S3 method for class 'Seurat'
VariableFeatures(
  object,
 method = NULL,
  assay = NULL,
 nfeatures = NULL,
  layer = NA,
  simplify = TRUE,
  selection.method = deprecated(),
)
## S3 replacement method for class 'Seurat'
VariableFeatures(object, assay = NULL, ...) <- value</pre>
## S3 method for class 'Seurat'
SVFInfo(
  object,
 method = c("markvariogram", "moransi"),
  status = FALSE,
 assay = NULL,
  selection.method = deprecated(),
)
## S3 method for class 'Seurat'
SpatiallyVariableFeatures(
 object,
 method = "moransi",
  assay = NULL,
  decreasing = TRUE,
  selection.method = deprecated(),
)
## S3 method for class 'Assay'
HVFInfo(object, method, status = FALSE, selection.method = deprecated(), ...)
## S3 method for class 'Assay'
SpatiallyVariableFeatures(
 object,
 method = "moransi",
 decreasing = TRUE,
  selection.method = deprecated(),
)
```

HVFInfo 61

```
## S3 method for class 'Assay'
   SVFInfo(
      object,
     method = c("markvariogram", "moransi"),
     status = FALSE,
      selection.method = deprecated(),
   )
   ## S3 method for class 'Assay'
   VariableFeatures(object, method = NULL, selection.method = deprecated(), ...)
   ## S3 replacement method for class 'Assay'
   VariableFeatures(object, ...) <- value
   ## S3 method for class 'Assay5'
   HVFInfo(object, method = NULL, status = FALSE, layer = NULL, strip = TRUE, ...)
   ## S3 method for class 'Assay5'
   VariableFeatures(
     object,
     method = NULL,
     layer = NA,
      simplify = TRUE,
     nfeatures = Inf,
      selection.method = deprecated(),
   )
   ## S3 replacement method for class 'Assay5'
   VariableFeatures(object, method = "custom", layer = NULL, ...) <- value</pre>
Arguments
   object
                    An object
   method
                    Which method to pull. For HVFInfo and VariableFeatures, choose one from
                    one of the following:
                      • "vst"
                      • "sctransform" or "sct"
                      • "mean.var.plot", "dispersion", "mvp", or "disp"
                    For SVFInfo and Spatially Variable Features, choose from:
                      · "markvariogram"
                      · "moransi"
                    Add variable status to the resulting data frame
   status
                    Arguments passed to other methods
```

62 Idents

value A character vector of variable features

assay Name of assay to pull highly variable feature information for

selection.method

[Deprecated]

nfeatures Maximum number of features to select when simplifying

layer Layer to pull variable features for

simplify When pulling for multiple layers, combine into a single vector and select a com-

mon set of variable features for all layers

decreasing Return features in decreasing order (most spatially variable first).

strip Remove method/layer identifiers from highly variable data frame

#### Value

HVFInfo: A data frame with feature means, dispersion, and scaled dispersion

VariableFeatures: a vector of the variable features

SVFInfo: a data frame with the spatially variable features

SpatiallyVariableFeatures: a character vector of the spatially variable features

## **Examples**

```
# Get the HVF info from a specific Assay in a Seurat object
HVFInfo(object = pbmc_small, assay = "RNA")[1:5, ]
# Get the HVF info directly from an Assay object
HVFInfo(pbmc_small[["RNA"]], method = 'vst')[1:5, ]
```

Idents

Get, set, and manipulate an object's identity classes

### Description

Get, set, and manipulate an object's identity classes

### Usage

```
Idents(object, ...)
Idents(object, ...) <- value
RenameIdents(object, ...)
ReorderIdent(object, var, ...)
SetIdent(object, ...)</pre>
```

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```
StashIdent(object, save.name, ...)
## S3 method for class 'Seurat'
Idents(object, ...)
## S3 replacement method for class 'Seurat'
Idents(object, cells = NULL, drop = FALSE, replace = FALSE, ...) <- value</pre>
## S3 method for class 'Seurat'
ReorderIdent(
  object,
 var,
  reverse = FALSE,
  afxn = mean,
  reorder.numeric = FALSE,
)
## S3 method for class 'Seurat'
RenameIdents(object, ...)
## S3 method for class 'Seurat'
SetIdent(object, cells = NULL, value, ...)
## S3 method for class 'Seurat'
StashIdent(object, save.name = "orig.ident", ...)
## S3 method for class 'Seurat'
droplevels(x, ...)
## S3 method for class 'Seurat'
levels(x)
## S3 replacement method for class 'Seurat'
levels(x) <- value
```

#### **Arguments**

• • •	Arguments passed to other methods; for RenameIdents: named arguments as old.ident = new.ident; for ReorderIdent: arguments passed on to FetchData
value	The name of the identities to pull from object metadata or the identities themselves
var	Feature or variable to order on
save.name	Store current identity information under this name
cells	Set cell identities for specific cells
drop	Drop unused levels

64 Idents

replace Replace identities for unset cells with NA

reverse Reverse ordering

afxn Function to evaluate each identity class based on; default is mean

reorder.numeric

Rename all identity classes to be increasing numbers starting from 1 (default is

FALSE)

x, object An object

#### Value

Idents: The cell identities

Idents<-: object with the cell identities changed

RenameIdents: An object with selected identity classes renamed

ReorderIdent: An object with

SetIdent: An object with new identity classes set StashIdent: An object with the identities stashed

```
# Get cell identity classes
Idents(pbmc_small)
# Set cell identity classes
# Can be used to set identities for specific cells to a new level
Idents(pbmc_small, cells = 1:4) <- 'a'</pre>
head(Idents(pbmc_small))
# Can also set idents from a value in object metadata
colnames(pbmc_small[[]])
Idents(pbmc_small) <- 'RNA_snn_res.1'</pre>
levels(pbmc_small)
# Rename cell identity classes
# Can provide an arbitrary amount of idents to rename
levels(pbmc_small)
pbmc_small <- RenameIdents(pbmc_small, '0' = 'A', '2' = 'C')</pre>
levels(pbmc_small)
## Not run:
head(Idents(pbmc_small))
pbmc_small <- ReorderIdent(pbmc_small, var = 'PC_1')</pre>
head(Idents(pbmc_small))
## End(Not run)
# Set cell identity classes using SetIdent
cells.use <- WhichCells(pbmc_small, idents = '1')</pre>
pbmc_small <- SetIdent(pbmc_small, cells = cells.use, value = 'B')</pre>
```

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```
head(pbmc_small[[]])
pbmc_small <- StashIdent(pbmc_small, save.name = 'idents')
head(pbmc_small[[]])

# Get the levels of identity classes of a Seurat object
levels(x = pbmc_small)

# Reorder identity classes
levels(x = pbmc_small)
levels(x = pbmc_small) <- c('C', 'A', 'B')
levels(x = pbmc_small)</pre>
```

**Images** 

Pull spatial image names

# Description

List the names of SpatialImage objects present in a Seurat object. If assay is provided, limits search to images associated with that assay

# Usage

```
Images(object, assay = NULL)
```

# Arguments

object A Seurat object

assay Name of assay to limit search to

## Value

A list of image names

```
## Not run:
Images(object)
## End(Not run)
```

66 Indices

Index

Get Neighbor algorithm index

## **Description**

Get Neighbor algorithm index

## Usage

```
Index(object, ...)
Index(object, ...) <- value
## S3 method for class 'Neighbor'
Index(object, ...)
## S3 replacement method for class 'Neighbor'
Index(object, ...) <- value</pre>
```

## **Arguments**

object An object

... Arguments passed to other methods

value The index to store

### Value

Returns the value in the alg.idx slot of the Neighbor object

Idents<-: A Neighbor object with the index stored

Indices

Get Neighbor nearest neighbor index matrices

## **Description**

Get Neighbor nearest neighbor index matrices

# Usage

```
Indices(object, ...)
## S3 method for class 'Neighbor'
Indices(object, ...)
```

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### **Arguments**

object An object
... Arguments passed to other methods

## Value

A matrix with the nearest neighbor indices

intersect.LogMap

Find Common Logical Map Values

## **Description**

Identify values in a logical map that are common to every observation

## Usage

```
## S3 method for class 'LogMap'
intersect(x, y = missing_arg(), ...)
```

## **Arguments**

```
x A LogMap objecty Ignored... Ignored
```

### Value

The values of x that are present in **every** observation

### See Also

```
Logical map objects, validity, and interaction methods: LogMap-validity, LogMap, as.matrix.LogMap(), droplevels.LogMap(), labels.LogMap()
```

```
map <- LogMap(letters[1:10])
map[['obs']] <- c(1, 3, 7)
map[['entry']] <- c(2, 7, 10)

# Identify values that are present in every observation
intersect(map)</pre>
```

68 IsMatrixEmpty

IsGlobal

Is an object global/persistent?

## **Description**

Typically, when removing Assay objects from an Seurat object, all associated objects (eg. DimReduc, Graph, and SeuratCommand objects) are removed as well. If an associated object is marked as global/persistent, the associated object will remain even if its original assay was deleted

### Usage

```
IsGlobal(object, ...)
## Default S3 method:
IsGlobal(object, ...)
## S3 method for class 'DimReduc'
IsGlobal(object, ...)
```

# Arguments

object An object

... Arguments passed to other methods

### Value

TRUE if the object is global/persistent otherwise FALSE

# **Examples**

```
IsGlobal(pbmc_small[['pca']])
```

IsMatrixEmpty

Check if a matrix is empty

# Description

Takes a matrix and asks if it's empty (either 0x0 or 1x1 with a value of NA)

# Usage

```
IsMatrixEmpty(x)
## Default S3 method:
IsMatrixEmpty(x)
```

IsNamedList 69

## **Arguments**

x A matrix

#### Value

Whether or not x is empty

#### **Examples**

```
IsMatrixEmpty(new("matrix"))
IsMatrixEmpty(matrix())
IsMatrixEmpty(matrix(1:3))
```

IsNamedList

Check List Names

## **Description**

Check to see if a list has names; also check to enforce that all names are present and unique

### Usage

```
IsNamedList(x, all.unique = TRUE, allow.empty = FALSE, pass.zero = FALSE)
```

## Arguments

```
x A list

all.unique Require that all names are unique from one another

allow.empty Allow empty (nchar = 0) names

pass.zero Pass on zero-length lists
```

## Value

```
TRUE if ..., otherwise FALSE
```

```
IsNamedList(list())
IsNamedList(list(), pass.zero = TRUE)
IsNamedList(list(1, 2, 3))
IsNamedList(list(a = 1, b = 2, c = 3))
IsNamedList(list(a = 1, 2, c = 3))
IsNamedList(list(a = 1, 2, c = 3), allow.empty = TRUE)
IsNamedList(list(a = 1, a = 2, a = 3))
IsNamedList(list(a = 1, a = 2, a = 3), all.unique = FALSE)
```

70 JackStrawData-methods

JackStrawData-class The JackStrawData Class

## **Description**

The JackStrawData is used to store the results of a JackStraw computation.

#### **Slots**

```
empirical.p.values Empirical p-values
fake.reduction.scores Fake reduction scores
empirical.p.values.full Empirical p-values on full
overall.p.values Overall p-values from ScoreJackStraw
```

JackStrawData-methods JackStrawData Methods

# Description

Methods for JackStrawData objects for generics defined in other packages

## Usage

```
## S3 method for class 'JackStrawData'
.DollarNames(x, pattern = "")
## S3 method for class 'JackStrawData'
x$i, ...
## S3 method for class 'JackStrawData'
as.logical(x, ...)
## S4 method for signature 'JackStrawData'
show(object)
```

# Arguments

```
    x, object A JackStrawData object
    pattern A regular expression. Only matching names are returned.
    i A JackStrawData slot name
    ... Ignored
```

JoinLayers 71

## Value

```
$: Slot i from x as.logical: TRUE if empirical p-values have been calculated otherwise FALSE show: Prints summary to stdout and invisibly returns NULL
```

#### **Functions**

- .DollarNames(JackStrawData): Autocompletion for \$ access on a JackStrawData object
- \$: Access data from a JackStrawData object
- as.logical(JackStrawData): Have empirical p-values for a JackStrawData object been calculated
- show(JackStrawData): Overview of a JackStrawData object

JoinLayers

Split and Join Layers Together

### **Description**

Split and Join Layers Together

# Usage

```
JoinLayers(object, ...)
## S3 method for class 'Assay5'
JoinLayers(object, layers = NULL, new = NULL, ...)
## S3 method for class 'Seurat'
JoinLayers(object, assay = NULL, layers = NULL, new = NULL, ...)
```

### **Arguments**

object An object
... Arguments passed to other methods
layers Names of layers to split or join
new Name of new layers
assay Name of assay to split layers

#### Value

object with the layers specified joined

Get and set JackStraw information

JS

# Description

Get and set JackStraw information

# Usage

```
JS(object, ...)
JS(object, ...) <- value

## S3 method for class 'JackStrawData'
JS(object, slot, ...)

## S3 replacement method for class 'JackStrawData'
JS(object, slot, ...) <- value

## S3 method for class 'DimReduc'
JS(object, slot = NULL, ...)

## S3 replacement method for class 'DimReduc'
JS(object, slot = NULL, ...) <- value</pre>
```

## Arguments

object	An object
	Arguments passed to other methods
value	JackStraw information
slot	Name of slot to store JackStraw scores to Can shorten to 'empirical', 'fake', 'full', or 'overall'

## Value

```
JS: either a JackStrawData object or the specified jackstraw data JS<-: object with the update jackstraw information
```

Key 73

Key

Get and set object keys

# **Description**

Get and set object keys

### Usage

```
Key(object, ...)
Keys(object, ...)
Key(object, ...) <- value</pre>
## S3 method for class 'Assay'
Key(object, ...)
## S3 replacement method for class 'Assay'
Key(object, ...) <- value</pre>
## S3 method for class 'Assay5'
Key(object, ...)
## S3 replacement method for class 'Assay5'
Key(object, ...) <- value</pre>
## S3 method for class 'DimReduc'
Key(object, ...)
## S3 replacement method for class 'DimReduc'
Key(object, ...) <- value</pre>
## S3 method for class 'Seurat'
Key(object, ...)
## S3 method for class 'Seurat'
Keys(object, ...)
```

# Arguments

object An object
... Arguments passed to other methods
value Key value

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# Value

```
Key: the object key

Keys: a named vector of keys of sub-objects

Key<-: object with an updated key
```

# **Examples**

```
# Get an Assay key
Key(pbmc_small[["RNA"]])

# Set the key for an Assay
Key(pbmc_small[["RNA"]]) <- "newkey_"
Key(pbmc_small[["RNA"]])

# Get a DimReduc key
Key(object = pbmc_small[["pca"]])

# Set the key for DimReduc
Key(object = pbmc_small[["pca"]]) <- "newkey2_"
Key(object = pbmc_small[["pca"]])

# Show all keys associated with a Seurat object
Key(object = pbmc_small)
Keys(object = pbmc_small)</pre>
```

labels.LogMap

Find Observations by Value

# Description

Identify the observations that contain a specific value in a logical map

```
## S3 method for class 'LogMap'
labels(
  object,
  values,
  select = c("first", "last", "common", "all"),
  simplify = TRUE,
  ...
)
```

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# Arguments

#### Value

labels: A list, or vector if simplify is TRUE, of all values and the observations they're found in, according to the value of select

### See Also

Logical map objects, validity, and interaction methods: LogMap-validity, LogMap, as.matrix.LogMap(), droplevels.LogMap(), intersect.LogMap()

### **Examples**

```
map <- LogMap(letters[1:10])
map[['obs']] <- c(1, 3, 7)
map[['entry']] <- c(2, 7, 10)

# Find observations for a set of values labels(map, c('a', 'b', 'g'))</pre>
```

LayerData

Query and Manipulate Assay Layers

# **Description**

Query and Manipulate Assay Layers

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```
LayerData(object, layer, ...)
LayerData(object, layer, ...) <- value
Layers(object, ...)
## S3 method for class 'Assay'
LayerData(
 object,
  layer = NULL,
  cells = NULL,
  features = NULL,
  slot = deprecated(),
)
## S3 replacement method for class 'Assay'
LayerData(object, layer, ...) <- value
## S3 method for class 'Assay'
Layers(object, search = NA, ...)
## S3 method for class 'Assay5'
LayerData(
  object,
  layer = NULL,
  cells = NULL,
  features = NULL,
  fast = FALSE,
  slot = deprecated(),
  . . .
)
## S3 replacement method for class 'Assay5'
LayerData(object, layer, features = NULL, cells = NULL, ...) <- value
## S3 method for class 'Assay5'
Layers(object, search = NA, ...)
## S3 method for class 'Seurat'
LayerData(object, layer = NULL, assay = NULL, slot = deprecated(), ...)
## S3 replacement method for class 'Seurat'
LayerData(object, layer, assay = NULL, ...) <- value
## S3 method for class 'Seurat'
Layers(object, search = NA, assay = NULL, ...)
```

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### **Arguments**

object An object

layer Name of layer to fetch or set

... Arguments passed to other methods

value New two-dimensional data to be added as a layer

features, cells

Vectors of features/cells to include

slot [Deprecated]

search A pattern to search layer names for; pass one of:

• "NA" to pull all layers

• "NULL" to pull the default layer(s)

• a regular expression that matches layer names

fast Determine how to return the layer data; choose from:

FALSE Apply any transpositions and attempt to add feature/cell names (if sup-

ported) back to the layer data

NA Attempt to add feature/cell names back to the layer data, skip any transposi-

tions

TRUE Do not apply any transpositions or add feature/cell names to the layer data

assay Name of assay to fetch layer data from or assign layer data to

### Value

LayerData: the layer data for layer from object

Layer<-: object with value added as a layer named layer

Layers: the names of the layers present in object

Loadings

Get and set feature loadings

#### **Description**

Get and set feature loadings

```
Loadings(object, ...)
Loadings(object, ...) <- value
## S3 method for class 'DimReduc'
Loadings(object, projected = FALSE, ...)</pre>
```

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```
## S3 replacement method for class 'DimReduc'
Loadings(object, projected = TRUE, ...) <- value
## S3 method for class 'Seurat'
Loadings(object, reduction = "pca", projected = FALSE, ...)</pre>
```

# **Arguments**

object An object

... Arguments passed to other methods

value Feature loadings to add

projected Pull the projected feature loadings?

reduction Name of reduction to pull feature loadings for

#### Value

```
Loadings: the feature loadings for object
Loadings<-: object with the updated loadings
```

# **Examples**

```
# Get the feature loadings for a given DimReduc
Loadings(object = pbmc_small[["pca"]])[1:5,1:5]

# Set the feature loadings for a given DimReduc
new.loadings <- Loadings(object = pbmc_small[["pca"]])
new.loadings <- new.loadings + 0.01
Loadings(object = pbmc_small[["pca"]]) <- new.loadings
# Get the feature loadings for a specified DimReduc in a Seurat object
Loadings(object = pbmc_small, reduction = "pca")[1:5,1:5]</pre>
```

LogMap

A Logical Map

# Description

A simple container for storing mappings of values using logical matrices. Keeps track of which values (rows) are present in which observations (columns). LogMap objects can be created with LogMap(); queries can be performed with [[ and observations can be added or removed with [[ <-

LogMap 79

### Usage

```
LogMap(y)

## S4 method for signature 'LogMap,character,missing'
x[[i, j, ...]]

## S4 method for signature 'LogMap,missing,missing'
x[[i, j, ...]]

## S4 method for signature 'LogMap,`NULL`,missing'
x[[i, j, ...]]

## S4 replacement method for signature 'LogMap,character,missing,character'
x[[i, j, ...]] <- value

## S4 replacement method for signature 'LogMap,character,missing,integer'
x[[i, j, ...]] <- value

## S4 replacement method for signature 'LogMap,character,missing,`NULL`'
x[[i, j, ...]] <- value

## S4 replacement method for signature 'LogMap,character,missing,`NULL`'
x[[i, j, ...]] <- value
```

# **Arguments**

У	A character vector
X	A LogMap object
i	A character vector of length 1, or NULL
j	Not used
	Ignored
value	A character or integer vector of values to record in the map for $i$ , or NULL to remove the record for $i$

### Value

LogMap: A new LogMap object with zero columns and length(x = x) rows; rownames are set to x [[: if i is a character vector, the rownames that are mapped to i; otherwise the rownames of x [[<-: If value is NULL, then x without the observations for i; otherwise, x with a new column for i recording a TRUE for all values present in value

#### **Slots**

.Data A logical matrix with at least one row

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### See Also

Logical map objects, validity, and interaction methods: LogMap-validity, as.matrix.LogMap(), droplevels.LogMap(), intersect.LogMap(), labels.LogMap()

### **Examples**

```
# Create a LogMap
map <- LogMap(letters[1:10])</pre>
# Get the names of values in the LogMap
map[[NULL]]
rownames(map)
# Add an observation to the LogMap
map[['obs']] \leftarrow c(1, 3, 7)
map[['entry']] \leftarrow c(2, 7, 10)
map
# Get the names of observations in the LogMap
colnames(map)
# Fetch an observation from the LogMap
map[['obs']]
# Get the full logical matrix
map[[]]
# Remove an observation from the LogMap
map[['obs']] <- NULL</pre>
map[['entry']] <- NULL</pre>
map
```

LogMap-validity

Logical Map Validity

# Description

Validation of LogMap objects is handled by validObject

### **Data Validation**

Logical maps must be a logical matrix containing only TRUE or FALSE values

#### Value Validation

All values must be named within the rownames of the object. Duplicate or empty ("") values are not allowed

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### **Observation Validation**

All observations must be named within the column names of the object. Duplicate or empty ("") observations are not allowed

#### See Also

```
validObject
```

Logical map objects, validity, and interaction methods: LogMap, as.matrix.LogMap(), droplevels.LogMap(), intersect.LogMap(), labels.LogMap()

# **Examples**

```
map <- LogMap(letters[1:10])
map[['obs']] <- c(1, 3, 7)
map[['entry']] <- c(2, 7, 10)
validObject(map)</pre>
```

LogSeuratCommand

Log a command

# **Description**

Logs command run, storing the name, timestamp, and argument list. Stores in the Seurat object

### Usage

```
LogSeuratCommand(object, return.command = FALSE)
```

# **Arguments**

```
object Name of Seurat object
return.command Return a SeuratCommand object instead
```

#### Value

If return.command, returns a SeuratCommand object; otherwise, returns the Seurat object with command stored

### See Also

# Command

```
Command log object and interaction methods $.SeuratCommand(), .DollarNames.SeuratCommand(), SeuratCommand-class, [.SeuratCommand(), as.list.SeuratCommand()
```

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merge.Assay

Merge Assays

# **Description**

Merge one or more v3 assays together

# Usage

```
## S3 method for class 'Assay'
merge(
  x = NULL,
  y = NULL,
  add.cell.ids = NULL,
  merge.data = TRUE,
  labels = NULL,
  collapse = TRUE,
  ...
)
```

### Arguments

```
x An Assay object
y One or more Assay objects
add.cell.ids A character vector of length(x = c(x, y)); appends the corresponding values to the start of each objects' cell names

merge.data Merge the data slots instead of just merging the counts (which requires renormalization); this is recommended if the same normalization approach was applied to all objects

labels, collapse

Currently unused
... Ignored
```

# Value

A new assay with data merged from c(x, y)

# See Also

```
v3 Assay object, validity, and interaction methods: $.Assay(), Assay-class, Assay-validity, CreateAssayObject(), [.Assay(), [.Assay(), dim.Assay(), dimnames.Assay(), split.Assay(), subset.Assay()
```

merge.Assay5

# Description

Merge one or more v5 assays together

# Usage

```
## S3 method for class 'Assay5'
merge(x, y, labels = NULL, add.cell.ids = NULL, collapse = FALSE, ...)
```

# Arguments

X	An Assay5 object
У	One or more Assay5 objects
labels	A character vector equal to the number of objects; defaults to as.character(seq_along(c(x, y)))
add.cell.ids	A character vector equal to the number of objects provided to append to all cell names; if TRUE, uses labels as add.cell.ids
collapse	If TRUE, merge layers of the same name together; if FALSE, appends labels to the layer name
	Ignored

# **Details**

Note: collapsing layers is currently not supported

# Value

A new v5 assay with data merged from c(x, y)

# See Also

```
v5 Assay object, validity, and interaction methods: $.Assay5(), Assay5-class, Assay5-validity, [.Assay5(), [[.Assay5(), dim.Assay5(), dimnames.Assay5(), split.Assay5(), subset.Assay5()
```

84 merge.Seurat

merge.DimReduc

Merge Dimensional Reductions

### **Description**

Merge two or more dimensional reduction together

### Usage

```
## S3 method for class 'DimReduc'
merge(x = NULL, y = NULL, add.cell.ids = NULL, ...)
```

# **Arguments**

```
    x A DimReduc object
    y One or more DimReduc objects
    add.cell.ids A character vector equal to the number of objects provided to append to all cell names; if TRUE, uses labels as add.cell.ids
    ... Ignored
```

#### Value

A new DimReduc object with data merged from c(x, y)

# See Also

Dimensional reduction object, validity, and interaction methods CreateDimReducObject(), DimReduc-class, DimReduc-validity, [.DimReduc(), [[.DimReduc(), dim.DimReduc(), print.DimReduc(), subset.DimReduc()

merge.Seurat

Merge Seurat Objects

### **Description**

Merge Seurat Objects

```
## S3 method for class 'Seurat'
merge(
  x = NULL,
  y = NULL,
  add.cell.ids = NULL,
  collapse = FALSE,
  merge.data = TRUE,
```

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```
merge.dr = FALSE,
project = getOption(x = "Seurat.object.project", default = "SeuratProject"),
...
)
```

#### **Arguments**

X	A Seurat object
У	A single Seurat object or a list of Seurat objects
add.cell.ids	A character vector of length( $x = c(x, y)$ ); appends the corresponding values to the start of each objects' cell names
collapse	If TRUE, merge layers of the same name together; if FALSE, appends labels to the layer name $$
merge.data	Merge the data slots instead of just merging the counts (which requires renormalization); this is recommended if the same normalization approach was applied to all objects
merge.dr	Choose how to handle merging dimensional reductions:
	• "TRUE": merge dimensional reductions with the same name across objects; dimensional reductions with different names are added as-is
	<ul> <li>"NA": keep dimensional reductions from separate objects separate; will append the project name for duplicate reduction names</li> </ul>
	"FALSE": do not add dimensional reductions
project	Project name for the Seurat object
	Arguments passed to other methods

#### Value

merge: Merged object

### **Merge Details**

When merging Seurat objects, the merge procedure will merge the Assay level counts and potentially the data slots (depending on the merge.data parameter). It will also merge the cell-level meta data that was stored with each object and preserve the cell identities that were active in the objects pre-merge. The merge will optionally merge reductions depending on the values passed to merge.dr if they have the same name across objects. Here the embeddings slots will be merged and if there are differing numbers of dimensions across objects, only the first N shared dimensions will be merged. The feature loadings slots will be filled by the values present in the first object. The merge will not preserve graphs, logged commands, or feature-level metadata that were present in the original objects. If add.cell.ids isn't specified and any cell names are duplicated, cell names will be appended with X, where X is the numeric index of the object in C(X, Y).

#### See Also

```
Seurat object, validity, and interaction methods $.Seurat(), Seurat-class, Seurat-validity,
[[.Seurat(), [[<-,Seurat, NULL, [[<-,Seurat, dim.Seurat(), dimnames.Seurat(), names.Seurat(),
subset.Seurat()</pre>
```

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### **Examples**

```
# `merge' examples
# merge two objects
merge(pbmc_small, y = pbmc_small)
# to merge more than two objects, pass one to x and a list of objects to y
merge(pbmc_small, y = c(pbmc_small, pbmc_small))
```

Misc

Get and set miscellaneous data

# Description

Get and set miscellaneous data

```
Misc(object, ...)
Misc(object, ...) <- value
## S3 method for class 'Assay'
Misc(object, slot = NULL, ...)
## S3 replacement method for class 'Assay'
Misc(object, slot, ...) <- value
## S3 method for class 'Assay5'
Misc(object, slot = NULL, ...)
## S3 replacement method for class 'Assay5'
Misc(object, slot, ...) <- value
## S3 method for class 'DimReduc'
Misc(object, slot = NULL, ...)
## S3 replacement method for class 'DimReduc'
Misc(object, slot, ...) <- value
## S3 method for class 'Seurat'
Misc(object, slot = NULL, ...)
## S3 replacement method for class 'Seurat'
Misc(object, slot, ...) <- value
```

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### **Arguments**

object An object

... Arguments passed to other methods

value Data to add

slot Name of specific bit of meta data to pull

### Value

Miscellaneous data

An object with miscellaneous data added

# **Examples**

```
# Get the misc info
Misc(object = pbmc_small, slot = "example")
# Add misc info
Misc(object = pbmc_small, slot = "example") <- "testing_misc"</pre>
```

Molecules-class

The Spatial Molecules Class

# Description

The Spatial Molecules Class

### **Slots**

```
.Data A list of SpatialPoints objects key The key for the Molecules
```

### See Also

Molecules methods: Molecules-methods

 $Segmentation\ layer\ classes:\ Centroids-class,\ Centroids-methods,\ Molecules-methods,\ Segmentation-class,\ Segmentation-methods$ 

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Molecules-methods

Molecules Methods

### **Description**

Methods for Molecules objects

### Usage

```
## S3 method for class 'Molecules'
Features(x, ...)

## S3 method for class 'Molecules'
GetTissueCoordinates(object, features = NULL, ...)

## S3 method for class 'Molecules'
subset(x, features = NULL, ...)

## S4 method for signature 'Molecules'
show(object)
```

#### **Arguments**

x, object A Molecules object

... Arguments passed to other methods

features A vector of molecule names to keep; if NULL, defaults to all molecules

### **Details**

Features: Get spatially-resolved molecule names

GetTissueCoordinates: Get spatially-resolved molecule coordinates

subset: Subset a Molecules object to certain molecules

show: Display an object summary to stdout

### Value

Features: A vector of spatially-resolved molecule names; if no molecular information present, returns NULL

GetTissueCoordinates: A data frame with three columns:

- "x": the x-coordinate of a molecule
- "y": the y-coordinate of a molecule
- "molecule": the molecule name

subset: x subsetted to the features specified by features

show: Invisibly returns NULL

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# See Also

```
Molecules-class
```

Segmentation layer classes: Centroids-class, Centroids-methods, Molecules-class, Segmentation-class, Segmentation-methods

names.Seurat

Subobject Names

# **Description**

Get the names of subobjects within a Seurat object

# Usage

```
## S3 method for class 'Seurat'
names(x)
```

# Arguments

Χ

A Seurat object

# Value

The names of all of the following subobjects within x:

- v3 and v5 assays
- dimensional reductions
- images and FOVs
- nearest-neighbor graphs

### See Also

```
Seurat object, validity, and interaction methods $.Seurat(), Seurat-class, Seurat-validity,
[[.Seurat(), [[<-,Seurat, NULL, [[<-,Seurat, dim.Seurat(), dimnames.Seurat(), merge.Seurat(),
subset.Seurat()</pre>
```

# **Examples**

```
names(pbmc_small)
```

90 Neighbor-methods

Neighbor-class

The Neighbor class

### **Description**

The Neighbor class is used to store the results of neighbor finding algorithms

#### **Slots**

```
nn.idx Matrix containing the nearest neighbor indices
```

nn.dist Matrix containing the nearest neighbor distances

alg.idx The neighbor finding index (if applicable). E.g. the annoy index

alg.info Any information associated with the algorithm that may be needed downstream (e.g. distance metric used with annoy is needed when reading in from stored file).

cell. names Names of the cells for which the neighbors have been computed.

Neighbor-methods

Neighbor Methods

# **Description**

Methods for Neighbor objects for generics defined in other packages

# Usage

```
## S3 method for class 'Neighbor'
dim(x)
## S4 method for signature 'Neighbor'
show(object)
```

# Arguments

```
x, object A Neighbor object
```

### Value

dim Dimensions of the indices matrix

show: Prints summary to stdout and invisibly returns NULL

### **Functions**

- dim(Neighbor): Dimensions of the neighbor indices
- show(Neighbor): Overview of a Neighbor object

Overlay 91

**Overlay** 

Overlay Spatial Objects Over One Another

### **Description**

Create an overlay of some query spatial object (x) against some target object (y). Basically, find all components of a query that fall within the bounds of a target spatial region

# Usage

```
Overlay(x, y, invert = FALSE, ...)
## S4 method for signature 'Centroids,SpatialPolygons'
Overlay(x, y, invert = FALSE, ...)
## S4 method for signature 'Segmentation,SpatialPolygons'
Overlay(x, y, invert = FALSE, ...)
## S4 method for signature 'Molecules,SpatialPolygons'
Overlay(x, y, invert = FALSE, ...)
## S4 method for signature 'FOV,Spatial'
Overlay(x, y, invert = FALSE, ...)
## S4 method for signature 'FOV,SpatialPolygons'
Overlay(x, y, invert = FALSE, ...)
## S4 method for signature 'FOV,FOV'
Overlay(x, y, invert = FALSE, ...)
```

# Arguments

X	Query Spatial object
У	Target Spatial object
invert	Invert the overlay and return only the components of $x$ that fall $outside$ the bounds of $y$
	Ignored

#### Value

x with only the components that fall within the bounds of y

#### Note

This function requires the sf package to be installed

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PackageCheck

Check the existence of a package

# Description

Check the existence of a package

# Usage

```
PackageCheck(..., error = TRUE)
```

# **Arguments**

.. Package names

error If true, throw an error if the package doesn't exist

### Value

Invisibly returns boolean denoting if the package is installed

# Lifecycle

# [Deprecated]

PackageCheck was deprecated in version 5.0.0; please use rlang::check\_installed() instead

# **Examples**

```
PackageCheck("SeuratObject", error = FALSE)
```

pbmc\_small

A small example version of the PBMC dataset

# Description

A subsetted version of 10X Genomics' 3k PBMC dataset

```
pbmc_small
```

print.DimReduc 93

### **Format**

A Seurat object with the following slots filled

assays Currently only contains one assay ("RNA" - scRNA-seq expression data)

counts - Raw expression data

- data Normalized expression data
- scale.data Scaled expression data
- var.features names of the current features selected as variable
- meta.features Assay level metadata such as mean and variance

meta.data Cell level metadata

active.assay Current default assay

active.ident Current default idents

graphs Neighbor graphs computed, currently stores the SNN

reductions Dimensional reductions: currently PCA and tSNE

version Seurat version used to create the object

**commands** Command history

#### Source

https://support.10xgenomics.com/single-cell-gene-expression/datasets/1.1.0/pbmc3k

print.DimReduc

Print Top Feature Loadings

# Description

Prints a set of features that most strongly define a set of components; **note**: requires feature loadings to be present in order to work

### Usage

```
## S3 method for class 'DimReduc'
print(x, dims = 1:5, nfeatures = 20, projected = FALSE, ...)
```

### **Arguments**

x A DimReduc object

dims Number of dimensions to display

nfeatures Number of genes to display

projected Use projected slot

... Ignored

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### Value

Displays set of features defining the components and invisibly returns x

#### See Also

cat

Dimensional reduction object, validity, and interaction methods CreateDimReducObject(), DimReduc-class, DimReduc-validity, [.DimReduc(), [[.DimReduc(), dim.DimReduc(), merge.DimReduc(), subset.DimReduc()

# **Examples**

```
pca <- pbmc_small[["pca"]]
print(pca)</pre>
```

Project

Get and set project information

# **Description**

Get and set project information

# Usage

```
Project(object, ...)
Project(object, ...) <- value

## S3 method for class 'Seurat'
Project(object, ...)

## S3 replacement method for class 'Seurat'
Project(object, ...) <- value</pre>
```

### **Arguments**

object An object

... Arguments passed to other methods

value Project information to set

#### Value

Project information

An object with project information added

Radius 95

Radius

Get the spot radius from an image

# Description

Get the spot radius from an image

# Usage

```
Radius(object)
```

# Arguments

object

An image object

# Value

The radius size

RandomName

Generate a random name

# **Description**

Make a name from randomly sampled characters, pasted together with no spaces

# Usage

```
RandomName(length = 5L, chars = letters, ...)
```

# **Arguments**

length How long should the name be

chars A vector of 1-length characters to use to generate the name

... Extra parameters passed to sample

# Value

A character with nchar == length of randomly sampled letters

# See Also

sample

96 RenameAssays

# **Examples**

```
set.seed(42L)
RandomName()
RandomName(7L, replace = TRUE)
```

RenameAssays

Rename assays in a Seurat object

# Description

Rename assays in a Seurat object

# Usage

```
RenameAssays(
  object,
  assay.name = NULL,
  new.assay.name = NULL,
  verbose = TRUE,
  ...
)
```

# Arguments

```
object A Seurat object
assay.name original name of assay
new.assay.name new name of assay
verbose Whether to print messages
... Named arguments as old.assay = new.assay
```

#### Value

object with assays renamed

# **Examples**

```
RenameAssays(object = pbmc_small, RNA = 'rna')
```

RenameCells 97

RenameCells Rename cells

### **Description**

Change the cell names in all the different parts of an object. Can be useful before combining multiple objects.

# Usage

```
RenameCells(object, ...)
## S3 method for class 'Assay'
RenameCells(object, new.names = NULL, ...)
## S3 method for class 'Assay5'
RenameCells(object, new.names = NULL, ...)
## S3 method for class 'DimReduc'
RenameCells(object, new.names = NULL, ...)
## S3 method for class 'Neighbor'
RenameCells(object, old.names = NULL, new.names = NULL, ...)
## S3 method for class 'Seurat'
RenameCells(
 object,
  add.cell.id = missing_arg(),
 new.names = missing_arg(),
  for.merge = deprecated(),
)
```

#### **Arguments**

object	An object
	Arguments passed to other methods
new.names	vector of new cell names
old.names	vector of old cell names
add.cell.id	prefix to add cell names
for.merge	Deprecated

#### **Details**

If add.cell.id is set a prefix is added to existing cell names. If new.names is set these will be used to replace existing names.

### Value

An object with new cell names

# Examples

```
# Rename cells in an Assay
head(x = colnames(x = pbmc_small[["RNA"]]))
renamed.assay <- RenameCells(</pre>
    pbmc_small[["RNA"]],
    new.names = paste0("A_", colnames(x = pbmc_small[["RNA"]]))
head(x = colnames(x = renamed.assay))
# Rename cells in a DimReduc
head(x = Cells(x = pbmc_small[["pca"]]))
renamed.dimreduc <- RenameCells(</pre>
    object = pbmc_small[["pca"]],
    new.names = paste0("A_", Cells(x = pbmc_small[["pca"]]))
head(x = Cells(x = renamed.dimreduc))
# Rename cells in a Seurat object
head(x = colnames(x = pbmc_small))
pbmc_small <- RenameCells(object = pbmc_small, add.cell.id = "A")</pre>
head(x = colnames(x = pbmc_small))
```

RowMergeSparseMatrices

Merge Sparse Matrices by Row

### **Description**

Merge two or more sparse matrices by rowname.

#### **Usage**

```
RowMergeSparseMatrices(mat1, mat2)
```

### **Arguments**

mat1 First matrix

mat2 Second matrix or list of matrices

#### **Details**

Shared matrix rows (with the same row name) will be merged, and unshared rows (with different names) will be filled with zeros in the matrix not containing the row.

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# Value

Returns a sparse matrix

SaveSeuratRds Save and Load Seurat Objects from Rds files

# **Description**

Save and Load Seurat Objects from Rds files

# Usage

```
SaveSeuratRds(object, file = NULL, destdir = NULL, relative = FALSE, ...)
LoadSeuratRds(file, ...)
```

# Arguments

object	A Seurat object
file	Path to save object to; defaults to file.path(getwd(), paste0(Project(object), ".Rds"))
destdir	Destination directory for on-disk layers saved in "/tmp/RtmpeZpjRg"
relative	Save relative paths instead of absolute ones
	Arguments passed on to base::saveRDS, base::readRDS
	ascii a logical. If TRUE or NA, an ASCII representation is written; otherwise (default), a binary one is used. See the comments in the help for save.
	version the workspace format version to use. NULL specifies the current default version (3). The only other supported value is 2, the default from R 1.4.0 to R 3.5.0.
	compress a logical specifying whether saving to a named file is to use "gzip" compression, or one of "gzip", "bzip2" or "xz" to indicate the type of compression to be used. Ignored if file is a connection.
	refhook a hook function for handling reference objects.

### Value

Invisibly returns file

# **Progress Updates with progressr**

This function uses **progressr** to render status updates and progress bars. To enable progress updates, wrap the function call in with\_progress or run handlers(global = TRUE) before running this function. For more details about **progressr**, please read vignette("progressr-intro")

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# Note

This function requires the fs package to be installed

#### See Also

```
saveRDS() readRDS()
```

# **Examples**

```
if (requireNamespace("fs", quietly = TRUE)) {
 # Write out with DelayedArray
 if (requireNamespace("HDF5Array", quietly = TRUE)) {
    pbmc <- pbmc_small</pre>
   pbmc[["disk"]] <- CreateAssay50bject(list(</pre>
      mem = LayerData(pbmc, "counts"),
      disk = as(LayerData(pbmc, "counts"), "HDF5Array")
    ))
    # Save `pbmc` to an Rds file
    out <- tempfile(fileext = ".Rds")</pre>
    SaveSeuratRds(pbmc, file = out)
    # Object cache
    obj <- readRDS(out)</pre>
    Tool(obj, "SaveSeuratRds")
    # Load the saved object with on-disk layers back into memory
    pbmc2 <- LoadSeuratRds(out)</pre>
   pbmc2
   pbmc2[["disk"]]
 # Write out with BPCells
 if (requireNamespace("BPCells", quietly = TRUE)) {
    pbmc <- pbmc_small</pre>
    bpm <- BPCells::write_matrix_dir(LayerData(pbmc, "counts"), dir = tempfile())</pre>
   bph <- BPCells::write_matrix_hdf5(</pre>
      LayerData(pbmc, "counts"),
      path = tempfile(fileext = ".h5"),
      group = "counts"
   pbmc[["disk"]] <- CreateAssay50bject(list(dir = bpm, h5 = bph))</pre>
    # Save `pbmc` to an Rds file
    out <- tempfile(fileext = ".Rds")</pre>
    SaveSeuratRds(pbmc, file = out)
    # Object cache
    obj <- readRDS(out)</pre>
    Tool(obj, "SaveSeuratRds")
```

Segmentation-class 101

```
# Load the saved object with on-disk layers back into memory
pbmc2 <- LoadSeuratRds(out)
pbmc2
pbmc2[["disk"]]
}</pre>
```

Segmentation-class

The Segmentation Class

# **Description**

The Segmentation Class

### See Also

Segmentation methods: Segmentation-methods

 $Segmentation\ layer\ classes:\ Centroids-methods,\ Molecules-class,\ Molecules-methods,\ Segmentation-methods$ 

Segmentation-methods Segmentation Methods

# **Description**

Methods for Segmentation objects

```
## S3 method for class 'Segmentation'
Cells(x, ...)
## S3 method for class 'Segmentation'
GetTissueCoordinates(object, full = TRUE, ...)
## S3 method for class 'Segmentation'
RenameCells(object, new.names = NULL, ...)
## S3 method for class 'Segmentation'
lengths(x, use.names = TRUE)
## S3 method for class 'Segmentation'
subset(x, cells = NULL, ...)
```

```
## S4 method for signature 'Segmentation, ANY, ANY'
x[i, j, ..., drop = TRUE]

## S4 method for signature 'Segmentation'
coordinates(obj, full = TRUE, ...)

## S4 method for signature 'Segmentation'
show(object)
```

### **Arguments**

x, object, obj A Segmentation object
... Arguments passed to other methods
full Expand the coordinates to the full polygon

new.names vector of new cell names

use.names Ignored

i, cells A vector of cells to keep; if NULL, defaults to all cells

j, drop Ignored

#### **Details**

Cells: Get cell names

GetTissueCoordinates, coordinates: Get tissue coordinates

RenameCells: Update cell names

lengths: Generate a run-length encoding of the cells present subset, [: Subset a Segmentation object to certain cells

show: Display an object summary to stdout

### Value

Cells: A vector of cell names

GetTissueCoordinates, coordinates: A data frame with three columns:

- "x": the x-coordinate
- "y": the y-coordinate
- "cell" or "ID": the cell name

If full is TRUE, then each coordinate will indicate a vertex for the cell polygon; otherwise, each coordinate will indicate a centroid for the cell. Note: GetTissueCoordinates ....

RenameCells: object with the cells renamed to new.names

lengths: An rle object for the cells

subset, [: x subsetted to the cells specified by cells/i

show: Invisibly returns NULL

set-if-null 103

### **Progress Updates with progressr**

The following methods use **progressr** to render status updates and progress bars:

• RenameCells

To enable progress updates, wrap the function call in with\_progress or run handlers(global = TRUE) before running this function. For more details about **progressr**, please read vignette("progressr-intro")

### Parallelization with future

The following methods use **future** to enable parallelization:

• RenameCells

Parallelization strategies can be set using plan. Common plans include "sequential" for non-parallelized processing or "multisession" for parallel evaluation using multiple R sessions; for other plans, see the "Implemented evaluation strategies" section of ?future::plan. For a more thorough introduction to **future**, see vignette("future-1-overview")

### See Also

Segmentation-class

Segmentation layer classes: Centroids-class, Centroids-methods, Molecules-class, Molecules-methods, Segmentation-class

set-if-null

Set If or If Not NULL

### **Description**

Set a default value depending on if an object is NULL

### Usage

```
x %||% y
x %iff% y
```

# Arguments

```
x An object to testy A default value
```

#### Value

```
For %||%: y if x is NULL; otherwise x
For %iff%: y if x is not NULL; otherwise x
```

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#### Author(s)

For % | |%: rlang developers

#### See Also

```
rlang::%||%
```

### **Examples**

```
# Set if NULL
1 %||% 2
NULL %||% 2
# Set if *not* NULL
1 %iff% 2
NULL %iff% 2
```

Seurat-class

The Seurat Class

### **Description**

The Seurat object is a representation of single-cell expression data for R; each Seurat object revolves around a set of cells and consists of one or more Assay objects, or individual representations of expression data (eg. RNA-seq, ATAC-seq, etc). These assays can be reduced from their high-dimensional state to a lower-dimension state and stored as DimReduc objects. Seurat objects also store additional metadata, both at the cell and feature level (contained within individual assays). The object was designed to be as self-contained as possible, and easily extendable to new methods.

### **Slots**

Seurat-validity 105

### See Also

Seurat-validity

Seurat Object Validity

### **Description**

Validation of Seurat objects is handled by validObject

#### See Also

```
validObject
```

Seurat object, validity, and interaction methods \$.Seurat(), Seurat-class, [[.Seurat(), [[<-, Seurat, NULL, [[<-, Seurat(), dimnames.Seurat(), merge.Seurat(), names.Seurat(), subset.Seurat()

SeuratCommand-class

The SeuratCommand Class

### **Description**

The SeuratCommand is used for logging commands that are run on a Seurat object; it stores parameters and timestamps

# Slots

```
name Command name
```

 $\label{time.stamp} \mbox{time.stamp of when command was tun}$ 

assay.used Optional name of assay used to generate SeuratCommand object

call.string String of the command call

params List of parameters used in the command call

#### See Also

```
Command log object and interaction methods $.SeuratCommand(), .DollarNames.SeuratCommand(), LogSeuratCommand(), [.SeuratCommand(), as.list.SeuratCommand()
```

106 Simplify

show, LogMap-method

LogMap Object Overview

# **Description**

Overview of a LogMap object

# Usage

```
## S4 method for signature 'LogMap'
show(object)
```

# Arguments

object

A LogMap object

### Value

Prints summary to stdout and invisibly returns NULL

Simplify

Simplify Geometry

# **Description**

Simplify Geometry

Simplify segmentations by reducing the number of vertices

# Usage

```
Simplify(coords, tol, topologyPreserve = TRUE)
## S3 method for class 'Spatial'
Simplify(coords, tol, topologyPreserve = TRUE)
```

# Arguments

coords

A 'Segmentation' object

tol

Numerical tolerance value to be used by the Douglas-Peuker algorithm

topologyPreserve

Logical determining if the algorithm should attempt to preserve the topology of

the original geometry

### Value

A simplified version of coords

A 'Segmentation' object with simplified segmentation vertices

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SpatialImage-class The SpatialImage class

### **Description**

The SpatialImage class is a virtual class representing spatial information for Seurat. All spatial image information must inherit from this class for use with Seurat objects

#### Slots

assay Name of assay to associate image data with; will give this image priority for visualization when the assay is set as the active/default assay in a Seurat object

key A one-length character vector with the object's key; keys must be one or more alphanumeric characters followed by an underscore "\_" (regex pattern "^[a-zA-Z][a-zA-Z0-9]\*\_\$")

#### See Also

SpatialImage-methods for a list of required and provided methods

SpatialImage-methods SpatialImage methods

# **Description**

Methods defined on the SpatialImage class. Some of these methods must be overridden in order to ensure proper functionality of the derived classes (see **Required methods** below). Other methods are designed to work across all SpatialImage-derived subclasses, and should only be overridden if necessary

```
## S3 method for class 'SpatialImage'
Cells(x, ...)
## S3 method for class 'SpatialImage'
DefaultAssay(object, ...)
## S3 replacement method for class 'SpatialImage'
DefaultAssay(object, ...) <- value
## S3 method for class 'SpatialImage'
GetImage(object, mode = c("grob", "raster", "plotly", "raw"), ...)
## S3 method for class 'SpatialImage'
GetTissueCoordinates(object, ...)</pre>
```

```
## S3 method for class 'SpatialImage'
IsGlobal(object, ...)
## S3 method for class 'SpatialImage'
Key(object, ...)
## S3 replacement method for class 'SpatialImage'
Key(object, ...) <- value</pre>
## S3 method for class 'SpatialImage'
Radius(object)
## S3 method for class 'SpatialImage'
RenameCells(object, new.names = NULL, ...)
## S3 method for class 'SpatialImage'
x[i, ...]
## S3 method for class 'SpatialImage'
dim(x)
## S3 method for class 'SpatialImage'
subset(x, cells, ...)
## S4 method for signature 'SpatialImage'
show(object)
```

#### **Arguments**

x, object A SpatialImage-derived object
... Arguments passed to other methods

value Depends on the method:

DefaultAssay<- Assay that the image should be associated with

Key<- New key for the image

mode How to return the image; should accept one of "grob", "raster", "plotly", or

"raw"

new.names vector of new cell names
i, cells A vector of cells to keep

#### Value

[Override] Cells: should return cell names

DefaultAssay: The associated assay of a SpatialImage-derived object

DefaultAssay<-: object with the associated assay updated

[Override] GetImage: The image data from a SpatialImage-derived object

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[Override] GetTissueCoordinates: ...

IsGlobal: returns TRUE as images are, by default, global

Key: The key for a SpatialImage-derived object

Key<-: object with the key set to value

Radius: The spot radius size; by default, returns NULL

[Override] RenameCells: object with the new cell names

[, subset: x/object for only the cells requested

[Override] dim: The dimensions of the image data in (Y, X) format

show: Prints summary to stdout and invisibly returns NULL

#### **Functions**

• Cells(SpatialImage): Get the cell names from an image ([Override])

- DefaultAssay(SpatialImage): Get the associated assay of a SpatialImage-derived object
- DefaultAssay(SpatialImage) <- value: Set the associated assay of a SpatialImage-derived object
- GetImage(SpatialImage): Get the image data from a SpatialImage-derived object
- GetTissueCoordinates(SpatialImage): Get tissue coordinates for a SpatialImage-derived object ([Override])
- IsGlobal(SpatialImage): Globality test for SpatialImage-derived object
- Key(SpatialImage): Get the key for a SpatialImage-derived object
- Key(SpatialImage) <- value: Set the key for a SpatialImage-derived object
- Radius(SpatialImage): Get the spot radius size
- RenameCells(SpatialImage): Rename cells in a SpatialImage-derived object([Override])
- [: Subset a SpatialImage-derived object
- dim(SpatialImage): Get the plotting dimensions of an image ([Override])
- subset(SpatialImage): Subset a SpatialImage-derived object ([Override])
- show(SpatialImage): Overview of a SpatialImage-derived object

#### Provided methods

These methods are defined on the SpatialImage object and should not be overridden without careful thought

- DefaultAssay and DefaultAssay<-
- Key and Key<-</li>
- GetImage; this method *can* be overridden to provide image data, normally returns empty image data. If overridden, should default to returning a grob object
- IsGlobal
- Radius; this method can be overridden to provide a spot radius for image objects
- [; this method *can* be overridden to change default subset behavior, normally returns subset(x = x, cells = i). If overridden, should only accept i

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#### Required methods

All subclasses of the SpatialImage class must define the following methods; simply relying on the SpatialImage method will result in errors. For required parameters and their values, see the Usage and Arguments sections

Cells Return the cell/spot barcodes associated with each position

dim Return the dimensions of the image for plotting in (Y, X) format

GetTissueCoordinates Return tissue coordinates; by default, must return a two-column data. frame with x-coordinates in the first column and y-coordinates in the second

Radius Return the spot radius; returns NULL by default for use with non-spot image technologies

RenameCells Rename the cell/spot barcodes for this image

subset Subset the image data by cells/spots

These methods are used throughout Seurat, so defining them and setting the proper defaults will allow subclasses of SpatialImage to work seamlessly

#### See Also

```
DefaultAssay
GetImage
GetTissueCoordinates
IsGlobal
Key
RenameCells
```

split.Assay

Split an Assay

## **Description**

Split an Assay

# Usage

```
## S3 method for class 'Assay'
split(x, f, drop = FALSE, layers = NA, ...)
```

split.Assay5

# Arguments

Χ	An Assay object
f	a 'factor' in the sense that as.factor(f) defines the grouping, or a list of such factors in which case their interaction is used for the grouping. If $x$ is a data frame, f can also be a formula of the form $\sim$ g to split by the variable g, or more generally of the form $\sim$ g1 + + gk to split by the interaction of the variables g1, , gk, where these variables are evaluated in the data frame $x$ using the usual non-standard evaluation rules.
drop	logical indicating if levels that do not occur should be dropped (if f is a factor or a list).
layers	Names of layers to include in the split; pass NA for all layers; pass NULL for the default layer
	Ignored

## Value

Returns a v5 assay with splitted layers

## See Also

```
v3 Assay object, validity, and interaction methods: $.Assay(), Assay-class, Assay-validity, CreateAssayObject(), [.Assay(), [.Assay(), dim.Assay(), dimnames.Assay(), merge.Assay(), subset.Assay()
```

split.Assay5

Split an Assay

# Description

Split an Assay

# Usage

```
## S3 method for class 'Assay5'
split(
    x,
    f,
    drop = FALSE,
    layers = c("counts", "data"),
    ret = c("assay", "multiassays", "layers"),
    ...
)
```

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## Arguments

x	An Assay5 object
f	a 'factor' in the sense that <code>as.factor(f)</code> defines the grouping, or a list of such factors in which case their interaction is used for the grouping. If $x$ is a data frame, $f$ can also be a formula of the form $\sim g$ to split by the variable $g$ , or more generally of the form $\sim g1 + \ldots + gk$ to split by the interaction of the variables $g1, \ldots, gk$ , where these variables are evaluated in the data frame $x$ using the usual non-standard evaluation rules.
drop	logical indicating if levels that do not occur should be dropped (if $f$ is a factor or a list).
layers	Names of layers to include in the split; pass NA for all layers; pass NULL for the default layer
ret	Type of return value; choose from:
	• "assay": a single Assay5 object
	• "multiassay": a list of Assay5 objects
	"layers": a list of layer matrices
	Ignored

#### Value

Depends on the value of ret:

- "assay": x with the layers requested in layers split based on f; all other layers are left as-is
- "multiassay": a list of Assay5 objects; the list contains one value per split and each assay contains only the layers requested in layers with the key set to the split
- "layers": a list of matrices of length length(assays) \* length(unique(f)); the list is named as "layer.split"

#### **Progress Updates with progressr**

This function uses **progressr** to render status updates and progress bars. To enable progress updates, wrap the function call in with\_progress or run handlers(global = TRUE) before running this function. For more details about **progressr**, please read vignette("progressr-intro")

#### See Also

```
v5 Assay object, validity, and interaction methods: $.Assay5(), Assay5-class, Assay5-validity, [.Assay5(), [[.Assay5(), dim.Assay5(), dimnames.Assay5(), merge.Assay5(), subset.Assay5()
```

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Stdev

Get the standard deviations for an object

## **Description**

Get the standard deviations for an object

## Usage

```
Stdev(object, ...)
## S3 method for class 'DimReduc'
Stdev(object, ...)
## S3 method for class 'Seurat'
Stdev(object, reduction = "pca", ...)
```

## **Arguments**

object An object

... Arguments passed to other methods

reduction Name of reduction to use

#### Value

The standard deviations

#### **Examples**

```
# Get the standard deviations for each PC from the DimReduc object
Stdev(object = pbmc_small[["pca"]])
# Get the standard deviations for each PC from the Seurat object
Stdev(object = pbmc_small, reduction = "pca")
```

StitchMatrix

Stitch Matrices Together

## **Description**

Stitch Matrices Together

## Usage

```
StitchMatrix(x, y, rowmap, colmap, ...)
```

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## **Arguments**

x A matrix

y One or more matrices of the same class or coercible to the same class as x

rowmap, colmap LogMaps describing the row and cell membership of each matrix; the LogMap entries are assumed to be in the order of c(x, y)

... Arguments passed to other methods

#### Value

A single matrix of type class(x) consisting of all values in component matrices

subset. Assay Subset an Assay

#### **Description**

Subset an Assay

## Usage

```
## S3 method for class 'Assay'
subset(x, cells = NULL, features = NULL, ...)
```

# Arguments

```
x An Assay objectcells Cell namesfeatures Feature names... Ignored
```

## Value

x with just the cells and features specified by cells and features

#### See Also

```
v3 Assay object, validity, and interaction methods: $.Assay(), Assay-class, Assay-validity, CreateAssayObject(), [.Assay(), [.Assay(), dim.Assay(), dimnames.Assay(), merge.Assay(), split.Assay()
```

```
rna <- pbmc_small[["RNA"]]
rna2 <- subset(rna, features = VariableFeatures(rna))
rna2</pre>
```

subset.Assay5

subset.Assay5

Subset an Assay

## **Description**

Subset an Assay

## Usage

```
## S3 method for class 'Assay5'
subset(x, cells = NULL, features = NULL, layers = NULL, ...)
```

# Arguments

x An Assay5 object

cells Cell names
features Feature names

layers Layer to keep; defaults to all layers

... Ignored

#### Value

x with just the cells and features specified by cells and features for the layers specified by layers

## See Also

```
v5 Assay object, validity, and interaction methods: $.Assay5(), Assay5-class, Assay5-validity, [.Assay5(), [[.Assay5(), dim.Assay5(), dimnames.Assay5(), merge.Assay5(), split.Assay5()
```

subset.DimReduc

Subset a Dimensional Reduction

# Description

Subset a DimReduc object

## Usage

```
## S3 method for class 'DimReduc'
subset(x, cells = NULL, features = NULL, ...)
```

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#### **Arguments**

```
x A DimReduc object cells, features

Cells and features to keep during the subset

Ignored
```

#### Value

x for cells cells and features features

#### See Also

Dimensional reduction object, validity, and interaction methods CreateDimReducObject(), DimReduc-class, DimReduc-validity, [.DimReduc(), [[.DimReduc(), dim.DimReduc(), merge.DimReduc(), print.DimReduc()

subset.Seurat

Subset Seurat Objects

#### **Description**

Subset Seurat Objects

#### Usage

```
## S3 method for class 'Seurat'
subset(
    x,
    subset,
    cells = NULL,
    features = NULL,
    idents = NULL,
    return.null = FALSE,
    ...
)

## S3 method for class 'Seurat'
x[i, j, ...]
```

#### **Arguments**

```
x A Seurat object
subset Logical expression indicating features/variables to keep
cells, j A vector of cell names or indices to keep
features, i A vector of feature names or indices to keep
idents A vector of identity classes to keep
return.null If no cells are requested, return a NULL; by default, throws an error
... Arguments passed to WhichCells
```

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## Value

```
subset: A subsetted Seurat object
[: object x with features i and cells j
```

#### See Also

```
WhichCells
```

```
Seurat object, validity, and interaction methods $.Seurat(), Seurat-class, Seurat-validity,
[[.Seurat(), [[<-,Seurat, NULL, [[<-,Seurat, dim.Seurat(), dimnames.Seurat(), merge.Seurat(),
names.Seurat()</pre>
```

## **Examples**

```
# `subset` examples
subset(pbmc_small, subset = MS4A1 > 4)
subset(pbmc_small, subset = `DLGAP1-AS1` > 2)
subset(pbmc_small, idents = '0', invert = TRUE)
subset(pbmc_small, subset = MS4A1 > 3, slot = 'counts')
subset(pbmc_small, features = VariableFeatures(object = pbmc_small))
# `[` examples
pbmc_small[VariableFeatures(object = pbmc_small), ]
pbmc_small[, 1:10]
```

Theta

Get the offset angle

# Description

Get the offset angle

## Usage

Theta(object)

#### **Arguments**

object An object

Tool

Tool

Get and Set Additional Tool Data

#### **Description**

Use Tool to get tool data. If no additional arguments are provided, will return a vector with the names of tools in the object.

#### Usage

```
Tool(object, ...)
Tool(object, ...) <- value
## S3 method for class 'Seurat'
Tool(object, slot = NULL, ...)
## S3 replacement method for class 'Seurat'
Tool(object, ...) <- value</pre>
```

## Arguments

object	An object
	Arguments passed to other methods
value	Information to be added to tool list
slot	Name of tool to pull

#### Value

If no additional arguments, returns the names of the tools in the object; otherwise returns the data placed by the tool requested

#### Note

For developers: set tool data using Tool<-. Tool<- will automatically set the name of the tool to the function that called Tool<-, so each function gets one entry in the tools list and cannot overwrite another function's entry. The automatic naming will also remove any method identifiers (eg. RunPCA. Seurat will become RunPCA); please plan accordingly

```
# Example function that adds unstructured data to tools
MyTool <- function(object) {
   sample.tool.output <- matrix(rnorm(n = 16), nrow = 4)
   # Note: `Tool<-` must be called from within a function
   # and the name of the tool will be generated from the function name
   Tool(object) <- sample.tool.output</pre>
```

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```
return(object)
}

# Run our tool
set.seed(42L)
pbmc_small <- MyTool(pbmc_small)

# Get a list of tools run
Tool(pbmc_small)

# Access specific tool data
Tool(pbmc_small, slot = "MyTool")</pre>
```

UpdateSeuratObject

Update old Seurat object to accommodate new features

# Description

Updates Seurat objects to new structure for storing data/calculations. For Seurat v3 objects, will validate object structure ensuring all keys and feature names are formed properly.

## Usage

```
UpdateSeuratObject(object)
```

## **Arguments**

object

Seurat object

#### Value

Returns a Seurat object compatible with latest changes

```
## Not run:
updated_seurat_object = UpdateSeuratObject(object = old_seurat_object)
## End(Not run)
```

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UpdateSlots

Update slots in an object

# Description

Update slots in an object

## Usage

```
UpdateSlots(object)
```

## Arguments

object

An object to update

# Value

object with the latest slot definitions

Version

Get Version Information

# Description

Get Version Information

# Usage

```
Version(object, ...)
## S3 method for class 'Seurat'
Version(object, ...)
```

# Arguments

object An object

... Arguments passed to other methods

```
Version(pbmc_small)
```

WhichCells 121

WhichCells

Identify cells matching certain criteria

## **Description**

Returns a list of cells that match a particular set of criteria such as identity class, high/low values for particular PCs, etc.

# Usage

```
WhichCells(object, ...)
## S3 method for class 'Assay'
WhichCells(object, cells = NULL, expression, invert = FALSE, ...)
## S3 method for class 'Seurat'
WhichCells(
  object,
  cells = NULL,
  idents = NULL,
  expression,
  slot = "data",
  invert = FALSE,
  downsample = Inf,
  seed = 1,
  ...
)
```

# Arguments

object	An object
	Arguments passed on to CellsByIdentities
	return.null If no cells are requested, return a NULL; by default, throws an error
cells	Subset of cell names
expression	A predicate expression for feature/variable expression, can evaluate anything that can be pulled by FetchData; please note, you may need to wrap feature names in backticks (``) if dashes between numbers are present in the feature name
invert	Invert the selection of cells
idents	A vector of identity classes to keep
slot	Slot to pull feature data for
downsample	Maximum number of cells per identity class, default is Inf; downsampling will happen after all other operations, including inverting the cell selection
seed	Random seed for downsampling. If NULL, does not set a seed

[.Assay

## Value

A vector of cell names

## See Also

FetchData

## **Examples**

```
WhichCells(pbmc_small, idents = 2)
WhichCells(pbmc_small, expression = MS4A1 > 3)
levels(pbmc_small)
WhichCells(pbmc_small, idents = c(1, 2), invert = TRUE)
```

[.Assay

Layer Data

# Description

Get and set layer data

# Usage

```
## S3 method for class 'Assay'
x[i = missing_arg(), j = missing_arg(), ...]
## S4 replacement method for signature 'Assay, character, ANY, ANY'
x[i, j, ...] <- value</pre>
```

# Arguments

X	An Assay object
i	Name of layer data to get or set
j	Ignored
	Arguments passed to LayerData
value	A matrix-like object to add as a new layer

## Value

```
[: The layer data for layer i
[<-: x with layer data value saved as i
```

[.Assay5

## See Also

```
LayerData
```

```
v3 Assay object, validity, and interaction methods: $.Assay(), Assay-class, Assay-validity, CreateAssayObject(), [[.Assay(), dim.Assay(), dimnames.Assay(), merge.Assay(), split.Assay(), subset.Assay()
```

# **Examples**

```
rna <- pbmc_small[["RNA"]]
# Get a vector of layer names in this assay
rna[]
# Fetch layer data
rna["data"][1:10, 1:4]
# Set layer data
rna["data"] <- rna["counts"]
rna["data"][1:10, 1:4]</pre>
```

[.Assay5

Layer Data

## **Description**

Get and set layer data

# Usage

```
## S3 method for class 'Assay5'
x[i = missing_arg(), j = missing_arg(), ...]
## S4 replacement method for signature 'Assay5, character, ANY, ANY'
x[i, j, ...] <- value</pre>
```

## **Arguments**

X	An Assay5 object
i	Name of layer data to get or set
j	Ignored
	Arguments passed to LayerData
value	A matrix-like object to add as a new layer

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## Value

```
[: The layer data for layer i
[<-: x with layer data value saved as i
```

#### See Also

#### LayerData

```
v5 Assay object, validity, and interaction methods: $.Assay5(), Assay5-class, Assay5-validity, [[.Assay5(), dim.Assay5(), dimnames.Assay5(), merge.Assay5(), split.Assay5(), subset.Assay5()
```

[.DimReduc

Get Feature Loadings

#### **Description**

Pull feature loadings from a dimensional reduction

#### **Usage**

```
## S3 method for class 'DimReduc'
x[i, j, drop = FALSE, ...]
```

#### **Arguments**

Х	A DimReduc object
i	Feature identifiers or indices
j	Dimension identifiers or indices
drop	Coerce the result to the lowest possible dimension; see drop for further details
	Arguments passed to other methods

#### **Details**

[ does not distinguish between projected and unprojected feature loadings; to select whether projected or unprojected loadings should be pulled, please use Loadings

#### Value

Feature loadings for features i and dimensions j

#### See Also

## Loadings

```
Dimensional reduction object, validity, and interaction methods CreateDimReducObject(), DimReduc-class, DimReduc-validity, [[.DimReduc(), dim.DimReduc(), merge.DimReduc(), print.DimReduc(), subset.DimReduc()
```

[.SeuratCommand 125

## **Examples**

```
pca <- pbmc_small[["pca"]]
pca[1:10, 1:5]</pre>
```

[.SeuratCommand

Command Log Data Access

# Description

Access data from a SeuratCommand object

# Usage

```
## S3 method for class 'SeuratCommand' x[i, \ldots]
```

# Arguments

x A SeuratCommand object

i The name of a command log slot

... Ignored

## Value

[: Slot i from x

## See Also

Command log object and interaction methods \$. SeuratCommand(), .DollarNames.SeuratCommand(), LogSeuratCommand(), SeuratCommand-class, as.list.SeuratCommand()

```
cmd <- pbmc_small[["NormalizeData.RNA"]]
cmd["call.string"]</pre>
```

[[.Assay

[[.Assay

Feature-Level Meta Data

#### **Description**

Get and set feature-level meta data

#### Usage

```
## S3 method for class 'Assay'
x[[i, ..., drop = FALSE]]

## S4 replacement method for signature 'Assay,ANY,ANY,ANY'
x[[i, j, ...]] <- value

## S3 method for class 'Assay'
head(x, n = 10L, ...)

## S3 method for class 'Assay'
tail(x, n = 10L, ...)

## S4 replacement method for signature 'Assay,missing,missing,data.frame'
x[[i, j, ...]] <- value</pre>
```

#### **Arguments**

X	An Assay object
i	Name of feature-level meta data to fetch or add
	Ignored
drop	See drop
j	Ignored
value	Feature-level meta data to add
n	Number of meta data rows to show

#### Value

```
[[: The feature-level meta data for i
[[<-: x with value added as i in feature-level meta data
head: The first n rows of feature-level meta data
tail: the last n rows of feature-level meta data
```

#### See Also

```
v3 Assay object, validity, and interaction methods: $.Assay(), Assay-class, Assay-validity, CreateAssayObject(), [.Assay(), dim.Assay(), dimnames.Assay(), merge.Assay(), split.Assay(), subset.Assay()
```

[[.Assay5

## **Examples**

```
rna <- pbmc_small[["RNA"]]

# Pull the entire feature-level meta data frame
head(rna[[]])

# Pull a specific column of feature-level meta data
head(rna[["vst.mean"]])
head(rna[["vst.mean", drop = TRUE]])

# `head` and `tail` can be used to quickly view feature-level meta data
head(rna)

tail(rna)</pre>
```

[[.Assay5

Feature-Level Meta Data

## **Description**

Get and set feature-level meta data

#### Usage

```
## S3 method for class 'Assay5'
x[[i, j, ..., drop = FALSE]]

## S4 replacement method for signature 'Assay5,ANY,ANY,ANY'
x[[i, j, ...]] <- value

## S3 method for class 'Assay5'
head(x, n = 10L, ...)

## S3 method for class 'Assay5'
tail(x, n = 10L, ...)</pre>
```

## **Arguments**

X	An Assay5 object
i	Name of feature-level meta data to fetch or add
j	Ignored
• • •	Ignored
drop	See drop
value	Feature-level meta data to add
n	Number of meta data rows to show

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#### Value

```
[[: The feature-level meta data for i
```

[[<-: x with value added as i in feature-level meta data

head: The first n rows of feature-level meta data

tail: the last n rows of feature-level meta data

#### See Also

```
v5 Assay object, validity, and interaction methods: $.Assay5(), Assay5-class, Assay5-validity, [.Assay5(), dim.Assay5(), dimnames.Assay5(), merge.Assay5(), split.Assay5(), subset.Assay5()
```

[[.DimReduc

Get Cell Embeddings

#### **Description**

Pull cell embeddings from a dimensional reduction

#### Usage

```
## S3 method for class 'DimReduc'
x[[i, j, drop = FALSE, ...]]
```

## Arguments

X	A DimReduc object
i	Cell names or indices

j Dimension identifiers or indices

drop Coerce the result to the lowest possible dimension; see drop for further details

... Arguments passed to other methods

#### Value

Cell embeddings for cells i and dimensions j

#### See Also

#### **Embeddings**

```
Dimensional reduction object, validity, and interaction methods CreateDimReducObject(), DimReduc-class, DimReduc-validity, [.DimReduc(), dim.DimReduc(), merge.DimReduc(), print.DimReduc(), subset.DimReduc()
```

```
pca <- pbmc_small[["pca"]]
pca[[1:10, 1:5]]</pre>
```

[[.Seurat 129

[[.Seurat

Subobjects and Cell-Level Meta Data

#### **Description**

The [[ operator pulls either subobjects (eg. v3 or v5 assays, dimensional reduction information, or nearest-neighbor graphs) or cell-level meta data from a Seurat object

#### Usage

```
## $3 method for class 'Seurat'
x[[i = missing_arg(), ..., drop = FALSE, na.rm = FALSE]]
## $3 method for class 'Seurat'
head(x, n = 10L, ...)
## $3 method for class 'Seurat'
tail(x, n = 10L, ...)
```

#### **Arguments**

X	A Seurat object
i	Name of cell-level meta data
	Ignored
drop	See drop
na.rm	Remove cells where meta data is all NA
n	Number of meta data rows to show

#### Value

Varies based on the value of i:

- If i is missing, a data frame with cell-level meta data
- If i is a vector with cell-level meta data names, a data frame (or vector of drop = TRUE) with cell-level meta data requested
- If i is a one-length character with the name of a subobject, the subobject specified by i

head: The first n rows of cell-level metadata tail: The last n rows of cell-level metadata

#### See Also

See here for adding meta data with [[<-, here for adding subobjects with [[<-, and here for removing subobjects and cell-level meta data with [[<-

```
Seurat object, validity, and interaction methods $.Seurat(), Seurat-class, Seurat-validity,
[[<-,Seurat,NULL,[[<-,Seurat,dim.Seurat(),dimnames.Seurat(),merge.Seurat(),names.Seurat(),
subset.Seurat()</pre>
```

[[<-,Seurat

#### **Examples**

```
# Get the cell-level metadata data frame
head(pbmc_small[[]])

# Pull specific metadata information
head(pbmc_small[[c("letter.idents", "groups")]])
head(pbmc_small[["groups", drop = TRUE]])

# Get a sub-object (eg. an `Assay` or `DimReduc`)
pbmc_small[["RNA"]]
pbmc_small[["pca"]]

# Get the first 10 rows of cell-level metadata
head(pbmc_small)

# Get the last 10 rows of cell-level metadata
tail(pbmc_small)
```

[[<-,Seurat</pre>

Add Subobjects

## Description

Add subobjects containing expression, dimensional reduction, or other containerized data to a Seurat object. Subobjects can be accessed with [[ and manipulated directly within the Seurat object or used independently

#### Usage

```
## S4 replacement method for signature 'Seurat,character,missing,Assay'
x[[i, j, ...]] <- value

## S4 replacement method for signature 'Seurat,character,missing,Assay5'
x[[i, j, ...]] <- value

## S4 replacement method for signature 'Seurat,character,missing,DimReduc'
x[[i, j, ...]] <- value

## S4 replacement method for signature 'Seurat,character,missing,Graph'
x[[i, j, ...]] <- value

## S4 replacement method for signature 'Seurat,character,missing,Neighbor'
x[[i, j, ...]] <- value

## S4 replacement method for signature 'Seurat,character,missing,Neighbor'
x[[i, j, ...]] <- value</pre>
```

[[<-,Seurat,NULL 131

```
## S4 replacement method for signature 'Seurat, character, missing, Spatial I mage' x[[i, j, ...]] \leftarrow value
```

## Arguments

```
x A Seurat object
i Name to add subobject as
j Ignored
... Ignored
```

value A valid subobject (eg. a v3 or v5 assay, or a dimensional reduction)

#### Value

x with value added as i

#### See Also

See here for pulling subobjects using [[, here for adding metadata with [[<-, and here for removing subobjects and cell-level meta data with [[<-

```
Seurat object, validity, and interaction methods $.Seurat(), Seurat-class, Seurat-validity,
[[.Seurat(), [[<-, Seurat, NULL, dim.Seurat(), dimnames.Seurat(), merge.Seurat(), names.Seurat(),
subset.Seurat()</pre>
```

[[<-,Seurat,NULL R

Remove Subobjects and Cell-Level Meta Data

## **Description**

Remove Subobjects and Cell-Level Meta Data

## Usage

```
## S4 replacement method for signature 'Seurat, character, missing, `NULL`' x[[i, j, ...]] \leftarrow value
```

# Arguments

х	A Seurat object
i	Name(s) of $subobject(s)$ or cell-level meta data to remove
j	Ignored
• • •	Ignored
value	NULL

## Value

x with i removed from the object

\$.Assay

#### See Also

See here for pulling subobjects using [[, here for adding metadata with [[<-, and here for adding subobjects with [[<-

Seurat object, validity, and interaction methods \$.Seurat(), Seurat-class, Seurat-validity,
[[.Seurat(), [[<-, Seurat, dim.Seurat(), dimnames.Seurat(), merge.Seurat(), names.Seurat(),
subset.Seurat()</pre>

\$.Assay

Layer Data

#### **Description**

Get and set layer data

#### Usage

```
## S3 method for class 'Assay'
x$i
## S3 replacement method for class 'Assay'
x$i <- value</pre>
```

# **Arguments**

x An Assay object

i Name of layer data to get or set

value A matrix-like object to add as a new layer

#### Value

```
$: Layer data for layer i
```

\$<-: x with layer data value saved as i</pre>

#### See Also

```
v3 Assay object, validity, and interaction methods: Assay-class, Assay-validity, CreateAssayObject(), [.Assay(), [[.Assay(), dim.Assay(), dimnames.Assay(), merge.Assay(), split.Assay(), subset.Assay()
```

\$.Assay5

## **Examples**

```
rna <- pbmc_small[["RNA"]]
# Fetch a layer with `$`
rna$data[1:10, 1:4]
# Add a layer with `$`
rna$data <- rna$counts
rna$data[1:10, 1:4]</pre>
```

\$.Assay5

Layer Data

# Description

Get and set layer data

# Usage

```
## S3 method for class 'Assay5'
x$i
## S3 replacement method for class 'Assay5'
x$i <- value</pre>
```

# **Arguments**

x An Assay5 object

i Name of layer data to get or set

value A matrix-like object to add as a new layer

#### Value

```
$: Layer data for layer i
```

\$<-: x with layer data value saved as i</pre>

## See Also

```
v5 Assay object, validity, and interaction methods: Assay5-class, Assay5-validity, [.Assay5(), [[.Assay5(), dim.Assay5(), dimnames.Assay5(), merge.Assay5(), split.Assay5(), subset.Assay5()
```

\$.Seurat

\$.Seurat

Cell-Level Meta Data

#### **Description**

Get and set cell-level meta data

## Usage

```
## S3 method for class 'Seurat'
x$i

## S3 replacement method for class 'Seurat'
x$i, ... <- value

## S4 replacement method for signature 'Seurat, character, missing, data. frame'
x[[i, j, ...]] <- value

## S4 replacement method for signature 'Seurat, missing, missing, data. frame'
x[[i, j, ...]] <- value

## S4 replacement method for signature 'Seurat, character, missing, factor'
x[[i, j, ...]] <- value

## S4 replacement method for signature 'Seurat, character, missing, list'
x[[i, j, ...]] <- value

## S4 replacement method for signature 'Seurat, missing, missing, list'
x[[i, j, ...]] <- value

## S4 replacement method for signature 'Seurat, character, missing, vector'
x[[i, j, ...]] <- value</pre>
```

#### **Arguments**

X	A Seurat object
i	Name of cell-level meta data
	Ignored
value	A vector to add as cell-level meta data
j	Ignored

## Value

\$: Metadata column i for object x; **note**: unlike [[, \$ drops the shape of the metadata to return a vector instead of a data frame

```
$<-: x with metadata value saved as i</pre>
```

\$.SeuratCommand 135

#### See Also

# Examples

```
# Get metadata using `$'
head(pbmc_small$groups)

# Add metadata using the `$' operator
set.seed(42)
pbmc_small$value <- sample(1:3, size = ncol(pbmc_small), replace = TRUE)
head(pbmc_small[["value"]])</pre>
```

\$.SeuratCommand

Command Log Parameter Access

# Description

Pull parameter values from a SeuratCommand object

#### Usage

```
## S3 method for class 'SeuratCommand' x$i
```

## **Arguments**

- x A SeuratCommand object
- i A parameter name

#### Value

The value for parameter i

#### See Also

```
Command log object and interaction methods .DollarNames.SeuratCommand(), LogSeuratCommand(), SeuratCommand-class, [.SeuratCommand(), as.list.SeuratCommand()
```

```
cmd <- pbmc_small[["NormalizeData.RNA"]]
cmd$normalization.method</pre>
```

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