

Package ‘imcExperiment’

August 19, 2021

Title Mass Cytometry S4 Class Structure Pipeline for Images

Version 0.99.0

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Description

Containerizes cytometry data and allows for S4 class structure to extend slots related to cell morphology, spatial coordinates, phenotype network information, and unique cellular labeling.

Depends R (>= 4.0), SingleCellExperiment, methods

Imports SummarizedExperiment, S4Vectors, spatstat.geom, stats

Suggests knitr, rmarkdown

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biocViews Software, WorkflowStep, MultipleComparison

Encoding UTF-8

RoxygenNote 7.1.1

VignetteBuilder knitr

NeedsCompilation no

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Repository CRAN

Date/Publication 2021-08-19 09:20:02 UTC

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`.checkSpatialDimension`

the rows are the panel names, the columns are the single cells, the column are the single cells to match the SCE designs (scRNA)

Description

the rows are the panel names, the columns are the single cells, the column are the single cells to match the SCE designs (scRNA)

Usage

`.checkSpatialDimension(object)`

Arguments

object imcExperiment object, class imcExperiment container

Value

imcExperiment container that has proper dimensions

`.imcExperimentToPPP` *map to point pattern from imcExperiment class.*

Description

map to point pattern from imcExperiment class.

Usage

`.imcExperimentToPPP(caseExperiment = NULL, phenotypeToUse = 1)`

Arguments

caseExperiment the subset IMC experiment to cast into a point pattern

phenotypeToUse the cluster id to annotate the pattern

Value

imcExperiment container converted to a point pattern set

cellIntensity	<i>finds the intensities getter.</i>
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Description

finds the intensities getter.

sets cell Intensity slot to a new matrix. rows protein, columns are cells.

Usage

```
cellIntensity(object, ...)

## S4 method for signature 'imcExperiment'
cellIntensity(object)

cellIntensity(object) <- value

## S4 replacement method for signature 'imcExperiment,matrix'
cellIntensity(object) <- value
```

Arguments

object	IMC container
...	additional arguments
value	matrix rows protein, columns are cells

Value

imcExperiment container
imcExperiment container
imcExperiment container
imcExperiment container

Examples

```
data(imcdata)
dim(cellIntensity(imcdata))
data(imcdata);dim(cellIntensity(imcdata))
head(t(cellIntensity(imcdata)))
data(imcdata)
x<-asinh(counts(imcdata))
cellIntensity(imcdata)<-x
```

data

data

Description

Data set containing 1,000 cells and 73 features which include panel antibody, neighborhood computations, and phenograph clustering.

Usage

```
data(data)
```

Format

A data frame of 1,000 cells and histoCAT features

ImageId feature from histoCAT

CellId feature from histoCAT

marker1 feature from histoCAT

marker2 feature from histoCAT

marker3 feature from histoCAT

marker4 feature from histoCAT

marker5 feature from histoCAT

marker6 feature from histoCAT

marker7 feature from histoCAT

marker8 feature from histoCAT

marker9 feature from histoCAT

marker10 feature from histoCAT

marker11 feature from histoCAT

marker12 feature from histoCAT

marker13 feature from histoCAT

marker14 feature from histoCAT

marker15 feature from histoCAT

marker16 feature from histoCAT

marker17 feature from histoCAT

marker18 feature from histoCAT

marker19 feature from histoCAT

marker20 feature from histoCAT

marker21 feature from histoCAT

marker22 feature from histoCAT

marker23 feature from histoCAT
marker24 feature from histoCAT
marker25 feature from histoCAT
marker26 feature from histoCAT
marker27 feature from histoCAT
marker28 feature from histoCAT
marker29 feature from histoCAT
marker30 feature from histoCAT
marker31 feature from histoCAT
marker32 feature from histoCAT
marker33 feature from histoCAT
marker34 feature from histoCAT
Area feature from histoCAT
Eccentricity feature from histoCAT
Solidity feature from histoCAT
Extent feature from histoCAT
EulerNumber feature from histoCAT
Perimeter feature from histoCAT
MajorAxisLength feature from histoCAT
MinorAxisLength feature from histoCAT
Orientation feature from histoCAT
X_position feature from histoCAT
Y_position feature from histoCAT
Percent_Touching feature from histoCAT
Number_Neighbors feature from histoCAT
neighbour_4_CellId1 feature from histoCAT
neighbour_4_CellId2 feature from histoCAT
neighbour_4_CellId3 feature from histoCAT
neighbour_4_CellId4 feature from histoCAT
neighbour_4_CellId5 feature from histoCAT
neighbour_4_CellId6 feature from histoCAT
neighbour_4_CellId7 feature from histoCAT
neighbour_4_CellId8 feature from histoCAT
neighbour_4_CellId9 feature from histoCAT
neighbour_4_CellId10 feature from histoCAT
Phenograph7851534969 feature from histoCAT
tSNE4148542692_1 feature from histoCAT
tSNE4148542692_2 feature from histoCAT

getCoordinates *finds the spatial coords, getter.*

Description

finds the spatial coords, getter.

Usage

```
getCoordinates(object)

## S4 method for signature 'imcExperiment'
getCoordinates(object)

## S4 replacement method for signature 'imcExperiment,matrix'
getCoordinates(object) <- value
```

Arguments

object	is IMC container
value	matrix rows cells, columns are x,y

Value

imcExperiment container
imcExperiment container
imcExperiment container

Examples

```
data(imcdata)
getCoordinates(imcdata)
data(imcdata)
getCoordinates(imcdata)
data(imcdata)
x<-getCoordinates(imcdata)
getCoordinates(imcdata)<-as.matrix(x)
```

getCoordinates<- *Sets the coordinate positions of each cell (matrix), columns are X,Y positions.*

Description

Sets the coordinate positions of each cell (matrix), columns are X,Y positions.

Usage

```
getCoordinates(object) <- value
```

Arguments

object	is IMC container
value	matrix rows cells, columns are x,y

Value

imcExperiment container

Examples

```
data(imcdata)
x<-getCoordinates(imcdata)
getCoordinates(imcdata)<-as.matrix(x)
```

getDistance<- *re-assigns the distance matrix (rows are cells)*

Description

re-assigns the distance matrix (rows are cells)

Usage

```
getDistance(object) <- value
```

Arguments

object	is IMC container
value	matrix rows cells, columns are distance measurements

Value

imcExperiment container

Examples

```
data(imcdata)
newD<-matrix(1,nrow=ncol(imcdata),ncol=1)
getDistance(imcdata)<-newD
```

getMorphology<- *re-assigns morphological features can be stored (matrix) rows are cells and columns are Area, etc.*

Description

re-assigns morphological features can be stored (matrix) rows are cells and columns are Area, etc.

Usage

```
getMorphology(object) <- value
```

Arguments

object	is IMC container
value	matrix rows cells, columns are Area, Eccentricity, etc.

Value

imcExperiment container

Examples

```
data(imcdata)
x<-matrix(1,nrow=ncol(imcdata),ncol=4)
getMorphology(imcdata)<-x
```

getNeighborhood *finds the neighborhood information.*

Description

finds the neighborhood information.

slow assignment for the histoCAT neighborhood data (matrix) columns are the neighbors

Usage

```
getNeighborhood(object, ...)  
  
## S4 method for signature 'imcExperiment'  
getNeighborhood(object)  
  
getNeighborhood(object) <- value  
  
## S4 replacement method for signature 'imcExperiment,matrix'  
getNeighborhood(object) <- value
```

Arguments

object	is IMC container
...	additional arguments
value	matrix rows cells, columns are neighborhood histoCAT output

Value

imcExperiment container
imcExperiment container data(imcdata) getNeighborhood(imcdata)
imcExperiment container
imcExperiment container

Examples

```
data(imcdata)  
getNeighborhood(imcdata)  
data(imcdata)  
x<-matrix(1,nrow=ncol(imcdata),ncol=2)  
getNeighborhood(imcdata)<-x  
data(imcdata)  
x<-matrix(1,nrow=ncol(imcdata),ncol=2)  
getNeighborhood(imcdata)<-x
```

getNetwork<- *re-assigns the network assignment (matrix)*

Description

re-assigns the network assignment (matrix)

Usage

```
getNetwork(object) <- value
```

Arguments

object	is IMC container
value	data.frame rows cells, columns are phenograph network ID

Value

imcExperiment container

Examples

```
data(imcdata)
x<-data.frame(ID=seq_len(ncol(imcdata)))
getNetwork(imcdata)<-x
```

imcdata	<i>imcdata</i>
---------	----------------

Description

histoCAT output containerized as IMC container. IMC S4 data set containing 2,452 cells and 44 antibody features which include panel antibody.

Usage

```
data(imcdata)
```

Format

A data frame of 2,452 cells and histoCAT features which are containerized into the imcExperiment

imcExperiment	<i>Initializes a imcExperiment and performs some rudimentary checks. Many of the arguments CAN be NULL; determination of which is required is done at run-time. A imcExperiment must contain at least the expressions and spatial/coordinate assays.</i>
---------------	--

Description

Initializes a imcExperiment and performs some rudimentary checks. Many of the arguments CAN be NULL; determination of which is required is done at run-time. A imcExperiment must contain at least the expressions and spatial/coordinate assays.

Usage

```
imcExperiment(
  coordinates = matrix(1, 3, 3),
  cellIntensity = matrix(1, 3, 3),
  neighborHood = matrix(1, 3, 3),
  network = data.frame(matrix(1, 3, 3)),
  distance = matrix(1, 3, 3),
  morphology = matrix(1, 3, 3),
  uniqueLabel = rep("A", 3),
  panel = as.character(seq_len(3)),
  ROIID = data.frame(ROIID = rep("A", 3)),
  ...
)
```

Arguments

coordinates	matrix of spatial coordinates (x,y)
cellIntensity	matrix of counts
neighborHood	neighborhood results
network	network assignments for each cell
distance	distances for each cell, can be square
morphology	morphology features for each cell, can be square
uniqueLabel	character class each cell is assigned a uniqueLabel
panel	antibody panel rownames set to rowData
ROIID	character for ROI
...	additional arguments

Value

imcExperiment container

Examples

```
x<-imcExperiment(cellIntensity=matrix(1,nrow=10,ncol=10),
  coordinates=matrix(1,nrow=10,ncol=2),
  neighborHood=matrix(1,nrow=10,ncol=10),
  network=data.frame(matrix(1,nrow=10,ncol=10)),
  distance=matrix(1,nrow=10,ncol=10),
  morphology=matrix(1,nrow=10,ncol=10),
  uniqueLabel=paste0("A",seq_len(10)),
  panel=letters[1:10],
  ROIID=data.frame(ROIID=rep("A",10)))
```

```
imcExperiment-class  a summarized experiment of IMC runs, dimensions of the spatial and
intensity data are regulated.#'
```

Description

a summarized experiment of IMC runs, dimensions of the spatial and intensity data are regulated.#'

finds the network information.

assigns cell cluster assignment to the container. rows are cells and column is the cluster ID

finds the distance information.

distance matrix can be stored in the distance slot for pairwise distance

finds the morphology information.

morphological features can be stored (matrix) rows are cells and columns are Area, etc.

finds the label information.

unique cell labels can be assigned (vector)

Usage

```
getNetwork(object)

## S4 method for signature 'imcExperiment'
getNetwork(object)

## S4 replacement method for signature 'imcExperiment,data.frame'
getNetwork(object) <- value

getDistance(object)

## S4 method for signature 'imcExperiment'
getDistance(object)

## S4 replacement method for signature 'imcExperiment,matrix'
getDistance(object) <- value

getMorphology(object)

## S4 method for signature 'imcExperiment'
getMorphology(object)

## S4 replacement method for signature 'imcExperiment,matrix'
getMorphology(object) <- value

getLabel(object)
```

```
## S4 method for signature 'imcExperiment'  
getLabel(object)
```

Arguments

object	imcExperiment
value	matrix rows cells, columns are Area, etc.

Value

imcExperiment container
imcExperiment container
imcExperiment container
imcExperiment container
imcExperiment container
imcExperiment container
imcExperiment container
imcExperiment container
imcExperiment container
imcExperiment container
imcExperiment container

Slots

coordinates matrix class containing x,y coordinates
cellIntensity matrix class containing intensity
neighborHood matrix class containing x,y neighbor
network data frame class containing network
distance matrix class containing x,y distances
morphology matrix class containing morphology
uniqueLabel labels

Examples

```
x<-imcExperiment(cellIntensity=matrix(1,nrow=10,ncol=10),  
coordinates=matrix(1,nrow=10,ncol=2),  
neighborHood=matrix(1,nrow=10,ncol=10),  
network=data.frame(matrix(1,nrow=10,ncol=10)),  
distance=matrix(1,nrow=10,ncol=10),  
morphology=matrix(1,nrow=10,ncol=10),  
uniqueLabel=paste0("A",seq_len(10)),  
panel=letters[1:10],  
ROIID=data.frame(ROIID=rep("A",10)))  
data(imcdata)  
getNetwork(imcdata)
```

```

data(imcdata)
getNetwork(imcdata)
data(imcdata)
x<-data.frame(ID=seq_len(ncol(imcdata)))
getNetwork(imcdata)<-x
data(imcdata)
getDistance(imcdata)
data(imcdata)
getDistance(imcdata)
data(imcdata)
newD<-matrix(1,nrow=ncol(imcdata),ncol=1)
getDistance(imcdata)<-newD
data(imcdata)
getMorphology(imcdata)
data(imcdata)
getMorphology(imcdata)
data(imcdata)
x<-matrix(1,nrow=ncol(imcdata),ncol=4)
getMorphology(imcdata)<-x
data(imcdata)
getLabel(imcdata)
data(imcdata)
getLabel(imcdata)

```

imcExperimentToHyperFrame

map to point pattern from imcExperiment class.

Description

map to point pattern from imcExperiment class.

Usage

```
imcExperimentToHyperFrame(imcExperiment = NULL, phenotypeToUse = 1)
```

Arguments

`imcExperiment` imcExperiment class
`phenotypeToUse` the network slot can often have many columns, this is the ID for the column number to use in the network slot.

Value

a hyperframe of point patterns

Examples

```

data(imcdata)
H<-imcExperimentToHyperFrame(imcExperiment=imcdata,phenotypeToUse = 1)

```

percentilenormalize *given a matrix of intensity counts, perform min/max norm.*

Description

given a matrix of intensity counts, perform min/max norm.

Usage

```
percentilenormalize(data = NULL, percentile = NULL)
```

Arguments

data matrix of numeric data only
percentile numeric value 0.99 default.

Value

normalized data, each column on [0,1] scale.

Examples

```
data(data)
dim(data)
expr<-data[,3:36]
normExp<-percentilenormalize(data=expr,percentile=0.99)
normExp<-as.matrix(normExp)
```

selectCases *subsets the imcExperiment to a case along with all slots for a selected multiple ROIs.*

Description

subsets the imcExperiment to a case along with all slots for a selected multiple ROIs.
method to subset the slots, requires colData with column "ROIID"

Usage

```
selectCases(object, value, ...)
```

S4 method for signature 'imcExperiment'
selectCases(object, value)

Arguments

object	IMC container
value	this is ROIID vector
...	additional parameters

Value

imcExperiment container of selected cases
imcExperiment container of selected cases

Examples

```
data(imcdata)
myCases<-selectCases(imcdata,c("30-BM16-202_7Pre_s1_p1_r4_a4_ac", "B17_350_14post_s1_p1_r5_a5_ac"))
myCases
table(colData(myCases)$ROIID)
data(imcdata)
myCases<-selectCases(imcdata,c("30-BM16-202_7Pre_s1_p1_r4_a4_ac", "B17_350_14post_s1_p1_r5_a5_ac"))
myCases
table(colData(myCases)$ROIID)
```

subsetCase	<i>subsets the imcExperiment to a case along with all slots for a single ROI, using for distance analysis</i>
------------	---

Description

subsets the imcExperiment to a case along with all slots for a single ROI, using for distance analysis method to subset the slots, requires colData with column "ROIID"

Usage

```
subsetCase(object, value, ...)

## S4 method for signature 'imcExperiment'
subsetCase(object, value)
```

Arguments

object	IMC container
value	this is ROIID a single character ID
...	additional parameters

Value

returns IMC object of a single case
roi imcExperiment

Examples

```
data(imcdata)
myCase<-subsetCase(imcdata,"30-BM16-202_7Pre_s1_p1_r4_a4_ac")
myCase
data(imcdata)
myCase<-subsetCase(imcdata,"30-BM16-202_7Pre_s1_p1_r4_a4_ac")
myCase
```

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