## Package 'MoleculeExperiment'

October 24, 2025

**Title** Prioritising a molecule-level storage of Spatial Transcriptomics

Data

Version 1.9.0

Description MoleculeExperiment contains functions to create and work with objects from the new MoleculeExperiment class. We introduce this class for analysing molecule-based spatial transcriptomics data (e.g., Xenium by 10X, Cosmx SMI by Nanostring, and Merscope by Vizgen). This allows researchers to analyse spatial transcriptomics data at the molecule level, and to have standardised data formats accross vendors.

```
License MIT + file LICENSE
Encoding UTF-8
Roxygen list(markdown = TRUE)
RoxygenNote 7.3.1
URL https://github.com/SydneyBioX/MoleculeExperiment
BugReports https://github.com/SydneyBioX/MoleculeExperiment/issues
Imports SpatialExperiment, Matrix, purrr, data.table, dplyr (>=
     1.1.1), magrittr, rjson, utils, methods, terra, ggplot2, rlang,
     cli, EBImage, rhdf5, BiocParallel, S4Vectors, stats
Suggests knitr, BiocStyle, testthat (>= 3.0.0)
VignetteBuilder knitr
biocViews DataImport, DataRepresentation, Infrastructure, Software,
     Spatial, Transcriptomics
Config/testthat/edition 3
Depends R (>= 2.10)
git url https://git.bioconductor.org/packages/MoleculeExperiment
git_branch devel
git_last_commit 3dd8051
git_last_commit_date 2025-04-15
Repository Bioconductor 3.23
```

2 .generateBPParam

## **Date/Publication** 2025-10-24

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## Description

Utility function to generate BPPARAM object.

## Usage

```
.generateBPParam(cores = 1)
```

## Arguments

cores

Desired number of cores for BPPARAM object.

## Value

A BPPPARAM object.

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accessors	Accessor functions to work with MoleculeExperiment objects

#### **Description**

Accessor functions to work with MoleculeExperiment objects

### Usage

```
## S4 method for signature 'MoleculeExperiment'
molecules(x, assayName = NULL, flatten = FALSE)

## S4 method for signature 'MoleculeExperiment'
boundaries(object, assayName = NULL, flatten = FALSE)

## S4 method for signature 'MoleculeExperiment'
features(object, assayName = NULL)

## S4 method for signature 'MoleculeExperiment'
segmentIDs(object, assayName = NULL)

## S4 replacement method for signature 'MoleculeExperiment'
molecules(x, assayName = NULL) <- value

## S4 replacement method for signature 'MoleculeExperiment'
boundaries(object, assayName = NULL) <- value</pre>
```

## Arguments

Character string specifying the name of the assay from which to retrieve or set information in the slot of interest.  Logical value specifying whether to flatten the ME list into a data.frame or not. Defaults to FALSE.  The MoleculeExperiment to access.  value  New value to be added to the slot and assay of interest.	X	The Molecule Experiment to access.
Defaults to FALSE.  object The MoleculeExperiment to access.	assayName	
1	flatten	
value New value to be added to the slot and assay of interest.	object	The MoleculeExperiment to access.
	value	New value to be added to the slot and assay of interest.

#### Value

A MoleculeExperiment object slot.

#### getters

Accessor functions to get data from the MoleculeExperiment object. These include:

• molecules() to retrieve information from the molecules slot.

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- boundaries() to retrieve information from the boundaries slot.
- features() to retrieve feature names from the molecules slot.
- segmentIDs() to retrieve segment ids from the boundaries slot.

#### setters

The molecules<- setter accesses the molecules slot, whereas the boundaries slot can be accessed with boundaries<-.

```
# get example data
repoDir <- system.file("extdata", package = "MoleculeExperiment")</pre>
repoDir <- paste0(repoDir, "/xenium_V1_FF_Mouse_Brain")</pre>
me <- readXenium(repoDir,</pre>
   keepCols = "essential".
   addBoundaries = "cell"
)
# get insight into MoleculeExperiment object (e.g., see assay names)
# get insight into molecules slot (e.g., see the assay names)
showMolecules(me)
# for developers, use molecules() getter
# expect a large output from call below
# molecules(me, assayName = "detected")
# alternatively, return rectangular data structure with flatten = TRUE
molecules(me, assayName = "detected", flatten = TRUE)
# get insight into boundaries slot (e.g., see the assay names)
showBoundaries(me)
# for developers, use boundaries() getter
# expect a large output from call below
# boundaries(me, assayName = "cell")
# alternatively, return rectangular data structure with flatten = TRUE
boundaries(me, assayName = "cell", flatten = TRUE)
# features() getter
features(me, assayName = "detected")
# segmentIDs() getter
segmentIDs(me, assayName = "cell")
# setter example
# read in and standardise nucleus boundaries too
nucleiMEList <- readBoundaries(</pre>
   dataDir = repoDir,
   pattern = "nucleus_boundaries.csv",
   segmentIDCol = "cell_id",
```

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```
xCol = "vertex_x",
yCol = "vertex_y",
keepCols = "essential",
boundariesAssay = "nucleus",
scaleFactorVector = 1
)

# use `boundaries<-` setter to add nucleus boundaries to the boundaries slot
boundaries(me, assayName = "nucleus") <- nucleiMEList
me</pre>
```

bufferBoundaries

Create a new boundaries assay with buffers

#### Description

This function takes in an existing MoleculeExperiment object and generates a new boundaries assay with added buffers. This can be useful for visualisation and for countMolecules.

#### Usage

```
bufferBoundaries(me, assayName = "cell", ...)
```

#### **Arguments**

me A MoleculeExperiment object.

assayName Character string (default "cell") specifying the existing boundaries assay that should have buffer added.

... Arguments that pass to internal functions. The most relevant parameter is buffer (default 0).

#### Value

A boundaries assay with essential columns and vertices with added buffer.

```
repoDir <- system.file("extdata", package = "MoleculeExperiment")
repoDir <- paste0(repoDir, "/xenium_V1_FF_Mouse_Brain")
me <- readXenium(repoDir,
   keepCols = "essential"
)
MoleculeExperiment::boundaries(me, "cell_buffer") <- bufferBoundaries(
   me,
   assayName = "cell", buffer = 1
)
library(ggplot2)
ggplot_me() +</pre>
```

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```
geom_polygon_me(me, assayName = "cell", fill = "grey") +
geom_polygon_me(me, assayName = "cell_buffer", fill = NA, colour = "red") +
geom_point_me(me) +
coord_cartesian(
    xlim = c(4900, 4919.98),
    ylim = c(6400.02, 6420)
)
```

countMolecules

Count molecules per region of interest (e.g., cell)

#### **Description**

This function takes the information from the molecules and boundaries slot, and counts the molecules per region of interest. Its input is a MoleculeExperiment object, and its output a SpatialExperiment object. That way, if one is interested in doing downstream analyses at the cell level, one can do so.

#### Usage

```
countMolecules(
   me,
   moleculesAssay = "detected",
   boundariesAssay = "cell",
   buffer = 0,
   matrixOnly = FALSE,
   nCores = 1
)
```

### **Arguments**

me

MoleculeExperiment object containing both the transcript data as well as the boundaries data. I.e., the "molecules" and "boundaries" slots need to be filled. See MoleculeExperiment() for more information.

moleculesAssay

Character string naming the list of the molecules slot from which transcript information should be retrieved from. The default is the detected transcript data that is read in when creating a MoleculeExperiment object. It is possible to change it to another mode, e.g., "high\_threshold" will access the transcript information that has been stored in the "high\_threshold" element of the list in the molecules slot.

boundariesAssay

Character string naming the list of the boundaries slot from which boundary information should be retrieved from. For example, for counting transcripts per cell, the list containing the cell boundaries (e.g., "cell") should be selected.

buffer

Single numeric value (default 0) indicating value to buffer beyond segment boundaries, i.e. to count molecules just outside of a segment boundary

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matrixOnly Logical value indicating whether to return a matrix of the counted molecules per

segment (e.g., cell). Is FALSE by default, i.e., the default output is a SpatialEx-

periment object.

nCores Number of cores to use for the operation.

#### Value

A SpatialExperiment object derived from a MoleculeExperiment object. Alternatively, a matrix with the counted molecules per segment.

## **Examples**

dataframeToMEList

Convert a transcript (molecule) or boundary dataframe to the ME list format

## Description

The goal of this function is to standardise transcript and boundary files for input to a MoleculeExperiment object.

### Usage

```
dataframeToMEList(
   df,
   dfType = NULL,
   assayName = NULL,
   sampleCol = "sample_id",
   factorCol = NULL,
   xCol = "x_location",
   yCol = "y_location",
   keepCols = "essential",
   scaleFactor = 1
)
```

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## **Arguments**

df	A data frame containing the transcript information or the boundary information. NOTE: this dataframe should, at a minimum, have the following 4 columns: sample_id, factorCol (e.g., feature_id in transcripts, or cell_id in boundaries), x_location and y_location.
dfType	Character string specifying contents of the dataframe. Can be either "molecules" or "boundaries".
assayName	Character string specifying the name with which to identify the information later on in an ME object.
sampleCol	Character string specifying the name of the column with the sample id.
factorCol	Character string specifying the name of the column with the factors with which to group the data in the lists. When working with molecules, this column would be e.g., "feature_id" in xenium. When working with boundaries, this column would be e.g., "cell_id" in xenium.
xCol	Character string specifying the name of the column with global x coordinates.
yCol	Character string specifying the name of the column with global y coordinates.
keepCols	Character string which can be either "essential" or "all". If "essential", the function will only work with the x and y location information.
scaleFactor	Integer specifying the scale factor by which to change the scale of the $x$ and $y$ locations (e.g., to change from pixel to micron). The default value is 1.

#### Value

A list with the format required to input it into slots of a MoleculeExperiment object.

## **Examples**

molecules MEList

MoleculeExperiment-class

MoleculeExperiment class: An S4 class container to store imaging-based spatial transcriptomics data.

#### Description

This class enables the analysis of imaging-based ST data at the molecule level, and standardises data across vendors. The aim of this class is to facilitate ST data integration and comparison and, importantly, facilitate common analytical and visualisation workflows.

#### Usage

```
MoleculeExperiment(molecules, boundaries = NULL)
```

#### **Arguments**

molecules Detected transcripts information in a standardised ME list format, as is generated

by dataframeToMEList() and readMolecules() functions.

boundaries Slot with boundary information in a standardised ME list format, as is generated

by dataframeToMEList() and readBoundaries() functions.

#### Value

A MoleculeExperiment object

#### **Slots**

molecules Slot containing information about the detected transcripts. This slot is designed as a list of lists, where each sample contains a list of tibbles with information for each gene. The basic information required for this slot are the gene names of the transcripts, as well as their x and y locations.

boundaries Slot containing the boundaries defining each segmented cell. The slot is designed as a list of lists, where each sample contains a list of tibbles for each cell, consisting of the x and y coordinates of the polygon vertices defining the cell boundary.

```
# creating a simple ME object from toy data
moleculesDf <- data.frame(
    sample_id = rep(c("sample1", "sample2"), times = c(30, 20)),
    features = rep(c("gene1", "gene2"), times = c(20, 30)),
    x_coords = runif(50),
    y_coords = runif(50)
)
boundariesDf <- data.frame(
    sample_id = rep(c("sample1", "sample2"), times = c(16, 6)),
    cell_id = rep(c("cell1", "cell2", "cell3", "cell4",</pre>
```

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```
"cell1", "cell2"),
                   times = c(4, 4, 4, 4, 3, 3)),
    vertex_x = rnorm(22),
    vertex_y = rnorm(22)
)
moleculesMEList <- dataframeToMEList(moleculesDf,</pre>
                                    dfType = "molecules",
                                    assayName = "detected",
                                    sampleCol = "sample_id",
                                    factorCol = "features",
                                    xCol = "x_coords",
                                   yCol = "y_coords")
boundariesMEList <- dataframeToMEList(boundariesDf,</pre>
                                     dfType = "boundaries",
                                     assayName = "cell",
                                     sampleCol = "sample_id",
                                     factorCol = "cell_id",
                                     xCol = "vertex_x",
                                     yCol = "vertex_y")
toyME <- MoleculeExperiment(molecules = moleculesMEList,</pre>
                              boundaries = boundariesMEList)
toyME
```

plotting-functions

Plotting functions for SpatialUtils

## **Description**

A set of ggplot functions to build customized plots for imaging based spatial transcriptomics data.

## Usage

```
ggplot_me()
geom_point_me(
    me,
    assayName = "detected",
    byColour = NULL,
    selectFeatures = NULL,
    ...
)
geom_polygon_me(me, assayName = "cell", byFill = NULL, ...)
geom_raster_img(
```

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```
path = NULL,
image = NULL,
displacement = c(0, 0),
pixelSize = 1,
...
```

## Arguments

me	MoleculeExperiment object.
assayName	Character string specifying name of assay from which to get data.
byColour	Character string specifying the column name to colour by.
selectFeatures	character vector of features to keep for geom_point_me.
• • •	Additional parameters to be passed to ggplot.
byFill	Character string specifying the column name to fill by.
path	Path of the image. Default: NULL
image	Image object to be plotted as raster. Default: NULL
displacement	the x-y coordinate of the top-left pixel of the image. Default: $c(0,0)$
pixelSize	the pixel size in micron, Default: 1

#### Value

A plot with transcripts and/or segmentation information for imaging based spatial transcriptomics data.

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readBoundaries

Read in csv boundary information and convert to ME list format.

#### **Description**

This function reads in csv boundary files and converts them to the ME list format, so that they can be added to an ME object later on. To account for different coordinate scales possible being used by the boundary versus transcript information, this function scales the coordinate values of the boundaries to match the unit of the detected transcript locations. The various arguments offer flexibility to standardise data from different molecule-based ST technologies into the ME list format.

#### Usage

```
readBoundaries(
  dataDir,
  pattern = NULL,
  segmentIDCol = NULL,
  xCol = NULL,
  yCol = NULL,
  keepCols = "essential",
  boundariesAssay = NULL,
  scaleFactorVector = 1
)
```

#### Arguments

dataDir Path of the directory containing the boundary csv files.

pattern Character string specifying the unique pattern with which to identify the files of

interest in the directory. This is useful to work with multiple samples. Defaults

to NULL.

segment IDCol Character string specifying the name of the column containing the segment IDs.

Defaults to NULL.

xCol Character string specifying the name of the column containing the x coordinates

of the vertices defining the boundaries. Defaults to NULL.

yCol Character string specifying the name of the column containing the y coordinates

of the vertices defining the boundaries. Defaults to NULL.

keepCols Character string specifying which columns to keep. Defaults to "essential". The

other option is to select "all", or custom columns by specifying their names in a

vector.

boundariesAssay

Character string specifying the name with which to identify the boundary data in the ME object later on. Defaults to NULL.

scaleFactorVector

Vector containing the scale factor/s with which to change the coordinate data from pixel to micron. It can be either a single integer, or multiple scale factors for the different samples. The default value is 1.

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#### Value

An ME list containing the boundary information. This can be used as input to the boundaries slot of an ME object.

## **Examples**

readCosmx

Read in Cosmx data (Nanostring) as an ME object.

## **Description**

This function is a wrapper around the readMolecules function. It can read both molecule and mask information. The segmentation masks are converted to boundaries, and these are added to the boundaries slot of the MoleculeExperiment object.

#### Usage

```
readCosmx(dataDir, keepCols = "essential", addBoundaries = "cell")
```

## **Arguments**

dataDir Character string specifying the directory with the Cosmx output files.

keepCols Character string specifying which columns to keep. Defaults to "essential". The

other option is to select "all", or custom columns by specifying their names in a

vector.

addBoundaries A string with which to specify the name of the boundary assay to be added to

the me object. Can be a string, or NULL. If NULL, a simple ME object with no

boundaries will be created.

#### Value

A MoleculeExperiment object

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#### **Examples**

readMerscope

Read in Merscope data to an ME object

#### **Description**

Reads in Merscope (Vizgen) molecule and boundary data from a directory, and standardises it into a MoleculeExperiment object.

#### Usage

```
readMerscope(dataDir, keepCols = "essential", addBoundaries = "cell")
```

#### **Arguments**

dataDir

Character string specifying the directory with the Cosmx output files.

keepCols

Vector of characters specifying the columns of interest from the transcripts file. "essential" selects columns with gene names, x and y locations. "all" will select all columns. Alternatively, specific columns of interest can be selected by specifying them as characters in a vector. Note that this personalised vector needs to

contain the essential columns.

addBoundaries

A string with which to specify the name of the boundary assay to be added to the me object. Can be a string, or NULL. If NULL, a simple ME object with no

boundaries will be created.

#### Value

A MoleculeExperiment object

```
repoDir <- system.file("extdata", package = "MoleculeExperiment")
repoDir <- paste0(repoDir, "/vizgen_HumanOvarianCancerPatient2Slice2")
meMerscope <- readMerscope(repoDir,
    keepCols = "essential",
    addBoundaries = "cell"
)
meMerscope</pre>
```

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readMolecules	Read in detected transcripts file/s into a	a MoleculeExperiment object
---------------	--	-----------------------------

## Description

A function to standardise transcripts.csv files across different molecule- based ST technologies, and store them into an ME object. It is technology agnostic, so it is accompanied with wrappers for the specific technologies (e.g., see readXenium).

## Usage

```
readMolecules(
  dataDir,
  pattern = NULL,
  featureCol = NULL,
  xCol = NULL,
  yCol = NULL,
  keepCols = "essential",
  moleculesAssay = NULL,
  scaleFactorVector = 1
)
```

#### **Arguments**

dataDir	Character string specifying the directory with the file/s containing detected transcripts for different runs/samples.
pattern	Character string specifying the pattern with which to find the transcripts files. For example, in Xenium data, the pattern would be "transcripts.csv". In contrast, in Cosmx data, the pattern would be "tx_file".
featureCol	Character string specifying the name of the column with feature names. For example, "feature_name" in xenium transcripts.csv files.
xCol	Character string specifying the name of the column with the x locations of the transcripts.
yCol	Character string specifying the name of the column with the y locations of the transcripts.
keepCols	Vector of characters specifying the columns of interest from the transcripts file. "essential" selects columns with gene names, x and y locations. "all" will select all columns. Alternatively, specific colums of interest can be selected by specifying them as characters in a vector. Note that this personalised vector needs to contain the essential columns.
moleculesAssay	Character string specifying the name of the list in which the transcript informa-

tected transcript information.

tion is going to be stored in the molecules slot. The default name is "detected", as we envision that a MoleculeExperiment will usually be created with raw de-

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scaleFactorVector

Vector containing the scale factor/s with which to change the coordinate data from pixel to micron. It can be either a single integer, or multiple scale factors for the different samples. The default value is 1.

#### Value

A simple MoleculeExperiment object with a filled molecules slot.

#### **Examples**

```
repoDir <- system.file("extdata", package = "MoleculeExperiment")
repoDir <- paste0(repoDir, "/xenium_V1_FF_Mouse_Brain")
simple_me <- readMolecules(repoDir,
    pattern = "transcripts.csv",
    featureCol = "feature_name",
    xCol = "x_location",
    yCol = "y_location",
    keepCols = "essential"
)
simple_me</pre>
```

readSegMask

Read a segmentation mask

## **Description**

Reads a segmentation mask TIFF and transforms it into a ME boundaries object. One must provide either the path or the loaded image object.

#### Usage

```
readSegMask(
  extent,
  path = NULL,
  image = NULL,
  assayName = "cell",
  background_value = NULL,
  sample_id = NULL
)
```

#### **Arguments**

extent The extent of the loaded segmentation mask in micrometers. Used to align the mask with the transcripts. This must be of the form c(xmin, xmax, ymin, ymax).

path The path of the segmenation mask, Default: NULL

image The loaded image object, Default: NULL

assayName The name of the segmentation (e.g. cell, or nucleus), Default: 'cell'

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background\_value

The value corresponding to the background in the segmentation, Default: NULL

sample\_id What the sample should be named, Default: NULL

#### Value

A boundaries object.

#### **Examples**

```
repoDir <- system.file("extdata", package = "MoleculeExperiment")</pre>
segMask <- paste0(repoDir, "/BIDcell_segmask.tif")</pre>
data <- paste0(repoDir, "/xenium_V1_FF_Mouse_Brain/sample1")</pre>
me <- readXenium(data,</pre>
   keepCols = "essential",
    addBoundaries = NULL
)
boundaries(me, "BIDcell_segmentation") <- readSegMask(</pre>
    # use the molecule extent to define the boundary extent
   extent(me, assayName = "detected"),
    path = segMask, assayName = "BIDcell_segmentation",
    sample_id = "sample1", background_value = 0
)
ggplot_me() +
   geom_polygon_me(
        me,
        assayName = "BIDcell_segmentation", fill = NA, colour = "black"
    geom_point_me(me, byColour = "feature_id", size = 0.1) +
   geom_polygon_me(
        me.
        assayName = "BIDcell_segmentation", fill = NA, colour = "red"
    )
```

readXenium

Read in Xenium data into a MoleculeExperiment object

#### **Description**

Function to read in, and standardise, Xenium output data into an ME object. Detected transcripts files are required. Additionally, it is also possible to read in boundary files ("cell", "nuclei", or both). This function is a wrapper around readMolecules and readBoundaries functions.

#### Usage

```
readXenium(dataDir, keepCols = "essential", addBoundaries = "cell")
```

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#### **Arguments**

dataDir Character string specifying the directory with the xenium output files.

keepCols Vector of characters specifying the columns of interest from the transcripts file

and boundaries file. Can be "all" or "essential". "essential" selects columns with gene names, x and y locations in the transcripts file; "essential" selects columns with cell ids, and x and y locations for the vertices defining the boundaries in

the boundaries file.

addBoundaries Vector with which to specify the names of the boundary assays to be added to

the me object. Can be "cell", "nucleus", both, or NULL. The latter will lead to

the creation of a simple ME object with just the molecules slot filled.

#### Value

A MoleculeExperiment object containing xenium data.

#### **Examples**

 $small\_me$ 

A subsetted Xenium dataset include for demostration purposes.

#### **Description**

A subsetted Xenium dataset include for demostration purposes.

#### Usage

```
data(small_me)
```

#### Format

An object of class MoleculeExperiment of length 1.

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subset	hv	extent
Subset	υv	extent

Subset functions for MoleculeExperiment objects

#### **Description**

A set of functions to subset MoleculeExperiment objects by different factors

#### Usage

```
subset_by_extent(me, extent)
```

#### **Arguments**

me

MoleculeExperiment object.

extent

The extent in micrometers to subset the me object. This must be of the form

c(xmin, xmax, ymin, ymax).

#### Value

A subsetted MoleculeExperiment object

#### **Examples**

```
data(small_me)
subset_extent <- c(xmin = 3000, xman = 4000, ymin = 2000, ymax = 3000)
subset_small_me <- subset_by_extent(small_me, subset_extent)
# check the extent after subsetting
extent(subset_small_me, assayName = "detected")</pre>
```

summarization

Summarization methods to get insights into a MoleculeExperiment object

## Description

The following methods are useful to get quick view of the contents in a MoleculeExperiment object. For example, showMolecules and showBoundaries summarise the large nested ME list of lists in the molecules and boundaries slots. nFeatures and nTranscripts get the numbers of features or transcripts, respectively.

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#### Usage

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

## S4 method for signature 'MoleculeExperiment'
showMolecules(object)

## S4 method for signature 'MoleculeExperiment'
showBoundaries(object)

## S4 method for signature 'MoleculeExperiment'
extent(object, assayName = NULL)

## S4 method for signature 'MoleculeExperiment'
nFeatures(object)

## S4 method for signature 'MoleculeExperiment'
nTranscripts(object)
```

#### Arguments

object Name of MoleculeExperiment object of interest.

assayName Character string specifying the name of the assay from which to view a summary

of the contents.

#### Value

A MoleculeExperiment object summary.

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