# Package 'Cardinal'

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**Description** Implements statistical & computational tools for analyzing mass spectrometry imaging datasets, including methods for efficient pre-processing, spatial segmentation, and classification.

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2 Contents

# **Contents**

Cardinal-package	3
bin	4
colocalized	5
deprecated	7
estimateDomain	7
features	8
findNeighbors	9
MassDataFrame-class	10
MeansTest	11
MSImagingArrays-class	13
MSImagingExperiment-class	15
normalize	17
peakAlign	18
peakPick	20
peakProcess	22
pixels	24
plot-image	25
plot-spectra	27
PositionDataFrame-class	29
process	30
readMSIData	32
recalibrate	34
reduceBaseline	35
ResultsList-class	37
selectROI	38
simulateSpectra	39
slice	42
smooth	43
SpatialCV	45
SpatialDGMM	46
SpatialFastmap	49
SpatialKMeans	51
SpatialNMF	53
SpatialPCA	54
SpatialPLS	56
SpatialResults-class	58
SpatialShrunkenCentroids	60
spatialWeights	63
SpectraArrays-class	65
SpectralImagingArrays-class	
SpectralImagingData-class	66
	67
SpectralImagingExperiment-class	69
spectrapply	70
subsetFeatures	72
summarizeFeatures	73 75
WILLENGALLIZE	17

Index		XDataFrame-class	76
	Index		78

3

Cardinal-package Mass spectrometry imaging tools

### **Description**

Cardinal-package

Implements statistical & computational tools for analyzing mass spectrometry imaging datasets, including methods for efficient pre-processing, spatial segmentation, and classification.

#### **Details**

Cardinal provides an abstracted interface to manipulating mass spectrometry imaging datasets, simplifying most of the basic programmatic tasks encountered during the statistical analysis of imaging data. These include image manipulation and processing of both images and mass spectra, and dynamic plotting of both.

While pre-processing steps including normalization, baseline correction, and peak-picking are provided, the core functionality of the package is statistical analysis. The package includes classification and clustering methods based on nearest shrunken centroids, as well as traditional tools like PCA and PLS.

Type browseVignettes("Cardinal") to view a user's guide and vignettes of common workflows.

### **Options**

The following options can be set:

- getCardinalgetCardinalBPPARAM(), setCardinalBPPARAM(BPPARAM=NULL): The default backend to use for parallel processing. By default, this is initially set to NULL (no parallelization). Otherwise, it must be a BiocParallelParam instance. See documentation for bplapply.
- getCardinalVerbose(), setCardinalVerbose(verbose=interactive()): Should progress messages be printed?
- getCardinalNChunks(), setCardinalNChunks(n=20L): The default number of data chunks used when iterating over large datasets. Used by many methods internally.

Additionally, graphical parameters can be set:

- vizi\_style(): Set the default plotting style.
- vizi\_engine(): Set the default plotting engine.
- vizi\_par(): Set default graphical parameters.

# Author(s)

Kylie A. Bemis

Maintainer: Kylie A. Bemis <k.bemis@northeastern.edu>

4 bin

bin

Bin spectra

# **Description**

Apply on-the-fly binning to spectra.

### Usage

```
## S4 method for signature 'MSImagingExperiment'
bin(x, ref,
    spectra = "intensity", index = "mz",
    method = c("sum", "mean", "max", "min",
        "linear", "cubic", "gaussian", "lanczos"),
    resolution = NA, units = c("ppm", "mz"), ...)
## S4 method for signature 'MSImagingArrays'
bin(x, ref,
    spectra = "intensity", index = "mz",
    method = c("sum", "mean", "max", "min",
        "linear", "cubic", "gaussian", "lanczos"),
    resolution = NA, units = c("ppm", "mz"), \ldots)
## S4 method for signature 'SpectralImagingExperiment'
bin(x, ref,
    spectra = "intensity", index = NULL,
    method = c("sum", "mean", "max", "min",
        "linear", "cubic", "gaussian", "lanczos"),
    resolution = NA, units = c("relative", "absolute"),
    verbose = getCardinalVerbose(), ...)
## S4 method for signature 'SpectralImagingArrays'
bin(x, ref,
    spectra = "intensity", index = NULL,
    method = c("sum", "mean", "max", "min",
        "linear", "cubic", "gaussian", "lanczos"),
    resolution = NA, units = c("relative", "absolute"),
    verbose = getCardinalVerbose(), ...)
```

### Arguments

x A spectral imaging dataset.

ref Optional. The bin centers, or their range if resolution is specified. Created from resolution if not provided.

spectra The name of the array in spectraData() to bin.

colocalized 5

column in featureData() (for SpectralImagingExperiment) to use for the

bins.

method The peak picking method to use. See approx1 for details.

resolution The bin widths.

units The units for the above resolution.
verbose Should progress messages be printed?

... Ignored.

#### **Details**

The binning is applied but not processed immediately. It is performed on-the-fly whenever the spectra are accessed.

#### Value

A new object derived from SpectralImagingExperiment with the binned spectra.

### Author(s)

Kylie A. Bemis

#### See Also

approx1, estimateDomain, estimateReferenceMz

# **Examples**

```
set.seed(1)
mse <- simulateImage(preset=1, npeaks=10, dim=c(3,3))
# bin to unit resolution
mse2 <- bin(mse, resolution=1, units="mz")
# bin to a specific range and resolution
mse3 <- bin(mse, ref=c(800, 1000), resolution=1, units="mz")</pre>
```

colocalized

Colocalized features

### Description

Find colocalized features in an imaging dataset.

6 colocalized

### Usage

```
## S4 method for signature 'MSImagingExperiment'
colocalized(object, mz, ...)
## S4 method for signature 'SpectralImagingExperiment'
colocalized(object, i, ref,
    threshold = median, n = Inf,
    sort.by = c("cor", "MOC", "M1", "M2", "Dice", "none"),
    nchunks = getCardinalNChunks(),
    verbose = getCardinalVerbose(),
    BPPARAM = getCardinalBPPARAM(), ...)
## S4 method for signature 'SpatialDGMM'
colocalized(object, ref,
    threshold = median, n = Inf,
    sort.by = c("MOC", "M1", "M2", "Dice", "none"),
    nchunks = getCardinalNChunks(),
    verbose = getCardinalVerbose(),
    BPPARAM = getCardinalBPPARAM(), ...)
```

### **Arguments**

object	An imaging experiment.
mz	An m/z value of a feature in object to use as a reference.
i	The index of a feature in object to use as a reference.
ref	Either a flattened image (i.e., a numeric vector) or a logical mask of a region-of-interest to use as a reference.
threshold	Either a function that returns the cutoff to use for creating logical masks of numeric references, or a numeric threshold to use.
n	The number of top-ranked colocalized features to return.
sort.by	The colocalization measure used to rank colocalized features. Possible options include Pearson's correlation ("cor"), Manders overlap coefficient ("MOC"), Manders colocalization coefficients ("M1" and "M2"), and Dice similarity coefficient ("Dice".
nchunks	The number of chunks to use for processing.
verbose	Should progress messages be printed?
BPPARAM	An optional instance of BiocParallelParam. See documentation for bplapply.
	Options passed to chunkApply.

#### Value

A data frame with the colocalized features, or a list of data frames if multiple references are given.

# Author(s)

Kylie A. Bemis

deprecated 7

# **Examples**

```
set.seed(1)
x <- simulateImage(preset=1, dim=c(10,10), representation="centroid")
# find features colocalized with first feature
colocalized(x, i=1)
```

deprecated

Deprecated and defunct objects in Cardinal

### **Description**

These functions are provided for compatibility with older versions of Cardinal, and will be removed in the future.

estimateDomain

Estimate shared domain

### **Description**

For unaligned spectral data, it is often necessary to estimate a suitable shared domain in order to calculate statistical summaries like the mean spectrum.

```
estimateDomain(xlist,
    units = c("relative", "absolute"),
    nchunks = getCardinalNChunks(),
    verbose = getCardinalVerbose(),
    BPPARAM = getCardinalBPPARAM(), ...)

estimateReferenceMz(object,
    units = c("ppm", "mz"),
    nchunks = getCardinalNChunks(),
    verbose = getCardinalVerbose(),
    BPPARAM = getCardinalBPPARAM(), ...)

estimateReferencePeaks(object, SNR = 2,
    method = c("diff", "sd", "mad", "quantile", "filter", "cwt"),
    nchunks = getCardinalNChunks(),
    verbose = getCardinalVerbose(),
    BPPARAM = getCardinalBPPARAM(), ...)
```

8 features

#### **Arguments**

xlist	A list of the domain values (e.g., m/z values) for each spectrum.
object	A mass spectral imaging dataset.
units	Should the spacing between domain values use absolute or relative units?
method	The peak picking method to use. See findpeaks for details.
SNR	The signal-to-noise threshold to use to determine a peak.
nchunks	The number of chunks to use for processing.
verbose	Should progress messages be printed?
BPPARAM	$An optional instance of {\tt BiocParallelParam}. See documentation for {\tt bplapply}.$
	Options passed to chunkLapply.

#### **Details**

For estimateDomain, the domain is estimated by first finding the resolution of each spectrum's individual domain values (e.g., the spacing between m/z values), and then creating a sequence of domain values using the median resolution of all spectra.

The estimateReferenceMz function simply calls estimateDomain on the appropriate components of a mass spectral imaging dataset to estimate profile m/z bins.

The estimateReferencePeaks function calculates the mean spectrum (or looks for a "mean" column in featureData()) and performs peak picking on the mean spectrum. It can be used to create a set of reference peaks if all relevant peaks appear in the mean spectrum.

# Value

A vector of domain values, m/z values, or peaks.

### Author(s)

Kylie A. Bemis

### See Also

summarizeFeatures, recalibrate, peakAlign, peakProcess

features	Find feature indices
----------	----------------------

# **Description**

Search for the row indices of a spectral imaging dataset that correspond to specificor features, based on a set of conditions.

findNeighbors 9

### Usage

```
## S4 method for signature 'MSImagingExperiment'
features(object, ..., mz, tolerance = NA, units = c("ppm", "mz"),
        env = NULL)
## S4 method for signature 'SpectralImagingExperiment'
features(object, ..., env = NULL)
```

# Arguments

object A spectral imaging dataset.

... Expressions that evaluate to logical vectors in the environment of featureData().

mz The m/z values of features to include.

tolerance The tolerance for matching features to m/z values.

units The units for the above tolerance.

env The enclosing environment for evaluating . . . .

### Author(s)

Kylie A. Bemis

# **Examples**

```
set.seed(1)
mse <- simulateImage(preset=1, npeaks=10, dim=c(10,10))
features(mse, mz > 800, mz < 1800)
features(mse, mz=metadata(mse)$design$featureData$mz)</pre>
```

findNeighbors

Find spatial neighbors

### **Description**

Find the indices of spatial neighbors for all observations in a dataset.

```
## S4 method for signature 'ANY'
findNeighbors(x, r = 1, groups = NULL,
    metric = "maximum", p = 2, matrix = FALSE, ...)
## S4 method for signature 'SpectralImagingData'
findNeighbors(x, r = 1, groups = run(x), ...)
## S4 method for signature 'PositionDataFrame'
findNeighbors(x, r = 1, groups = run(x), ...)
```

10 MassDataFrame-class

### **Arguments**

X	An imaging dataset or data frame with spatial dimensions.
r	The spatial maximum distance for an observation to be considered a neighbor.
groups	A vector coercible to a factor giving which observations should be treated as spatially-independent. Observations in the same group are assumed to have a spatial relationship.
metric	Distance metric to use when finding the nearest neighbors. Supported metrics include "euclidean", "maximum", "manhattan", and "minkowski".
р	The power for the Minkowski distance.
matrix	Should the neighbors be returned as a sparse adjacency matrix instead of a list?
	Additional arguments passed to the next call.

### Value

Either a list of indices of neighbors or a sparse adjacency matrix (sparse\_mat).

# Author(s)

Kylie A. Bemis

# See Also

```
spatialWeights
```

# **Examples**

```
pdata <- PositionDataFrame(coord=expand.grid(x=1:8, y=1:8))
# find spatial neighbors
findNeighbors(pdata, r=1)</pre>
```

MassDataFrame-class

MassDataFrame: Extended data frame with key columns

# Description

A data frame for mass spectrometry feature metadata with a required column for m/z values.

```
MassDataFrame(mz, ..., row.names = FALSE)
```

MeansTest 11

### **Arguments**

mz A sorted vector of m/z values.

... Arguments passed to the DataFrame().

Either a vector of row names or a logical value indicating whether row names

should be generated automatically (from the m/z values).

### Methods

```
mz(object, ...), mz(object, ...) <- value: Get or set the m/z values.
```

#### Author(s)

Kylie A. Bemis

#### See Also

XDataFrame, PositionDataFrame

### **Examples**

```
## Create an MassDataFrame object
MassDataFrame(mz=sort(500 * runif(10)), label=LETTERS[1:10])
```

MeansTest

Linear model-based testing for summarized imaging experiments

### **Description**

Performs hypothesis testing for imaging experiments by fitting linear mixed models to summarizations or segmentations.

```
## S4 method for signature 'ANY'
meansTest(x, data, fixed, random, samples,
    response = "y", reduced = ~ 1, byrow = FALSE,
    nchunks = getCardinalNChunks(),
    verbose = getCardinalVerbose(),
    BPPARAM = getCardinalBPPARAM(), ...)

## S4 method for signature 'SpectralImagingExperiment'
meansTest(x, fixed, random, samples = run(x),
    response = "intensity", ...)

## S4 method for signature 'SpatialDGMM'
meansTest(x, fixed, random, class = 1L,
    response = "intensity", reduced = ~ 1,
```

12 MeansTest

# **Arguments**

x A dataset in P x N matrix format or a set of spatially segmented images.

data A data frame of additional variables parallel to x.

fixed A one-sided formula giving the fixed effects of the model on the RHS. The

response will added to the LHS, and the formula will be passed to the underlying

modeling function.

random A one-sided formula giving the random effects of the model on the RHS. See

1me for the allowed specifications.

samples A vector coercible to a factor giving the observational unit (i.e., the samples and

replicates).

class For SpatialDGMM, the class (segment) from the Gaussian mixture models that

should be used for the comparison. By default, compare the classes (segments)

with the highest means in each sample.

response The name of to assign the response variable in the fitted models.

reduced A one-sided formula specifying the reduced model for the null hypothesis. The

default is an intercept-only model.

byrow For the default method, are the rows or the columns the x.

nchunks The number of chunks to use for processing.

verbose Should progress messages be printed?

BPPARAM An optional instance of BiocParallelParam. See documentation for bplapply.

... Passed to internal linear modeling methods. Either 1m if only fixed effects are

given or lme if random effects are given.

object A fitted model object to summarize.

n, sort.by For topFeatures, the number of top features to return and how to sort them.

i The index of the model(s)/feature(s) to plot.

type The type of plot to display.

show.obs Should individual observations (i.e., the summarized mean for each sample) be

plotted too?

fill Should the boxplots be filled?

layout A vector of the form c(nrow, ncol) specifying the number of rows and columns

in the facet grid.

### Value

An object of class MeansTest derived from ResultsList, where each element contains a linear model.

### Author(s)

Dan Guo and Kylie A. Bemis

#### See Also

```
lm, lme, spatialDGMM
```

# **Examples**

MSImagingArrays-class MSImagingArrays: MS imaging data with arbitrary m/z values

#### **Description**

The MSImagingArrays class provides a list-like container for high-throughput mass spectrometry imaging data where every mass spectrum may have its own m/z values. It is designed for easy access to raw mass spectra for the purposes of pre-processing.

It can be converted to a MSImagingExperiment object for easier image slicing and for applying statistical models and machine learning methods.

```
## Instance creation
MSImagingArrays(spectraData = SimpleList(),
    pixelData = PositionDataFrame(), experimentData = NULL,
    centroided = NA, continuous = NA, metadata = list())
## Additional methods documented below
```

### Arguments

spectraData	Either a list-like object with lists of individual spectra and lists of their domain values, or a SpectraArrays instance.
pixelData	A PositionDataFrame with pixel metadata, with a row for each spectrum.
experimentData	Either NULL or a ImzMeta object with MS-specific experiment-level metadata.
centroided	A logical value indicated whether the spectra have been centroided.
continuous	A logical value indicated whether the spectra all have the same m/z values.
metadata	A list of arbitrary metadata.

#### Slots

spectraData: A SpectraArrays object storing one or more array-like data elements with conformable dimensions.

elementMetadata: A PositionDataFrame containing spectrum-level metadata, including each spectrum's pixel coordinates and experimental run information.

processing: A list containing unexecuted ProcessingStep objects.

experimentData: Either NULL or an ImzMeta object containing experiment-level metadata (necessary for writing the data to imzML).

centroided: A logical value indicated whether the spectra have been centroided (if known).

continuous: A logical value indicated whether the spectra all have the same m/z values (if known).

### Methods

All methods for SpectralImagingData and SpectralImagingArrays also work on MSImagingArrays objects. Additional methods are documented below:

```
mz(object, i = NULL, ...), mz(object, i = NULL, ...) <- value: Get or set the m/z arrays in
    the spectraData slot.

intensity(object, i = NULL, ...), intensity(object, i = NULL, ...) <- value: Get or set
    the intensity arrays in the spectraData slot.

centroided(object, ...), centroided(object, ...) <- value: Get or set the centroided slot.

isCentroided(object, ...): Equivalent to isTRUE(centroided(object)).

experimentData(object), experimentData(object) <- value: Get or set the experimentData
    slot.</pre>
```

### Author(s)

Kylie A. Bemis

#### See Also

SpectralImagingArrays, MSImagingExperiment

### **Examples**

```
set.seed(1)
x <- replicate(9, rlnorm(10), simplify=FALSE)
mz <- replicate(9, 500 * sort(runif(10)), simplify=FALSE)
coord <- expand.grid(x=1:3, y=1:3)

msa <- MSImagingArrays(
    spectraData=list(intensity=x, mz=mz),
    pixelData=PositionDataFrame(coord))</pre>
```

MSImagingExperiment-class

MSImagingExperiment: MS imaging data with shared m/z values

### **Description**

The MSImagingExperiment class provides a matrix-like container for high-throughput mass spectrometry imaging data where every mass spectrum shares the same m/z values. It is designed to provide easy access to both the spectra (as columns) and sliced images (as rows).

It can be converted from a MSImagingArrays object which is designed for representing raw mass spectra.

# Usage

```
## Instance creation
MSImagingExperiment(spectraData = SimpleList(),
    featureData = MassDataFrame(), pixelData = PositionDataFrame(),
    experimentData = NULL, centroided = NA, metadata = list())
## Additional methods documented below
```

#### **Arguments**

	spectraData	Either a matrix-like object with number of rows equal to the number of features and number of columns equal to the number of pixels, a list of such objects, or a SpectraArrays instance.
	featureData	A MassDataFrame with feature metadata, with a row for each feature.
pixelData A PositionDataFrame with pixel metadata, with a row for each spectrum		A PositionDataFrame with pixel metadata, with a row for each spectrum.
	${\tt experimentData}$	$Either\ NULL\ or\ a\ {\tt ImzMeta}\ object\ with\ MS-specific\ experiment-level\ metadata.$
	centroided	A logical value indicated whether the spectra have been centroided.
	metadata	A list of arbitrary metadata.

#### Slots

spectraData: A SpectraArrays object storing one or more array-like data elements with conformable dimensions.

featureData: A MassDataFrame containing feature-level metadata.

elementMetadata: A PositionDataFrame containing spectrum-level metadata, including each spectrum's pixel coordinates and experimental run information.

processing: A list containing unexecuted ProcessingStep objects.

experimentData: Either NULL or an ImzMeta object containing experiment-level metadata (necessary for writing the data to imzML).

centroided: A logical value indicated whether the spectra have been centroided (if known).

#### Methods

All methods for SpectralImagingData and SpectralImagingExperiment also work on MSImagingExperiment objects. Additional methods are documented below:

### Author(s)

Kylie A. Bemis

# See Also

SpectralImagingExperiment, MSImagingArrays

# **Examples**

```
set.seed(1)
x <- matrix(rlnorm(81), nrow=9, ncol=9)
mz <- sort(runif(9))
coord <- expand.grid(x=1:3, y=1:3)

mse <- MSImagingExperiment(
    spectraData=x,
    featureData=MassDataFrame(mz=mz),
    pixelData=PositionDataFrame(coord))

print(mse)</pre>
```

normalize 17

normalize

Normalize spectra

# Description

Apply deferred normalization to spectra.

### Usage

```
## S4 method for signature 'SpectralImagingData'
normalize(object,
    method = c("tic", "rms", "reference"), ...)
```

# Arguments

object A spectral imaging dataset.

method The normalization method to use. See rescale for details.

... Additional arguments passed to the normalization function.

### **Details**

The supported normalization methods are:

- "tic": Total ion current normalization using rescale\_sum.
- "rms": Root-mean-squared normalization using rescale\_rms.
- "reference": Normalization according to a reference feature using rescale\_ref.

### Value

An object of the same class with the processing step queued.

# Note

The normalization is deferred until process() is called.

### Author(s)

```
Kylie A. Bemis
```

### See Also

```
smooth, recalibrate, reduceBaseline, peakPick, process
```

18 peakAlign

### **Examples**

```
set.seed(1)
mse <- simulateImage(preset=1, npeaks=10, dim=c(3,3))
# queue normalization
mse2 <- normalize(mse, method="tic")
# apply normalization
mse2 <- process(mse2)</pre>
```

peakAlign

Align peaks across spectra

# **Description**

Align peaks across spectra in a spectral imaging dataset.

```
## S4 method for signature 'MSImagingExperiment'
peakAlign(object, ref,
    spectra = "intensity", index = "mz",
   tolerance = NA, units = c("ppm", "mz"), ...)
## S4 method for signature 'MSImagingArrays'
peakAlign(object, ref,
    spectra = "intensity", index = "mz",
   tolerance = NA, units = c("ppm", "mz"), ...)
## S4 method for signature 'SpectralImagingExperiment'
peakAlign(object, ref,
    spectra = "intensity", index = NULL,
   tolerance = NA, units = c("relative", "absolute"),
   nchunks = getCardinalNChunks(),
   verbose = getCardinalVerbose(),
   BPPARAM = getCardinalBPPARAM(), ...)
## S4 method for signature 'SpectralImagingArrays'
peakAlign(object, ref,
   spectra = "intensity", index = NULL,
   tolerance = NA, units = c("relative", "absolute"),
   nchunks = getCardinalNChunks(),
   verbose = getCardinalVerbose(),
   BPPARAM = getCardinalBPPARAM(), ...)
```

peakAlign 19

#### **Arguments**

object A spectral imaging dataset.

ref The locations of reference peaks to use for the alignment.

spectra The name of the array in spectraData() to use for the peak intensities.

index The name of the array in spectraData() (for SpectralImagingArrays) or

column in featureData() (for SpectralImagingExperiment) to use for the

peak locations.

tolerance The tolerance for matching a detected peak to the reference. If NA, then the

tolerance is automatically determined as half the minimum distance between

locations in the estimated spectral domain (see "Details").

units The units for the above tolerance.

nchunks The number of chunks to use for processing.

verbose Should progress messages be printed?

BPPARAM An optional instance of BiocParallelParam. See documentation for bplapply.

... Options passed to process().

#### **Details**

Before peak alignment, process() is called to apply any queued pre-processing steps. It is assumed that peakPick() has either been queued or already applied to the data.

If ref is provided, then the aligned peaks are returned immediately without additional processing. (Peaks are binned on-the-fly to the reference peak locations.)

If ref is not provided, then the shared peaks must be determined automatically. This starts with creation of a shared domain giving a list of possible peak locations.

For SpectralImagingArrays, estimateDomain() is used to create the shared domain from the index array. For SpectralImagingExperiment, the index column of featureData() is used directly.

Next, binpeaks is used to bin the observed peaks to the shared domain. Then, mergepeaks is used to merge peaks that are separated by a distance less than the given tolerance.

The averaged locations of the merged peaks in each bin are used as the shared peaks for the full dataset, and the aligned peaks are returned. (Peaks are binned on-the-fly to the shared peak locations.)

### Value

A new object derived from SpectralImagingExperiment with the aligned peaks.

#### Author(s)

Kylie A. Bemis

#### See Also

process peakPick, peakProcess

20 peakPick

### **Examples**

```
set.seed(1)
mse <- simulateImage(preset=1, npeaks=10, dim=c(3,3))
# queue peak picking
mse2 <- peakPick(mse, method="diff", SNR=6)
# align peaks
mse2 <- peakAlign(mse2)
plot(mse2, i=4)</pre>
```

peakPick

Peak pick spectra

#### **Description**

Apply deferred peak picking to spectra.

# Usage

```
## S4 method for signature 'MSImagingExperiment'
peakPick(object, ref,
    method = c("diff", "sd", "mad", "quantile", "filter", "cwt"),
    SNR = 2, type = c("height", "area"),
    tolerance = NA, units = c("ppm", "mz"), ...)

## S4 method for signature 'MSImagingArrays'
peakPick(object, ref,
    method = c("diff", "sd", "mad", "quantile", "filter", "cwt"),
    SNR = 2, type = c("height", "area"),
    tolerance = NA, units = c("ppm", "mz"), ...)

## S4 method for signature 'SpectralImagingData'
peakPick(object, ref,
    method = c("diff", "sd", "mad", "quantile", "filter", "cwt"),
    SNR = 2, type = c("height", "area"),
    tolerance = NA, units = c("relative", "absolute"), ...)
```

# **Arguments**

object A spectral imaging dataset.

ref Optional vector giving locations of reference peaks to extract from the dataset.

method The peak picking method to use. See findpeaks for details.

SNR The signal-to-noise threshold to use to determine a peak.

type The type of value to use to summarize the peak.

peakPick 21

tolerance	If ref is specified, the tolerance to use when deciding if a local peak in a spectrum matches a reference peak. If NA, then the tolerance is automatically determined as half the minimum distance between peaks in the reference.
units	The units for the above tolerance.
	Additional arguments passed to the peak picking function.

#### **Details**

Unless otherwise specified, peaks are detected as local maxima which are then compared to the estimated noise level to determine a signal-to-noise ratio for each peak. Most of the peak detection methods below are differentiated by how they estimate the noise in the specturm.

The supported peak picking methods are:

- "diff": Estimate noise based on the derivative of the signal using estnoise\_diff.
- "sd": Estimate noise from standard deviation using estnoise\_sd.
- "mad": Estimate noise from mean absolute deviation using estnoise\_mad.
- "quantile": Estimate noise from a rolling quantile of the difference between the raw signal and a smoothed signal using estnoise\_quant.
- "filter": Estimate noise using dynamic filtering of the local peaks using estnoise\_filt.
- "cwt": Detect peaks based on continuous wavelet transform (CWT) using findpeaks\_cwt.

If ref is provided, then the signal-to-noise ratio is not determined, and any detected local maxima are summarized as long as they match to a reference peak.

#### Value

An object of the same class with the processing step queued.

#### Note

The peak picking is deferred until process() is called.

### Author(s)

```
Kylie A. Bemis
```

### See Also

```
process, peakAlign, peakProcess, estimateReferencePeaks
```

### **Examples**

```
set.seed(1)
mse <- simulateImage(preset=1, npeaks=10, dim=c(3,3))
# queue peak picking
mse2 <- peakPick(mse, method="diff", SNR=6)
plot(mse2, i=4)</pre>
```

22 peakProcess

```
# apply peak picking
mse2 <- process(mse2)</pre>
```

peakProcess

Process peaks in mass spectra

# **Description**

Apply peak picking and alignment to a mass spectrometry imaging dataset.

### Usage

```
## S4 method for signature 'MSImagingExperiment_OR_Arrays'
peakProcess(object, ref,
    spectra = "intensity", index = "mz",
    method = c("diff", "sd", "mad", "quantile", "filter", "cwt"),
    SNR = 2, type = c("height", "area"),
    tolerance = NA, units = c("ppm", "mz"),
    sampleSize = NA, filterFreq = TRUE, outfile = NULL,
    nchunks = getCardinalNChunks(),
    verbose = getCardinalVerbose(),
    BPPARAM = getCardinalBPPARAM(), ...)
```

# Arguments

object	A spectral imaging dataset.
ref	The locations of reference peaks to use for the alignment.
spectra	The name of the array in spectraData() to use for the peak intensities.
index	The name of the array in spectraData() (for MSImagingArrays) or column in featureData() (for MSImagingExperiment) to use for the peak locations.
method	The peak picking method to use. See findpeaks for details.
SNR	The signal-to-noise threshold to use to determine a peak.
type	The type of value to use to summarize the peak.
tolerance	The tolerance for matching a detected peak to the reference peaks or the shared m/z values. Passed to peakPick and peakAlign.
units	The units for the above tolerance.
sampleSize	The count or proportion giving a subset of spectra to use to determine reference peaks.
filterFreq	Either a logical value indicating whether singleton peaks should be removed, or a count or frequency used as a threshold to filter the peaks.
outfile	Optional. The name of a file to write the resulting dataset as imzML.
nchunks	The number of chunks to use for processing.
verbose	Should progress messages be printed?
BPPARAM	An optional instance of BiocParallelParam. See documentation for bplapply.
	Options passed to process().

peakProcess 23

#### **Details**

This method provides a combined interface for peakPick and peakAlign for the most common approaches to peak processing.

If peakPick() has been queued already, then it will be applied. Otherwise, it will be called internally with the provided arguments.

There are two main paths depending on whether (1) peaks should be extracted based on a reference or (2) peak picking should be performed on the full dataset and then aligned.

If either ref is provided or sampleSize is finite, then (1) is chosen and peaks are extracted based on the reference. If the reference is not provided, then peak picking and alignment performed on a subset of spectra (according to sampleSize) to create the reference peaks. The reference peaks are then used to extract peaks from the full dataset.

Otherwise, (2) is chosen and peaks are picked and aligned across all spectra.

The advantage of (1) is that all reference peaks will be summarized even they would not have a high enough signal-to-noise ratio to be detected in some spectra. The disadvantage is that rare peaks that do not appear in the sampled subset of spectra will not be included in the process peaks.

The advantage of (2) is that rare peaks will be included because peak detection is performed on all spectra. The disadvantage is that some peaks may be missing from some spectra despite having nonzero intensities, because they did not have a high enough signal-to-noise ratio to be detected as peaks.

Setting sampleSize to 1 will balance these advantages and disadvantages because the reference will be based on all spectra. However, this means the full dataset must be processed at least twice (possibly more if intermediate calculations are necessary), so it will be more time-consuming.

#### Value

A new object derived from MSImagingExperiment with the processed peaks.

#### Author(s)

Kylie A. Bemis

#### See Also

```
process peakPick, peakAlign
```

# Examples

```
set.seed(1)
mse <- simulateImage(preset=1, npeaks=10, dim=c(3,3))
# process peaks
mse2 <- peakProcess(mse, method="diff", SNR=3)
plot(mse2, i=4)</pre>
```

24 pixels

pixels

Find pixel indices

# **Description**

Search for the column indices of a spectral imaging dataset that correspond to specific pixels, based on a set of conditions.

# Usage

# Arguments

object A spectral imaging dataset.

Expressions that evaluate to logical vectors in the environment of pixelData().

The coordinates of the pixels to include.

The run of the pixels to include.

The tolerance for matching pixels to coordinates.

env The enclosing environment for evaluating . . . .

# Author(s)

Kylie A. Bemis

# **Examples**

```
set.seed(1)
mse <- simulateImage(preset=1, npeaks=10, dim=c(10,10))
pixels(mse, x > 6, y > 6)
pixels(mse, coord=expand.grid(x=1:3, y=1:3))
```

plot-image 25

plot-image

Plot images from a spectral imaging dataset

### **Description**

Create and display images from the spectra or pixel data of a spectral imaging dataset using a formula interface.

#### **Usage**

```
## S4 method for signature 'MSImagingExperiment'
image(x,
    formula = intensity \sim x * y,
    i = features(x, mz=mz),
   mz = NULL,
    tolerance = NA,
    units = c("ppm", "mz"),
    xlab, ylab)
## S4 method for signature 'SpectralImagingExperiment'
image(x,
    formula,
    i = 1L
    run = NULL,
    groups = NULL,
    superpose = FALSE,
    key = TRUE,
    enhance = NULL,
    smooth = NULL,
    scale = NULL,
    subset = TRUE)
```

### **Arguments**

formula A formula of the form vals  $\sim x * y$  giving the image values and the pixel co-

ordinates. The LHS is taken to be the name of an array in spectraData() and the RHS is taken to be columns of pixelData(). Alternatively, if formula is a string or if i is NULL, then the LHS is interpreted as the name of a column of

pixelData() as well.

i The index of the feature(s) to plot for the image(s).

mz The m/z value(s) to plot for the image(s).

tolerance If specified, the tolerance to consider a feature as being equal to the given mz

value.

26 plot-image

The units for the above tolerance. units Additional arguments passed to plot\_image. Plotting labels. xlab, ylab The names of experimental runs to include, or the index of the levels of the runs run to include. A vector coercible to a factor indicating which of the specified features should groups be plotted with the same color. If multiple images are plotted, should they be superposed on top of each other, superpose or plotted seperately? Should a legend or colorkey be plotted? key enhance The name of a contrast enhancement method, such as "hist" or "adapt" for enhance\_hist() and enhance\_adapt(), etc. See enhance for details. smooth The name of a smoothing method, such as "gauss" or "bi" for filt2\_gauss() and filt2\_bi(), etc. See filt2 for details. If multiple images are plotted, should they be scaled to the same intensity scale? scale

A logical vector indicating which pixels to include in the image.

### Author(s)

subset

Kylie A. Bemis

# See Also

```
image, plot_image, selectROI
```

### **Examples**

```
set.seed(1)
x <- simulateImage(preset=2, npeaks=10, dim=c(16,16))
peaks <- mz(metadata(x)$design$featureData)
image(x, mz=peaks[1L], tolerance=0.5, units="mz")
image(x, mz=peaks[1L], smooth="gaussian")
image(x, mz=peaks[1:9], smooth="adaptive")

x <- summarizePixels(x, stat=c(TIC="mean"))
image(x, "TIC")</pre>
```

plot-spectra 27

plot-spectra

Plot spectra from a spectral imaging dataset

### **Description**

Create and display sliced images from a spectral imaging dataset using a formula interface.

```
## S4 method for signature 'MSImagingExperiment, missing'
plot(x,
    formula = intensity ~ mz,
    i = pixels(x, coord=coord, run=run),
    coord = NULL,
    run = NULL,
    . . . ,
    xlab, ylab,
    isPeaks = isCentroided(x))
## S4 method for signature 'MSImagingArrays,missing'
plot(x,
    formula = intensity ~ mz,
    i = pixels(x, coord=coord, run=run),
    coord = NULL,
    run = NULL,
    . . . ,
    xlab, ylab,
    isPeaks = isCentroided(x))
## S4 method for signature 'SpectralImagingExperiment,missing'
plot(x,
    formula,
    i = 1L,
    groups = NULL,
    superpose = FALSE,
    key = TRUE,
    n = Inf,
    downsampler = "lttb",
    isPeaks = FALSE,
    annPeaks = 0)
## S4 method for signature 'SpectralImagingArrays,missing'
plot(x,
    formula,
    i = 1L,
    groups = NULL,
```

28 plot-spectra

```
superpose = FALSE,
key = TRUE,
...,
n = Inf,
downsampler = "lttb",
isPeaks = FALSE,
annPeaks = 0)
```

#### **Arguments**

x A spectral imaging dataset.

formula A formula of the form vals ~ t giving the spectra values and their domain loca-

tions. The LHS is taken to be the name of an array in spectraData() and the RHS is either an array in spectraData() for SpectralImagingArrays-derived classes or a column of featureData() for SpectralImagingExperiment-derived classes. Alternatively, if formula is a string or if i is NULL, then the LHS is in-

terpreted as the name of a column of featureData() for SpectralImagingExperiment

as well.

i The index of the spectrum to plot.

coord The coordinates of the spectrum to plot.

run The run of the spectrum to plot.

... Additional arguments passed to plot\_signal.

xlab, ylab Plotting labels.

isPeaks Should the spectrum be plotted as peaks or as a continuous signal?

annPeaks If isPeaks is TRUE, either an integer giving the number of peaks to annotate

(i.e., label with their location), or a plotting symbol (e.g., "circle", "cross", etc.)

to indicate the peak locations.

groups A vector coercible to a factor indicating which of the specified spectra should be

plotted with the same color.

superpose If multiple spectra are plotted, should they be superposed on top of each other,

or plotted seperately?

key Should a legend or colorkey be plotted?

n, downsampler A spectrum can contain far more data points than are needed to visualize it,

potentially making the plotting unnecessarily slow. Downsampling can be performed to improve plotting speed while maintaining the visual integrity of the

spectrum. See downsample for details.

#### Author(s)

Kylie A. Bemis

#### See Also

```
plot, plot_signal
```

PositionDataFrame-class 29

### **Examples**

PositionDataFrame-class

PositionDataFrame: Extended data frame with key columns

# **Description**

A data frame for metadata with spatial coordinates and multiple experimental runs.

### Usage

```
PositionDataFrame(coord, run, ..., row.names = FALSE)
```

### **Arguments**

coord A data frame or matrix of coordinates.

run A factor giving the experimental runs.

... Arguments passed to the DataFrame().

Either a vector of row names or a logical value indicating whether row names should be generated automatically (from the m/z values).

#### Methods

```
coord(object), coord(object) <- value: Get or set the coordinate columns.
coordNames(object), coordNames(object) <- value: Get or set the names of the coordinate columns.
run(object), run(object) <- value: Get or set the experimental run column.
runNames(object), runNames(object) <- value: Get or set the experimental run levels.
nrun(object): Get the number of experimental runs.
is3D(object): Check if the number of spatial dimensions is greater than 2.</pre>
```

### Author(s)

Kylie A. Bemis

30 process

### See Also

XDataFrame, MassDataFrame

# **Examples**

```
## Create an PositionDataFrame object
coord <- expand.grid(x=1:3, y=1:3)
PositionDataFrame(coord=coord, label=LETTERS[1:9])</pre>
```

process

Apply queued processing to spectra

# Description

Queue pre-processing steps on an imaging dataset and apply them, possibly writing out the processed data to a file.

```
## S4 method for signature 'MSImagingExperiment'
process(object, spectra = "intensity", index = "mz",
   domain = NULL, outfile = NULL, ...)
## S4 method for signature 'MSImagingArrays'
process(object, spectra = "intensity", index = "mz",
    domain = NULL, outfile = NULL, ...)
## S4 method for signature 'SpectralImagingExperiment'
process(object, spectra = "intensity", index = NULL,
   domain = NULL, outfile = NULL,
   nchunks = getCardinalNChunks(),
   verbose = getCardinalVerbose(),
   BPPARAM = getCardinalBPPARAM(), ...)
## S4 method for signature 'SpectralImagingArrays'
process(object, spectra = "intensity", index = NULL,
   domain = NULL, outfile = NULL,
   nchunks = getCardinalNChunks(),
   verbose = getCardinalVerbose(),
   BPPARAM = getCardinalBPPARAM(), ...)
## S4 method for signature 'SpectralImagingData'
addProcessing(object, FUN, label, verbose = getCardinalVerbose(), ...)
reset(object, ...)
```

process 31

### **Arguments**

object A spectral imaging dataset. The name of the array in spectraData() to use for the peak intensities. spectra The name of the array in spectraData() (for MSImagingArrays) or column in index featureData() (for MSImagingExperiment) to use for the peak locations. Optional. The name of the array in spectraData() (for MSImagingArrays) domain or column in featureData() (for MSImagingExperiment) to use for output domain (if known). outfile Optional. The name of a file to write the resulting dataset. Creates an imzML file for MSImagingExperiment or MSImagingArrays. The "continuous" format will be written if domain is specified; otherwise the "processed" format will be used in most cases. nchunks The number of chunks to use for processing. verbose Should progress messages be printed? **BPPARAM** An optional instance of BiocParallelParam. See documentation for bplapply. Options passed to chunk\_mapply or chunk\_colapply. . . . FUN A user-specified processing function. label The name of the processing step.

### **Details**

This method allows queueing of delayed processing to an imaging dataset. All of the queued processing steps will be applied in sequence whenever process() is called next. Use reset() to remove all queued processing steps.

Typically, processing steps are queued using methods like normalize, smooth, peakPick, etc.

However, a processing step can be queued manually with addProcessing.

In this case, the user-specified function *must* accept (1) a first argument giving the spectral intensities as a numeric vector and (2) a second argument giving the intensity locations (e.g., m/z values) as a numeric vector.

The value returned by a user-specified function must return either (1) a numeric vector of the same length as the input intensities or (2) a 2-column matrix where the first column is the new locations (e.g., m/z values of peaks) and the second column is the new intensities.

### Value

An object of the same class as the original object, with all processing steps applied.

#### Author(s)

Kylie A. Bemis

#### See Also

normalize, smooth, recalibrate, reduceBaseline, peakPick

32 readMSIData

### **Examples**

```
set.seed(1)
mse <- simulateImage(preset=1, dim=c(3,3), baseline=1)
mse2 <- smooth(mse, width=11)
mse2 <- reduceBaseline(mse2)
plot(mse2, i=4)
mse2 <- process(mse2)</pre>
```

readMSIData

Read mass spectrometry imaging data files

### Description

Read supported mass spectrometry imaging data files, including imzML and Analyze 7.5.

```
## Read any supported MS imaging file
readMSIData(file, ...)
## Read imzML file
readImzML(file, memory = FALSE, check = FALSE,
mass.range = NULL, resolution = NA, units = c("ppm", "mz"),
guess.max = 1000L, as = "auto", parse.only=FALSE,
nchunks = getCardinalNChunks(),
verbose = getCardinalVerbose(),
BPPARAM = getCardinalBPPARAM(), ...)
## Read Analyze 7.5 file
readAnalyze(file, memory = FALSE, as = "auto",
verbose = getCardinalVerbose(), ...)
## Convert from MSImagingExperiment to MSImagingArrays
convertMSImagingExperiment2Arrays(object)
## Convert from MSImagingArrays to MSImagingExperiment
convertMSImagingArrays2Experiment(object, mz = NULL,
mass.range = NULL, resolution = NA, units = c("ppm", "mz"),
guess.max = 1000L, tolerance = 0.5 * resolution,
nchunks = getCardinalNChunks(),
verbose = getCardinalVerbose(),
BPPARAM = getCardinalBPPARAM(), ...)
```

readMSIData 33

#### **Arguments**

file The absolute or relative file path. The file extension must be included for readMSIData. Should the spectra be loaded into memory? If FALSE, the spectra are attached memory as an out-of-memory matrix. check Should the UUID and checksum of the binary data file be checked against the corresponding imzML tags? The mass range to use when converting the data to an MSImagingExperiment. mass.range resolution The mass resolution to use when converting the data to an MSImagingExperiment. This is the inverse of the instrument resolution, if known. It is the width of the m/z bins when converting the data to an MSImagingExperiment. units The units for the above resolution. The number of spectra to use when guessing the mass range and resolution, if guess.max they are not provided. as After reading in the data, what class of object should be returned? The data is initially loaded as an MSImagingArrays object. It may be converted to an MSImagingExperiment object. Setting to "auto" means to determine whichever is more appropriate depending on whether the spectra appear to have been processed and centroided. If TRUE, return only the parsed imzML metadata without creating a new MSImagingArrays parse.only or MSImagingExperiment object. nchunks The number of chunks to use for processing. verbose Should progress messages be printed? **BPPARAM** An optional instance of BiocParallelParam. See documentation for bplapply. Additional arguments passed to parseImzML or parseAnalyze. . . . object A mass spectrometry imaging dataset to convert from one class to another. A vector of shared m/z values for converting to MSImagingExperiment, if not mΖ to be determined automatically. tolerance If the spectra have been centroided but the peaks are unaligned, then this is passed to peakAlign.

#### **Details**

The spectra are initially loaded into a MSImagingArrays object before conversion to MSImagingExperiment (if applicable).

This conversion can be sped up by specifying the mass.range and resolution so they do not have to be determined from the spectra directly. Using a larger value of guess.max can improve the accuracy of the m/z binning for downstream analysis at the expense of a longer conversion time.

If greater control is desired, spectra should be imported as MSImagingArrays, and processing to MSImagingExperiment can be performed manually.

If problems are encountered while trying to import imzML files, the files should be verified and fixed with imzMLValidator.

A Java version of imzML validator can be found at: https://gitlab.com/imzML/imzMLValidator.

A web-based version of imzML validator can be found at: https://imzml.github.io.

34 recalibrate

#### Value

A MSImagingExperiment or MSImagingArrays object.

#### Author(s)

Kylie A. Bemis

#### References

Schramm T, Hester A, Klinkert I, Both J-P, Heeren RMA, Brunelle A, Laprevote O, Desbenoit N, Robbe M-F, Stoeckli M, Spengler B, Rompp A (2012) imzML - A common data format for the flexible exchange and processing of mass spectrometry imaging data. Journal of Proteomics 75 (16):5106-5110. doi:10.1016/j.jprot.2012.07.026

### See Also

```
parseImzML, parseAnalyze writeMSIData
```

recalibrate

Recalibrate spectra

### **Description**

Apply deferred recalibration to spectra.

#### Usage

```
## S4 method for signature 'MSImagingExperiment_OR_Arrays'
recalibrate(object, ref,
    method = c("locmax", "dtw", "cow"),
    tolerance = NA, units = c("ppm", "mz"), ...)

## S4 method for signature 'SpectralImagingData'
recalibrate(object, ref,
    method = c("locmax", "dtw", "cow"),
    tolerance = NA, units = c("relative", "absolute"), ...)
```

# Arguments

object	A spectral	imaging	dataset.
--------	------------	---------	----------

ref The domain (m/z) values or indices of reference peaks to use for the recalibra-

tion.

method The recalibration method to use. See warp1 for details.

tolerance The tolerance for how much a peak can be shifted in either direction.

units The units for the above tolerance.

. . . Additional arguments passed to the recalibration function.

reduceBaseline 35

### **Details**

The supported recalibration methods are:

- "locmax": Align to local maxima using warp1\_loc.
- "dtw": Dynamic time warping using warp1\_dtw.
- "cow": Correlation optimized warping using warp1\_cow.

#### Value

An object of the same class with the processing step queued.

### Note

The recalibration is deferred until process() is called.

### Author(s)

Kylie A. Bemis

#### See Also

```
normalize, smooth, recalibrate, peakPick, process
```

# **Examples**

```
set.seed(1)
mse <- simulateImage(preset=1, npeaks=10, dim=c(3,3), sdmz=250)
plot(mse, i=c(2,4,5), superpose=TRUE, xlim=c(1260,1320))

# queue recalibration
peaks <- estimateReferencePeaks(mse)
mse2 <- recalibrate(mse, ref=peaks, method="locmax", tolerance=500)

# apply recalibration
mse2 <- process(mse2)
plot(mse2, i=c(2,4,5), superpose=TRUE, xlim=c(1260,1320))</pre>
```

reduceBaseline

Reduce baselines in spectra

# **Description**

Apply deferred baseline reduction to spectra.

```
## S4 method for signature 'SpectralImagingData'
reduceBaseline(object,
    method = c("locmin", "hull", "snip", "median"), ...)
```

36 reduceBaseline

# **Arguments**

object A spectral imaging dataset.

method The baseline estimation method to use. See estbase for details.

... Additional arguments passed to the baseline estimation function.

### **Details**

The supported baseline estimation methods are:

- "locmin": Interpolate from local minima using estbase\_loc.
- "hull": Convex hull estimation using estbase\_hull.
- "snip": Sensitive nonlinear iterative peak (SNIP) clipping using estbase\_snip.
- "median": Running medians using estbase\_med.

#### Value

An object of the same class with the processing step queued.

### Note

The baseline reduction is deferred until process() is called.

# Author(s)

```
Kylie A. Bemis
```

#### See Also

```
normalize, smooth, reduceBaseline, peakPick, process
```

# **Examples**

```
set.seed(1)
mse <- simulateImage(preset=1, npeaks=10, dim=c(3,3), baseline=1)
# queue baseline reduction
mse2 <- reduceBaseline(mse, method="locmin")
plot(mse2, i=4)
# apply baseline reduction
mse2 <- process(mse2)</pre>
```

ResultsList-class 37

ResultsList-class

ResultsList: List of modeling results

# **Description**

The ResultsList class provides a container for modeling results with spatial metadata.

#### Usage

```
## Instance creation
ResultsList(..., mcols = NULL)
## Additional methods documented below
```

# Arguments

... The modeling results.

mcols The metadata columns.

## Methods

All methods for SimpleList also work on ResultsList objects. Additional methods are documented below:

```
fitted(object, ...): Extract fitted values from each modeling results object in the list.
predict(object, ...): Predict on each modeling results object in the list.
topFeatures(object, ...): Rank top features for each modeling results object in the list.
plot(x, i = 1L, ...): Plot the ith modeling results.
image(x, i = 1L, ...): Display images for the ith modeling results.
```

#### Author(s)

Kylie A. Bemis

## See Also

SpatialResults

38 selectROI

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Select regions-of-interest in an image

## **Description**

Manually select regions-of-interest or pixels on an imaging dataset. The selectROI method uses the built-in locator function. It can be used with an existing image plot, or a new image will be plotted if image arguments are passed via . . . .

The regions of interest are returned as logical vectors indicating which pixels have been selected. These logical vectors can be combined into factors using the makeFactor function.

# Usage

```
## $4 method for signature 'SpectralImagingExperiment'
selectROI(object, ..., mode = c("region", "pixels"))
makeFactor(..., ordered = FALSE)
```

# **Arguments**

object	A spectral imaging dataset.
mode	The mode of selection: "region" to select a region-of-interest as a polygon, or "pixels" to select individual pixels.
	Additional arguments to be passed to image for selectROI, or name-value pairs of logical vectors to be combined by makeFactor.
ordered	Should the resulting factor be ordered or not?

### Value

A logical vector of length equal to the number of pixels for selectROI.

A factor of the same length as the logical vectors for makeFactor.

## Author(s)

```
Kylie A. Bemis
```

#### See Also

image

simulateSpectra 39

simulateSpectra

Simulate a mass spectrum or MS imaging experiment

#### **Description**

Simulate mass spectra or complete MS imaging experiments, including a possible baseline, spatial and spectral noise, mass drift, mass resolution, and multiplicative variation, etc.

A number of preset imaging designs are available for quick-and-dirty simulation of images.

These functions are designed for small proof-of-concept examples and testing, and may not scale well to simulating larger datasets.

#### Usage

```
simulateSpectra(n = 1L, npeaks = 50L,
    mz = rlnorm(npeaks, 7, 0.3), intensity = rlnorm(npeaks, 1, 0.9),
    from = 0.9 * min(mz), to = 1.1 * max(mz), by = 400,
    sdpeaks = sdpeakmult * log1p(intensity), sdpeakmult = 0.2,
    sdnoise = 0.1, sdmz = 10, resolution = 1000, fmax = 0.5,
    baseline = 0, decay = 10, units=c("ppm", "mz"),
    representation = c("profile", "centroid"), ...)
simulateImage(pixelData, featureData, preset,
    from = 0.9 \times \min(mz), to = 1.1 \times \max(mz), by = 400,
    sdrun = 1, sdpixel = 1, spcorr = 0.3, units=c("ppm", "mz"),
    representation = c("profile", "centroid"),
    nchunks = getCardinalNChunks(),
    verbose = getCardinalVerbose(),
    BPPARAM = getCardinalBPPARAM(), ...)
addShape(pixelData, center, size, shape=c("circle", "square"), name=shape)
presetImageDef(preset = 1L, nrun = 1, npeaks = 30L,
    dim = c(20L, 20L), peakheight = 1, peakdiff = 1,
    sdsample = 0.2, jitter = TRUE, ...)
```

## **Arguments**

n	The number of spectra to simulate.
npeaks	The number of peaks to simulate. Not used if mz and intensity are provided.
mz	The theoretical m/z values of the simulated peaks.
intensity	The mean intensities of the simulated peaks.
from	The minimum m/z value used for the mass range.
to	The maximum m/z value used for the mass range.
by	The step-size used for the observed m/z-values of the profile spectrum.

40 simulateSpectra

sdpeaks The standard deviation(s) for the distributions of observed peak intensities on

the log scale.

sdpeakmult A multiplier used to calculate sdpeaks based on the mean intensities of peaks;

used to simulate multiplicative variance. Not used if sdpeaks is provided.

sdnoise The standard deviation of the random noise in the spectrum on the log scale.

sdmz The standard deviation of the mass error in the observed m/z values of peaks, in

units indicated by units.

resolution The mass resolution as defined by m / dm, where m is the observed mass and dm

is the width of the peak at a proportion of its maximum height defined by fmax (defaults to full-width-at-half-maximum – FWHM – definition). Note that this is NOT the same as the definition of resolution in the readImzML function.

fmax The fraction of the maximum peak height to use when defining the mass resolu-

tion.

baseline The maximum intensity of the baseline. Note that baseline=0 means there is

no baseline.

decay A constant used to calculate the exponential decay of the baseline. Larger values

mean the baseline decays more sharply at the lower mass range of the spectrum.

units The units for by and sdmz. Either parts-per-million or absolute m/z units.

representation Should a profile spectrum be returned or only the centroided peaks?

nchunks The number of chunks to use for processing.

verbose Should progress messages be printed?

BPPARAM An optional instance of BiocParallelParam. See documentation for bplapply.

pixelData A PositionDataFrame giving the pixel design of the experiment. The names

of the columns should match the names of columns in featureData. Each column should be a logical vector corresponding to a morphological substructure,

indicate which pixels belong to that substructure.

featureData A MassDataFrame giving the feature design of the experiment. Each row should

correspond to an expected peak. The names of the columns should match the names of columns in pixelData. Each column should be a numeric vector corresponding to a morphological substructure, giving the mean intensity of that

peak for that substructure.

preset A number indicating a preset image definition to use.

The number of runs to simulate for each condition.

Sdrun A standard deviation giving the run-to-run variance.

sdpixel A standard deviation giving the pixel-to-pixel variance.

spcorr The spatial autocorrelation. Must be between 0 and 1, where spcorr=0 indicates

no spatial autocorrelation.

... Additional arguments to pass to simulateSpectra or presetImageDef.

dim The dimensions of the preset image.

peakheight Reference intensities used for peak heights by the preset.

peakdiff A reference intensity difference used for the mean peak height difference be-

tween conditions, for presets that simulate multiple conditions.

simulateSpectra 41

A standard deviation giving the amount of variation from the true peak heights for this simulated sample.

jitter Should random noise be added to the location and size of the shapes?

center The center of the shape.

size The size of the shape (from the center).

shape What type of shape to add.

name The name of the added column.

#### **Details**

The simulateSpectra() and simulateImage() functions are used to simulate mass spectra and MS imaging experiments. They provide a great deal of control over the parameters of the simulation, including all sources of variation.

For simulateImage(), the user should provide the design of the simulated experiment via matching columns in pixelData and featureData, where each column corresponds to different morphological substructures or differing conditions. These design data frames are returned in the metadata() of the returned object for later reference.

A number of presets are defined by presetImageDef(), which returns only the pixelData and featureData necessary to define the experiment for simulateImage(). These can be referenced for help in understanding how to define experiments for simulateImage().

The preset images are:

- 1: a centered circle
- 2: a topleft circle and a bottomright square
- 3: two corner squares and a centered circle
- 4: a centered circle with conditions A and B in different runs
- 5: a topleft circle and a bottomright square with conditions A and B in different runs
- 6: two corner squares and a centered circle; the circle has conditions A and B in different runs
- 7: matched pairs of circles with conditions A and B within the same runs; includes reference peaks
- 8: matched pairs of circles inside squares with conditions A and B within the same runs; includes reference peaks
- 9: a small sphere inside a larger sphere (3D)

The addShape() function is provided for convenience when generating the pixelData for simulateImage(), as a simple way of adding morphological substructures using basic shapes such as squares and circles.

#### Value

For simulateSpectra, a list with elements:

- mz: a numeric vector of the observed m/z values
- intensity: a numeric vector or matrix of the intensities

42 slice

For simulateImage, a MSImagingExperiment object.

For addShape, a new PositionDataFrame with a logical column added for the corresponding shape.

For presetImageDef, a list with two elements: the pixelData and featureData to be used as input to simulateImage().

## Author(s)

Kylie A. Bemis

#### See Also

simspec

## **Examples**

```
# generate a spectrum
s <- simulateSpectra(1)
plot(s$intensity ~ s$mz, type="1")

# generate a noisy low-resolution spectrum with a baseline
s <- simulateSpectra(1, baseline=2, sdnoise=0.3, resolution=100)
plot(s$intensity ~ s$mz, type="1")

# generate a high-resolution spectrum
s <- simulateSpectra(1, npeaks=100, resolution=10000)
plot(s$intensity ~ s$mz, type="1")

# generate an image
mse <- simulateImage(preset=1, npeaks=10, dim=c(10,10))
peaks <- mz(metadata(mse)$design$featureData)

image(mse, mz=peaks[c(1,4,5,6)])
plot(mse, coord=c(x=3,y=3))</pre>
```

slice

Slice an image

## **Description**

Slice a spectral imaging dataset as a "data cube".

#### Usage

```
slice(x, i = features(x, ...), ..., run = NULL,
    simplify = TRUE, drop = TRUE)
```

smooth 43

# **Arguments**

X	A spectral imaging dataset.
i	The indices of features to slice for the images.
	Conditions describing features to slice, passed to features().
run	The names of experimental runs to include, or the index of the levels of the runs to include.
simplify	The image slices be returned as a list, or simplified to an array?
drop	Should redundant array dimensions be dropped? If TRUE, dimensions with only one level are dropped using drop.

#### Value

A list or array of the sliced image(s). If multiple images are sliced and simplify=TRUE, then the *last* dimension will be the features.

## Author(s)

Kylie A. Bemis

# **Examples**

```
set.seed(1)
mse <- simulateImage(preset=1, npeaks=10, dim=c(10,10), representation="centroid")
peaks <- mz(metadata(mse)$design$featureData)

# slice image for first feature
slice(mse, 1)

# slice by m/z-value
slice(mse, mz=peaks[1])

# slice multiple
slice(mse, mz=peaks[1:3])</pre>
```

smooth

Smooth spectra

# Description

Apply deferred smoothing to spectra.

# Usage

```
## S4 method for signature 'SpectralImagingData'
smooth(x,
    method = c("gaussian", "bilateral", "adaptive",
        "diff", "guide", "pag", "sgolay", "ma"), ...)
```

44 smooth

# **Arguments**

x A spectral imaging dataset.
 method The smoothing method to use. See filt1 for details.
 ... Additional arguments passed to the smoothing function.

#### **Details**

The supported smoothing methods are:

- "gaussian": Gaussian smoothing using filt1\_gauss.
- "bilateral": Bilateral filter using filt1\_bi.
- "adaptive": Adaptive bilateral filter using filt1\_adapt.
- "diff": Nonlinear diffusion smoothing using filt1\_diff.
- "guide": Guided filter using filt1\_guide.
- "pag": Peak-aware guided filter using filt1\_pag.
- "sgolay": Savitzky-Golar filter using filt1\_sg.
- "ma": Moving average filter using filt1\_ma.

#### Value

An object of the same class with the processing step queued.

## Note

The smoothing is deferred until process() is called.

### Author(s)

```
Kylie A. Bemis
```

# See Also

```
normalize, recalibrate, reduceBaseline, peakPick, process
```

## **Examples**

```
set.seed(1)
mse <- simulateImage(preset=1, npeaks=10, dim=c(3,3))
# queue smoothing
mse2 <- smooth(mse, method="gaussian", width=11)
plot(mse2, i=4)
# apply smoothing
mse2 <- process(mse2)</pre>
```

SpatialCV 45

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Cross-validation for spectral imaging data

## **Description**

Apply cross-validation with an existing or a user-specified modeling function over folds of a spectral imaging dataset.

# Usage

#### **Arguments**

fit.	The function used to fit the model.
x, y	The data and response variable, where $x$ is assumed to be an $P$ $x$ $N$ dataset such as a SpectralImagingExperiment
folds	A vector coercible to a factor giving the fold for each row or column of x.
	Additional arguments passed to fit. and predict
predict.	The function used to predict on new data from the fitted model. The fitted model is passed as the 1st argument and the test data is passed as the 2nd argument.
keep.models	Should the models be kept and returned?
trainProcess, t	rainArgs
	A function and arguments used for processing the training sets. The training set

A function and arguments used for processing the training sets. The training set is passed as the 1st argument to trainProcess.

testProcess, testArgs

A function and arguments used for processing the test sets. The test set is passed as the 1st argument to trainProcess, and the processed training set is passed as the 2nd argument.

verbose Should progress be printed for each iteration?

nchunks The number of chunks to use. *Passed* to fit., predict., trainProcess and

testProcess.

46 SpatialDGMM

BPPARAM	An optional instance of BiocParallelParam. See documentation for bplapply. <i>Passed</i> to fit., predict., trainProcess and testProcess.
object	An object inheriting from SpatialCV.
type	The type of prediction, where "response" means the fitted response matrix and "class" will be the vector of class predictions (only for classification).
i	If predictions are made for multiple sets of parameters, which set of parameters (i.e., which element of the fitted.values list) should be plotted?
layout	A vector of the form c(nrow, ncol) specifying the number of rows and columns in the facet grid.
free	A string specifying the free spatial dimensions during faceting. E.g., "", "x", "y", "xy", "yx".

#### **Details**

This method is designed to be used with the provided classification methods, but can also be used with user-provided functions and methods as long as they conform to certain expectations. Internally, cv\_do from the matter package is used to perform the cross-validation. See ?cv\_do for details.

#### Value

An object of class SpatialCV derived from SpatialResults and containing accuracies for each fold, the predictions for each fold, and (optionally) the fitted models.

#### Author(s)

Kylie A. Bemis

## See Also

cv\_do, spatialShrunkenCentroids, PLS, OPLS

SpatialDGMM	Spatially-aware Dirichlet Gaussian mixture model	

# **Description**

Fit a spatially-aware Gaussian mixture models to each feature. The model uses Dirichlet prior is used to achieve spatial smoothing. The means and standard deviations of the Gaussian components are estimated using gradient descent. Simulated annealing is used to avoid local optimia and achieve better parameter estimates.

SpatialDGMM 47

#### Usage

```
## S4 method for signature 'ANY'
spatialDGMM(x, coord, i, r = 1, k = 2, groups = NULL,
   weights = c("gaussian", "adaptive"),
   neighbors = findNeighbors(coord, r=r, groups=groups),
   annealing = TRUE, compress = TRUE, byrow = FALSE,
   nchunks = getCardinalNChunks(),
   verbose = getCardinalVerbose(),
   BPPARAM = getCardinalBPPARAM(), ...)
## S4 method for signature 'SpectralImagingExperiment'
spatialDGMM(x, i, r = 1, k = 2, groups = run(x),
   weights = c("gaussian", "adaptive"),
    neighbors = findNeighbors(coord(x), r=r, groups=groups), ...)
## S4 method for signature 'SpatialDGMM'
logLik(object, ...)
## S4 method for signature 'SpatialDGMM, missing'
plot(x, i = 1L, type = "density",
   layout = NULL, free = "", ...)
## S4 method for signature 'SpatialDGMM'
image(x, i = 1L, type = "class",
   layout = NULL, free = "", ...)
```

# Arguments

X	A spatial dataset in P x N matrix format.
i	The rows/columns of x to segment (if not all of them).
coord	The spatial coordinates of the rows/columns of x. Ignored if neighbors is provided.
r	The spatial maximum distance for an observation to be considered a neighbor. Ignored if neighbors is provided.
k	The number of Gaussian components.
groups	Observations belonging to the different groups will be segmented independently. This should be set to the samples if statistic testing (via meansTest is to be performed.)
weights	The type of spatial weights to use for the smoothing. Gaussian weights are weighted only by distance, while adaptive weights also consider the dissimilarity between neighboring observations.
neighbors	A factor giving which observations should be treated as spatially-independent. Observations in the same group are assumed to have a spatial relationship.
annealing	Should simulated annealing be used?
compress	Should the results be compressed? The results can be larger than the original dataset, so compressing them is useful. If this option is used, then the class

48 SpatialDGMM

probabilities are not returned, and the class assignments are compressed using

	drle.
byrow	Should the rows or columns of x be segmented?
nchunks	The number of chunks to use for processing.
verbose	Should progress messages be printed?
BPPARAM	$An optional \ instance \ of \ BiocParallel Param. \ See \ documentation \ for \ bplapply.$
	Additional arguments passed to the next method.
object	A SpatialDGMM object.
type	The type of plot to display.
layout	A vector of the form $c(nrow, ncol)$ specifying the number of rows and columns in the facet grid.
free	A string specifying the free spatial dimensions during faceting. E.g., "", "x", "y", "xy", "yx".

#### Value

An object of class SpatialDGMM derived from SpatialResults, containing the fitted sgmixn object and the spatial metadata.

# Author(s)

Dan Guo and Kylie A. Bemis

## References

Guo, D., Bemis, K., Rawlins, C., Agar, J., and Vitek, O. (2019.) Unsupervised segmentation of mass spectrometric ion images characterizes morphology of tissues. Proceedings of ISMB/ECCB, Basel, Switzerland, 2019.

## **Examples**

SpatialFastmap 49

SpatialFastmap

Spatially-aware FastMap projection

### **Description**

Compute spatially-aware FastMap projection.

## Usage

```
## S4 method for signature 'ANY'
spatialFastmap(x, coord, r = 1, ncomp = 3,
   weights = c("gaussian", "adaptive"),
   neighbors = findNeighbors(coord, r=r),
   transpose = TRUE, niter = 3L,
   nchunks = getCardinalNChunks(),
   verbose = getCardinalVerbose(),
   BPPARAM = getCardinalBPPARAM(), ...)
## S4 method for signature 'SpectralImagingExperiment'
spatialFastmap(x, r = 1, ncomp = 3,
   weights = c("gaussian", "adaptive"),
   neighbors = findNeighbors(x, r=r), ...)
## S4 method for signature 'SpatialFastmap'
predict(object, newdata,
    neighbors = findNeighbors(newdata, r=object$r),
   nchunks = getCardinalNChunks(),
   BPPARAM = getCardinalBPPARAM(), ...)
## S4 method for signature 'SpatialFastmap,missing'
plot(x, type = c("scree", "x"), ..., xlab, ylab)
## S4 method for signature 'SpatialFastmap'
image(x, type = "x", ...)
```

#### **Arguments**

X	A spatial dataset in P x N matrix format.
coord	The spatial coordinates of the rows/columns of $\boldsymbol{x}$ . Ignored if neighbors is provided.
r	The spatial maximum distance for an observation to be considered a neighbor. Ignored if neighbors is provided.
ncomp	The number of FastMap components.
weights	The type of spatial weights to use for the smoothing. Gaussian weights are weighted only by distance, while adaptive weights also consider the dissimilarity between neighboring observations.

50 SpatialFastmap

A factor giving which observations should be treated as spatially-independent. neighbors Observations in the same group are assumed to have a spatial relationship. Should x be considered P x N? transpose The number of iterations used to calculate the pivots for each FastMap componiter The number of chunks to use for processing. nchunks verbose Should progress messages be printed? **BPPARAM** An optional instance of BiocParallelParam. See documentation for bplapply. Additional arguments passed to the next method. object A SpatialFastmap object. A new SpectralImagingExperiment for which to calculate the scores. newdata The type of plot to display.

type

Plotting labels. xlab, ylab

#### Value

An object of class SpatialFastmap derived from SpatialResults, containing the fitted fastmap object and the spatial metadata.

#### Author(s)

Kylie A. Bemis

#### References

Alexandrov, T., & Kobarg, J. H. (2011). Efficient spatial segmentation of large imaging mass spectrometry datasets with spatially aware clustering. Bioinformatics, 27(13), i230-i238. doi:10.1093/bioinformatics/btr246 Faloutsos, C., & Lin, D. (1995). FastMap: A Fast Algorithm for Indexing, Data-Mining and Visualization of Traditional and Multimedia Datasets. Presented at the Proceedings of the 1995 ACM SIGMOD international conference on Management of data.

#### See Also

```
PCA, NMF, spatialKMeans
```

#### **Examples**

```
set.seed(1)
mse <- simulateImage(preset=2, npeaks=20, dim=c(10,10),</pre>
    representation="centroid")
# project to FastMap components
fm <- spatialFastmap(mse, r=1, ncomp=2, weights="adaptive")</pre>
# visualize first 2 components
image(fm)
```

SpatialKMeans 51

SpatialKMeans

Spatially-aware K-means clustering

## **Description**

Perform spatially-aware k-means clustering. First the data is projected to a reduced dimension space using spatialFastmap. Then ordinary k-means clustering is applied to the projected data.

### Usage

```
## S4 method for signature 'ANY'
spatialKMeans(x, coord, r = 1, k = 2, ncomp = max(k),
       weights = c("gaussian", "adaptive"),
       neighbors = findNeighbors(coord, r=r),
       transpose = TRUE, niter = 2L,
       centers = TRUE, correlation = TRUE,
       nchunks = getCardinalNChunks(),
       verbose = getCardinalVerbose(),
       BPPARAM = getCardinalBPPARAM(), ...)
## S4 method for signature 'SpectralImagingExperiment'
spatialKMeans(x, r = 1, k = 2, ncomp = max(k),
       weights = c("gaussian", "adaptive"),
       neighbors = findNeighbors(x, r=r), ...)
## S4 method for signature 'SpatialKMeans'
topFeatures(object, n = Inf, sort.by = "correlation", ...)
## S4 method for signature 'SpatialKMeans,missing'
plot(x, type = c("correlation", "centers"), ..., xlab, ylab)
## S4 method for signature 'SpatialKMeans'
image(x, type = "cluster", ...)
```

# Arguments

Χ	A spatial dataset in P x N matrix format.
coord	The spatial coordinates of the rows/columns of x. Ignored if neighbors is provided.
r	The spatial maximum distance for an observation to be considered a neighbor. Ignored if neighbors is provided.
k	The number of clusters.
ncomp	The number of FastMap components.
weights	The type of spatial weights to use for the smoothing. Gaussian weights are weighted only by distance, while adaptive weights also consider the dissimilarity

between neighboring observations.

52 SpatialKMeans

neighbors A factor giving which observations should be treated as spatially-independent.

Observations in the same group are assumed to have a spatial relationship.

transpose Should x be considered P x N?

niter The number of iterations used to calculate the pivots for each FastMap compo-

nent.

centers Should the cluster centers be re-calculated on the original data?

correlation Should the correlations between features and the clusters be calculated?

nchunks The number of chunks to use for processing. verbose Should progress messages be printed?

BPPARAM An optional instance of BiocParallelParam. See documentation for bplapply.

... Additional arguments passed to the next method.

object A SpatialKMeans object.

n, sort.by For topFeatures, the number of top features to return and how to sort them.

type The type of plot to display.

xlab, ylab Plotting labels.

#### Value

An object of class SpatialKMeans derived from SpatialResults, containing the fitted kmeans object and the spatial metadata.

#### Author(s)

Kylie A. Bemis

#### References

Alexandrov, T., & Kobarg, J. H. (2011). Efficient spatial segmentation of large imaging mass spectrometry datasets with spatially aware clustering. Bioinformatics, 27(13), i230-i238. doi:10.1093/bioinformatics/btr246 Faloutsos, C., & Lin, D. (1995). FastMap: A Fast Algorithm for Indexing, Data-Mining and Visualization of Traditional and Multimedia Datasets. Presented at the Proceedings of the 1995 ACM SIGMOD international conference on Management of data.

#### See Also

spatialKMeans spatialShrunkenCentroids

## **Examples**

SpatialNMF 53

|--|

#### **Description**

Compute nonnegative matrix factorization using alternating least squares or multiplicative updates.

# Usage

```
## S4 method for signature 'ANY'
NMF(x, ncomp = 3, method = c("als", "mult"),
    verbose = getCardinalVerbose(), ...)

## S4 method for signature 'SpectralImagingExperiment'
NMF(x, ncomp = 3, method = c("als", "mult"), ...)

## S4 method for signature 'SpatialNMF'
predict(object, newdata, ...)

## S4 method for signature 'SpatialNMF,missing'
plot(x, type = c("activation", "x"), ..., xlab, ylab)

## S4 method for signature 'SpatialNMF'
image(x, type = "x", ...)
```

## **Arguments**

X	A dataset in P x N matrix format.
ncomp	The number of components to calculate.
method	The method to use. Alternating least squares ("als") tends to be faster and potentially more accurate, but can be numerically unstable for data with high correlated features. Multiplicative updates ("mult") can be slower, but is more numerically stable.
verbose	Should progress messages be printed?
	Options passed to irlba.
object	A SpatialNMF object.
newdata	A new SpectralImagingExperiment for which to calculate the scores.
type	The type of plot to display.
xlab, ylab	Plotting labels.

#### Value

An object of class SpatialNMF derived from SpatialResults, containing the fitted nnmf object and the spatial metadata.

54 SpatialPCA

#### Author(s)

```
Kylie A. Bemis
```

#### See Also

```
nnmf_als, nnmf_mult, PCA, spatialFastmap
```

### **Examples**

```
set.seed(1)
mse <- simulateImage(preset=2, npeaks=20, dim=c(10,10),
    representation="centroid")

# project to principal components
mf <- NMF(mse, ncomp=2)

# visualize first 2 components
image(mf, superpose=FALSE, scale=TRUE)</pre>
```

SpatialPCA

Principal components analysis

## Description

Compute principal components efficiently using implicitly restarted Lanczos bi-diagonalization (IRLBA) algorithm for approximate singular value decomposition.

### Usage

SpatialPCA 55

```
## S4 method for signature 'SpatialPCA'
image(x, type = "x", ...)
```

## **Arguments**

X	A dataset in P x N matrix format.
ncomp	The number of principal components to calculate.
center	Should the data be centered?
scale	Shoud the data be scaled?
nchunks	The number of chunks to use for processing.
verbose	Should progress messages be printed?
BPPARAM	$An optional instance of {\tt BiocParallelParam}. See documentation for {\tt bplapply}.$
	Options passed to irlba.
object	A SpatialPCA object.
newdata	A new SpectralImagingExperiment for which to calculate the scores.
type	The type of plot to display.

## Value

An object of class SpatialPCA derived from SpatialResults, containing the fitted prcomp\_lanczos object and the spatial metadata.

## Author(s)

```
Kylie A. Bemis
```

xlab, ylab

## See Also

```
prcomp_lanczos, NMF, spatialFastmap, irlba, svd
```

Plotting labels.

# **Examples**

```
set.seed(1)
mse <- simulateImage(preset=2, npeaks=20, dim=c(10,10),
    representation="centroid")

# project to principal components
pc <- PCA(mse, ncomp=2)

# visualize first 2 components
image(pc, superpose=FALSE, scale=TRUE)</pre>
```

56 SpatialPLS

SpatialPLS

Partial least squares (projection to latent structures)

### Description

Compute partial least squares (also called projection to latent structures or PLS). This will also perform discriminant analysis (PLS-DA) if the response is a factor. Orthogonal partial least squares options (O-PLS and O-PLS-DA) is also supported; in this case, O-PLS step is a pre-processing step to remove noise orthogonal to the response, before fitting a PLS model with a single component.

## Usage

```
## S4 method for signature 'ANY'
PLS(x, y, ncomp = 3,
    method = c("nipals", "simpls", "kernel1", "kernel2"),
    center = TRUE, scale = FALSE, bags = NULL,
    nchunks = getCardinalNChunks(),
    verbose = getCardinalVerbose(),
    BPPARAM = getCardinalBPPARAM(), ...)
## S4 method for signature 'SpectralImagingExperiment'
PLS(x, y, ncomp = 3,
    method = c("nipals", "simpls", "kernel1", "kernel2"),
    center = TRUE, scale = FALSE, ...)
## S4 method for signature 'SpatialPLS'
fitted(object, type = c("response", "class"), ...)
## S4 method for signature 'SpatialPLS'
predict(object, newdata, ncomp,
        type = c("response", "class"), simplify = TRUE, ...)
## S4 method for signature 'SpatialPLS'
topFeatures(object, n = Inf, sort.by = c("vip", "coefficients"), ...)
## S4 method for signature 'SpatialPLS, missing'
plot(x, type = c("coefficients", "vip", "scores"), ..., xlab, ylab)
## S4 method for signature 'SpatialPLS'
image(x, type = c("response", "class"), ...)
## S4 method for signature 'ANY'
OPLS(x, y, ncomp = 3, retx = TRUE,
    center = TRUE, scale = FALSE, bags = NULL,
    nchunks = getCardinalNChunks(),
    verbose = getCardinalVerbose(),
    BPPARAM = getCardinalBPPARAM(), ...)
```

SpatialPLS 57

```
## S4 method for signature 'SpectralImagingExperiment'
    OPLS(x, y, ncomp = 3, retx = FALSE,
        center = TRUE, scale = FALSE, ...)
    ## S4 method for signature 'SpatialOPLS'
    coef(object, ...)
    ## S4 method for signature 'SpatialOPLS'
    residuals(object, ...)
    ## S4 method for signature 'SpatialOPLS'
    fitted(object, type = c("response", "class"), ...)
    ## S4 method for signature 'SpatialOPLS'
    predict(object, newdata, ncomp,
             type = c("response", "class"), simplify = TRUE, ...)
    ## S4 method for signature 'SpatialOPLS'
    topFeatures(object, n = Inf, sort.by = c("vip", "coefficients"), ...)
    ## S4 method for signature 'SpatialOPLS, missing'
    plot(x, type = c("coefficients", "vip", "scores"), ..., xlab, ylab)
    ## S4 method for signature 'SpatialOPLS'
    image(x, type = c("response", "class"), ...)
Arguments
                     A dataset in P x N matrix format.
    Х
                     The response variable.
    У
                     The number of principal components to calculate.
    ncomp
                     The method used for calculating the principal components. See pls for details.
    method
    center
                     Should the data be centered?
                     Shoud the data be scaled?
    scale
                     Bags for multiple instance learning. If provided, then it is assumed all observa-
    bags
                     tions within a bag have the same label, and if a single observation is "positive"
                     then all observations in the bag are "positive". Multiple instance learning is
                     performed using mi_learn.
    retx
                     Should the (potentially large) processed data matrix be included in the result?
                     The number of chunks to use for processing.
    nchunks
    verbose
                     Should progress messages be printed?
    BPPARAM
                     An optional instance of BiocParallelParam. See documentation for bplapply.
                     Options passed to irlba.
```

A SpatialPLS or SpatialOPLS object.

object

58 SpatialResults-class

newdata	A new SpectralImagingExperiment for which to make predictions.
type	The type of fitted values to extract or the type of predictions to make.
simplify	If predictions are made using multiple numbers of components, should they be returned as a list, or simplified to an array?
n, sort.by	For topFeatures, the number of top features to return and how to sort them.
xlab, ylab	Plotting labels.

#### Value

An object of class SpatialPLS or SpatialOPLS derived from SpatialResults, containing the fitted pls or opls model and the spatial metadata.

#### Author(s)

Kylie A. Bemis

#### References

Trygg, J., & Wold, S. (2002). Orthogonal projections to latent structures (O-PLS). Journal of Chemometrics, 16(3), 119-128. doi:10.1002/cem.695

#### See Also

PCA, spatialShrunkenCentroids,

# **Examples**

```
set.seed(1)
mse <- simulateImage(preset=2, npeaks=20, dim=c(10,10), representation="centroid")
cls <- makeFactor(circle=pData(mse)$circle, square=pData(mse)$square)

# fit a PLS model with 3 components
pls <- PLS(mse, cls, ncomp=1:3)
plot(pls, type="coefficients", annPeaks="circle")

# visualize predictions
image(pls)</pre>
```

SpatialResults-class SpatialResults: Modeling results with spatial metadata

### **Description**

The SpatialResults class provides a container for modeling results with spatial metadata. Most modeling functions applied to a SpectralImagingExperiment will return a SpatialResults-derived model object.

SpatialResults-class 59

#### Usage

```
## Instance creation
SpatialResults(model, data,
    featureData = if (!missing(data)) fData(data) else NULL,
    pixelData = if (!missing(data)) pData(data) else NULL)

## S4 method for signature 'SpatialResults,ANY'
plot(x, y, ...,
    select = NULL, groups = NULL,
    superpose = TRUE, reducedDims = FALSE)

## S4 method for signature 'SpatialResults'
image(x, y, ...,
    select = NULL, subset = TRUE,
    superpose = TRUE)

## Additional methods documented below
```

#### **Arguments**

model	The model object.
data	An object (typically the original dataset) with featureData and pixelData components.
featureData	A DataFrame with feature metadata, with a row for each feature.
pixelData	A PositionDataFrame with pixel metadata, with a row for each spectrum.
x, y	The model object and results to plot. (Not typically called directly.)
	Additional options passed to plotting methods.
select	Select elements of the results to plot. For example, this selects a subset of matrix columns or a subset of factor levels to plot.
subset	A logical vector indicating which pixels to include in the image.
groups	A vector coercible to a factor indicating which of the specified spectra should be plotted with the same color.
superpose	If multiple results are plotted, should they be superposed on top of each other, or plotted seperately?
reducedDims	Does this results component represent reduced dimensions (e.g., from PCA)?

## Slots

```
model: The model.
```

featureData: A DataFrame containing feature-level metadata (e.g., a color channel, a molecular analyte, or a mass-to-charge ratio).

pixelData: A PositionDataFrame containing spatial metadata, including each observations's pixel coordinates and experimental run information.

#### Methods

```
modelData(object), modelData(object) <- value: Get or set the model slot.</pre>
```

featureData(object), featureData(object) <- value: Get or set the featureData slot.

fData(object), fData(object) <- value: Get or set the featureData slot.

featureNames(object), featureNames(object) <- value: Get or set the feature names (i.e., the row names of the featureData slot).

pixelData(object), pixelData(object) <- value: Get or set the elementMetadata slot.</pre>

pData(object), pData(object) <- value: Get or set the elementMetadata slot.

pixelNames(object), pixelNames(object) <- value: Get or set the pixel names (i.e., the row names of the elementMetadata slot).

coord(object), coord(object) <- value: Get or set the pixel coordinate columns in pixelData.</pre>

coordNames(object), coordNames(object) <- value: Get or set the names of the pixel coordinate columns in pixelData.

run(object), run(object) <- value: Get or set the experimental run column from pixelData.</pre>

runNames(object), runNames(object) <- value: Get or set the experimental run levels from
pixelData.</pre>

nrun(object): Get the number of experimental runs.

#### Author(s)

Kylie A. Bemis

#### See Also

ResultsList

SpatialShrunkenCentroids

Spatially-aware shrunken centroid clustering and classification

#### **Description**

Perform spatially-aware nearest shrunken centroid clustering or classification. These methods use statistical regularization to shrink the t-statistics of the features toward 0 so that unimportant features are removed from the model. The dissimilarity to class centroids are spatially smoothed.

# Usage

```
## S4 method for signature 'ANY,ANY'
spatialShrunkenCentroids(x, y, coord, r = 1, s = 0,
    weights = c("gaussian", "adaptive"),
    neighbors = findNeighbors(coord, r=r), bags = NULL,
    priors = table(y), center = NULL, transpose = TRUE,
```

```
nchunks = getCardinalNChunks(),
   verbose = getCardinalVerbose(),
   BPPARAM = getCardinalBPPARAM(), ...)
## S4 method for signature 'SpectralImagingExperiment, ANY'
spatialShrunkenCentroids(x, y, r = 1, s = 0,
   weights = c("gaussian", "adaptive"),
   neighbors = findNeighbors(x, r=r), ...)
## S4 method for signature 'ANY, missing'
spatialShrunkenCentroids(x, coord, r = 1, k = 2, s = 0,
    weights = c("gaussian", "adaptive"),
   neighbors = findNeighbors(coord, r=r),
   init = NULL, threshold = 0.01, niter = 10L,
   center = NULL, transpose = FALSE,
   nchunks = getCardinalNChunks(),
   verbose = getCardinalVerbose(),
   BPPARAM = getCardinalBPPARAM(), ...)
## S4 method for signature 'SpectralImagingExperiment, missing'
spatialShrunkenCentroids(x, r = 1, k = 2, s = 0,
   weights = c("gaussian", "adaptive"),
   neighbors = findNeighbors(x, r=r), ...)
## S4 method for signature 'SpatialShrunkenCentroids'
fitted(object, type = c("response", "class"), ...)
## S4 method for signature 'SpatialShrunkenCentroids'
predict(object, newdata,
        type = c("response", "class"),
        neighbors = findNeighbors(newdata, r=object$r),
       nchunks = getCardinalNChunks(),
       BPPARAM = getCardinalBPPARAM(), ...)
## S4 method for signature 'SpatialShrunkenCentroids'
logLik(object, ...)
## S4 method for signature 'SpatialShrunkenCentroids'
topFeatures(object, n = Inf, sort.by = c("statistic", "centers"), ...)
## S4 method for signature 'SpatialShrunkenCentroids, missing'
plot(x, type = c("statistic", "centers"), ..., xlab, ylab)
## S4 method for signature 'SpatialShrunkenCentroids'
image(x, type = c("probability", "class"), ...)
```

# Arguments

A spatial dataset in P x N matrix format.

у	The response variable.
coord	The spatial coordinates of the rows/columns of x. Ignored if neighbors is provided.
r	The spatial maximum distance for an observation to be considered a neighbor. Ignored if neighbors is provided.
k	The number of classes for clustering.
S	The sparsity parameter.
weights	The type of spatial weights to use for the smoothing. Gaussian weights are weighted only by distance, while adaptive weights also consider the dissimilarity between neighboring observations.
neighbors	A factor giving which observations should be treated as spatially-independent. Observations in the same group are assumed to have a spatial relationship.
bags	Bags for multiple instance learning. If provided, then it is assumed all observations within a bag have the same label, and if a single observation is "positive" then all observations in the bag are "positive". Multiple instance learning is performed using mi_learn.
priors	The (unnormalized) prior probabilities for each class.
priors center	The (unnormalized) prior probabilities for each class.  The global centroid (if known).
center	The global centroid (if known).
center transpose	The global centroid (if known).  Should x be considered P x N?
center transpose init	The global centroid (if known).  Should x be considered P x N?  A list of initial cluster configurations. (Should resemble the output of kmeans.)  Stop iteration when the proportion of cluster assignment updates is less than this
center transpose init threshold	The global centroid (if known).  Should x be considered P x N?  A list of initial cluster configurations. (Should resemble the output of kmeans.)  Stop iteration when the proportion of cluster assignment updates is less than this threshold.
center transpose init threshold	The global centroid (if known).  Should x be considered P x N?  A list of initial cluster configurations. (Should resemble the output of kmeans.)  Stop iteration when the proportion of cluster assignment updates is less than this threshold.  The maximum number of iterations.
center transpose init threshold niter nchunks	The global centroid (if known).  Should x be considered P x N?  A list of initial cluster configurations. (Should resemble the output of kmeans.)  Stop iteration when the proportion of cluster assignment updates is less than this threshold.  The maximum number of iterations.  The number of chunks to use for processing.
center transpose init threshold niter nchunks verbose	The global centroid (if known).  Should x be considered P x N?  A list of initial cluster configurations. (Should resemble the output of kmeans.)  Stop iteration when the proportion of cluster assignment updates is less than this threshold.  The maximum number of iterations.  The number of chunks to use for processing.  Should progress messages be printed?
center transpose init threshold niter nchunks verbose BPPARAM	The global centroid (if known).  Should x be considered P x N?  A list of initial cluster configurations. (Should resemble the output of kmeans.)  Stop iteration when the proportion of cluster assignment updates is less than this threshold.  The maximum number of iterations.  The number of chunks to use for processing.  Should progress messages be printed?  An optional instance of BiocParallelParam. See documentation for bplapply.
center transpose init threshold niter nchunks verbose BPPARAM	The global centroid (if known).  Should x be considered P x N?  A list of initial cluster configurations. (Should resemble the output of kmeans.)  Stop iteration when the proportion of cluster assignment updates is less than this threshold.  The maximum number of iterations.  The number of chunks to use for processing.  Should progress messages be printed?  An optional instance of BiocParallelParam. See documentation for bplapply.  Additional arguments passed to the next method.

# Value

An object of class SpatialShrunkenCentroids derived from SpatialResults, containing the fitted nscentroids object and the spatial metadata.

For topFeatures, the number of top features to return and how to sort them.

# Author(s)

Kylie A. Bemis

n, sort.by

xlab, ylab

Plotting labels.

spatialWeights 63

#### References

Bemis, K., Harry, A., Eberlin, L. S., Ferreira, C., van de Ven, S. M., Mallick, P., Stolowitz, M., and Vitek, O. (2016.) Probabilistic segmentation of mass spectrometry images helps select important ions and characterize confidence in the resulting segments. Molecular & Cellular Proteomics. doi:10.1074/mcp.O115.053918

Tibshirani, R., Hastie, T., Narasimhan, B., & Chu, G. (2003). Class Prediction by Nearest Shrunken Centroids, with Applications to DNA Microarrays. Statistical Science, 18, 104-117.

Alexandrov, T., & Kobarg, J. H. (2011). Efficient spatial segmentation of large imaging mass spectrometry datasets with spatially aware clustering. Bioinformatics, 27(13), i230-i238. doi:10.1093/bioinformatics/btr246

#### See Also

```
spatialKMeans
```

#### **Examples**

spatialWeights

Calculate spatial weights

#### **Description**

Calculate weights for neighboring observations based on either the spatial distance between the neighbors or the dissimilarity between the observations.

#### Usage

```
## S4 method for signature 'ANY'
spatialWeights(x, coord = x, r = 1, byrow = TRUE,
    neighbors = findNeighbors(coord, r=r),
    weights = c("gaussian", "adaptive"),
    sd = ((2 * r) + 1) / 4, matrix = FALSE,
    nchunks = getCardinalNChunks(),
    verbose = getCardinalVerbose(),
    BPPARAM = getCardinalBPPARAM(), ...)
```

spatialWeights

# Arguments

X	Either matrix or data frame of spatial coordinates, or a data matrix with rows or columns located at the coordinates given by coord.
coord	The spatial coordinates of the rows/columns of x. Ignored if neighbors is provided.
r	The spatial maximum distance for an observation to be considered a neighbor. Ignored if neighbors is provided.
byrow	If x is a data matrix, then are the weights calculated based on the dissimilarity between the rows (TRUE) or the columns (FALSE).
neighbors	A factor giving which observations should be treated as spatially-independent. Observations in the same group are assumed to have a spatial relationship.
weights	The type of weights to calculate. Either Gaussian weights with a constant standard deviation, or adaptive weights with a standard deviation based on the dissimilarity between the neighboring observations.
sd	The standard deviation for the Gaussian weights. Ignored with $\texttt{weights="adaptive"}$ .
matrix	Should the weights be returned as a sparse adjacency matrix instead of a list?
nchunks	The number of chunks to use for processing.
verbose	Should progress messages be printed?
BPPARAM	An optional instance of BiocParallelParam. See documentation for bplapply.
	Additional arguments passed to the next method.

# Value

Either a list of weights of neighbors or a sparse adjacency matrix (sparse\_mat).

#### Author(s)

Kylie A. Bemis

#### See Also

findNeighbors

SpectraArrays-class 65

#### **Examples**

```
set.seed(1)
mse <- simulateImage(preset=1, dim=c(10,10))

# calculate weights based on distance
spatialWeights(pixelData(mse), r=1)

# calculate weights based on spectral dissimilarity
spatialWeights(mse, r=1)</pre>
```

SpectraArrays: List of spectra arrays

# **Description**

The SpectraArrays class provides a list-like container for spectra arrays of conformable dimensions.

#### Usage

```
## Instance creation
SpectraArrays(arrays = SimpleList())
## Additional methods documented below
```

## **Arguments**

arrays A

A list of arrays.

# Details

The SpectraArrays class is intended to be flexible and the arrays do not need to be "array-like" (i.e., have non-NULL dim().) One dimensional arrays and lists are allowd. Every array must have the same NROW() and NCOL().

It supports lossless coercion to and from SimpleList.

#### Methods

All methods for SpectralImagingData also work on SpectraArrays objects. Additional methods are documented below:

```
length(object): Get the number of spectra in the object.
names(object), names(object) <- value: Get or set the names of spectra arrays in the object.
object[[i]], object[[i]] <- value: Get or set an array in the object.
object[i, j, ..., drop]: Subset as a list or array, depending on the number of dimensions of the stored spectra arrays. The result is the same class as the original object.
rbind(...), cbind(...): Combine SpectraArrays objects by row or column.
c(...): Combine SpectraArrays objects as lists.</pre>
```

#### Author(s)

Kylie A. Bemis

#### See Also

SpectralImagingData, MSImagingArrays

## **Examples**

```
set.seed(1)
x <- matrix(rlnorm(128), nrow=16, ncol=8)
y <- matrix(rlnorm(128), nrow=16, ncol=8)
s <- SpectraArrays(list(x=x, y=y))
print(s)</pre>
```

SpectralImagingArrays-class

SpectralImagingArrays: Spectral imaging data with arbitrary domain

#### **Description**

The SpectralImagingArrays class provides a list-like container for high-dimensional spectral imaging data where every spectrum may have its own domain values. It is designed to provide easy access to raw individual spectra, but images cannot be easily reconstructed.

The MSImagingArrays class extends SpectralImagingArrays for mass spectrometry-based imaging experiments with unaligned mass features.

# Usage

# Arguments

spectraData Either a list-like object with lists of individual spectra and lists of their domain values, or a SpectraArrays instance.

pixelData A PositionDataFrame with pixel metadata, with a row for each spectrum.

metadata A list with experimental-level metadata.

# Slots

spectraData: A SpectraArrays object storing one or more array-like data elements with conformable dimensions.

elementMetadata: A PositionDataFrame containing spectrum-level metadata, including each spectrum's pixel coordinates and experimental run information.

processing: A list containing unexecuted ProcessingStep objects.

#### Methods

All methods for SpectralImagingData also work on SpectralImagingArrays objects. Additional methods are documented below:

```
length(object): Get the number of spectra in the object.
```

object[i, ..., drop]: Subset as a list based on the spectra. The result is the same class as the original object.

rbind(...), cbind(...): Combine SpectralImagingArrays objects by row or column.

#### Author(s)

Kylie A. Bemis

### See Also

SpectralImagingData, MSImagingArrays

# **Examples**

```
set.seed(1)
x <- replicate(9, rlnorm(10), simplify=FALSE)
t <- replicate(9, sort(runif(10)), simplify=FALSE)
coord <- expand.grid(x=1:3, y=1:3)

sa <- SpectralImagingArrays(
    spectraData=list(intensity=x, wavelength=t),
    pixelData=PositionDataFrame(coord))

print(sa)</pre>
```

SpectralImagingData-class

SpectralImagingData: Abstract class for spectral imaging data

#### **Description**

The SpectralImagingData class is an abstract container for high-dimensional spectral imaging data. Every spectrum is associated with spatial coordinates so that an image can be constructed from the spectral intensities.

The SpectralImagingArrays and SpectralImagingExperiment classes directly extend this class, where SpectralImagingArrays is primarily intended for unprocessed spectra with unaligned features, and SpectralImagingExperiment is intended for processed spectra with aligned features.

The MSImagingArrays and MSImagingExperiment classes further extend these classes for mass spectrometry imaging data.

#### Slots

spectraData: A SpectraArrays object storing one or more array-like data elements with conformable dimensions.

elementMetadata: A PositionDataFrame containing spectrum-level metadata, including each spectrum's pixel coordinates and experimental run information.

processing: A list containing unexecuted ProcessingStep objects.

#### Methods

```
spectraData(object, ...), spectraData(object, ...) <- value: Get or set the spectraData
slot.</pre>
```

spectraNames(object, ...), spectraNames(object, ...) <- value: Get or set the names of
 the spectra in the spectraData slot.</pre>

spectra(object, i = 1L, ...), spectra(object, i = 1L, ...) <- value: Get or set a specific spectra array in the spectraData slot.

pixelData(object), pixelData(object) <- value: Get or set the elementMetadata slot.</pre>

pData(object), pData(object) <- value: Get or set the elementMetadata slot.

pixelNames(object), pixelNames(object) <- value: Get or set the pixel names (i.e., the row names of the elementMetadata slot).

spectraVariables(object, ...): Get the names of the spectrum-level variables (i.e., the columns of the elementMetadata slot).

coord(object), coord(object) <- value: Get or set the pixel coordinate columns in pixelData.</pre>

coordNames(object), coordNames(object) <- value: Get or set the names of the pixel coordinate columns in pixelData.

run(object), run(object) <- value: Get or set the experimental run column from pixelData.

runNames(object), runNames(object) <- value: Get or set the experimental run levels from pixelData.

nrun(object): Get the number of experimental runs.

is3D(object): Check if the number of spatial dimensions is greater than 2.

#### Author(s)

Kylie A. Bemis

#### See Also

SpectralImagingExperiment, SpectralImagingArrays, MSImagingExperiment, MSImagingArrays

```
SpectralImagingExperiment-class
```

SpectralImagingExperiment: Spectral imaging data with shared domain

## Description

The SpectralImagingExperiment class provides a matrix-like container for high-dimensional spectral imaging data where every spectrum shares the same domain values. It is designed to provide easy access to both the spectra (as columns) and sliced images (as rows).

The MSImagingExperiment class extends SpectralImagingExperiment for mass spectrometry-based imaging experiments with aligned mass features.

## Usage

```
## Instance creation
SpectralImagingExperiment(spectraData = SimpleList(),
    featureData = DataFrame(), pixelData = PositionDataFrame(),
    metadata = list())
## Additional methods documented below
```

#### **Arguments**

spectraData Either a matrix-like object with number of rows equal to the number of features

and number of columns equal to the number of pixels, a list of such objects, or

a SpectraArrays instance.

featureData A DataFrame with feature metadata, with a row for each feature.

pixelData A PositionDataFrame with pixel metadata, with a row for each spectrum.

metadata A list with experimental-level metadata.

#### **Slots**

spectraData: A SpectraArrays object storing one or more array-like data elements with conformable dimensions.

featureData: A DataFrame containing feature-level metadata (e.g., a color channel, a molecular analyte, or a mass-to-charge ratio).

elementMetadata: A PositionDataFrame containing spectrum-level metadata, including each spectrum's pixel coordinates and experimental run information.

processing: A list containing unexecuted ProcessingStep objects.

70 spectrapply

#### Methods

All methods for SpectralImagingData also work on SpectralImagingExperiment objects. Additional methods are documented below:

```
featureData(object), featureData(object) <- value: Get or set the featureData slot.</pre>
```

fData(object), fData(object) <- value: Get or set the featureData slot.

featureNames(object), featureNames(object) <- value: Get or set the feature names (i.e., the row names of the featureData slot).

length(object): Get the number of spectra in the object.

nrow(object), ncol(object): Get the number of rows (features) or the number of columns (pixels) in the object.

object[i, j, ..., drop]: Subset based on the rows (featureData) and the columns (pixelData). The result is the same class as the original object.

rbind(...), cbind(...): Combine SpectralImagingExperiment objects by row or column.

## Author(s)

Kylie A. Bemis

#### See Also

SpectralImagingData, MSImagingExperiment

#### **Examples**

```
set.seed(1)
x <- matrix(rlnorm(81), nrow=9, ncol=9)
index <- 1:9
coord <- expand.grid(x=1:3, y=1:3)

se <- SpectralImagingExperiment(
    spectraData=x,
    featureData=DataFrame(index=1:9),
    pixelData=PositionDataFrame(coord))

print(se)</pre>
```

spectrapply

Apply functions over spectra

#### Description

Apply a user-specified function over all spectra in a spectral imaging dataset.

spectrapply 71

#### Usage

```
## S4 method for signature 'MSImagingExperiment'
spectrapply(object, FUN, ...,
   spectra = "intensity", index = "mz")
## S4 method for signature 'MSImagingArrays'
spectrapply(object, FUN, ...,
    spectra = "intensity", index = "mz")
## S4 method for signature 'SpectralImagingExperiment'
spectrapply(object, FUN, ...,
       spectra = "intensity", index = NULL,
       simplify = TRUE, outpath = NULL,
       nchunks = getCardinalNChunks(),
       verbose = getCardinalVerbose(),
       BPPARAM = getCardinalBPPARAM())
## S4 method for signature 'SpectralImagingArrays'
spectrapply(object, FUN, ...,
       spectra = "intensity", index = NULL,
        simplify = TRUE, outpath = NULL,
       nchunks = getCardinalNChunks(),
       verbose = getCardinalVerbose(),
       BPPARAM = getCardinalBPPARAM())
```

# Arguments

object	A spectral imaging dataset.
FUN	A function to be applied. The function <i>must</i> accept (1) a first argument giving the spectral intensities as a numeric vector and (2) a second argument giving the intensity locations (e.g., m/z values) as a numeric vector.
	Options passed to chunkMapply or chunkApply.
spectra	The name of the array in spectraData() to use for the spectral intensities.
index	The name of the array in spectraData() (for SpectralImagingArrays) or column in featureData() (for SpectralImagingExperiment) to use for the spectral locations.
simplify	Should the result be simplified to an array if possible?
outpath	Optional. The name of a file to write the resulting data.
nchunks	The number of chunks to use for processing.
verbose	Should progress messages be printed?
BPPARAM	$An optional instance of {\tt BiocParallelParam}. See documentation for {\tt bplapply}.$

## Value

A list if simplify=FALSE. Otherwise, a vector or matrix, or a higher-dimensional array if the attempted simplification is successful.

72 subsetFeatures

#### Author(s)

```
Kylie A. Bemis
```

#### See Also

```
summarizeFeatures, summarizePixels
```

# **Examples**

```
set.seed(1)
mse <- simulateImage(preset=1, npeaks=10, dim=c(10,10))
# find m/z locations of peaks in each spectrum
peaks <- spectrapply(mse, function(x, mz) mz[matter::findpeaks(x)])
head(peaks[[1L]])
head(peaks[[2L]])</pre>
```

subsetFeatures

Subset a spectral imaging dataset

## **Description**

Returns a subset of the dataset that meets the conditions.

## Usage

```
## S4 method for signature 'SpectralImagingArrays'
subset(x, subset, ...)

## S4 method for signature 'SpectralImagingExperiment'
subset(x, select, subset, ...)

subsetFeatures(x, ...)

subsetPixels(x, ...)
```

## **Arguments**

X	A spectral imaging dataset.
select	Logical expression to be evaluated in the object's featureData() indicating which rows (features) to keep.
subset	Logical expression to be evaluated in the object's pixelData() indicating which columns (pixels) to keep.
	Conditions describing rows (features) or columns (pixels) to be retained. Passed to features() and pixels() methods to obtain the subset indices.

summarizeFeatures 73

#### Value

An object of the same class as x with the appropriate subsetting applied to it.

## Author(s)

Kylie A. Bemis

# **Examples**

```
set.seed(1)
mse <- simulateImage(preset=1, npeaks=10, dim=c(10,10))
# subset features to mass range 1000 - 1500
subsetFeatures(mse, 1000 < mz, mz < 1500)
# select pixels to coordinates x = 1..3, y = 1..3
subsetPixels(mse, x <= 3, y <= 3)
# subset both features + pixels
subset(mse, 1000 < mz & mz < 1500, x <= 3 & y <= 3)</pre>
```

summarizeFeatures

Summarize a spectral imaging dataset

# **Description**

Summarizes over the rows or columns of the dataset.

# Usage

74 summarizeFeatures

```
nchunks = getCardinalNChunks(),
    verbose = getCardinalVerbose(),
    BPPARAM = getCardinalBPPARAM())

## S4 method for signature 'SpectralImagingExperiment'
rowSums(x, na.rm = FALSE, dims = 1, ...)

## S4 method for signature 'SpectralImagingExperiment'
colSums(x, na.rm = FALSE, dims = 1, ...)

## S4 method for signature 'SpectralImagingExperiment'
rowMeans(x, na.rm = FALSE, dims = 1, ...)

## S4 method for signature 'SpectralImagingExperiment'
colMeans(x, na.rm = FALSE, dims = 1, ...)
```

# Arguments

X	A spectral imaging dataset.
stat	The name of summary statistics to compute over the rows or columns of a matrix. Allowable values include: "min", "max", "prod", "sum", "mean", "var", "sd", "any", "all", and "nnzero".
groups	A vector coercible to a factor giving groups to summarize.
na.rm	If TRUE, remove NA values before summarizing.
dims	Ignored.
nchunks	The number of chunks to use for processing.
verbose	Should progress messages be printed?
BPPARAM	An optional instance of BiocParallelParam. See documentation for bplapply.
	Additional arguments passed to rowStats or colStats, such as the number of chunks.

# Value

For summarizeFeatures and summarizePixels, an object of the same class as x with the statistical summaries added as columns in the featureData() or pixelData(), respectively.

For rowStats, colStats, etc., a vector, matrix, or array with the summary statistics.

# Author(s)

Kylie A. Bemis

# **Examples**

```
set.seed(1)
mse <- simulateImage(preset=1, npeaks=10, dim=c(10,10))
# summarize mean spectrum</pre>
```

writeMSIData 75

```
mse <- summarizeFeatures(mse, stat="mean")
plot(mse, "mean")

# summarize total ion current
mse <- summarizePixels(mse, stat=c(TIC="sum"))
image(mse, "TIC")</pre>
```

writeMSIData

Write mass spectrometry imaging data files

## **Description**

Write supported mass spectrometry imaging data files, including imzML and Analyze 7.5.

#### Usage

```
writeMSIData(object, file, ...)
## S4 method for signature 'MSImagingExperiment_OR_Arrays'
writeImzML(object, file, bundle = TRUE,
verbose = getCardinalVerbose(), ...)
## S4 method for signature 'MSImagingExperiment'
writeAnalyze(object, file, verbose = getCardinalVerbose(), ...)
## S4 method for signature 'SpectralImagingExperiment'
writeAnalyze(object, file, verbose = getCardinalVerbose(), ...)
```

# **Arguments**

object	A spectral imaging dataset.
file	The absolute or relative file path. The file extension must be included for $\verb write  MSIData$ .
bundle	Should the ".imzML" and ".ibd" files be bundled into a new directory of the same name?
verbose	Should progress messages be printed?
	Additional arguments passed to writeImzML or writeAnalyze.

# **Details**

The writeImzML function supports writing both the "continuous" and "processed" formats.

Exporting the experimental metadata to cvParam tags is lossy, and not all metadata will be preserved. If exporting an object that was originally imported from an imzML file, only metadata that appears in experimentData() will be preserved when writing.

Datasets with multiple experimental runs will be merged into a single file. The object's pixelData() and featureData() will also be written to tab-delimted files if appropriate. These will be read back in by readImzML().

The imzML files can be modified after writing (such as to add additional experimental metadata) using the Java-based imzMLValidator application: https://gitlab.com/imzML/imzMLValidator/.

#### Value

TRUE if the file was written successfully, with the output file paths attached as an attribute.

#### Author(s)

Kylie A. Bemis

#### References

Schramm T, Hester A, Klinkert I, Both J-P, Heeren RMA, Brunelle A, Laprevote O, Desbenoit N, Robbe M-F, Stoeckli M, Spengler B, Rompp A (2012) imzML - A common data format for the flexible exchange and processing of mass spectrometry imaging data. Journal of Proteomics 75 (16):5106-5110. doi:10.1016/j.jprot.2012.07.026

## See Also

readMSIData

XDataFrame-class

XDataFrame: Extended data frame with key columns

# **Description**

The XDataFrame extends the DataFrame class from the S4Vectors package with support for columns (or sets of columns) designated as keys.

#### Usage

```
XDataFrame(..., keys = list())
```

# Arguments

.. Arguments passed to the DataFrame().

keys A named list of character vectors giving the names of key columns. The names of the list become the names of the keys (which may be different from the

of the list become the names of the keys (which may be different from the columns). The character vectors specify the names of columns that compose

that key.

#### **Details**

For the most part, XDataFrame behaves identically to DataFrame, and key columns can be get or set as usual. The XDataFrame class is primarily intended as a way to enforce additional requirements or constraints on specific sets of columns in a structured way. It provides an abstracted way of manipulating sets of columns that are expected to follow certain rules. The keys remain consistent and accessible even if the columns of the data frame are renamed.

XDataFrame-class 77

The base class currently has only minimal requirements for keys (i.e., that they are valid columns in the data frame). Additionally, keys are checked for compatibility when combining data frames. Uniqueness is *not* checked.

Subclasses can enforce additional constraints on key columns. For example, the PositionDataFrame and MassDataFrame classes.

# Methods

```
keys(object, i = NULL, ..., drop = TRUE), keys(object, i = NULL, ...) <- value: Get or set the key columns. By default, this gets or sets the keys slot. Provide i to get or set specific keys.
```

# Author(s)

Kylie A. Bemis

# See Also

DataFrame, MassDataFrame, PositionDataFrame

# **Examples**

```
## Create an XDataFrame object
XDataFrame(id=1:10, letter=LETTERS[1:10], keys=list(index="id"))
```

# **Index**

* IO	colocalized, 5
readMSIData, 32	* models
writeMSIData, 75	MeansTest, 11
* classes	* multivariate
MassDataFrame-class, 10	SpatialFastmap, 49
MSImagingArrays-class, 13	SpatialNMF, 53
MSImagingExperiment-class, 15	SpatialPCA, 54
PositionDataFrame-class, 29	SpatialPLS, 56
ResultsList-class, 37	* package
SpatialResults-class, 58	Cardinal-package, 3
SpectraArrays-class, 65	* regression
SpectralImagingArrays-class, 66	MeansTest, 11
SpectralImagingData-class, 67	SpatialCV, 45
SpectralImagingExperiment-class,	* smooth
69	smooth, 43
XDataFrame-class, 76	* spatial
* classif	findNeighbors,9
SpatialCV, 45	SpatialDGMM, 46
SpatialPLS, 56	SpatialFastmap, 49
SpatialShrunkenCentroids, 60	SpatialKMeans, 51
* clustering	SpatialShrunkenCentroids, 60
SpatialDGMM, 46	spatialWeights, 63
SpatialKMeans, 51	* <b>ts</b>
SpatialShrunkenCentroids, 60	bin, 4
* datagen	estimateDomain,7
simulateSpectra, 39	normalize, 17
* hplot	peakAlign, 18
plot-image, 25	peakPick, 20
plot-spectra, 27	peakProcess, 22
* htest	process, 30
MeansTest, 11	recalibrate, 34
* iplot	reduceBaseline, 35
selectROI, 38	smooth, 43
* manip	* univar
process, 30	summarizeFeatures, 73
slice, 42	* utilities
spectrapply, 70	estimateDomain,7
subsetFeatures, 72	features, 8
* methods	pixels, 24

SpatialCV, 45	(XDataFrame-class), 76
[,MassDataFrame,ANY,ANY,ANY-method	
(MassDataFrame-class), 10	addProcessing,SpectralImagingData-method
[,PositionDataFrame,ANY,ANY,ANY-method	(process), 30
(PositionDataFrame-class), 29	addShape(simulateSpectra), 39
[,SpectraArrays,ANY,ANY,ANY-method	aggregate, SpectralImagingExperiment-method
(SpectraArrays-class), 65	(deprecated), 7
[,SpectralImagingArrays,ANY,ANY,ANY-method	alpha.colors (deprecated), 7
(SpectralImagingArrays-class), 66	approx1, 5
$\hbox{\tt [,SpectralImagingExperiment,ANY,ANY,ANY-meth]}$	106aselineReduction(deprecated) 7
(SpectralImagingExperiment-class),	baselineReduction<- (deprecated), 7
69	bin, 4
[,XDataFrame,ANY,ANY,ANY-method	bin, MSImagingArrays-method (bin), 4
(XDataFrame-class), 76	bin, MSImagingExperiment-method (bin), 4
[<-, SpectraArrays, ANY, ANY, ANY-method	bin, SpectralImagingArrays-method (bin),
(SpectraArrays-class), 65	4
[<-,SpectralImagingArrays,ANY,ANY,ANY-method	bin,SpectralImagingExperiment-method
(SpectralImagingArrays-class),	(bin), 4
66	binpeaks, 19
[<-, Spectral Imaging Experiment, ANY, ANY, ANY-meanth of the property of	ethetannly, 3, 6, 8, 12, 19, 22, 31, 33, 40, 46.
(SpectralImagingExperiment-class),	48, 50, 52, 55, 57, 62, 64, 71, 74
69	bw.colors (deprecated), 7
<pre>[&lt;-,XDataFrame,ANY,ANY,ANY-method</pre>	Switter of (depressated), /
(XDataFrame-class), 76	c,MSImagingArrays-method
[[,SpatialResults-method	(MSImagingArrays-class), 13
(SpatialResults-class), 58	c,SpectraArrays-method
[[,SpectraArrays-method	(SpectraArrays-class), 65
(SpectraArrays-class), 65	c,SpectralImagingArrays-method
[[,SpectralImagingData-method	(SpectralImagingArrays-class),
(SpectralImagingData-class), 67	66
[[<-,SpectraArrays-method	Cardinal (Cardinal-package), 3
(SpectraArrays-class), 65	Cardinal-package, 3
[[<-,SpectralImagingData-method	CardinalVersion (Cardinal-package), 3
(SpectralImagingData-class), 67	cbind, MSImagingExperiment-method
[[<-,XDataFrame-method	(MSImagingExperiment-class), 15
(XDataFrame-class), 76	cbind, SpectraArrays-method
<pre>\$,SpatialResults-method</pre>	(SpectraArrays-class), 65
(SpatialResults-class), 58	cbind, SpectralImagingArrays-method
\$,SpectraArrays-method	(SpectralImagingArrays-class),
(SpectraArrays-class), 65	66
<pre>\$,SpectralImagingData-method</pre>	cbind, SpectralImagingExperiment-method
(SpectralImagingData-class), 67	(SpectralImagingExperiment-class),
\$<-,SpectraArrays-method	69
(SpectraArrays-class), 65	cbind,XDataFrame-method
<pre>\$&lt;-,SpectralImagingData-method</pre>	(XDataFrame-class), 76
(SpectralImagingData-class), 67	centroided, MSImagingExperiment_OR_Arrays-method
\$<-,XDataFrame-method	(MSImagingExperiment-class), 15

<pre>centroided&lt;-,MSImagingExperiment_OR_Arrays</pre>	s-me <b>thas</b> s:SpectralImagingData
(MSImagingExperiment-class), 15	(SpectralImagingData-class), 67
chunk_colapply, 31	class:SpectralImagingExperiment
chunk_mapply, 31	(SpectralImagingExperiment-class),
chunkApply, 6, 71	69
chunkLapply, 8	<pre>class:XDataFrame (XDataFrame-class), 76</pre>
chunkMapply, 71	class: XDFrame (XDataFrame-class), 76
class:Hashmat (deprecated), 7	classNameForDisplay,XDFrame-method
class:IAnnotatedDataFrame (deprecated),	(XDataFrame-class), 76
7	coef, SpatialOPLS-method (SpatialPLS), 56
class:ImageData(deprecated),7	coerce, DataFrame, XDataFrame-method
class:iSet (deprecated), 7	(XDataFrame-class), 76
class:MassDataFrame	coerce, DFrame, MassDataFrame-method
(MassDataFrame-class), 10	(MassDataFrame-class), 10
class: MeansTest (MeansTest), 11	coerce, DFrame, PositionDataFrame-method
class:MIAPE-Imaging (deprecated), 7	(PositionDataFrame-class), 29
class: MSImageData (deprecated), 7	coerce,list,SpectraArrays-method
class:MSImageProcess (deprecated), 7	(SpectraArrays-class), 65
class: MSImageSet (deprecated), 7	coerce, MSImageSet, MSImagingExperiment-method
class:MSImagingArrays	(MSImagingExperiment-class), 15
(MSImagingArrays-class), 13	coerce, MSImagingArrays, MSImagingExperiment-method
class:MSImagingExperiment	(readMSIData), 32
(MSImagingExperiment-class), 15	coerce, MSImagingExperiment, MSImagingArrays-method
class:PositionDataFrame	(readMSIData), 32
(PositionDataFrame-class), 29	coerce, SimpleList, SpectraArrays-method
class:ResultSet (deprecated), 7	(SpectraArrays-class), 65
class:ResultsList (ResultsList-class),	coerce, SpectraArrays, list-method
37	(SpectraArrays-class), 65
class:SImageData (deprecated), 7	coerce, SpectraArrays, SimpleList-method
class:SImageSet (deprecated), 7	(SpectraArrays-class), 65
class:SpatialCV (SpatialCV), 45	col.map (deprecated), 7
class:SpatialDGMM (SpatialDGMM), 46	colMeans, SpectralImagingExperiment-method
class:SpatialFastmap(SpatialFastmap),	(summarizeFeatures), 73
49	colnames, SpectralImagingExperiment-method
class: SpatialKMeans (SpatialKMeans), 51	(SpectralImagingExperiment-class),
class:SpatialNMF (SpatialNMF), 53	69
class:SpatialOPLS (SpatialPLS), 56	colnames<-, SpectralImagingExperiment-method
class:SpatialPCA (SpatialPCA), 54	(SpectralImagingExperiment-class),
class:SpatialPLS (SpatialPLS), 56	69
class:SpatialResults	colocalized, 5
(SpatialResults-class), 58	colocalized,MSImagingExperiment-method
class:SpatialShrunkenCentroids	(colocalized), 5
(SpatialShrunkenCentroids), 60	colocalized,SpatialDGMM-method
class:SpectraArrays	(colocalized), 5
(SpectraArrays-class), 65	colocalized,SpectralImagingExperiment-method
class:SpectralImagingArrays	(colocalized), 5
(SpectralImagingArrays-class),	<pre>color.map (deprecated), 7</pre>
66	colStats, 74

colStats,SpectralImagingExperiment-method	cv_do, 46
(summarizeFeatures), 73	cvApply (deprecated), 7
colSums, SpectralImagingExperiment-method	
(summarizeFeatures), 73	darkmode (deprecated), 7
combine, SpectraArrays, ANY-method	DataFrame, 59, 69, 76, 77
(SpectraArrays-class), 65	deprecated, 7
combine, SpectralImagingArrays, ANY-method	dim,SpectraArrays-method
(SpectralImagingArrays-class),	(SpectraArrays-class), 65
66	dim, SpectralImagingArrays-method
<pre>combine,SpectralImagingExperiment,ANY-method</pre>	(SpectralImagingArrays-class),
(SpectralImagingExperiment-class),	66
69	dim,SpectralImagingExperiment-method
convertMSImagingArrays2Experiment	(SpectralImagingExperiment-class),
(readMSIData), 32	69
convertMSImagingExperiment2Arrays	discrete.colors (deprecated), 7
(readMSIData), 32	divergent.colors (deprecated), 7
coord (PositionDataFrame-class), 29	downsample, 28
coord, PositionDataFrame-method	dpal (plot-image), 25
(PositionDataFrame-class), 29	drle, 48
coord, SpatialResults-method	ui 1e, 40
(SpatialResults-class), 58	h 26
coord, SpectralImagingData-method	enhance, 26
(SpectralImagingData-class), 67	estbase, 36
	estbase_hull, 36
coord<- (PositionDataFrame-class), 29	estbase_loc, 36
coord<-,PositionDataFrame-method	estbase_med, 36
(PositionDataFrame-class), 29	estbase_snip, 36
coord<-,SpatialResults-method	estimateDomain, 5, 7
(SpatialResults-class), 58	estimateReferenceMz, 5
coord<-,SpectralImagingData-method	estimateReferenceMz (estimateDomain), 7
(SpectralImagingData-class), 67	estimateReferencePeaks, 21
coordNames (PositionDataFrame-class), 29	estimateReferencePeaks
coordNames,PositionDataFrame-method	(estimateDomain), 7
(PositionDataFrame-class), 29	estnoise_diff, 21
coordNames,SpatialResults-method	estnoise_filt, 21
(SpatialResults-class), 58	estnoise_mad, 21
coordNames,SpectralImagingData-method	estnoise_quant, 21
(SpectralImagingData-class), 67	estnoise_sd, 21
<pre>coordNames&lt;- (PositionDataFrame-class),</pre>	experimentData,MSImagingExperiment_OR_Arrays-method (MSImagingExperiment-class), 15
coordNames<-,PositionDataFrame-method	<pre>experimentData&lt;-,MSImagingExperiment_OR_Arrays,ANY-method</pre>
(PositionDataFrame-class), 29	(MSImagingExperiment-class), 15
coordNames<-,SpatialResults-method	
(SpatialResults-class), 58	fastmap, 50
coordNames<-,SpectralImagingData-method	fData, Spatial Results-method
(SpectralImagingData-class), 67	(SpatialResults-class), 58
coregister (colocalized), 5	fData, Spectral Imaging Experiment-method
cpal (plot-image), 25	(SpectralImagingExperiment-class),
crossValidate (SpatialCV), 45	69
0. 000, GIIIGGC   OPGCIGIC   /. TJ	

fData<-,SpatialResults,ANY-method	$\verb findNeighbors,SpectralImagingData-method \\$
(SpatialResults-class), 58	(findNeighbors), 9
fData<-,SpectralImagingExperiment,ANY-method	findpeaks, 8, 20, 22
(SpectralImagingExperiment-class),	findpeaks_cwt, 21
69	fitted,ResultsList-method
featureApply(deprecated),7	(ResultsList-class), 37
${\sf feature Apply, Spectral Imaging Experiment-method}$	
(deprecated), 7	fitted, SpatialOPLS-method (SpatialPLS),
featureData,SpatialResults-method	56
(SpatialResults-class), 58	fitted, SpatialPLS-method (SpatialPLS),
$feature {\tt Data}, {\tt SpectralImaging Experiment-method}$	56
(SpectralImagingExperiment-class),	fitted, SpatialResults-method
69	(SpatialResults-class), 58
featureData<-,SpatialResults,ANY-method	fitted, Spatial Shrunken Centroids-method
(SpatialResults-class), 58	(SpatialShrunkenCentroids), $60$
featureData <- , $SpectralImagingExperiment$ , $ANY-mathright -mathrix$	method
(SpectralImagingExperiment-class),	getCardinalBPPARAM (Cardinal-package), 3
69	getCardinalNChunks (Cardinal-package), 3
featureNames,SpatialResults-method	getCardinalNumBlocks (deprecated), 7
(SpatialResults-class), 58	getCardinalVerbose (Cardinal-package), 3
featureNames, SpectralImagingExperiment-method	gridded (depresented) 7
$({\tt SpectralImagingExperiment-class}),$	gridded (deprecated), 7
69	gridded<- (deprecated), 7
featureNames<-,SpatialResults-method	Hashmat (deprecated), 7
(SpatialResults-class), 58	Hashmat-class (denrecated) 7
reacurenames<-, spectrall maging experiment-metr	height (deprecated), 7
(SpectralImagingExperiment-class),	height<- (deprecated), 7
69	(35)
features, 8	<pre>IAnnotatedDataFrame (deprecated), 7</pre>
features, MSI maging Experiment-method	<pre>IAnnotatedDataFrame-class (deprecated),</pre>
(features), 8	7
features, SpectralImagingExperiment-method	iData (deprecated), 7
(features), 8	iData, ANY-method (deprecated), 7
filt1, 44	iData<- (deprecated), 7
filt1_adapt, 44	iData<-, ANY-method (deprecated), 7
filt1_bi, 44	image, $26, 38$
filt1_diff, 44	<pre>image (plot-image), 25</pre>
filt1_gauss, 44	<pre>image,MSImagingExperiment-method</pre>
filt1_guide, <i>44</i>	(plot-image), 25
filt1_ma, <i>44</i>	<pre>image,ResultsList-method</pre>
filt1_pag, <i>44</i>	(ResultsList-class), 37
filt1_sg, <i>44</i>	<pre>image,SpatialCV-method(SpatialCV), 45</pre>
filt2, <u>26</u>	${\tt image,SpatialDGMM-method}~({\tt SpatialDGMM}),$
findNeighbors, 9, 64	46
findNeighbors,ANY-method	<pre>image,SpatialFastmap-method</pre>
(findNeighbors), 9	(SpatialFastmap), 49
findNeighbors,PositionDataFrame-method	<pre>image,SpatialKMeans-method</pre>
(findNeighbors),9	(SpatialKMeans), 51

image, SpatialNMF-method (SpatialNMF), 53	kmeans, $52$
<pre>image,SpatialOPLS-method(SpatialPLS),</pre>	
56	length, SpatialResults-method
<pre>image,SpatialPCA-method(SpatialPCA), 54</pre>	(SpatialResults-class), 58
image, SpatialPLS-method (SpatialPLS), 56	length, SpectraArrays-method
image,SpatialResults-method	(SpectraArrays-class), 65
(SpatialResults-class), 58	length,SpectralImagingArrays-method
<pre>image,SpatialShrunkenCentroids-method</pre>	(SpectralImagingArrays-class),
(SpatialShrunkenCentroids), 60	66
<pre>image,SpectralImagingExperiment-method</pre>	length,SpectralImagingExperiment-method
(plot-image), 25	(SpectralImagingExperiment-class),
image3D (plot-image), 25	69
<pre>image3D,MSImagingExperiment-method</pre>	lightmode (deprecated), 7
(plot-image), 25	lineScanDirection (deprecated), 7
ImageData (deprecated), 7	1m, 12, 13
<pre>ImageData-class (deprecated), 7</pre>	lme, 12, 13
ImzMeta, <i>14–16</i>	locator, 38
<pre>instrumentVendor(deprecated), 7</pre>	logLik,SpatialDGMM-method
intensity,MSImagingArrays-method	(SpatialDGMM), 46
(MSImagingArrays-class), 13	logLik, Spatial Shrunken Centroids-method
<pre>intensity,MSImagingExperiment-method</pre>	(SpatialShrunkenCentroids), 60
(MSImagingExperiment-class), 15	mala Fastan (aslast DOI) 20
intensity.colors(deprecated),7	makeFactor (selectROI), 38
intensity<-,MSImagingArrays-method	massAnalyzerType (deprecated), 7
(MSImagingArrays-class), 13	MassDataFrame, 15, 16, 30, 40, 77
<pre>intensity&lt;-,MSImagingExperiment-method</pre>	MassDataFrame (MassDataFrame-class), 10
(MSImagingExperiment-class), 15	MassDataFrame-class, 10
intensityData (deprecated), 7	matrixApplication (deprecated), 7
<pre>intensityData&lt;- (deprecated), 7</pre>	MeansTest, 11
<pre>ionizationType (deprecated), 7</pre>	meansTest, 47
irlba, 53, 55, 57	meansTest (MeansTest), 11
is3D (deprecated), 7	meansTest, ANY-method (MeansTest), 11
is3D,PositionDataFrame-method	meansTest, SpatialDGMM-method
(PositionDataFrame-class), 29	(MeansTest), 11
is3D,SpectralImagingData-method	meansTest, SpectralImagingExperiment-method
(SpectralImagingData-class), 67	(MeansTest), 11
isCentroided, MSImagingExperiment_OR_Arrays-r	merganaska 10
(MSImagingExperiment-class), 15	mergepeaks, 19
iSet (deprecated), 7	mi_learn, 57, 62
iSet-class (deprecated), 7	MIAPE-Imaging (deprecated), 7
	MIAPE-Imaging-class (deprecated), 7
jet.colors(deprecated),7	modelData (SpatialResults-class), 58
Leave (VDataFrame alleas) 76	modelData, SpatialResults-method
keys (XDataFrame-class), 76	(SpatialResults-class), 58
keys, XDataFrame-method	modelData<- (SpatialResults-class), 58
(XDataFrame-class), 76	modelData<-,SpatialResults-method
keys<- (XDataFrame-class), 76	(SpatialResults-class), 58
keys<-,XDataFrame-method	msiInfo (deprecated), 7
(XDataFrame-class), 76	MSImageData (deprecated), 7

MSImageData-class (deprecated), 7	66
MSImageProcess (deprecated), 7	names, Spectral Imaging Experiment-method
MSImageProcess-class (deprecated), 7	(SpectralImagingExperiment-class)
MSImageSet (deprecated), 7	69
MSImageSet-class (deprecated), 7	names<-,SpectraArrays-method
MSImagingArrays, 15, 16, 34, 66–69	(SpectraArrays-class), 65
MSImagingArrays	names<-,SpectralImagingArrays-method
(MSImagingArrays-class), 13	(SpectralImagingArrays-class),
MSImagingArrays-class, 13	66
MSImagingExperiment, <i>13</i> , <i>14</i> , <i>34</i> , <i>42</i> , <i>68</i> – <i>70</i>	names<-,SpectralImagingExperiment-method
MSImagingExperiment	(SpectralImagingExperiment-class)
(MSImagingExperiment-class), 15	69
MSImagingExperiment-class, 15	names<-,XDataFrame-method
mz,MassDataFrame-method	(XDataFrame-class), 76
(MassDataFrame-class), 10	NMF, 50, 55
mz,missing-method	NMF (SpatialNMF), 53
(MSImagingExperiment-class), 15	NMF, ANY-method (SpatialNMF), 53
mz,MSImagingArrays-method	NMF,SpectralImagingExperiment-method
(MSImagingArrays-class), 13	(SpatialNMF), 53
mz,MSImagingExperiment-method	nnmf, <i>53</i>
(MSImagingExperiment-class), 15	nnmf_als, 54
mz<-,MassDataFrame-method	nnmf_mult, <i>54</i>
(MassDataFrame-class), 10	normalization (deprecated), $7$
mz<-,MSImagingArrays-method	normalization<- (deprecated), 7
(MSImagingArrays-class), 13	normalize, 17, <i>31</i> , <i>35</i> , <i>36</i> , <i>44</i>
<pre>mz&lt;-,MSImagingExperiment-method</pre>	normalize,SpectralImagingData-method
(MSImagingExperiment-class), 15	(normalize), 17
mzAlign (deprecated), 7	nrun,PositionDataFrame-method
<pre>mzAlign,MSImagingExperiment,missing-method</pre>	(PositionDataFrame-class), 29
(deprecated), 7	nrun,SpatialResults-method
<pre>mzAlign,MSImagingExperiment,numeric-method</pre>	(SpatialResults-class), 58
(deprecated), 7	nrun,SpectralImagingData-method
mzBin (deprecated), 7	(SpectralImagingData-class), 67
mzBin,MSImagingExperiment,missing-method	nscentroids, 62
(deprecated), 7	
mzBin,MSImagingExperiment,numeric-method	OPLS, <i>46</i>
(deprecated), 7	OPLS (SpatialPLS), 56
mzData (deprecated), 7	opls, 58
<pre>mzData&lt;- (deprecated), 7</pre>	OPLS, ANY-method (SpatialPLS), 56
mzFilter (deprecated), 7	OPLS,SpectralImagingExperiment-method
mzFilter,MSImagingExperiment-method	(SpatialPLS), 56
(deprecated), 7	
	parseAnalyze, 33, 34
names,SpatialResults-method	parseImzML, <i>33</i> , <i>34</i>
(SpatialResults-class), 58	PCA, 50, 54, 58
names,SpectraArrays-method	PCA (SpatialPCA), 54
(SpectraArrays-class), 65	PCA, ANY-method (SpatialPCA), 54
names,SpectralImagingArrays-method	PCA, SpectralImagingExperiment-method
(SpectralImagingArrays-class).	(SpatialPCA), 54

pData,SpatialResults-method	<pre>pixelData&lt;-,SpectralImagingData-method</pre>
(SpatialResults-class), 58	(SpectralImagingData-class), 67
pData,SpectralImagingData-method	<pre>pixelNames (SpectralImagingData-class),</pre>
(SpectralImagingData-class), 67	67
pData<-,SpatialResults,ANY-method	pixelNames,SpatialResults-method
(SpatialResults-class), 58	(SpatialResults-class), 58
pData<-,SpectralImagingData,ANY-method	pixelNames,SpectralImagingData-method
(SpectralImagingData-class), 67	(SpectralImagingData-class), 67
peakAlign, 8, 18, 21–23, 33	pixelNames<-
peakAlign,MSImagingArrays-method	(SpectralImagingData-class), 67
(peakAlign), 18	pixelNames<-,SpatialResults-method
peakAlign,MSImagingExperiment-method	(SpatialResults-class), 58
(peakAlign), 18	<pre>pixelNames&lt;-,SpectralImagingData-method</pre>
peakAlign,SpectralImagingArrays-method	(SpectralImagingData-class), 67
(peakAlign), 18	pixels, 24
<pre>peakAlign,SpectralImagingExperiment-method</pre>	pixels, SpectralImagingArrays-method
(peakAlign), 18	(pixels), 24
peakBin (deprecated), 7	pixels, SpectralImagingData-method
peakData (deprecated), 7	(pixels), 24
peakData<- (deprecated), 7	pixels, SpectralImagingExperiment-method
peakFilter (deprecated), 7	(pixels), 24
peakFilter,MSImagingExperiment-method	pixelSize (deprecated), 7
(deprecated), 7	plot, 28
peakPick, 17, 19, 20, 22, 23, 31, 35, 36, 44	plot (plot-spectra), 27
peakPick,MSImagingArrays-method	plot, MeansTest, missing-method
(peakPick), 20	(MeansTest), 11
peakPick,MSImagingExperiment-method	plot,MSImagingArrays,formula-method
(peakPick), 20	(plot-spectra), 27
peakPick, SpectralImagingData-method	plot, MSImagingArrays, missing-method
(peakPick), 20	(plot-spectra), 27
peakPicking (deprecated), 7	plot, MSImagingArrays, numeric-method
peakPicking<- (deprecated), 7	(plot-spectra), 27
peakProcess, 8, 19, 21, 22	plot, MSImaging Experiment, character-method
peakProcess,MSImagingExperiment_OR_Arrays-me	
(peakProcess), 22	plot, MSImaging Experiment, formula-method
pixelApply (deprecated), 7	(plot-spectra), 27
<pre>pixelApply,SpectralImagingExperiment-method</pre>	
(deprecated), 7	(plot-spectra), 27
pixelData (SpectralImagingData-class),	plot, MSImagingExperiment, numeric-method
67	(plot-spectra), 27
pixelData, SpatialResults-method	plot,ResultsList,ANY-method
(SpatialResults-class), 58	(ResultsList-class), 37
pixelData, SpectralImagingData-method	plot, ResultsList, missing-method
(SpectralImagingData-class), 67	(ResultsList-class), 37
pixelData<-	plot,SpatialDGMM,missing-method
(SpectralImagingData-class), 67	(SpatialDGMM), 46
pixelData<-,SpatialResults-method	plot,SpatialFastmap,missing-method
(SpatialResults-class), 58	(SpatialFastmap), 49
( - i	\ -

olot,SpatialKMeans,missing-method	<pre>predict,SpatialNMF-method(SpatialNMF),</pre>
(SpatialKMeans), 51	53
olot,SpatialNMF,missing-method	predict,SpatialOPLS-method
(SpatialNMF), 53	(SpatialPLS), 56
olot,SpatialOPLS,missing-method	<pre>predict,SpatialPCA-method(SpatialPCA),</pre>
(SpatialPLS), 56	54
olot,SpatialPCA,missing-method	<pre>predict,SpatialPLS-method(SpatialPLS),</pre>
(SpatialPCA), 54	56
olot,SpatialPLS,missing-method	<pre>predict,SpatialShrunkenCentroids-method</pre>
(SpatialPLS), 56	(SpatialShrunkenCentroids), 60
olot,SpatialResults,ANY-method	<pre>presetImageDef, 40</pre>
(SpatialResults-class), 58	<pre>presetImageDef (simulateSpectra), 39</pre>
olot,SpatialShrunkenCentroids,missing-method	process, 17, 19, 21, 23, 30, 35, 36, 44
(SpatialShrunkenCentroids), 60	process,MSImagingArrays-method
olot,SpectralImagingArrays,formula-method	(process), 30
(plot-spectra), 27	<pre>process,MSImagingExperiment-method</pre>
olot,SpectralImagingArrays,missing-method	(process), 30
(plot-spectra), 27	process, SpectralImagingArrays-method
olot,SpectralImagingArrays,numeric-method	(process), 30
(plot-spectra), 27	<pre>process,SpectralImagingExperiment-method</pre>
olot,SpectralImagingExperiment,character-met	nod (process), 30
(plot-spectra), 27	processingData,SpectralImagingData-method
olot,SpectralImagingExperiment,formula-method	(SpectralImagingData-class), 67
(plot-spectra), 27	processingData<-,SpectralImagingData-method
olot,SpectralImagingExperiment,missing-method	(SpectralImagingData-class), 67
(plot-spectra), 27	ProcessingStep, <i>14</i> , <i>16</i> , <i>67–69</i>
olot,SpectralImagingExperiment,numeric-method	1
(plot-spectra), 27	rbind, MSimagingExperiment-method
olot-image, 25	(MSImagingExperiment-class), 15
olot-spectra, 27	rbind, SpectraArrays-method
olot_image, 26	(SpectraArrays-class), 65
olot_signal, 28	rbind, SpectralImagingArrays-method
PLS, 46	(SpectralImagingArrays-class),
PLS (SpatialPLS), 56	66
	rbind,SpectralImagingExperiment-method
ols, 57, 58	(SpectralImagingExperiment-class),
PLS, ANY-method (SpatialPLS), 56	69
PLS, SpectralImagingExperiment-method	rbind, XDataFrame-method
(SpatialPLS), 56	(XDataFrame-class), 76
PositionDataFrame, 11, 14–16, 40, 42, 59,	readAnalyze (readMSIData), 32
66–69, 77	readImzML, 40
PositionDataFrame	readImzML (readMSIData), 32
(PositionDataFrame-class), 29	readMSIData, 32, 76
PositionDataFrame-class, 29	recalibrate, 8, 17, 31, 34, 35, 44
orcomp_lanczos, 55	recalibrate, MSImagingExperiment_OR_Arrays-method
oredict,ResultsList-method	(recalibrate), 34
(ResultsList-class), 37	recalibrate, SpectralImagingData-method
oredict,SpatialFastmap-method	(recalibrate), 34
(SpatialFastmap), 49	reduceBaseline, 17, 31, 35, 36, 44

reduceBaseline,SpectralImagingData-method	runNames (PositionDataFrame-class), 29
(reduceBaseline), 35	runNames,PositionDataFrame-method
rescale, 17	(PositionDataFrame-class), 29
rescale_ref, 17	runNames,SpatialResults-method
rescale_rms, 17	(SpatialResults-class), 58
rescale_sum, 17	runNames,SpectralImagingData-method
reset (process), 30	(SpectralImagingData-class), 67
residuals, SpatialOPLS-method	runNames<- (PositionDataFrame-class), 29
(SpatialPLS), 56	runNames<-,PositionDataFrame-method
resolution (deprecated), 7	(PositionDataFrame-class), 29
resolution<- (deprecated), 7	runNames<-,SpatialResults-method
resultData (deprecated), 7	(SpatialResults-class), 58
resultData<- (deprecated), 7	runNames<-,SpectralImagingData-method
resultNames (deprecated), 7	(SpectralImagingData-class), 67
resultNames<- (deprecated), 7	
ResultSet (deprecated), 7	scanDirection (deprecated), 7
ResultSet-class (deprecated), 7	scanPattern (deprecated), 7
• • • • • • • • • • • • • • • • • • • •	scanPolarity (deprecated), 7
ResultsList, 60	scanType (deprecated), 7
ResultsList (ResultsList-class), 37	segmentationTest (MeansTest), 11
ResultsList-class, 37	selectROI, 26, 38
risk.colors (deprecated), 7	<pre>selectROI,SpectralImagingExperiment-method</pre>
rowMeans, SpectralImagingExperiment-method	(selectROI), 38
(summarizeFeatures), 73	setCardinalBPPARAM(Cardinal-package), 3
rownames, SpectralImagingExperiment-method	setCardinalNChunks (Cardinal-package), 3
(SpectralImagingExperiment-class),	setCardinalNumBlocks (deprecated), 7
69	setCardinalVerbose (Cardinal-package), 3
<pre>rownames&lt;-,SpectralImagingExperiment-method</pre>	sgmixn, 48
(SpectralImagingExperiment-class),	show, MSImagingArrays-method
69	(MSImagingArrays-class), 13
rowStats, 74	show, MSImagingExperiment-method
rowStats,SpectralImagingExperiment-method	(MSImagingExperiment-class), 15
(summarizeFeatures), 73	show, ResultsList-method
rowSums,SpectralImagingExperiment-method	(ResultsList-class), 37
(summarizeFeatures), 73	show, SpatialResults-method
run (PositionDataFrame-class), 29	(SpatialResults-class), 58
run,PositionDataFrame-method	show, SpectraArrays-method
(PositionDataFrame-class), 29	(SpectraArrays-class), 65
run, SpatialResults-method	show, SpectralImagingArrays-method
(SpatialResults-class), 58	(SpectralImagingArrays-class),
run, Spectral Imaging Data-method	66
(SpectralImagingData-class), 67	show, SpectralImagingData-method
run<- (PositionDataFrame-class), 29	(SpectralImagingData-class), 67
run<-,PositionDataFrame-method	show, SpectralImagingExperiment-method
(PositionDataFrame-class), 29	(SpectralImagingExperiment-class),
run<-,SpatialResults-method	69
(SpatialResults-class), 58	show,XDataFrame-method
run<-,SpectralImagingData-method	(XDataFrame-class), 76
(SpectralImagingData-class), 67	SImageData (deprecated), 7
(50000) 4111114011160404 61455/, 67	

SImageData-class (deprecated), 7	SpatialNMF-class(SpatialNMF), 53
SImageSet (deprecated), 7	SpatialOPLS (SpatialPLS), 56
SImageSet-class (deprecated), 7	SpatialOPLS-class (SpatialPLS), 56
SimpleList, 37, 65	SpatialPCA, 54
simspec, 42	SpatialPCA-class (SpatialPCA), 54
<pre>simulateImage (simulateSpectra), 39</pre>	SpatialPLS, 56
simulateSpectra, 39, 40	SpatialPLS-class (SpatialPLS), 56
simulateSpectrum (deprecated), 7	SpatialResults, 37
slice, 42	SpatialResults (SpatialResults-class),
smooth, 17, 31, 35, 36, 43	58
smooth, SpectralImagingData-method	SpatialResults-class, 58
(smooth), 43	SpatialShrunkenCentroids, 60
smoothing (deprecated), 7	spatialShrunkenCentroids, 46, 52, 58
<pre>smoothing&lt;- (deprecated), 7</pre>	spatialShrunkenCentroids
<pre>smoothSignal (deprecated), 7</pre>	(SpatialShrunkenCentroids), 60
<pre>smoothSignal,SpectralImagingExperiment-metho</pre>	dspatialShrunkenCentroids, ANY, ANY-method
(deprecated), 7	(SpatialShrunkenCentroids), 60
sparse_mat, 10, 64	spatialShrunkenCentroids, ANY, missing-method
spatialApply (deprecated), 7	(SpatialShrunkenCentroids), 60
<pre>spatialApply,SpectralImagingExperiment-metho</pre>	dspatialShrunkenCentroids,SpectralImagingExperiment,ANY-met
(deprecated), 7	(SpatialShrunkenCentroids), 60
SpatialCV, 45	spatialShrunkenCentroids,SpectralImagingExperiment,missing
SpatialCV-class (SpatialCV), 45	(SpatialShrunkenCentroids), 60
SpatialDGMM, 46	SpatialShrunkenCentroids-class
spatialDGMM, 13	(SpatialShrunkenCentroids), 60
spatialDGMM (SpatialDGMM), 46	spatialWeights, 10,63
spatialDGMM, ANY-method (SpatialDGMM), 46	spatialWeights, ANY-method
<pre>spatialDGMM,SpectralImagingExperiment-method</pre>	(spatialWeights), 63
(SpatialDGMM), 46	spatialWeights,PositionDataFrame-method
SpatialDGMM-class (SpatialDGMM), 46	(spatialWeights), 63
SpatialFastmap, 49	spatialWeights,SpectralImagingExperiment-method
spatialFastmap, <i>51</i> , <i>54</i> , <i>55</i>	(spatialWeights), 63
spatialFastmap (SpatialFastmap), 49	spectra,SpectralImagingData-method
spatialFastmap, ANY-method	(SpectralImagingData-class), 67
(SpatialFastmap), 49	<pre>spectra&lt;-,SpectralImagingData-method</pre>
<pre>spatialFastmap,SpectralImagingExperiment-met</pre>	hod (SpectralImagingData-class), 67
(SpatialFastmap), 49	SpectraArrays, <i>14–16</i> , <i>66–69</i>
SpatialFastmap-class(SpatialFastmap),	SpectraArrays (SpectraArrays-class), 65
49	SpectraArrays-class, 65
SpatialKMeans, 51	spectraData,SpectralImagingData-method
spatialKMeans, <i>50</i> , <i>52</i> , <i>63</i>	(SpectralImagingData-class), 67
spatialKMeans (SpatialKMeans), 51	<pre>spectraData&lt;-,SpectralImagingData-method</pre>
spatialKMeans, ANY-method	(SpectralImagingData-class), 67
(SpatialKMeans), 51	SpectralImagingArrays, 14, 68, 69
spatialKMeans,SpectralImagingExperiment-methospectralImagingArrays	
(SpatialKMeans), 51	(SpectralImagingArrays-class),
SpatialKMeans-class (SpatialKMeans), 51	66
SpatialNMF, 53	SpectralImagingArrays-class, 66

SpectralImagingData, <i>14</i> , <i>16</i> , <i>65–67</i> , <i>70</i>	topFeatures,SpatialShrunkenCentroids-method
SpectralImagingData	(SpatialShrunkenCentroids), 60
(SpectralImagingData-class), 67	
SpectralImagingData-class, 67	vizi_engine (Cardinal-package), 3
SpectralImagingExperiment, 16, 58, 68, 69	vizi_par(Cardinal-package), 3
SpectralImagingExperiment	vizi_style (Cardinal-package), 3
(SpectralImagingExperiment-class),	vm_used,SpectraArrays-method
69	(SpectraArrays-class),65
SpectralImagingExperiment-class, 69	
spectraNames, SpectralImagingData-method	warp1, <i>34</i>
(SpectralImagingData-class), 67	warp1_cow, <i>35</i>
spectraNames<-,SpectralImagingData-method	warp1_dtw, <i>35</i>
(SpectralImagingData-class), 67	warp1_loc, <i>35</i>
spectrapply, 70	writeAnalyze, 75
spectrapply, MSImagingArrays-method	writeAnalyze (writeMSIData), 75
(spectrapply), 70	writeAnalyze,MSImagingExperiment-method
spectrapply, MSImagingExperiment-method	(writeMSIData), 75
(spectrapply), 70	writeAnalyze,SpectralImagingExperiment-method
	(writeMSIData), 75
spectrapply, SpectralImagingArrays-method	writeImzML, 75
(spectrapply), 70	writeImzML (writeMSIData), 75
spectrapply, SpectralImagingExperiment-method	writeImzML,MSImagingExperiment_OR_Arrays-method
(spectrapply), 70	(writeMSIData), 75
spectraVariables, SpectralImagingData-method	writeMSIData, 34, 75
(SpectralImagingData-class), 67	
spectrumRepresentation (deprecated), 7	XDataFrame, <i>11</i> , <i>30</i>
spectrumRepresentation<- (deprecated), 7	XDataFrame (XDataFrame-class), 76
subset, Spectral Imaging Arrays - method	XDataFrame-class, 76
(subsetFeatures), 72	XDFrame (XDataFrame-class), 76
subset, SpectralImagingExperiment-method	XDFrame-class (XDataFrame-class), 76
(subsetFeatures), 72	, , , , , , , , , , , , , , , , , , , ,
subsetFeatures, 72	
subsetPixels (subsetFeatures), 72	
summarizeFeatures, 8, 72, 73	
summarizePixels, 72	
summarizePixels (summarizeFeatures), 73	
svd, <i>55</i>	
. 5	
topFeatures (SpatialShrunkenCentroids),	
60	
topFeatures,MeansTest-method	
(MeansTest), 11	
topFeatures,ResultsList-method	
(ResultsList-class), 37	
topFeatures,SpatialKMeans-method	
(SpatialKMeans), 51	
topFeatures,SpatialOPLS-method	
(SpatialPLS), 56	
topFeatures,SpatialPLS-method	
(SpatialPLS), 56	