

Package ‘imcRtools’

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Title Methods for imaging mass cytometry data analysis

Description This R package supports the handling and analysis of imaging mass cytometry and other highly multiplexed imaging data. The main functionality includes reading in single-cell data after image segmentation and measurement, data formatting to perform channel spillover correction and a number of spatial analysis approaches. First, cell-cell interactions are detected via spatial graph construction; these graphs can be visualized with cells representing nodes and interactions representing edges. Furthermore, per cell, its direct neighbours are summarized to allow spatial clustering. Per image/grouping level, interactions between types of cells are counted, averaged and compared against random permutations. In that way, types of cells that interact more (attraction) or less (avoidance) frequently than expected by chance are detected.

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aggregateNeighbors *Function to aggregate all neighbors of each cell.*

Description

Function to summarize categorical or expression values of all neighbors of each cell.

Usage

```
aggregateNeighbors(
  object,
  colPairName,
  aggregate_by = c("metadata", "expression"),
  count_by = NULL,
  proportions = TRUE,
  assay_type = NULL,
```

```

subset_row = NULL,
statistic = c("mean", "median", "sd", "var"),
name = NULL
)

```

Arguments

object	a SingleCellExperiment or SpatialExperiment object
colPairName	single character indicating the colPair(object) entry containing the neighbor information.
aggregate_by	character specifying whether the neighborhood should be summarized by cellular features stored in colData(object) (aggregate_by = "metadata") or by marker expression of the neighboring cells (aggregate_by = "expression").
count_by	for summarize_by = "metadata", a single character specifying the colData(object) entry containing the cellular metadata that should be summarized across each cell's neighborhood.
proportions	single logical indicating whether aggregated metadata should be returned in form of proportions instead of absolute counts.
assay_type	for summarize_by = "expression", single character indicating the assay slot to use.
subset_row	for summarize_by = "expression", an integer, logical or character vector specifying the features to use. If NULL, defaults to all features.
statistic	for summarize_by = "expression", a single character specifying the statistic to be used for summarizing the expression values across all neighboring cells. Supported entries are "mean", "median", "sd", "var". Defaults to "mean" if not specified.
name	single character specifying the name of the data frame to be saved in the colData(object). Defaults to "aggregatedNeighbors" when summarize_by = "metadata" or "statistic_aggregatedExpression" when summarize_by = "expression".

Value

returns an object of class(object) containing the aggregated values in form of a DataFrame object in colData(object)[[name]].

Author(s)

Daniel Schulz (<daniel.schulz@uzh.ch>)

Examples

```

library(cytomapper)
data(pancreasSCE)

sce <- buildSpatialGraph(pancreasSCE, img_id = "ImageNb",
                        type = "knn", k = 3)

```

```
# Aggregating neighboring cell-types
sce <- aggregateNeighbors(sce, colPairName = "knn_interaction_graph",
  aggregate_by = "metadata",
  count_by = "CellType")
sce$aggregatedNeighbors

# Aggregating neighboring expression values
sce <- aggregateNeighbors(sce, colPairName = "knn_interaction_graph",
  aggregate_by = "expression",
  assay_type = "exprs",
  statistic = "mean")
sce$mean_aggregatedExpression
```

binAcrossPixels	<i>Aggregate consecutive pixels per single-metal spot</i>
-----------------	---

Description

Helper function for estimating the spillover matrix. Per metal spot, consecutive pixels are aggregated (default: summed).

Usage

```
binAcrossPixels(
  object,
  bin_size,
  spot_id = "sample_id",
  assay_type = "counts",
  statistic = "sum",
  ...
)
```

Arguments

object	a SingleCellExperiment object containing pixel intensities for all channels. Individual pixels are stored as columns and channels are stored as rows.
bin_size	single numeric indicating how many consecutive pixels per spot should be aggregated.
spot_id	character string indicating which colData(object) entry stores the isotope names of the spotted metal.
assay_type	character string indicating which assay to use.
statistic	character string indicating the statistic to use for aggregating consecutive pixels.
...	additional arguments passed to aggregateAcrossCells

Value

returns the binned pixel intensities in form of a SingleCellExperiment object

Author(s)

Nils Eling (<nils.eling@dqbm.uzh.ch>)

See Also

[aggregateAcrossCells](#) for the aggregation function

Examples

```
path <- system.file("extdata/spillover", package = "imcRtools")
# Read in .txt files
sce <- readSCEfromTXT(path)
dim(sce)

# Visualizes heatmap before aggregation
plotSpotHeatmap(sce)

# Sum consecutive pixels
sce <- binAcrossPixels(sce, bin_size = 10)
dim(sce)

# Visualizes heatmap after aggregation
plotSpotHeatmap(sce)
```

buildSpatialGraph *Builds an interaction graph based on the cells' locations*

Description

Function to define cell-cell interactions via distance-based expansion, delaunay triangulation or k nearest neighbor detection.

Usage

```
buildSpatialGraph(
  object,
  img_id,
  type = c("expansion", "knn", "delaunay"),
  k = NULL,
  directed = TRUE,
  max_dist = NULL,
  threshold = NULL,
  coords = c("Pos_X", "Pos_Y"),
  name = NULL,
  BNPARAM = KmknParam(),
  BPPARAM = SerialParam(),
  ...
)
```

Arguments

object	a SingleCellExperiment or SpatialExperiment object
img_id	single character indicating the colData(object) entry containing the unique image identifiers.
type	single character specifying the type of graph to be build. Supported entries are "expansion" (default) to find interacting cells via distance thresholding; "delaunay" to find interactions via delaunay triangulation; "knn" to find the k nearest neighboring cells.
k	(when type = "knn") single numeric integer defining the number of nearest neighbors to search for.
directed	(when type = "knn") should the returned graph be directed? (see details).
max_dist	(when type = "knn" or type = "delaunay") the maximum distance at which to consider neighboring cells. All neighbors within a distance larger than max_dist will be excluded from graph construction.
threshold	(when type = "expansion") single numeric specifying the maximum distance for considering neighbors
coords	character vector of length 2 specifying the names of the colData (for a SingleCellExperiment object) or the spatialCoords entries of the cells' x and y locations.
name	single character specifying the name of the graph.
BNPARAM	a BiocNeighborParam object defining the algorithm to use.
BPPARAM	a BiocParallelParam-class object defining how to parallelize computations.
...	additional parameters passed to the findNeighbors function (type = "expansion"), the triangulate function (type = "delaunay") or the findKNN function (type = "knn").

Value

returns a SpatialExperiment or SingleCellExperiment containing the graph in form of a SelfHits object in colPair(object, name).

Building an interaction graph

This function defines interacting cells in different ways. They are based on the cells' centroids and do not incorporate cell shape or area.

1. When type = "expansion", all cells within the radius threshold are considered interacting cells.
2. When type = "delaunay", interacting cells are found via a delaunay triangulation of the cells' centroids.
3. When type = "knn", interacting cells are defined as the k nearest neighbors in the 2D spatial plane.

The directed parameter only affects graph construction via k nearest neighbor search. For directed = FALSE, each interaction will be stored as mutual edge (e.g. node 2 is connected to node 10 and vice versa). For type = "expansion" and type = "delaunay", each edge is stored as mutual edge by default.

The graph is stored in form of a SelfHits object in `colPair(object, name)`. This object can be regarded as an edgelist and coerced to an igraph object via `graph_from_edgelist(as.matrix(colPair(object, name)))`.

Choosing the graph construction method

When finding interactions via expansion or knn, the `findNeighbors` or `findKNN` functions are used, respectively. Both functions accept the `BNPARAM` parameter via which the graph construction method can be defined (default `KmknnParam`). For an overview on the different algorithms, see `BiocNeighborParam`. Within the `BiocNeighborParam` object, distance can be set to "Euclidean" (default), "Manhattan" or "Cosine".

Author(s)

Nils Eling (<nils.eling@dqbm.uzh.ch>)

See Also

`findNeighbors` for the function finding interactions via expansion

`findKNN` for the function finding interactions via nearest neighbor search

`triangulate` for the function finding interactions via delaunay triangulation

`plotSpatial` for visualizing spatial graphs

Examples

```
path <- system.file("extdata/mockData/steinbock", package = "imcRtools")
spe <- read_steinbock(path)

# Constructing a graph via expansion
spe <- buildSpatialGraph(spe, img_id = "sample_id",
                        type = "expansion", threshold = 10)
colPair(spe, "expansion_interaction_graph")

# Constructing a graph via delaunay triangulation
spe <- buildSpatialGraph(spe, img_id = "sample_id",
                        type = "delaunay")
colPair(spe, "delaunay_interaction_graph")

# Constructing a graph via k nearest neighbor search
spe <- buildSpatialGraph(spe, img_id = "sample_id",
                        type = "knn", k = 5)
colPair(spe, "knn_interaction_graph")
```

countInteractions *Summarizes cell-cell interactions within grouping levels (e.g. images)*

Description

Function to calculate the average number of neighbors B that a cell of type A has using different approaches.

Usage

```
countInteractions(
  object,
  group_by,
  label,
  colPairName,
  method = c("classic", "histocat", "patch"),
  patch_size = NULL
)
```

Arguments

object	a SingleCellExperiment or SpatialExperiment object .
group_by	a single character indicating the colData(object) entry by which interactions are grouped. This is usually the image ID or patient ID.
label	single character specifying the colData(object) entry which stores the cell labels. These can be cell-types labels or other metadata.
colPairName	single character indicating the colPair(object) entry containing cell-cell interactions in form of an edge list.
method	which cell-cell interaction counting method to use (see details)
patch_size	if method = "patch", a single numeric specifying the minimum number of neighbors of the same type to be considered a patch (see details)

Value

a DataFrame containing one row per group_by entry and unique label entry combination (from_label, to_label). The ct entry stores the interaction count as described in the details. NA is returned if a certain label is not present in this grouping level.

Counting and summarizing cell-cell interactions

In principle, the countInteractions function counts the number of edges (interactions) between each set of unique entries in colData(object)[[label]]. Simplified, it counts for each cell of type A the number of neighbors of type B. This count is averaged within each unique entry colData(object)[[group_by]] in three different ways:

1. method = "classic": The count is divided by the total number of cells of type A. The final count can be interpreted as "How many neighbors of type B does a cell of type A have on average?"

2. `method = "histocat"`: The count is divided by the number of cells of type A that have at least one neighbor of type B. The final count can be interpreted as "How many many neighbors of type B has a cell of type A on average, given it has at least one neighbor of type B?"

3. `method = "patch"`: For each cell, the count is binarized to 0 (less than `patch_size` neighbors of type B) or 1 (more or equal to `patch_size` neighbors of type B). The binarized counts are averaged across all cells of type A. The final count can be interpreted as "What fraction of cells of type A have at least a given number of neighbors of type B?"

Author(s)

Vito Zanotelli

Jana Fischer

adapted by Nils Eling (<nils.eling@dqbm.uzh.ch>)

References

Schulz, D. et al., Simultaneous Multiplexed Imaging of mRNA and Proteins with Subcellular Resolution in Breast Cancer Tissue Samples by Mass Cytometry., *Cell Systems* 2018 6(1):25-36.e5

Shapiro, D. et al., histoCAT: analysis of cell phenotypes and interactions in multiplex image cytometry data, *Nature Methods* 2017 14, p. 873–876

See Also

[testInteractions](#) for testing cell-cell interactions per grouping level.

Examples

```
library(cytomapper)
data(pancreasSCE)

pancreasSCE <- buildSpatialGraph(pancreasSCE, img_id = "ImageNb",
                                type = "knn", k = 3)

# Classic style calculation
(out <- countInteractions(pancreasSCE,
                          group_by = "ImageNb",
                          label = "CellType",
                          method = "classic",
                          colPairName = "knn_interaction_graph"))

# Histocat style calculation
(out <- countInteractions(pancreasSCE,
                          group_by = "ImageNb",
                          label = "CellType",
                          method = "histocat",
                          colPairName = "knn_interaction_graph"))

# Patch style calculation
(out <- countInteractions(pancreasSCE,
                          group_by = "ImageNb",
```

```
label = "CellType",
method = "patch",
patch_size = 3,
colPairName = "knn_interaction_graph"))
```

filterPixels

Filter pixels based on their assigned masses

Description

Helper function for estimating the spillover matrix. After assigning each pixel to a spotted mass, this function will filter incorrectly assigned pixels and remove small pixel sets.

Usage

```
filterPixels(
  object,
  bc_id = "bc_id",
  spot_mass = "sample_mass",
  minevents = 40,
  correct_pixels = TRUE
)
```

Arguments

object	a SingleCellExperiment object containing pixel intensities per channel. Individual pixels are stored as columns and channels are stored as rows.
bc_id	character string indicating which colData(object) entry stores the estimated mass
spot_mass	character string indicating which colData(object) entry stores the true isotope mass of the spotted metal.
minevents	single numeric indicating the threshold under which pixel sets are excluded from spillover estimation.
correct_pixels	logical indicating if incorrectly assigned pixels should be excluded from spillover estimation.

Value

returns a SingleCellExperiment object in which colData(object)\$bc_id has been adjusted based on the filter criteria.

Author(s)

Vito Zanotelli, adapted by Nils Eling (<nils.eling@dqbm.uzh.ch>)

Examples

```
path <- system.file("extdata/spillover", package = "imcRtools")
sce <- readSCEfromTXT(path)
assay(sce, "exprs") <- asinh(counts(sce)/5)

# Pre-process via CATALYST
library(CATALYST)

bc_key <- as.numeric(unique(sce$sample_mass))
sce <- assignPrelim(sce, bc_key = bc_key)
sce <- estCutoffs(sce)
sce <- applyCutoffs(sce)

sce <- filterPixels(sce)

table(sce$sample_mass, sce$bc_id)
```

findBorderCells	<i>Find cells at the image border</i>
-----------------	---------------------------------------

Description

Detection of cells close to the image border for subsequent exclusion from downstream analyses.

Usage

```
findBorderCells(object, img_id, border_dist, coords = c("Pos_X", "Pos_Y"))
```

Arguments

object	a SingleCellExperiment or SpatialExperiment object.
img_id	single character indicating the colData(object) entry containing the unique image identifiers.
border_dist	single numeric defining the distance to the image border. The image border here is defined as the minimum and maximum among the cells' x and y location.
coords	character vector of length 2 specifying the names of the colData (for a SingleCellExperiment object) or the spatialCoords entries indicating the cells' x and y locations.

Value

an object of class(object) containing the logical border_cells entry in the colData slot.

Author(s)

Nils Eling (<nils.eling@dqbm.uzh.ch>)

Examples

```
library(cytomapper)
data("pancreasSCE")

sce <- findBorderCells(pancreasSCE, img_id = "ImageNb",
                      border_dist = 10)

plotSpatial(sce, img_id = "ImageNb", node_color_by = "border_cells")
```

patchDetection	<i>Function to detect patches containing defined cell types</i>
----------------	---

Description

Function to detect spatial clusters of defined types of cells. By defining a certain distance threshold, all cells within the vicinity of these clusters are detected as well.

Usage

```
patchDetection(
  object,
  patch_cells,
  colPairName,
  min_patch_size = 1,
  name = "patch_id",
  expand_by = 0,
  coords = c("Pos_X", "Pos_Y"),
  convex = FALSE,
  img_id = NULL,
  BPPARAM = SerialParam()
)
```

Arguments

<code>object</code>	a <code>SingleCellExperiment</code> or <code>SpatialExperiment</code> object
<code>patch_cells</code>	logical vector of length equal to the number of cells contained in <code>object</code> . TRUE entries define the cells to consider for patch detection (see Details).
<code>colPairName</code>	single character indicating the <code>colPair(object)</code> entry containing the neighbor information.
<code>min_patch_size</code>	single integer indicating the minimum number of connected cells that make up a patch before expansion.
<code>name</code>	single character specifying the <code>colData</code> entry storing the patch IDs in the returned object.
<code>expand_by</code>	single numeric indicating in which vicinity range cells should be considered as belonging to the patch (see Details).

coords	character vector of length 2 specifying the names of the colData (for a SingleCellExperiment object) or the spatialCoords entries of the cells' x and y locations.
convex	should the convex hull be computed before expansion? Default: the concave hull is computed.
img_id	single character indicating the colData(object) entry containing the unique image identifiers.
BPPARAM	a BiocParallelParam-class object defining how to parallelize computations.

Value

An object of class(object) containing a patch ID for each cell in colData(object)[[name]].

Detecting patches of defined cell types

This function works as follows:

1. Only cells defined by patch_cells are considered for patch detection.
2. Patches of connected cells are detected. Here, cell-to-cell connections are defined by the interaction graph stored in colPair(object, colPairName). At this point, patches that contain fewer than min_patch_size cells are removed.
3. If expand_by > 0, a concave (default) or convex hull is constructed around each patch. This is then expanded by expand_by and cells within the expanded hull are detected and assigned to the patch. This expansion only works if a patch contains at least 3 cells.

The returned object contains an additional entry colData(object)[[name]], which stores the patch ID per cell. NA indicate cells that are not part of a patch.

Author(s)

Tobias Hoch

adapted by Nils Eling (<nils.eling@dqbm.uzh.ch>)

References

Hoch, T. et al., [Multiplexed Imaging Mass Cytometry of Chemokine Milieus in Metastatic Melanoma Characterizes Features of Response to Immunotherapy.](#), bioRxiv 2021

Examples

```
library(cytomapper)
data(pancreasSCE)

# Visualize cell types
plotSpatial(pancreasSCE, img_id = "ImageNb", node_color_by = "CellType")

# Build interaction graph
pancreasSCE <- buildSpatialGraph(pancreasSCE, img_id = "ImageNb",
                                type = "expansion", threshold = 20)

# Detect patches of "celltype_B" cells
```

```

pancreasSCE <- patchDetection(pancreasSCE,
                             patch_cells = pancreasSCE$CellType == "celltype_B",
                             colPairName = "expansion_interaction_graph")

plotSpatial(pancreasSCE, img_id = "ImageNb", node_color_by = "patch_id")

# Include cells in vicinity
pancreasSCE <- patchDetection(pancreasSCE,
                             patch_cells = pancreasSCE$CellType == "celltype_B",
                             colPairName = "expansion_interaction_graph",
                             expand_by = 20,
                             img_id = "ImageNb")

plotSpatial(pancreasSCE, img_id = "ImageNb", node_color_by = "patch_id")

```

plotSpatial

Visualizes the spatial locations and interactions of cells

Description

A general function to plot spatial locations of cells while specifying color, shape, size. Cell-cell interactions can be visualized in form of edges between points.

Usage

```

plotSpatial(
  object,
  img_id,
  coords = c("Pos_X", "Pos_Y"),
  node_color_by = NULL,
  node_shape_by = NULL,
  node_size_by = NULL,
  node_color_fix = NULL,
  node_shape_fix = NULL,
  node_size_fix = NULL,
  assay_type = NULL,
  draw_edges = FALSE,
  directed = TRUE,
  edge_color_by = NULL,
  edge_width_by = NULL,
  edge_color_fix = NULL,
  edge_width_fix = NULL,
  arrow = NULL,
  end_cap = NULL,
  colPairName = NULL,
  nodes_first = TRUE,
  ncols = NULL,

```

```

    nrows = NULL,
    scales = "free",
    flip_x = FALSE,
    flip_y = TRUE
  )

```

Arguments

object	a SingleCellExperiment or SpatialExperiment object.
img_id	single character indicating the colData(object) entry containing the unique image identifiers.
coords	character vector of length 2 specifying the names of the colData (for a SingleCellExperiment object) or the spatialCoords entries indicating the the cells' x and y locations.
node_color_by	single character indicating the colData(object) entry or marker name by which the nodes (cell locations) should be colored.
node_shape_by	single character indicating the colData(object) entry by which the shape of the nodes are defined.
node_size_by	single character indicating the colData(object) entry by which the size of the nodes are defined.
node_color_fix	single character or numeric specifying the color of all nodes.
node_shape_fix	single numeric or character specifying the shape of all nodes.
node_size_fix	single numeric specifying the size of all nodes
assay_type	single character indicating the assay slot from which to extract the expression data when node_color_by is set to one of rownames(object).
draw_edges	should cell-cell interactions be drawn as edges between nodes?
directed	should cell-cell interactions be handled as a directed graph?
edge_color_by	single character indicating by which to color the edges. See details for more information.
edge_width_by	single character determining the size of the edges. See details for more information.
edge_color_fix	single character or numeric specifying the color of all edges.
edge_width_fix	single numeric specifying the size of all edges.
arrow	an arrow object specifying how to draw arrows between cells.
end_cap	a geometry object specifying how long the edges are. This only takes effect when drawing arrows. Default: end_cap = circle(0.1, 'cm')
colPairName	single character specifying the colPair(object) slot to retrieve the cell-cell pairings.
nodes_first	should the nodes be plotted first and then the edges?
ncols	number of columns of the grid to arrange individual images.
nrows	number of rows of the grid to arrange individual images.
scales	one of "free", "fixed", "free_x" or "free_y" indicating if x- and y-axis ranges should be fixed across all images.
flip_x	flip the x-axis?
flip_y	flip the y-axis?

Value

returns a ggplot object.

Visualizing cell locations and cell-cell interactions

By default, the cells' locations are visualized in form of points (here also referred to as "nodes") on a 2-dimensional plane. The cells' coordinates are extracted either from `colData(object)` slot (for a `SingleCellExperiment` input object) or from the `spatialCoords(object)` slot (for a `SpatialExperiment` input object). Node aesthetics are controlled by setting `node_color_by`, `node_shape_by` and `node_size_by` for associating the aesthetics with variables. If node aesthetics should be the same for all nodes, `node_color_fix`, `node_shape_fix` and `node_size_fix` can be set.

When `draw_edges = TRUE`, cell-cell interactions are visualized in form of edges between nodes. For this, object needs to contain column pairings in `colPair(object, colPairName)`. Edge color and size can be set by specifying either an entry in `mcols(colPair(object, colPairName))` (edge attributes) or in `colData(object)`. In the latter case, edges are colored by attributes associated to the "from" node. Variable aesthetics can be set using `edge_color_by` and `edge_width_by`. If all edges should have the same width or color, `edge_color_fix` and `edge_width_fix` can be set.

Arrows for displaying directed graphs can be drawn by supplying a `arrow` object. Arrow attributes can be set within this class. To cap the edge before it reaches the next node, the `end_cap` parameter can be used.

Author(s)

Nils Eling (<nils.eling@dqbm.uzh.ch>)

See Also

[buildSpatialGraph](#) for constructing interaction graphs

[ggraph](#) for handling graph aesthetics

Examples

```
library(cytomapper)
data(pancreasSCE)

sce <- buildSpatialGraph(pancreasSCE, img_id = "ImageNb",
                        type = "knn", k = 3, directed = FALSE)

# Only nodes
plotSpatial(sce, img_id = "ImageNb",
            node_color_by = "CellType",
            node_shape_by = "ImageNb",
            node_size_by = "Area")

# With edges and nodes colored by expression
plotSpatial(sce, img_id = "ImageNb",
            node_color_by = "PIN",
            assay_type = "exprs",
```



```

        node_shape_by = "ImageNb",
        node_size_by = "Area",
        draw_edges = TRUE,
        colPairName = "knn_interaction_graph",
        edge_color_by = "Pattern")

# With arrows
plotSpatial(sce, img_id = "ImageNb",
            node_color_by = "CellType",
            node_shape_by = "ImageNb",
            node_size_by = "Area",
            draw_edges = TRUE,
            colPairName = "knn_interaction_graph",
            edge_color_fix = "green",
            arrow = grid::arrow(length = grid::unit(0.1, "inch")),
            end_cap = ggraph::circle(0.2, "cm"))

```

plotSpotHeatmap	<i>Summarizes and visualizes the pixel intensities per spot and channel</i>
-----------------	---

Description

Helper function for estimating the spillover matrix. This function visualizes the median pixel intensities per spot (rows) and per channel (columns) in form of a heatmap.

Usage

```

plotSpotHeatmap(
  object,
  spot_id = "sample_id",
  channel_id = "channel_name",
  assay_type = "counts",
  statistic = "median",
  log = TRUE,
  threshold = NULL,
  order_metals = TRUE,
  color = viridis(100),
  breaks = NA,
  legend_breaks = NA,
  cluster_cols = FALSE,
  cluster_rows = FALSE,
  ...
)

```

Arguments

object	a SingleCellExperiment object containing pixel intensities per channel. Individual pixels are stored as columns and channels are stored as rows.
--------	--

spot_id	character string indicating which colData(object) entry stores the isotope names of the spotted metal. Entries should be of the form (mt)(mass) (e.g. Sm152 for Samarium isotope with the atomic mass 152).
channel_id	character string indicating which rowData(object) entry contains the isotope names of the acquired channels.
assay_type	character string indicating which assay to use (default counts).
statistic	the statistic to use when aggregating channels per spot (default median)
log	should the aggregated pixel intensities be $\log_{10}(x + 1)$ transformed?
threshold	single numeric indicating a threshold after pixel aggregation. All aggregated values larger than threshold will be labeled as 1.
order_metals	should the metals be ordered based on spotted mass?
color	see parameter in pheatmap
breaks	see parameter in pheatmap
legend_breaks	see parameter in pheatmap
cluster_cols	see parameter in pheatmap
cluster_rows	see parameter in pheatmap
...	other arguments passed to pheatmap.

Value

a [pheatmap](#) object

Quality control for spillover estimation

Visualizing the aggregated pixel intensities serves two purposes:

1. Small median pixel intensities (< 200 counts) might hinder the robust estimation of the channel spillover. In that case, consecutive pixels can be summed (see [binAcrossPixels](#)).
2. Each spotted metal (row) should show the highest median pixel intensity in its corresponding channel (column). If this is not the case, either the naming of the .txt files was incorrect or the incorrect metal was spotted.

By setting the threshold parameter, the user can easily identify spots where pixel intensities are too low for robust spillover estimation.

Author(s)

Nils Eling (<nils.eling@dqbm.uzh.ch>)

See Also

[pheatmap](#) for visual modifications

[aggregateAcrossCells](#) for the aggregation function

Examples

```

path <- system.file("extdata/spillover", package = "imcRtools")
# Read in .txt files
sce <- readSCEfromTXT(path)

# Visualizes heatmap
plotSpotHeatmap(sce)

# Visualizes thresholding results
plotSpotHeatmap(sce, log = FALSE, threshold = 200)

```

readImagefromTXT	<i>Reads one or multiple .txt files into a CytoImageList object</i>
------------------	---

Description

Reader function to generate [Image](#) objects in form of a [CytoImageList](#) container from .txt files.

Usage

```

readImagefromTXT(
  path,
  pattern = ".txt$",
  channel_pattern = "[A-Za-z]{1,2}[0-9]{2,3}Di",
  index_names = c("X", "Y"),
  BPPARAM = SerialParam()
)

```

Arguments

path	Full path to where the individual .txt files are located. This is usually the path where the .mcd file is located.
pattern	pattern to select which files should be read in (default ".txt\$").
channel_pattern	regular expression to select the channel names from the files.
index_names	exact names of the columns storing the x and y coordinates of the image
BPPARAM	parameters for parallelized reading in of images. This is only recommended for very large images.

Value

returns a [CytoImageList](#) object containing one [Image](#) object per .txt file.

Imaging mass cytometry .txt files

As part of the raw data folder, the Hyperion imaging system writes out one .txt file per acquisition. These files store the ion counts per pixel and channel.

This function reads these .txt files into a single [CytoImageList](#) object for downstream analysis. The `pattern` argument allows selection of all .txt files or a specific subset of files. The `channelNames` of the [CytoImageList](#) object are determined by the `channel_pattern` argument.

Author(s)

Nils Eling (<nils.eling@dqbm.uzh.ch>)

See Also

[CytoImageList](#) for the container

[MulticoreParam](#) for parallelized processing

[Image](#) for the multi-channel image object

`vignette("cytomapper")` for visualization of multi-channel images

Examples

```
path <- system.file("extdata/mockData/raw", package = "imcRtools")

# Read in all images
x <- readImagefromTXT(path)
x

# Read in specific files
y <- readImagefromTXT(path, pattern = "ROI_002")
y

# Read in other channelNames
z <- readImagefromTXT(path, channel_pattern = "[A-Za-z]{2}[0-9]{3}")
z
```

readSCEfromTXT

Generates a SingleCellExperiment from .txt files

Description

Helper function to process raw .txt files acquired by the Hyperion imaging system into a [SingleCellExperiment](#) object. This function is mainly used to read-in data generated from a "spillover slide". Here, each .txt file contains the measurements of multiple pixels for a single stain across all open channels.

Usage

```
readSCEfromTXT(
  x,
  pattern = ".txt$",
  metadata_cols = c("Start_push", "End_push", "Pushes_duration", "X", "Y", "Z"),
  verbose = TRUE,
  read_metal_from_filename = TRUE
)
```

Arguments

x	input can be of different types: A path Full path to where the single stain .txt files are located. A list object A named list object where each entry is a data.frame or coercible to one. The names of each entry indicate the spotted metals (see details).
pattern	pattern to select which files should be read in (default ".txt\$"). Only used when x is a path.
metadata_cols	character vector indicating which column entries of the .txt files should be saved in the colData(sce) slot.
verbose	logical indicating if additional information regarding the spotted and acquired masses should be shown.
read_metal_from_filename	should the sample metal and mass be extracted from the file/object names?

Value

returns a SCE object where pixels are stored as columns and acquired channels are stored as rows.

Reading in .txt files for spillover correction

As described in the original publication, single metal spots are acquired using the Hyperion imaging system. Each acquisition corresponds to one spot. All acquisitions are stored in a single .mcd file and individual acquisitions are stored in single .txt files.

This function aggregates these measurements into a single SingleCellExperiment object. For this, two inputs are possible:

1. x is a path: By default all .txt files are read in from the specified path. Here, the path should indicate the location of the spillover slide measurement. The file names of the .txt file must contain the spotted metal isotope name in the format (mt)(mass) (e.g. Sm152 for Samarium isotope with the atomic mass 152). Internally, the last occurrence of such a pattern is read in as the metal isotope name and stored in the colData(sce)\$sample_id slot.
2. x is a named list: If there are issues with reading in the metal isotope names from the .txt file names, the user can provide a list for which each entry contains the contents of a single .txt file. The names of the list must indicate the spotted metal in the format (mt)(mass). These names will be stored in the colData(sce)\$sample_id slot.

When `read_metal_from_filename = FALSE`, the function will not attempt to read in the spotted metal isotopes from the file or list names. Therefore, only the `sample_id` will be set based on the file/list names.

Author(s)

Nils Eling (<nils.eling@dqbm.uzh.ch>)

References

Chevrier, S. et al. 2017. "Compensation of Signal Spillover in Suspension and Imaging Mass Cytometry." *Cell Systems* 6: 612–20.

Examples

```
# Read files from path
path <- system.file("extdata/spillover", package = "imcRtools")

sce <- readSCEfromTXT(path)
sce

# Read files as list
cur_file_names <- list.files(path, pattern = ".txt", full.names = TRUE)
cur_files <- lapply(cur_file_names, read.delim)
names(cur_files) <- sub(".txt", "", basename(cur_file_names))

sce <- readSCEfromTXT(cur_files)
sce
```

read_cpout

Reads in single-cell data generated by the ImcSegmentationPipeline

Description

Reader function to generate a [SpatialExperiment](#) or [SingleCellExperiment](#) object from single-cell data obtained by the [ImcSegmentationPipeline](#) pipeline.

Usage

```
read_cpout(
  path,
  object_file = "cell.csv",
  image_file = "Image.csv",
  panel_file = "panel.csv",
  graph_file = "Object relationships.csv",
  object_feature_file = "var_cell.csv",
  intensities = "Intensity_MeanIntensity_FullStack",
  extract_imgid_from = "ImageNumber",
```

```

extract_cellid_from = "ObjectNumber",
extract_coords_from = c("Location_Center_X", "Location_Center_Y"),
extract_cellmetadata_from = c("AreaShape_Area", "Neighbors_NumberOfNeighbors_8",
  "AreaShape_Eccentricity", "AreaShape_MajorAxisLength", "AreaShape_MinorAxisLength",
  "AreaShape_MeanRadius"),
extract_imagemetadadata_from = c("Metadata_acname", "Metadata_acid",
  "Metadata_description"),
extract_graphimageid_from = "First Image Number",
extract_graphcellids_from = c("First Object Number", "Second Object Number"),
extract_metal_from = "Metal Tag",
scale_intensities = TRUE,
extract_scalingfactor_from = "Scaling_FullStack",
return_as = c("spe", "sce")
)

```

Arguments

path	full path to the CellProfiler output folder.
object_file	single character indicating the file name storing the object/cell-specific intensities and metadata.
image_file	single character indicating the file name storing meta data per image (can be NULL).
panel_file	single character indicating the file name storing the panel information (can be NULL).
graph_file	single character indicating the file name storing the object/cell interaction information (can be NULL).
object_feature_file	single character indicating the file name storing object/cell feature information.
intensities	single character indicating which column entries of the object_file contain the intensity features of interest. See details.
extract_imgid_from	single character indicating which column entries of the object_file and image_file contain the image integer ID.
extract_cellid_from	single character indicating which column entry of the object_file contains the object/cell integer ID.
extract_coords_from	character vector indicating which column entries of the object_file contain the x and y location of the objects/cells.
extract_cellmetadata_from	character vector indicating which additional object/cell specific metadata to extract from the object_file.
extract_imagemetadadata_from	character vector indicating which additional image specific metadata to extract from the image_file. These will be stored in the colData(x) slot as object/cell-specific entries.

extract_graphimageid_from	single character indicating which column entries of the graph_file contain the image integer ID.
extract_graphcellids_from	character vector indicating which column entries of the graph_file contain the first and second object/cell integer IDs. These will be stored as the from and to entry of the SelfHits object in colPair(x, "neighborhood").
extract_metal_from	single character indicating which column entry of the panel_file contains the metal isotopes of the used antibodies. This entry is used to match the panel information to the acquired channel information.
scale_intensities	single logical. Should the measured intensity features be scaled by extract_scalingfactor_from.
extract_scalingfactor_from	single character indicating which column entries of the image_file contain the image specific scaling factor.
return_as	should the object be returned as <code>SpatialExperiment</code> (return_as = "spe") or <code>SingleCellExperiment</code> (return_as = "sce").

Value

returns a `SpatialExperiment` or `SingleCellExperiment` object with markers in rows and cells in columns.

The returned data container

In the case of both containers `x`, intensity features (as selected by the `intensities` parameter) are stored in the `counts(x)` slot. Cell metadata (e.g morphological features) are stored in the `colData(x)` slot. The interaction graphs are stored as `SelfHits` object in the `colPair(x, "neighborhood")` slot.

Intensity features are extracted via partial string matching. Internally, the `read_cpout` function checks if per channel a single intensity feature is read in (by checking the `_cXY` ending where `XY` is the channel number).

In the case of a returned `SpatialExperiment` object, the cell coordinates are stored in the `spatialCoords(x)` slot.

In the case of a returned `SingleCellExperiment` object, the cell coordinates are stored in the `colData(x)` slot named as `Pos_X` and `Pos_Y`.

Author(s)

Tobias Hoch

Nils Eling (<nils.eling@dqbm.uzh.ch>)

See Also

<https://github.com/BodenmillerGroup/ImcSegmentationPipeline> for the pipeline

`read_steinbock` for reading in single-cell data as produced by the steinbock pipeline

`colPair` for information on how to work with the cell-cell interaction graphs

Examples

```

path <- system.file("extdata/mockData/cpout", package = "imcRtools")

# Read in as SpatialExperiment object
x <- read_cpout(path, graph_file = "Object_relationships.csv")
x

# Read in as SingleCellExperiment object
x <- read_cpout(path, graph_file = "Object_relationships.csv",
                return_as = "sce")
x

```

read_steinbock	<i>Reads in single-cell data generated by the steinbock pipeline</i>
----------------	--

Description

Reader function to generate a [SpatialExperiment](#) or [SingleCellExperiment](#) object from single-cell data obtained by the [steinbock](#) pipeline.

Usage

```

read_steinbock(
  path,
  intensities_folder = "intensities",
  regionprops_folder = "regionprops",
  graphs_folder = "neighbors",
  pattern = NULL,
  extract_cellid_from = "Object",
  extract_coords_from = c("centroid-1", "centroid-0"),
  image_file = "images.csv",
  extract_imagemetadadata_from = c("width_px", "height_px"),
  panel_file = "panel.csv",
  extract_names_from = "name",
  return_as = c("spe", "sce"),
  BPPARAM = SerialParam()
)

```

Arguments

path	full path to the steinbock output folder
intensities_folder	name of the folder containing the intensity measurements per image
regionprops_folder	name of the folder containing the cell-specific morphology and spatial measurements per image. Can be set to NULL to exclude reading in morphology measures.

graphs_folder	name of the folder containing the spatial connectivity graphs per image. Can be set to NULL to exclude reading in graphs.
pattern	regular expression specifying a subset of files that should be read in.
extract_cellid_from	single character indicating which column entry in the intensity files contains the integer cell id.
extract_coords_from	character vector indicating which column entries in the regionprops files contain the x (first entry) and y (second entry) coordinates.
image_file	single character indicating the file name storing meta data per image (can be NULL).
extract_imagemetadata_from	character vector indicating which additional image specific metadata to extract from the image_file. These will be stored in the colData(x) slot as object/cell-specific entries.
panel_file	single character containing the name of the panel file. This can either be inside the steinbock path (recommended) or located somewhere else.
extract_names_from	single character indicating the column of the panel file containing the channel names.
return_as	should the object be returned as SpatialExperiment (return_as = "spe") or SingleCellExperiment (return_as = "sce").
BPPARAM	parameters for parallelised processing.

Value

returns a [SpatialExperiment](#) or [SingleCellExperiment](#) object markers in rows and cells in columns.

The returned data container

In the case of both containers x, intensity features are stored in the counts(x) slot. Morphological features are stored in the colData(x) slot. The graphs are stored as [SelfHits](#) object in the colPair(x, "neighborhood") slot.

In the case of a returned [SpatialExperiment](#) object, the cell coordinates are stored in the spatialCoords(x) slot.

In the case of a returned [SingleCellExperiment](#) object, the cell coordinates are stored in the colData(x) slot named as Pos_X and Pos_Y.

Author(s)

Nils Eling (<nils.eling@dqbm.uzh.ch>)

See Also

<https://github.com/BodenmillerGroup/steinbock> for the pipeline
`read_cpout` for reading in single-cell data as produced by the `ImcSegmentationPipeline`
`SingleCellExperiment` and `SpatialExperiment` for the constructor functions.
`colPair` for information on how to work with the cell-cell interaction graphs
`bpparam` for the parallelised backend

Examples

```
path <- system.file("extdata/mockData/steinbock", package = "imcRtools")

# Read in as SpatialExperiment object
x <- read_steinbock(path)
x

# Read in as SingleCellExperiment object
x <- read_steinbock(path, return_as = "sce")
x

# Read in a subset of files
x <- read_steinbock(path, pattern = "mockData1")
x

# Only read in intensities
x <- read_steinbock(path, graphs_folder = NULL, regionprops_folder = NULL)
x

# Parallelisation
x <- read_steinbock(path, BPPARAM = BiocParallel::bpparam())
```

`show_cpout_features` *Display all features measured by CellProfiler.*

Description

Searchable datatable object of cell and image features as extracted by CellProfiler.

Usage

```
show_cpout_features(  
  path,  
  display = c("cell_features", "image_features"),  
  cell_features = "var_cell.csv",  
  image_features = "var_Image.csv"  
)
```

Arguments

path	full path to the CellProfiler output folder
display	single character indicating which features to display. Accepted entries are <code>cell_features</code> to display extracted single-cell features or <code>image_features</code> to display extracted image-level features.
cell_features	single character indicating the name of the file storing the extracted cell features.
image_features	single character indicating the name of the file storing the extracted image features.

Value

a `datatable` object

Author(s)

Nils Eling (<nils.eling@dqbm.uzh.ch>)

See Also

[read_cpout](#) for the CellProfiler reader function

Examples

```
path <- system.file("extdata/mockData/cpout", package = "imcRtools")

# Display cell features
show_cpout_features(path)

# Display image features
show_cpout_features(path, display = "image_features")
```

testInteractions *Tests if cell types interact more or less frequently than random*

Description

Cell-cell interactions are summarized in different ways and the resulting count is compared to a distribution of counts arising from random permutations.

Usage

```
testInteractions(
  object,
  group_by,
  label,
  colPairName,
```

```

method = c("classic", "histocat", "patch"),
patch_size = NULL,
iter = 1000,
p_threshold = 0.01,
BPPARAM = SerialParam()
)

```

Arguments

object	a SingleCellExperiment or SpatialExperiment object.
group_by	a single character indicating the colData(object) entry by which interactions are grouped. This is usually the image ID or patient ID.
label	single character specifying the colData(object) entry which stores the cell labels. These can be cell-types labels or other metadata entries.
colPairName	single character indicating the colPair(object) entry containing cell-cell interactions in form of an edge list.
method	which cell-cell interaction counting method to use (see details)
patch_size	if method = "patch", a single numeric specifying the minimum number of neighbors of the same type to be considered a patch (see details)
iter	single numeric specifying the number of permutations to perform
p_threshold	single numeric indicating the empirical p-value threshold at which interactions are considered to be significantly enriched or depleted per group.
BPPARAM	parameters for parallelized processing.

Value

a DataFrame containing one row per group_by entry and unique label entry combination (from_label, to_label). The object contains following entries:

- ct: stores the interaction count as described in the details
- p_gt: stores the fraction of perturbations equal or greater than ct
- p_lt: stores the fraction of perturbations equal or less than ct
- interaction: is there the tendency for a positive interaction (attraction) between from_label and to_label? Is p_lt greater than p_gt?
- p: the smaller value of p_gt and p_lt.
- sig: is p smaller than p_threshold?
- sigval: Combination of interaction and sig.
 - -1: interaction == FALSE and sig == TRUE
 - 0: sig == FALSE
 - 1: interaction == TRUE and sig == TRUE

NA is returned if a certain label is not present in this grouping level.

Counting and summarizing cell-cell interactions

In principle, the `countInteractions` function counts the number of edges (interactions) between each set of unique entries in `colData(object)[[label]]`. Simplified, it counts for each cell of type A the number of neighbors of type B. This count is averaged within each unique entry `colData(object)[[group_by]]` in three different ways:

1. `method = "classic"`: The count is divided by the total number of cells of type A. The final count can be interpreted as "How many neighbors of type B does a cell of type A have on average?"
2. `method = "histocat"`: The count is divided by the number of cells of type A that have at least one neighbor of type B. The final count can be interpreted as "How many many neighbors of type B has a cell of type A on average, given it has at least one neighbor of type B?"
3. `method = "patch"`: For each cell, the count is binarized to 0 (less than `patch_size` neighbors of type B) or 1 (more or equal to `patch_size` neighbors of type B). The binarized counts are averaged across all cells of type A. The final count can be interpreted as "What fraction of cells of type A have at least a given number of neighbors of type B?"

Testing for significance

Within each unique entry to `colData(object)[[group_by]]`, the entries of `colData(object)[[label]]` are randomized `iter` times. For each iteration, the interactions are counted as described above. The result is a distribution of the interaction count under spatial randomness. The observed interaction count is compared against this Null distribution to derive empirical p-values:

`p_gt`: fraction of perturbations equal or greater than the observed count

`p_lt`: fraction of perturbations equal or less than the observed count

Based on these empirical p-values, the interaction score (attraction or avoidance), overall p value and significance by comparison to `p_threshold` (`sig` and `signal`) are derived.

Author(s)

Vito Zanotelli

Jana Fischer

adapted by Nils Eling (<nils.eling@dqbm.uzh.ch>)

References

Schulz, D. et al., Simultaneous Multiplexed Imaging of mRNA and Proteins with Subcellular Resolution in Breast Cancer Tissue Samples by Mass Cytometry., *Cell Systems* 2018 6(1):25-36.e5

Shapiro, D. et al., histoCAT: analysis of cell phenotypes and interactions in multiplex image cytometry data, *Nature Methods* 2017 14, p. 873–876

See Also

`countInteractions` for counting (but not testing) cell-cell interactions per grouping level.

`bpparam` for the parallelised backend

Examples

```
library(cytomapper)
data(pancreasSCE)

pancreasSCE <- buildSpatialGraph(pancreasSCE, img_id = "ImageNb",
                                type = "knn", k = 3)

# Classic style calculation
(out <- testInteractions(pancreasSCE,
                        group_by = "ImageNb",
                        label = "CellType",
                        method = "classic",
                        colPairName = "knn_interaction_graph",
                        iter = 1000))

# HistoCAT style calculation
(out <- testInteractions(pancreasSCE,
                        group_by = "ImageNb",
                        label = "CellType",
                        method = "histocat",
                        colPairName = "knn_interaction_graph",
                        iter = 1000))

# Patch style calculation
(out <- testInteractions(pancreasSCE,
                        group_by = "ImageNb",
                        label = "CellType",
                        method = "patch",
                        patch_size = 3,
                        colPairName = "knn_interaction_graph",
                        ))
```

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