

Package: MOCHA (via r-universe)

March 20, 2025

Type Package

Title Modeling for Single-Cell Open Chromatin Analysis

Version 1.1.0

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Description A statistical framework and analysis tool for open chromatin analysis designed specifically for single cell ATAC-seq (Assay for Transposase-Accessible Chromatin) data, after cell type/cluster identification. These novel modules remove unwanted technical variation, identify open chromatin, robustly models repeated measures in single cell data, implement advanced statistical frameworks to model zero-inflation for differential and co-accessibility analyses, and integrate with existing databases and modules for downstream analyses to reveal biological insights. MOCHA provides a statistical foundation for complex downstream analysis to help advance the potential of single cell ATAC-seq for applied studies. Methods for zero-inflated statistics are as described in: Ghazanfar, S., Lin, Y., Su, X. et al. (2020) <doi:10.1038/s41592-020-0885-x>. Pimentel, Ronald Silva, ``Kendall's Tau and Spearman's Rho for Zero-Inflated Data'' (2009) <<https://scholarworks.wmich.edu/dissertations/721/>>.

License GPL (>= 3)

Encoding UTF-8

LazyData true

RoxygenNote 7.2.3

Depends R (>= 4.1.0)

Imports data.table, plyranges (>= 1.14.0), dplyr, GenomicRanges, RaggedExperiment, MultiAssayExperiment, SummarizedExperiment, stringr, ggbio, wCorr, magrittr, rlang, AnnotationDbi, BiocGenerics, GenomeInfoDb, GenomicFeatures, IRanges, S4Vectors, assertthat, ensemblDb, ggplot2, ggrepel, matrixStats, methods, qvalue, scales, tidyr, ggridges, pbapply, BSgenome, tidyselect, lifecycle

Suggests ArchR, RMariaDB, motifmatchr, BiocManager,
 TxDb.Hsapiens.UCSC.hg38.refGene,
 TxDb.Hsapiens.UCSC.hg19.knownGene, org.Hs.eg.db,
 BSgenome.Hsapiens.UCSC.hg19, withr, knitr, rmarkdown, chromVAR,
 testthat (>= 3.0.0), uwot, irlba, glmmTMB, Matrix, waldo,
 purrr, lmerTest, Biobase, lme4, zip, rtracklayer, cowplot,
 mixtools, zoo

Additional_repositories <https://imran-aifi.github.io/drat>,
<https://cranhaven.r-universe.dev>

VignetteBuilder knitr

Config/testthat/edition 3

NeedsCompilation no

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Date/Publication 2024-01-25 12:20:12 UTC

Config/pak/sysreqs make libicu-dev libpng-dev libxml2-dev libssl-dev

Repository <https://cranhaven.r-universe.dev>

RemoteUrl <https://github.com/cranhaven/cranhaven.r-universe.dev>

RemoteRef package/MOCHA

RemoteSha b07f58b7dc71d127bc9b549150dd00936b49c87a

RemoteSubdir MOCHA

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.counts_plot_default_theme

Default ggplot theme for counts plot

Description

Default ggplot theme for counts plot

Usage

```
.counts_plot_default_theme
```

Format

An object of class `list` of length 10.

```
.gene_plot_theme
```

Common theme for gene plots

Description

Common theme for gene plots

Usage

```
.gene_plot_theme
```

Format

An object of class `list` of length 5.

```
addAccessibilityShift addAccessibilityShift
```

Description

`addAccessibilityShift` will add a new condition to the `SummarizedExperiment` output of `extractRegion`, which will contain the difference in accessibility between two conditions

Usage

```
addAccessibilityShift(countSE, foreground, background, assayName = NULL)
```

Arguments

<code>countSE</code>	The <code>SummarizedExperiment</code> object output from <code>extractRegion</code>
<code>foreground</code>	Group that will be used as the foreground for the subtraction of accessibility
<code>background</code>	Group that will be used as the background for the subtraction of accessibility
<code>assayName</code>	The name given to the new assay that is difference in accessibility between foreground and background.

Value

`countSE` a `SummarizedExperiment` containing coverage for the given input cell populations.

Examples

```
## Not run:
# CountSE is a SummarizedExperiment generated by extractRegion()
countSE <- MOCHA::addAccessibilityShift(
  CountSE = CountSE,
  foreground = "Condition1",
  background = "Condition2",
  assayName = "AccessibilityChanges"
)

## End(Not run)
```

```
addMotifSet      addMotifSet
```

Description

addMotifSet Identify motifs within your peakset.

Usage

```
addMotifSet(
  SampleTileObj,
  motifPWMs,
  w = 7,
  returnSTM = TRUE,
  motifSetName = "Motifs"
)
```

Arguments

SampleTileObj	A SummarizedExperiment, specifically the output of getSampleTileMatrix
motifPWMs	A pwms object for the motif database. Either PFMMatrix, PFMMatrixList, PWMMatrix, or PWMMatrixList
w	Parameter for motifmatchr controlling size in basepairs of window for filtration. Default is 7.
returnSTM	If TRUE, return the modified SampleTileObj with motif set added to metadata (default). If FALSE, return the motifs from motifmatchr as a GRanges.
motifSetName	Name to give motifList in the SampleTileObj's metadata if 'returnSTM=TRUE'. Default is 'Motifs'.

Value

the modified SampleTileObj with motifs added to the metadata

Examples

```
## Not run:
# load a curated motif set from library(chromVARmotifs)
# included with ArchR installation
data(human_pwm_v2)
SE_with_motifs <- addMotifSet(
  SampleTileObj,
  motifPWMs = human_pwm_v2,
  returnSTM = TRUE, motifSetName = "Motifs", w = 7
)

## End(Not run)
```

annotateTiles	annotateTiles
---------------	---------------

Description

annotateTiles annotates a set of sample-tile matrices given with gene annotations. Details on TxDb and Org annotation packages and available annotations can be found at Bioconductor: <https://bioconductor.org/packages>

Usage

```
annotateTiles(Obj, TxDb = NULL, Org = NULL, promoterRegion = c(2000, 100))
```

Arguments

Obj	A RangedSummarizedExperiment generated from getSampleTileMatrix, containing TxDb and Org in the metadata. This may also be a GRanges object.
TxDb	The annotation package for TxDb object for your genome. Optional, only required if Obj is a GRanges.
Org	The genome-wide annotation for your organism. Optional, only required if Obj is a GRanges.
promoterRegion	Optional list containing the window size in basepairs defining the promoter region. The format is (upstream, downstream). Default is (2000, 100).

Value

Obj, the input data structure with added gene annotations (whether GRanges or SampleTileObj)

Examples

```
## Not run:
library(TxDb.Hsapiens.UCSC.hg38.refGene)
library(org.Hs.eg.db)
SampleTileMatricesAnnotated <- MOCHA::annotateTiles(
  SampleTileMatrices,
```

```

    Txdb = TxDb.Hsapiens.UCSC.hg38.refGene,
    Org = org.Hs.eg.db
)

## End(Not run)

```

```

bulkDimReduction      bulkDimReduction

```

Description

bulkDimReduction runs dimensionality reduction (either PCA or LSI). We adapt Signac's

Usage

```

bulkDimReduction(
  SampleTileObj,
  cellType = "All",
  componentNumber = 30,
  method = "LSI",
  verbose = FALSE
)

```

Arguments

SampleTileObj	The SummarizedExperiment object output from getSampleTileMatrix
cellType	vector of strings. Cell subsets for which to call peaks. This list of group names must be identical to names that appear in the SampleTileObj. Optional, if cellPopulations='ALL', then peak calling is done on all cell populations. Default is 'ALL'.
componentNumber	integer. Number of components to include in LSI, or PCA This must be strictly less than
method	a string. Represents the method to use. Includes LSI or PCA, but we do not recommend PCA for scATAC pseudobulk.
verbose	Set TRUE to display additional messages. Default is FALSE.

Value

SEObj a SummarizedExperiment containing PC components from dimensionality reduction and metadata from the SampleTileObj

References

LSI method adapted from Andrew Hill: <http://andrewjohnhill.com/blog/2019/05/06/dimensionality-reduction-for-scatac-data/>

Examples

```
## Not run:
LSIObj <- MOCHA::bulkDimReduction(SampleTileObj, cellType = "CD16_Mono")

## End(Not run)
```

bulkUMAP

bulkUMAP

Description

bulkUMAP generates UMAP from pseudobulk LSIObj object, and merges in metadata.

Usage

```
bulkUMAP(
  SEObj,
  assay = "LSI",
  components = c(1:30),
  nNeighbors = 15,
  returnModel = FALSE,
  seed = 1,
  ...
)
```

Arguments

SEObj	The SummarizedExperiment object output from bulkDimReduction, or an STM, subsetted down to just one cell type.
assay	A string, describing the name of the assay within SEObj to run UMAP ('PCA', 'LSI', or 'counts').
components	A vector of integers. Number of components to include in LSI (1:30 typically).
nNeighbors	See umap . The size of local neighborhood (in terms of number of neighboring sample points) used for manifold approximation. Default is 15.
returnModel	A boolean. Default is FALSE. If set to true, it will return a list, where the first is the UMAP coordinates with metadata for plotting, and the second is the full UMAP model so further projection can occur.
seed	an integer. Represents the random seed to pass to the UMAP. Default seed is 1.
...	Additional arguments to be passed to umap .

Value

fullUMAP data.frame of UMAP values with metadata attached.

Examples

```
## Not run:  
UMAPvalues <- MOCHA::bulkUMAP(LSIObj)  
  
## End(Not run)
```

callOpenTiles	callOpenTiles <i>Perform peak-calling on a set of fragments or an ArchR Project.</i>
---------------	--

Description

callOpenTiles is the main peak-calling function in MOCHA that serves as a wrapper function to call peaks provided a set of fragment files and an ArchR Project for meta-data purposes

Usage

```
callOpenTiles(  
  ATACFragments,  
  cellColData,  
  blackList,  
  genome,  
  cellPopLabel,  
  cellPopulations = "ALL",  
  studySignal = NULL,  
  generalizeStudySignal = FALSE,  
  cellCol = "RG",  
  TxDb,  
  OrgDb,  
  outDir,  
  numCores = 30,  
  verbose = FALSE,  
  force = FALSE  
)  
  
## S4 method for signature 'GRangesList'  
callOpenTiles(  
  ATACFragments,  
  cellColData,  
  blackList,  
  genome,  
  cellPopLabel,  
  cellPopulations = "ALL",  
  studySignal = NULL,  
  generalizeStudySignal = FALSE,  
  cellCol = "RG",  
  TxDb,
```

```

    OrgDb,
    outDir,
    numCores = 30,
    verbose = FALSE,
    force = FALSE
)

## S4 method for signature 'list'
callOpenTiles(
  ATACFragments,
  cellColData,
  blackList,
  genome,
  cellPopLabel,
  cellPopulations = "ALL",
  studySignal = NULL,
  generalizeStudySignal = FALSE,
  cellCol = "RG",
  TxDb,
  OrgDb,
  outDir,
  numCores = 30,
  verbose = FALSE,
  force = FALSE
)

.callOpenTiles_ArchR(
  ATACFragments,
  cellPopLabel,
  cellPopulations = "ALL",
  studySignal = NULL,
  generalizeStudySignal = FALSE,
  TxDb,
  OrgDb,
  outDir = NULL,
  numCores = 30,
  verbose = FALSE,
  force = FALSE
)

```

Arguments

ATACFragments	an ArchR Project, or a GRangesList of fragments. Each GRanges in the GRanges list must have unique cell IDs in the column given by 'cellCol'.
cellColData	A DataFrame containing cell-level metadata. This must contain both a column 'Sample' with unique sample IDs and the column specified by 'cellPopLabel'.
blackList	A GRanges of blacklisted regions
genome	A BSgenome object, or the full name of an installed BSgenome data package,

or a short string specifying the name of an NCBI assembly (e.g. "GRCh38", "TAIR10.1", etc...) or UCSC genome (e.g. "hg38", "bosTau9", "galGal6", "ce11", etc...). The supplied short string must refer unambiguously to an installed BSgenome data package. See [getBSgenome](#).

cellPopLabel	string indicating which column in the ArchRProject metadata contains the cell population label.
cellPopulations	vector of strings. Cell subsets for which to call peaks. This list of group names must be identical to names that appear in the ArchRProject metadata. Optional, if cellPopulations='ALL', then peak calling is done on all cell populations in the ArchR project metadata. Default is 'ALL'.
studySignal	The median signal (number of fragments) in your study. If not set, this will be calculated using the input ArchR project but relies on the assumption that the ArchR project encompasses your whole study (i.e. is not a subset).
generalizeStudySignal	If 'studySignal' is not provided, calculate the signal as the mean of the mean & median number of fragments for of individual samples within each cell population. This may improve MOCHA's ability to generalize to datasets with XXXXXX #TODO. Default is FALSE, use the median number of fragments.
cellCol	The column in cellColData specifying unique cell ids or barcodes. Default is "RG", the unique cell identifier used by ArchR.
TxDB	The exact package name of a TxDb-class transcript annotation package for your organism (e.g. "TxDb.Hsapiens.UCSC.hg38.refGene"). This must be installed. See Bioconductor AnnotationData Packages .
OrgDb	The exact package name of a OrgDb-class genome wide annotation package for your organism (e.g. "org.Hs.eg.db"). This must be installed. See Bioconductor AnnotationData Packages
outDir	is a string describing the output directory for coverage files. Must be a complete directory string. With ArchR input, set outDir to NULL to create a directory within the input ArchR project directory named MOCHA for saving files.
numCores	integer. Number of cores to parallelize peak-calling across multiple cell populations.
verbose	Set TRUE to display additional messages. Default is FALSE.
force	Optional, whether to force creation of coverage files if they already exist. Default is FALSE.

Value

tileResults A MultiAssayExperiment object containing ranged data for each tile

Examples

```
## Not run:
# Starting from an ArchR Project:
tileResults <- MOCHA::callOpenTiles(
  ArchRProj = myArchRProj,
```

```

    cellPopLabel = "celltype_labeling",
    cellPopulations = "CD4",
    TxDb = "TxDb.Hsapiens.UCSC.hg38.refGene",
    OrgDb = "org.Hs.eg.db",
    numCores = 1
  )

## End(Not run)

# Starting from GRangesList
if (
  requireNamespace("BSgenome.Hsapiens.UCSC.hg19") &&
  requireNamespace("TxDb.Hsapiens.UCSC.hg38.refGene") &&
  requireNamespace("org.Hs.eg.db")
) {
  tiles <- MOCHA::callOpenTiles(
    ATACFragments = MOCHA::exampleFragments,
    cellColData = MOCHA::exampleCellColData,
    blacklist = MOCHA::exampleBlackList,
    genome = "BSgenome.Hsapiens.UCSC.hg19",
    TxDb = "TxDb.Hsapiens.UCSC.hg38.refGene",
    OrgDb = "org.Hs.eg.db",
    outDir = tempdir(),
    cellPopLabel = "Clusters",
    cellPopulations = c("C2", "C5"),
    numCores = 1
  )
}

```

```

combineSampleTileMatrix
      combineSampleTileMatrix

```

Description

combineSampleTileMatrix combines all celltypes in a SampleTileMatrix object into a SummarizedExperiment with one single matrix across all cell types and samples,

Usage

```
combineSampleTileMatrix(SampleTileObj, NAtoZero = TRUE, verbose = FALSE)
```

Arguments

SampleTileObj	The SummarizedExperiment object output from getSampleTileMatrix containing your sample-tile matrices
NAtoZero	Set NA values in the sample-tile matrix to zero
verbose	Set TRUE to display additional messages. Default is FALSE.

Value

TileCorr A data.table correlation matrix

differentialsToGRanges

differentialsToGRanges Converts a data.frame matrix to a GRanges, preserving additional columns as GRanges metadata

Description

differentialsToGRanges Converts a data.frame matrix to a GRanges, preserving additional columns as GRanges metadata

Usage

```
differentialsToGRanges(differentials, tileColumn = "Tile")
```

Arguments

`differentials` a matrix/data.frame with a column `tileColumn` containing region strings in the format "chr:start-end"

`tileColumn` name of column containing region strings. Default is "Tile".

Value

a GRanges containing all original information

`exampleBlackList` *exampleBlackList*

Description

Example input of a blackList extracted from the PBMC_Small dataset consisting of 2k cells and spanning chr1 and 2 (~2-300MB). The data is publicly available with the ArchR package at <<https://www.archrproject.com/re>

Usage

```
exampleBlackList
```

Format

A GRanges object with 210 ranges and 2 metadata columns

exampleCellColData *exampleCellColData*

Description

Example input of cellColData extracted from the PBMC_Small dataset consisting of 2k cells and spanning chr1 and 2 (~2-300MB). The data is publicly available with the ArchR package at <<https://www.archrproject.com/re>

Usage

```
exampleCellColData
```

Format

A DataFrame with 2217 rows and 3 columns

exampleFragments *exampleFragments*

Description

Example input of ATAC fragments extracted from the PBMC_Small dataset consisting of 2k cells and spanning chr1 and 2 (~2-300MB). This subset consists of two cell populations: Clusters C2 and C5. The data is publicly available with the ArchR package at <<https://www.archrproject.com/reference/getTestProject.html>>

Usage

```
exampleFragments
```

Format

A list of 2 GRanges objects

exportCoverage	exportCoverage
----------------	----------------

Description

exportCoverage will export normalized coverage files to BigWig files, either as sample-specific or sample-averaged files, for visualization in genome browsers.

Usage

```
exportCoverage(  
  SampleTileObject,  
  dir = getwd(),  
  type = TRUE,  
  cellPopulations = "ALL",  
  groupColumn = NULL,  
  subGroups = NULL,  
  sampleSpecific = FALSE,  
  saveFile = TRUE,  
  numCores = 1,  
  verbose = FALSE  
)
```

Arguments

SampleTileObject	The SummarizedExperiment object output from getSampleTileMatrix
dir	string. Directory to save files to.
type	Boolean. Default is TRUE, and exports Coverage. If set to FALSE, exports Insertions.
cellPopulations	vector of strings. Cell subsets for which to call peaks. This list of group names must be identical to names that appear in the SampleTileObject. Optional, if cellPopulations='ALL', then peak calling is done on all cell populations. Default is 'ALL'.
groupColumn	Optional, the column containing sample group labels for returning coverage within sample groups. Default is NULL, all samples will be used.
subGroups	a list of subgroup(s) within the groupColumn from the metadata. Optional, default is NULL, all labels within groupColumn will be used.
sampleSpecific	If TRUE, a BigWig will export for each sample-cell type combination.
saveFile	Boolean. If TRUE, it will save to a BigWig. If FALSE, it will return the GRangesList without writing a BigWig.
numCores	integer. Number of cores to parallelize peak-calling across multiple cell populations
verbose	Set TRUE to display additional messages. Default is FALSE.

Value

countSE a SummarizedExperiment containing coverage for the given input cell populations.

Examples

```
## Not run:
MOCHA::exportCoverage(
  SampleTileObject = SampleTileMatrices,
  cellPopulations = "ALL",
  numCores = 30,
  sampleSpecific = FALSE
)

## End(Not run)
```

```
exportDifferentials  exportDifferentials
```

Description

exportDifferentials exports the differential peaks output GRangesList output from getDifferentialAccessibleTiles to bigBed format for visualization in genome browsers.

Usage

```
exportDifferentials(
  SampleTileObject,
  DifferentialsGRLList,
  outDir,
  verbose = FALSE
)
```

Arguments

SampleTileObject	The SummarizedExperiment object output from getSampleTileMatrix
DifferentialsGRLList	GRangesList output from getDifferentialAccessibleTiles
outDir	Desired output directory where bigBed files will be saved
verbose	Set TRUE to display additional messages. Default is FALSE.

Value

outList A List of output filepaths

Examples

```
## Not run:
MOCHA::exportDifferentials(
  SampleTileObject = SampleTileMatrices,
  DifferentialsGRList,
  outDir = tempdir(),
  verbose = TRUE
)

## End(Not run)
```

```
exportMotifs
```

```
exportMotifs
```

Description

exportMotifs exports a motif set GRanges from running addMotifSet(returnSTM=FALSE) to bigBed file files for visualization in genome browsers.

Usage

```
exportMotifs(
  SampleTileObject,
  motifsGRanges,
  motifSetName = "motifs",
  filterByOpenTiles = FALSE,
  outDir,
  verbose = FALSE
)
```

Arguments

SampleTileObject	The SummarizedExperiment object output from getSampleTileMatrix
motifsGRanges	A GRanges containing motif annotations, typically from addMotifSet(returnSTM=FALSE)
motifSetName	Optional, a name indicating the motif set. Used to name files in the specified outdir. Default is "motifs".
filterByOpenTiles	Boolean. If TRUE, a bigBed file will be exported for each cell population with motifs filtered to those occurring only in open tiles.
outDir	Desired output directory where bigBed files will be saved
verbose	Set TRUE to display additional messages. Default is FALSE.

Value

outList A List of output filepaths

Examples

```
## Not run:
MOCHA::exportMotifs(
  SampleTileObject = SampleTileMatrices,
  motifsGRanges,
  motifSetName = "CISBP",
  filterByOpenTiles = FALSE,
  outDir = tempdir(),
  verbose = TRUE
)

## End(Not run)
```

exportOpenTiles	exportOpenTiles
-----------------	-----------------

Description

exportOpenTiles exports the open tiles of a given cell population to bigBed file for visualization in genome browsers.

Usage

```
exportOpenTiles(SampleTileObject, cellPopulation, outDir, verbose = FALSE)
```

Arguments

SampleTileObject	The SummarizedExperiment object output from getSampleTileMatrix
cellPopulation	The name of the cell population to export
outDir	Desired output directory where bigBed files will be saved
verbose	Set TRUE to display additional messages. Default is FALSE.

Value

outList A List of output filepaths

Examples

```
## Not run:
MOCHA::exportOpenTiles(
  SampleTileObject = SampleTileObject,
  cellPopulation,
  outDir = tempdir(),
  verbose = TRUE
)

## End(Not run)
```

```
exportSmoothedInsertions
      exportSmoothedInsertions
```

Description

exportSmoothedInsertions Takes a SampleTileMatrix with linked insertion files and applies a smoothing filter (a rolling sum then rolling median) to the insertions, finally exporting the smoothed insertion files to bigwig format.

Usage

```
exportSmoothedInsertions(
  SampleTileObj,
  cellPopulation,
  outDir = NULL,
  sumWidth = 10,
  medianWidth = 11,
  force = FALSE,
  slow = FALSE,
  verbose = FALSE
)
```

Arguments

SampleTileObj	A MultiAssayExperiment or RangedSummarizedExperiment from MOCHA
cellPopulation	A string denoting the cell population of interest
outDir	Directory to write output bigwig files. Default is NULL, where the directory in 'SampleTileObj@metadata\$Directory' will be used.
sumWidth	Window size for rolling sum in basepairs. Default is 10.
medianWidth	Window size for rolling median in basepairs. Must be odd. Default is 11.
force	Set TRUE to overwrite existing files. Default is FALSE.
slow	Set TRUE to bypass optimisations and compute smoothing filter directly on the whole genome. May run slower and consume more RAM. Default is FALSE.
verbose	Set TRUE to display additional messages. Default is FALSE.

Value

outPaths List of paths of exported insertion files

Examples

```
## Not run:
# Depends on and manipulates files on filesystem
outPath <- MOCHA::exportSmoothedInsertions(
```


subGroups	a list of subgroup(s) within the groupColumn from the metadata. Optional, default is NULL, all labels within groupColumn will be used.
sampleSpecific	If TRUE, get a sample-specific count dataframe out. Default is FALSE, average across samples and get a dataframe out.
approxLimit	Optional limit to region size, where if region is larger than approxLimit basepairs, binning will be used. Default is 100000.
binSize	Optional numeric, size of bins in basepairs when binning is used. Default is 250.
sliding	Optional numeric. Default is NULL. This number is the size of the sliding window for generating average intensities.
numCores	integer. Number of cores to parallelize peak-calling across multiple cell populations
verbose	Set TRUE to display additional messages. Default is FALSE.

Value

countSE a SummarizedExperiment containing coverage for the given input cell populations.

Examples

```
## Not run:
countSE <- MOCHA::extractRegion(
  SampleTileObj = SampleTileMatrices,
  cellPopulations = "ALL",
  region = "chr1:18137866-38139912",
  numCores = 30,
  sampleSpecific = FALSE
)

## End(Not run)
```

filterCoAccessibleLinks

filterCoAccessibleLinks

Description

filterCoAccessibleLinks will filter the output from getCoAccessibleLinks by a threshold, retaining links with a absolute correlation greater than the threshold. This function also adds the chr, start, and end site of each link to the output table.

Usage

```
filterCoAccessibleLinks(TileCorr, threshold = 0.5)
```

Arguments

TileCorr	The correlation table output from getCoAccessibleLinks
threshold	Keep

Value

FilteredTileCorr The filtered correlation table with chr, start, and end site of each link

Examples

```
## Not run:
# links is the output of MOCHA::getCoAccessibleLinks
MOCHA::filterCoAccessibleLinks(links, threshold = 0.5)

## End(Not run)
```

<code>finalModelObject</code>	<i>finalModelObject</i>
-------------------------------	-------------------------

Description

Trained MOCHA models - LOESS and linear regression

Usage

```
finalModelObject
```

Format

A list of lists containing 2 items: "Loess" and "Linear" each with "Total" "Max" and "Intercept"

Loess LOESS model

Linear Linear model

getAltTSS	<i>Annotate Peaks falling in Transcription Start Sites (TSS) and identify alternatively regulated TSSs for each gene.</i>
-----------	---

Description

getAltTSS Pulls out all peaks that fall in TSS, annotates them with the name of gene, and identifies genes that have evidence for alternatively regulated TSSs, including both type i (only some of the open TSSs for a gene are significantly more (or less) accessible), and type ii (multiple TSSs are significant different, with some being more accessible and others less). Alternatively, this function will return all open TSSs with differential measurements if the returnAllTSS flag is set to TRUE.

Usage

```
getAltTSS(
  completeDAPs,
  returnAllTSS = FALSE,
  nuancedTSS = TRUE,
  nuancedTSSGap = 150,
  threshold = 0.2,
  TxDb,
  OrgDb
)
```

Arguments

completeDAPs	GRanges object that contains the differential measurements across all peaks (unfiltered DAPs). Will also work with data.frame or data.table version of a GRanges object. If you want alternatively regulated TSSs, the object must include a column names 'FDR', and 'Log2FC_C', which is standard for MOCHA differentials.
returnAllTSS	Flag to return all TSSs with DAPs measurements, without filtering for alternative TSS usage. If multiple TSSs fall within the same tile, then that tile will be repeated for each TSS.
nuancedTSS	True/False flag to determine if alternative TSS genes should be filtered out if all their differential TSS usage falls within too small of a range. Default is TRUE
nuancedTSSGap	Minimum distance between TSSs needed for them to considered distinctly regulated TSSs. If two TSSs are too close, it is unclear and highly unlikely that ATAC data can distinguish between them. Default is 150 bp.
threshold	FDR Threshold for determining significant vs non-significant changes in accessibility. Following MOCHA's standards, default is 0.2.
TxDb	The TxDb-class transcript annotation package for your organism (e.g. "TxDb.Hsapiens.UCSC.hg38.refGene"). This must be installed. See Bioconductor AnnotationData Packages .
OrgDb	The OrgDb-class genome wide annotation package for your organism (e.g. "org.Hs.eg.db"). This must be installed. See Bioconductor AnnotationData Packages

Value

tpeaks A GRanges containing annotated peaks falling in TSS

getAnnotationDbFromInstalledPkgrname

getAnnotationDbFromInstalledPkgrname *Loads and attaches an installed TxDb or OrgDb-class Annotation database package.*

Description

See [getBSgenome](#)

Usage

```
getAnnotationDbFromInstalledPkgrname(dbName, type)
```

Arguments

dbName Exact name of installed annotation data package.
 type Expected class of the annotation data package, must be either "OrgDb" or "TxDb".

Value

the loaded Annotation database object.#' @noRd

getCellPopMatrix getCellPopMatrix

Description

getCellPopMatrix pulls out the SampleTileMatrix of tiles called in one given cell population.

Usage

```
getCellPopMatrix(  
  SampleTileObj,  
  cellPopulation,  
  dropSamples = TRUE,  
  NtoZero = TRUE  
)
```


Arguments

SampleTileObj	The output from getSampleTileMatrix, a SummarizedExperiment of pseudobulk intensities across all tiles & cell types.
cellPopulation	The cell population you want to pull out.
dropSamples	Boolean flag to determine whether to drop samples that were too small for peak calling.
NAtoZero	Boolean flag to determine whether to replace NAs with zero

Value

sampleTileMatrix a matrix of samples by called tiles for a given cell population.

getCellTypes	<i>getCellTypes Extract cell type names from a Tile Results or Sample Tile object.</i>
--------------	--

Description

getCellTypes Returns a vector of cell names from a Tile Results or Sample Tile object.

Usage

```
getCellTypes(object)
```

Arguments

object	tileResults object from callOpenTiles or SummarizedExperiment from getSampleTileMatrix
--------	--

Value

a vector of cell type names.

getCellTypeTiles	<i>getCellTypeTiles Extract the GRanges for a particular cell type</i>
------------------	--

Description

getCellTypeTiles Returns a GRanges object of all tiles called for a certain cell type

Usage

```
getCellTypeTiles(object, cellType)
```

Arguments

object	A SampleTileObject.
cellType	A string describing one cell type.

Value

a vector of cell type names.

```
getCoAccessibleLinks  getCoAccessibleLinks
```

Description

getCoAccessibleLinks takes an input set of regions (tiles) and finds co-accessible neighboring regions within a window. Co-accessibility is defined as the correlation between two region intensity (openness) across samples.

Usage

```
getCoAccessibleLinks(
  SampleTileObj,
  cellPopulation = "All",
  regions,
  chrChunks = 1,
  windowSize = 1 * 10^6,
  numCores = 1,
  ZI = TRUE,
  approximateTile = FALSE,
  verbose = FALSE
)
```

Arguments

SampleTileObj	The SummarizedExperiment object output from getSampleTileMatrix containing your sample-tile matrices
cellPopulation	A string denoting the cell population of interest, which must be present in SampleTileObj
regions	a GRanges object or vector or strings containing the regions on which to compute co-accessible links. Strings must be in the format "chr:start-end", e.g. "chr4:1300-2222". Can be the output from getDifferentialAccessibleTiles.
chrChunks	This functions subsets by groups of chromosome, and then parallelizes within each group of chromosomes when running correlations. This method keeps memory low. To speed things up on high performing platforms, you can chunk out more than one chromosome at a time. Default is chrChunks = 1, so only one chromosome at a time.

windowSize	the size of the window, in basepairs, around each input region to search for co-accessible links
numCores	Optional, the number of cores to use with multiprocessing. Default is 1.
ZI	boolean flag that enables zero-inflated (ZI) Spearman correlations to be used. Default is TRUE. If FALSE, skip zero-inflation and calculate the normal Spearman.
approximateTile	If set to TRUE, it will use all tiles that overlap with the regions given, instead of finding an exact match to the regions variable. Default is FALSE.
verbose	Set TRUE to display additional messages. Default is FALSE.

Details

The technical details of the zero-inflated correlation can be found here:

Pimentel, Ronald Silva, "Kendall's Tau and Spearman's Rho for Zero-Inflated Data" (2009). Dissertations.

while the implementation (schOT R package), can be found here: <http://www.bioconductor.org/packages/release/bioc/html/schOT/>

Value

TileCorr A data.table correlation matrix

getCoverage	<i>Get sample-specific coverage files for each sample-cell population.</i>
-------------	--

Description

getCoverage takes the output of MOCHA::getPopFragments and returns a GRanges of single-basepair resolution coverage.

Usage

```
getCoverage(
  popFragments,
  normFactor,
  TxDb,
  cl,
  filterEmpty = FALSE,
  verbose = FALSE
)
```

Arguments

popFrag	GRangesList of fragments for all sample/cell populations
normFactor	Normalization factor. Can be either be one, in which case all coverage files will be normalized by the same value, or the same length as the GRangesList
TxDb	The TxDb-class transcript annotation package for your organism (e.g. "TxDb.Hsapiens.UCSC.hg38.refGene"). This must be installed. See Bioconductor AnnotationData Packages .
cl	cl argument to pblapply
filterEmpty	True/False flag on whether or not to carry forward regions without coverage.
verbose	Boolean variable to determine verbosity of output.

Value

popCounts A GRangesList of coverage for each sample and cell population

```
getDifferentialAccessibleTiles
      getDifferentialAccessibleTiles
```

Description

`getDifferentialAccessibleTiles` allows you to determine whether regions of chromatin are differentially accessible between groups by conducting a test

Usage

```
getDifferentialAccessibleTiles(
  SampleTileObj,
  cellPopulation,
  groupColumn,
  foreground,
  background,
  signalThreshold = 12,
  minZeroDiff = 0.5,
  fdrToDisplay = 0.2,
  outputGRanges = TRUE,
  numCores = 1,
  verbose = FALSE
)
```

Arguments

SampleTileObj	The SummarizedExperiment object output from <code>getSampleTileMatrix</code>
cellPopulation	A string denoting the cell population of interest
groupColumn	The column containing sample group labels

foreground	The foreground group of samples for differential comparison
background	The background group of samples for differential comparison
signalThreshold	Minimum median intensity required to keep tiles for differential testing to increase statistical power in small sample cohorts. Default is 12.
minZeroDiff	Minimum difference in average dropout rates across groups require to keep tiles for differential testing. Default is 0.5 (50%).
fdrToDisplay	False-discovery rate used only for standard output messaging. Default is 0.2.
outputGRanges	Outputs a GRanges if TRUE and a data.frame if FALSE. Default is TRUE.
numCores	The number of cores to use with multiprocessing. Default is 1.
verbose	Set TRUE to display additional messages. Default is FALSE.

Value

full_results The differential accessibility results as a GRanges or matrix data.frame depending on the flag 'outputGRanges'.

Examples

```
## Not run:
cellPopulation <- "MAIT"
foreground <- "Positive"
background <- "Negative"
# Standard output will display the number of tiles found below a false-discovery rate threshold.
# This parameter does not filter results and only affects the aforementioned message.
fdrToDisplay <- 0.2
# Choose to output a GRanges or data.frame.
# Default is TRUE
outputGRanges <- TRUE
# SampleTileMatrices is the output of MOCHA::getSampleTileMatrix
differentials <- MOCHA::getDifferentialAccessibleTiles(
  SampleTileObj = SampleTileMatrices,
  cellPopulation = cellPopulation,
  groupColumn = groupColumn,
  foreground = foreground,
  background = background,
  fdrToDisplay = fdrToDisplay,
  outputGRanges = outputGRanges,
  numCores = numCores
)
## End(Not run)
```

```
getIntensityThreshold  getIntensityThreshold
```

Description

getIntensityThreshold takes the output of peak calling with callOpenTiles and creates sample-tile matrices containing the signal intensity at each tile.

Usage

```
getIntensityThreshold(
  TSAM,
  cellPopulations = "all",
  type = "mean",
  returnPlots = TRUE,
  verbose = FALSE
)
```

Arguments

TSAM	a SummarizedExperiment object generated by MOCHA
cellPopulations	vector of strings. Cell subsets found in the TSAM, or the word 'All' if all should be used.
type	string. Describes the type of metric to be used. Options include median or mean.
returnPlots	Boolean. Default is TRUE and returns a plot of
verbose	Set TRUE to display additional messages. Default is FALSE.

Value

plot object

```
getModelValues          getModelValues from runZIGLMM output.
```

Description

‘r lifecycle::badge("deprecated")’ This function is deprecated - improved modeling functions can be found in the package "ChAI" at <https://github.com/aifimmunology/ChAI> getModelValues Pull out a data.frame of model values (slope, significance, and std.error) for a given factor from the SummarizedExperiment output of runZIGLMM.

Usage

```
getModelValues(object, specificVariable)
```

Arguments

object A SummarizedExperiment object generated from runZIGLMM.
 specificVariable A string, describing the factor of influence.

Value

A data.frame of slopes, significance, and standard error for one factor.

Examples

```
## Not run:
age_df <- getModelValues(runZIGLMM_output, "Age")

## End(Not run)
```

 getPopFrag

Extract fragments by populations from an ArchR Project

Description

getPopFrag returns a list of sample-specific fragments per cell population as a GRangesList.

Usage

```
getPopFrag(
  ArchRProj,
  cellPopLabel,
  cellSubsets = "ALL",
  poolSamples = FALSE,
  numCores = 1,
  verbose = FALSE
)
```

Arguments

ArchRProj The ArchR Project.
 cellPopLabel The name of the metadata column of the ArchR Project that contains the populations of cells you want to extract fragments from.
 cellSubsets Default is 'ALL'. If you want to export only some populations, then give it a list of group names. This needs to be unique - no duplicated names. This list of group names must be identical to names that appear in the given cellPopLabel metadata column of the ArchR Project.
 poolSamples Set TRUE to pool sample-specific fragments by cell population. By default this is FALSE and sample-specific fragments are returned.
 numCores Number of cores to use.
 verbose Set TRUE to display additional messages. Default is FALSE.

Value

A list of GRanges containing fragments. Each GRanges corresponds to a population defined by cellSubsets and sample.

getPromoterGenes	getPromoterGenes
------------------	------------------

Description

getPromoterGenes Takes a rowRanges from annotateTiles and extracts a unique list of genes.

Usage

```
getPromoterGenes(GRangesObj)
```

Arguments

GRangesObj a GRanges object with a metadata column for tileType and Gene.

Value

vector of strings with gene names.

getSampleCellTypeMetadata	
	getSampleCellTypeMetadata <i>Extract Sample-celltype specific metadata</i>

Description

getSampleCellTypeMetadata Extract Sample-celltype specific metadata like fragment number, cell counts, and

Usage

```
getSampleCellTypeMetadata(object)
```

Arguments

object tileResults object from callOpenTiles or SummarizedExperiment from getSampleTileMatrix

Value

a SummarizedExperiment where each assay is a different type of metadata.

```
getSampleTileMatrix  getSampleTileMatrix
```

Description

getSampleTileMatrix takes the output of peak calling with callOpenTiles and creates sample-tile matrices containing the signal intensity at each tile.

Usage

```
getSampleTileMatrix(  
  tileResults,  
  cellPopulations = "ALL",  
  groupColumn = NULL,  
  threshold = 0.2,  
  numCores = 1,  
  verbose = FALSE  
)
```

Arguments

tileResults	a MultiAssayExperiment returned by callOpenTiles containing containing peak calling results.
cellPopulations	vector of strings. Cell subsets in TileResults for which to generate sample-tile matrices. This list of group names must be identical to names that appear in the ArchRProject metadata. If cellPopulations='ALL', then peak calling is done on all cell populations in the ArchR project metadata. Default is 'ALL'.
groupColumn	Optional, the column containing sample group labels for determining consensus tiles within sample groups. Default is NULL, all samples will be used for determining consensus tiles.
threshold	Threshold for consensus tiles, the minimum % of samples (within a sample group, if groupColumn is set) that a peak must be called in to be retained. If set to 0, retain the union of all samples' peaks (this is equivalent to a threshold of 1/numSamples). It is recommended to tune this parameter to omit potentially spurious peaks.
numCores	Optional, the number of cores to use with multiprocessing. Default is 1.
verbose	Set TRUE to display additional messages. Default is FALSE.

Value

SampleTileMatrices a MultiAssayExperiment containing a sample-tile intensity matrix for each cell population

Examples

```

# Starting from GRangesList
if (
  require(BSgenome.Hsapiens.UCSC.hg19) &&
  require(TxDb.Hsapiens.UCSC.hg38.refGene) &&
  require(org.Hs.eg.db)
) {
  tiles <- MOCHA::callOpenTiles(
    ATACFragments = MOCHA::exampleFragments,
    cellColData = MOCHA::exampleCellColData,
    blackList = MOCHA::exampleBlackList,
    genome = "BSgenome.Hsapiens.UCSC.hg19",
    TxDb = "TxDb.Hsapiens.UCSC.hg38.refGene",
    Org = "org.Hs.eg.db",
    outDir = tempdir(),
    cellPopLabel = "Clusters",
    cellPopulations = c("C2", "C5"),
    numCores = 1
  )

  SampleTileMatrices <- MOCHA::getSampleTileMatrix(
    tiles,
    cellPopulations = c("C2", "C5"),
    threshold = 0 # Take union of all samples' open tiles
  )
}

```

```

getSequencingBias      getSequencingBias

```

Description

getSequencingBias takes the output of peak calling with callOpenTiles and creates sample-tile matrices containing the signal intensity at each tile.

Usage

```

getSequencingBias(
  SampleTileObj,
  cellPopulations = "all",
  cellPopulation,
  groupColumn,
  foreground,
  background,
  verbose = TRUE
)

```

Arguments

SampleTileObj	a SummarizedExperiment object generated by MOCHA
cellPopulations	vector of strings. Cell subsets found in the TSAM, or the word 'All' if all should be used.
cellPopulation	A string denoting the cell population of interest
groupColumn	The column containing sample group labels
foreground	The foreground group of samples for differential comparison
background	The background group of samples for differential comparison
verbose	Set TRUE to display additional messages. Default is FALSE.

Value

plot object

GRangesToString	GRangesToString <i>Converts a GRanges object to a string in the format 'chr1:100-200'</i>
-----------------	---

Description

GRangesToString Turns a GRanges Object into a list of strings in the format chr1:100-200

Usage

```
GRangesToString(GR_obj)
```

Arguments

GR_obj the GRanges object to convert to a string

Value

A string or list of strings in the format 'chr1:100-200' representing ranges in the input GRanges

mergeTileResults	mergeTileResults
------------------	------------------

Description

mergeTileResults merges a list of tileResults that each contain unique samples into a single object encompassing all samples. Only cell populations shared among all input tileResults will be retained. This function can merge MultiAssayExperiment objects from callOpenTiles that are created with the same TxDb, OrgDb, and Genome assembly.

Usage

```
mergeTileResults(tileResultsList, numCores = 1, verbose = TRUE)
```

Arguments

tileResultsList	List of MultiAssayExperiments objects returned by callOpenTiles containing containing peak calling results.
numCores	Optional, the number of cores to use with multiprocessing. Default is 1.
verbose	Set TRUE to display additional messages. Default is FALSE.

Value

tileResults a single MultiAssayExperiment containing a sample-tile intensity matrix for each sample and common cell population in the input tileResultsList.

Examples

```
## Not run:  
# Depends on local MOCHA tileResults  
MOCHA::mergeTileResults(  
  list(tileResultsCelltypesABC, tileResultsCelltypesBCD)  
)  
  
## End(Not run)
```

MotifEnrichment	MotifEnrichment
-----------------	-----------------

Description

Test for enrichment of motifs within Group1 against a background Group2 using a hypergeometric t-test.

Usage

```
MotifEnrichment(Group1, Group2, motifPosList, type = NULL)
```

Arguments

Group1	A GRanges object, such as a set of significant differential tiles.
Group2	A GRanges object containing background regions, non-overlapping with Group1
motifPosList	A GRangesList of motifs and positions for each motif. Must be named for each motif.
type	Optional, name of a metadata column in Group1 and Group2 to test for enrichment the number of unique entries in column given by 'type'. Default is NULL, which tests the number of Ranges.

Value

A data.frame containing enrichment for each group

MotifSetEnrichmentAnalysis	MotifSetEnrichmentAnalysis
----------------------------	----------------------------

Description

This analogous to Gene Set Enrichment Analysis. Instead of testing for enrichment of a geneset with a given gene set in a pathway, we are testing the enrichment of a given TF motif set against a motif set downstream of a multiple ligands. If there is enrichment, it's a sign that that ligand could drive that set of motifs.

Usage

```
MotifSetEnrichmentAnalysis(
  ligandTFMatrix,
  motifEnrichmentDF,
  motifColumn,
  ligands,
  statColumn,
```

```

    statThreshold,
    annotationName = "CellType",
    annotation = "none",
    numCores = 1,
    verbose = FALSE
  )

```

Arguments

ligandTFMatrix	NicheNet Ligand-TF matrix
motifEnrichmentDF	Dataframe (unfiltered) from ArchR's peakAnnoEnrich step. Expected to have a column with motif names, and a column with the -log10 adjusted p-values.
motifColumn	Column name within the motifEnrichmentDF that has motif names.
ligands	Vector of ligands to test
statColumn	Column name in motifEnrichmentDF containing the statistic to test
statThreshold	Significance threshold used to select significant motif set
annotationName	Optional column name for the annotation. Default is "CellType".
annotation	Optional annotation value added to all rows of the output motif dataframe. Can be character vector or numeric. Default is "none".
numCores	The number of cores to use with multiprocessing. Default is 1.
verbose	Set TRUE to display additional messages. Default is FALSE.

Value

specDF A dataframe containing enrichment analysis results

packMOCHA

packMOCHA

Description

packMOCHA combines a MOCHA object (Sample-Tile Matrix or tileResults) with its saved coverage tracks into a single zip archive. This allows MOCHA objects and the necessary coverage files for plotting to be shared to other file systems. See also: [unpackMOCHA](#)

Usage

```
packMOCHA(MOCHAObj, zipfile, verbose = FALSE)
```

Arguments

MOCHAObj	A MultiAssayExperiment or RangedSummarizedExperiment, from MOCHA
zipfile	Filename and path of the zip archive.
verbose	Set TRUE to display additional messages. Default is FALSE.

Value

zipfile Path to zip archive.

Examples

```
## Not run:
# Depends on and manipulates files on filesystem
myOutputDir <- "/home/documents/MOCHA_out"
zipPath <- MOCHA::packMOCHA(
  tileResults, zipfile = file.path(myOutputDir, "testzip.zip")
)

## End(Not run)
```

pilotLMEM

Execute a pilot run of single linear model on a subset of data

Description

`'r lifecycle::badge("deprecated")'` This function is deprecated - improved modeling functions can be found in the package "ChAI" at <https://github.com/aifimmunology/ChAI> pilotLMEM Runs linear mixed-effects modeling for continuous, non-zero inflated data using [lmer](#)

Usage

```
pilotLMEM(
  ExperimentObj,
  assayName,
  modelFormula,
  pilotIndices = 1:10,
  verbose = FALSE
)
```

Arguments

ExperimentObj	A SummarizedExperiment-type object generated from chromVAR, makePseudobulkRNA, or other. Objects from getSampleTileMatrix can work, but we recommend runZIGLMM for those objects, not runLMEM>
assayName	a character string, matching the name of an assay within the SummarizedExperiment. The assay named will be used for modeling.
modelFormula	The formula to use with lmerTest::lmer, in the format (exp ~ factors). All factors must be found in column names of the ExperimentObj metadata.
pilotIndices	A vector of integers defining the subset of the ExperimentObj matrix. Default is 1:10.
verbose	Set TRUE to display additional messages. Default is FALSE.

Value

modelList a list of outputs from lmerTest::lmer

pilotZIGLMM	<i>Execute a pilot run of model on a subset of data</i>
-------------	---

Description

`'r lifecycle::badge("deprecated")'` This function is deprecated - improved modeling functions can be found in the package "ChAI" at <https://github.com/aifimmunology/ChAI> pilotLMEM Runs linear mixed-effects modeling for zero inflated data using [glmTMB](#). TryCatch will catch errors, and return the error and dataframe for troubleshooting.

Usage

```
pilotZIGLMM(
  TSAM_Object,
  cellPopulation = NULL,
  continuousFormula = NULL,
  ziformula = NULL,
  zi_threshold = 0,
  verbose = FALSE,
  pilotIndices = 1:10
)
```

Arguments

TSAM_Object	A SummarizedExperiment object generated from getSampleTileMatrix, chromVAR, or other.
cellPopulation	A single cell population on which to run this pilot model
continuousFormula	The formula, see glmTMB . Combined fixed and random effects formula, following lme4 syntax.
ziformula	The zero-inflated formula, see glmTMB . a one-sided (i.e., no response variable) formula for zero-inflation combining fixed and random effects: the default <code>~0</code> specifies no zero-inflation. Specifying <code>~.</code> sets the zero-inflation formula identical to the right-hand side of formula (i.e., the conditional effects formula); terms can also be added or subtracted. When using <code>~.</code> as the zero-inflation formula in models where the conditional effects formula contains an offset term, the offset term will automatically be dropped. The zero-inflation model uses a logit link.
zi_threshold	Zero-inflated threshold (range = 0-1), representing the fraction of samples with zeros. When the percentage of zeros in the tile is between 0 and <code>zi_threshold</code> , samples with zeroes are dropped and only the continuous formula is used. Use this parameter at your own risk. Default is 0.
verbose	Set TRUE to display additional messages. Default is FALSE.
pilotIndices	A vector of integers defining the subset of the ExperimentObj matrix. Default is 1:10.

Value

modelList a list of outputs from glmmTMB::glmmTMB

plotConsensus	plotConsensus
---------------	---------------

Description

plotConsensus Extracts the peak reproducibility and generates a heuristic plots that can be used to determine the reproducibility threshold used within getSampleTileMatrix.

Usage

```
plotConsensus(
  tileObject,
  cellPopulations = "All",
  groupColumn = NULL,
  returnPlotList = FALSE,
  returnDFs = FALSE,
  numCores = 1
)
```

Arguments

tileObject	A MultiAssayExperiment object from callOpenTiles,
cellPopulations	the cell populations you want to visualize.
groupColumn	Optional parameter, same as in getSampleTileMatrix, which defines whether you want to plot reproducibility within each
returnPlotList	Instead of one plot with all celltypes/conditions, it returns a list of plots for each cell types
returnDFs	Instead of a plot, returns a data.frame of the reproducibility across samples. If set to false, then it plots the data.frame instead of returning it.
numCores	Number of cores to multithread over.

Value

SampleTileObj the input data structure with added gene annotations.

```
plotIntensityDistribution
      plotIntensityDistribution
```

Description

`plotIntensityDistribution` Plots the distribution of sample-tile intensities for a give cell type

`plotIntensityDistribution` Plots the distribution of sample-tile intensities for a give cell type

Usage

```
plotIntensityDistribution(
  TSAM_object,
  cellPopulation,
  returnDF = FALSE,
  density = TRUE
)
```

```
plotIntensityDistribution(
  TSAM_object,
  cellPopulation,
  returnDF = FALSE,
  density = TRUE
)
```

Arguments

<code>TSAM_object</code>	SummarizedExperiment from <code>getSampleTileMatrix</code>
<code>cellPopulation</code>	Cell type names (assay name) within the <code>TSAM_object</code>
<code>returnDF</code>	If TRUE, return the data frame without plotting. Default is FALSE.
<code>density</code>	Boolean to determine whether to plot density or histogram. Default is TRUE (plots density).

Value

data.frame or ggplot histogram.

data.frame or ggplot histogram.

plotRegion	plotRegion
------------	------------

Description

plotRegion Plots the region that you've summarized across all cell groupings (groups=initial getPopFrag(s) split) with optional motif overlay, chromosome position ideogram, and additional GRanges tracks. If plotting motif overlay, ensure that motif annotations have been added to your counts SummarizedExperiment. A basic plot can be rendered with just a counts SummarizedExperiment, but additional formatting arguments allow for further customization. Note that to show specific genes with the option 'whichGenes' the **RMariaDB** package must be installed.

Usage

```
plotRegion(
  countSE,
  plotType = "area",
  base_size = 12,
  counts_color = NULL,
  range_label_size = 2,
  legend.position = NULL,
  legendRatio = 0.25,
  facet_label_side = "top",
  counts_color_var = "Groups",
  counts_group_colors = NULL,
  counts_theme_ls = NULL,
  motifSetName = NULL,
  motif_y_space_factor = 4,
  motif_stagger_labels_y = FALSE,
  motif_weights = NULL,
  motif_weight_name = "Motif Weight",
  motif_weight_colors = c(darkblue = -10, gray = 0, darkred = 10),
  motif_lab_size = 1,
  motif_lab_alpha = 0.25,
  motif_line_alpha = 0.25,
  motif_line_size = 0.75,
  showGene = TRUE,
  whichGenes = NULL,
  monotoneGenes = FALSE,
  db_id_col = "REFSEQ",
  collapseGenes = FALSE,
  gene_theme_ls = NULL,
  additionalGRangesTrack = NULL,
  linkdf = NULL,
  showIdeogram = TRUE,
  ideogram_genome = "hg19",
  relativeHeights = c(Chr = 0.9, `Normalized Counts` = 7, Links = 1.5, Genes = 2,
```

```

    AdditionalGRanges = 4.5),
    verbose = FALSE
  )

```

Arguments

countSE	A SummarizedExperiment from MOCHA::getCoverage
plotType	Options include 'overlaid', 'area', 'line', or 'RidgePlot'. default is 'area', which will plot a separate track for each group with the area filled in under the curve. Setting plotType to 'overlaid' will overlay count plot histograms across samples, instead of faceting out separately. Setting plotType to 'RidgePlot' will generate a RidgePlot across all groups.
base_size	Numeric, default 12. Global plot base text size parameter
counts_color	Optional color palette. A named vector of color values where names are unique values in the 'color_var' column
range_label_size	Numeric value, default 4. Text size for the y-axis range label
legend.position	Any acceptable 'legend.position' argument to theme(). Default NULL will place legend for overlaid plots at (0.8,0.8), or to the "right" for faceted plots.
legendRatio	Ratio of width or height of the main plot to the legend. Useful if the legend is to large. If only used when legend.position is set to top, bottom, left, or right.
facet_label_side	Direction character value, default "top". Can also be "right", "left", or "bottom". Position of facet label.
counts_color_var	Character value, default "Groups". Column name from countdf to use to color counts plots. Only used if counts_group_colors provided
counts_group_colors	Optional named color vector. Values as colors, names are levels of 'counts_color_var'. If provided, will color the plots specifically using 'scale_color_manual()'
counts_theme_ls	A list of named theme arguments passed to theme(). For example, 'list(axis.ticks = element_blank())'. Default NULL will use 'counts_plot_default_theme'.
motifSetName	The name of the motif set in ArchRProj to use for annotation. Example: 'JasparMotifs'
motif_y_space_factor	A factor for vertical spacing between motif labels. Default 4. Increase to make labels farther apart, decrease to make labels closer.
motif_stagger_labels_y	= FALSE Logical value, default FALSE. If TRUE, will stagger motif labels in adjacent columns in the vertical direction
motif_weights	Optional numeric vector, default NULL. If provided will be used to color motif labels by the weighted values

motif_weight_name	Character value, default "Motif Weight". Used to label the legend for motif colors
motif_weight_colors	Named numeric vector. Names should be color values and breaks should be the corresponding values of motif_weights. Values outside the highest and lowest value will appear as max or min defined color value.
motif_lab_size	Numeric value, default 1. Size of motif labels.
motif_lab_alpha	Numeric value, default 0.25. Alpha for motif labels.
motif_line_alpha	Numeric value, default 0.25. Alpha for motif lines.
motif_line_size	Numeric value, default 1. Size of motif lines.
showGene	Logical value, default TRUE. Whether or not the gene track should be plotted.
whichGenes	Name of gene for plotting this specific gene in region.
monotoneGenes	Boolean. Determines whether to color-code genes by gene name, or to set them all to dark gray.
db_id_col	Character value. Column in 'OrgDb' containing the output id for 'whichGenes' plotting. Default "REFSEQ".
collapseGenes	Options include 'collapseAll', 'longestTx', or 'None' Default 'None' will plot the expanded view of the reference genes, 'collapseAll' if you want collapse the gene tracks into one, and 'longestTx' will only plot the longest transcript of each gene.
gene_theme_ls	Named list of parameters passed to 'theme()' for the gene plot. Default NULL will use '.gene_plot_theme'
additionalGRangesTrack	A GRanges object containing additional track plot data
linkdf	A dataframe with co-accessible links to display as an additional track
showIdeogram	Logical value, default TRUE. If TRUE plots the chromosome ideogram at the top of the multi-track plot
ideogram_genome	Character value, a genome name for the ideogram plot. Default 'hg19'.
relativeHeights	Named numeric vector of relative heights for each of the 4 track plots to enable clean visualization when there are many tracks. Unused tracks will be ignored. Default value = c('Chr' = 0.9, 'Normalized Counts' = 7, 'Genes' = 2, 'AdditionalGRanges' = 4.5)
verbose	Set TRUE to display additional messages. Default is FALSE.

Value

The input ggplot object with motif labels overlaid

Examples

```
## Not run:
# my_count_SE is a counts data frame generated by extractRegion()

# Simple counts + ideogram + all genes:
plotRegion(countSE = my_count_SE)

# Motif overlay for a project my_proj containing "JasparMotifs" annotations:
plotRegion(
  countSE = my_count_SE, motifSetName = "JasparMotifs",
  motif_lab_alpha = 1, motif_line_alpha = 1
)

# Motif overlay w/ weights:
plotRegion(
  countSE = my_count_SE, motifSetName = "JasparMotifs", motif_lab_alpha = 1,
  motif_line_alpha = 1, motif_weights = my_enrichment_weights
)

## End(Not run)
```

renameCellTypes	renameCellTypes
-----------------	-----------------

Description

renameCellTypes Allows you to modify the cell type names for a MOCHA SampleTileObject, from the assay names, GRanges column names, and summarizedData (within the metadata), all at once.

Usage

```
renameCellTypes(MOCHAObject, oldNames, newNames)
```

Arguments

MOCHAObject	A RangedSummarizedExperiment,
oldNames	A list of cell type names that you want to change.
newNames	A list of new cell type names to replace the old names with.

Value

A MOCHA SampleTile object with new cell types.

runLMEM	<i>Run Linear Mixed-Effects Modeling for continuous, non-zero inflated data</i>
---------	---

Description

runLMEM Runs linear mixed-effects modeling for continuous, non-zero inflated data using [lmer](#)

Usage

```
runLMEM(  
  ExperimentObj,  
  assayName,  
  modelFormula,  
  initialSampling = 5,  
  verbose = FALSE,  
  numCores = 2  
)
```

Arguments

ExperimentObj	A SummarizedExperiment object generated from <code>getSampleTileMatrix</code> , <code>chromVAR</code> , or other. It is expected to contain only one assay, or only the first assay will be used for the model. Data should not be zero-inflated.
assayName	The name of the assay to model within the SummarizedExperiment.
modelFormula	The formula to use with <code>lmerTest::lmer</code> , in the format (<code>exp ~ factors</code>). All factors must be found in column names of the ExperimentObj metadata. modelFormula must start with 'exp' as the response. See lmer .
initialSampling	Size of data to use for pilot
verbose	Set TRUE to display additional messages. Default is FALSE.
numCores	integer. Number of cores to parallelize across.

Value

results a SummarizedExperiment containing LMEM results. Assays are metrics related to the model coefficients, including the Estimate, Std_Error, df, t_value, p_value. Within each assay, each row corresponds to each row of the SummarizedExperiment and columns correspond to each fixed effect variable within the model. Any row metadata from the ExperimentObject (see `rowData(ExperimentObj)`) is preserved in the output. The Residual matrix and the variance of the random effects are saved in the metadata slot of the output.

Examples

```
## Not run:
modelList <- runLMEM(ExperimentObj,
  assayName = names(ExperimentObj)[[1]]
  modelFormula = NULL,
  initialSampling = 5,
  verbose = FALSE,
  numCores = 1
)

## End(Not run)
```

runZIGLMM

Run Zero-inflated Generalized Linear Mixed Modeling on pseudobulked scATAC data

Description

`'r lifecycle::badge("deprecated")'` This function is deprecated - improved modeling functions can be found in the package "ChAI" at <https://github.com/aifimmunology/ChAI> runZIGLMM Runs linear mixed-effects modeling for zero-inflated data using [glmmTMB](#).

Usage

```
runZIGLMM(
  TSAM_Object,
  cellPopulation = "all",
  continuousFormula = NULL,
  ziformula = NULL,
  zi_threshold = 0,
  initialSampling = 5,
  verbose = FALSE,
  numCores = 1
)
```

Arguments

TSAM_Object A SummarizedExperiment object generated from `getSampleTileMatrix`.

cellPopulation Name of a cell type(s), or 'all'. The function will combine the cell types mentioned into one matrix before running the model.

continuousFormula The formula for the continuous data that should be used within `glmmTMB`. It should be in the format `(exp ~ factors)`. All factors must be found in column names of the `TSAM_Object` metadata, except for `CellType`, `FragNumber` and `CellCount`, which will be extracted from the `TSAM_Object`. `modelFormula` must start with 'exp' as the response. See [glmmTMB](#).

ziformula	The formula for the zero-inflated data that should be used within glmmTMB. It should be in the format (~ factors). All factors must be found in column names of the TSAM_Object colData metadata, except for CellType, FragNumber and CellCount, which will be extracted from the TSAM_Object.
zi_threshold	Zero-inflated threshold (range = 0-1), representing the fraction of samples with zeros. When the percentage of zeros in the tile is between 0 and zi_threshold, samples with zeroes are dropped and only the continuous formula is used. Use this parameter at your own risk. Default is 0.
initialSampling	Size of data to use for pilot
verbose	Set TRUE to display additional messages. Default is FALSE.
numCores	integer. Number of cores to parallelize across.

Value

results a SummarizedExperiment containing LMEM results

Examples

```
## Not run:
modelList <- runZIGLMM(STM[c(1:1000)], [,
  cellPopulation = "CD16 Mono",
  continuousFormula = exp ~ Age + Sex + days_since_symptoms + (1 | PTID),
  ziformula = ~ FragNumber + Age,
  verbose = TRUE,
  numCores = 35
)

## End(Not run)
```

StringsToGRanges	StringsToGRanges
------------------	------------------

Description

StringsToGRanges Turns a list of strings in the format chr1:100-200 into a GRanges object

Usage

```
StringsToGRanges(regionString)
```

Arguments

regionString A string or list of strings each in the format chr1:100-200

Value

a GRanges object with ranges representing the input string(s)

```
subsetMOCHAObject    subsetMOCHAObject
```

Description

subsetMOCHAObject subsets a tileResults-type object (from callOpenTiles), or a SummarizedExperiment-type object (from getSampleTileMatrix), either by cell type or sample metadata.

Usage

```
subsetMOCHAObject(
  Object,
  subsetBy,
  groupList,
  removeNA = TRUE,
  subsetPeaks = TRUE,
  verbose = FALSE
)
```

Arguments

Object	A MultiAssayExperiment or RangedSummarizedExperiment,
subsetBy	The variable to subset by. Can either be 'celltype', or a column from the sample metadata (see 'colData(Object)').
groupList	the list of cell type names or sample-associated data that should be used to subset the Object
removeNA	If TRUE, removes groups in groupList that are NA. If FALSE, keep groups that are NA.
subsetPeaks	If 'subsetBy' = 'celltype', subset the tile set to tiles only called in those cell types. Default is TRUE.
verbose	Set TRUE to display additional messages. Default is FALSE.

Value

Object the input Object, filtered down to either the cell type or samples desired.

```
testCoAccessibility testCoAccessibility
```

Description

testCoAccessibility takes an input set of tile pairs and tests whether they are significantly different compared to random, non-overlapping background set.

Usage

```
testCoAccessibility(
  SampleTileObj,
  tile1,
  tile2,
  numCores = 1,
  ZI = TRUE,
  backNumber = 1000,
  calcPValue = TRUE,
  returnBackGround = FALSE,
  verbose = TRUE
)
```

Arguments

SampleTileObj	The SummarizedExperiment object output from getSampleTileMatrix containing your sample-tile matrices
tile1	vector of indices or tile names (chrX:100-2000) for tile pairs to test (first tile in each pair)
tile2	vector of indices or tile names (chrX:100-2000) for tile pairs to test (second tile in each pair)
numCores	Optional, the number of cores to use with multiprocessing. Default is 1.
ZI	boolean flag that enables zero-inflated (ZI) Spearman correlations to be used. Default is TRUE. If FALSE, skip zero-inflation and calculate the normal Spearman.
backNumber	number of background pairs. Default is 1000.
calcPValue	Boolean, if TRUE calculate p-values. Default is TRUE.
returnBackGround	Boolean, if TRUE return the background correlations as well as foreground. Default is FALSE.
verbose	Set TRUE to display additional messages. Default is FALSE.

Value

foreGround A data.frame with Tile1, Tile2, Correlation, and p-value for that correlation compared to the background

unpackMOCHA	unpackMOCHA
-------------	-------------

Description

unpackMOCHA will unpack a zip archive created by [unpackMOCHA](#), setting the stored MOCHA object's stored directory path to the new location. See also: [packMOCHA](#)

Usage

```
unpackMOCHA(zipfile, exdir, verbose = FALSE)
```

Arguments

zipfile	Filepath to the packed MOCHA object.
exdir	The path to the external directory where you want to unpack the MOCHA object.
verbose	Display additional messages. Default is FALSE.

Value

MOCHAObj the MOCHA object (tileResults or Sample-Tile Matrix)

Examples

```
## Not run:
# Depends on files existing on your system
MOCHA::unpackMOCHA(zipfile = "./mochaobj.zip", exdir = "./newMOCHAdir")

## End(Not run)
```

varZIGLMM	<i>Zero-inflated Variance Decomposition for pseudobulked scATAC data</i>
-----------	--

Description

`'r lifecycle::badge("deprecated")'` This function is deprecated - improved modeling functions can be found in the package "ChAI" at <https://github.com/aifimmunology/ChAI> varZIGLMM Identified variance decomposition on a given cell type across both zero-inflated and continuous space using a zero-inflated general linear mixed model [glmTMB](#)

Usage

```
varZIGLMM(
  TSAM_Object,
  cellPopulation = NULL,
  continuousRandom = NULL,
  ziRandom = NULL,
  zi_threshold = 0.1,
  verbose = FALSE,
  numCores = 1
)
```

Arguments

TSAM_Object	A SummarizedExperiment object generated from <code>getSampleTileMatrix</code> .
cellPopulation	Name of a cell type(s), or 'all'. The function will combine the cell types mentioned into one matrix before running the model.
continuousRandom	Random effects to test in the continuous portion. All factors must be found in column names of the TSAM_Object metadata, except for <code>FragNumber</code> and <code>CellCount</code> , which will be extracted from the TSAM_Object's metadata.
ziRandom	Random effects to test in the zero-inflated portion. All factors must be found in column names of the TSAM_Object <code>colData</code> metadata, except for <code>FragNumber</code> and <code>CellCount</code> , which will be extracted from the TSAM_Object's metadata.
zi_threshold	Zero-inflated threshold (range = 0-1), representing the fraction of samples with zeros. When the percentage of zeros in the tile is between 0 and <code>zi_threshold</code> , samples with zeroes are dropped and only the continuous formula is used. Use this parameter at your own risk. Default is 0.
verbose	Set TRUE to display additional messages. Default is FALSE.
numCores	integer. Number of cores to parallelize across.

Value

results a SummarizedExperiment containing results from ZIGLMM (Fixed effect estimates, P-values, and Std Error)

Examples

```
## Not run:
modelList <- runZIGLMM(STM[c(1:1000)], [,
  cellPopulation = "CD16 Mono",
  continuousRandom = c("Age", "Sex", "Days"),
  ziRandom = c("FragNumber", "Days"),
  verbose = TRUE,
  numCores = 35
)

## End(Not run)
```

<code>youden_threshold</code>	<i>youden_threshold</i>
-------------------------------	-------------------------

Description

Trained regression model for predicting a cutoff threshold for peak calling. Call: `loess(formula = OptimalCutpoint ~ Ncells, data = thresh_df)`

Usage

`youden_threshold`

Format

A list of 18 regression variables

Details

Number of Observations: 27 Equivalent Number of Parameters: 5.98 Residual Standard Error: 0.02121