Package 'CRMetrics'

September 1, 2023

Title Cell Ranger Output Filtering and Metrics Visualization

Version 0.3.0

Description Sample and cell filtering as well as visualisation of output metrics from 'Cell Ranger' by Grace X.Y. Zheng et al. (2017) <doi:10.1038/ncomms14049>. 'CR-Metrics' allows for easy plotting of output metrics across multiple samples as well as comparative plots including statistical assessments of these. 'CRMetrics' allows for easy removal of ambient RNA using 'SoupX' by Matthew D Young and Sam Behjati (2020) <doi:10.1093/gigascience/giaa151> or 'CellBender' by Stephen J Fleming et al. (2022) <doi:10.1101/791699>. Furthermore, it is possible to preprocess data using 'Pagoda2' by Nikolas Barkas et al. (2021) <https://github.com/kharchenkolab/pagoda2> or 'Seurat' by Yuhan Hao et al. (2021) <doi:10.1016/j.cell.2021.04.048> followed by embedding of cells using 'Conos' by Nikolas Barkas et al. (2019) <doi:10.1038/s41592-019-0466-z>. Finally, doublets can be detected using 'scrublet' by Samuel L. Wolock et al. (2019) <doi:10.1016/j.cels.2018.11.005> or 'Doublet-Detection' by Gayoso et al. (2020) <doi:10.5281/zenodo.2678041>. In the end, cells are fil-

License GPL-3 **Encoding** UTF-8 **Depends** R (>= 4.0.0)

biocViews

Imports cowplot, dplyr, ggbeeswarm, ggplot2, ggpmisc, ggpubr, ggrepel, magrittr, Matrix, methods, R6, scales, sccore, sparseMatrixStats, stats, tibble, tidyr, utils

tered based on user input for use in downstream applications.

Suggests conos, data.table, markdown, pagoda2, reticulate, rhdf5, Seurat, SoupX, testthat (>= 3.0.0)

RoxygenNote 7.2.3

URL https://github.com/khodosevichlab/CRMetrics

BugReports https://github.com/khodosevichlab/CRMetrics/issues

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Config/testthat/edition 3

NeedsCompilation no Author Rasmus Rydbirk [aut, cre], Fabienne Kick [aut], Henrietta Holze [aut], Xian Xin [ctb] Repository CRAN Date/Publication 2023-09-01 09:00:06 UTC R topics documented:		
Inde	X	43
CR	RMetrics	CRMetrics class object
]	•	e Cell Ranger count data. To initialize a new object, 'data.path' or 'cms' is s also recommended, but not required.
ı	Metadata mu	ne or character Path to metadata file or name of metadata data.frame object. contain a column named 'sample' containing sample names that must match a 'data.path' (default = NULL)
(er $Path(s)$ to Cell Ranger count data, one directory per sample. If multiple paths, path2") (default = $NULL$)
		unt matrices (default = NULL) list List with preprocessed count matrices after \$doPreprocessing() (default =
	NULL)	ith raw, unfiltered count matrices, i.e., including all CBs detected also empty
,		data.frame Summary metrics from Cell Ranger (default = NULL)
	-	data.frame Detailed metrics, i.e., no. genes and UMIs per cell (default =
(ter A group present in the metadata to compare the metrics by, can be added arison (default = NULL)

n.cores numeric Number of cores for calculations (default = 1) Initialize a CRMetrics object

verbose logical Print messages or not (default = TRUE)

 $\label{eq:continuous_problem} \begin{tabular}{ll} theme & ggplot2 & theme & (default: theme_bw()) \\ pal & Plotting & palette & (default = NULL) \\ \end{tabular}$

Methods

Public methods:

```
• CRMetrics$new()
• CRMetrics$addDetailedMetrics()
• CRMetrics$addComparison()
• CRMetrics$plotSamples()
• CRMetrics$plotSummaryMetrics()
• CRMetrics$plotDetailedMetrics()
• CRMetrics$plotEmbedding()
• CRMetrics$plotDepth()
• CRMetrics$plotMitoFraction()
• CRMetrics$detectDoublets()
• CRMetrics$doPreprocessing()
• CRMetrics$createEmbedding()
• CRMetrics$filterCms()
• CRMetrics$selectMetrics()
• CRMetrics$plotFilteredCells()
• CRMetrics$getDepth()
• CRMetrics$getMitoFraction()
• CRMetrics$prepareCellbender()
• CRMetrics$saveCellbenderScript()
• CRMetrics$getExpectedCells()
• CRMetrics$getTotalDroplets()
• CRMetrics$addCms()
• CRMetrics$plotCbTraining()
• CRMetrics$plotCbCellProbs()
• CRMetrics$plotCbAmbExp()
• CRMetrics$plotCbAmbGenes()
• CRMetrics$addSummaryFromCms()
• CRMetrics$runSoupX()
• CRMetrics$plotSoupX()
```

Method new(): To initialize new object, 'data.path' or 'cms' is needed. 'metadata' is also recommended, but not required.

```
Usage:
CRMetrics$new(
  data.path = NULL,
 metadata = NULL,
  cms = NULL,
  samples = NULL,
```

• CRMetrics\$clone()

• CRMetrics\$plotCbCells() • CRMetrics\$addDoublets()

```
unique.names = TRUE,
    sep.cells = "!!",
    comp.group = NULL,
    verbose = TRUE,
    theme = theme_bw(),
    n.cores = 1,
    sep.meta = ",",
    raw.meta = FALSE,
    pal = NULL
 )
 Arguments:
 data.path character Path to directory with Cell Ranger count data, one directory per sample
     (default = NULL).
 metadata data.frame or character Path to metadata file (comma-separated) or name of meta-
     data dataframe object. Metadata must contain a column named 'sample' containing sample
     names that must match folder names in 'data.path' (default = NULL)
 cms list List with count matrices (default = NULL)
 samples character Sample names. Only relevant is cms is provided (default = NULL)
 unique.names logical Create unique cell names. Only relevant if cms is provided (default =
     TRUE)
 sep.cells character Sample-cell separator. Only relevant if cms is provided and unique.names=TRUE
     (default = "!!")
 comp.group character A group present in the metadata to compare the metrics by, can be added
     with addComparison (default = NULL)
 verbose logical Print messages or not (default = TRUE)
 theme ggplot2 theme (default: theme_bw())
 n.cores integer Number of cores for the calculations (default = self$n.cores)
 sep.meta character Separator for metadata file (default = ",")
 raw.meta logical Keep metadata in its raw format. If FALSE, classes will be converted using
     "type.convert" (default = FALSE)
 pal character Plotting palette (default = NULL)
 Returns: CRMetrics object
 Examples:
 \dontrun{
 crm <- CRMetrics$new(data.path = "/path/to/count/data/")</pre>
Method addDetailedMetrics(): Function to read in detailed metrics. This is not done upon
initialization for speed.
 CRMetrics$addDetailedMetrics(
    cms = self$cms,
   min.transcripts.per.cell = 100,
   n.cores = self$n.cores,
    verbose = self$verbose
 )
```

```
Arguments:
 cms list List of (sparse) count matrices (default = self$cms)
 min.transcripts.per.cell numeric Minimal number of transcripts per cell (default = 100)
 n.cores integer Number of cores for the calculations (default = self$n.cores).
 verbose logical Print messages or not (default = self$verbose).
 Returns: Count matrices
 Examples:
 # Simulate data
 testdata.cms <- lapply(seq_len(2), \(x) {
 out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)</pre>
 out[out < 0] <- 1
 dimnames(out) <- list(sapply(seq_len(2e3), \(x) paste0("gene",x)),</pre>
 sapply(seq_len(1e3), \(x) paste0("cell",x)))
 return(out)
 })
 # Initialize
 crm <- CRMetrics$new(cms = testdata.cms, samples = c("sample1", "sample2"), n.cores = 1)</pre>
 # Run function
 crm$addDetailedMetrics()
Method addComparison(): Add comparison group for statistical testing.
 Usage:
 CRMetrics$addComparison(comp.group, metadata = self$metadata)
 Arguments:
 comp.group character Comparison metric (default = self$comp.group).
 metadata data.frame Metadata for samples (default = self$metadata).
 Returns: Vector
 Examples:
 # Simulate data
 testdata.cms <- lapply(seq_len(2), \(x) {
 out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)</pre>
 out[out < 0] <- 1
 dimnames(out) \leftarrow list(sapply(seq_len(2e3), \xspace ("gene",x)),
 sapply(seq_len(1e3), \(x) paste0("cell",x)))
 return(out)
 })
 # Initialize
 crm <- CRMetrics$new(cms = testdata.cms, samples = c("sample1", "sample2"), n.cores = 1)</pre>
 # Add metadata
 crm$metadata <- data.frame(sex = c("male", "female"))</pre>
```

```
# Add comparison group
 crm$addComparison(comp.group = "sex")
Method plotSamples(): Plot the number of samples.
 Usage:
 CRMetrics$plotSamples(
   comp.group = self$comp.group,
   h.adj = 0.05,
   exact = FALSE,
   metadata = self$metadata,
   second.comp.group = NULL,
   pal = self$pal
 )
 Arguments:
 comp.group character Comparison metric, must match a column name of metadata (default =
     self$comp.group).
 h.adj numeric Position of statistics test p value as \% of max(y) (default = 0.05).
 exact logical Whether to calculate exact p values (default = FALSE).
 metadata data.frame Metadata for samples (default = self$metadata).
 second.comp.group character Second comparison metric, must match a column name of meta-
     data (default = NULL).
 pal character Plotting palette (default = self$pal)
 Returns: ggplot2 object
 Examples:
 samples <- c("sample1", "sample2")</pre>
 # Simulate data
 testdata.cms <- lapply(seq_len(2), \(x) {
 out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)</pre>
 out[out < 0] <- 1
 dimnames(out) <- list(sapply(seq_len(2e3), \(x) paste0("gene",x)),</pre>
 sapply(seq_len(1e3), \(x) paste0("cell",x)))
 return(out)
 })
 names(testdata.cms) <- samples</pre>
 # Create metadata
 metadata <- data.frame(sample = samples,</pre>
 sex = c("male", "female"),
 condition = c("a","b"))
 # Initialize
 crm <- CRMetrics$new(cms = testdata.cms, metadata = metadata, n.cores = 1)</pre>
 # Plot
 crm$plotSamples(comp.group = "sex", second.comp.group = "condition")
```

```
Method plotSummaryMetrics(): Plot all summary stats or a selected list.
 CRMetrics$plotSummaryMetrics(
    comp.group = self$comp.group,
    second.comp.group = NULL,
   metrics = NULL,
   h.adj = 0.05,
    plot.stat = TRUE,
    stat.test = c("non-parametric", "parametric"),
    exact = FALSE,
   metadata = self$metadata,
    summary.metrics = self$summary.metrics,
    plot.geom = "bar",
    se = FALSE,
    group.reg.lines = FALSE,
    secondary.testing = TRUE,
    pal = self$pal
 Arguments:
 comp.group character Comparison metric (default = self$comp.group).
 second.comp.group character Second comparison metric, used for the metric "samples per
     group" or when "comp.group" is a numeric or an integer (default = NULL).
 metrics character Metrics to plot (default = NULL).
 h.adj numeric Position of statistics test p value as % of max(y) (default = 0.05)
 plot.stat logical Show statistics in plot. Will be FALSE if "comp.group" = "sample" or if
     "comp.group" is a numeric or an integer (default = TRUE)
 stat.test character Statistical test to perform to compare means. Can either be "non-parametric"
     or "parametric" (default = "non-parametric").
 exact logical Whether to calculate exact p values (default = FALSE).
 metadata data.frame Metadata for samples (default = self$metadata).
 summary.metrics data.frame Summary metrics (default = self$summary.metrics).
 plot.geom character Which geometric is used to plot the data (default = "point").
 se logical For regression lines, show SE (default = FALSE)
 group.reg.lines logical For regression lines, if FALSE show one line, if TRUE show line per
     group defined by second.comp.group (default = FALSE)
 secondary.testing logical Whether to show post hoc testing (default = TRUE)
 pal character Plotting palette (default = self$pal)
 Returns: ggplot2 object
 Examples:
 \donttest{
 # Simulate data
 testdata.cms <- lapply(seq_len(2), \(x) {</pre>
 out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)</pre>
 out[out < 0] <- 1
 dimnames(out) <- list(sapply(seq_len(2e3), \(x) paste0("gene",x)),</pre>
```

```
sapply(seq_len(1e3), \(x) paste0("cell",x)))
 return(out)
 })
 # Initialize
 crm <- CRMetrics$new(cms = testdata.cms, samples = c("sample1", "sample2"), n.cores = 1)</pre>
 # Add summary metrics
 crm$addSummaryFromCms()
 crm$plotSummaryMetrics(plot.geom = "point")
 }
Method plotDetailedMetrics(): Plot detailed metrics from the detailed.metrics object
 Usage:
 CRMetrics$plotDetailedMetrics(
    comp.group = self$comp.group,
   detailed.metrics = self$detailed.metrics,
   metadata = self$metadata,
   metrics = NULL,
   plot.geom = "violin",
   hline = TRUE,
   pal = self*pal
 )
 Arguments:
 comp.group character Comparison metric (default = self$comp.group).
 detailed.metrics data.frame Object containing the count matrices (default = self$detailed.metrics).
 metadata data.frame Metadata for samples (default = self$metadata).
 metrics character Metrics to plot. NULL plots both plots (default = NULL).
 plot.geom character How to plot the data (default = "violin").
 hline logical Whether to show median as horizontal line (default = TRUE)
 pal character Plotting palette (default = self$pal)
 data.path character Path to Cell Ranger count data (default = self$data.path).
 Returns: ggplot2 object
 Examples:
 \donttest{
 # Simulate data
 testdata.cms <- lapply(seq_len(2), \(x) {
 out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)</pre>
 out[out < 0] <- 1
 dimnames(out) <- list(sapply(seq_len(2e3), \(x) paste0("gene",x)),</pre>
 sapply(seq_len(1e3), \(x) paste0("cell",x)))
 return(out)
 })
 # Initialize
```

```
crm <- CRMetrics$new(cms = testdata.cms, samples = c("sample1", "sample2"), n.cores = 1)</pre>
 # Add detailed metrics
 crm$addDetailedMetrics()
 # Plot
 crm$plotDetailedMetrics()
Method plotEmbedding(): Plot cells in embedding using Conos and color by depth and dou-
blets.
 Usage:
 CRMetrics$plotEmbedding(
   depth = FALSE,
    doublet.method = NULL,
    doublet.scores = FALSE,
    depth.cutoff = 1000,
   mito.frac = FALSE,
   mito.cutoff = 0.05,
   species = c("human", "mouse"),
   size = 0.3,
   sep = "!!",
   pal = NULL,
 )
 Arguments:
 depth logical Plot depth or not (default = FALSE).
 doublet.method character Doublet detection method (default = NULL).
 doublet.scores logical Plot doublet scores or not (default = FALSE).
 depth.cutoff numeric Depth cutoff (default = 1e3).
 mito.frac logical Plot mitochondrial fraction or not (default = FALSE).
 mito.cutoff numeric Mitochondrial fraction cutoff (default = 0.05).
 species character Species to calculate the mitochondrial fraction for (default = c("human", "mouse")).
 size numeric Dot size (default = 0.3)
 sep character Separator for creating unique cell names (default = "!!")
 pal character Plotting palette (default = NULL)
 ... Plotting parameters passed to sccore::embeddingPlot.
 Returns: ggplot2 object
 Examples:
 \donttest{
 if (requireNamespace("pagoda2", quietly = TRUE)) {
 if (requireNamespace("conos", quietly = TRUE)) {
 # Simulate data
 testdata.cms <- lapply(seq_len(2), \(x) {
 out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)</pre>
```

```
out[out < 0] <- 1
 dimnames(out) \leftarrow list(sapply(seq_len(2e3), \xspace ("gene",x)),
 sapply(seq_len(1e3), \(x) paste0("cell",x)))
 return(out)
 })
 # Initialize
 crm <- CRMetrics$new(cms = testdata.cms, samples = c("sample1", "sample2"), n.cores = 1)</pre>
 # Create embedding
 crm$doPreprocessing()
 crm$createEmbedding()
 crm$plotEmbedding()
 } else {
 message("Package 'conos' not available.")
 }
 } else {
 message("Package 'pagoda2' not available.")
 }
 }
Method plotDepth(): Plot the sequencing depth in histogram.
 Usage:
 CRMetrics$plotDepth(
   cutoff = 1000,
   samples = self$metadata$sample,
    sep = "!!",
   keep.col = "#E7CDC2",
    filter.col = "#A65141"
 )
 Arguments:
 cutoff numeric The depth cutoff to color the cells in the embedding (default = 1e3).
 samples character Sample names to include for plotting (default = $metadata$sample).
 sep character Separator for creating unique cell names (default = "!!")
 keep.col character Color for density of cells that are kept (default = "#E7CDC2")
 filter.col Character Color for density of cells to be filtered (default = "#A65141")
 Returns: ggplot2 object
 Examples:
 \donttest{
 if (requireNamespace("pagoda2", quietly = TRUE)) {
 if (requireNamespace("conos", quietly = TRUE)) {
 # Simulate data
 testdata.cms <- lapply(seq_len(2), \(x) {
 out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)</pre>
 out[out < 0] <- 1
```

```
dimnames(out) <- list(sapply(seq_len(2e3), \(x) paste0("gene",x)),</pre>
 sapply(seq_len(1e3), \(x) paste0("cell",x)))
 return(out)
 })
 # Initialize
 crm <- CRMetrics$new(cms = testdata.cms, samples = c("sample1", "sample2"), n.cores = 1)</pre>
 # Create embedding
 crm$doPreprocessing()
 crm$createEmbedding()
 # Plot
 crm$plotDepth()
 } else {
 message("Package 'conos' not available.")
 }
 } else {
 message("Package 'pagoda2' not available.")
 }
 }
Method plotMitoFraction(): Plot the mitochondrial fraction in histogram.
 Usage:
 CRMetrics$plotMitoFraction(
   cutoff = 0.05,
   species = c("human", "mouse"),
   samples = self$metadata$sample,
   sep = "!!",
   keep.col = "#E7CDC2",
    filter.col = "#A65141"
 )
 Arguments:
 cutoff numeric The mito. fraction cutoff to color the embedding (default = 0.05)
 species character Species to calculate the mitochondrial fraction for (default = "human")
 samples character Sample names to include for plotting (default = $metadata$sample)
 sep character Separator for creating unique cell names (default = "!!")
 keep.col character Color for density of cells that are kept (default = "#E7CDC2")
 filter.col Character Color for density of cells to be filtered (default = "#A65141")
 Returns: ggplot2 object
 Examples:
 \donttest{
 if (requireNamespace("pagoda2", quietly = TRUE)) {
 if (requireNamespace("conos", quietly = TRUE)) {
 # Simulate data
 testdata.cms <- lapply(seq_len(2), \(x) {
```

```
out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)</pre>
 out[out < 0] <- 1
 dimnames(out) \leftarrow list(sapply(seq_len(2e3), \xspace ("gene", x)),
 sapply(seq_len(1e3), \(x) paste0("cell",x)))
 return(out)
 })
 # Initialize
 crm <- CRMetrics$new(cms = testdata.cms, samples = c("sample1", "sample2"), n.cores = 1)</pre>
 # Create embedding
 crm$doPreprocessing()
 crm$createEmbedding()
 # Plot
 crm$plotMitoFraction()
 } else {
 message("Package 'conos' not available.")
 }
 } else {
 message("Package 'pagoda2' not available.")
 }
 }
Method detectDoublets(): Detect doublet cells.
 Usage:
 CRMetrics$detectDoublets(
   method = c("scrublet", "doubletdetection"),
   cms = self$cms,
   samples = self$metadata$sample,
   env = "r-reticulate",
   conda.path = system("whereis conda"),
   n.cores = self$n.cores,
   verbose = self$verbose,
   args = list(),
   export = FALSE,
    data.path = self$data.path
 )
 Arguments:
 method character Which method to use, either scrublet or doubletdetection (default="scrublet").
 cms list List containing the count matrices (default=self$cms).
 samples character Vector of sample names. If NULL, samples are extracted from cms (default
     = self$metadata$sample)
 env character Environment to run python in (default="r-reticulate").
 conda.path character Path to conda environment (default=system("whereis conda")).
 n.cores integer Number of cores to use (default = self$n.cores)
 verbose logical Print messages or not (default = self$verbose)
```

```
args list A list with additional arguments for either DoubletDetection or scrublet. Please
     check the respective manuals.
 export boolean Export CMs in order to detect doublets outside R (default = FALSE)
 data.path character Path to write data, only relevant if export = TRUE. Last character must be
     / (default = self$data.path)
 Returns: data.frame
 Examples:
 \dontrun{
 # Simulate data
 testdata.cms <- lapply(seq_len(2), \(x) {
 out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)</pre>
 out[out < 0] <- 1
 dimnames(out) <- list(sapply(seq_len(2e3), \(x) paste0("gene",x)),</pre>
 sapply(seq_len(1e3), \(x) paste0("cell",x)))
 return(out)
 })
 # Initialize
 crm <- CRMetrics$new(cms = testdata.cms, samples = c("sample1", "sample2"), n.cores = 1)</pre>
 # Detect doublets
 crm$detectDoublets(method = "scrublet",
 conda.path = "/opt/software/miniconda/4.12.0/condabin/conda")
 }
Method doPreprocessing(): Perform conos preprocessing.
 Usage:
 CRMetrics$doPreprocessing(
   cms = self$cms,
   preprocess = c("pagoda2", "seurat"),
   min.transcripts.per.cell = 100,
   verbose = self$verbose,
   n.cores = self$n.cores,
   get.largevis = FALSE,
    tsne = FALSE,
   make.geneknn = FALSE,
   cluster = FALSE,
 )
 Arguments:
 cms list List containing the count matrices (default = self$cms).
 preprocess character Method to use for preprocessing (default = c("pagoda2", "seurat")).
 min.transcripts.per.cell numeric Minimal transcripts per cell (default = 100)
 verbose logical Print messages or not (default = self$verbose).
 n.cores integer Number of cores for the calculations (default = self$n.cores).
```

```
get.largevis logical For Pagoda2, create largeVis embedding (default = FALSE)
 tsne logical Create tSNE embedding (default = FALSE)
 make.geneknn logical For Pagoda2, estimate gene kNN (default = FALSE)
 cluster logical For Seurat, estimate clusters (default = FALSE)
 ... Additional arguments for Pagaoda2::basicP2Proc or conos:::basicSeuratProc
 Returns: Conos object
 Examples:
 \donttest{
 if (requireNamespace("pagoda2", quietly = TRUE)) {
 # Simulate data
 testdata.cms <- lapply(seq_len(2), \(x) {
 out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)</pre>
 out[out < 0] <- 1
 dimnames(out) <- list(sapply(seq_len(2e3), \(x) paste0("gene",x)),</pre>
 sapply(seq_len(1e3), \(x) paste0("cell",x)))
 return(out)
 })
 # Initialize
 crm <- CRMetrics$new(cms = testdata.cms, samples = c("sample1", "sample2"), n.cores = 1)</pre>
 # Perform preprocessing
 crm$doPreprocessing(preprocess = "pagoda2")
 message("Package 'pagoda2' not available.")
 }
 }
Method createEmbedding(): Create Conos embedding.
 Usage:
 CRMetrics$createEmbedding(
   cms = self$cms.preprocessed,
   verbose = self$verbose,
   n.cores = self$n.cores,
   arg.buildGraph = list(),
   arg.findCommunities = list(),
   arg.embedGraph = list(method = "UMAP")
 )
 Arguments:
 cms list List containing the preprocessed count matrices (default = self$cms.preprocessed).
 verbose logical Print messages or not (default = self$verbose).
 n.cores integer Number of cores for the calculations (default = self$n.cores).
 arg.buildGraph list A list with additional arguments for the buildGraph function in Conos
     (default = list())
 arg.findCommunities list A list with additional arguments for the findCommunities function
     in Conos (default = list())
```

```
arg.embedGraph list A list with additional arguments for the embedGraph function in Conos
     (default = list(method = "UMAP))
 Returns: Conos object
 Examples:
 \donttest{
 if (requireNamespace("pagoda2", quietly = TRUE)) {
 if (requireNamespace("conos", quietly = TRUE)) {
 # Simulate data
 testdata.cms <- lapply(seq_len(2), \(x) {
 out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)</pre>
 out[out < 0] <- 1
 dimnames(out) <- list(sapply(seq_len(2e3), \(x) paste0("gene",x)),</pre>
 sapply(seq_len(1e3), \(x) paste0("cell",x)))
 return(out)
 })
 # Initialize
 crm <- CRMetrics$new(cms = testdata.cms, samples = c("sample1", "sample2"), n.cores = 1)</pre>
 # Create embedding
 crm$doPreprocessing()
 crm$createEmbedding()
 } else {
 message("Package 'conos' not available.")
 }
 } else {
 message("Package 'pagoda2' not available.")
 }
 }
Method filterCms(): Filter cells based on depth, mitochondrial fraction and doublets from the
count matrix.
 Usage:
 CRMetrics$filterCms(
   depth.cutoff = NULL,
   mito.cutoff = NULL,
   doublets = NULL,
   species = c("human", "mouse"),
   samples.to.exclude = NULL,
   verbose = self$verbose,
   sep = "!!",
   raw = FALSE
 )
 Arguments:
 depth.cutoff numeric Depth (transcripts per cell) cutoff (default = NULL).
 mito.cutoff numeric Mitochondrial fraction cutoff (default = NULL).
 doublets character Doublet detection method to use (default = NULL).
```

```
species character Species to calculate the mitochondrial fraction for (default = "human").
 samples.to.exclude character Sample names to exclude (default = NULL)
 verbose logical Show progress (default = self$verbose)
 sep character Separator for creating unique cell names (default = "!!")
 raw boolean Filter on raw, unfiltered count matrices. Usually not intended (default = FALSE)
 Returns: list of filtered count matrices
 Examples:
 \donttest{
 if (requireNamespace("pagoda2", quietly = TRUE)) {
 if (requireNamespace("conos", quietly = TRUE)) {
 # Simulate data
 testdata.cms <- lapply(seq_len(2), \(x) {
 out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)</pre>
 out[out < 0] <- 1
 dimnames(out) <- list(sapply(seq_len(2e3), \(x) paste0("gene",x)),</pre>
 sapply(seq_len(1e3), \(x) paste0("cell",x)))
 return(out)
 })
 # Initialize
 crm <- CRMetrics$new(cms = testdata.cms, samples = c("sample1", "sample2"), n.cores = 1)</pre>
 # Create embedding
 crm$doPreprocessing()
 crm$createEmbedding()
 # Filter CMs
 crm$filterCms(depth.cutoff = 1e3, mito.cutoff = 0.05)
 } else {
 message("Package 'conos' not available.")
 }
 } else {
 message("Package 'pagoda2' not available.")
 }
Method selectMetrics(): Select metrics from summary.metrics
 Usage:
 CRMetrics$selectMetrics(ids = NULL)
 ids character Metric id to select (default = NULL).
 Returns: vector
 Examples:
```

```
# Simulate data
 testdata.cms <- lapply(seq_len(2), \(x) {
 out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)</pre>
 out[out < 0] <- 1
 dimnames(out) <- list(sapply(seq_len(2e3), \(x) paste0("gene",x)),</pre>
 sapply(seq_len(1e3), \(x) paste0("cell",x)))
 return(out)
 })
 # Initialize
 crm <- CRMetrics$new(cms = testdata.cms, samples = c("sample1", "sample2"), n.cores = 1)</pre>
 # Select metrics
 crm$selectMetrics()
 selection.metrics <- crm$selectMetrics(c(1:4))</pre>
Method plotFilteredCells(): Plot filtered cells in an embedding, in a bar plot, on a tile or
export the data frame
 Usage:
 CRMetrics$plotFilteredCells(
    type = c("embedding", "bar", "tile", "export"),
    depth = TRUE,
    depth.cutoff = 1000,
    doublet.method = NULL,
   mito.frac = TRUE,
   mito.cutoff = 0.05,
    species = c("human", "mouse"),
    size = 0.3,
    sep = "!!",
   cols = c("grey80", "red", "blue", "green", "yellow", "black", "pink", "purple"),
 )
 Arguments:
 type character The type of plot to use: embedding, bar, tile or export (default = c("embedding", "bar", "tile", "export")).
 depth logical Plot the depth or not (default = TRUE).
 depth.cutoff numeric Depth cutoff, either a single number or a vector with cutoff per sample
     and with sampleIDs as names (default = 1e3).
 doublet.method character Method to detect doublets (default = NULL).
 mito. frac logical Plot the mitochondrial fraction or not (default = TRUE).
 mito.cutoff numeric Mitochondrial fraction cutoff, either a single number or a vector with
     cutoff per sample and with sampleIDs as names (default = 0.05).
 species character Species to calculate the mitochondrial fraction for (default = c("human", "mouse")).
 size numeric Dot size (default = 0.3)
 sep character Separator for creating unique cell names (default = "!!")
 cols character Colors used for plotting (default = c("grey80","red","blue","green","yellow","black","pink","purple"))
 ... Plotting parameters passed to sccore::embeddingPlot.
```

```
Returns: ggplot2 object or data frame
 Examples:
 \donttest{
 if (requireNamespace("pagoda2", quietly = TRUE)) {
 if (requireNamespace("conos", quietly = TRUE)) {
 # Simulate data
 testdata.cms <- lapply(seq_len(2), \(x) {</pre>
 out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)</pre>
 out[out < 0] <- 1
 dimnames(out) <- list(sapply(seq_len(2e3), \(x) paste0("gene",x)),</pre>
 sapply(seq_len(1e3), \(x) paste0("cell",x)))
 return(out)
 })
 # Initialize
 crm <- CRMetrics$new(cms = testdata.cms, samples = c("sample1", "sample2"), n.cores = 1)</pre>
 # Create embedding
 crm$doPreprocessing()
 crm$createEmbedding()
 # Plot and extract result
 crm$plotFilteredCells(type = "embedding")
 filtered.cells <- crm$plotFilteredCells(type = "export")</pre>
 message("Package 'conos' not available.")
 }
 } else {
 message("Package 'pagoda2' not available.")
 }
 }
Method getDepth(): Extract sequencing depth from Conos object.
 Usage:
 CRMetrics$getDepth(cms = self$cms)
 cms list List of (sparse) count matrices (default = self$cms)
 Returns: data frame
 Examples:
 \donttest{
 if (requireNamespace("pagoda2", quietly = TRUE)) {
 if (requireNamespace("conos", quietly = TRUE)) {
 # Simulate data
 testdata.cms <- lapply(seq_len(2), \(x) {
 out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)</pre>
 out[out < 0] <- 1
```

```
dimnames(out) <- list(sapply(seq_len(2e3), \(x) paste0("gene",x)),</pre>
 sapply(seq_len(1e3), \(x) paste0("cell",x)))
 return(out)
 })
 # Initialize
 crm <- CRMetrics$new(cms = testdata.cms, samples = c("sample1", "sample2"), n.cores = 1)</pre>
 # Create embedding
 crm$doPreprocessing()
 crm$createEmbedding()
 # Get depth
 crm$getDepth()
 } else {
 message("Package 'conos' not available.")
 }
 } else {
 message("Package 'pagoda2' not available.")
 }
Method getMitoFraction(): Calculate the fraction of mitochondrial genes.
 Usage:
 CRMetrics$getMitoFraction(species = c("human", "mouse"), cms = self$cms)
 Arguments:
 species character Species to calculate the mitochondrial fraction for (default = "human").
 cms list List of (sparse) count matrices (default = self$cms)
 Returns: data frame
 Examples:
 \donttest{
 if (requireNamespace("pagoda2", quietly = TRUE)) {
 if (requireNamespace("conos", quietly = TRUE)) {
 # Simulate data
 testdata.cms <- lapply(seq_len(2), \(x) {
 out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)
 out[out < 0] <- 1
 dimnames(out) <- list(sapply(seq_len(2e3), \(x) paste0("gene",x)),</pre>
 sapply(seq_len(1e3), \(x) paste0("cell",x)))
 return(out)
 })
 # Initialize
 crm <- CRMetrics$new(cms = testdata.cms, samples = c("sample1", "sample2"), n.cores = 1)</pre>
 # Create embedding
 crm$doPreprocessing()
```

crm\$createEmbedding()

```
# Get mito. fraction
 crm$getMitoFraction(species = c("human", "mouse"))
 } else {
 message("Package 'conos' not available.")
 }
 } else {
 message("Package 'pagoda2' not available.")
 }
 }
Method prepareCellbender(): Create plots and script call for CellBender
 Usage:
 CRMetrics$prepareCellbender(
    shrinkage = 100,
    show.expected.cells = TRUE,
    show.total.droplets = TRUE,
    expected.cells = NULL,
    total.droplets = NULL,
    cms.raw = self$cms.raw,
    umi.counts = self$cellbender$umi.counts,
    data.path = self$data.path,
    samples = self$metadata$sample,
    verbose = self$verbose,
    n.cores = self$n.cores,
   unique.names = FALSE,
    sep = "!!"
 )
 Arguments:
 shrinkage integer Select every nth UMI count per cell for plotting. Improves plotting speed
     drastically. To plot all cells, set to 1 (default = 100)
 show.expected.cells logical Plot line depicting expected number of cells (default = TRUE)
 show.total.droplets logical Plot line depicting total droplets included for CellBender run
     (default = TRUE)
 expected.cells named numeric If NULL, expected cells will be deduced from the number of
     cells per sample identified by Cell Ranger. Otherwise, a named vector of expected cells with
     sample IDs as names. Sample IDs must match those in summary metrics (default: stored
     named vector)
 total.droplets named numeric If NULL, total droplets included will be deduced from ex-
     pected cells multiplied by 3. Otherwise, a named vector of total droplets included with
     sample IDs as names. Sample IDs must match those in summary.metrics (default: stored
     named vector)
 cms.raw list Raw count matrices from HDF5 Cell Ranger outputs (default = self$cms.raw)
 umi.counts list UMI counts calculated as column sums of raw count matrices from HDF5 Cell
     Ranger outputs (default: stored list)
 data.path character Path to Cell Ranger outputs (default = self$data.path)
```

```
samples character Sample names to include (default = self$metadata$sample)
 verbose logical Show progress (default: stored vector)
 n.cores integer Number of cores (default: stored vector)
 unique.names logical Create unique cell names (default = FALSE)
 sep character Separator for creating unique cell names (default = "!!")
 Returns: ggplot2 object and bash script
 Examples:
 \dontrun{
 crm <- CRMetrics$new(data.path = "/path/to/count/data")</pre>
 crm$prepareCellbender()
Method saveCellbenderScript():
 Usage:
 CRMetrics$saveCellbenderScript(
    file = "cellbender_script.sh",
    fpr = 0.01,
    epochs = 150,
    use.gpu = TRUE,
    expected.cells = NULL,
    total.droplets = NULL,
    data.path = self$data.path,
    samples = self$metadata$sample,
    args = NULL
 Arguments:
 file character File name for CellBender script. Will be stored in data.path (default: "cell-
     bender_script.sh")
 fpr numeric False positive rate for CellBender (default = 0.01)
 epochs integer Number of epochs for CellBender (default = 150)
 use.gpu logical Use CUDA capable GPU (default = TRUE)
 expected.cells named numeric If NULL, expected cells will be deduced from the number of
     cells per sample identified by Cell Ranger. Otherwise, a named vector of expected cells with
     sample IDs as names. Sample IDs must match those in summary.metrics (default: stored
     named vector)
 total.droplets named numeric If NULL, total droplets included will be deduced from ex-
     pected cells multiplied by 3. Otherwise, a named vector of total droplets included with
     sample IDs as names. Sample IDs must match those in summary.metrics (default: stored
     named vector)
 data.path character Path to Cell Ranger outputs (default = self$data.path)
 samples character Sample names to include (default = self$metadata$sample)
 args character (optional) Additional parameters for CellBender
 Returns: bash script
 Examples:
```

```
\dontrun{
 crm <- CRMetrics$new(data.path = "/path/to/count/data/")</pre>
 crm$prepareCellbender()
 crm$saveCellbenderScript()
 }
Method getExpectedCells(): Extract the expected number of cells per sample based on the
Cell Ranger summary metrics
 Usage:
 CRMetrics$getExpectedCells(samples = self$metadata$sample)
 Arguments:
 samples character Sample names to include (default = self$metadata$sample)
 Returns: A numeric vector
 Examples:
 # Simulate data
 testdata.cms <- lapply(seq_len(2), \(x) {
 out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)</pre>
 out[out < 0] <- 1
 dimnames(out) <- list(sapply(seq_len(2e3), \(x) paste0("gene",x)),</pre>
 sapply(seq_len(1e3), \(x) paste0("cell",x)))
 return(out)
 })
 # Initialize
 crm <- CRMetrics$new(cms = testdata.cms, samples = c("sample1", "sample2"), n.cores = 1)</pre>
 # Get summary
 crm$addSummaryFromCms()
 # Get no. cells
 crm$getExpectedCells()
Method getTotalDroplets(): Get the total number of droplets included in the CellBender
estimations. Based on the Cell Ranger summary metrics and multiplied by a preset multiplier.
 Usage:
 CRMetrics$getTotalDroplets(samples = self$metadata$sample, multiplier = 3)
 Arguments:
 samples character Samples names to include (default = self$metadata$sample)
 multiplier numeric Number to multiply expected number of cells with (default = 3)
 Returns: A numeric vector
 Examples:
 # Simulate data
 testdata.cms <- lapply(seq_len(2), \(x) {
 out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)</pre>
 out[out < 0] <- 1
```

```
dimnames(out) <- list(sapply(seq_len(2e3), \(x) paste0("gene",x)),</pre>
 sapply(seq_len(1e3), \(x) paste0("cell",x)))
 return(out)
 })
 # Initialize
 crm <- CRMetrics$new(cms = testdata.cms, samples = c("sample1", "sample2"), n.cores = 1)</pre>
 # Add summary
 crm$addSummaryFromCms()
 # Get no. droplets
 crm$getTotalDroplets()
Method addCms(): Add a list of count matrices to the CRMetrics object.
 Usage:
 CRMetrics$addCms(
    cms = NULL,
    data.path = self$data.path,
    samples = self$metadata$sample,
    cellbender = FALSE,
    raw = FALSE,
    symbol = TRUE,
    unique.names = TRUE,
    sep = "!!",
    add.metadata = TRUE,
   n.cores = self$n.cores,
    verbose = self$verbose
 )
 Arguments:
 cms list List of (sparse) count matrices (default = NULL)
 data.path character Path to cellranger count data (default = self$data.path).
 samples character Vector of sample names. If NULL, samples are extracted from cms (default
     = self$metadata$sample)
 cellbender logical Add CellBender filtered count matrices in HDF5 format. Requires that
     "cellbender" is in the names of the files (default = FALSE)
 raw logical Add raw count matrices from Cell Ranger output. Cannot be combined with
     cellbender=TRUE (default = FALSE)
 symbol character The type of gene IDs to use, SYMBOL (TRUE) or ENSEMBLE (default =
     TRUE)
 unique.names logical Make cell names unique based on sep parameter (default = TRUE)
 sep character Separator used to create unique cell names (default = "!!")
 add.metadata boolean Add metadata from cms or not (default = TRUE)
 n.cores integer Number of cores to use (default = self$n.cores)
 verbose boolean Print progress (default = self$verbose)
 Returns: Add list of (sparse) count matrices to R6 class object
```

```
Examples:
 \dontrun{
 crm <- CRMetrics$new(data.path = "/path/to/count/data/")</pre>
 # Simulate data
 testdata.cms <- lapply(seq_len(2), \xspace (x)) {
 out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)</pre>
 out[out < 0] <- 1
 dimnames(out) <- list(sapply(seq_len(2e3), \(x) paste0("gene",x)),</pre>
 sapply(seq_len(1e3), \(x) paste0("cell",x)))
 return(out)
 })
 crm$addCms(cms = testdata.cms)
 }
Method plotCbTraining(): Plot the results from the CellBender estimations
 Usage:
 CRMetrics$plotCbTraining(
   data.path = self$data.path,
   samples = self$metadata$sample,
   pal = self$pal
 )
 Arguments:
 data.path character Path to Cell Ranger outputs (default = self$data.path)
 samples character Sample names to include (default = self$metadata$sample)
 pal character Plotting palette (default = self$pal)
 Returns: A ggplot2 object
 Examples:
 \dontrun{
 crm <- CRMetrics$new(data.path = "/path/to/count/data/")</pre>
 crm$prepareCellbender()
 crm$saveCellbenderScript()
 ## Run CellBender script
 crm$plotCbTraining()
 }
Method plotCbCellProbs(): Plot the CellBender assigned cell probabilities
 Usage:
 CRMetrics$plotCbCellProbs(
   data.path = self$data.path,
   samples = self$metadata$sample,
   low.col = "gray",
   high.col = "red"
 )
 Arguments:
```

```
data.path character Path to Cell Ranger outputs (default = self$data.path)
 samples character Sample names to include (default = self$metadata$sample)
 low.col character Color for low probabilities (default = "gray")
 high.col character Color for high probabilities (default = "red")
 Returns: A ggplot2 object
 Examples:
 \dontrun{
 crm <- CRMetrics$new(data.path = "/path/to/count/data/")</pre>
 crm$prepareCellbender()
 crm$saveCellbenderScript()
 ## Run the CellBender script
 crm$plotCbCellProbs()
Method plotCbAmbExp(): Plot the estimated ambient gene expression per sample from Cell-
Bender calculations
 Usage:
 CRMetrics$plotCbAmbExp(
   cutoff = 0.005,
   data.path = self$data.path,
    samples = self$metadata$sample
 )
 Arguments:
 cutoff numeric Horizontal line included in the plot to indicate highly expressed ambient genes
     (default = 0.005)
 data.path character Path to Cell Ranger outputs (default = self$data.path)
 samples character Sample names to include (default = self$metadata$sample)
 Returns: A ggplot2 object
 Examples:
 \dontrun{
 crm <- CRMetrics$new(data.path = "/path/to/count/data/")</pre>
 crm$prepareCellbender()
 crm$saveCellbenderScript()
 ## Run CellBender script
 crm$plotCbAmbExp()
 }
Method plotCbAmbGenes(): Plot the most abundant estimated ambient genes from the Cell-
Bender calculations
 Usage:
 CRMetrics$plotCbAmbGenes(
   cutoff = 0.005,
   data.path = self$data.path,
   samples = self$metadata$sample,
   pal = self$pal
 )
```

```
Arguments:
 cutoff numeric Cutoff of ambient gene expression to use to extract ambient genes per sample
 data.path character Path to Cell Ranger outputs (default = self$data.path)
 samples character Sample names to include (default = self$metadata$sample)
 pal character Plotting palette (default = self$pal)
 Returns: A ggplot2 object
 Examples:
 \dontrun{
 crm <- CRMetrics$new(data.path = "/path/to/count/data/")</pre>
 crm$prepareCellbender()
 crm$saveCellbenderScript()
 ## Run CellBender script
 crm$plotCbAmbGenes()
 }
Method addSummaryFromCms(): Add summary metrics from a list of count matrices
 Usage:
 CRMetrics$addSummaryFromCms(
   cms = self$cms.
   n.cores = self$n.cores,
   verbose = self$verbose
 )
 Arguments:
 cms list A list of filtered count matrices (default = self$cms)
 n.cores integer Number of cores to use (default = self$n.cores)
 verbose logical Show progress (default = self$verbose)
 Returns: data.frame
 Examples:
 # Simulate data
 testdata.cms <- lapply(seq_len(2), \(x) {
 out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)
 out[out < 0] <- 1
 dimnames(out) <- list(sapply(seq_len(2e3), \(x) paste0("gene",x)),</pre>
 sapply(seq_len(1e3), \x) paste0("cell",x)))
 return(out)
 })
 # Initialize
 crm <- CRMetrics$new(cms = testdata.cms, samples = c("sample1", "sample2"), n.cores = 1)</pre>
 # Add summary
 crm$addSummaryFromCms()
Method runSoupX(): Run SoupX ambient RNA estimation and correction
 Usage:
```

```
CRMetrics$runSoupX(
   data.path = self$data.path,
    samples = self$metadata$sample,
   n.cores = self$n.cores,
   verbose = self$verbose,
   arg.load10X = list(),
   arg.autoEstCont = list(),
    arg.adjustCounts = list()
 )
 Arguments:
 data.path character Path to Cell Ranger outputs (default = self$data.path)
 samples character Sample names to include (default = self$metadata$sample)
 n.cores numeric Number of cores (default = self$n.cores)
 verbose logical Show progress (default = self$verbose)
 arg.load10X list A list with additional parameters for SoupX::load10X (default = list())
 arg.autoEstCont list A list with additional parameters for SoupX::autoEstCont (default =
     list())
 arg.adjustCounts list A list with additional parameters for SoupX::adjustCounts (default
     = list()
 Returns: List containing a list with corrected counts, and a data frame containing plotting
 estimations
 Examples:
 \dontrun{
 crm <- CRMetrics$new(data.path = "/path/to/count/data/")</pre>
 crm$runSoupX()
 }
Method plotSoupX(): Plot the results from the SoupX estimations
 CRMetrics$plotSoupX(plot.df = self$soupx$plot.df)
 Arguments:
 plot.df data.frame SoupX estimations (default = self$soupx$plot.df)
 Returns: A ggplot2 object
 Examples:
 \dontrun{
 crm <- CRMetrics$new(data.path = "/path/to/count/data/")</pre>
 crm$runSoupX()
 crm$plotSoupX()
Method plotCbCells(): Plot CellBender cell estimations against the estimated cell numbers
from Cell Ranger
```

Usage:

```
CRMetrics$plotCbCells(
    data.path = self$data.path,
   samples = self$metadata$sample,
   pal = self$pal
 )
 Arguments:
 data.path character Path to Cell Ranger outputs (default = self$data.path)
 samples character Sample names to include (default = self$metadata$sample)
 pal character Plotting palette (default = self$pal)
 Returns: A ggplot2 object
 Examples:
 \dontrun{
 crm <- CRMetrics$new(data.path = "/path/to/count/data/")</pre>
 crm$prepareCellbender()
 crm$saveCellbenderScript()
 ## Run CellBender script
 crm$plotCbCells()
 }
Method addDoublets(): Add doublet results created from exported Python script
 Usage:
 CRMetrics$addDoublets(
   method = c("scrublet", "doubletdetection"),
   data.path = self$data.path,
   samples = self$metadata$sample,
    cms = self$cms,
    verbose = self$verbose
 )
 Arguments:
 method character Which method to use, either scrublet or doubletdetection (default is
 data.path character Path to Cell Ranger outputs (default = self$data.path)
 samples character Sample names to include (default = self$metadata$sample)
 cms list List containing the count matrices (default = self$cms).
 verbose boolean Print progress (default = self$verbose)
 Returns: List of doublet results
 Examples:
 \dontrun{
 crm <- CRMetrics$new(data.path = "/path/to/count/data/")</pre>
 crm$detectDoublets(export = TRUE)
 ## Run Python script
 crm$addDoublets()
 }
```

Method clone(): The objects of this class are cloneable with this method.

```
Usage:
CRMetrics$clone(deep = FALSE)
Arguments:
deep Whether to make a deep clone.
```

Examples

```
## Method `CRMetrics$new`
## -----
## Not run:
crm <- CRMetrics$new(data.path = "/path/to/count/data/")</pre>
## End(Not run)
## -----
## Method `CRMetrics$addDetailedMetrics`
## -----
# Simulate data
testdata.cms <- lapply(seq_len(2), \(x) {
out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)</pre>
out[out < 0] <- 1
dimnames(out) <- list(sapply(seq_len(2e3), \(x) paste0("gene",x)),</pre>
sapply(seq_len(1e3), \(x) paste0("cell",x)))
return(out)
})
# Initialize
crm <- CRMetrics$new(cms = testdata.cms, samples = c("sample1", "sample2"), n.cores = 1)</pre>
# Run function
crm$addDetailedMetrics()
## -----
## Method `CRMetrics$addComparison`
# Simulate data
testdata.cms <- lapply(seq_len(2), \(x) {
out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)</pre>
out[out < 0] <- 1
dimnames(out) <- list(sapply(seq_len(2e3), \(x) paste0("gene",x)),</pre>
sapply(seq\_len(1e3), \xspace("cell",x)))
return(out)
})
# Initialize
crm <- CRMetrics$new(cms = testdata.cms, samples = c("sample1", "sample2"), n.cores = 1)</pre>
```

```
# Add metadata
crm$metadata <- data.frame(sex = c("male", "female"))</pre>
# Add comparison group
crm$addComparison(comp.group = "sex")
## -----
## Method `CRMetrics$plotSamples`
samples <- c("sample1", "sample2")</pre>
# Simulate data
testdata.cms <- lapply(seq_len(2), \(x) {
out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)</pre>
out[out < 0] <- 1
dimnames(out) \leftarrow list(sapply(seq_len(2e3), \xspace ("gene",x)),
sapply(seq\_len(1e3), \x) paste0("cell",x)))
return(out)
})
names(testdata.cms) <- samples</pre>
# Create metadata
metadata <- data.frame(sample = samples,</pre>
sex = c("male","female"),
condition = c("a","b"))
# Initialize
crm <- CRMetrics$new(cms = testdata.cms, metadata = metadata, n.cores = 1)</pre>
# Plot
crm$plotSamples(comp.group = "sex", second.comp.group = "condition")
## -----
## Method `CRMetrics$plotSummaryMetrics`
# Simulate data
testdata.cms <- lapply(seq_len(2), \(x) {
out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)</pre>
out[out < 0] <- 1
dimnames(out) \leftarrow list(sapply(seq_len(2e3), \xspace ("gene",x)),
sapply(seq\_len(1e3), \x) paste0("cell",x)))
return(out)
})
# Initialize
crm <- CRMetrics$new(cms = testdata.cms, samples = c("sample1", "sample2"), n.cores = 1)</pre>
# Add summary metrics
crm$addSummaryFromCms()
```

```
crm$plotSummaryMetrics(plot.geom = "point")
## -----
## Method `CRMetrics$plotDetailedMetrics`
# Simulate data
testdata.cms <- lapply(seq_len(2), \(x) {
out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)</pre>
out[out < 0] <- 1
dimnames(out) <- list(sapply(seq_len(2e3), \(x) paste0("gene",x)),</pre>
sapply(seq_len(1e3), \x) paste0("cell",x)))
return(out)
})
# Initialize
crm <- CRMetrics$new(cms = testdata.cms, samples = c("sample1", "sample2"), n.cores = 1)</pre>
# Add detailed metrics
crm$addDetailedMetrics()
# Plot
crm$plotDetailedMetrics()
## -----
## Method `CRMetrics$plotEmbedding`
if (requireNamespace("pagoda2", quietly = TRUE)) {
if (requireNamespace("conos", quietly = TRUE)) {
# Simulate data
testdata.cms <- lapply(seq_len(2), \xspace (x) {
out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)</pre>
out[out < 0] <- 1
dimnames(out) <- list(sapply(seq_len(2e3), \(x) paste0("gene",x)),</pre>
sapply(seq_len(1e3), \x) paste0("cell",x)))
return(out)
})
# Initialize
crm <- CRMetrics$new(cms = testdata.cms, samples = c("sample1", "sample2"), n.cores = 1)</pre>
# Create embedding
crm$doPreprocessing()
crm$createEmbedding()
crm$plotEmbedding()
} else {
message("Package 'conos' not available.")
```

```
}
} else {
message("Package 'pagoda2' not available.")
## -----
## Method `CRMetrics$plotDepth`
if (requireNamespace("pagoda2", quietly = TRUE)) {
if (requireNamespace("conos", quietly = TRUE)) {
# Simulate data
testdata.cms <- lapply(seq_len(2), \(x) {
out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)</pre>
out[out < 0] <- 1
dimnames(out) \leftarrow list(sapply(seq_len(2e3), \xspace ("gene",x)),
sapply(seq\_len(1e3), \x) paste0("cell",x)))
return(out)
})
# Initialize
crm <- CRMetrics$new(cms = testdata.cms, samples = c("sample1", "sample2"), n.cores = 1)</pre>
# Create embedding
crm$doPreprocessing()
crm$createEmbedding()
# Plot
crm$plotDepth()
} else {
message("Package 'conos' not available.")
}
} else {
message("Package 'pagoda2' not available.")
## Method `CRMetrics$plotMitoFraction`
if (requireNamespace("pagoda2", quietly = TRUE)) {
if (requireNamespace("conos", quietly = TRUE)) {
# Simulate data
testdata.cms <- lapply(seq_len(2), \(x) {
out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)</pre>
out[out < 0] <- 1
dimnames(out) \leftarrow list(sapply(seq_len(2e3), \xspace ("gene",x)),
sapply(seq\_len(1e3), \x) paste0("cell",x)))
return(out)
```

```
})
# Initialize
crm <- CRMetrics$new(cms = testdata.cms, samples = c("sample1", "sample2"), n.cores = 1)</pre>
# Create embedding
crm$doPreprocessing()
crm$createEmbedding()
# Plot
crm$plotMitoFraction()
} else {
message("Package 'conos' not available.")
} else {
message("Package 'pagoda2' not available.")
## -----
## Method `CRMetrics$detectDoublets`
## Not run:
# Simulate data
testdata.cms <- lapply(seq_len(2), \(x) {
out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)</pre>
out[out < 0] <- 1
\label{eq:dimnames} \mbox{dimnames(out)} <- \mbox{list(sapply(seq_len(2e3), \xspace), paste0("gene",x)),}
sapply(seq\_len(1e3), \x) paste0("cell",x)))
return(out)
})
# Initialize
crm <- CRMetrics$new(cms = testdata.cms, samples = c("sample1", "sample2"), n.cores = 1)</pre>
# Detect doublets
crm$detectDoublets(method = "scrublet",
conda.path = "/opt/software/miniconda/4.12.0/condabin/conda")
## End(Not run)
## -----
## Method `CRMetrics$doPreprocessing`
## -----
if (requireNamespace("pagoda2", quietly = TRUE)) {
# Simulate data
testdata.cms <- lapply(seq_len(2), \(x) {
out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)</pre>
out[out < 0] <- 1
```

```
dimnames(out) <- list(sapply(seq_len(2e3), \(x) paste0("gene",x)),</pre>
sapply(seq_len(1e3), \(x) paste0("cell",x)))
return(out)
})
# Initialize
crm <- CRMetrics$new(cms = testdata.cms, samples = c("sample1", "sample2"), n.cores = 1)</pre>
# Perform preprocessing
crm$doPreprocessing(preprocess = "pagoda2")
} else {
message("Package 'pagoda2' not available.")
## Method `CRMetrics$createEmbedding`
## -----
if (requireNamespace("pagoda2", quietly = TRUE)) {
if (requireNamespace("conos", quietly = TRUE)) {
# Simulate data
testdata.cms <- lapply(seq_len(2), \xspace(x) {
out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)</pre>
out[out < 0] <- 1
dimnames(out) <- list(sapply(seq_len(2e3), \(x) paste0("gene",x)),</pre>
sapply(seq\_len(1e3), \x) paste0("cell",x)))
return(out)
})
# Initialize
crm <- CRMetrics$new(cms = testdata.cms, samples = c("sample1", "sample2"), n.cores = 1)</pre>
# Create embedding
crm$doPreprocessing()
crm$createEmbedding()
} else {
message("Package 'conos' not available.")
}
} else {
message("Package 'pagoda2' not available.")
## -----
## Method `CRMetrics$filterCms`
## -----
if (requireNamespace("pagoda2", quietly = TRUE)) {
if (requireNamespace("conos", quietly = TRUE)) {
# Simulate data
```

```
testdata.cms <- lapply(seq_len(2), \(x) {
out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)</pre>
out[out < 0] <- 1
dimnames(out) <- list(sapply(seq_len(2e3), \(x) paste0("gene",x)),</pre>
sapply(seq\_len(1e3), \x) paste0("cell",x)))
return(out)
})
# Initialize
crm <- CRMetrics$new(cms = testdata.cms, samples = c("sample1", "sample2"), n.cores = 1)</pre>
# Create embedding
crm$doPreprocessing()
crm$createEmbedding()
# Filter CMs
crm$filterCms(depth.cutoff = 1e3, mito.cutoff = 0.05)
message("Package 'conos' not available.")
}
} else {
message("Package 'pagoda2' not available.")
## Method `CRMetrics$selectMetrics`
# Simulate data
testdata.cms <- lapply(seq_len(2), \(x) {
out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)</pre>
out[out < 0] <- 1
dimnames(out) <- list(sapply(seq_len(2e3), \(x) paste0("gene",x)),</pre>
sapply(seq_len(1e3), \(x) paste0("cell",x)))
return(out)
})
crm <- CRMetrics$new(cms = testdata.cms, samples = c("sample1", "sample2"), n.cores = 1)</pre>
# Select metrics
crm$selectMetrics()
selection.metrics <- crm$selectMetrics(c(1:4))</pre>
## -----
## Method `CRMetrics$plotFilteredCells`
if (requireNamespace("pagoda2", quietly = TRUE)) {
if (requireNamespace("conos", quietly = TRUE)) {
```

```
# Simulate data
testdata.cms <- lapply(seq_len(2), \(x) {
out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)</pre>
out[out < 0] <- 1
dimnames(out) <- list(sapply(seq_len(2e3), \(x) paste0("gene",x)),</pre>
sapply(seq\_len(1e3), \x) paste0("cell",x)))
return(out)
})
# Initialize
crm <- CRMetrics$new(cms = testdata.cms, samples = c("sample1", "sample2"), n.cores = 1)</pre>
# Create embedding
crm$doPreprocessing()
crm$createEmbedding()
# Plot and extract result
crm$plotFilteredCells(type = "embedding")
filtered.cells <- crm$plotFilteredCells(type = "export")</pre>
message("Package 'conos' not available.")
}
} else {
message("Package 'pagoda2' not available.")
## Method `CRMetrics$getDepth`
if (requireNamespace("pagoda2", quietly = TRUE)) {
if (requireNamespace("conos", quietly = TRUE)) {
# Simulate data
testdata.cms <- lapply(seq_len(2), \xspace (x) {
out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)</pre>
out[out < 0] <- 1
dimnames(out) <- list(sapply(seq_len(2e3), \(x) paste0("gene",x)),</pre>
sapply(seq_len(1e3), \x) paste0("cell",x)))
return(out)
})
# Initialize
crm <- CRMetrics$new(cms = testdata.cms, samples = c("sample1", "sample2"), n.cores = 1)</pre>
# Create embedding
crm$doPreprocessing()
crm$createEmbedding()
# Get depth
crm$getDepth()
} else {
```

```
message("Package 'conos' not available.")
} else {
message("Package 'pagoda2' not available.")
}
## Method `CRMetrics$getMitoFraction`
if (requireNamespace("pagoda2", quietly = TRUE)) {
if (requireNamespace("conos", quietly = TRUE)) {
# Simulate data
testdata.cms <- lapply(seq_len(2), \(x) {
out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)</pre>
out[out < 0] <- 1
dimnames(out) <- list(sapply(seq_len(2e3), \(x) paste0("gene",x)),</pre>
sapply(seq_len(1e3), \(x) paste0("cell",x)))
return(out)
})
# Initialize
crm <- CRMetrics$new(cms = testdata.cms, samples = c("sample1", "sample2"), n.cores = 1)</pre>
# Create embedding
crm$doPreprocessing()
crm$createEmbedding()
# Get mito. fraction
crm$getMitoFraction(species = c("human", "mouse"))
message("Package 'conos' not available.")
} else {
message("Package 'pagoda2' not available.")
}
## Method `CRMetrics$prepareCellbender`
crm <- CRMetrics$new(data.path = "/path/to/count/data")</pre>
crm$prepareCellbender()
## End(Not run)
## -----
## Method `CRMetrics$saveCellbenderScript`
## -----
```

```
## Not run:
crm <- CRMetrics$new(data.path = "/path/to/count/data/")</pre>
crm$prepareCellbender()
crm$saveCellbenderScript()
## End(Not run)
## Method `CRMetrics$getExpectedCells`
# Simulate data
testdata.cms <- lapply(seq_len(2), \(x) {
out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)</pre>
out[out < 0] <- 1
dimnames(out) \leftarrow list(sapply(seq_len(2e3), \xspace ("gene",x)),
sapply(seq\_len(1e3), \x) paste0("cell",x)))
return(out)
})
# Initialize
crm <- CRMetrics$new(cms = testdata.cms, samples = c("sample1", "sample2"), n.cores = 1)</pre>
# Get summary
crm$addSummaryFromCms()
# Get no. cells
crm$getExpectedCells()
## -----
## Method `CRMetrics$getTotalDroplets`
# Simulate data
testdata.cms <- lapply(seq_len(2), \xspace (x) {
out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)</pre>
out[out < 0] <- 1
dimnames(out) <- list(sapply(seq_len(2e3), \(x) paste0("gene",x)),</pre>
sapply(seq_len(1e3), \(x) paste0("cell",x)))
return(out)
})
# Initialize
crm <- CRMetrics$new(cms = testdata.cms, samples = c("sample1", "sample2"), n.cores = 1)</pre>
# Add summary
crm$addSummaryFromCms()
# Get no. droplets
crm$getTotalDroplets()
## -----
```

```
## Method `CRMetrics$addCms`
## Not run:
crm <- CRMetrics$new(data.path = "/path/to/count/data/")</pre>
# Simulate data
testdata.cms <- lapply(seq_len(2), \(x) {
out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)</pre>
out[out < 0] <- 1
\label{eq:dimnames} \mbox{dimnames(out)} <- \mbox{list(sapply(seq_len(2e3), \xspace), paste0("gene",x)),}
sapply(seq\_len(1e3), \x) paste0("cell",x)))
return(out)
})
crm$addCms(cms = testdata.cms)
## End(Not run)
## -----
## Method `CRMetrics$plotCbTraining`
## Not run:
crm <- CRMetrics$new(data.path = "/path/to/count/data/")</pre>
crm$prepareCellbender()
crm$saveCellbenderScript()
## Run CellBender script
crm$plotCbTraining()
## End(Not run)
## -----
## Method `CRMetrics$plotCbCellProbs`
crm <- CRMetrics$new(data.path = "/path/to/count/data/")</pre>
crm$prepareCellbender()
crm$saveCellbenderScript()
## Run the CellBender script
crm$plotCbCellProbs()
## End(Not run)
## -----
## Method `CRMetrics$plotCbAmbExp`
## -----
## Not run:
crm <- CRMetrics$new(data.path = "/path/to/count/data/")</pre>
crm$prepareCellbender()
crm$saveCellbenderScript()
```

```
## Run CellBender script
crm$plotCbAmbExp()
## End(Not run)
## -----
## Method `CRMetrics$plotCbAmbGenes`
## Not run:
crm <- CRMetrics$new(data.path = "/path/to/count/data/")</pre>
crm$prepareCellbender()
crm$saveCellbenderScript()
## Run CellBender script
crm$plotCbAmbGenes()
## End(Not run)
## -----
## Method `CRMetrics$addSummaryFromCms`
## -----
# Simulate data
testdata.cms <- lapply(seq_len(2), \xspace (x) {
out <- Matrix::rsparsematrix(2e3, 1e3, 0.1)</pre>
out[out < 0] <- 1
dimnames(out) <- list(sapply(seq_len(2e3), \(x) paste0("gene",x)),</pre>
sapply(seq_len(1e3), \(x) paste0("cell",x)))
return(out)
})
# Initialize
crm <- CRMetrics$new(cms = testdata.cms, samples = c("sample1", "sample2"), n.cores = 1)</pre>
# Add summary
crm$addSummaryFromCms()
## -----
## Method `CRMetrics$runSoupX`
## -----
## Not run:
crm <- CRMetrics$new(data.path = "/path/to/count/data/")</pre>
crm$runSoupX()
## End(Not run)
## -----
## Method `CRMetrics$plotSoupX`
## Not run:
crm <- CRMetrics$new(data.path = "/path/to/count/data/")</pre>
```

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```
crm$runSoupX()
crm$plotSoupX()
## End(Not run)
## Method `CRMetrics$plotCbCells`
## Not run:
crm <- CRMetrics$new(data.path = "/path/to/count/data/")</pre>
crm$prepareCellbender()
crm$saveCellbenderScript()
## Run CellBender script
crm$plotCbCells()
## End(Not run)
## -----
## Method `CRMetrics$addDoublets`
## Not run:
crm <- CRMetrics$new(data.path = "/path/to/count/data/")</pre>
crm$detectDoublets(export = TRUE)
## Run Python script
crm$addDoublets()
## End(Not run)
```

read10xH5

Read 10x HDF5 files

Description

Read 10x HDF5 files

Usage

```
read10xH5(
  data.path,
  samples = NULL,
  type = c("raw", "filtered", "cellbender", "cellbender_filtered"),
  symbol = TRUE,
  sep = "!!",
  n.cores = 1,
  verbose = TRUE,
  unique.names = FALSE
)
```

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Arguments

data.path character samples character vector, select specific samples for processing (default = NULL) name of H5 file to search for, "raw" and "filtered" are Cell Ranger count outputs, type "cellbender" is output from CellBender after running script from saveCellbenderScript symbol logical Use gene SYMBOLs (TRUE) or ENSEMBL IDs (FALSE) (default = TRUE) character Separator for creating unique cell names from sample IDs and cell IDs sep (default = "!!") integer Number of cores (default = 1) n.cores verbose logical Print progress (default = TRUE)

logical Create unique cell IDs (default = FALSE)

Value

list with sparse count matrices

unique.names

Examples

```
## Not run:
cms.h5 <- read10xH5(data.path = "/path/to/count/data")
## End(Not run)</pre>
```

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