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velociraptor-package velociraptor: Toolkit for Single-Cell Velocity

Description

This package provides Bioconductor-friendly wrappers for RNA velocity calculations in single-cell RNA-seq data. We use the basilisk package to manage Conda environments, and the zellkonverter package to convert data structures between SingleCellExperiment (R) and AnnData (Python). The information produced by the velocity methods is stored in the various components of the Single-CellExperiment class.

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See Also

Useful links:

- https://github.com/kevinrue/velociraptor
- Report bugs at https://github.com/kevinrue/velociraptor/issues

embedVelocity

Description

Project the velocity vector for each cell onto an existing low-dimensional embedding.

Usage

```
embedVelocity(x, vobj, ...)
## S4 method for signature 'ANY'
embedVelocity(x, vobj, ...)
## S4 method for signature 'SingleCellExperiment'
embedVelocity(x, vobj, ..., use.dimred = 1)
```

Arguments

х	A numeric matrix of low-dimensional coordinates, e.g., after t-SNE. Alterna- tively, a SingleCellExperiment containing such coordinates in its reducedDims.
vobj	A SingleCellExperiment containing the output of the velocity calculations, typ- ically after running scvelo.
	For the generic, further arguments to pass to specific methods.
	For the ANY method, further arguments to pass to the velocity_embedding Python function from scVelo.
	For the SingleCellExperiment method, further arguments to pass to the ANY method.
use.dimred	String or integer scalar specifying the reduced dimensions to retrieve from x.

Details

This is a simple wrapper around the scvelo.tools.velocity_embedding function. Briefly, we construct a cell-cell transition matrix where a cell is more likely to transition to one of its neighbors if its velocity vector is pointing in the same direction as that neighbor. The resulting matrix is then used to compute a weighted average of the positions in x, allowing us to compute a velocity in the low-dimensional embedding.

Value

A numeric matrix of the same dimensions as x, containing the projected velocity vectors in that embedding.

Author(s)

Aaron Lun

Examples

```
example(scvelo, echo=FALSE) # recycling that example.
# Making up a new embedding.
tsne.results <- matrix(rnorm(2*ncol(out)), ncol=2)
# Projecting the future state of each cell:
projected <- embedVelocity(tsne.results, out)</pre>
```

gridVectors

Summarize vectors into a grid

Description

Summarize the velocity vectors into a grid, usually for easy plotting.

Usage

```
gridVectors(x, embedded, ...)
## S4 method for signature 'ANY'
gridVectors(
    x,
    embedded,
    resolution = 40,
    scale = TRUE,
    as.data.frame = TRUE,
    return.intermediates = FALSE
)
## S4 method for signature 'SingleCellExperiment'
```

gridVectors(x, embedded, ..., use.dimred = 1)

Arguments

x	A numeric matrix of low-dimensional coordinates, e.g., after t-SNE. Alterna- tively, a SingleCellExperiment containing such coordinates in its reducedDims.
embedded	A low-dimensional projection of the velocity vectors into the embedding of x. This should be of the same dimensions as x and is typically produced by embedVelocity.
	For the generic, further arguments to pass to specific methods. For the SingleCellExperiment method, further arguments to pass to the ANY
	method.
resolution	Integer scalar specifying the resolution of the grid, in terms of the number of grid intervals along each axis.

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gridVectors

scale	Logical scalar indicating whether the averaged vectors should be scaled by the grid resolution.
as.data.frame	Logical scalar indicating whether the output should be a data.frame. If FALSE, a list of two matrices is returned.
return.intermediates	
	Logical scalar indicating whether intermediate objects should also be returned. This enforces as.data.frame=FALSE and throws a warning if it is TRUE.
use.dimred	String or integer scalar specifying the reduced dimensions to retrieve from x.

Details

This partitions the bounding box of x into a grid with resolution units in each dimension. The locations and vectors of all cells in each block are averaged to obtain a representative of that block. This is most obviously useful for visualization to avoid overplotting of velocity vectors.

If scale=TRUE, per-block vectors are scaled so that the median vector length is comparable to the spacing between blocks. This improves visualization when the scales of x and embedded are not immediately comparable.

Value

If as.data.frame=FALSE, a list is returned containing start and end, two numeric matrices with one row per non-empty block in the grid and one column per column in x. start contains the mean location of all cells inside that block, and end contains the endpoint after adding the (scaled) average of the block's cell's velocity vectors.

If as.data.frame=TRUE, a data.frame is returned with numeric columns of the same contents as the list above. Column names are prefixed by start.* and end.*.

If return.intermediates=TRUE, a list is returned (irrespective of the value of as.data.frame) that in addition to start and end also contains intermediate objects limits (the ranges in x and y), delta (the grid intervals in x and y), categories (a DataFrame with integer row and column indices for each cell that specify the grid field that it is contained in), grp (numerical index of grid fields for each cell) and vec (velocity vectors for non-empty grid fields).

Author(s)

Aaron Lun

See Also

embedVelocity, to generate embedded.

Examples

```
tsne.results <- matrix(rnorm(10000), ncol=2)
tsne.vectors <- matrix(rnorm(10000), ncol=2)
out <- gridVectors(tsne.results, tsne.vectors)
# Demonstration for plotting.
plot(tsne.results[,1], tsne.results[,2], col='grey')</pre>
```

arrows(out\$start.1, out\$start.2, out\$end.1, out\$end.2, length=0.05)

plotVelocity

Phase and velocity graphs for a set of genes

Description

For a each gene in a set of genes, show the phase graph (spliced versus unspliced counts and fitted model) and reduced dimension graphs with cell colored by velocity and (spliced) expression.

Usage

```
plotVelocity(
  х,
  genes,
  use.dimred = 1,
  assay.splicedM = "Ms",
  assay.unsplicedM = "Mu",
 which.plots = c("phase", "velocity", "expression"),
  genes.per.row = 1,
  color_by = "#222222",
  color.alpha = 0.4,
 colors.velocity = c("#A50026", "#D73027", "#F46D43", "#FDAE61", "#FEE08B", "#FFFFBF",
    "#D9EF8B", "#A6D96A", "#66BD63", "#1A9850", "#006837"),
 colors.expression = c("#440154", "#482576", "#414487", "#35608D", "#2A788E", "#21908C",
    "#22A884", "#43BF71", "#7AD151", "#BBDF27", "#FDE725"),
 max.abs.velo = 0.001
)
```

Arguments

x	A SingleCellExperiment object with RNA velocity results as returned by scvelo, and low-dimensional coordinates, e.g., after t-SNE, in its reducedDims.
genes	A character vector with one or several genes for which to plot phase and velocity graphs. genes have to be in rownames(x).
use.dimred	String or integer scalar specifying the reduced dimensions to retrieve from x.
assay.splicedM	An integer scalar or string specifying the assay of x containing the moments of spliced abundances.
assay.unsplicedM	
	An integer scalar or string specifying the assay of x containing the moments unspliced abundances.
which.plots	A character vector specifying which plots to create for each gene. Possible values are "phase", "velocity", "expression" and correspond to the phase graph or reduced dimension graphs with cells colored by velocity or (spliced) expression.

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genes.per.row	An integer scalar with the numbers of genes to visualize per row of plots. For example, if which.plots = c("phase", "expression") and genes.per.row = 2, the resulting figure will have four plot panels per row.
color_by	A character scalar specifying a column in colData(x) to color cells in the phase graph. Alternatively, color_by can be set to vector of valid R colors, either of length one (recycled for all cells) or of length ncol(x), which will then be used to color cells in the phase graph.
color.alpha	An integer scalar giving the transparency of colored cells. Possible values are between 0 (fully transparent) and 1.0 (opaque).
colors.velocity	r, colors.expression
	Character vectors specifying the color ranges used for mapping velocities and expression values. The defaults are RColorBrewer::brewer.pal(11, "RdYlGn") for the velocities and viridisLite::viridis(11) for the expression values.
max.abs.velo	A numeric scalar greater than zero giving the maximum absolute velocity to limit the color scale for the "velocity" graph.

Details

Please note that plotVelocity will modify parameters of the current graphics device using layout and par, in order to create the layout for the generated graph panels.

Value

A patchwork object with the plots selected by which.plot for the genes in genes, arranged in a grid according to genes.per.row.

Author(s)

Michael Stadler

See Also

scvelo, to generate x, brewer.pal and viridis for creation of color palettes, packages **ggplot2** and **patchwork** used to generate and arrange the plots.

Examples

```
library(scuttle)
set.seed(42)
sce1 <- mockSCE(ncells = 100, ngenes = 500)
sce2 <- mockSCE(ncells = 100, ngenes = 500)
datlist <- list(X=counts(sce1), spliced=counts(sce1), unspliced=counts(sce2))
out1 <- scvelo(datlist, mode = "steady_state")
out2 <- scvelo(datlist, mode = "dynamical")
plotVelocity(out1, c("Gene_0031", "Gene_0268"))
plotVelocity(out2, c("Gene_0031", "Gene_0268"))</pre>
```

plotVelocityStream Velocity stream plot in low-dimensional space

Description

Plot velocities embedded into low-dimensional space as a stream plot. Stream lines are lines that follow the gradient in the velocity field and illustrate paths that cells could follow based on observed RNA velocities.

Usage

```
plotVelocityStream(
 sce,
 embedded,
 use.dimred = 1,
 color_by = "#444444",
 color.alpha = 0.2,
 grid.resolution = 60,
 scale = TRUE,
 stream.L = 10,
 stream.min.L = 0,
 stream.res = 4,
 stream.width = 8,
 color.streamlines = FALSE,
arrow.angle = 8,
 arrow.length = 0.8
)
```

Arguments

sce	A SingleCellExperiment object containing low-dimensional coordinates, e.g., after t-SNE, in its reducedDims.
embedded	A low-dimensional projection of the velocity vectors into the embedding of sce. This should be of the same dimensions as sce and is typically produced by embedVelocity.
use.dimred	String or integer scalar specifying the reduced dimensions to retrieve from sce.
color_by	A character scalar specifying a column in colData(sce) to color cells in the phase graph. Alternatively, color_by can be set to a valid R color to be used to color cells.
color.alpha	An integer scalar giving the transparency of colored cells. Possible values are between 0 (fully transparent) and 1.0 (opaque).
grid.resolution	
	Integer scalar specifying the resolution of the grid, in terms of the number of grid intervals along each axis.

scale	Logical scalar indicating whether the averaged vectors should be scaled by the grid resolution.
stream.L	Integer scalar giving the typical length of a streamline low-dimensional space units.
stream.min.L	A numeric scalar with the minimum length of segments to be shown.
stream.res	Numeric scalar specifying the resolution of estimated streamlines (higher numbers increase smoothness of lines but also the time for computation).
stream.width	A numeric scalar controlling the width of streamlines.
color.streamlines	
	Logical scalar. If TRUE streamlines will be colored by local velocity. Arrows cannot be shown in that case.
color.streamlines.map	
	A character vector specifying the color range used for mapping local velocities
	to streamline colors. The default is viridisLite::viridis(11).
arrow.angle, arrow.length	
	Numeric scalars giving the angle and length of arrowheads.

Details

grid.resolution and scale are passed to gridVectors, which is used to summarized the velocity vectors into an initial grid. A full regular grid is computed from that and used in geom_streamline to calculate streamlines. The following arguments are passed to the arguments given in parenthesis of geom_streamline: stream.L (L), stream.res (res), stream.min.L (min.L), arrow.angle (arrow.angle) and arrow.length (arrow.length). Streamlines are computed by simple integration with a forward Euler method, and stream.L and stream.res are used to compute the number of steps and the time interval between steps for the integration. stream.width is multiplied with ..step.. estimated by geom_streamline to control the width of streamlines.

Value

A ggplot2 object with the streamline plot.

Author(s)

Michael Stadler

See Also

gridVectors used to summarize velocity vectors into a grid (velocity field), the **ggplot2** package used for plotting, geom_streamline in package **metR** used to calculate and add streamlines from the RNA velocity field to the plot, viridis for creation of color palettes.

Examples

```
library(scuttle)
set.seed(42)
sce1 <- mockSCE(ncells = 100, ngenes = 500)
sce2 <- mockSCE(ncells = 100, ngenes = 500)</pre>
```

```
datlist <- list(X=counts(sce1), spliced=counts(sce1), unspliced=counts(sce2))
out <- scvelo(datlist, mode = "dynamical")
em <- embedVelocity(reducedDim(out, 1), out)[,1:2]
# https://github.com/kevinrue/velociraptor/issues/90
if (!basilisk.utils::isMacOSX()){
plotVelocityStream(out, em)
plotVelocityStream(out, em, color.streamlines = TRUE)
}</pre>
```

scvelo

RNA velocity with scVelo

Description

Perform RNA velocity calculations with the scVelo package.

Usage

```
scvelo(x, ...)
## S4 method for signature 'ANY'
scvelo(
 х,
  subset.row = NULL,
  sf.X = NULL,
  sf.spliced = NULL,
  sf.unspliced = NULL,
  use.theirs = FALSE,
 mode = c("steady_state", "deterministic", "stochastic", "dynamical"),
  scvelo.params = list(),
  dimred = NULL,
  ncomponents = 30,
 BPPARAM = SerialParam(),
  BSPARAM = bsparam()
)
## S4 method for signature 'SummarizedExperiment'
scvelo(
  х,
  ...,
  assay.X = "counts",
  assay.spliced = "spliced",
```

scvelo

```
assay.unspliced = "unspliced"
)
## S4 method for signature 'SingleCellExperiment'
scvelo(x, ..., sf.X = sizeFactors(x), dimred = NULL, use.dimred = NULL)
```

Arguments

x	A named list of three matrices of the same dimensions where genes are in rows and cells are in columns. The list should contain "spliced" and "unspliced" entries containing spliced and unspliced counts, respectively. It should also con- tain an "X" entry containing the "usual" count matrix, see details below. Alternatively, a SummarizedExperiment object containing three such matrices among its assays.
	For the generic, further arguments to pass to specific methods. For the Summa- rizedExperiment and SingleCellExperiment methods, further arguments to pass to the ANY method.
subset.row	A character, integer or logical vector specifying the genes to use for the velocity calculations. Defaults to all genes.
sf.X	A numeric vector containing size factors for usual count matrix. Defaults to librarySizeFactors on the "X" matrix in x.
sf.spliced	A numeric vector containing size factors for the spliced counts for each cell. Defaults to librarySizeFactors on the "spliced" matrix in x.
sf.unspliced	A numeric vector containing size factors for the unspliced counts for each cell. Defaults to librarySizeFactors on the "unspliced" matrix in x.
use.theirs	Logical scalar indicating whether scVelo 's gene filtering and normalization should be used.
mode	String specifying the method to use to estimate the transcriptional dynamics.
scvelo.params	List of lists containing arguments for individual scVelo functions, see details below.
dimred	A low-dimensional representation of the cells with number of rows equal to the number of cells in x, used to find the nearest neighbors.
ncomponents	Numeric scalar indicating the number of principal components to obtain. Only used if use.theirs=FALSE and dimred=NULL.
BPPARAM	A BiocParallelParam object specifying whether the PCA calculations should be parallelized. Only used if use.theirs=FALSE and dimred=NULL.
BSPARAM	A BiocSingularParam object specifying which algorithm should be used to per- form the PCA. Only used if use.theirs=FALSE and dimred=NULL.
assay.X	An integer scalar or string specifying the assay of x containing the usual count matrix.
assay.spliced	An integer scalar or string specifying the assay of x containing the spliced counts.
assay.unspliced	
	An integer scalar or string specifying the assay of x containing the unspliced counts.

use.dimred String naming the entry of reducedDims(x) to use for nearest neighbor calculations. Ignored if dimred is supplied.

Details

This function uses the **scVelo** Python package (https://pypi.org/project/scvelo/) for RNA velocity calculations. The main difference from the original **velocyto** approach is that the dynamical model of **scVelo** does not rely on the presence of observed steady-state populations, which should improve the reliability of the velocity calculations in general applications.

For consistency with other Bioconductor workflows, we perform as many standard steps in R as we can before starting the velocity calculations with **scVelo**. This involves:

- 1. Size factor-based normalization with sf.* values and normalizeCounts. For "X", log-transformation is performed as well, while for the others, only scaling normalization is performed.
- 2. Subsetting all matrices to subset.row, most typically to a subset of interest, e.g., highly variable genes. Note that, if set, any subsetting is done *after* normalization so that library sizes are correctly computed.
- 3. If dimred=NULL, the PCA step on the log-expression values derived from the "X" matrix, using the specified BSPARAM to obtain the first ncomponents PCs.

This allows us to guarantee that, for example, the log-expression matrix of HVGs or the PCA coordinates are the same as that used in other applications like clustering or trajectory reconstruction.

Nonetheless, one can set use.theirs=TRUE to directly use the entire **scVelo** normalization and filtering pipeline. This ignores all of the size factors arguments (sf.*), all of the PCA-related arguments (ncomponents, BSPARAM) and subset.row. However, if a low-dimensionality result is supplied via dimred or use.dimred, the **scVelo** PCA will always be omitted.

Upon first use, this function will instantiate a conda environment containing the **scVelo** package. This is done via the **basilisk** package - see the documentation for that package for trouble-shooting.

Value

A SingleCellExperiment is returned containing the output of the velocity calculations. Of particular interest are:

- the velocity_pseudotime field in the colData, containing the velocity pseudotime for each cell.
- the velocity entry of the assays, containing the velocity vectors for each cell.

The output will always have number of columns equal to the number of cells supplied in x, though the number of rows will depend on whether any subsetting (if subset.row is supplied) or feature selection (if use.theirs=TRUE) was performed.

Comments on the three matrices

Strictly speaking, only the spliced and unspliced matrices are necessary for the velocity calculations. However, it is often the case that the spliced matrix is not actually the same as a "usual" count matrix (e.g., generated by summing counts across all exons for all mapped genes). This is due to differences in the handling of ambiguous reads that map across exon-intron boundaries, or to genomic regions scvelo

that can be either exonic or intronic depending on the isoform; the spliced count matrix is more likely to exclude such reads.

We request the usual count matrix as the "X" entry of x to ensure that the PCA and nearest neighbor detection in **scVelo** are done on the same data as that used in other steps of the large analysis (e.g., clustering, visualization, trajectory reconstruction). In practice, if the usual count matrix is not available, one can often achieve satisfactory results by simply re-using the spliced count matrix as both the "X" and "spliced" entries of x.

Note that if reduced dimensions are supplied in dimred, any "X" entry is only used to create the AnnData object and is not used in any actual calculations.

Additional arguments to Python

Additional arguments to **scVelo** functions are provided via scvelo.params. This is a named list where each entry is named after a function and is itself a named list of arguments for that function. The following function names are currently recognized:

- "filter_and_normalize", for gene selection and normalization. This is not used unless use.theirs=TRUE.
- "moments", for PCA and nearest neighbor detection. The PCA is not performed if dimred or use.dimred is already supplied.
- "recover_dynamics"
- "velocity"
- "velocity_graph"
- "velocity_pseudotime"
- "latent_time"
- "velocity_confidence"

See the **scVelo** documentation for more details about the available arguments and the examples below for a syntax example.

Supported operating systems and architectures

scVelo dependencies are pinned in a Conda environment to ensure reproducibility.

Differences in packages and versions available from Conda require different environments for different operating systems and architectures. **basilisk.utils** is used to determine the operating system and architecture of the computer used to run scvelo(), using to the appropriate Conda environment.

As of the latest velociraptor update (24 May 2024):

- All environments tqdm and ipywidgets are installed to suppress the message "Unable to create progress bar".
- Linux scVelo v0.3.2 from conda-forge is used. This is the latest version available to date. libtiff is pinned to v4.5.1 and pillow is pinned to v10.0.0 (https://github.com/conda-forge/ libtiff-feedstock/issues/104#issuecomment-2375893029), scipy is pinned to v1.13.1 (https://github.com/theislab/scvelo/issues/1260).

- Linux AArch64 scVelo v0.3.2 from conda-forge is used. This is the latest version available to date. libtiff is pinned to v4.5.1 and pillow is pinned to v10.0.0 (https://github.com/ conda-forge/libtiff-feedstock/issues/104#issuecomment-2375893029), scipy is pinned to v1.13.1 (https://github.com/theislab/scvelo/issues/1260). Please note that this environment has not been validated yet; it is derived from the environment from Linux (above) and requires additional testing to identify a working environment before pinning all the packages and versions in the environment.
- **macOS** scVelo v0.3.2 from conda-forge is used. This is the latest version available to date. scipy is pinned to v1.13.1 (https://github.com/theislab/scvelo/issues/1260).
- **macOS ARM scVelo** v0.3.2 from conda-forge is used. This is the latest version available to date. **scipy** is pinned to v1.13.1 (https://github.com/theislab/scvelo/issues/1260).
- Windows scVelo v0.2.5 from bioconda is used. Later versions of scVelo depend on jaxlib which is not supported on Windows (https://github.com/google/jax/issues/438). matplotlib is pinned to v3.6.3 (https://github.com/scverse/scanpy/issues/2411), pandas is pinned to v1.5.2 (https://stackoverflow.com/questions/76234312/importerror-cannot-import-name-is-categor and numpy is pinned to v1.21.1 (https://github.com/theislab/scvelo/issues/1109).

Author(s)

Aaron Lun, Charlotte Soneson, Kevin Rue-Albrecht

References

Bergen, V., Lange, M., Peidli, S. et al. Generalizing RNA velocity to transient cell states through dynamical modeling. Nat Biotechnol 38, 1408–1414 (2020). https://doi.org/10.1038/s41587-020-0591-3

Examples

```
# Using mock data to demonstrate the process:
library(scuttle)
sce1 <- mockSCE()
sce2 <- mockSCE()
spliced <- counts(sce1)
unspliced <- counts(sce2)</pre>
```

out <- scvelo(list(X=spliced, spliced=spliced, unspliced=unspliced))</pre>

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