Package 'cellTree'

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Title Inference and visualisation of Single-Cell RNA-seq data as a hierarchical tree structure

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Description This packages computes a Latent Dirichlet Allocation (LDA) model of single-cell RNA-seq data and builds a compact tree modelling the relationship between individual cells over time or space.

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Description

This packages computes a Latent Dirichlet Allocation (LDA) model of single-cell RNA-seq data and build a compact tree modelling the relationship between individual cells over time or space.

Details

A typical use-case will require you to run [compute.lda](#page-6-1) on your expression data, to fit an LDA model, followed by [compute.backbone.tree](#page-2-1) to generate a tree structure from the LDA model.

Plotting functions [ct.plot.grouping](#page-9-1) and [ct.plot.topics](#page-11-1) will then help you plot the tree structure with the desired annotations, while function [cell.ordering.table](#page-2-2) will output an ordered table of cells, ranked by their position in the tree.

To get further information on each topic, you can run Gene Ontology enrichment tests, using [com](#page-5-1)[pute.go.enrichment,](#page-5-1) plot the result tables as a graph using [ct.plot.go.dag](#page-8-1) or render it with LaTeX, using [go.results.to.latex.](#page-13-1)

cell.ordering.table *Ranking of cells according to backbone tree structure*

Description

Produces a table of input cells ranked by their position in the backbone tree.

Usage

```
cell.ordering.table(b.tree, write.to.tex.file = NULL)
```
Arguments

b.tree An [igraph](#page-0-0) backbone tree, as returned by [compute.backbone.tree](#page-2-1). write.to.tex.file

Boolean (optional). If not NULL, outputs LaTeX version of table to file write.to.tex.file.

Value

List of all cells, ranked by position in backbone tree, along with topic information.

Examples

```
# Load pre-computed LDA model for skeletal myoblast RNA-Seq data from HSMMSingleCell package:
data(HSMM_lda_model)
```

```
# Recover sampling time (in days) for each cell:
library(HSMMSingleCell)
data(HSMM_sample_sheet)
days.factor = HSMM_sample_sheet$Hours
days = as.numeric(levels(days.factor))[days.factor]
```

```
# Compute near-optimal backbone tree:
b.tree = compute.backbone.tree(HSMM_lda_model, days)
# Load pre-computed LDA model for skeletal myoblast RNA-Seq data from HSMMSingleCell package:
temp.output = tempfile()
cell.ordering.table(b.tree, write.to.tex.file = temp.output)
```
compute.backbone.tree *Backbone Tree construction*

Description

Builds a 'backbone tree' from a fitted LDA model.

```
compute.backbone.tree(lda.results, grouping = NULL,
  start.group.label = NULL, absolute.width = 0, width.scale.factor = 1.2,
 outlier.tolerance.factor = 0.1, rooting.method = NULL, only.mst = FALSE,
 grouping.colors = NULL, merge.sequential.backbone = FALSE)
```


Details

In order to easily visualise the structural and temporal relationship between cells, we introduced a special type of tree structure dubbed 'backbone tree', defined as such:

Considering a set of vertices V and a distance function over all pairs of vertices: $d: V \times V \to R^+$, we call *backbone tree* a graph, T with backbone B, such that:

- T is a tree with set of vertices V and edges E .
- B is a tree with set of vertices $V_B \subseteq V$ and edges $E_B \subseteq E$.
- All 'vertebrae' vertices of $T: v \in V \setminus V_B$ are connected by a single edge to the closest vertex in the set of backbone vertices $v_B^* \in V_B$. I.e: $v_B^* = argmin_{v_B \in V_B} d(v_B, v)$.
- For all vertices in $V \setminus V_B$ are less than distance δ to a vertex in the backbone tree B: $\forall v \in$ $V \setminus V_B$, $\exists v_B \in V_B$ such that $d(v, v_b) \leq \delta$.

In this instance, we relax the last condition to cover only 'most' non-backbone vertices, allowing for a variable proportion of outliers at distance $\geq \delta$ from any vertices in V_B .

We can then define the 'optimal' backbone tree to be a backbone tree such that the sum of weighted edges in the backbone subtree E_B is minimal. Finding such a tree can be easily shown to be NP-Complete (by reduction to the Vertex Cover problem), but we developed a fast heuristic relying on Minimum Spanning Tree to produce a reasonable approximation.

The resulting quasi-optimal backbone tree (simply referred to as 'the' backbone tree thereafter) gives a clear hierarchical representation of the cell relationship: the objective function puts pressure on finding a (small) group of prominent cells (the backbone) that are good representatives of major steps in the cell evolution (in time or space), while remaining cells are similar enough to their closest representative for their difference to be ignored. Such a tree provides a very clear visualisation of overall cell differentiation paths (including potential differentiation into sub-types).

Value

A [igraph](#page-0-0) object with either a minimum rooted spanning-tree (if only.mst is TRUE) or a quasioptimal backbone tree connecting all input cells. Cell topic distribution, distances and branch order are added as vertex/edge/graph attributes.

Examples

Load pre-computed LDA model for skeletal myoblast RNA-Seq data from HSMMSingleCell package: data(HSMM_lda_model)

```
# Recover sampling time (in days) for each cell:
library(HSMMSingleCell)
data(HSMM_sample_sheet)
days.factor = HSMM_sample_sheet$Hours
days = as.numeric(levels(days.factor))[days.factor]
```

```
# Compute near-optimal backbone tree:
b.tree = compute.backbone.tree(HSMM_lda_model, days)
# Plot resulting tree with sampling time as a vertex group colour:
ct.plot.grouping(b.tree)
```
compute.go.enrichment *Gene Ontology enrichment analysis*

Description

Computes enrichment scores for Gene Ontology terms associated with genes in each topic.

Usage

```
compute.go.enrichment(lda.results, go.db, ontology.type = "BP",
  reformat.gene.names = FALSE, bonferroni.correct = TRUE,
 p.val.threshold = if (bonferroni.correct) 0.05 else 0.01,
 go.score.class = "weight01Score", dag.file.prefix = FALSE)
```
Arguments

Value

Returns a named list object with ranked tables of significantly enriched GO terms for each topic ('all'), terms that only appear in each topic ('unique') and terms that appear in less than half of the other topics ('rare'). In addition the list object contains an [igraph](#page-0-0) object with the full GO DAG, annotated with each term's p-value and the significance threshold adjusted for multiple testing (Bonferroni method).

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Examples

Load pre-computed LDA model for skeletal myoblast RNA-Seq data from HSMMSingleCell package: data(HSMM_lda_model)

Load GO mapping database for 'homo sapiens': library(org.Hs.eg.db) # Compute Cellular Component GO enrichment sets for each topic: go.results = compute.go.enrichment(HSMM_lda_model, org.Hs.eg.db, ontology.type="CC", bonferroni.correct=TRUE, p. # Print table of terms that are only significantly enriched in each topic: print(go.results\$unique)

compute.lda *LDA model inference*

Description

This function fits a Latent Dirichlet Allocation (LDA) to single-cell RNA-seq data.

Usage

```
compute.lda(data, method = "maptpx", k.topics = if (method == "maptpx") 2:15
 else 4, log.scale = TRUE, sd.filter = 0.5, tot.iter = if (method ==
  "Gibbs") 200 else 1e+06, tol = if (method == "maptpx") 0.05 else 10^-5)
```


Details

Latent Dirichlet allocation (LDA) is a generative model that allows sets of observations to be explained by unobserved groups (topics) that explain why some parts of the data are similar [Blei, 2003]. Each topic is modelled as a (Dirichlet) distribution over observations and each set of observations is also modelled as a (Dirichlet) distribution over topics. In lieu of the traditional NLP context of word occurence counts in documents, our model uses RNA-seq observation counts in single cells. Three separate LDA inference methods can be used at the moment:

- Gibbs uses Collapsed Gibbs Sampling method (implemented by Xuan-Hieu Phan and coauthors in the topicmodels package [Phan, 2008]) to infer the parameters of the Dirichlet distributions for a given number of topics. It gives high accuracy but is very time-consuming to run on large number of cells and genes.
- VEM uses Variational Expectation-Maximisation (as described in [Hoffman, 2010]). This method tends to converge faster than Gibbs collapsed sampling, albeit with lower accuracy.
- maptpx uses the method described in [Taddy, 2011] and implemented in package maptpx to estimate the parameters of the topic model for increasing number of topics (using previous estimates as a starting point for larger topic numbers). The best model (/number of topics) is selected based on Bayes factor over the Null model. Although potentially less accurate, this method provides the fastest way to train and select from a large number of models, when the number of topics is not well known.

When in doubt, the function can be ran with its default parameter values and should produce a usable LDA model in reasonable time (using the 'maptpx' inference method). The model can be further refined for a specific number of topics with slower methods. While larger models (using large number of topics) might fit the data well, there is a high risk of overfitting and it is recommended to use the smallest possible number of topics that still explains the observations well. Anecdotally, a typical number of topics for cell differentiation data (from pluripotent to fully specialised) would seem to be around 4 or 5.

Value

A LDA model fitted for data, of class [LDA-class](#page-0-0) (for methods 'Gibbs' or 'VEM') or [topics](#page-0-0) (for 'maptpx')

References

- Blei, Ng, and Jordan. "Latent dirichlet allocation." the Journal of machine Learning research 3 (2003): 993-1022.
- Hoffman, Blei and Bach (2010). "Online Learning for Latent Dirichlet Allocation." In J Lafferty, CKI Williams, J Shawe-Taylor, R Zemel, A Culotta (eds.), Advances in Neural Information Processing Systems 23, pp. 856-864. MIT Press, Cambridge, MA.
- Hornik and Grün. "topicmodels: An R package for fitting topic models." Journal of Statistical Software 40.13 (2011): 1-30.
- Phan, Nguyen and Horiguchi. "Learning to classify short and sparse text & web with hidden topics from large-scale data collections." Proceedings of the 17th international conference on World Wide Web. ACM, 2008.
- Taddy. "On estimation and selection for topic models." arXiv preprint arXiv:1109.4518 (2011).

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See Also

[LDA,](#page-0-0) [topics,](#page-0-0) [LDA_Gibbscontrol-class,](#page-0-0) [CTM_VEMcontrol-class](#page-0-0)

Examples

```
# Load skeletal myoblast RNA-Seq data from HSMMSingleCell package:
library(HSMMSingleCell)
data(HSMM_expr_matrix)
# Run LDA inference using 'maptpx' method for k = 4:
lda.results = compute.lda(HSMM_expr_matrix, k.topics=4, method="maptpx")
# Run LDA inference using 'maptpx' method for number of topics k = 3 to 6:
lda.results = compute.lda(HSMM_expr_matrix, k.topics=3:6, method="maptpx")
# Run LDA inference using 'Gibbs' [collapsed sampling] method for number of k = 4 topics:
lda.results = compute.lda(HSMM_expr_matrix, k.topics=4, method="Gibbs")
```
ct.plot.go.dag *Gene Ontology enrichment sets plotting*

Description

Plots DAG of significantly enriched terms for all topics, along with ancestor nodes.

Usage

```
ct.plot.go.dag(go.results, up.generations = 2, only.topics = NULL,
  file.output = NULL, p.val.threshold = go.results$adjusted.p.threshold,
  only.unique = FALSE, topic.colors = rainbow(length(go.results$results)))
```


Value

An [igraph](#page-0-0) object with the annotated GO DAG.

Examples

```
# Load pre-computed LDA model for skeletal myoblast RNA-Seq data from HSMMSingleCell package:
data(HSMM_lda_model)
```

```
# Load GO mapping database for 'homo sapiens':
library(org.Hs.eg.db)
```

```
# Compute GO enrichment sets for each topic:
go.results = compute.go.enrichment(HSMM_lda_model, org.Hs.eg.db, bonferroni.correct=TRUE)
```

```
go.dag.subtree = ct.plot.go.dag(go.results, up.generations = 2)
```
ct.plot.grouping *Plot cell tree with grouping information*

Description

Plots a backbone tree (or MST) that was computed with [compute.backbone.tree](#page-2-1), displaying each cell's grouping.

Usage

```
ct.plot.grouping(tree, file.output = NULL, show.labels = FALSE,
  force.recompute.layout = FALSE, height = 20, width = 10,
  vertebrae.distance = 0, backbone.vertex.size = 0, vert.vertex.size = 0)
```


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```
backbone.vertex.size
                  Numeric (optional). Diameter (in pixels) of backbone and vertebrae cell ver-
                  tices.
vert.vertex.size
                  Numeric (optional). Diameter (in pixels) of backbone and vertebrae cell ver-
                  tices.
```
Value

An updated [igraph](#page-0-0) object with x and y vertex coordinate attributes.

Examples

```
# Load pre-computed LDA model for skeletal myoblast RNA-Seq data from HSMMSingleCell package:
data(HSMM_lda_model)
```

```
# Recover sampling time (in days) for each cell:
library(HSMMSingleCell)
data(HSMM_sample_sheet)
days.factor = HSMM_sample_sheet$Hours
days = as.numeric(levels(days.factor))[days.factor]
```

```
# Compute near-optimal backbone tree:
b.tree = compute.backbone.tree(HSMM_lda_model, days)
# Plot resulting tree with sampling time as a vertex group colour:
ct.plot.grouping(b.tree)
```
ct.plot.heatmap *Gene Expression Heatmap*

Description

Plots a heatmap of gene expression, with cells ordered according to the structure computed by [compute.backbone.tree](#page-2-1).

Usage

```
ct.plot.heatmap(data, b.tree, log.scale = TRUE, sd.filter = 0.7,
  reorder.genes = TRUE)
```


Value

data object reordered according to the backbone tree, such as used to plot the heatmap.

Examples

```
# Load pre-computed LDA model for skeletal myoblast RNA-Seq data from HSMMSingleCell package:
data(HSMM_lda_model)
```

```
# Recover sampling time (in days) for each cell:
library(HSMMSingleCell)
data(HSMM_sample_sheet)
days.factor = HSMM_sample_sheet$Hours
days = as.numeric(levels(days.factor))[days.factor]
# Compute near-optimal backbone tree:
b.tree = compute.backbone.tree(HSMM_lda_model, days)
# Plot heatmap:
data(HSMM_expr_matrix)
```

```
ct.plot.heatmap(HSMM_expr_matrix[1:2000,], b.tree, reorder.genes=FALSE)
```
ct.plot.topics *Plot cell tree with topic distributions*

Description

Plots a backbone tree (or MST) that was computed with [compute.backbone.tree](#page-2-1), displaying each cell's topic distribution as a pie chart.

Usage

```
ct.plot.topics(tree, file.output = NULL, show.labels = FALSE,
  force.recompute.layout = FALSE, height = 20, width = 10,
  vertebrae.distance = 0, backbone.vertex.size = 0, vert.vertex.size = 0)
```


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```
backbone.vertex.size, vert.vertex.size
                 Numeric (optional). Diameter (in pixels) of backbone and vertebrae cell ver-
                 tices.
```
Value

An updated [igraph](#page-0-0) object with x and y vertex coordinate attributes.

Examples

Load pre-computed LDA model for skeletal myoblast RNA-Seq data from HSMMSingleCell package: data(HSMM_lda_model)

```
# Recover sampling time (in days) for each cell:
library(HSMMSingleCell)
data(HSMM_sample_sheet)
days.factor = HSMM_sample_sheet$Hours
days = as.numeric(levels(days.factor))[days.factor]
```

```
# Compute near-optimal backbone tree:
b.tree = compute.backbone.tree(HSMM_lda_model, days)
# Plot resulting tree with sampling time as a vertex group colour:
ct.plot.grouping(b.tree)
```
get.cell.dists *Cell Pairwise-Distance Matrix*

Description

Computes the pairwise distance between cells, based on the topic histograms form a fitted LDA model.

Usage

```
get.cell.dists(lda.results)
```
Arguments

lda.results A fitted LDA model, as returned by [compute.lda](#page-6-1)

Details

Distances between histograms are computed using the Chi-square distance

Value

A square matrix of pairwise distance between cells in the input model.

Examples

Load pre-computed LDA model for skeletal myoblast RNA-Seq data from HSMMSingleCell package: data(HSMM_lda_model)

```
# Compute cell pairwise distances:
b.tree = get.cell.dists(HSMM_lda_model)
```
go.results.to.latex *LaTeX output for Gene Ontology Enrichment results*

Description

Outputs or writes result tables of Gene Ontology enrichment testing in LaTeX format.

Usage

```
go.results.to.latex(go.results, tex.file.name = "go_terms.tex",
  topic.colors = rainbow(length(go.results$all)))
```
Arguments

Value

GO enrichment results in LaTeX format

Examples

Load pre-computed LDA model for skeletal myoblast RNA-Seq data from HSMMSingleCell package: data(HSMM_lda_model)

```
# Load GO mapping database for 'homo sapiens':
library(org.Hs.eg.db)
# Compute GO enrichment sets for each topic:
go.results = compute.go.enrichment(HSMM_lda_model, org.Hs.eg.db, bonferroni.correct=TRUE)
# Output LaTeX tables for GO results
latex.table = go.results.to.latex(go.results, tex.file.name='go_results.tex')
# [Optional] compile LaTeX to PDF (need to have LaTeX binaries installed):
library('tools')
texi2pdf('go_results.tex')
library('Biobase')
openPDF('go_results.pdf')
```


Description

Pre-computed LDA model for HSMMSingleCell data

Usage

HSMM_lda_model

Format

An object of class list of length 3.

Value

LDA model obtained by running [compute.lda](#page-6-1) on the HSMMSingleCell package's data. This demo model can be used with all functions in this package that require a fitted LDA model, such as [compute.backbone.tree](#page-2-1).

See examples for [compute.lda](#page-6-1) for more details.

See Also

[compute.lda](#page-6-1)

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