Package ‘DDRTree’
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Title Learning Principal Graphs with DDRTree
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Description Provides an implementation of the framework of reversed graph embedding (RGE) which projects data into a reduced dimensional space while constructs a principal tree which passes through the middle of the data simultaneously. DDRTree shows superiority to alternatives (Wishbone, DPT) for inferring the ordering as well as the intrinsic structure of the single cell genomics data. In general, it could be used to reconstruct the temporal progression as well as bifurcation structure of any datatype.
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Perform DDRTree construction

This is an R and C code implementation of the DDRTree algorithm from Qi Mao, Li Wang et al.

Qi Mao, Li Wang, Steve Goodison, and Yijun Sun. Dimensionality Reduction via Graph Structure Learning. The 21st ACM SIGKDD Conference on Knowledge Discovery and Data Mining (KDD’15), 2015 http://dl.acm.org/citation.cfm?id=2783309

to perform dimension reduction and principal graph learning simultaneously. Please cite this package and KDD’15 paper if you found DDRTree is useful for your research.

Usage

DDRTree(X, dimensions = 2, initial_method = NULL, maxIter = 20, sigma = 0.001, lambda = NULL, ncenter = NULL, param.gamma = 10, tol = 0.001, verbose = F, ...)

Arguments

X a matrix with $D \times N$ dimension which is needed to perform DDRTree construction

dimensions reduced dimension

initial_method a function to take the data transpose of X as input and then output the reduced dimension. row number should not larger than observation and column number should not be larger than variables (like isomap may only return matrix on valid sample sets). Sample names of returned reduced dimension should be preserved.

maxIter maximum iterations

sigma bandwidth parameter

lambda regularization parameter for inverse graph embedding

ncenter number of nodes allowed in the regularization graph

param.gamma regularization parameter for k-means (the prefix of 'param' is used to avoid name collision with gamma)

tol relative objective difference

verbose emit extensive debug output

... additional arguments passed to DDRTree
**DDRTree**

**Value**

a list with $W$, $Z$, stree, $Y$, history $W$ is the orthogonal set of $d$ (dimensions) linear basis vector $Z$ is the reduced dimension space stree is the smooth tree graph embedded in the low dimension space $Y$ represents latent points as the center of $Z$

**Introduction**

The unprecedented increase in big-data causes a huge difficulty in data visualization and downstream analysis. Conventional dimension reduction approaches (for example, PCA, ICA, Isomap, LLE, etc.) are limited in their ability to explicitly recover the intrinsic structure from the data as well as the discriminative feature representation, both are important for scientific discovery. The DDRTree algorithm is a new algorithm to perform the following three tasks in one setting:

1. Reduce high dimension data into a low dimension space
2. Recover an explicit smooth graph structure with local geometry only captured by distances of data points in the low dimension space.
3. Obtain clustering structures of data points in reduced dimension

**Dimensionality reduction via graph structure learning**

Reverse graph embedding is previously applied to learn the intrinsic graph structure in the original dimension. The optimization of graph inference can be represented as:

$$\min_{f_{g} \in \mathcal{F}} \min_{\{z_1, \ldots, z_M\}} \sum_{(V_i, V_j) \in \mathcal{E}} b_{i,j} ||f_{g}(z_i) - f_{g}(z_j)||^2$$

where $f_{g}$ is a function to map the intrinsic data space $\mathcal{Z} = \{z_1, \ldots, z_M\}$ back to the input data space (reverse embedding) $\mathcal{X} = \{x_1, \ldots, x_N\}$. $V_i$ is the vertex of the intrinsic undirected graph $\mathcal{G} = (\mathcal{V}, \mathcal{E})$. $b_{i,j}$ is the edge weight associates with the edge set $\mathcal{E}$. In order to learn the intrinsic structure from a reduced dimension, we need also to consider a term which includes the error during the learning of the intrinsic structure. This strategy is incorporated as the following:

$$\min_{\mathcal{G} \in \hat{\mathcal{G}}_b} \min_{f_{g} \in \mathcal{F}} \sum_{i=1}^{N} ||x_i - f_{g}(z_i)||^2 + \frac{\lambda}{2} \sum_{(V_i, V_j) \in \mathcal{E}} b_{i,j} ||f_{g}(z_i) - f_{g}(z_j)||^2$$

where $\lambda$ is a non-negative parameter which controls the tradeoff between the data reconstruction error and the reverse graph embedding.

**Dimensionality reduction via learning a tree**

The general framework for reducing dimension by learning an intrinsic structure in a low dimension requires a feasible set $\hat{\mathcal{G}}_b$ of graph and a mapping function $f_{g}$. The algorithm uses minimum spanning tree as the feasible tree graph structure, which can be solved by Kruskal’ algorithm. A linear projection model $f_{g}(z) = Wz$ is used as the mapping function. Those setting results in the
following specific form for the previous framework:

$$\min_{W,Z,B} \sum_{i=1}^{N} ||x_i - Wz_i||^2 + \frac{\lambda}{2} \sum_{i,j} b_{i,j} ||Wz_i - Wz_j||^2$$

where $W = [w_1, ..., w_d] \in \mathcal{R}^{D \times d}$ is an orthogonal set of $d$ linear basis vectors. We can group tree graph $B$, the orthogonal set of linear basis vectors and projected points in reduced dimension $W$, $Z$ as two groups and apply alternative structure optimization to optimize the tree graph. This method is defined as DRtree (Dimension Reduction tree) as discussed by the authors.

**Discriminative dimensionality reduction via learning a tree**

In order to avoid the issues where data points scattered into different branches (which leads to lose of cluster information) and to incorporate the discriminative information, another set of points $\{y_k\}_{K=1}^{K}$ as the centers of $\{z_i\}_{N=1}^{N}$ can be also introduced. By so doing, the objective functions of K-means and the DRtree can be simultanously minimized. The author further proposed a soft partition method to account for the limits from K-means and proposed the following objective function:

$$\min_{W,Z,Y,B,R} \sum_{i=1}^{N} ||x_i - Wz_i||^2 + \frac{\lambda}{2} \sum_{k,k'} b_{k,k'} ||W_{k} - W_{k'}||^2 + \gamma \left[ \sum_{k=1}^{K} \sum_{i=1}^{N} r_{i,k} ||z_i - y_k||^2 + \sigma \Omega(R) \right]$$

$$s.t. \ W^T W = I, B \in B, \sum_{k=1}^{K} r_{i,k} = 1, r_{i,k} \leq 0, \forall i, \forall k$$

where $R \in \mathcal{R}^{N \times N}$, $\Omega(R) = \sum_{i=1}^{N} \sum_{k=1}^{K} r_{i,k} \log r_{i,k}$ is the negative entropy regularization which transforms the hard assignments used in K-means into soft assignments and $\sigma > 0$ is the reg-ularization parameter. Alternative structure optimization is again used to solve the above problem by separately optimize each group $W$, $Z$, $Y$, $B$, $R$ until convergence.

**The actual algorithm of DDRTree**

1. **Input**: Data matrix $X$, parameters $\lambda, \sigma, \gamma$
2. Initialize $Z$ by PCA
3. $K = N$, $Y = Z$
4. repeat:
5. $d_{k,k'} = ||y_k - y_{k'}||^2, \forall k, \forall k'$
6. Obtain $B$ via Kruskal’s algorithm
7. $L = diag(B1) - B$
8. Compute $R$ with each element
9. $\tau = diag(1^T R)$
10. $Q = \frac{1}{1+\gamma}\left[I + R(\frac{1+\gamma}{\gamma}L + \tau)^{-1}\right]$
11. $C = XQX^T$
12. Perform eigen-decomposition on $C$ such that $C = U \wedge U^T$ and $diag(\wedge)$ is sorted in a descending order
13. $W = U(:, 1 : d)$
14. $Z = W^T X Q$
15. $Y = Z R(\frac{1}{5}L + \tau)^{-1}$
16. Until Convergence
Implementation of DDRTree algorithm

We implemented the algorithm mostly in Rcpp for the purpose of efficiency. It also has extensive optimization for sparse input data. This implementation is originally based on the matlab code provided from the author of DDRTree paper.

Examples

data('iris')
subset_iris_mat <- as.matrix(t(iris[c(1, 2, 52, 103), 1:4])) #subset the data
#run DDRTree with ncenters equal to species number
DDRTree_res <- DDRTree(subset_iris_mat, dimensions = 2, maxIter = 5, sigma = 1e-2, lambda = 1, ncenter = 3, param.gamma = 10, tol = 1e-2, verbose = FALSE)
Z <- DDRTree_res$Z #obtain matrix
Y <- DDRTree_res$Y
stree <- DDRTree_res$stree
plot(Z[, 1], Z[, 2], col = iris$c(1, 2, 52, 103), 'Species') #reduced dimension
legend("center", legend = unique(iris[c(1, 2, 52, 103), 'Species']), cex=0.8, col=unique(iris[c(1, 2, 52, 103), 'Species']), pch = 1) #legend
title(main="DDRTree reduced dimension", col.main="red", font.main=4)
dev.off()
plot(Y[, 1], Y[, 2], col = 'blue', pch = 17) #center of the Z
title(main="DDRTree smooth principal curves", col.main="red", font.main=4)

#run DDRTree with ncenters equal to species number
DDRTree_res <- DDRTree(subset_iris_mat, dimensions = 2, maxIter = 5, sigma = 1e-3, lambda = 1, ncenter = NULL, param.gamma = 10, tol = 1e-2, verbose = FALSE)
Z <- DDRTree_res$Z #obtain matrix
Y <- DDRTree_res$Y
stree <- DDRTree_res$stree
plot(Z[, 1], Z[, 2], col = iris$c(1, 2, 52, 103), 'Species') #reduced dimension
legend("center", legend = unique(iris[c(1, 2, 52, 103), 'Species']), cex=0.8, col=unique(iris[c(1, 2, 52, 103), 'Species']), pch = 1) #legend
title(main="DDRTree reduced dimension", col.main="red", font.main=4)
dev.off()
plot(Y[, 1], Y[, 2], col = 'blue', pch = 2) #center of the Z
title(main="DDRTree smooth principal graphs", col.main="red", font.main=4)

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get_major_eigenvalue  

Get the top $L$ eigenvalues

Description

Get the top $L$ eigenvalues

Usage

get_major_eigenvalue(C, L)
Arguments
  C  data matrix used for eigendecomposition
  L  number for the top eigenvalues

pca_projection_R  Compute the PCA projection

Description
  Compute the PCA projection

Usage
  pca_projection_R(C, L)

Arguments
  C  data matrix used for PCA projection
  L  number for the top principal components

sqdist_R  calculate the square distance between a, b

Description
  calculate the square distance between a, b

Usage
  sqdist_R(a, b)

Arguments
  a  a matrix with $D \times N$ dimension
  b  a matrix with $D \times N$ dimension

Value
  a numeric value for the different between a and b
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