Scientific Computing for Biologists

Lecture 8: (Dis)similarity and clustering

Instructor: Paul M. Magwene

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Outline of Lecture

- Distance and dissimilarity measures
 - Quantitative data
 - Dichotomous data
 - Qualitative data
- Hierarchical clustering
- Neighbor-joining
- Multidimensional scaling (MDS)
- Minimum Spanning Tree (MST)

Similarity/Dissimilarity

Intuition

Similarity is a measure of "likeness" between two entities of interest. Dissimilarity is the complement of similarity.

Dissimilarities may be converted to similarities (and vise versa) by taking any monotonically decreasing function. For example:

$$s = 1 - d_{ij}$$
 (for $0 \le d_{ij} \le 1$)

- Dissimilarities are usually in range $0 \le d_{ij} \le C$ where C is the maximum dissimilarity
- Distances are one measure of dissimilarity but distances are unbounded to the right

$$d_{ij} \in [0, \infty]$$

Dissimilarty Measures for Quantitative Data

· Gudidean Distance dij = { \(\xi_{ix} - \times_{jk}\)^2 \(\xi_{2}\)

$$d_{ij} = \sqrt{a^2 + b^2}$$

· Marhattan (tax;-cab) distance $d_{i} = \sum_{k=1}^{k} |X_{ik} - X_{jik}|$

· Scaled Euclidean Distance

Scaled Euclidean Distance
$$d_{ij} = \left\{ \stackrel{\mathcal{E}}{\xi} \; \mathcal{W}_{\kappa}^{2} \left(\chi_{i\kappa} - \chi_{j\kappa} \right)^{2} \right\}^{1/2}$$

where W_{K} are surtable weights e.g. (Std. dev of) -1 or (range of)-1 (variable R)

Metric Vs. Non-metric

A non-regulive function, g(x,y), is metric if:
i) Satisfies the triangle inequality:

 $g(x,y) \leq g(x,z) + g(y,z)$

ii) Symmetric:

g(x,y) = g(y,x)

iii) g(xy) = 0 only if x = y

Euclidean Dist. is a metric function (as 15 monhottem distance)

Other Quantitative Measures of Discountanty

· Minkowski Metrz

$$di = \left\{ \sum_{k=1}^{r} |x_{ik} - x_{jk}|^{2} \right\}^{r} \quad \text{for integers } \lambda$$

7=1 is Manhatten distance, 71=2 is Euclidean Dist.

• Can berra Metric
$$d_{ij} = \underbrace{\frac{1}{K-1}} \frac{1}{(X_{iK} - X_{jK})}$$
• Can berra Metric
$$d_{ij} = \underbrace{\frac{1}{K-1}} \frac{1}{(X_{iK} + X_{jK})}$$
• Can berra Metric
points of relations up to points.

· CZeKanowski Coefficient

$$d_{ij} = 1 - \frac{2 \sum_{k=1}^{k} m_{in}(x_{ik}, x_{jk})}{\sum_{k=1}^{k} (x_{ik} + x_{jk})} \left[\begin{array}{c} % \text{ dissimulative } \\ \text{over all variables} \end{array} \right]$$

are treated equivalently (predictive due =
$$1 - \sum_{i=1}^{R} X_{i,K} X_{i,L}$$
 = uncentered correlation

Dissimilanty for Dichotomous Data

Simple Matching:
$$d_{ij} = 1 - \frac{a+d}{p} = \frac{b+c}{p}$$

Vaccard Coefficient: $d_{ij} = \frac{b+c}{a+b+c}$ (joint absence does)

(ZeKanowski Colf: dij = b+c
2a +b+c

Dissimilarly bother Variable a = # of objects showing + for both

varieties, k of l

t = t

c d

$$a + b + c + d = n (\# q) objects/indu/iduals)$$

 $\chi^{2} = \frac{\left(ad - bc\right)^{2} \left(a + b + c + d\right)}{\left(a + b\right)\left(a + c\right)\left(c + d\right)\left(b + d\right)}$

 $d_{kl} = 1 - \sqrt{\frac{\chi^2}{h}}$

Dissimilarities for mixed data types

Gower (1971) suggests:

$$S_{ij} = \underbrace{\sum_{K=1}^{P} W_{ijK} S_{ijK}}_{F}$$

Wix= 0 when k mssng an i

Wijk = WK otherwise (fren 1)

Define dissimilarly as:

$$d_{ij} = \left(l - s_{ij}\right)^{1/2}$$

where S_{ijk} is the similarity for i of based on variable k

-reconnereds

Sijk = 1 for brong data w/ps, tive whatch of categorical data when i and in same

Sisk = 1 - 1Xin-Xin

for continuos variables where Rx is range of variable K Introduction to Clusterny

Goal	of Clusting
· F:	nd "matural groups" in data
	ore definition:
	patches of high dimensity surrounded by patches of lower density in the p-dimensional space defined by the varieties
-	<u> </u>

Hierarchical Clustenus Agglomeratur/Divisive methods
. In practice almost always agglomerative For n data points define a set of n-1 joins that represent groupings of objects a different berelo of similarity

os + A R C D E

Simple Algorithm for Hrearchical Chusterry

- 1) Calulate a dissimilarly water for the n items
- 2) Join the two nearest items, i 4;
- 3) Delete the ith of jth row and column of the dissimilarly matrix; add a vew row/column * that represents dissimilarly of new group (; j) to all other items
 - 4) Repeat from steep 2 until there is a single group

Methods of Hierarchical Chuokenny

The different methods are determined by the function used to determine the distance between groups

Some Common Group Distance Contria

Single linkage (nearest neighbor)

Complete linkage (furthest neighbor)

Group average

Central

Single Larrage Clustering Ni, nj are # of objects in groups i of j A Dij is the smallest of the nin; dissimilantier between each element of i of each element - hvariant under monotonic transformation of the > Unaffected by fies -> Provably nice assymptotiz properties -> susceptible to "chaining"

Complete Linkage

Dij is the maximum of the ninj dissimbation between the two groups

also invariant under himotoniz transformation

Group average

Dij is the arrage of the Ninj dissimalorguer between the two groups (UPGMA, WPGMA)

Centroid method

Dis is the squared hudiden distance between the centrids of groups i of j

Hierarchical Clustering, A worked Example Single Linkage D|4 2 4 0 E|5 5 3 4 0 (A,C) B D E (A,C) (A,c) (B,0) E (B,0) 4 0 E (3) 4

Worked Example, cont.

Single Linkage Clusterne

Neighbor Joining

Originally described by Saitou and Nei, 1987.

Goal

Tries to create the (unrooted) tree topology with the least branch length (minimum-evolution criterion).

Basic algorithm:

- 1 Calculate matrix Q (next slide) from the distance matrix
- 2 Find the pair of taxa in *Q* with the lowest value; create a node on the tree that joins these two taxa (i.e. the closest neighbors)
- 3 Calculate the distance of each of the taxa in the pair to this new node
- 4 Calculate the distance of all taxa outside of this pair to the new node
- Repeat from step 1 using the distances calculated in the previous step

Neighbor Joining, cont.

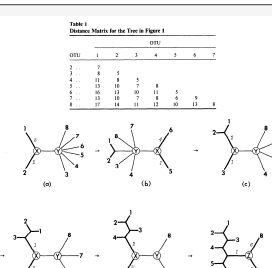
$$Q_{ij} = (r-2)d_{ij} - (R_i + R_j)$$

where r is the number of taxa, d_{ij} is the distance between taxa i and j and R_k is the row sum over row k of the distance matrix $(R_k = \sum_i d_{ik})$.

When nodes i and j are joined they are replaced by a node, A, with distance to a remaining node k given by:

$$d_{Ak} = \frac{1}{2}(d_{ik} + d_{jk} - d_{ij})$$

NJ example from Saitou and Nei 1987



(e) FIG. 3.—Application of the neighbor-joining method to the distance matrix of table 1. Italic numbers

(f)

(d)

Multidimensional Scaling (MDS)

Goal

Given dissimilarities between objects, d_{ij} , estimate a k-dimensional set of points, X, such that $|x_i - x_j| \approx d_{ij}$.

Derivation of MDS

Motivation

If we know the coordinates of n points in p-dimensional space, we can easily calculate the Euclidean distances between every pair of points. Can we reverse this process, starting with the distances and getting back the coordinates points?

Consider a data matrix X ($n \times p$). Let Q = XX' be a $n \times n$ matrix, where

$$q_{rs} = \sum_{j=1}^{p} x_{rj} x_{sj}$$

If d_{rs}^2 is the squared Euclidean distance between points r and s then we can write this as:

$$d_{rs}^{2} = \sum_{j=1}^{p} (x_{rj} - x_{sj})^{2}$$
$$= q_{rr} + q_{ss} - 2q_{rs}$$

Derivation of MDS, cont.

With a little bit of simple algebra we can show that:

$$q_{rs} = -\frac{1}{2}(d_{rs}^2 - d_{r.}^2 - d_{.s}^2 - d_{..}^2)$$

where a dot represent the average of values over the corresponding suffix: $d_{r.}^2$ is the average over the rth row of matrix $D = (d_{ij}^2)$, $d_{.s}^2$ is the average over the sth column of D, and $d_{..}^2$ is the average of all elements of D.So, given D, the squared interpoint distances, we can regenerate Q.

Since Q is symmetric, we can use eigendecomposition to write $Q=T\Lambda T'$ where Λ is a diagonal matrix of eigenvalues of Q and T is the matrix of eigenvectors. Furthermore we can write $Q=T\Lambda T'=T\Lambda^{\frac{1}{2}}\Lambda^{\frac{1}{2}}T'=XX'$ where $X=T\Lambda^{\frac{1}{2}}$.

Thus we've found how to get X from the squared distances.

See Krzanowski, W. J. (2000) Principles of multivariate analysis, for full details.

Algorithm for MDS

Given an $n \times n$ matrix of dissimilarities, D, with elements d_{ij} :

- 1 Form matrix, E, where $e_{ij} = -\frac{1}{2}d_{ij}^2$
- Subtract from each element of E the means of the row and column in which it is located and the mean of all elements of E; call the resulting matrix F
- 3 Calculate the eigenvalues (λ_i) and eigenvectors v_i of F, sorted in decreasing order. Eigenvectors should be normalized (i.e. $v_i \cdot v_i = 1$).
- The coordinates of the n point on the j-th axis are given $\sqrt{\lambda_j} v_j$

Potential MDS Complications

If the d_{ij} are metric (i.e. $d_{ij} \le d_{ik} + d_{kj}$) than F is always positive semidefinite (psd; i.e. eigenvalues ≥ 0).

If F is not psd than how do you handle negative eigenvalues?

- Most common approach is only to consider positive eigenvalues
- This is OK if negative eigenvalues have small magnitude
- If negative eigenvalues are large than approximation tends to be poor

Multidimensional Scaling: Keep in mind...

■ The configuration produced by any MDS method is indeterminate with respect to translation, rotation, and reflection.

Relationship between metric MDS and PCA

If the d_{ij} are Euclidean distances from a data matrix, X, then metric MDS of D yields the PC scores obtained by PCA of X.

Interpretation

PCA and MDS are dual methods:

- One operates on variable space (PCA)
- The other operates on subject space (MDS)

Other Metric MDS Approaches

Classical MDS minimizes:

$$\sum_{i}\sum_{j}(\delta_{ij}^{2}-d_{ij}^{2})$$

where δ_{ij} is the distance between observations i and j in the MDS approximation.

Alternates approaches try to minimize other measures of discrepancy. For example, "Sammon MDS" minimizes:

$$\sum_{i}\sum_{j}(\delta_{ij}-d_{ij})^{2}$$

Non-Metric MDS

Non-metric MDS approaches try to preserve only the rank order of the distances.

lf

$$d_{i1,j1} < d_{i2,j2} < \cdots < d_{im,jm}$$

then

$$\delta_{i1,j1} < \delta_{i2,j2} < \cdots < \delta_{im,jm}$$

Shepard-Kruskal solution:

Find \hat{d}_{ij} that minimizes:

$$\mathsf{STRESS} = \sqrt{\{\frac{\sum \sum_{i < j} (d_{ij} - \hat{d}_{ij})^2}{\sum \sum d_{ij}^2}\}}$$

MDS Example: Road Distances

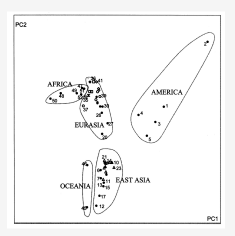
Input D: road distances between U.S. cities



More MDS Examples I

Source: Zhivotovsky et al. (2003). Features of evolution and expansion of modern humans, inferred from genomwide microsatellite markers. Am J Hum Genet 72: 1171Đ1186.

Dissimilarities: F_{ST} 's between population samples.



Good MDS References

Kenkel, N. C. and L. Oroloci (1986). Applying metric and nonmetric multidimensional scaling to ecological studies: Some new results. Ecology 67:919-928

Minimum Spanning Tree

Goal

Construct a tree that connects all points in the data set and whose total length is minimized.

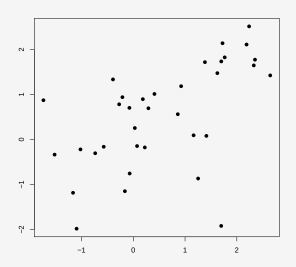
Statistical applications

- highlights close neighbors in a data set
- useful check for distortions produced by projection techniques
- tests of normality

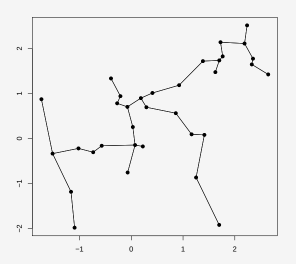
Other applications

- urban planning/engineering
- circuit design

Example Data Set



Minimum Spanning Tree: Example



Relationship between MST and Single Linkage Clustering

- Cut a single linkage dendrogram at height, δ → clusters
- Remove all edges in the MST with length $\geq \delta$ → subgraphs corresponding to the same clusters

A Generic MST Algorithm

Input: dissimilarity matrix, D, between each object (point) of interest

- **I** Create a graph, G, where $V = \{v_1, \dots, v_n\}$ and $E = \{\}$ (*E* initially empty)
- **2** Find the smallest dissimilarity, d_{ij} where (i,j) is not in E.
- 3 Add (i,j) to E if (i,j) does not create a cycle
- 4 Repeat from step 2 until every vertex is included in at least one edge

Not particularly efficient algorithm, but simple. More efficient algorithms for finding MSTs include Kruskal's Algorithm and Prim's algorithm.

Applications of the MST

MST tends to highlight close neighbors; can be used to look for distortions associated with projections to lower dimensional spaces.

Using the MST to look for Projection Distortion

- Calculate the MST based on dissimilarity in a high-dimensional space
- Draw the MST edges among points in the projection space (e.g. MDS or PCA)
- MST edges that cross highlight geometric relationships among points that are not well represented by the projection