HOMOGENEITY ANALYSIS USING EUCLIDEAN MINIMUM SPANNING TREES

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1. INTRODUCTION

In homogeneity analysis the data are a system of $m$ subsets of a finite set of $n$ objects. The purpose of the technique is to locate the objects in low-dimensional Euclidean space in such a way that the points in each of the $m$ subsets are close together. Thus we want the subsets to be compact or homogeneous, relative to the overall size of the configuration of the $n$ points.

In the usual forms of homogeneity analysis (which are also known as multiple correspondence analysis or MCA) the size of a point set is defined as the size of the star plot of the set. The star plot is the graph defined by connecting all points in the set with their centroid, and the size is the total squared length of the edges. In order to prevent trivial solutions the configuration $X$ is usually centered and normalized such that $X'X = I$. Minimizing total size of the star plots under the normalization constraints amounts to solving a (spare) singular value decomposition problem.

Recently, there have been a number of proposals to use different measures of point set size [De Leeuw, 2003]. One reason for looking at alternatives is the well known sensitivity of squared distances to outliers. Ordinary homogeneity analysis tends to locate subsets with only a few points way on the outside of the plot, which means that they dominate the solution by determining the scale. A second reason is that ordinary homogeneity analysis tends to create horseshoes for many types of data sets, i.e. two dimensional representations in which the second dimension is a quadratic function of the first [Schriever, 1985; Rijckevelopsel, 1987; Bekker and Leeuw, 1988]. This is generally seen as wasteful, because we use two dimensions to represent an essentially one dimensional structure.

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The most straightforward modification of MCA is to measure the size of the star plots by using distances instead of squared distances [Heiser, 1987; Deleeuw and Michailides, 2003; Michailides and Deleeuw, 2003]. If we continue to apply the normalization $X'X = I$ then the solutions to this modified problem map all $n$ objects into $p + 1$ points if we compute a representation in $p$ dimensions. This means the solutions are only marginally dependent on the data and do not show sufficient detail to be interesting. From the data analysis point of view they are close to worthless.

In this paper we study another measure of size, which is quite different from the star plot based ones. The size of a point set is defined as the length of the Euclidean minimum spanning tree, and our loss function is the sum of these lengths over all $m$ trees. We continue to use the same normalization as in MCA, i.e. we only consider centered and orthonormal $X$.

2. Algorithm

The problem we study is to minimize

$$\sigma(X) = \sum_{j=1}^{m} \min_{W \in W_j} \text{tr} \ W_j D(X)$$

over $X'X = I$, where $W_j$ is the set of adjacency matrices for the spanning trees of the subset $j$ and $D(X)$ is the matrix of Euclidean distances.

One obvious algorithm is to apply block relaxation [De Leeuw, 1994] to

$$\sigma(X, W_1, \cdots, W_m) = \sum_{j=1}^{m} \text{tr} \ W_j D(X).$$

Thus we minimize our loss function by alternating minimization over $X$ with minimization over the $W_j$.

It is easy to deal with the subproblem of minimizing over $W_j \in W_j$ for $X$ fixed at its current value, because that is simply computation of the minimum spanning tree, for which we can use Kruskal’s or Prim’s algorithm [Graham and Hell, 1985]. The problem of minimizing loss over normalized $X$ for fixed $W_j$ is somewhat more complicated, but we can solve it using majorization [De Leeuw, 1994; Heiser, 1995; Lange et al., 2000].
We first regularize the problem to deal with the possibility of zero distances, because in such configurations our loss function is not differentiable. Suppose $\epsilon > 0$ is a fixed small number. For each pair of points $i$ and $k$ we define

$$d_{ik}(X, \epsilon) = \sqrt{(x_i - x_k)'(x_i - x_k) + \epsilon}.$$  

Clearly this regularized distance is everywhere positive and continuously differentiable in $X$.

As in [Heiser 1987] we now apply the Arithmetic-Geometric mean inequality to obtain, for two different configurations $X$ and $Y$,

$$d_{ik}(X, \epsilon) \leq \frac{1}{2}d_{ik}^2(X, \epsilon) + \frac{1}{2}d_{ik}^2(Y, \epsilon).$$

We use the representation, which is standard in multidimensional scaling literature,

$$d_{ik}^2(X, \epsilon) = d_{ik}^2(X) + \epsilon = \text{tr} X'A_{ik}X + \epsilon,$$

where $A_{ik}$ is defined in terms of the unit vectors $e_i$ and $e_k$ as

$$A_{ik} = (e_i - e_k)(e_i - e_k)' .$$

Using this notation

$$\sigma(X, \epsilon) = \sum_{j=1}^m \text{tr} W_j D(X, \epsilon) \leq$$

$$\leq \frac{1}{2} (\text{tr} X'B(Y, \epsilon)X + \text{tr} Y'B(Y, \epsilon)Y + \epsilon \phi(Y, \epsilon)),$$

where we define the matrix

$$B(Y, \epsilon) = \sum_{i=1}^n \sum_{k=1}^n \frac{1}{d_{ik}(Y, \epsilon)} A_{ik},$$

and the scalar

$$\phi(Y, \epsilon) = \sum_{i=1}^n \sum_{k=1}^n \frac{w_{jik}}{d_{ik}(Y, \epsilon)} .$$

Given an intermediate solution $X^{(v)}$, the majorization algorithm finds an update $X^{(v+1)}$ by minimizing $\text{tr} X'B(X^{(v)}, \epsilon)X$ over $X'X = I$.

Convergence follows from general results on majorization methods, and in the particular case we can use the fact that $\sigma(X, \epsilon)$ is decreasing in $\epsilon$ to show, in addition, that letting $\epsilon \to 0$ gives convergence to the solution of the original non-regularized problem.
In general we do not iterate the majorization algorithm, which finds an optimal $X$ for given $W_j$, until convergence, but we just take a single step before computing new minimum spanning trees. An implementation of the algorithm, in the R programming language, is given in the Appendix. For the MST calculation it depends on the ape package from the CRAN repository, written by Paradis, Strimmer, Claude, Jobb, Noel, and Bolker.

3. Examples

4. A Modification

The algorithm simplifies greatly if we define our minimum spanning trees by using the square of the Euclidean distance. In fact, we need neither regularization nor majorization. We have

$$\sigma(X, W_1, \cdots, W_m) = \sum_{j=1}^{m} \text{tr} \ W_j D^2(X) = \text{tr} \ X'VX,$$

where

$$V = \sum_{i=1}^{n} \sum_{k=1}^{n} \sum_{j=1}^{m} w_{ji} A_{ik}.$$

Now minimizing over normalized $X$ for fixed $W_j$ means finding eigenvectors corresponding with the smallest non-zero eigenvalues of $V$. And computing the minimum over the $W_j$ for fixed $X$ means computing the MST for the squared distances.

Because there are only a finite number of spanning trees, and the algorithm stops if the loss does not decrease, it is clear that in this case we have convergence in a finite number of steps. This sounds much better than it is, because examples show us that the finite convergence gets us to different stationary points from different random starts.

5. Discussion

There are many variations possible. For instance travelling salesman tour. For instance squared distance MST.
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APPENDIX A. CODE

```r
require(bomals)
require(ape)

hommst <- function (g, pow = 1, eps = 1e-10) {
  nc <- dim(g)[1]; mc <- dim(g)[2]
  x <- matrix(rnorm(2*nc, n, 2))
  repeat(
    d <- euclidist(x, pow, eps)
    cc <- matrix(0, n, n); s <- 0.0
    for (j in 1:m)
      tmp <- spanner(as.factor(g[, j]), d)
      s <- s + tmp$s; cc <- tmp$e
  )
  bc <- bmat(g, d, pow)
  xc <- eigen(b$c$vectors[, c(n-1, n-2)]
  print(s)
  for (j in 1:m) spanplot(as.factor(g[, j]), x, euclidist(x, pow, eps))
}

spanner <- function (g, d)
{
  lev <- levels(g); nn <- length(g)
  s <- 0.0; cc <- matrix(0, nn, nn)
  for (k in lev) {
    ind <- which(k == g)
    nc <- length(ind)
    if (n == 1) dd <- mm <- matrix(0, 1, 1)
    else {
      dd <- d[ind, ind]
      mm <- mst(dd)
      e[ind, ind] <- mm
      s <- s + sum(dd*mm)
    }
    list(c = cc, s = s)
  }
}

spanplot <- function (g, x, d)
{
  plot(x, col="GREEN", pch=8)
  lev <- levels(g); nn <- length(g)
  rb <- rainbow(length(lev))
  for (k in lev) {
    ind <- which(k == g)
    nc <- length(ind)
    if (n == 1) dd <- mm <- matrix(0, 1, 1)
    else {
      dd <- d[ind, ind]
      mm <- mst(dd)
    }
    for (i in 1:n)
      jind <- which(1 == as.vector(mm[i, ]))
  }
}
```
```r
sapply(ind, function(r) lines(rbind(x[ind[i,j], x[ind[r,] ], col=r, which(lev==k))))
}

eudist <- function(x, pow, eps)
  c <- crossprod(t(x)); s <- diag(c)
  dd <- outer(s, s, "*") - 2*c
  if (pow==2) return(dd)
  else return(sqrt(dd+eps))

bmat <- function(c, d, pow)
  if (pow==2) b <- c
  else b <- c/d
  r <- diag(rowSums(b))
  return(r-b)
```

REFERENCES


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