Smacof at 50: A Manual Part 1: Smacof Basics

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Abstract

TBD

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1 Introduction

In Multidimensional Scaling (MDS) the data consists of information about the similarity or dissimilarity between pairs of objects selected from a finite set $\mathcal{O} = \{o_1, \cdots, o_n\}$.

In metric MDS we have numerical dissimilarity measures δ on a subset of $\mathcal{O} \times \mathcal{O}$ and we want to map the objects o_i into n points x_i of some metric space in such a way that the distances between the points approximate the dissimilarities between the objects. In smacof, our framework for MDS theory, algorithms, and computer programs, the metric space is \mathbb{R}^p , the space of all p-tuples of real numbers, and the distance is the usual Euclidean distance. Thus we are looking for x_i , with $i=1,\cdots,n$, such that

$$\delta_{ij} := \delta(o_i, o_j) \approx d_{ij}(X) := \sqrt{\sum_{s=1}^{p} (x_{is} - x_{js})^2}. \tag{1}$$

(the symbol := is used for definitions). Note that *distance matrix completion* (Fang and O'Leary (2012)) is an example of metric MDS.

Besides metric MDS there is also something called *non-metric MDS*. The terms is ambiguous and can mean either one of two things. One definition is that we do not have numerical information about the dissimilarities, but only ordinal of nominal information. In the ordinal case we know that some dissimilarities are larger or smaller than others, in the nominal case that the objects are partitioned into groups and within-group distances are smaller than between-group distances. An MDS analysis is also non-metric if we have numerical dissimilarities but we decide to use only the ordinal or nominal information in the dissimilarities as data for our MDS analysis. The data are a binary relation \leq_{δ} on a subset of $\mathcal{O} \times \mathcal{O}$. There is a second binary relation \leq_{d} on $\mathbb{R}_{+} \times \mathbb{R}_{+}$. We want to find the configuration X such that

$$(o_i, o_i) \leq_{\delta} (o_k, o_l) \quad \Rightarrow \quad d_{ij}(X) \leq_d d_{kl}(X). \tag{2}$$

For most non-metric MDS problems both \leq_{δ} and \prec_d are partial orders, but for nominal data they can be equivalence relations.

The term "non-metric" is sometimes also used in the case in which we do have numerical dissimilarities δ_{ij} , but the MDS problem is to find Euclidean distances which approximate some partially unknown function of the dissimilarities. The function should be member of a well-defined class of functions, for instance a third degree polynomial or a piecewise linear spline on a given knot sequence. The MDS technique not only finds the map of $\mathcal O$ into $\mathbb R^p$, but also chooses a function f from the set $\mathfrak D$ to improve the approximation.

$$f(\delta_{ij}) \approx d_{ij}(X).$$
 (3)

Note that (3) implies that f is applied to each dissimilarity separately. Thus \mathfrak{D} consists of real-valued functions of a single non-negative real variable.

To avoid confusion, we will refer to this class of MDS techniques as *non-linear MDS*. This choice of terminology has the unfortunately side-effect that \mathfrak{D} in non-linear MDS can be the set of all linear transformations. One important example of this linear non-linear MDS is classical MDS with an

additive constant (Messick and Abelson (1956)). And, as another consequence, \mathfrak{D} can also be the set of all monotone or all one-one transformations, in which case the non-linear MDS technique is also a non-metric MDS technique. And finally the set \mathfrak{D} can have only a single element, the identity transformation, in which case non-linear MDS is metric MDS.

One way to solve this terminological dilemma is to simply define MDS as finding an approximate solution (X, Δ) to

$$\Delta \in \mathfrak{D},$$
 (4a)

$$X \in \mathfrak{X},$$
 (4b)

$$D(X) = \Delta, \tag{4c}$$

where D(X) is the matrix of Euclidean distances and $\mathfrak D$ is the set of *transformed dissimilarities* in the non-linear case or the set of *quantified dissimilarities* in the non-metric case. We can then freely discuss metric MDS, linear MDS, non-linear MDS, and non-metric MDS as special cases, defined by different sets $\mathfrak D$.

2 Types of Data

- 1. \mathcal{R} is a set of n objects called *rows*.
- 2. \mathcal{C} is a set of m objects called *columns*.
- 3. Numerical
- $\mathcal{R}\mathcal{R}$ data, $\langle \mathcal{R} \times \mathcal{R}, \delta \rangle$, real-valued function δ on a subset of $\mathcal{R} \times \mathcal{R}$.
- \mathcal{CC} data, $\langle \mathcal{C} \times \mathcal{C}, \delta \rangle$, real-valued function δ on a subset of $\mathcal{C} \times \mathcal{C}$.
- \mathcal{RC} data, $\langle \mathcal{R} \times \mathcal{C}, \delta \rangle$, real-valued function δ on a subset of $\mathcal{R} \times \mathcal{C}$.
- 2. Ordinal
- XX data, partial order on a subset of $X \times X$
- XY data, partial order \leq on the $n \times m$ -element set $X \times Y$
- 3. Nominal
- XX data, equivalence relation on a subset of $X \times X$
- XY data, equivalence relation on a subset of $X \times Y$

Conditional Data + XX data, n partial \leq is the union of n partial orders \leq_i on the n disjoint n-element sets $x_i \times X + XY$ data, the partial order \leq is the union of n partial orders \leq_i on the n disjoint m-element sets $x_i \times Y$

Replicated Data

As ordinal, but the partial orders are equivalence relations.

3 Loss Function

In the pioneering papers Kruskal (1964a) and Kruskal (1964b) the MDS problem was formulated for the first time as minimization of an explicit *loss function* or *badness-of-fit function*, which measures the quality of the approximation of the dissimilarities by the distances. To be historically accurate, we should mention that the non-metric MDS technique proposed by Shepard (1962a) and Shepard (1962b) can be reformulated as minimization of an explicit loss function (see, for example, De Leeuw (2017b)). And the classical Young-Householder-Torgerson MDS technique (Torgerson (1952)) for metric MDS can be reformulated as minimizing an explicit least squares loss function (De Leeuw and Heiser (1982)) as well. But neither of these two predecessors was formulated originally as an explicit minimization problem for a specific loss function

3.1 Metric MDS

The loss function in least squares metric Euclidean MDS is called *raw stress* and is defined as

$$\sigma_R(X) := \frac{1}{2} \sum_{1 \le j < i \le n} w_{ij} (\delta_{ij} - d_{ij}(X))^2. \tag{5}$$

The subscript R in σ_R stands for "raw", because we will discuss other least squares loss functions for which we will also use the symbol σ , but with other subscripts.

In definition (5) the w_{ij} are known non-negative weights, the δ_{ij} are the known non-negative dissimilarities between objects o_i and o_j , and the $d_{ij}(X)$ are the distances between the corresponding points x_i and x_j . The summation is over all pairs (i,j) with $w_{ij} > 0$. From now on we use "metric MDS" to mean the minimization of σ_R .

The $n \times p$ matrix X, which has the coordinates x_i of the n points as its rows, is called the *configuration*, where p is the *dimension* of the Euclidean space in which we make the map. The metric MDS problem (of dimension p, for given W and Δ) is the minimization of (5) over the $n \times p$ configurations X.

The weights w_{ij} can be used to quantify information about the precision or importance of the corresponding dissimilarities. Some of the weights may be zero, which can be used to code *missing data*. If all weights are positive we have *complete data*. If we have complete data, and all weights are equal to one, we have *unweighted* metric MDS. The pioneering papers by Shepard, Kruskal, and Guttman only consider the unweighted case. Weights were only introduced in MDS in De Leeuw (1977).

We assume throughout that the weights are *irreducible* (De Leeuw (1977)). This means there is no partitioning of the index set $I_n := \{1, 2, \cdots, n\}$ into subsets for which all between-subset weights are zero. A reducible metric MDS problems decomposes into a number of smaller independent metric MDS problems, so the irreducibility assumption causes no real loss of generality.

The fact that the summation in (5) is over all j < i indicates that the diagonal elements of Δ are not used (they are assumed to be zero) and the elements above the diagonal are not used either (they are assumed to be equal to the corresponding elements below the diagonal). The somewhat mysterious

factor $\frac{1}{2}$ in definition (5) is there because it simplifies some of the formulas in later sections of this paper.

3.2 Non-linear MDS

Kruskal was not really interested in metric MDS and the "raw" loss function (5). His papers are really about non-metric MDS, by which we mean least squares non-metric Euclidean MDS. Non-metric MDS differs from metric MDS because we have incomplete information about the dissimilarities. As we have seen, that if some dissimilarities are missing metric MDS can handle this by using zero weights. In some situations, however, we only know the rank order of the non-missing dissimilarities. We do not know, or we refuse to use, their actual numeric values. Or, to put it differently, even if we have numerical dissimilarities we are looking for a *transformation* of the non-missing dissimilarities, where the transformation is chosen from a set of admissible transformations (for instance from all linear or monotone transformations). If the dissimilarities are non-numerical, for example rank orders or partitionings, we choose from the set of admissible *quantifications*.

In non-metric MDS raw stress becomes

$$\sigma_R(X,\Delta) := \frac{1}{2} \sum w_{ij} (\delta_{ij} - d_{ij}(X))^2, \tag{6} \label{eq:sigma_R}$$

where Δ varies over the quantified or transformed dissimilarities. In MDS parlance they are also called *pseudo-distances* or *disparities*. Loss function (6) must be minimized over both configurations and disparities, with the condition that the disparities Δ are an admissible transformation or quantification of the data. In Kruskal's non-metric MDS this means requiring monotonicity. In this paper we will consider various other choices for the set of admissible transformations. We will use the symbol $\mathfrak D$ for the set of admissible transformations

The most familiar examples of $\mathfrak D$ (linear, polynomial, splines, monotone) define convex cones with apex at the origin. This means that if $\Delta \in \mathfrak D$ then so is $\lambda \Delta$ for all $\lambda \geq 0$. But consequently minimizing (6) over all $\Delta \in \mathfrak D$ and over all configurations has the trivial solution $\Delta = 0$ and X = 0, corresponding with the global minimum $\sigma(X, \Delta) = 0$. We need additional constraints to rule out this trivial solution, and in non-metric MDS this is done by choosing a *normalization* that keeps the solution away from zero.

Kruskal's original solution is to define normalized stress as

$$\sigma(X,\Delta) := \frac{\sum w_{ij}(\delta_{ij} - d_{ij}(X))^2}{\sum w_{ij}d_{ij}^2(X)}.$$
 (7)

To be precise, in Kruskal's formulation there are no weights, and he actually takes the square root of (7) to define *Kruskal's stress*. The non-metric Euclidean MDS problem now is to minimize loss function (7) over all $n \times p$ configurations X and all admissible disparities Δ .

3.3 Non-metric MDS

3.4 Normalization

Equation (7) is only one way to normalize raw stress. Some obvious alternatives are discussed in detail in Kruskal and Carroll (1969) and De Leeuw (1975). In the terminology of De Leeuw (1975) there are *explicit* and *implicit* normalizations.

In implicit normalization we minimize either

$$\sigma(X, \hat{D}) := \frac{\sum w_{ij} (\hat{d}_{ij} - d_{ij}(X))^2}{\sum w_{ij} \hat{d}_{ij}^2} \tag{8}$$

or

$$\sigma(X, \hat{D}) := \frac{\sum w_{ij} (\hat{d}_{ij} - d_{ij}(X))^2}{\sum w_{ij} d_{ij}^2(X)} \tag{9}$$

over X and $\Delta \in \mathfrak{D}$.

As we have seen, Kruskal (1964a) chooses definition (9) and calls the explicitly normalized loss function *normalized stress*. Note that we overload the symbol σ to denote any one of the least squares loss functions. It will always be clear from the text which σ we are talking about.

In explicit normalization we minimize the raw stress $\sigma_R(X,\hat{D})$ from (6), but we add the explicit constraint

$$\sum w_{ij}d_{ij}^{2}(X) = 1, (10)$$

or the constraint

$$\sum w_{ij}\hat{d}_{ij}^2 = 1. \tag{11}$$

Kruskal and Carroll (1969) and De Leeuw (2019) show that these four normalizations all lead to essentially the same solution for X and \hat{D} , up to scale factors dictated by the choice of the particular normalization. It is also possible to normalize both X and \hat{D} , either explicitly or implicitly, and again this will give the same solutions, suitably normalized. These invariance results assume the admissible transformations form a closed cone with apex at the origin, i.e. if \hat{D} is admissible and $\lambda \geq 0$ then $\lambda \hat{D}$ is admissible as well. The matrices of Euclidean distances D(X) form a similar closed cone as well. The non-metric MDS problem is to find an element of the \hat{D} cone \mathcal{D} and an element of the D(X) cone where the angle between the two is a small as possible.

In the R version of smacof (De Leeuw and Mair (2009), Mair, Groenen, and De Leeuw (2022)) we use explicit normalization (11). This is supported by the result, also due to De Leeuw (1975), that projection on the intersection of the cone of disparities and the sphere defined by (11) is equivalent to first projecting on the cone and then normalizing the projection (see also Bauschke, Bui, and Wang (2018)).

In the version of non-metric MDS discussed in this manual we need more flexibility. For algorithmic reasons that may become clear later on, we will go with the original (7), i.e. with the implicitly normalized Kruskal's stress. For the final results the choice between normalizations should not make a difference, but the iterative computations will be different for the different choices.

3.5 Some thoughts on ALS

The formulation in equations (4a) and (4b) neatly separates the metric MDS part (4a) and the transformation/quantification part (4b). This second part is also often called the *optimal scaling* part.

Equations (4a) and (4b) corresponds with the way most iterative non-linear and non-metric MDS techniques are implemented. The algorithms use Alternating Least Squares (ALS). There have been quite a few ALS algorithms avant-la-lettre, but as far as I know both the name and ALS as a general approach to algorithm construction were first introduced in De Leeuw (1968), and then widely disseminated in a series of papers by De Leeuw, Young, and Takane in the 1970's (work summarized in Young, De Leeuw, and Takane (1980) and Young (1981)).

In the ALS implementation of MDS two sub-algorithms are used in each iteration: one to improve the fit of the distances to the current disparities Δ and one to improve the fit of the disparities to the current distances. The two sub-algorithms define one major iteration of the MDS technique. In formulas (using superscript (k) for major iteration number) we start with $(X^{(0)}, \Delta^{(0)})$ and then alternate the mimization problems

$$X^{(k+1)} \ni \{ \sigma(X^{(k+1)}, \Delta^{(k)}) = \min_{X \in \mathfrak{X}} \sigma(X, \Delta^{(k)}) \}, \tag{12a}$$

$$\begin{split} X^{(k+1)} &\ni \{\sigma(X^{(k+1)}, \Delta^{(k)}) = \min_{X \in \mathfrak{X}} \sigma(X, \Delta^{(k)})\}, \\ \Delta^{(k+1)} &\ni \{\sigma(X^{(k+1)}, \Delta^{(k+1)}) = \min_{\Delta \in \mathfrak{D}} \sigma(X^{(k+1)}, \Delta)\}, \end{split} \tag{12a}$$

where \ni is short for "such that". In MDS it is more realistic not to minimize loss in the sub-steps but merely to decrease it. Minimization in one or both of the two subproblems may itself require an infinite iterative method, which we have to truncate anyway. Thus

$$X^{(k+1)} \in \mathfrak{X} \ni \{ \sigma(X^{(k+1)}, \Delta^{(k)}) < \sigma(X^{(k)}, \Delta^{(k)}) \}, \tag{13a}$$

$$\Delta^{(k+1)} \in \mathfrak{D} \ni \{ \sigma(X^{(k+1)}, \Delta^{(k+1)}) < \sigma(X^{(k+1)}, \Delta^{(k)}) \}. \tag{13b}$$

The Single-Phase approach

In Kruskal (1964a) defines

$$\sigma(X) := \min_{\hat{D} \in \mathfrak{D}} \ \sigma(\hat{D}, X) = \sigma(X, \hat{D}(X)), \tag{14}$$

where $\sigma(\hat{D},X)$ is defined by (9). The minimum in (14) is over admissible transformations. In definition (14)

$$\hat{D}(X) := \underset{\hat{D} \in \mathfrak{D}}{\operatorname{argmin}} \, \sigma(X, \hat{D}). \tag{15}$$

Normalized stress defined by (14) is now a function of X only. Under some conditions, which are true in Kruskal's definition of non-metric MDS, there is a simple relation between the partials of (9) and those of (14).

$$\mathcal{D}\sigma(X) = \mathcal{D}_1\sigma(X, \hat{D}(X)),\tag{16}$$

where $\mathcal{D}\sigma(X)$ are the derivatives of σ from (14) and $\mathcal{D}_1\sigma(X,\hat{D}(X))$ are the partial derivatives of σ from (9) with respect to X. Thus the partials of σ from (14) can be computed by evaluating the

partials of σ from (9) with respect to X at $(X, \hat{D}(X))$. This has created much confusion in the past. The non-metric MDS problem in Kruskal's original formulation is now to minimize σ from (14), which is a function of X alone.

Guttman (1968) calls this the *single-phase approach*. A variation of Kruskal's single-phase approach defines

$$\sigma(X) = \sum w_{ij} (d_{ij}^{\#}(X) - d_{ij}(X))^2, \tag{17}$$

where the $d_{ij}^\#(X)$ are *Guttman's rank images*, i.e. the permutation of the $d_{ij}(X)$ that makes them monotone with the δ_{ij} (Guttman (1968)). Or, alternatively, define

$$\sigma(X) := \sum w_{ij} (d_{ij}^{\%}(X) - d_{ij}(X))^2, \tag{18}$$

where the $\hat{d}_{ij}^{\%}(X)$ are *Shepard's rank images*, i.e. the permutation of the δ_{ij} that makes them monotone with the $d_{ij}(X)$ (Shepard (1962a), Shepard (1962b), De Leeuw (2017b)).

Minimizing the Shepard or Guttman single-phase loss functions is computationally more complicated than Kruskal's *monotone regression* approach, mostly because the rank-image transformations are not differentiable, and there is no analog of (16) and of the equivalence of the different implicit and explicit normalizations.

The Two-Phase Approach

The two-phase approach or alternating least squares (ALS) approach alternates minimization of $\sigma(\hat{D}, X)$ over X for our current best estimate of \hat{D} with minimization of $\sigma(\hat{D}, X)$ over $\Delta \in \mathfrak{D}$ for our current best value of X. Thus an update from iteration k to iteration k+1 looks like

$$\begin{split} \hat{D}^{(k)} &= \mathop{\mathrm{argmin}}_{\hat{D} \in \mathfrak{D}} \sigma(\hat{D}, X^{(k)}), \\ X^{(k+1)} &= \mathop{\mathrm{argmin}}_{Y} \sigma(\hat{D}^{(k)}, X). \end{split} \tag{19a}$$

$$X^{(k+1)} = \operatorname*{argmin}_{X} \sigma(\hat{D}^{(k)}, X). \tag{19b}$$

This ALS approach to MDS was in the air since the early (unsuccessful) attempts around 1968 of Young and De Leeuw to combine Torgerson's classic metric MDS method with Kruskal's monotone regression transformation. All previous implementations of non-metric smacof use the two-phase approach, and we will do the same in this paper.

As formulated, however, there are some problems with the ALS algorithm. Step (19a) is easy to carry out, using monotone regression. Step (19b) means solving a metric scaling problem, which is an iterative proces that requires an infinite number of iterations. Thus, in the usual implementations, step (19a) is combined with one of more iterations of a convergent iterative procedure for metric MDS, such as smacof. If we take only one of these *inner iterations* the algorithm becomes indistinguishable from Kruskal's single-phase method. This has also created much confusion in the past.

In the usual implementations of the ALS approach we solve the first subproblem (19a) exactly, while we take only a single step towards the solution for given \hat{D} in the second phase (19b). If we have an infinite iterative procedure to compute the optimal $\hat{D} \in \mathfrak{D}$ for given X, then a more balanced approach would be to take several inner iterations in the first phase and several inner

iterations in the second phase. How many of each, nobody knows. In our current implementation of smacof we take several inner iteration steps in the first phase and a single inner iteration step in the second phase.

Smacof Notation and Terminology 4

We discuss some standard MDS notation, first introduced in De Leeuw (1977). This notation is useful for the second phase of the ALS algorithm, in which solve the metric MDS problem of we minimizing unnormalized $\sigma(X, \hat{D})$ over X for fixed \hat{D} . We will discuss the first ALS phase later in the paper.

Start with the unit vectors e_i of length n. They have a non-zero element equal to one in position i, all other elements are zero. Think of the e_i as the columns of the identity matrix.

Using the e_i we define for all $i \neq j$ the matrices

$$A_{ij} := (e_i - e_j)(e_i - e_j)'. \tag{20}$$

The A_{ij} are of order n, symmetric, doubly-centered, and of rank one. They have four non-zero elements. Elements (i, i) and (j, j) are equal to +1, elements (i, j) and (j, i) are -1.

The importance of A_{ij} in MDS comes from the equation

$$d_{ij}^2(X) = \operatorname{tr} X' A_{ij} X. \tag{21}$$

In addition we use the fact that the A_{ij} form a basis for the binomn2-dimensional linear space of all doubly-centered symmetric matrices.

Expanding the square in the definition of stress gives

$$\sigma(X) = \frac{1}{2} \{ \sum w_k \delta_k^2 - 2 \sum w_k \delta_k d_k(X) + \sum w_k d_k^2(X) \}. \tag{22}$$

It is convenient to have notation for the three separate components of stress from equation (22). Define

$$\eta_{\hat{D}}^2 = \sum w_{ij} \hat{d}_{ij}^2, \tag{23}$$

$$\rho(X) = \sum w_{ij} \hat{d}_{ij} d_{ij}(X), \tag{24}$$

$$\begin{split} \eta_{\hat{D}}^2 &= \sum w_{ij} \hat{d}_{ij}^2, \\ \rho(X) &= \sum w_{ij} \hat{d}_{ij} d_{ij}(X), \\ \eta^2(X) &= \sum w_{ij} d_{ij}(X)^2. \end{split} \tag{23}$$

which lead to

$$\sigma(X) = \frac{1}{2} \left\{ \eta_{\hat{D}}^2 - 2\rho(X) + \eta^2(X) \right\}. \tag{26}$$

We also need

$$\lambda(X) = \frac{\rho(X)}{\eta(X)}. (27)$$

Using the A_{ij} makes it possible to give matrix expressions for ρ and η^2 . First

$$\eta^2(X) = \operatorname{tr} X' V X, \tag{28}$$

with

$$V := \sum w_{ij} A_{ij}. \tag{29}$$

In the same way

$$\rho(X) = \operatorname{tr} X' B(X) X, \tag{30}$$

with

$$B(X) := \sum w_{ij} r_{ij}(X) A_{ij}, \tag{31}$$

with

$$r_{ij}(X) := \begin{cases} \frac{\delta_{ij}}{d_{ij}(X)} & \text{if } d_{ij}(X) > 0, \\ 0 & \text{if } d_{ij}(X) = 0. \end{cases} \tag{32}$$

Note that B is a function from the set of $n \times p$ configurations into the set of symmetric doubly-dentered matrices of order n. All matrices of the form $\sum x_{ij}A_{ij}$, where summation is over all pairs (i,j) with j < i, are symmetric and doubly-centered. They have $-x_{ij}$ as off-diagonal elements while the diagonal elements (i,i) are $\sum_{j=1}^n x_{ij}$.

Because B(X) and V are non-negative linear combinations of the A_{ij} they are both positive semi-definite. Because W is assumed to be irreducible the matrix V has rank n-1, with only vectors proportional to the vector e with all elements equal to one in its null-space (De Leeuw (1977)).

Summarizing the results so far we have

$$\sigma(X) = \frac{1}{2} \{ \eta_{\hat{D}}^2 - \operatorname{tr} X' B(X) X + \operatorname{tr} X' V X \}. \tag{33} \label{eq:33}$$

Next we define the Guttman transform of a configuration X, for given W and Δ , as

$$G(X) = V^+ B(X)X, (34)$$

with V^+ the Moore-Penrose inverse of V. In our computations we use

$$V^{+} = (V + \frac{1}{n}ee')^{-1} - \frac{1}{n}ee'$$
(35)

Also note that in the unweighted case with complete data V=nJ, where J is the centering matrix $I-\frac{1}{n}ee'$, and thus $V^+=\frac{1}{n}J$. The Guttman transform is then simply $G(X)=n^{-1}B(X)X$.

5 Intermezzo: Explicit Normalization

$$\sigma(X,\hat{D}) = \frac{1}{2} \frac{\sum w_{ij} (\hat{d}_{ij} - d_{ij}(X))^2}{\sum w_{ij} d_{ij}^2(X)} \label{eq:sigma}$$

Majorize

$$\sigma(X,\hat{D}) \leq \frac{1}{2} \frac{\eta^2(\hat{D}) - 2 \mathrm{tr} \; X' V \overline{Y} + \mathrm{tr} \; X' V X}{\mathrm{tr} \; X' V X} = \frac{\omega(X,Y)}{\eta^2(X)}$$

Stationary equations

$$\eta^2(X)(VX-VG(Y)) - \omega(X,Y)VX = V\{(\eta^2(X)-\omega(X,Y))X - \eta^2(X)\overline{Y}\}$$

So at a minimum X is proportional to \overline{Y} or $X = \alpha \overline{Y}$ for some α . For ... to be zero we must have

$$\alpha(\alpha^2\eta^2(\overline{Y}) - (\eta^2(\hat{D}) - 2\alpha\eta^2(\overline{Y}) + \alpha^2\eta^2(\overline{Y})) = \alpha^2\eta^2(\overline{Y})$$

which works out to be

$$\alpha = \frac{\eta^2(\hat{D})}{\eta^2(\overline{Y})}$$

$$\hat{X} = \frac{\eta^2(\hat{D})}{\eta^2(\overline{Y})} \, \overline{Y}$$

The minimum is equal to

$$\frac{-\frac{(\eta^2(\overline{Y}))^2}{\eta^2(\widehat{D})} + \eta^2(\overline{Y})}{\eta^2(\overline{Y})} = 1 - \frac{\eta^2(\overline{Y})}{\eta^2(\widehat{D})}$$

Use homogeneity of the Guttman transform.

More generally suppose we update with

$$X = \overline{Y} + \alpha(Y - \overline{Y})$$

Write

$$\omega(X,Y) = \eta^2(\hat{D}) + \operatorname{tr}\,(X - \overline{Y})'V(X - \overline{Y}) - \eta^2(\overline{Y})$$

Thus if $X(\alpha) = \overline{Y} + \alpha(Y - \overline{Y})$ we have

$$\omega(\alpha) = \eta^2(\hat{D}) + \alpha^2 \mathrm{tr} \; (Y - \overline{Y})' V (Y - \overline{Y}) - \eta^2(\overline{Y})$$

and

$$\begin{split} \eta^2(\alpha) &= \eta^2(\overline{Y}) + 2\alpha \mathrm{tr}\; (Y - \overline{Y})' V \overline{Y} + \alpha^2 \mathrm{tr}\; (Y - \overline{Y})' V (Y - \overline{Y}) \\ \omega(Y,Y) &= \eta^2(\hat{D}) + \mathrm{tr}\; (Y - \overline{Y})' V (Y - \overline{Y}) - \eta^2(\overline{Y}) \\ \frac{\omega(\alpha)}{\eta^2(\alpha)} &\leq \sigma(Y) \end{split}$$

6 Properties of Smacof Loss

6.1 Derivatives

The Euclidean distance function d_{ij} from ... is not differentiable at configurations X with $x_i = x_j$. If $d_{ij}(X) > 0$ then

$$\mathcal{D}\sigma(X) = \frac{1}{d_{ij}(X)}A_{ij}X$$

If $d_{ij}(X) = 0$ then

$$D_+d_{ij}(X,Y) = \lim_{\epsilon \downarrow 0} \frac{d_{ij}(X+\epsilon Y) - d_{ij}(X)}{\epsilon} = d_{ij}(Y)$$

which is non-linear in Y, showing non-differentiability.

$$D_+\sigma(X,Y)=\operatorname{tr} Y'(V-B(X))X+\sum\{w_{ij}\delta_{ij}d_{ij}(Y)\mid d_{ij}(X)=0\}$$

This form of the directional derivative is used by De Leeuw (1984) to show that two independent necessary conditions for a local minimum are (V-B(X))X=0 and $d_{ij}(X)>0$ for all (i,j) with $w_{ij}\delta_{ij}>0$.

6.1.1 Gradient

$$\mathcal{D}\sigma(X) = (V - B(X))X$$

At a stationary point B(X)X = VX or $V^+B(X)X = X$. Thus a necessary condition for a local minimum is that $V^+B(X)$ has at least p eigenvalues equal to one. De Leeuw (2014) has shown that if $V^+B(X) \lesssim I$ then actually X is a global minimizer of stress.

$$\rho(X) = \sum w_{ij} \delta_{ij}(X)$$

$$\nabla d_{ij}(X) = \begin{bmatrix} 0 \\ \frac{x_i - x_j}{d_{ij}(X)} \\ 0 \\ -\frac{x_i - x_j}{d_{ij}(X)} \\ 0 \end{bmatrix}$$

$$\partial d_{ij}(X) = \left\{ \begin{bmatrix} 0 \\ y \\ 0 \\ -y \\ 0 \end{bmatrix} \mid y'y \le 1 \right\}.$$

6.1.2 Hessian

The results on the Hessian of stress are largely unpublished. So we summarize them here in this manual, so they'll be even more unpublished.

$$\begin{split} H_{st}(X) &:= \sum w_{ij} \frac{\delta_{ij}}{d_{ij}(X)} \left\{ \frac{A_{ij} x_s x_t' A_{ij}}{d_{ij}^2(X)} \right\} \\ H_{st}(X) &= \sum w_{ij} \frac{\delta_{ij}}{d_{ij}^3(X)} (x_{is} - x_{js}) (x_{it} - x_{jt}) A_{ij} \\ \mathcal{D}_{st} \sigma(X) &= \begin{cases} H_{st}(X) & \text{if } s \neq t, \\ V - B(X) + H_{st} & \text{if } s = t. \end{cases} \end{split}$$

If I_p is the identity matrix of order p, and \otimes is the Kronecker product, then

$$\mathcal{D}^2\sigma(X) = I_p \otimes (V - B(X)) + H(X)$$

$$\sum_{s=1}^p \sum_{t=1}^p y_s' H_{st} y_t = \sum w_{ij} \frac{\delta_{ij}}{d_{ij}(X)} \left\{ \frac{(\operatorname{tr} Y' A_{ij} X)^2}{d_{ij}^2(X)} \right\} \leq \sum w_{ij} \frac{\delta_{ij}}{d_{ij}(X)} \operatorname{tr} Y' A_{ij} Y = \operatorname{tr} Y' B(X) Y.$$

Thus

$$0\lesssim H\lesssim I_p\otimes B(X),$$

and

$$I_p \otimes (V - B(X)) \lesssim \mathcal{D}^2 \sigma(X) \lesssim I_p \otimes V$$

At a local minimum of σ

$$0 \lesssim \mathcal{D}^2 \sigma(X) \lesssim I_p \otimes V$$

In comparing the lower bounds on $\mathcal{D}^2\sigma(X)$ in ... and ... De Leeuw (2014) shows that $V-B(X)\gtrsim 0$ is sufficient for a *global* minimum of stress (but far from necessary).

Also

$$\sum_{t=1}^p H_{st} y_t = \sum w_{ij} \frac{\delta_{ij}}{d_{ij}(X)} \left\{ \frac{\operatorname{tr} Y' A_{ij} X}{d_{ij}^2(X)} \right\} A_{ij} x_s$$

If Y=X then $H(X)y=(I_p\otimes B(X))x$ and thus

$$\mathcal{D}^2\sigma(X)x=(I_p\otimes V)x.$$

In the unweighted case this means that X is an eigenvector of $\mathcal{D}^2\sigma(X)$ with eigenvalue n. Inequalities ... show that this is actually the largest eigenvalue. Or $(I_p\otimes V)^+\mathcal{D}^2\sigma(X)\lesssim I$.

If Y = XT with T anti-symmetric then $\operatorname{tr} Y'A_{ij}X = 0$ then thus H(X)y = 0. Thus

$$\sum_{t=1}^p \mathcal{D}_{st} \sigma(X) y_t = (V - B(X)) y_t$$

which is zero if $\mathcal{D}\sigma(X)$ is zero. Thus at a stationary point of stress $\mathcal{D}^{\sigma}(X)$ has $\frac{1}{2}p(p-1)$ zero eigenvalues.

There are several ways to think of the Hessian. The simplest one (perhaps) is as an $np \times np$ symmetric matrix (corresponding to

column-major R vector of length $\frac{1}{2}np(np+1)$). This is what we would use for a straightforward version of Newton-Raphson.

It is more elegant, however, to think of H as a symmetric super-matrix of order p, with as elements $n \times n$ matrices. And, for some purposes, such as the pseudo-confidence ellipsoids in De Leeuw (2017a), as a super-matrix of order n with as elements $p \times p$ matrices. Both the super-matrix interpretations lead to four-dimensional arrays, the first a $p \times p \times n \times n$ array, the second an $n \times n \times p \times p$ array. The different interpretations lead to different ways to store the Hessian in memory, and to different ways to retrieve its elements. Of course we can write routines to transform from one interpretation to another.

6.2 Lagrangian

In our implementation of the smacof algorithm we minimize stress over configurations with $\eta(X)=1$, or, equivalently, $\sum w_{ij}d_{ij}^2(X)=1$. This means we do not look for X with $\mathcal{D}\sigma(X)=(V-B(X))X=0$, but we look for solutions of

$$(V - B(X))X - \lambda VX = 0, \text{tr } X'VX = 1.$$

At the solution

$$\lambda = 1 - \rho(X)$$

and

$$X = \frac{\Gamma(X)}{\eta(\Gamma(X))}$$

Also it is necessary for a local minimum that

$$\Gamma(X) = \rho(X)X$$

Because the Guttman transform is homogeneous of degree zero this implies

$$\Gamma(\Gamma(X) = \Gamma(X),$$

so although X is not a fixed point of the Guttman transform, $\Gamma(X)$ is.

The second order necessary condition is that

$$H(X)\gtrsim I_p\otimes (\rho(X)V-B(X))$$

is positive

6.2.1 Kuhn-Tucker Points

bozo

Smacof Algorithm 7

Introduction to Majorization

Majorization, these days better known as MM (Lange (2016)), is a general approach for the construction of minimization algorithms. There is also minorization, which leads to maximization algorithms, which explains the MM acronym: minorization for maximization and majorization for minimization.

Before the MM principle was formulated as a general approach to algorithm construction there were some important predecessors. Major classes of MM algorithms avant la lettre were the EM Algorithm for maximum likelihood estimation of Dempster, Laird, and Rubin (1977), the Smacof Algorithm for MDS of De Leeuw (1977), the Generalized Weiszfeldt Method of Vosz and Eckhardt (1980), and the Quadratic Approximation Method of Böhning and Lindsay (1988). The first formulation of the general majorization principle seems to be De Leeuw (1994).

Let's start with a brief introduction to majorization. Minimize a real valued function σ over $x \in \mathbb{S}$, where \mathbb{S} is some subset of \mathbb{R}^n . There are obvious extensions of majorization to functions defined on more general spaces, with values in any partially ordered set, but we do not need that level of generality in this manual. Also majorization applied to σ is minorization applied to $-\sigma$, so concentrating on majorization-minimization and ignoring minorization-maximization causes no loss of generality

Suppose there is a real-valued function η on $\mathbb{S} \otimes \mathbb{S}$ such that

$$\sigma(x) \le \eta(x, y) \qquad \forall x, y \in \mathbb{S},
\sigma(x) = \eta(x, x) \qquad \forall x \in \mathbb{S}.$$
(36)

$$\sigma(x) = \eta(x, x) \qquad \forall x \in \mathbb{S}.$$
 (37)

The function η is called a *majorization scheme* for σ on S. A majorization scheme is *strict* if $\sigma(x) < \eta(x,y)$ for all $x,y \in S$ with $x \neq y$.

Define

$$x^{(k+1)} \in \operatorname*{argmin}_{x \in \mathbb{S}} \eta(x, x^{(k)}), \tag{38}$$

assuming that $\eta(\bullet,y)$ attains its (not necessarily unique) minimum over $x\in\mathbb{S}$ for each y. If $x^{(k)} \in \operatorname{argmin}_{x \in \mathbb{S}} \eta(x, x^{(k)})$ then we stop.

By majorization property (36)

$$\sigma(x^{(k+1)}) \le \eta(x^{(k+1)}, x^{(k)}). \tag{39}$$

Because we did not stop update rule (38) implies

$$\eta(x^{(k+1)}, x^{(k)}) < \eta(x^{(k)}, x^{(k)}).$$
(40)

and finally by majorization property (36)

$$\eta(x^{(k)}, x^{(k)}) = \sigma(x^{(k)}). \tag{41}$$

If the minimum in (38) is attained for a unique x then $\eta(x^{(k+1)},x^{(k)})<\eta(x^{(k)},x^{(k)})$. If the majorization scheme is strict then $\sigma(x^{(k+1)})<\eta(x^{(k+1)},x^{(k)})$. Under either of these two additional conditions $\sigma(x^{(k+1)})<\sigma(x^{(k)})$, which means that the majorization algorithm is a monotone descent algorithm, and if σ is bounded below on $\mathbb S$ the sequence $\sigma(x^{(k)})$ converges.

Note that we only use the order relation to prove convergence of the sequence of function values. To prove convergence of the $x^{(k)}$ we need stronger compactness and continuity assumptions to apply the general theory of Zangwill (1969). For such a proof the argmin in update formula (38) can be generalized to $x^{(k+1)} = \phi(x^{(k)})$, where ϕ maps $\mathbb S$ into $\mathbb S$ such that $\eta(\phi(x), x) \leq \sigma(x)$ for all x.

We give a small illustration in which we minimize σ with $\sigma(x) = \sqrt{x} - \log x$ over x > 0. Obviously we do not need majorization here, because solving $\mathcal{D}\sigma(x) = 0$ immediately gives x = 4 as the solution we are looking for.

To arrive at this solution using majorization we start with

$$\sqrt{x} \le \sqrt{y} + \frac{1}{2} \frac{x - y}{\sqrt{y}},\tag{42}$$

which is true because a differentiable concave function such as the square root is majorized by its tangent everywhere. Inequality (42) implies

$$\sigma(x) \le \eta(x,y) := \sqrt{y} + \frac{1}{2} \frac{x-y}{\sqrt{y}} - \log x. \tag{43}$$

Note that $\eta(\bullet,y)$ is convex in its first argument for each y. We have $\mathcal{D}_1\eta(x,y)=0$ if and only if $x=2\sqrt{y}$ and thus the majorization algorithm is

$$x^{(k+1)} = 2\sqrt{x^{(k)}} \tag{44}$$

The sequence $x^{(k)}$ converges monotonically to the fixed point $x=2\sqrt{x}$, i.e. to x=4. If $x^{(0)}<4$ the sequence is increasing, if $x^{(0)}<4$ it is decreasing. Also, by l'Hôpital,

$$\lim_{x \to 4} \frac{2\sqrt{x} - 4}{x - 4} = \frac{1}{2} \tag{45}$$

and thus convergence to the minimizer is linear with asymptotic convergence rate $\frac{1}{2}$. By another application of l'Hôpital

$$\lim_{x \to 4} \frac{\sigma(2\sqrt{x}) - \sigma(4)}{\sigma(x) - \sigma(4)} = \frac{1}{4},\tag{46}$$

and convergence to the minimum is linear with asymptotic convergence rate $\frac{1}{4}$. Linear convergence to the minimizer is typical for majorization algorithms, as is the twice-as-fast linear convergence to the minimum value.

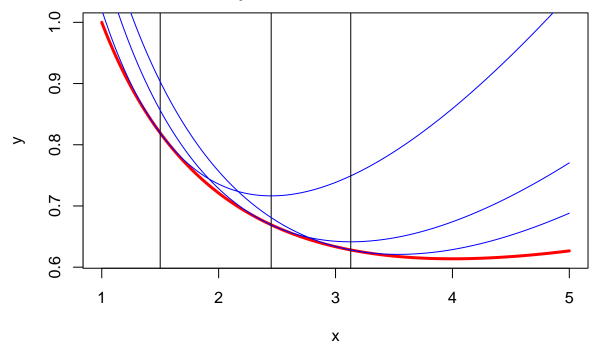
This small example is also of interest, because we minimize a *DC function*, the difference of two convex functions. In our example the convex functions are minus the square root and minus the logarithm. Algorithms for minimizing DC functions define other important subclasses of MM algorithms, the *DC Algorithm* of Tao Pham Dinh (see Le Thi and Tao (2018) for a recent overview),

the *Concave-Convex Procedure* of Yuille and Rangarajan (2003), and the *Half-Quadratic Method* of Donald Geman (see Niikolova and Ng (2005) for a recent overview). For each of these methods there is a huge literature, with surprisingly little non-overlapping literatures. The first phase of the smacof algorithm, in which we improve the configuration for given disparities, is DC, concave-convex, and half-quadratic.

In the table below we show convergence of (44) starting at x=1.5. The first column show how far $x^{(k)}$ deviates from the minimizer (i.e. from 4), the second shows how $and and and and and and and an article from the minimum (i.e. from <math>2-\log 4$). We clearly see the convergence rates $\frac{1}{2}$ and $\frac{1}{4}$ in action.

```
## itel
          1 2.5000000000 0.2055741244
## itel
          2 1.5505102572 0.0554992066
## itel
          3 0.8698308399 0.0144357214
## itel
          4 0.4615431837 0.0036822877
          5 0.2378427379 0.0009299530
## itel
## itel
          6 0.1207437506 0.0002336744
## itel
          7 0.0608344795 0.0000585677
## itel
          8 0.0305337787 0.0000146606
## itel
          9 0.0152961358 0.0000036675
## itel
         10 0.0076553935 0.0000009172
## itel
         11 0.0038295299 0.0000002293
         12 0.0019152235 0.0000000573
## itel
## itel
         13 0.0009577264 0.0000000143
## itel
         14 0.0004788919 0.0000000036
## itel
         15 0.0002394531 0.0000000009
```

The first three iterations are shown in the figure below. The vertical lines indicate the value of x, function is in red, and the first three majorizations are in blue.



7.0.2 Majorizing Stress

8 MDS topics not covered

- asymmetric MDS
- Non-Euclidean Distances
- Non-Least-Squares Loss Functions
- Rank Images
- Shepard
- McGee

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