Multi-sample comparison of detrital age distributions

Pieter Vermeesch *

Department of Earth Sciences, University College London, Gower Street, London WC1E 6BT, UK

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A B S T R A C T

The petrography and geochronology of detrital minerals form rich archives of information pertaining to the provenance of siliclastic sediments. The composition and age spectra of multi-sample datasets can be used to trace the flow of sediments through modern and ancient sediment routing systems. Such studies often involve dozens of samples comprising thousands of measurements. Objective interpretation of such large datasets can be challenging and greatly benefits from dimension-reducing exploratory data analysis tools. Principal components analysis (PCA) is a proven method that has been widely used in the context of compositional data analysis and traditional heavy mineral studies. Unfortunately, PCA cannot be readily applied to geochronological data, which are rapidly overtaking petrographic techniques as the method of choice for large scale provenance studies. This paper proposes another standard statistical technique called multidimensional scaling (MDS) as an appropriate tool to fill this void. MDS is a robust and flexible superset of PCA which makes fewer assumptions about the data. Given a table of pairwise ‘dissimilarities’ between samples, MDS produces a ‘map’ of points on which ‘similar’ samples cluster closely together, and ‘dissimilar’ samples plot far apart. It is shown that the statistical effect size of the Kolmogorov-Smirnov test is a viable dissimilarity measure. This is not the case for the p-values of this and other tests. To aid in the adoption of the method by the geochronological community, this paper includes some simple code using the statistical programming language R. More extensive software tools are provided on http://mudisc.london-geochron.com.

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1. Introduction

Ever since the development of single grain U-Pb dating by (ion and laser) microprobe analysis, the method has been applied to detrital zircon (DZ) as a means of reconstructing the provenance of siliciclastic rocks. Initially, DZ geochronology was primarily used to trace the provenance of such rocks back to individual ‘protosources’ or source terranes (Gehrels et al., 1995; Pell et al., 1997). But in recent years, the ever-increasing throughput and ever decreasing cost of DZ geochronology have enabled a more sophisticated kind of application, in which the U-Pb age distributions of multiple samples are used as a characteristic ‘fingerprint’ to trace the flow of zircon grains through the sediment routing system.

This paper introduces methods that make the interpretation of such datasets more objective, using a recently published provenance study from China as an example. Stevens et al. (in press) present a dataset comprising ten sandstone samples from the Mu Us desert, a Quaternary loess sample, a modern fluvial sand sample from the Yellow River, and a dataset of DZ ages from the Tibetan headwaters of the Yellow River taken from Pullen et al. (2011). The degree of similarity between these samples can be assessed on a qualitative basis by jointly plotting their respective age spectra (Fig. 1). Another commonly used visual aid is the so-called ‘QQ plot’, in which various quantiles of the samples are plotted against each other, the idea being that two samples follow an identical distribution if and only if their quantiles plot on a 1:1-line (Fig. 2).

Both the QQ plots and the age spectra can become unwieldy if they contain more than a dozen or so samples. For example, Fig. 1 contains $n = 13$ kernel density estimates (KDEs, Vermeesch, 2012) showing the probability distributions of 2025 single grain age estimates, while the QQ-plots in Fig. 2 form an upper triangular matrix with $n(n-1)/2 = 78$ pairwise comparisons. This is simply too much information for the human eye to process. To solve this problem, we need a ‘filter’ removing the redundant features of the individual distributions while preserving and amplifying the significant differences between them. This paper makes the case that a standard statistical technique called multidimensional scaling (MDS) can be used effectively for this purpose (Sections 3 and 4).

In addition to the DZ ages, all but one (T) of the samples in the Chinese study were subjected to heavy mineral (HM) analysis. With the exception of samples 1 and 8, the HM analyses were performed on separate aliquots from the U-Pb measurements. For samples 1 and 8, the HM mounts were prepared by mixing leftover mineral separates from the DZ study. Between 201 and 419 grains were counted in the 63–250 μm size fraction of each sample, resulting in an additional 2901 datapoints. Part of the aim of this paper is to treat these categorical data on an equal footing with...
the continuous age data in a consistent mathematical framework (Section 5). All the analyses presented in this paper can be reproduced using the software discussed in Section 6 and made available on http://mudisc.london-geochron.com.

Fig. 1. Sample locations and kernel density estimates (KDEs, Vermeesch, 2012) of the Chinese loess study (N = number of concordant ages). Samples 1–5, 7 and 9–10 — dune sand; samples 6 and 8 — Cretaceous sandstone; sample L — Quaternary loess; sample Y — fluvial sand from the Yellow River; sample T — compilation of proposed sources in Yellow River headwaters from Pullen et al. (2011).

2. Measuring the dissimilarity between two samples

Before discussing multi-sample comparisons, it is useful to review the underlying principles for measuring the dissimilarity ($\delta_{ij}$) between two samples ($i$ and $j$, say). It is desirable for any dissimilarity measure to fulfill the following four requirements:

1. $\delta_{ij}$ should independent of sample size $N$

2. $\delta_{ij} = 0$ if $i = j$ and $\delta_{ij} > 0$ otherwise (nonnegativity)

3. $\delta_{ij} = \delta_{ji}$ (symmetry)
The first requirement is important for the Chinese loess study, as it would be impossible to compare samples 6 (N = 58) and T (N = 772) otherwise. If a dissimilarity measure fulfills the remaining three conditions, then it is said to be a ‘metric’. One example of such a metric is the Euclidean distance (d_{ij}). Given two R-dimensional points x_i = (x_{i1}, x_{i2}, ..., x_{iR}) and x_j = (x_{j1}, x_{j2}, ..., x_{jR}), the Euclidean distance is given by:

\[ d_{ij} = \sqrt{\sum_{k=1}^{R} (x_{ik} - x_{jk})^2}. \]  

Unfortunately, the space of detrital age distributions (which is a ‘function space’) is not a Euclidean space. And neither is the space of HM compositions (which is a ‘simplex’). Therefore, the Euclidean distance cannot be directly applied to either DZ or HM data. However, several alternatives are available that approximate the disparities between samples i and j:

The triangle inequality (Eq. (4)), causing some limitations. Therefore, the remainder of this paper will use the KS effect size as a dissimilarity measure.

3. Multidimensional scaling

Let \( \hat{\delta} \) be a matrix of pairwise dissimilarities between n samples:

\[ \hat{\delta} = \begin{pmatrix} \hat{\delta}_{1,1} & \hat{\delta}_{1,2} & \ldots & \hat{\delta}_{1,n} \\ \hat{\delta}_{2,1} & \hat{\delta}_{2,2} & \ldots & \hat{\delta}_{2,n} \\ \vdots & \vdots & \ddots & \vdots \\ \hat{\delta}_{n,1} & \hat{\delta}_{n,2} & \ldots & \hat{\delta}_{n,n} \end{pmatrix} \]  

where \( \hat{\delta}_{ij} \) is the dissimilarity between samples i and j, as before. And let \( f(\hat{\delta}_{ij}) \) be a monotonically increasing function transforming the dissimilarities into ‘distances’. Then MDS produces an R-dimensional configuration of points \( x = (x_1, x_2, ..., x_n) \) with \( x_i = (x_{i1}, x_{i2}, ..., x_{iR}) \) and \( x_j = (x_{j1}, x_{j2}, ..., x_{jR}) \) for \( 1 \leq i, j \leq n \). The Euclidean distance between any two points \( x_i \) and \( x_j \) in this configuration approximates the disparities between samples i and j:

\[ d_{ij} \approx f(\hat{\delta}_{ij}). \]  

with \( d_{ij} \) as defined by Eq. (5). In this paper, we will only consider the case where \( R = 2 \), so that the configuration \( x \) can be plotted on a sheet of paper, like a map. Therefore, we will use the terms ‘configuration’ and ‘map’ interchangeably. In fact, one of the classical applications of MDS involves the reconstruction of a topographic map from a table of pairwise (Euclidean) distances between cities with the minimum possible distortion (e.g., p. 8 of Kruskal and Wish, 1978).

If the disparity transformation is the identity matrix (i.e., \( f(\hat{\delta}_{ij}) = \hat{\delta}_{ij} \)), and if \( \hat{\delta}_{ij} \) is a metric, then the configuration \( x \) can be found analytically by relatively straightforward linear algebra (Cox and Cox, 2000, p.23-25). This is called classical MDS. If the triangle inequality (Eq. (4)) is violated, then \( \hat{\delta}_{ij} \) is not a metric but a ‘semi-metric’. In this case, it may still be possible to perform an MDS analysis by considering a disparity transformation such as:

\[ f(\hat{\delta}_{ij}) = a + b\hat{\delta}_{ij} \]  
or any other parametric equation (e.g. exponential), so long as it monotonically increases. This superset of classical MDS is (somewhat confusingly) called metric MDS. It solves for the configuration that simultaneously determines both the distances \( d_{ij} \) and the fit to the disparity transformation \( f(\hat{\delta}_{ij}) \). In the most general case, \( f \) is allowed to take any nonparametric monotonically increasing form such as a step function. This method is called nonmetric MDS, and allows not only violations of the triangle inequality but also of the condition of symmetry (Eq. (3)). The goal of nonmetric MDS is not to approximate the dissimilarities themselves, but rather their relative ranks. In the case of non-metric MDS, the solution is not found analytically but numerically. This is done by minimizing a so-called ‘stress parameter’ S, the most common form (so-called ‘Stress-1’) of which is (Kruskal, 1964):

\[ S = \sqrt{\frac{\sum_{i=1}^{n} \sum_{j=1}^{n} \left[ f(\hat{\delta}_{ij}) - d_{ij} \right]^2}{\sum_{i=1}^{n} \sum_{j=i+1}^{n} d_{ij}^2}}. \]  

The final stress value can be used to evaluate the quality of the MDS fit (Table 1).

MDS is generally insensitive to the choice of S and f. Therefore, metric and nonmetric MDS will generally give similar-looking results, and even the choice of \( \hat{\delta}_{ij} \) will often have only a minor effect. This robustness is one of the major strengths of the method.
which is again a symmetric matrix with zero diagonal because the Euclidean distance is a metric and therefore obeys Eqs. (2)–(4). Plotting the (upper- or lower triangular parts of the) distance matrix \(d\) (Eq. (14)) against the corresponding dissimilarities \(\delta\) (Eq. (12)) yields a so-called ‘Shepard plot’, allowing a graphical assessment of the quality of the model fit (Fig. 3b). Doing the same exercise with nonmetric MDS yields a very similar looking map (Fig. 3c), with the fitted distances on the Shepard plot, of course, not following a linear but a step function (Fig. 3d). Both Shepard plots show a tight fit of the distances around the disparities. The relatively minor amount of scatter is reflected in the stress values of 0.064 and 0.025. Using the rules of thumb given in Table 1, this qualifies as a ‘good’ fit for the metric, and an ‘excellent’ fit for the nonmetric MDS, respectively.

The MDS maps group samples with similar age spectra, and pull apart samples with different spectra. For example, samples T (Tibet) and Y (Yellow River) nearly overlap on the MDS maps because the KS dissimilarity between them is only 0.07, which is the smallest value in Eq. (12). On the other hand, samples L (loess) and B (northeastern Mu-Us) plot very far apart on the MDS map because the KS dissimilarity between them is 0.63, which is the largest value in Eq. (12). One simple but effective way to aid in the interpretation of MDS maps is to draw a solid line from each point in the configuration to its ‘closest’ neighbour in dissimilarity-space, and a dotted line to the second closest neighbour. For example, sample 8 is the closest (or ‘least dissimilar’) to sample 7 (KS = 0.16), and the second closest to sample 1 (KS = 0.22), while sample 7 is the closest to sample 6 (KS = 0.14) and second closest to sample 2 (KS = 0.15). These connecting lines define two distinct groups dividing the field area into a southwestern area containing sediments of ‘Yellow River affinity’ (samples 3, 4, 9, 10, L, T and Y) and a northeastern area containing sediments of a different origin (samples 1, 2, 5, 6, 7 and 8). The MDS maps reveal a pronounced East-West dichotomy within the Mu-Us sand desert, in which sands from the eastern part of the desert are locally derived, whereas the western sands are far travelled and exhibit virtually identical age spectra to the fluvial sands of the Yellow River, which in turn are indistinguishable from Quaternary loess of the Chinese Loess Plateau. This leads to two new conclusions. First, the northern parts of the Tibet Plateau are the ‘ultimate source’ of the Chinese loess. Second, fluvial transport is a more significant supplier of silt-sized particles to the Chinese loess deposits than was previously recognised (Stevens et al., in review).

### 5. Relationship with principal components analysis (PCA)

Principal components analysis (PCA) is one of the most widely used dimension-reducing exploratory data analysis tools for compositional data such as HM counts. Given a multi-dimensional dataset, PCA finds an orthogonal transformation (i.e., a rotation, reflection or both) to define a lower-dimensional set of new variables explaining most of the scatter in the input data. PCA is closely related to MDS. In fact, it can be shown that PCA is a special case of classical MDS in which the dissimilarities are Euclidean distances. This means that \(\delta_{ij}\) can be calculated in exactly the same way as \(d_{ij}\) in Eq. (5). The best way to measure the dissimilarity between two HM compositions is the so-called ‘Aitchison distance’. As mentioned in Section 2 and explained in Appendix A, this distance is calculated by first applying a data transformation and then calculating the Euclidean distance on the transformed data. Therefore, an MDS analysis of the Aitchison distance is identical to a PCA of the transformed data (Fig. 4). One common way to visualise PCA results is as a ‘biplot’ which jointly shows the configuration and the endmember compositions. This reveals that the most important differences between the two groups are found in their epidote and amphibole contents. The MDS/PCA map of the HM counts has a striking resemblance to the MDS map of the DZ ages (Fig. 3). Both maps are organised into the same two groups. Therefore, not only the zircons but also the other heavy minerals exhibit the aforementioned East-West dichotomy, indicating
limited aeolian mixing within the Mu-Us desert. Only sample 1 has 'switched groups', possibly due in part to the fact that it was 'reconstructed' from mineral separates.

6. Software

Given that MDS is a standard statistical technique, numerous software tools are available that can easily be adopted for detrital geochronology, including several open source options, such as R (http://www.r-project.org). In R, classical MDS is implemented by the cmdscale function of the stats package, while nonmetric MDS is implemented by the isoMDS function of the MASS package. Appendix C contains a simple snippet of code using the Chinese DZ dataset as an example. PCA of compositional data is implemented in R by the princomp function of the compositions package. In Matlab, both classical, metric and non-metric MDS are implemented by the mdscalc function of the Statistics Toolbox. A graphical user interface has been created that implements this function in a user-friendly way and can be downloaded from http://mudisc.london-geochron.com along with further details about the R code, input files and so forth.

7. Concluding remarks

This paper introduced MDS as a simple yet powerful tool providing informative, mutually consistent measures of difference for a group of observations taken together. It is important to note that, just like any other dimension-reducing technique, MDS makes an abstraction of the data and may not always represent all the details of complex datasets. One problem with dissimilarity measures such as the KS effect size is that they are simple scalars which cannot capture the full richness of entire probability distributions. For example, the KS test cannot see the difference between a 'comb-like' and a 'flat' density. Despite these limitations, the MDS map often captures the main features of detrital zircon datasets, as illustrated in the Chinese case study.

It was shown that PCA is a special case of classical MDS, using Euclidean distance for dissimilarities and thus making restrictive assumptions about the structure of the data, which are relaxed by metric and non-metric MDS. Thus, MDS is a flexible technique which makes fewer assumptions about the data than PCA. The user is not restricted to the KS effect size used in this paper. Other statistic-based dissimilarities such as Cramér–von Mises and Anderson–Darling would also work, although it is advised to use nonmetric MDS in these cases as

![MDS plots](image-url)
Principal components analysis (PCA) of the logratio-transformed HM data from China. This plot is identical to the classical MDS map of the Aitchison distances. The blue lines show the endmember compositions resulting from the PCA. The combination of the endmember compositions and the configuration is called a 'biplot'. Asterisks mark compositions measured on mixtures of mineral separates rather than separate aliquots.

...this is the most robust method. Nonmetric MDS usually yields better fits than metric MDS. In the Chinese case study, this is reflected in the stress value of S = 0.025 for the former and S = 0.064 for the latter (Fig. 3). The stress of the corresponding classical MDS analysis is even higher at S = 0.78. It would therefore seem that nonmetric MDS is the best method. However, the nonmetric MDS algorithm may fail for small datasets. In these cases, metric and classical MDS provide a more stable backup solution.

One aspect of detrital geochronology which this paper has ignored is analytical uncertainty. An implicit assumption behind the use of the KS effect size as a measure of dissimilarity is that all the samples under consideration are characterised by the same analytical precision. This is a valid assumption for the Chinese data, all of which were measured by LA-ICP-MS (and all but one of which were analysed in the same lab). But it is not necessarily true for datasets combining age spectra from different laboratories using a combination of LA-ICP-MS and SIMS, say. In such cases, it would be advised to remove the analytical effects first. Sircombe and Hazelton (2004) show that this can be achieved by adding an arbitrary variance term to all the measurements. The resulting ‘kernel functional estimates’ (KFE) are, essentially, over-smoothed KDEs that can be objectively compared using the methods described in this paper. Finally, although this paper focused on detrital zircon U–Pb age spectra, the conclusions follow for any other mineral (such as rutile, apatite, or monazite) or dating technique (such as $^{40}\text{Ar}/^{39}\text{Ar}$, fission tracks, or U–Th–He). In the rapidly changing field of detrital geo- and thermochronology, the multi-sample comparison method described here will be able to inform more evolved efforts of the future.

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Appendix A. Brief summary of the Aitchison geometry applied to PCA

Let $y = [y_1, y_2, \ldots, y_k]$ be a set of k-dimensional point-counting data

$y_a = [y_a^1, y_a^2, \ldots, y_a^k]$, so that $y_a^b$ represents the integer number of grains of mineral $b$ counted in sample $a$. The most straightforward way to measure the dissimilarity between two samples ($i$ and $j$, say) would appear to be the Euclidean distance between the corresponding proportions:

$$d_{ij} = \sqrt{\sum_{b=1}^{k} \left( \frac{y_i^b}{\sum_{c=1}^{k} y_i^c} - \frac{y_j^b}{\sum_{c=1}^{k} y_j^c} \right)^2}.$$  \hfill (A-1)

Unfortunately, this approach is wrong because the proportions are subject to a constant sum constraint (all fractions must add up to 100%) and are therefore not mutually independent. This restriction produces geometric artefacts (curvature) which may negatively affect linear procedures such as PCA. Note that this also invalidates prior attempts to apply PCA to detrital age histograms (Sircombe, 2000). To solve this problem, Aitchison (1982, 1983, 1986) developed a mathematical formalism in which the compositional data are transformed to an ordinary Euclidean space using a (centred) log-ratio transformation ($y_a^b \rightarrow z_a^b$), by normalising the compositions to their respective geometric means and taking logarithms:

$$z_a^b = \ln \left( \frac{y_a^b}{\sqrt{\prod_{c=1}^{k} y_a^c}} \right)$$  \hfill (A-2)

with $a$ and $b$ referring to the sample and mineral, as before. This logratio transformation removes the geometric artefacts caused by the closure operation so that the transformed data can be subjected to a PCA. Instead of the centred logratio transformation, other options are available such as the isometric logratio transformation (ilr), which may actually be even better suited for PCA/MDS (Egozcue et al., 2003; Filzmoser et al., 2009).

Appendix B. Why p-values are unsuitable as a measure of dissimilarity

Some workers have used statistical tests such as Kolmogorov–Smirnov to make the comparison of detrital age distributions more ‘objective’ than mere visual inspection of age spectra or QQ-plots. Such statistical tests require the formulation of a so-called ‘null hypothesis’ ($H_0$, e.g., “two detrital age distributions were drawn from the same population”), and the selection of a ‘test statistic’ (e.g., Eq. (6)). If the observed value of the test statistic is ‘unlikely’ to occur under the null hypothesis, then the latter is rejected in favour of the alternative hypothesis ($H_a$: “two detrital age distributions were drawn from different populations”). The probability of observing a value at least as extreme as the observed statistic under the null hypothesis is called the ‘p-value’. An arbitrary cut-off value $\alpha$ may be used to make a decision on a 100(1–$\alpha$)% confidence level. For example, if $\alpha = 0.05$ and $p < \alpha$, then $H_0$ is rejected in favour of $H_a$ with 95% confidence. The probability of erroneously rejecting a true null hypothesis, which is given by $\alpha$, is also called a ‘Type-I error’. In other words, the $\alpha$-value expresses the risk we are willing to take of committing a Type-I error. The p-value is a poor measure of dissimilarity between samples, because it lumps together two factors: the ‘effect size’ and the ‘sample size’. The sample size is simply the total number of grains analysed ($N$). The effect size expresses “the degree to which the null hypothesis is false” (Cohen, 1977). To illustrate this concept, consider the case of two Normally distributed detrital age populations with means $\mu_1$ and $\mu_2$, variances $\sigma_1^2$ and $\sigma_2^2$, and sample sizes $N_1$ and $N_2$, respectively. Assume that $N_1 = N_2 = N$ and $\sigma_1 = \sigma_2 = \sigma$. Let $X_1$ and $X_2$ be the sample means and $S_1^2$ and $S_2^2$...
Appendix C. Multidimensional scaling with R

This section presents a simple example of computer code for MDS analysis with the open source statistical programming language R. The example input file DZages.Rdata containing the data shown in Fig. 1 can be downloaded from http://mudisc.london-geochron.com.

Table 2
Statistic (t) and p-value (p) of a t-test with sample effect size \(d = 0.1\) as a function of sample size (N). The strong dependence of t and p on N indicates that both parameters are unsuitable as dissimilarity measures for MDS analysis.

<table>
<thead>
<tr>
<th>N</th>
<th>10</th>
<th>100</th>
<th>1000</th>
<th>10,000</th>
</tr>
</thead>
<tbody>
<tr>
<td>t</td>
<td>1/\sqrt{10}</td>
<td>1</td>
<td>\sqrt{10}</td>
<td>100</td>
</tr>
<tr>
<td>p</td>
<td>0.41</td>
<td>0.24</td>
<td>0.013</td>
<td>8 \times 10^{-13}</td>
</tr>
</tbody>
</table>

The sample variances. We can test the null hypothesis \(H_0: \mu_1 = \mu_2\) with the t-test, using the t-statistic:

\[
t = \frac{X_1 - X_2}{\sqrt{(S_1^2 + S_2^2)/N}}
\]

where the denominator is the standard error of the difference between the two means. The sample effect size of the t-test is given by:

\[
d = \frac{X_1 - X_2}{\sqrt{(S_1^2 + S_2^2)/2}}
\]

where the denominator is the ‘pooled standard deviation’ of the two samples. By definition, \(d = 0\) under the null hypothesis. Let us now consider the situation where \(d \neq 0\). Suppose that \(\mu_1 = 100.1\) Ma, \(\mu_2 = 100.0\) Ma, and \(\sigma = 1\) Ma so that the population effect size is 0.1. Suppose for the sake of simplicity that these population characteristics are reflected in the sample so that \(d = 0.1\). Table 2 shows some key values of the t-statistic and corresponding p-values for different sample sizes. Not surprisingly, a sample size of \(N = 10\) is insufficient to resolve the 0.1 Ma ‘offset’ between the two peaks. In this case, the p-value is 0.41, which is greater than the cutoff value of \(\alpha = 0.05\), leading us to erroneously accept the false null hypothesis. This means that we have committed a ‘Type-II error’. It is only when sample size increases to \(N = 1000\) that the t-test has sufficient ‘power’ to reject the null hypothesis \((0.013 < \alpha)\). In conclusion, both the statistic and the p-value of the t-test are sensitive functions of sample size. Both are therefore unsuitable as measures of dissimilarity in an MDS analysis.

These same arguments are valid for other statistical tests such as Chi-square and Kolmogorov–Smirnov, with the caveat that the effect size of the KS-test is given by the KS-statistic itself (Eq. (6)). Nevertheless, the p-value of the KS-test is still a function of sample size and is therefore still useless as a dissimilarity measure.

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