Minimum Covariance Determinant and Extensions

Mia Hubert*, Michiel Debruyne†, Peter J. Rousseeuw*

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Abstract

The Minimum Covariance Determinant (MCD) method is a highly robust estimator of multivariate location and scatter, for which a fast algorithm is available. Since estimating the covariance matrix is the cornerstone of many multivariate statistical methods, the MCD is an important building block when developing robust multivariate techniques. It also serves as a convenient and efficient tool for outlier detection.

The MCD estimator is reviewed, along with its main properties such as affine equivariance, breakdown value, and influence function. We discuss its computation, and list applications and extensions of the MCD in applied and methodological multivariate statistics. Two recent extensions of the MCD are described. The first one is a fast deterministic algorithm which inherits the robustness of the MCD while being almost affine equivariant. The second is tailored to high-dimensional data, possibly with more dimensions than cases, and incorporates regularization to prevent singular matrices.

*Department of Mathematics, KU Leuven, Celestijnenlaan 200B, BE-3001 Leuven, Belgium
†Dexia Bank, Belgium
INTRODUCTION

The Minimum Covariance Determinant (MCD) estimator is one of the first affine equivariant and highly robust estimators of multivariate location and scatter\cite{1,2}. Being resistant to outlying observations makes the MCD very useful for outlier detection. Although already introduced in 1984, its main use has only started since the construction of the computationally efficient FastMCD algorithm of\cite{3} in 1999. Since then, the MCD has been applied in numerous fields such as medicine, finance, image analysis and chemistry. Moreover the MCD has also been used to develop many robust multivariate techniques, among which robust principal component analysis, factor analysis and multiple regression. Recent modifications of the MCD include a deterministic algorithm and a regularized version for high-dimensional data.

DESCRIPTION OF THE MCD ESTIMATOR

Motivation

In the multivariate location and scatter setting the data are stored in an $n \times p$ data matrix $X = (x_1, \ldots, x_n)'$ with $x_i = (x_{i1}, \ldots, x_{ip})'$ the $i$-th observation, so $n$ stands for the number of objects and $p$ for the number of variables. We assume that the observations are sampled from an elliptically symmetric unimodal distribution with unknown parameters $\mu$ and $\Sigma$, where $\mu$ is a vector with $p$ components and $\Sigma$ is a positive definite $p \times p$ matrix. To be precise, a multivariate distribution is called elliptically symmetric and unimodal if there exists a strictly decreasing real function $g$ such that the density can be written in the form

$$f(x) = \frac{1}{\sqrt{\det|\Sigma|}} g(d^2(x, \mu, \Sigma))$$  \hspace{1cm} (1)

in which the statistical distance $d(x, \mu, \Sigma)$ is given by

$$d(x, \mu, \Sigma) = \sqrt{(x - \mu)'\Sigma^{-1}(x - \mu)}.$$  \hspace{1cm} (2)

To illustrate the MCD, we first consider the wine data set available in\cite{4} and also analyzed in\cite{5}. This data set contains the quantities of 13 constituents found in three types of Italian wines. We consider the first group containing 59 wines, and focus on the constituents ‘Malic...
acid’ and ‘Proline’. This yields a bivariate data set, i.e. \( p = 2 \). A scatter plot of the data is shown in Figure 1, in which we see that the points on the lower right hand side of the plot are outlying relative to the majority of the data.

![Classical and robust tolerance ellipse](image)

Figure 1: Bivariate wine data: tolerance ellipse of the classical mean and covariance matrix (red), and that of the robust location and scatter matrix (blue).

In the figure we see two ellipses. The classical tolerance ellipse is defined as the set of \( p \)-dimensional points \( \mathbf{x} \) whose Mahalanobis distance

\[
\text{MD}(\mathbf{x}) = d(\mathbf{x}, \bar{x}, \text{Cov}(X)) = \sqrt{(\mathbf{x} - \bar{x})'\text{Cov}(X)^{-1}(\mathbf{x} - \bar{x})}
\]

equals \( \chi^2_{p,0.975} \). Here \( \bar{x} \) is the sample mean and \( \text{Cov}(X) \) the sample covariance matrix. The Mahalanobis distance \( \text{MD}(\mathbf{x}_i) \) should tell us how far away \( \mathbf{x}_i \) is from the center of the data cloud, relative to its size and shape. In Figure 1 we see that the red tolerance ellipse tries to encompass all observations. Therefore none of the Mahalanobis distances is exceptionally large, as we can see in Figure 2(a). Based on Figure 2(a) alone we would say there are only three mild outliers in the data (we ignore borderline cases).
On the other hand, the robust tolerance ellipse is based on the robust distances

$$\text{RD}(\mathbf{x}) = d(\mathbf{x}, \hat{\mu}_{MCD}, \hat{\Sigma}_{MCD})$$  \hspace{1cm} (4)$$

where $\hat{\mu}_{MCD}$ is the MCD estimate of location and $\hat{\Sigma}_{MCD}$ is the MCD covariance estimate, which we will explain soon. In Figure 1 we see that the robust ellipse (in blue) is much smaller and only encloses the regular data points. The robust distances shown in Figure 2(b) now clearly expose 8 outliers.

This illustrates the *masking effect*: the classical estimates can be so strongly affected by contamination that diagnostic tools such as the Mahalanobis distances become unable to detect the outliers. To avoid masking we instead need reliable estimators that can resist outliers when they occur. The MCD is such a robust estimator.

**Definition**

The raw Minimum Covariance Determinant (MCD) estimator with tuning constant $n/2 \leq h \leq n$ is $(\hat{\mu}_0, \hat{\Sigma}_0)$ where
1. The location estimate $\hat{\mu}_0$ is the mean of the $h$ observations for which the determinant of the sample covariance matrix is as small as possible;

2. The scatter matrix estimate $\hat{\Sigma}_0$ is the corresponding covariance matrix multiplied by a consistency factor $c_0$.

Note that the MCD estimator can only be computed when $h > p$, otherwise the covariance matrix of any $h$-subset has determinant zero, so we need at least $n > 2p$. To avoid excessive noise it is however recommended that $n > 5p$, so that we have at least 5 observations per dimension. (When this condition is not satisfied one can instead use the MRCD method (11) described near the end of this article.) To obtain consistency at the normal distribution, the consistency factor $c_0$ equals $\alpha / F_{\chi^2_p}(q_\alpha)$ with $\alpha = \lim_{n \to \infty} h(n) / n$, and $q_\alpha$ the $\alpha$-quantile of the $\chi^2_p$ distribution. Also a finite-sample correction factor can be incorporated.

Consistency of the raw MCD estimator of location and scatter at elliptical models, as well as asymptotic normality of the MCD location estimator has been proved in. Consistency and asymptotic normality of the MCD covariance matrix at a broader class of distributions is derived in.

The MCD estimator is the most robust when taking $h = [(n + p + 1)/2]$ where $[a]$ is the largest integer $\leq a$. At the population level this corresponds to $\alpha = 0.5$. But unfortunately the MCD then suffers from low efficiency at the normal model. For example, if $\alpha = 0.5$ the asymptotic relative efficiency of the diagonal elements of the MCD scatter matrix relative to the sample covariance matrix is only 6% when $p = 2$, and 20.5% when $p = 10$. This efficiency can be increased by considering a higher $\alpha$ such as $\alpha = 0.75$. This yields relative efficiencies of 26.2% for $p = 2$ and 45.9% for $p = 10$ (see). On the other hand this choice of $\alpha$ diminishes the robustness to possible outliers.

In order to increase the efficiency while retaining high robustness one can apply a weighting step. For the MCD this yields the estimates

$$\hat{\mu}_{MCD} = \frac{\sum_{i=1}^{n} W(d_i^2) x_i}{\sum_{i=1}^{n} W(d_i^2)}$$

$$\hat{\Sigma}_{MCD} = c_1 n \frac{1}{n} \sum_{i=1}^{n} W(d_i^2) (x_i - \hat{\mu}_{MCD})(x_i - \hat{\mu}_{MCD})'$$

(5)
with \( d_i = d(x, \hat{\mu}_0, \hat{\Sigma}_0) \) and \( W \) an appropriate weight function. The constant \( c_1 \) is again a consistency factor. A simple yet effective choice for \( W \) is to set it to 1 when the robust distance is below the cutoff \( \sqrt{\chi^2_{p,0.975}} \) and to zero otherwise, that is, \( W(d^2) = I(d^2 \leq \chi^2_{p,0.975}) \). This is the default choice in the current implementations in R, SAS, Matlab and S-PLUS. If we take \( \alpha = 0.5 \) this weighting step increases the efficiency to 45.5\% for \( p = 2 \) and to 82\% for \( p = 10 \). In the example of the wine data (Figure 1) we applied the weighted MCD estimator with \( \alpha = 0.75 \), but the results were similar for smaller values of \( \alpha \).

Note that one can construct a robust correlation matrix from the MCD scatter matrix. The robust correlation between variables \( X_i \) and \( X_j \) is given by

\[
    r_{ij} = \frac{s_{ij}}{\sqrt{s_{ii}s_{jj}}}
\]

with \( s_{ij} \) the \((i,j)\)-th element of the MCD scatter matrix. In Figure 1 the MCD-based robust correlation is \( 0.10 \approx 0 \) because the majority of the data do not show a trend, whereas the classical correlation of \( -0.37 \) was caused by the outliers in the lower right part of the plot.

**Outlier detection**

As already illustrated in Figure 2, the robust MCD estimator is very useful to detect outliers in multivariate data. As the robust distances (4) are not sensitive to the masking effect, they can be used to flag the outliers\(^{13,14} \). This is crucial for data sets in more than three dimensions, which are difficult to visualize.

We illustrate the outlier detection potential of the MCD on the full wine data set, with all \( p = 13 \) variables. The *distance-distance plot* of\(^3 \) in Figure 3 shows the robust distances based on the MCD versus the classical distances (3). From the robust analysis we see that seven observations clearly stand out (plus some mild outliers), whereas the classical analysis does not flag any of them.
Figure 3: Distance-distance plot of the full 13-dimensional wine data set.

Note that the cutoff value $\sqrt{\chi^2_{p,0.975}}$ is based on the asymptotic distribution of the robust distances, and often flags too many observations as outlying. For relatively small $n$ the true distribution of the robust distances can be better approximated by an $F$-distribution, see$^{15}$.

**PROPERTIES**

**Affine equivariance**

The MCD estimator of location and scatter is *affine equivariant*. This means that for any nonsingular $p \times p$ matrix $A$ and any $p$-dimensional column vector $b$ it holds that

$$
\hat{\mu}_{MCD}(XA' + 1_nb') = \hat{\mu}_{MCD}(X)A' + b
$$

(i)

$$
\hat{\Sigma}_{MCD}(XA' + 1_nb') = A\hat{\Sigma}_{MCD}(X)A'
$$

(ii)

where the vector $1_n$ is $(1, 1, \ldots, 1)'$ with $n$ elements. This property follows from the fact that for each subset $H$ of $\{1, 2, \ldots, n\}$ of size $h$ and corresponding data set $X_H$, the determinant
of the covariance matrix of the transformed data equals
\[ |S(X_H A')| = |A S(X_H) A'| = |A|^2 |S(X_H)|. \]

Therefore, transforming an \( h \)-subset with lowest determinant yields an \( h \)-subset \( X_H A' \) with lowest determinant among all \( h \)-subsets of the transformed data set \( X A' \), and its covariance matrix is transformed appropriately. The affine equivariance of the raw MCD location estimator follows from the equivariance of the sample mean. Finally we note that the robust distances \( d_i = d(x, \hat{\mu}_0, \hat{\Sigma}_0) \) are affine invariant, meaning they stay the same after transforming the data, which implies that the weighted estimator is affine equivariant too.

Affine equivariance implies that the estimator transforms well under any non-singular reparametrization of the space in which the \( x_i \) live. Consequently, the data might be rotated, translated or rescaled (for example through a change of measurement units) without affecting the outlier detection diagnostics.

The MCD is one of the first high-breakdown affine equivariant estimators of location and scatter, and was only preceded by the Stahel-Donoho estimator\(^{16,17}\). Together with the MCD also the Minimum Volume Ellipsoid estimator was introduced\(^{1,2}\) which is equally robust but not asymptotically normal, and is harder to compute than the MCD.

**Breakdown value**

The breakdown value of an estimator is the smallest fraction of observations that need to be replaced (by arbitrary values) to make the estimate useless. For a multivariate *location* estimator \( T_n \) the breakdown value is defined as
\[
\varepsilon_n(T_n; X_n) = \frac{1}{n} \min \{ m : \sup \| T_n(X_{n,m}) - T_n(X_n) \| = +\infty \}
\]

where \( 1 \leq m \leq n \) and the supremum is over all data sets \( X_{n,m} \) obtained by replacing any \( m \) data points \( x_{i_1}, \ldots, x_{i_m} \) of \( X_n \) by arbitrary points.

For a multivariate *scatter* estimator \( C_n \) we set
\[
\varepsilon_n(C_n; X_n) = \frac{1}{n} \min \{ m : \sup \max_i |\log(\lambda_i(C_{n,m}(X_n))) - \log(\lambda_i(C_n(X_n)))| = +\infty \}
\]

with \( \lambda_1(C_n) \geq \ldots \geq \lambda_p(C_n) > 0 \) the eigenvalues of \( C_n \). This means that we consider a scatter estimator to be broken when \( \lambda_1 \) can become arbitrarily large (‘explosion’) and/or \( \lambda_p \)
can become arbitrary close to 0 (‘implosion’). Implosion is a problem because it makes the scatter matrix singular whereas in many situations its inverse is required, e.g. in (4).

Let \( k(X_n) \) denote the highest number of observations in the data set that lie on an affine hyperplane in \( p \)-dimensional space, and assume \( k(X_n) < h \). Then the raw MCD estimator of location and scatter satisfies

\[
\varepsilon^*_n(\hat{\mu}_0; X_n) = \varepsilon^*_n(\hat{\Sigma}_0; X_n) = \frac{\min(n - h + 1, h - k(X_n))}{n}.
\]

If the data are sampled from a continuous distribution, then almost surely \( k(X_n) = p \) which is called general position. Then \( \varepsilon^*_n(\mu_0; X_n) = \varepsilon^*_n(\Sigma_0; X_n) = \min(n - h + 1, h - p)/n \), and consequently any \( [(n+p)/2] \leq h \leq [(n+p+1)/2] \) gives the breakdown value \( [(n-p+1)/2] \). This is the highest possible breakdown value for affine equivariant scatter estimators at data sets in general position. Also for affine equivariant location estimators the upper bound on the breakdown value is \( [(n - p + 1)/2] \) under natural regularity conditions. Note that in the limit \( \lim_{n \to \infty} \varepsilon^*_n = \min(1 - \alpha, \alpha) \) which is maximal for \( \alpha = 0.5 \).

Finally we note that the breakdown value of the weighted MCD estimators \( \hat{\mu}_{MCD} \) and \( \hat{\Sigma}_{MCD} \) is not lower than the breakdown value of the raw MCD estimator, as long as the weight function \( W \) used in (5) is bounded and becomes zero for large \( d_i \), see

\textbf{Influence function}

The influence function of an estimator measures the effect of a small (infinitesimal) fraction of outliers placed at a given point. It is defined at the population level hence it requires the functional form of the estimator \( T \), which maps a distribution \( F \) to a value \( T(F) \) in the parameter space. For multivariate location this parameter space is \( \text{IR}^p \), whereas for multivariate scatter the parameter space is the set of all positive semidefinite \( p \times p \) matrices. The influence function of the estimator \( T \) at the distribution \( F \) in a point \( \boldsymbol{x} \) is then defined as

\[
IF(\boldsymbol{x}, T, F) = \lim_{\varepsilon \to 0} \frac{T(F_{\varepsilon}) - T(F)}{\varepsilon}
\]

with \( F_{\varepsilon} = (1 - \varepsilon)F + \varepsilon \Delta_{\boldsymbol{x}} \) a contaminated distribution with point mass in \( \boldsymbol{x} \).

The influence function of the raw and the weighted MCD has been computed in\(^6\) and turns out to be bounded. This is a desirable property for robust estimators, as it limits
the effect of a small fraction of outliers on the estimate. At the standard multivariate normal distribution, the influence function of the MCD location estimator becomes zero for all \( x \) with \( \| x \|^2 > \chi^2_{p, \alpha} \) hence far outliers do not influence the estimates at all. The same happens with the off-diagonal elements of the MCD scatter estimator. On the other hand, the influence function of the diagonal elements remains constant (different from zero) when \( \| x \|^2 \) is sufficiently large. Therefore the outliers still have a bounded influence on the estimator. All these influence functions are smooth, except at those \( x \) with \( \| x \|^2 = \chi^2_{p, \alpha} \). The weighted MCD estimator has an additional jump in \( \| x \|^2 = \chi^2_{p, 0.975} \) due to the discontinuity of the weight function, but one could use a smooth weight function instead.

Univariate MCD

For univariate data \( x_1, \ldots, x_n \) the MCD estimates reduce to the mean and the standard deviation of the \( h \)-subset with smallest variance. They can be computed in \( O(n \log n) \) time by sorting the observations and only considering contiguous \( h \)-subsets so that their means and variances can be calculated recursively\(^{22}\). Their consistency and asymptotic normality is proved in\(^ {2,23}\). For \( h = \lfloor n/2 \rfloor + 1 \) the MCD location estimator has breakdown value \( [(n + 1)/2]/n \) and the MCD scale estimator has \( [n/2]/n \). These are the highest breakdown values that can be attained by univariate affine equivariant estimators\(^ {24}\). The univariate MCD estimators also have bounded influence functions, see\(^ 6\) for details. Their maximal asymptotic bias is studied in\(^ {25,26}\) as a function of the contamination fraction.

Note that in the univariate case the MCD estimator corresponds to the Least Trimmed Squares (LTS) regression estimator\(^ 1\), which is defined by

\[
\hat{\beta}_{LTS} = \arg\min_{\mu} \sum_{i=1}^{h} (r_{\beta}^2)_{i:n}
\]

where \( (r_{\beta}^2)_{1:n} \leq (r_{\beta}^2)_{2:n} \leq \ldots \leq (r_{\beta}^2)_{n:n} \) are the ordered squared residuals. For univariate data these residuals are simply \( (r_{\beta})_i = x_i - \beta \).
COMPUTATION

The exact MCD estimator is very hard to compute, as it requires the evaluation of all $\binom{n}{h}$ subsets of size $h$. Therefore one switches to an approximate algorithm such as the FastMCD algorithm of\(^3\) which is quite efficient. The key component of the algorithm is the C-step:

Theorem. Take $X = \{x_1, \ldots, x_n\}$ and let $H_1 \subset \{1, \ldots, n\}$ be a subset of size $h$. Put $\hat{\mu}_1$ and $\hat{\Sigma}_1$ the empirical mean and covariance matrix of the data in $H_1$. If $|\hat{\Sigma}_1| \neq 0$ define the relative distances $d_1(i) := d(x_i, \hat{\mu}_1, \hat{\Sigma}_1)$ for $i = 1, \ldots, n$. Now take $H_2$ such that $\{d_1(i); i \in H_2\} := \{(d_1)_1:n, \ldots, (d_1)_h:n\}$ where $(d_1)_1:n \leq (d_1)_2:n \leq \cdots \leq (d_1)_{n:n}$ are the ordered distances, and compute $\hat{\mu}_2$ and $\hat{\Sigma}_2$ based on $H_2$. Then

$$|\hat{\Sigma}_2| \leq |\hat{\Sigma}_1|$$

with equality if and only if $\hat{\mu}_2 = \hat{\mu}_1$ and $\hat{\Sigma}_2 = \hat{\Sigma}_1$.

If $|\hat{\Sigma}_1| > 0$, the C-step thus easily yields a new $h$-subset with lower covariance determinant. Note that the C stands for ‘concentration’ since $\hat{\Sigma}_2$ is more concentrated (has a lower determinant) than $\hat{\Sigma}_1$. The condition $|\hat{\Sigma}_1| \neq 0$ in the theorem is no real restriction because if $|\hat{\Sigma}_1| = 0$ the minimal objective value is already attained (and in fact the $h$-subset $H_1$ lies on an affine hyperplane).

C-steps can be iterated until $|\hat{\Sigma}_{\text{new}}| = |\hat{\Sigma}_{\text{old}}|$. The sequence of determinants obtained in this way must converge in a finite number of steps because there are only finitely many $h$-subsets, and in practice converges quickly. However, there is no guarantee that the final value $|\hat{\Sigma}_{\text{new}}|$ of the iteration process is the global minimum of the MCD objective function. Therefore an approximate MCD solution can be obtained by taking many initial choices of $H_1$ and applying C-steps to each, keeping the solution with lowest determinant.

To construct an initial subset $H_1$ one draws a random $(p + 1)$-subset $J$ and computes its empirical mean $\hat{\mu}_0$ and covariance matrix $\hat{\Sigma}_0$. (If $|\hat{\Sigma}_0| = 0$ then $J$ can be extended by adding observations until $|\hat{\Sigma}_0| > 0$.) Then the distances $d_0^2(i) := d^2(x_i, \hat{\mu}_0, \hat{\Sigma}_0)$ are computed for $i = 1, \ldots, n$ and sorted. The initial subset $H_1$ then consists of the $h$ observations with smallest distance $d_0$. This method yields better initial subsets than drawing random $h$-
subsets directly, because the probability of drawing an outlier-free \((p + 1)\)-subset is much higher than that of drawing an outlier-free \(h\)-subset.

The FastMCD algorithm contains several computational improvements. Since each C-step involves the calculation of a covariance matrix, its determinant and the corresponding distances, using fewer C-steps considerably improves the speed of the algorithm. It turns out that after two C-steps, many runs that will lead to the global minimum already have a rather small determinant. Therefore, the number of C-steps is reduced by applying only two C-steps to each initial subset and selecting the 10 subsets with lowest determinants. Only for these 10 subsets further C-steps are taken until convergence.

This procedure is very fast for small sample sizes \(n\), but when \(n\) grows the computation time increases due to the \(n\) distances that need to be calculated in each C-step. For large \(n\) FastMCD partitions the data set, which avoids doing all calculations on the entire data set.

Note that the FastMCD algorithm is itself affine equivariant. Implementations of the FastMCD algorithm are available in R (as part of the packages \texttt{rrcov}, \texttt{robust} and \texttt{robustbase}), in SAS/IML Version 7 and SAS Version 9 (in \texttt{PROC ROBUSTREG}), and in S-PLUS (as the built-in function \texttt{cov.mcd}). There is also a Matlab version in LIBRA, a LIBrary for Robust Analysis\textsuperscript{27,28} which can be downloaded from \url{http://wis.kuleuven.be/stat/robust}. Moreover, it is available in the PLS toolbox of Eigenvector Research (\url{http://www.eigenvector.com}). Note that some MCD functions use \(\alpha = 0.5\) by default, yielding a breakdown value of 50\%, whereas other implementations use \(\alpha = 0.75\). Of course \(\alpha\) can always be set by the user.

**APPLICATIONS**

There are many applications of the MCD, for instance in finance and econometrics\textsuperscript{29–31}, medicine\textsuperscript{32}, quality control\textsuperscript{33}, geophysics\textsuperscript{35}, geochemistry\textsuperscript{34}, image analysis\textsuperscript{36,37} and chemistry\textsuperscript{38}, but this list is far from complete.
MCD-BASED MULTIVARIATE METHODS

Many multivariate statistical methods rely on covariance estimation, hence the MCD estimator is well-suited for constructing robust multivariate techniques. Moreover, the trimming idea of the MCD and the C-step have been generalized to many new estimators. Here we list some applications and extensions.

The MCD analog in regression is the Least Trimmed Squares regression estimator which minimizes the sum of the $h$ smallest squared residuals (10). Equivalently, the LTS estimate corresponds to the least squares fit of the $h$-subset with smallest sum of squared residuals. The FastLTS algorithm uses techniques similar to FastMCD. The outlier map introduced in plots the robust regression residuals versus the robust distances of the predictors, and is very useful for classifying outliers, see also.

Moreover, MCD-based robust distances are also useful for robust linear regression, regression with continuous and categorical regressors, and for logistic regression. In the multivariate regression setting (that is, with several response variables) the MCD can be used directly to obtain MCD-regression, whereas MCD applied to the residuals leads to multivariate LTS estimation. Robust errors-in-variables regression is proposed in.

Covariance estimation is also important in principal component analysis and related methods. For low-dimensional data (with $n > 5p$) the principal components can be obtained as the eigenvectors of the MCD scatter matrix, and robust factor analysis based on the MCD has been studied in. The MCD was also used for invariant coordinate selection. Robust canonical correlation is proposed in. For high-dimensional data, projection pursuit ideas combined with the MCD results in the ROBPCA method for robust PCA. In turn ROBPCA has led to the construction of robust Principal Component Regression and robust Partial Least Squares Regression, together with appropriate outlier maps, see also. Also methods for robust PARAFAC and robust multilevel simultaneous component analysis are based on ROBPCA. The LTS subspace estimator generalizes LTS regression to subspace estimation and orthogonal regression.

An MCD-based alternative to the Hotelling test is provided in. A robust bootstrap for the MCD is proposed in and a fast cross-validation algorithm in. Computation of
the MCD for data with missing values is explored in\textsuperscript{65–67}. A robust Cronbach alpha is studied in\textsuperscript{68}. Classification (i.e. discriminant analysis) based on MCD is constructed in\textsuperscript{69,70}, whereas an alternative for high-dimensional data is developed in\textsuperscript{71}. Robust clustering is handled in\textsuperscript{72–74}.

The trimming procedure of the MCD has inspired the construction of maximum trimmed likelihood estimators\textsuperscript{75–78}, trimmed \textit{k}-means\textsuperscript{79–81}, least weighted squares regression\textsuperscript{82}, and minimum weighted covariance determinant estimation\textsuperscript{18}. The idea of the C-step in the FastMCD algorithm has also been extended to S-estimators\textsuperscript{83,84}.

**RECENT EXTENSIONS**

**Deterministic MCD**

As the FastMCD algorithm starts by drawing random subsets, it does not necessarily give the same result at multiple runs of the algorithm. (To address this, most implementations fix the seed of the random selection.) Moreover, FastMCD needs to draw many initial subsets in order to obtain at least one that is outlier-free. To circumvent both problems, a deterministic algorithm for robust location and scatter has been developed, denoted as DetMCD\textsuperscript{85}. It uses the same iteration steps as FastMCD but does not start from random subsets. Unlike FastMCD it is permutation invariant, i.e. the result does not depend on the order of the observations in the data set. Furthermore DetMCD runs even faster than FastMCD, and is less sensitive to point contamination.

DetMCD computes a small number of deterministic initial estimates, followed by concentration steps. Let \(X_j\) denote the columns of the data matrix \(X\). First each variable \(X_j\) is standardized by subtracting its median and dividing by the \(Q_n\) scale estimator of\textsuperscript{86}. This standardization makes the algorithm location and scale equivariant, i.e. equations (6) hold for any non-singular diagonal matrix \(A\). The standardized data set is denoted as the \(n \times p\) matrix \(Z\) with rows \(z_i'\) (\(i = 1, \ldots, n\)) and columns \(Z_j\) (\(j = 1, \ldots, p\)).

Next, six preliminary estimates \(S_k\) are constructed (\(k = 1, \ldots, 6\)) for the scatter or correlation of \(Z\):
1. $S_1 = \text{corr}(Y)$ with $Y_j = \tanh(Z_j)$ for $j = 1, \ldots, p$.

2. Let $R_j$ be the ranks of the column $Z_j$ and put $S_2 = \text{corr}(R)$. This is the Spearman correlation matrix of $Z$.

3. $S_3 = \text{corr}(T)$ with the normal scores $T_j = \Phi^{-1}((R_j - 1/3)/(n + 1/3))$.

4. The fourth scatter estimate is the spatial sign covariance matrix: define $k_i = z_i/\|z_i\|$ for all $i$ and let $S_4 = (1/n) \sum_{i=1}^n k_i k_i'$.

5. $S_5$ is the covariance matrix of the $\lfloor n/2 \rfloor$ standardized observations $z_i$ with smallest norm, which corresponds to the first step of the BACON algorithm.

6. The sixth scatter estimate is the raw orthogonalized Gnanadesikan-Kettenring (OGK) estimator.

As these $S_k$ may have very inaccurate eigenvalues, the following steps are applied to each of them:

1. Compute the matrix $E$ of eigenvectors of $S_k$ and put $V = ZE$.

2. Estimate the scatter of $Z$ by $S_k(Z) = E \Lambda E'$ where $\Lambda = \text{diag}(Q_n^2(V_1), \ldots, Q_n^2(V_p))$.

3. Estimate the center of $Z$ by $\hat{\mu}_k(Z) = S_k^{1/2}\text{comed}(ZS_k^{-1/2}))$ where comed is the coordinatewise median.

For the six estimates $(\hat{\mu}_k(Z), S_k(Z))$ the statistical distances $d_{ik} = d(z_i, \hat{\mu}_k(Z), S_k(Z))$ of all points are computed as in (2). For each initial estimate $k = 1, \ldots, 6$ we compute the mean and covariance matrix of the $h_0 = \lfloor n/2 \rfloor$ observations with smallest $d_{ik}$, and relative to those we compute statistical distances (denoted as $d_{ik}^*$) of all $n$ points. For each $k = 1, \ldots, 6$ the $h$ observations $x_i$ with smallest $d_{ik}^*$ are selected, and C-steps are applied to them until convergence. The solution with smallest determinant is called the raw DetMCD. Then a weighting step is applied as in (5), yielding the final DetMCD.

DetMCD has the advantage that estimates can be quickly computed for a whole range of $h$ values (and hence a whole range of breakdown values), as only the C-steps in the second part of the algorithm depend on $h$. Monitoring some diagnostics (such as the condition number
of the scatter estimate) can give additional insights in the underlying data structure, as in the example in

Note that even though DetMCD is not affine equivariant, it turns out that its deviation from affine equivariance is very small.

**Minimum regularized covariance determinant**

In high dimensions we need a modification of MCD, since the existing MCD algorithms take long and are less robust in that case. For large $p$ we can still make a rough estimate of the scatter as follows. First compute the first $q < p$ robust principal components of the data. For this we can use the MCD-based ROBPCA method\(^{53}\), which requires that the number of components $q$ be set rather low. The robust PCA yields a center $\hat{\mu}$ and $q$ loading vectors. Then form the $p \times q$ matrix $L$ with the loading vectors as columns. The principal component scores $t_i$ are then given by $t_i = L'(x_i - \mu)$. Now compute $\lambda_j$ for $j = 1, \ldots, q$ as a robust variance estimate of the $j$-th principal component, and gather all the $\lambda_j$ in a diagonal matrix $\Lambda$. Then we can robustly estimate the scatter matrix of the original data set $X$ by $\hat{\Sigma}(X) = L\Lambda L'$. Unfortunately, whenever $q < p$ the resulting matrix $\hat{\Sigma}(X)$ will have $p - q$ eigenvalues equal to zero, hence $\hat{\Sigma}(X)$ is singular.

If we require a nonsingular scatter matrix we need a different approach using regularization. The *minimum regularized covariance determinant* (MRCD) method\(^{90}\) was constructed for this purpose, and works when $n < p$ too. The MRCD minimizes

$$\det\{\rho T + (1 - \rho)\text{Cov}(X_H)\}$$

(11)

where $T$ is a positive definite ‘target’ matrix and $\text{Cov}(X_H)$ is the usual covariance matrix of an $h$-subset $X_H$ of $X$. Even when $\text{Cov}(X_H)$ is singular by itself, the combined matrix is always positive definite hence invertible. The target matrix $T$ depends on the application, and can for instance be the $p \times p$ identity matrix or an equicorrelation matrix in which the single bivariate correlation is estimated robustly from all the data. Perhaps surprisingly, it turns out that the C-step theorem can be extended to the MRCD. The MRCD algorithm is similar to the DetMCD described above, with deterministic starts followed by iterating these modified C-steps. The method simulates well even in 1000 dimensions.
Software for DetMCD and MRCD is available from http://wis.kuleuven.be/stat/robust.

CONCLUSIONS

In this paper we have reviewed the Minimum Covariance Determinant (MCD) estimator of multivariate location and scatter. We have illustrated its resistance to outliers on a real data example. Its main properties concerning robustness, efficiency and equivariance were described, as well as computational aspects. We have provided a detailed reference list with applications and generalizations of the MCD in applied and methodological research. Finally, two recent modifications of the MCD make it possible to save computing time and to deal with high-dimensional data.

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