Stochastic Optimization in Robust Statistic

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Abstract: The paper studies a stochastic optimization algorithm for computing of robust estimators of location proposed by Vandev (1992). A random approximation of the exact solution was proposed which is much cheaper in time and easy to program.

Two examples are presented. Besides standard estimators of location like trimmed mean also robust regressions (LMS and LTS) introduced by Rousseeuw and Leroy are considered. MATLAB programs are included.

Keywords: robust estimators of location, least median of squares, stochastic approximation algorithm, Monte – Carlo study.

1. Introduction

Many authors considered robust estimators of the covariance matrix and the location in the multidimensional case. When a high level of contamination is expected it is appropriate to use estimators with high breakdown point. Such estimators are the minimum volume ellipsoid (MVE) and the minimum covariance determinant (MCD), introduced by Rousseeuw and Leroy [10]. On the other hand in the robust regression literature very popular is the Least Median of the squares (LME) estimator which also has high breakdown point. Recently Neykov and Neytchev [7] proposed a robust alternative of the maximum likelihood estimators. Namely let $f(\theta, x)$ be the likelihood functions of the individual observation x. We denote by X the finite set of all observations. Here θ is the vector of unknown parameters. Let $A(\theta) = \{-log(f(\theta, x)), x \in X\}$ be the (increasingly) ordered set of the values of f at a fixed point θ . Denote by $M(k, \theta)$ the k-smallest and by $S(k, \theta)$ the sum of the k smallest numbers of the set $A(\theta)$. The minimizers of these two random functions are to be considered as estimators in statistical sense.

Vandev [12] has shown that MVE and MCD estimators may be extracted from this robustified version in the gaussean case. The same is true for LME in regression. It was also shown that in general all robustified maximum likelihood estimators have high break down point.

Computationally both (trimmed and least median) problems are not easy to solve in a conventional way because the functions involved have many local minima. Thus the minimization turns out to be a serious combinatorial problem. Up to now mainly the resampling technique is used for the purpose, see Rousseeuw & Leroy [10].

In this paper an algorithm is presented for approximate calculating of LME(k) and LTE(k). Hawkins [2] used a feasible set algorithm for exact calculation of the minima. Our proposition is based on the well known Robins-Monro [8] procedure for stochastic optimization, which was already successfully used by Martin and Masreliez [5] in the robust estimation. We will call the algorithm RM algorithm.

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2. Robust estimators in statistics

For modeling gross errors and outliers in the sample, the most popular is the Tukey supermodel [11] based on the Gaussean low:

$$\mathcal{F} = \left\{ F : F(x) = (1 - \varepsilon)\Phi(x) + \varepsilon\Phi(\frac{x - \theta}{k}), \quad 0 < \varepsilon < 1, 1 < k \right\}.$$
 (1)

Huber [3] considered more general model

$$\mathcal{F} = \{F : F(x) = (1 - \varepsilon)F_0(x) + \varepsilon H(x)\}, \qquad (2)$$

where F_0 is some given distribution (the ideal model) and H(x) is an arbitrary continuous distribution (contamination).

2.1. Break-down point

Since the general definition of a supermodel is based on the concept of a distance in the space of all distributions, the same concept is involved into the construction for a measure of the global robustness. Let d be such a distance. Then the breakdown point of the estimator $T_n = T(F_n)$ for the functional T(F) at \mathcal{F} is defined by

$$\varepsilon^*(T,\mathcal{F}) = \sup_{\varepsilon < 1} \{ \varepsilon : \sup_{F: d(F,F_0) < \varepsilon} |T(F) - T(F_0)| < \infty \}.$$

The breakdown point characterizes the maximal deviation (in the sense of a metric chosen) from the ideal model F_0 that provides the boundedness of the estimator bias.

Breakdown point as applied to the Huber supermodel

$$\varepsilon^*(T, \mathcal{F}) = \sup_{\varepsilon < 1} \{ \varepsilon : \sup_{F: F = (1 - \varepsilon)F_0 + \varepsilon H} |T(F) - T(F_0)| < \infty \}.$$
(3)

This notion defines the largest fraction of gross errors that still keeps the bias bounded. Here is the replacement variant of the finite sample breakdown point given by Hampel [1].

2.2. LMS and LTS

The multiple regression is probably most used statistical procedure in the statistics. Consider the model

$$y_i = x_i^T \beta + \varepsilon_i,$$

where y_i is an observed response, x_i is a $p \times 1$ -dimensional vector of explanatory variables and β is a $p \times 1$ vector of unknown parameters. Classically ε_i , $i = 1, \ldots, n$ are assumed to be i.i.d. $N(0, \sigma^2)$, for some $\sigma^2 > 0$.

The LMS (Least Median of Squares) and LTS (Least Trimmed Squares) estimators were proposed by Rousseeuw [9] as robust alternatives of the LSE

$$LMS(r_1, \dots, r_n) = \underset{\theta}{\operatorname{argmin}} \operatorname{med}\{r_i^2, i = 1, \dots, n\},$$
(4)

$$LTS(k)(r_1, \dots, r_n) = \operatorname{argmin}_{\theta} \sum_{i=1}^k r_{\nu(i,\theta)}^2.$$
 (5)

Here $\nu(i,\theta)$ is a permutation of the indices, such that $r_{\nu(i,\theta)}^2 \leq r_{\nu(i+1,\theta)}^2$. Thus the idea was to minimize the sum of squares using "smallest residuals" only.

Theorem 1 The breakdown point of the regression estimators (4 and 5) is equal to (n - k)/n if the index k is within the bounds $(n + p + 1)/2 \le k \le n - p - 1$, $n \ge 3(p + 1)$ and the data points $x_i \in \mathbb{R}^p$ for i = 1, ..., n are in general position.

This theorem was first proved by Rousseeuw [9] and then easily by Vandev [12] with different technique.

2.3. Robustified Maximum Likelihood

Neykov and Neytchev [7] proposed to replace in these estimators (LMS and LTS) the squared residuals with - log likelihood's of the individual observations and thus to obtain robustified likelihood.

Let the observations x_1, x_2, \ldots, x_n be generated by an arbitrary probability density function $\psi(x, \theta)$ with unknown vector parameter θ .

LME(k) =
$$\underset{\theta}{\operatorname{argmin}} \{-\log \psi(x_{\nu(k,\theta)}, \theta)\},$$
 (6)

$$LTE(k) = \operatorname{argmin}_{\theta} \sum_{i=1}^{k} \{-\log \psi(x_{\nu(i,\theta)}, \theta)\}.$$
(7)

Thus the idea was to maximize the likelihood over the best k observations (with "largest likelihood").

3. Stochastic Optimization

The famous Robins-Monro [8] procedure, later extended by Kiefer and Wolfowitz [4], when applied to the problem of minimizing the function $F(\theta)$ consists in the following. Let start with some $\theta = \theta_0$. Let now calculate the gradient $grad(F(\theta))$ at this point. It may be randomly disturbed by some random variable with zero expectation. At the step *i* the parameter will be changed according the following formula:

$$\theta_{i+1} = \theta_i - \gamma_i * \frac{grad(F(\theta_i))}{||grad(F(\theta_i))||}.$$
(8)

The sequence $\{\gamma_i, i = 1, 2, ...\}$ is chosen to satisfy the relations: $\sum_{i=1}^{\infty} \gamma_i^2 < \infty$, $\sum_{i=1}^{\infty} \gamma_i = \infty$. Here the only difference with the standard method as described in Wasan [6] is the normalizing of the gradient.

4. The Proposed Algorithm

Let F be set of functions of size n defined on p-dimensional Euclidean space E. Let $A(\theta) = \{f(\theta, x), x \in X\}$ be the (increasingly) ordered set of the values of f at a fixed point θ . Denote by $M(k, \theta)$ the k-smallest number in the set $A(\theta)$ and by $T(k, \theta)$ - the sum of k smallest numbers. Denote by :

$$LME(k) = \arg\min_{\theta} M(k, \theta) = \arg\min_{\theta} f_{(k)}(\theta), \tag{9}$$

$$LTE(k) = \arg\min_{\theta} T(k, \theta) = \arg\min_{\theta} \sum_{i=1}^{k} f_{(i)}(\theta),$$
(10)

where $f_{(1)}(\theta) \leq f_{(2)}(\theta) \leq \cdots \leq \cdots \leq f_{(n)}(\theta)$. As usual here the subindex denote the element of the corresponding permutation which depends on the value of θ .

```
Step 0. SET number of iterations maxi, set i=1, set \delta.
Step 1. Chose at random 10 indexes among the numbers from 1 to n.
Calculate these 10 functions. Sort their values.
Step 2. Chose the value j, such that (j/10=k/n)
and the function which produces that value (say f).
Step 3. Calculate the normalized gradient D(f) of f.
Step 4. SET B:=B - D(f)*\delta /i. Set i=i+1.
IF i < maxi THEN GOTO STEP 1.</pre>
```

5. MATLAB program

Here we present a MATLAB program able to handle the stochastic approximation algorithm in robust statistics.

```
function [theta] = soaml(x,theta0,FUN,pr,delta,iter)
[n,m]=size(x); theta=theta0;
for k=1:iter
   gama=delta/k;
                                         %
                                             new gama
    J=round(ones(kkk,1)/2+rand(kkk,1)*n); %
                                             10 random in (1:n) numbers
    eval(['[Y,X]=' FUN '(x(J,:),theta);']);% residuals, gradient
    [dum,list]=sort(Y);
                       % sort 25 values
%========LME or LTE ==============
    jj=list(pr,1);
                        % jj=list(1:pr,1);
                        % s=sum(X(jj,:))';
   s=X(jj,:)';
   w=sqrt(s'*s);
    theta=theta-s*(gama/w);
end
```

Pgm. 1: LME (and LTE) program

Note between commented lines the minor changes needed to transform this program for work in the LTE case.

The user defined function [Y,X]=FUN(x,theta) should produce in Y the values corresponding to observations in x and in X – corresponding gradients. Below we present some examples of such functions for various estimators:

```
function [Y,X]=gradmea(x,a) function [Y,X]=gradreg(x,a) function [Y,X]=gradnor(x,a)
[n,m]=size(x);
                             [n,m]=size(x);
                                                           [n,dum]=size(x);
                             xx=[ones(n,1),x(:,2:m)];
                                                           mu=a(ones(n,1),1);
aa=a';
                                                           si=exp(a(2,1));
X=x-aa(ones(n,1),:);
                             Y=x(:,1)-xx*a;
Y= diag(X*X');
                             X= -2*(Y(:,ones(1,m)).*xx); Y=(x-mu)/si;
                                                           X=[-Y/(2*si), (ones(n,1)-Y.*Y)];
X = -2 * X . * x;
                             Y=Y.*Y;
                                                           Y=Y.*Y/2+ones(n,1)*a(2);
                                                               Pgm. 4: N(\mu, \sigma) in \mathbb{R}^1
     Pgm. 2: Location
                                  Pgm. 3: Regression
```

6. Examples of application

Here we present several simulated examples. In all cases we use 1000 observations generated and 20% contamination, when not mentioned other.

6.1. Location

The 6-dimensional mean, 100 hundred repetitions, 6/10 LME:

Table 1: LME estimate of location

True	56.1761	0.9569	2.0455	3.0177	4.0263	4.9971
Est.	56.1019	0.9668	2.0476	2.9981	4.0225	4.9860
Err.	0.6054	0.0915	0.0936	0.0964	0.0870	0.0942

6.2. Simple regression

The first regression model was chosen to illustrate the robust properties of the used version of maximum likelihood. The response Y is generated by the following model:

$$y = 5 - 2 * x + e.$$

Here e is a standard normal random variable. The sample consists of 1000 observations. It was corrupted by destroying 30% of the observations. The algorithm was used with number of iterations equal to 150 and $\delta = 10$.

On Fig.1 a random solution is presented for the estimator 6/10. For a comparison the unique least squares solution is also plotted in red

Fig. 1: LME and LSQ regression

6.3. Multiple Regression

The model is:

$$y = 2 - 2 * x_1 + 5 * x_2 - 5 * x_3 + x_4 + e.$$

The aim was to test the performance of different estimators of the same kind (LME) when the percent of contamination changes.

In this case we each time generate totally new data set of 4000 uniform random numbers for x and 1000 normal for e. Each experiment was performed 100 times in order to estimate the variance.

The results are presented in the following table. The number of contaminated observations is shown in the first column. The form of used estimator is in the second column. Each cell in the table contains the average (with the sample standard error below) for 100 simulated with the same model data sets. In the next 4 columns are the results for the parameters of the model. The last column represents the obtained value of the functional.

	_	-	-			
Cont.	Est.	$a_0 = 2$	$a_1 = -2$	$a_2 = 5$	$a_3 = -5$	LME
100	9/10	1.9235	-1.9421	4.9492	-4.8944	3.3777
		.1149	.1569	.1249	.1265	2.4862
	8/10	1.9644	-1.9821	4.9029	-4.9164	1.2839
		.0990	.0987	.1401	.1136	.1512
	7/10	1.9390	-2.0412	4.9467	-4.8380	1.1343
		.1596	.1834	.1959	.1705	.2168
	6/10	1.9823	-1.9756	4.7355	-4.7443	.9773
		.2313	.2328	.3107	.2601	.2017
200	8/10	1.9664	-2.0136	4.7889	-4.7541	5.9930
		.1534	.1828	.1808	.2410	3.7446
	7/10	1.9103	-1.9337	4.9010	-4.8629	1.3670
		.2338	.2833	.2781	.2783	.5853
	6/10	1.9484	-1.9812	4.8867	-4.9113	1.0957
		.1811	.2229	.2409	.1701	.2407
300	7/10	1.7643	-1.7374	4.4975	-4.5186	7.7961
		.4012	.3970	.6973	.7480	4.5630
	6/10	1.8873	-1.8956	4.8093	-4.7899	1.5153
		.3159	.2421	.5369	.4467	.8834
400	6/10	1.5886	-1.6556	4.2696	-4.1614	9.5648
		.5058	.4803	.7968	.9176	4.7157

Table 2: Simulation results for multiple regression

What is easily seen in this table are the good results of 7/10 estimator for 10% contamination and 6/10 estimator – for 20%.

7. Mean and covariance

The estimating of variance needs special attention because it has to be positive. In the one-dimensional case the problem is solved using new parameter $\ln \sigma$ (see Pgm.4). In the multi-dimensional case however such approach is not easy. Before explaining difficulties let us present one unsuccessful example of two-dimensional estimate of mean and the covariance:

Table 3: Location and scale

Original mean	0.6242	2.5444
Estimated mean	0.9204	3.0970
Original Cova	37.0107	-20.4700
	-20.4700	51.5504
Estimated Cova	35.1363	-10.3004
	-10.3004	43.6298

Fig. 2: Location and scale

7.1. The problem of gradient

In the simultaneous estimation of the mean and covariance the main problem consists in calculation of the gradient of $Q = -\log L(x, m, \Sigma)$:

$$Q = \log \det(\Sigma)^{1/2} + (x - \mu)' \Sigma^{-1} (x - \mu).$$
(11)

Let denote $M = \Sigma^{-1}$. Then it is easy to show that

$$\frac{dQ}{dM} = -M^{-1} + (x - \mu)(x - \mu)'.$$
(12)

Let us replace $M = \exp(L)$ as in the univariate case and try to use the formal relation

$$\frac{dQ}{dL} = \frac{dQ}{dM} \bigotimes \frac{dM}{dL}.$$

Consider the standard expansion of $\exp(L)$

$$M = \exp L = I + L + L^2/2! + L^3/3! + \dots$$

The question now is how to represent $\frac{dM}{dL}$. We tried the following approximation of this $(m \times m)^2$ tensor:

$$\frac{dM}{dL} = (I + L/18) \bigotimes (I + L/18)$$

Thus we come to the result:

$$\frac{dQ}{dL} = (I + L/18)'((x - \mu)(x - \mu)' - M^{-1})(I + L/18)$$

Note that we are not sure how exact is this approximation.

7.2. The Simulation Results

These were obtained using 20% contamination of 1000 observations and MLE 6/10.

Table 4: Means

Original	18.0293	0.9973	-1.9745	3.0041	-6.0700	3.3209
Estimated	18.0253	1.0166	-2.0165	2.9931	-5.9780	3.3387
S.E.	0.1845	0.1340	0.1453	0.1536	0.1419	0.1077

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6.0332	0.4005	-0.6253	1.0875	-1.9673	1.0251
0.4005	1.0741	-0.0554	-0.0257	0.0529	0.0359
-0.6253	-0.0554	0.8846	-0.0332	0.0724	0.0294
1.0875	-0.0257	-0.0332	0.9997	-0.0110	0.0234
-1.9673	0.0529	0.0724	-0.0110	2.2645	0.0299
1.0251	0.0359	0.0294	0.0234	0.0299	0.4159

Table 6: Estimated covariance matrix

4.0793	0.2168	-0.3825	0.6135	-1.2302	0.6816
0.2168	0.6719	-0.0042	0.0088	-0.0132	0.0119
-0.3825	-0.0042	0.6703	-0.0101	0.0233	-0.0072
0.6135	0.0088	-0.0101	0.6832	-0.0281	0.0238
-1.2302	-0.0132	0.0233	-0.0281	1.4797	-0.0266
0.6816	0.0119	-0.0072	0.0238	-0.0266	0.3330

While the estimation of mean is excellent (see Table 4) the bias of the covariance is obvious on Table 6. Thus the proposed algorithm was not successful with estimation of covariance matrix. The reason is that the true unbiased gradient is not easy to obtain.

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