Greedy Approach for Low-Rank matrix recovery

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Abstract—We describe the Simple Greedy Matrix Completion Algorithm providing an efficient method for restoration of low-rank matrices from incomplete corrupted entries.

We provide numerical evidences that, even in the simplest implementation, the greedy approach may increase the recovery capability of existing algorithms significantly.

Keywords: Law-Rank Matrix Completion, Compressed Sensing, Image Inpainting, Motion Tracking, Face Recognition

1. Introduction

We consider a greedy strategy based algorithm for the recovery of the low-rank matrix from incomplete corrupted samples.

The problem of low-rank matrix completion is not new. However, it got a new impulse ([4], [2]) in connection with the development of the compressed sensing theory and algorithms and ideas to use the $\ell_1$ minimization as a surrogate for the sparsest solution ([3], [6], [12]).

This paper can be considered as a feasibility study for the methods inspired by ideas from both low-rank matrix completion and our compressed sensing oriented $\ell_1$-greedy algorithm ([7], [10], [11]).

The problem is set as follows. It is required to restore (complete) the matrix $A \in \mathbb{R}^{m \times n}$ of rank $r$, $r < \min\{m, n\}$, given by its $k$ entries, $k < mn$. The set of the given entries is $\Omega \subseteq \{1, \ldots, m\} \times \{1, \ldots, n\}$; $|\Omega|$ is the cardinality of $\Omega$ (which in our case is equal to $k$). We also introduce notation $d(\Omega)$ for the density of the set $\Omega$, $d(\Omega) := |\Omega|/(nm)$. The complementary set $\bar{\Omega}$ is a set of erasures, $1 - d(\Omega)$ represents the density of erasures. The theoretical bounds for recoverability of the matrix depends not only on the density of the samples but also on the matrix $A$ and on the 2D-geometry of $\Omega$. The matrix consisting of only one non-zero entry is the simplest example of a rank 1 matrix which can be restored only if the value at the non-zero entry is known. Anyway, it turned out (cf. [2]) that under quite mild conditions random matrices of size $n \times n$ and rank $r$ can be recovered from at most $O(rn^{1.2} \log n)$ entries as a matrix with the minimum of nuclear norm.

The popularity of that problem can be explained by an enormous number of applied problems which can be formulated in terms of matrix completion. Among many settings in different applied areas, we mention problems related to image processing. Image inpainting, including more particular image upsampling, face recognition technique, motion tracking and segmentation in video are most typical of those problems. While the problem of low-rank matrix completion is studied for a long time, the theory got a big push due to development of Compressed Sensing / Compressive Sampling (CS) technique. After some simplification, the CS data decoding goal can be reduced to solving underdetermined systems

\begin{equation}
Ax = y + e,
\end{equation}

where $x \in \mathbb{R}^n$ is a sparse vector of data "encoded" with the known to the decoder matrix $A \in \mathbb{R}^{m \times n}$, $m < n$; $y \in \mathbb{R}^m$ is a vector of measurements (of $x$) possibly corrupted by the vector $e \in \mathbb{R}^n$. Here and bellow we assume that the sparse solution $x$ exists and the vector of errors $e$ is also sparse. The sparsity of $a \in \mathbb{R}^N$ means that

$|a|_0 := |\{a_i \neq 0\}| < N.$

The value $|a|_0$ is called the Hamming weight of the vector $a$. Since the problem of finding sparse solutions has non-polynomial complexity ([9]), the mainstream CS researches suggested to use to replace the minimization of $|x|_0$ (or $|x|_0 + |e|_0$) with the minimization of $\ell_1$-norm. It turned out that such approach based on convex optimization gives the optimal sparse solution at least when $|x|_0$ is not very large (cf. [6], [3], [12] for the case $e = 0$). Thus, in some special cases the original non-convex problem can be reduced to convex programming. In what follows, like for the notion for the Hamming weight, we use notation $| \cdot |_p$, $0 < p < \infty$, for element-wise (quasi-)norms of vectors and matrices. Say, for the matrix $A$, $|A|_p := \left(\sum_{i,j} |A_{ij}|^p\right)^{1/p}$. In particular, $| \cdot |_2$ is the Frobenius norm. The inner product of 2 matrices $A$ and $B$ is defined as $\langle A, B \rangle := \text{trace}(A^TB)$. Thus, $\langle A, A \rangle = |A|^2_2$. The notation $\| \cdot \|_p$ is reserved for the operator norms of matrices.

CS results inspired the authors of [2] and [4] on replacing the minimum rank condition leading to non-polynomial complexity with the minimization of the nuclear norm $\|A\|_* := \sum \sigma_i$ of the matrix $A$, where $\sigma_i$ are singular values.
of $A$. To be more precise, the problem
$$\|A\|_* \rightarrow \text{min} \text{ subject to } A_{ij} = M_{ij}, \ (i,j) \in \Omega,$$
where $M_{i,j}$ are the known entries (measurements) of the matrix $A$, is considered as relaxation of the rank minimization problem above.

Many different settings giving a solution of the original problems have been studied for the last years. In most cases, the intention of those studies was to find the faster algorithms with the higher capability of the recovery. Typically, modifications of the problem leading to unconstrained optimization were introduced.

2. Basic Algorithm

For our experiments we need the algorithm providing convex minimization recovering low-rank matrices from incomplete corrupted samples. It is used as a basic constructive block in our algorithm. The problem of restoring a matrix from corrupted entries is less studied than the simpler matrix completion problem when the available entries are not corrupted at all or corrupted by noise with relatively low level magnitude. Anyway, there are a few computationally efficient algorithms solving that problem (e.g., [5], [8], [14]).

For our purposes, we selected the algorithm from [8] based on the method of Augmented Lagrange Multipliers (ALM) (e.g., [11]). Having corrupted samples, instead of finding the matrix with the sparsest set of singular values coinciding with the measurements on as large as possible set, the algorithm finds the minimum of the functional
$$L(A,E,Y,\mu) := \|A\|_* + \lambda |E|_1 + \langle Y,R \rangle + \frac{\mu}{2} \|R\|_2^2,$$  \hspace{1em} (2)
where $R = M - A - E$ is the residual of approximation of the measurements $M$ of the estimate of the unknown matrix $A$ and the estimate of the unknown matrix of errors $E$. The entries of the input matrix $M$ on the $\Omega$ are unknown. It is assumed that $R$ vanishes on $\bar{\Omega}$ and does not contribute into the third and the forth terms as well as $E$ does not contribute into the second term.

If it is known that the observed entries in $M$ are not corrupted, the second term can be omitted. However, we assume that we never know whether the entries are corrupted. So, in what follows, we minimize the 4-term functional given in (2).

We will need the following notation
$$S_\epsilon[x] := \begin{cases} x - \epsilon, & x > \epsilon, \\ x + \epsilon, & x < -\epsilon, \\ 0, & \text{otherwise}; \end{cases}$$
where $x$ can be either a number or a vector or a matrix. For vectors and matrices the operator is applied entrywise. The operator $S_\epsilon$ is called the shrinkage operator.

The minimization algorithm, as it is described in [8] and implemented in Matlab code, is as follows

Algorithm ALM.

Input. Observation matrix $M \in \mathbb{R}^{m \times n}$, defined on $\Omega$, and $\lambda > 0$.

Initialization. $Y^0 = \frac{1}{\max\{\|M\|_2,\|M\|_\infty/\lambda\}} M$, $E^0 = 0$, $\mu_0 > 0$, $\rho > 1$, $k = 0$;
1. while not converged do
2. $(U,S,V) := \text{svd}(M - E^k + \mu^{-1}Y^k)$;
3. $A^{k+1} := US_{\mu^{-1}}[S]V^T$;
4. $E^{k+1} := S_{\mu^{-1}}[M - A^{k+1} + \mu^{-1}Y^k]$;
5. $Y^{k+1} := Y^k + \mu_k(M - A^{k+1} - E^{k+1})$;
6. $\mu_{k+1} := \rho \mu_k$, $k := k + 1$;
7. end while.
Output. $A^{k+1}$, $E^{k+1}$.

3. Our algorithm

Our modification of the algorithm above is inspired by significant success reached by applying greedy ideas to solving underdetermined systems ([17], [10], [11]). The general greedy strategy in optimization algorithms consists in sequential finding a simple suboptimal solutions giving some (incomplete) information about the optimal solution. A greedy algorithm picks up the most obvious features or elements of those solutions and gives them a privilege to be pivot for the next iteration of the suboptimal algorithm. Each iteration brings new pivot elements.

In the matrix recovery algorithm, the erasures from the set $\bar{\Omega}$ forms such group from the beginning. Whereas, the elements of $\Omega$ are just suspicious to be erroneous. If we have sufficient evidence that some element in $\Omega$ contains a random error independent of the content of other entries from $\Omega$, that the decision to move this element to $\bar{\Omega}$ is quite justifiable. While the independence condition is not always accurate even in our experiments with artificially generated data, we use this strategy for estimation of the capability of the greedy ideas for matrix recovery.

Our greedy algorithm consists in iterating with updated (dilated) sets $\Omega_k$. We will call it the Simple Greedy Matrix Completion Algorithm (SGMCA). Generally speaking, any matrix recovery algorithm, including SGMCA itself, which is able to fight the mixture of erasures and errors can be used as a basic block of SGMCA.

Formally, all our experiments can be described in the following way.

Algorithm SGMCA.

Input. $M$, $\Omega$. Initialization. $\lambda > 1$, $\Omega_0 := \Omega$, $A^0 := M$, $E^0 := 0$, $0 < q_0, q_1 < 1$, $k = 0$.
1. Set $k := k + 1$;
2. $A^k := \text{ALM}(M, \Omega_k)$;
3. if $k = 1$ then $T_1 = q_0 \max_{i,j} \{|A^1_{ij} - M_{ij}|; \text{ else } T_k := q_k T_{k-1}$;
4. $\Omega_{k+1} := \Omega_k \backslash \{(i,j) \mid |A^k_{ij} - M_{ij}| > T_k\}$;
5. if not converged go to 2.
4. Numerical Experiments

Since our intention was to conduct an algorithm feasibility study, the goal of this section is to give comparison with the output of recently published algorithms and with pure ALM (one iteration of the algorithm above with no update). The parameters in the basic algorithm are selected as 

$$\mu_0 = 0.3/\|M\|_2, \quad \rho = 1.1 + 0.5\Omega/(mn), \quad q_0 = 0.3, \quad q_1 = 0.65.$$  

The parameter $\lambda$ is defined by the combination of $d(\Omega)$ and the density of errors in $\Omega$ samples. The general trend can be characterized as follows: the higher error rate, the less value of $\lambda$ has to be used. At the same time, too small value of $\lambda$ may bring the increased level of the false alarm in identification of error locations. We will use the least values of $\lambda$ in the cases when we have an additional mechanism for protection from the false alarm. One of such cases considered below is the case when the rank of the matrix $A$ is known in advance. In what follows, we do not use fine tuning of $\lambda$. The same $\lambda$ is used for big groups of our experiments. At the same time, tuning $\lambda$ may bring significant increase of the algorithm efficiency. In this paper, we use values of $\lambda$ in the range $0.02 \div 100$ (from the case of the known rank to the blind matrix completion).

For our experiments, we used Matlab implementation of ALM algorithm available at http://perception.csl.illinois.edu/matrix-rank/home.html We used the code for Matrix Completion via the inexact ALM Method with our adaptation to the input with errors.

In all our experiments with synthetic data, $A$ are square $m \times n$, $m = n$, matrices of rank $r$ obtained as $A = UV^T$, where $U, V \in \mathbb{R}^{n \times r}$. The matrices $U$ and $V$ consist of independent gaussian random values with zero expectation and the variance 1. The coordinates of erasures were selected randomly. The models of errors below were different for different experiments.

In the first experiment (Fig.1), we demonstrate advantage of the iterative SGMCA over ALM (one iteration of the same algorithm) for the matrix with fixed sizes $m \times n$, $m = n$, and the rank of matrices $r = 15$. We use the additive model of errors adding the random values from the standard normal distribution at the random available for reconstruction entries of the matrix $A$. The number of the corrupted entries is fixed in each experiment and does not exceed the value $|\Omega|$.

The solid curves on Fig. 1 correspond to SGMCA (up to 10 iterations) for $n = 128, 512, 1024$ (from the bottom to the top). The corresponding graphs for ALM algorithm are plotted with dashed curves.

The horizontal coordinate indicates the fraction of the matrix available for restoration (i.e., $d(\Omega)$), while the vertical coordinate is the fraction of randomly corrupted entries in $\Omega$. The magnitude of the corruption is randomly set from the standard normal distribution. The curves define "phase transition" bounds. In our experiments, we run 10 trials. At the points of curves as well as under each curve all 10 attempts were accomplished with success, i.e., for the obtained estimate $\hat{A}$, $|A - \hat{A}|_2/|A|_2 < 10^{-3}$, whereas for the points above the curves, at least one attempt failed. This means that the regions under the curves are regions of "success".

The second experiment (Fig.2 and Fig.3) is devoted to comparison with the results from [5]. Unfortunately, we do not have full information about the error model. So we use the same additive model of errors as above. While all other parameters are taken from [5]. The matrix of rank 2 is constructed as above. Its size changes from 100 to 3000.

The experiment consists of 2 parts. The first plot (Fig.2) contains the curves for the fixed erasure rate 0.1, i.e., $d(\Omega) = 0.9$. However, the probability of errors in those entries varies. There are 3 graphs on Fig. 2. The solid line corresponds to 10 iterations of SGMCA, the dashed line corresponds to ALM, and the dotted curve corresponds to the result from [5]. The values defined by curves give the maximum error probability admitting successful correction by the corresponding algorithm. If we were aware of the error model from [5], the dashed and dotted curves have to coincide up to statistical discrepancy.

On the second plot (Fig.3) the error rate is fixed and equal to 0.1. The graphs show dependence of maximum possible rate of erasures from the size of matrix.
The efficiency of the algorithms is defined by the distance of curve values on Fig. 2 and Fig. 3 to 1. On Fig. 2, this distance defines the fraction of uncorrupted entries among all available entries. Whereas that distance on Fig. 3 defines the fraction of entries available for the procedure of matrix completion among all uncorrupted entries. It is easy to see that, when the error rate is fixed, SGMCA curve is twice closer to 1 than ALM. Hence, SGMCA restores low rank matrices from only half of entries necessary for the ALM minimization. For the fixes erasure rate, the number of uncorrupted entries for SGMCA successful restoration can be only one third of that necessary for recovery with ALM.

The more precise value of the SGMCA graph at $m = 3000$ on Fig 3 is 0.986, i.e., having error probability 0.1, the matrix can be restored from 1.4% of random entries.

In our last experiment (Fig. 4–6), we compare the output of SGMCA with the results of RTRMC algorithm from [14] providing very impressive recovery. However, for such successful recovery it requires a priori knowledge of the rank of the matrix $A$. The ALM-based algorithm used as a basic algorithm in SGMCA does not require any knowledge about the rank while the rank knowledge is useful for it and can be incorporated. To provide equal opportunities for SGMCA and RTRMC we applied the internal fixation of the rank within the ALM procedure.

The results for ranks $r = 5, 15, 25$ are given on Fig. 4–6 correspondingly. The size of matrices is $512 \times 512$. The horizontal coordinate corresponds to $d(\Omega)$, whereas the vertical coordinate is the probability of errors in the coefficients available for reconstruction. In most of cases, SGMCA outperforms RTRMC by 15–25% in the maximum admissible probability of errors. The reason why SGMCA loses on interval $[0, 0.175]$ on Fig.4 is the parameter $\lambda = 0.02$ which was fixed for all 3 experiments. Setting $\lambda = 0.2$ on that interval, we would get the overwhelming advantage of SGMCA. We emphasize that when the rank is known in advance, the optimal parameter $\lambda$ can be computed for each $d(\Omega)$. This would not contradict the equal opportunity of the two algorithms. Thus, we can make conclusion that actually SGMCA outperforms RTRMC for all considered configurations of input data.

Optimal selection of $\lambda$ is a reserve not used in our experiments.

The model of data in [14] is identical to the model described above. At the same time, the error model is different. The values of the corrupted entries are randomly uniformly distributed between minimum and maximum of uncorrupted values. In this case, the average magnitude of errors is significantly higher than in 2 models considered above. We also use that error model in our experiments presented on Fig. 4–6. So we satisfy all experiment conditions described in [14].
The dashed line corresponding to RTRMC is shorter than our solid line since we used the data directly from [14].

5. Future Studies

This study shows that even the simplest implementation of the greedy idea in the form of SGMCA outperforms the recent state-of-the-art algorithms significantly in the recovering ability for very incomplete measurements with high level of corruption. The results above show the feasibility of the idea, its perspectives and a high level of expectation.

Because of its iterative nature, the algorithm has to repeat the basic step a few times. We restricted the number of iterations by 10. However, in most of cases, 5 iterations provide necessary precision. Thus, SGMCA has obvious increase of algorithm computational cost in 5-10 times against the basic algorithm. Among possible directions for improvement, acceleration ways are needed to be considered.

One of possible ways is to do not wait for the completion of each iteration, updating $\Omega$ within internal iterations of the basic algorithm. In the simplest, but maybe in less efficient form, it can be done even with no intrusion into the basic algorithm. For instance, when the greedy step looks for coordinates of large errors, we do not need high precision output of the basic algorithm. So an update of the precision on each iteration from low to high may accelerate the algorithm for the data close to the limits of the potential recovering ability (phase transition points). However, it should be mentioned that this modification may slightly slow down the recovery of the data located far from the phase transition points. Other way for acceleration skipped by us in this paper is to use the estimate of $A$ obtained on the previous iteration as a basic algorithm start point for the next iteration.

Now we discuss the ways for increasing the capability of SGMCA in the matrix recovery.

First of all, in our experiments, we practically did not use fine tuning of the weight $\lambda$. We used fixed $\lambda$ for big range of the parameters of input data. Whereas, just by replacing the value 0.02 with the value 0.2, the results presented on Fig. 4–6 can be significantly improved for the high level of erasures (on the left side of the graphs). Indeed, when the rank of $A$ and the model of errors is known, the optimal values of $\lambda$ admitting the maximum density of error can be found in advance for any number of erasures and used in the recovery procedure.

When the rank is unknown or nature of possible corruption is unknown in advance, adaptive finding $\lambda$ becomes a challenging problem. This problem has many common features with the problem of finding sparse solutions of underdetermined linear systems with corrupted input. In the mentioned problem, the weight $\lambda$ is defined by the interaction between the sparsity of the possible solution and the error vector. In the problem of matrix completion, the low rank plays a role of the solution sparsity in CS. So the methods (or at least principles) developed in CS can be applied to the matrix completion problem. In [11], we showed that the sparsity of the solution can be reliably estimated from the dynamic of change of the values $|x|_{0.5}/|x|_2$ and $|e|_{0.5}/|e|_2$ for the intermediate approximations $x$ and $e$ of the solution and errors. The same characteristic of the matrix $A$ intermediate estimates probably can be used for the matrix completion procedure.

We also have to say that, generally speaking, for finding $\lambda$ we do not need both the rank and the sparsity of errors estimates. Indeed, if we have the sparsity of errors, we can evaluate the potential maximum rank $r$ admitting recovery with the given algorithm provided that $\lambda$ is optimal. Then that optimal $\lambda$ will also provide recovery of matrices with the rank less than $r$. Thus, the adaptive $\lambda$ is one of quite realistic sources for increasing algorithm capability.

6. Conclusion

The paper presents a feasibility study for the Simple Greedy Matrix Completion algorithm. The considered al-
gorithm is based on the ALM algorithm from [8]. While a problem of resistance of the matrix completion algorithms to discrete errors in data is not well studied yet., some recent development in this direction took place.

We gave numerical evidence that SGMCA outperforms recently developed algorithms of matrix completion from [5], [8], [14] significantly. We also discussed the ways for further increase of SGMCA efficiency.

References