Tensor Completion for Estimating Missing Values in Visual Data

Ji Liu, Przemyslaw Musialski, Peter Wonka, and Jieping Ye, Senior Member, IEEE

Abstract—In this paper, we propose an algorithm to estimate missing values in tensors of visual data. The values can be missing due to problems in the acquisition process or because the user manually identified unwanted outliers. Our algorithm works even with a small amount of samples and it can propagate structure to fill larger missing regions. Our methodology is built on recent studies about matrix completion using the matrix trace norm. The contribution of our paper is to extend the matrix case to the tensor case by proposing the first definition of the trace norm for tensors and then by building a working algorithm. First, we propose a definition for the tensor trace norm that generalizes the established definition of the matrix trace norm. Second, similarly to matrix completion, the tensor completion is formulated as a convex optimization problem. Unfortunately, the straightforward problem extension is significantly harder to solve than the matrix case because of the dependency among multiple constraints. To tackle this problem, we developed three algorithms: simple low rank tensor completion (SiLRTC), fast low rank tensor completion (FaLRTC), and high accuracy low rank tensor completion (HaLRTC). The SiLRTC algorithm is simple to implement and employs a relaxation technique to separate the dependant relationships and uses the block coordinate descent (BCD) method to achieve a globally optimal solution; the FaLRTC algorithm utilizes a smoothing scheme to transform the original nonsmooth problem into a smooth one and can be used to solve a general tensor trace norm minimization problem; the HaLRTC algorithm applies the alternating direction method of multipliers (ADMMs) to our problem. Our experiments show potential applications of our algorithms and the quantitative evaluation indicates that our methods are more accurate and robust than heuristic approaches. The efficiency comparison indicates that FaLTRC and HaLRTC are more efficient than SiLRTC and between FaLRTC and HaLRTC the former is more efficient to obtain a low accuracy solution and the latter is preferred if a high-accuracy solution is desired.

| Index | Terms— | Tensor | completion, | trace norm, | sparse | learning |
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1 Introduction

In computer vision and graphics, many problems can be formulated as a missing value estimation problem, e.g., image in-painting [4], [22], video decoding, video in-painting [23], scan completion, and appearance acquisition completion. The core problem of the missing value estimation lies on how to build up the relationship between the known elements and the unknown ones. Some energy methods broadly used in image in-painting, e.g., PDEs [4] and belief propagation [22], mainly focus on the local relationship. The basic (implicit) assumption is that the missing entries mainly depend on their neighbors. The further apart two points are, the smaller their dependance is. However, sometimes the value of the missing entry depends on the entries which are far away. Thus, it is necessary to develop a tool to directly capture the global information in the data.

- J. Liu is with the University of Wisconsin–Madison, Madison, WI 53706.
 E-mail: ji-liu@cs.wisc.edu.
- P. Musialski is with the VRVis Research Center, Vienna, Austria. E-mail: musialski@vrvis.at.
- P. Wonka is with Arizona State University, Tempe, AZ 85287-8809, and King Abdullah University of Science and Technology (KAUST), Saudi Arabia. E-mail: pwonka@gmail.com.
- J. Ye is with Arizona State University, 699 South Mill Avenue, Tempe, AZ 85287-8809. E-mail: jieping.ye@asu.edu.

Manuscript received 22 Nov. 2010; revised 21 Nov. 2011; accepted 5 Jan. 2012; published online 19 Jan. 2012.

Recommended for acceptance by Y. Ma.

For information on obtaining reprints of this article, please send e-mail to: tpami@computer.org, and reference IEEECS Log Number TPAMI-2010-11-0888.

Digital Object Identifier no. 10.1109/TPAMI.2012.39.

In the 2D case, i.e., the matrix case, the "rank" is a powerful tool to capture some type of global information. In Fig. 1, we show a texture with 80 percent of its elements removed randomly on the left and its reconstruction using a low rank constraint on the right. This example illustrates the power of low rank approximation for missing data estimation. However, "rank(·)" is unfortunately not a convex function. Some heuristic algorithms were proposed to estimate the missing values iteratively [13], [24]. However, they are not guaranteed to find a globally optimal solution due to the nonconvexity of the rank constraint.

Recently, the trace norm of matrices was used to approximate the rank of matrices [30], [7], [37], which leads to a convex optimization problem. The trace norm has been shown to be the tightest convex approximation for the rank of matrices [37], and efficient algorithms for the matrix completion problem using the trace norm constraint were proposed in [30] and [7]. Recently, Candès and Recht [9], Recht et al. [37], and Candès and Tao [10] showed that under certain conditions, the minimum rank solution can be recovered by solving a convex optimization problem, namely, the minimization of the trace norm over the given affine space. Their work theoretically justified the validity of the trace norm to approximate the rank.

Although the low rank approximation problem has been well studied for matrices, there is not much work on tensors, which are a higher dimensional extension of matrices. One major challenge lies in an appropriate definition of the trace norm for tensors. To the best of our knowledge, this has been not addressed in the literature. In this paper, we make





Fig. 1. The left figure contains 80 percent missing entries, shown as white pixels, and the right figure shows its reconstruction using the low rank approximation.

two main contributions: 1) We lay the theoretical foundation of low rank tensor completion and propose the first definition of the trace norm for tensors. 2) We are the first to propose a solution for the low rank completion of tensors.

The challenge of the second part is to build a high-quality algorithm. Similarly to matrix completion, the tensor completion can be formulated as a convex optimization problem. Unfortunately, the straightforward problem extension is significantly harder to solve than the matrix case because of the dependency among multiple constraints. To tackle this problem, we developed three algorithms: simple low rank tensor completion (SiLRTC), fast low rank tensor completion (FaLRTC), and high accuracy low rank tensor completion (HaLRTC). The SiLRTC algorithm, a pretty simple and intuitive method, employs a relaxation technique to separate the dependant relationships and uses the block coordinate descent (BCD) method to achieve a globally optimal solution. It actually simplifies the LRTC algorithm proposed in our conference paper [29]. The FaLRTC algorithm utilizes a smoothing scheme to transform the original nonsmooth problem into a smooth problem. We also present a theoretical analysis of the convergence rate for the FaLRTC algorithm. The third method applies the alternating direction method of multipliers (ADMM) algorithm [5] to our problem. In addition, we present several heuristic models which involve nonconvex optimization problems. Our experiments show that our method is more accurate and robust than these heuristic approaches. We also give some potential applications in image in-painting, video compression, and BRDF data estimation, using our tensor completion technique. The efficiency comparison indicates that FaLRTC and HaLRTC are more efficient than SiLRTC and between FaLRTC and HaLRTC the former is more efficient to obtain a low accuracy solution and the latter is preferred if a high-accuracy solution is desired.

1.1 Notation

We use upper case letters for matrices, e.g., X, and lower case letters for the entries, e.g., x_{ij} . $\Sigma(X)$ is a vector, consisting of the singular values of X in descending order and $\sigma_i(X)$ denotes the ith largest singular value. The Frobenius norm of the matrix X is defined as $\|X\|_F := (\sum_{i,j} |x_{ij}|^2)^{\frac{1}{2}}$. The spectral norm is denoted as $\|X\| := \sigma_1(X)$ and the trace norm as $\|X\|_{tr} := \sum_i \sigma_i(X)$. Let $X = U \Sigma V^{\top}$ be the singular value decomposition for X. The "shrinkage" operator $\mathbf{D}_{\tau}(X)$ is defined as [7]

$$\mathbf{D}_{\tau}(X) = U \Sigma_{\tau} V^{\top}, \tag{1}$$

where $\Sigma_{\tau} = diag(\max(\sigma_i - \tau, 0))$. The "truncate" operation $\mathbf{T}_{\tau}(X)$ is defined as

$$\mathbf{T}_{\tau}(X) = U \Sigma_{\bar{\tau}} V^{\top}, \tag{2}$$

where $\Sigma_{\bar{\tau}} = diag(\min(\sigma_i, \tau))$. It is easy to verify that $X = \mathbf{T}_{\tau}(X) + \mathbf{D}_{\tau}(X)$. Let Ω be an index set, then X_{Ω} denotes the matrix copying the entries from X in the set Ω and letting the remaining entries be "0." A similar definition can be extended to the tensor case. The inner product of the matrix space is defined by $\langle X, Y \rangle = \sum_{i,j} X_{ij} Y_{ij}$.

We follow [11] to define the terminology of tensors used in the paper. An n-mode tensor (or n-order tensor) is defined as $\mathcal{X} \in \mathbb{R}^{I_1 \times I_2 \times \cdots \times I_n}$. Its elements are denoted as x_{i_1,\dots,i_n} , where $1 \leq i_k \leq I_k$, $1 \leq k \leq n$. For example, a vector is a 1-mode tensor and a matrix is a 2-mode tensor. It is sometimes convenient to unfold a tensor into a matrix. The "unfold" operation along the kth mode on a tensor \mathcal{X} is defined as $\mathrm{unfold}_k(\mathcal{X}) := X_{(k)} \in \mathbb{R}^{I_k \times (I_1 \dots I_{k-1} I_{k+1} \dots I_n)}$. The opposite operation "fold" is defined as $\mathrm{fold}_k(X_{(k)}) := \mathcal{X}$. Denote $\|\mathcal{X}\|_F := (\sum_{i_1,i_2,\dots i_n} |a_{i_1,i_2,\dots i_n}|^2)^{\frac{1}{2}}$ as the Frobenius norm of a tensor. It is clear that $\|\mathcal{X}\|_F = \|X_{(k)}\|_F$ for any $1 \leq k \leq n$. Please refer to [11] for a more extensive overview of tensors. In addition, we use a nonnegative superscript number to denote the iteration index, e.g., \mathcal{X}^k denotes the value of \mathcal{X} at the kth iteration; the superscript "-2" in K^{-2} denotes the power.

1.2 Organization

We review related work in Section 2, introduce a convex model and three heuristic models for the low rank tensor completion problem in Section 3, present the SiLRTC, FaLRTC, and HaLRTC algorithms to solve the convex model in Sections 4, 5, and 6, respectively, report empirical results in Section 7, and conclude this paper in Section 8. To increase the readability of this paper for the casual reader most technical details can be found in the Appendix, which can be found in the Computer Society Digital Library at http://doi.ieeecomputersociety.org/10.1109/TPAMI.2012.39.

2 RELATED WORK

The low rank or approximately low rank problem broadly occurs in science and engineering, e.g., computer vision [42], machine learning [1], [2], signal processing [26], and bioinformatics [44]. Fazel et al. [13], [12] introduced a low rank minimization problem in control system analysis and design. They heuristically used the trace norm to approximate the rank of the matrix. They showed that the trace norm minimization problem can be reformulated as a semidefinite programming (SDP) problem via its dual norm (spectral norm). Srebro et al. [39] employed second-order cone programming (SCOP) to formulate a trace norm related problem in matrix factorization. However, many existing optimization methods such as SDPT3 [41] and SeDuMi [40] cannot solve an SDP or SOCP problem when the size of the matrix is much larger than 100×100 [30], [37]. This limitation prevented the usage of the matrix completion technique in computer vision and image processing. Recently, to solve the rank minimization problem for large scale matrices, Ma et al. [30] applied the fixed point and Bregman iterative method and Cai et al. [7] proposed a singular value

thresholding algorithm. In both algorithms, one key building block is the existence of a closed form solution for the following optimization problem:

$$\min_{X \in \mathbb{R}^{p \times q}} : \frac{1}{2} \|X - M\|_F^2 + \tau \|X\|_{tr}, \tag{3}$$

where $M \in \mathbb{R}^{p \times q}$ and τ is a constant. Candès and Recht [9], Recht et al. [37], and Candès and Tao [10] theoretically justified the validity of the trace norm to approximate the rank of matrices. Recht [36] recently improved their result and also largely simplified the proof by using the golfing scheme from quantum information theory [15]. An alternative singular value-based method for matrix completion was recently proposed and justified by Keshavan et al. [21].

This paper builds on our own previous work [29] where we extended the matrix trace norm to the tensor case and proposed to recover the missing entries in a low rank tensor by solving a tensor trace norm minimization problem. We used a relaxation trick on the objective function such that the block coordinate descent algorithm can be employed to solve this problem [29]. Since this approach is not efficient enough, some recent papers tried to use the alternating direction method of multipliers to efficiently solve the tensor trace norm minimization problem. The ADMM algorithm was developed in the 1970s, but was successful in solving large scale problems and optimization problems with multiple nonsmooth terms in the objective function [28] recently. Signoretto et al. [38] and Gandy et al. [14] applied the ADMM algorithm to solve the tensor completion problem with Gaussian observation noise, i.e.,

$$\min_{\mathcal{X}} : \frac{\lambda}{2} \| \mathcal{X}_{\Omega} - \mathbf{T}_{\Omega} \|_F^2 + \| \mathcal{X} \|_*, \tag{4}$$

where $\|\mathcal{X}\|_*$ is the tensor trace norm defined in (8). The tensor completion problem without observation noise can be solved by optimizing (4) iteratively with an increasing value of λ [38], [14]. Tomioka et al. [43] proposed several slightly different models for the problem (4) by introducing dummy variables and also applied ADMM to solve them. Out of these three algorithms for tensor completion based on ADMM, we choose to compare to the algorithm by Gandy et al. because the problem statement is identical to ours. Our results will show that our adaption of ADMM and our proposed FaLRTC algorithm are more efficient.

Besides tensor completion, the tensor trace norm proposed in [26] can be applied in various other computer vision problems such as visual saliency detection [47], medical imaging [16], corrupted data correction [26], [27], data compression [25].

3 THE FORMULATION OF TENSOR COMPLETION

This section presents a convex model and three heuristic models for tensor completion.

3.1 Convex Formulation for Tensor Completion

Before introducing the low rank tensor completion problem, let us start from the well-known optimization problem [24] for low rank matrix completion:

$$\min_{X} : \operatorname{rank}(X)
s.t. : X_{\Omega} = M_{\Omega},$$
(5)

where $X, M \in \mathbb{R}^{p \times q}$ and the elements of M in the set Ω are given while the remaining elements are missing. The missing elements of X are determined such that the rank of the matrix X is as small as possible. The optimization problem in (5) is a nonconvex optimization problem since the function $\mathrm{rank}(X)$ is nonconvex. One common approach is to use the trace norm $\|.\|_*$ to approximate the rank of matrices. The advantage of the trace norm is that $\|.\|_*$ is the tightest convex envelop for the rank of matrices. This leads to the following convex optimization problem for matrix completion [3], [7], [30]:

$$\min_{X} : ||X||_{*}$$

$$s.t. : X_{\Omega} = M_{\Omega}.$$
(6)

The tensor is the generalization of the matrix concept. We generalize the completion algorithm for the matrix (i.e., 2-mode or 2-order tensor) case to higher order tensors by solving the following optimization problem:

$$\min_{\mathcal{X}} : \|\mathcal{X}\|_{*}
s.t. : \mathcal{X}_{\Omega} = \mathcal{T}_{\Omega},$$
(7)

where \mathcal{X} , \mathcal{T} are <u>n-mode tensors</u> with identical size in each mode. The first issue is the definition of the trace norm for the general tensor case. We propose the following definition for the tensor trace norm:

$$\|\mathcal{X}\|_* := \sum_{i=1}^n \alpha_i \|\underline{\mathcal{X}_{(i)}}\|_*, \tag{8}$$

where α_i s are constants satisfying $\alpha_i \geq 0$ and $\sum_{i=1}^n \alpha_i = 1$. In essence, the trace norm of a tensor is a convex combination of the trace norms of all matrices unfolded along each mode. Note that when the mode number n is equal to 2 (i.e., the matrix case), the definition of the trace norm of a tensor is consistent with the matrix case because the trace norm of a matrix is equal to the trace norm of its transpose. Under this definition, the optimization in (7) can be written as

$$\min_{\mathcal{X}} : \sum_{i=1}^{n} \alpha_i \|\mathcal{X}_{(i)}\|_*$$

$$s.t. : \mathcal{X}_{\Omega} = \mathcal{T}_{\Omega}.$$
(9)

Here one might ask why we do not define the tensor trace norm as the convex envelop of the tensor rank like in the matrix case. Unlike matrices, computing the rank of a general tensor (mode number > 2) is an NP hard problem [18]. Therefore, there is no explicit expression for the convex envelop of the tensor rank to the best of our knowledge.

3.2 Three Heuristic Algorithm

We introduce several heuristic models, which, unlike the one in the last section, involve nonconvex optimization problems. A goal of introducing the heuristic algorithms is to establish some basic methods that can be used for comparison.

Tucker. One natural approach is to use the Tucker model [46] for tensor factorization to the tensor completion problem as follows:

$$\min_{\mathcal{X}, C, U_1, \dots, U_n} : \frac{1}{2} \| \mathcal{X} - \underline{C} \times_1 U_1 \times_2 U_2 \times_3 \dots \times_n U_n \|_F^2
s.t. : \mathcal{X}_O = \mathcal{T}_O,$$
(10)

where $C \times_1 U_1 \times_2 U_2 \times_3 \cdots \times_n U_n$ is the Tucker model-based tensor factorization, $U_i \in {\rm I\!R}^{I_i \times r_i}$, $\mathcal{C} \in {\rm I\!R}^{r_1 \times \cdots \times r_n}$, and $\mathcal{T}, \mathcal{X} \in {\rm I\!R}^{I_1 \times \cdots \times I_n}$. One can simply use the block coordinate descent method to solve this problem by iteratively optimizing two blocks \mathcal{X} and C, U_1, \ldots, U_n , respectively, while fixing the other. \mathcal{X} can be computed by letting $\mathcal{X}_\Omega = \mathcal{T}_\Omega$ and $\mathcal{X}_{\bar{\Omega}} = (C \times_1 U_1 \times_2 U_2 \times_3 \cdots \times_n U_n)_{\bar{\Omega}}$. C, U_1, \ldots, U_n can be computed by any existing tensor factorization algorithm based on the Tucker model. The procedure can also be employed to solve the following two heuristic algorithms.

Parafac. Another natural approach is to use the parallel factor analysis (Parafac) model [17], resulting in the following optimization problem:

$$\min_{\mathcal{X}, U_1, U_2, \dots, U_n} : \frac{1}{2} \| \mathcal{X} - U_1 \circ U_2 \circ \dots \circ U_n \|_F^2$$

$$s.t. : \mathcal{X}_{\Omega} = \mathcal{T}_{\Omega}$$
(11)

where \circ denotes the outer product and $U_1 \circ U_2 \circ \cdots \circ U_n$ is the Parafac model-based decomposition, $U_i \in \mathbb{R}^{I_i \times r}$, and $\mathcal{T}, \mathcal{X} \in \mathbb{R}^{I_1 \times \cdots \times I_n}$.

SVD. The third alternative is to consider the tensor as multiple matrices and force the unfolding matrix along each mode of the tensor to be <u>low rank</u> as follows:

$$\min_{\mathcal{X}, M_1, M_2, \dots, M_n} : \frac{1}{2} \sum_{i=1}^n \|\mathcal{X}_{(i)} - M_i\|_F^2$$

$$s.t. : \mathcal{X}_{\Omega} = \mathcal{T}_{\Omega}$$

$$\operatorname{rank}(M_i) \leq r_i \ i = 1, \dots, n,$$
(12)

where $M_i \in {\rm I\!R}^{I_i imes (\prod_{k \neq i} I_k)}$ and $\mathcal{T}, \mathcal{X} \in {\rm I\!R}^{I_1 imes \cdots imes I_n}$.

4 A SIMPLE LOW RANK TENSOR COMPLETION ALGORITHM

In this section, we present the SiLRTC algorithm to solve the convex model in (9), which is simple to understand and to implement. In Section 4.1, we relax the original problem into a simple convex structure which can be solved by block coordinate descent. Section 4.2 presents the details of the proposed algorithm.

4.1 Simplified Formulation

The problem in (9) is difficult to solve due to the interdependent matrix trace norm terms, i.e., while we optimize the sum of multiple matrix trace norms, the matrices share the same entries and cannot be optimized independently. Hence, the existing result in (3) cannot be used directly. Our key motivation of simplifying this original problem is how to split these interdependent terms such that they can be solved independently. We introduce additional matrices M_1, \ldots, M_n and obtain the following equivalent formulation:

$$\min_{\mathcal{X}, M_i} : \sum_{i=1}^n \alpha_i || M_i ||_*
s.t. : \mathcal{X}_{(i)} = M_i \quad \text{for } i = 1, \dots, n
\mathcal{X}_{\Omega} = \mathcal{T}_{\Omega}.$$
(13)

In this formulation, the trace norm terms are still not independent because of the equality constraints $M_i = \mathcal{X}_{(i)}$ which forces all M_i s to be identical. Thus, we relax the

equality constraints $M_i = \mathcal{X}_{(i)}$ by $||M_i - \mathcal{X}_{(i)}||_F^2 \le d_i$ as (14) so that we can independently solve each subproblem later on:

$$\min_{\mathcal{X}, M_i} : \sum_{i=1}^{n} \alpha_i \|M_i\|_*$$

$$s.t. : \|\mathcal{X}_{(i)} - M_i\|_F^2 \le d_i \quad \text{for } i = 1, \dots, n$$

$$\mathcal{X}_{\Omega} = \mathcal{T}_{\Omega}, \tag{14}$$

 $d_i(>0)$ is a threshold that could be defined by the user, but we do not use d_i explicitly in our algorithm. This optimization problem can be converted to an equivalent formulation for certain positive values of β_i s:

$$\min_{\mathcal{X}, M_i} : \sum_{i=1}^{n} \alpha_i \underline{\|M_i\|}_* + \frac{\beta_i}{2} \|\mathcal{X}_{(i)} - M_i\|_F^2$$

$$s.t. : \mathcal{X}_{\Omega} = \mathcal{T}_{\Omega}.$$
(15)

This is a convex but nondifferentiable optimization problem. Next, we show how to solve the optimization problem in (15).

4.2 The Main Algorithm

We propose to employ block coordinate descent for the optimization. The basic idea of block coordinate descent is to optimize a group (block) of variables while fixing the other groups. We divide the variables into n + 1 blocks: \mathcal{X} , M_1, M_2, \ldots, M_n .

Computing \mathcal{X} . The optimal \mathcal{X} with all other variables fixed is given by solving the following subproblem:

$$\min_{\mathcal{X}} : \sum_{i=1}^{n} \frac{\beta_i}{2} \| M_i - \mathcal{X}_{(i)} \|_F^2$$

$$s.t. : \mathcal{X}_{\Omega} = \mathcal{T}_{\Omega}.$$
(16)

It is easy to check that the solution to (16) is given by

$$\mathcal{X}_{i_1,\dots,i_n} = \begin{cases}
\left(\frac{\sum_i \beta_i \operatorname{fold}_i(M_i)}{\sum_i \beta_i}\right)_{i_1,\dots,i_n} & (i_1,\dots,i_n) \notin \Omega; \\
\mathcal{T}_{i_1,\dots,i_n} & (i_1,\dots,i_n) \in \Omega.
\end{cases} (17)$$

Computing M_i . M_i is the optimal solution of the following problem:

$$\min_{M_{i}} : \frac{\beta_{i}}{2} \| M_{i} - \mathcal{X}_{(i)} \|_{F}^{2} + \alpha_{i} \| M_{i} \|_{*}
\equiv \frac{1}{2} \| M_{i} - \mathcal{X}_{(i)} \|_{F}^{2} + \frac{\alpha_{i}}{\beta_{i}} \| M_{i} \|_{*}.$$
(18)

This problem has been proven to lead to a closed form in recent papers like [30], [7]. Thus, the optimal M_i can be computed as $\mathbf{D}_{\tau}(\mathcal{X}_{(i)})$, where $\tau = \frac{\alpha_i}{\beta_i}$. We call the proposed algorithm "SiLRTC," which stands

We call the proposed algorithm "SiLRTC," which stands for Simple Low Rank Tensor Completion algorithm. The pseudocode of the SiLRTC algorithm is given in Algorithm 1 below. As convergence criteria we compare the difference of \mathcal{X} in subsequent iterations to a threshold. Since the objective in (15) is convex and the nonsmooth term is separable, BCD is guaranteed to find the global optimal solution [45]. Note that this SiLRTC algorithm actually simplifies the LRTC algorithm proposed in our conference paper [29] by removing a redundant variable \mathcal{Y} . SiLRTC and LRTC produce almost identical results.

Algorithm 1. SiLRTC: Simple Low Rank Tensor Completion **Input:** \mathcal{X} with $\mathcal{X}_{\Omega} = \mathcal{T}_{\Omega}$, β_i s, and K

Output: X

1: for k = 1 to K do

2: **for** i = 1 to n **do**

3: $M_i = \mathbf{D}_{\frac{\alpha_i}{4}}(\mathcal{X}_{(i)})$

4: end for

5: update \mathcal{X} by (17).

6: end for

5 A FAST LOW RANK TENSOR COMPLETION ALGORITHM

Although the proposed algorithm in Section 4 is easy to implement, its convergence speed is low in practice. In addition, SiLRTC is hard to extend to any tensor trace norm minimization problem, e.g., the formulation "logistic loss + tensor trace norm" is hard to minimize using the strategy above. In this section, we propose a new algorithm to significantly improve the convergence speed of SiLRTC and to solve a general tensor trace norm minimization problem defined below:

$$\min_{\mathcal{X} \in Q} : f(\mathcal{X}) := f_0(\mathcal{X}) + \sum_{i=1}^n \alpha_i \|\mathcal{X}_{(i)}\|_*, \tag{19}$$

where $\mathcal{X} \in \mathbb{R}^{I_1 \times I_2 \times \cdots \times I_n}$, Q is a convex set, and $f_0(\mathcal{X})$ is smooth and convex. One can easily verify that the low rank tensor completion problem in (9) is just a special case with $f_0(\mathcal{X}) = 0$ and $Q = \{\mathcal{X} \in \mathbb{R}^{I_1 \times I_2 \times \cdots \times I_n} | \mathcal{X}_\Omega = \mathcal{T}_\Omega \}$.

The difficulty in efficiently solving the tensor trace norm-related minimization problems lies in the fact that there exist multiple dependent nonsmooth terms in the objective function. Although one can use the subgradient information to replace the gradient information, the convergence rate is $O(K^{-1/2})$, where K is the iteration number [33]. In comparison, the optimal convergence rate for minimizing general smooth functions is $O(K^{-2})$ [33], [31]. Nesterov [34] proposed a general method to solve a nonsmooth optimization problem. The basic idea is to

- first convert the original nonsmooth problem into a smooth one;
- then solve the smooth problem and use its solution to approximate the original problem.

We will follow this procedure to solve the problem in (19).

Section 5.1 employs the smoothing scheme proposed by Nesterov [34] to convert the original nonsmooth objective function in (19) into a smooth one. Section 5.2 proposes an efficient updating rule to solve the smooth problem and analyzes the convergence rate of this method. Section 5.3 applies the smoothing scheme and the efficient updating rule to the low rank tensor completion problem.

5.1 Smoothing Scheme

Consider one nonsmooth term in (19), i.e., the matrix trace norm function $\|X\|_*$. Its dual version can be written as

$$g(X) := \|X\|_* = \max_{\|Y\| \le 1} \langle X, Y \rangle. \tag{20}$$

Its smooth version is

$$g_{\mu}(X) = \max_{\|Y\| \le 1} \langle X, Y \rangle - d_{\mu}(Y), \tag{21}$$

where $d_{\mu}(Y)$ is a strongly convex function with the parameter μ . Nesterov [34, Theorem 1] proves that $g_{\mu}(X)$ is smooth and its gradient can be computed by

$$\nabla g_{\mu}(X) = Y^*(X) := \arg \max_{\|Y\| \le 1} \langle X, Y \rangle - d_{\mu}(Y). \tag{22}$$

Although one can arbitrarily choose a strongly convex function $d_{\mu}(Y)$ to smooth the original function, a good choice can lead to a closed form for the dual variables Y. Otherwise, it involves a complicated min-max optimization problem. Here, we choose $d(Y) = \frac{\mu}{2} ||Y||_F^2$, where $\mu > 0$ as the strongly convex term and the gradient is

$$\nabla g_{\mu}(X) = Y^{*}(X) := \arg \max_{\|Y\| \le 1} \langle X, Y \rangle - \frac{\mu}{2} \|Y\|_{F}^{2}$$

$$= \arg \min_{\|Y\| \le 1} \|Y - \frac{1}{\mu} X\|_{F}^{2} = \mathbf{T}_{1} \left(\frac{1}{\mu} X\right). \tag{23}$$

The last equality is due to Lemma 1, available online in the supplemental material.

We apply this smoothing scheme to all nonsmooth terms in (19) by introducing n dual variables $\mathcal{Y}_1,\ldots,\mathcal{Y}_n\in\mathbb{R}^{I_1\times I_2\times\cdots\times I_n}$ and n positive constants μ_1,\ldots,μ_n . The objective function f(X) is converted into

$$f_{\mu}(\mathcal{X}) := f_{\mu_{1},\dots,\mu_{n}}(\mathcal{X})$$

$$= f_{0}(\mathcal{X}) + \sum_{i=1}^{n} \max_{\|\mathcal{Y}_{i(i)}\| \leq 1} \alpha_{i} \langle \mathcal{X}_{(i)}, \mathcal{Y}_{i(i)} \rangle - \frac{\mu_{i}}{2} \|\mathcal{Y}_{i(i)}\|_{F}^{2}$$

$$= f_{0}(\mathcal{X}) + \sum_{i=1}^{n} \max_{\|\mathcal{Y}_{i(i)}\| \leq 1} \alpha_{i} \langle \mathcal{X}, \mathcal{Y}_{i} \rangle - \frac{\mu_{i}}{2} \|\mathcal{Y}_{i}\|_{F}^{2}.$$

$$(24)$$

Its gradient can be computed by

$$\nabla f_{\mu}(\mathcal{X}) = \nabla f_0(\mathcal{X}) + \sum_{i=1}^n \alpha_i \mathbf{T}_1 \left(\frac{\alpha_i}{\mu_i} \mathcal{X}_{(i)} \right). \tag{25}$$

Finally, we obtain a smooth optimization problem as follows:

$$\min_{\mathcal{X} \in Q \subset \mathbb{R}^{I_1 \times \dots \times I_n}} : f_{\mu}(\mathcal{X}) \tag{26}$$

and will use its optimal solution to approximate the original problem in (19).

5.2 An Efficient Algorithm to Solve the Smooth Version

In essence, the problem in (26) is a smooth optimization. Nesterov [34] also proposed an algorithm to solve any smooth problem in a bounded domain and guaranteed two convergence rates $O(K^{-2})$ for the smooth problem and $O(K^{-1})$ for the original problem, i.e.,

$$f_{\mu}(\mathcal{X}^K) - f_{\mu}(\mathcal{X}_{\mu}^*) \le O(K^{-2}), \tag{27}$$

$$f(\mathcal{X}^K) - f(\mathcal{X}^*) \le O(K^{-1}),\tag{28}$$

where \mathcal{X}_{μ}^{*} and \mathcal{X}^{*} are, respectively, the optimal solutions of $f_{\mu}(\mathcal{X})$ and $f(\mathcal{X})$, \mathcal{X}^0 is the initial point, and \mathcal{X}^K is the output of our updating rule for $f_{\mu}(\mathcal{X})$. One is not surprised about the first convergence rate since, for a general smooth optimization problem, the rate $O(K^{-2})$ is guaranteed by Nesterov's popular accelerated algorithm [32]. The second rate is quite interesting. Note that \mathcal{X}^K is the output by minimizing $f_{\mu}(\mathcal{X})$ instead of $f(\mathcal{X})$. The second inequality indeed indicates how far the approximate solution is away from the true solution after *K* iterations.

However, the constant factor of $O(K^{-1})$ in (28) is proportional to the size of the domain set Q [34, Theorem 3]. For this reason, the domain *Q* is assumed to be bounded in [34]. Hence, if this algorithm is applied to solve (26) directly, the inequality in (28) cannot be guaranteed. Based on the algorithm in [34], we propose an improved efficient algorithm to solve the problem (26) in Algorithm 2 which allows the domain set Q to be unbound and can guarantee the results in (27) and (28). A more detailed explanation of the accelerated scheme in Algorithm 2 is provided in the supplemental material, available online.

Algorithm 2. An Efficient Algorithm

Input: $c \in (0,1)$, x^0 , K, and L^0 .

Output: x^K

- 1: Initialize $z^0 = w^0 = x^0$ and $B^0 = 0$
- 2: **for** k = 0 to K **do**
- 3: Find a L^{k+1} as small as possible from $\{\cdots, cL^k, L^k, L^k/c, \ldots\}$, such that Eq. (31) holds. Let

$$\theta^{k+1} = \frac{L^k}{2L^{k+1}} (1 + \sqrt{1 + 4B^k L^{k+1}}), \quad w^{k+1}$$
$$= \tau^{k+1} z^k + (1 - \tau^{k+1}) x^k$$
(29)

$$x' = \arg\min_{x \in Q} : p(x) = f_{\mu}(w^{k+1}) + \langle \nabla f_{\mu}(w^{k+1}),$$

$$x - w^{k+1} \rangle + \frac{L^{k+1}}{2} ||x - w^{k+1}||^{2}$$
(30)

where $au^{k+1}=(frac{ heta^{k+1}}{L^k})/B^{k+1}$ and $B^k=\sum_{i=1}^k frac{ heta^i}{L^{i-1}}.$ Test whether the following inequality holds:

$$f_{\mu}(x') \le p(x^{k+1}),$$
 (31)
4: Update x^{k+1} and z^{k+1} by

$$\begin{split} x^{k+1} &= \arg\min_{x \in \{x', x^k, z^k\}} : f_{\mu}(x), \\ z^{k+1} &= \arg\min_{z \in Q} : h(z) = \frac{1}{2} \|z - x^0\|^2 \\ &+ \sum_{i=1}^{k+1} \frac{\theta^i}{L^{i-1}} (f_{\mu}(w^i) + \langle \nabla f_{\mu}(w^i), z - w^i \rangle) \end{split} \tag{32}$$

5: end for

Unsurprisingly, the proposed algorithm can guarantee the convergence rate $O(K^{-2})$ for the smooth problem like many existing accelerated algorithms [33], [34], [19], [31], see Theorem 2, available online in the supplemental material.

At the same time, the following theorem shows that the convergence rate $O(K^{-1})$ for the original nonsmooth problem is also obtained by the proposed algorithm.

Theorem 1. Define D as any positive constant satisfying

$$D \ge \min_{\mathcal{X}^*} : \|\mathcal{X}^* - \mathcal{X}^0\|_F, \tag{34}$$

where \mathcal{X}^* is the optimal solution of the problem (19) and \mathcal{X}^0 is the starting point. Set the parameters in the problem (26) as

$$\mu_i = \frac{2\alpha_i D}{K\sqrt{cI_i}}.$$

After K iterations in Algorithm 2, the output \mathcal{X}^K for the problem in (26) satisfies

$$f(\mathcal{X}^K) - f(\mathcal{X}^*) \le \frac{2\bar{L}}{c} \left(\frac{D}{K}\right)^2 + \frac{2D}{K} \sum_i \alpha_i \sqrt{\frac{I_i}{c}},$$
 (35)

where \bar{L} is the Lipschitz constant of $f_0(\mathcal{X})$.

Theorems 1 and 2 extend the results in [34].

Although Nesterov's popular accelerated algorithm [32], [31], [33] can guarantee the convergence rate $O(K^{-2})$ for the smooth problem, there is no evidence showing that it can achieve $O(K^{-1})$ for the original problem. Ji et al. [19] used this smoothing scheme and Nesterov's popular accelerated method to solve a multiple kernel learning problem and also claimed the convergence rate $O(K^{-1})$ for the original problem. However, what they really guaranteed is $f(\mathcal{X}^K) - f(\mathcal{X}_u^*) \leq O(K^{-1})$, which can be obtained from (27).

Theorem 1 also implies that the reasonable parameters should satisfy

$$\mu_1: \mu_2: \dots: \mu_n = \frac{\alpha_1}{\sqrt{I_1}}: \frac{\alpha_2}{\sqrt{I_2}}: \dots: \frac{\alpha_n}{\sqrt{I_n}}.$$

The FaLRTC Algorithm

This section considers the specific tensor completion problem in (9). The smooth version of this problem is

$$\min_{\mathcal{X}} : \sum_{i=1}^{n} \max_{\|\mathcal{Y}_{i(i)}\| \le 1} : \alpha_{i} \langle \mathcal{X}, \mathcal{Y}_{i} \rangle - \frac{\mu_{i}}{2} \|\mathcal{Y}\|_{F}^{2}$$

$$st : \mathcal{X}_{O} = \mathcal{T}_{O} \tag{36}$$

Now we can use the proposed Algorithm 2 to solve this smooth minimization problem above. First it is easy to verify

$$\mathcal{Z}^{k+1} = \mathcal{Z}^k - \frac{\theta^{k+1}}{L^k} \nabla f_{\mu}(\mathcal{W}^{k+1}), \tag{37}$$

$$(\nabla f_{\mu}(\mathcal{W}^{k+1}))_{i_{1},\dots,i_{n}}$$

$$= \begin{cases} \left(\sum_{i} \frac{(\alpha_{i})^{2}}{\mu_{i}} \mathbf{T}_{\frac{\mu_{i}}{a_{i}}} (\mathcal{W}_{(i)}^{k+1})\right)_{i_{1},\dots,i_{n}}, & (i_{1},i_{2},\dots,i_{n}) \notin \Omega; \\ 0, & (i_{1},i_{2},\dots,i_{n}) \in \Omega. \end{cases}$$
(38)

The last equation is due to $W_{\Omega}^{k+1} = T_{\Omega}$. For a simple implementation, we require the sequence L^k to be nondecreasing, although the updating rule in Algorithm 2 allows that L^k (L' in Algorithm 3) becomes smaller (theoretically, the smaller, the better, see the proof of Theorem 2). Because $f(\mathcal{X})$ involves computing SVD (this is a big workload), we remove \mathcal{Z}^k from the candidate pool. Note that these slight changes do not change the properties of the output \mathcal{X}^K in Theorems 2 and 1. Finally, Algorithm 3 summarizes the fast low rank tensor completion algorithm, namely, FaLRTC. Lines 5 to 7 in Algorithm 3 are optional as they merely guarantee that the objective is nonincreasing. Lines 8 to 13 are the line search step. The main cost in this part is evaluating $f_{\mu}(\mathcal{X}')$, $f_{\mu}(\mathcal{W})$, and $\nabla f_{\mu}(\mathcal{W})$. In fact, one of $f_{\mu}(\mathcal{W})$ and $\nabla f_{\mu}(\mathcal{W})$ can be obtained without much additional cost while computing the other. Since all gradient algorithms have to compute the gradient, the additional cost in each iteration of the proposed algorithm is to compute $f_{\mu}(\mathcal{X}')$. One can avoid this extra cost by initialing L as the Lipschitz constant of the objective $f_{\mu}(\mathcal{W})$, i.e., $\sum_{i=1}^n \frac{1}{\mu_i}$ because it satisfies the condition in line 9. However, in practice a line search often improves the efficiency. In addition, the FaLRTC algorithm can be accelerated by decreasing μ_i iteratively. Specifically, we set $\mu_i^k = ak^{-p} + b$ for $k = 1, \ldots, K$ at the kth iteration where p is a constant in the range [1.1, 1.2] and a and b are determined by solving $\mu_i^0 = a + b$ and $\mu_i^K = aK^{-p} + b$ ($\mu_i^0 = 0.4\alpha_i \|\mathcal{X}_{(i)}\|$, $\mu_i^K = \mu_i$ is the input of Algorithm 3).

Algorithm 3. FaLRTC: Fast Low Rank Tensor Completion **Input:** $c \in (0,1)$, \mathcal{X} with $\mathcal{X}_{\Omega} = \mathcal{T}_{\Omega}$, K, μ_i s, and L. **Output:** \mathcal{X}

1: Initialize
$$\mathcal{Z} = \mathcal{W} = \mathcal{X}$$
, $L' = L$, and $B = 0$
2: **for** $k = 0$ to K **do**
3: **while** true **do**
4: $\theta = \frac{L}{2L'}(1 + \sqrt{1 + 4L'B});$ $\mathcal{W} = \frac{\theta/L}{B + \theta/L}\mathcal{Z} + \frac{B}{B + \theta/L}\mathcal{X};$ (39)
5: **if** $f_{\mu}(\mathcal{X}) \leq f_{\mu}(\mathcal{W}) - \|\nabla f_{\mu}(\mathcal{W})\|_F^2/2L'$ **then**
6: break;

6: break;
7: end if
8:
$$\mathcal{X}' = \mathcal{W} - \nabla f_{\mu}(\mathcal{W})/L';$$
9: if $f_{\mu}(\mathcal{X}') \leq f_{\mu}(\mathcal{W}) - \|\nabla f_{\mu}(\mathcal{W})\|_F^2/2L'$ then
10: $\mathcal{X} = \mathcal{X}';$
11: break;
12: end if
13: $L' = L'/c;$
14: end while
15:
$$L = L';$$

$$\mathcal{Z} = \mathcal{Z} - \frac{\theta}{L} \nabla f_{\mu}(\mathcal{W});$$

$$B = B + \frac{\theta}{L};$$
(40)

16: end for

6 A HIGH ACCURACY LOW RANK TENSOR COMPLETION ALGORITHM

The ADMM algorithm was developed in the 1970s, with roots in the 1950s, but received renewed interest due to the fact that it is efficient to tackle large scale problems and solve optimization problems with multiple nonsmooth terms in the objective [28]. This section follows the ADMM algorithm to solve the noiseless case in a direct way. Based on the SiLRTC algorithm, we also give a simple implementation using the ADMM framework. Recall that the formulation in (13) is an equivalent form of the original problem. We replace the dummy matrices M_i s by their tensor versions:

$$\min_{\mathcal{X}, \mathcal{M}_{1}, \dots, \mathcal{M}_{n}} : \sum_{i=1}^{n} \alpha_{i} \| \mathcal{M}_{i(i)} \|_{*}$$

$$s.t. : \mathcal{X}_{\Omega} = \mathcal{T}_{\Omega}$$

$$\mathcal{X} = \mathcal{M}_{i}, \quad i = 1, \dots, n.$$
(41)

We define the augmented Lagrangian function as follows:

$$L_{\rho}(\mathcal{X}, \mathcal{M}_{1}, \dots, \mathcal{M}_{n}, \mathcal{Y}_{1}, \dots, \mathcal{Y}_{n})$$

$$= \sum_{i=1}^{n} \alpha_{i} \|\mathcal{M}_{i(i)}\|_{*} + \langle \mathcal{X} - \mathcal{M}_{i}, \mathcal{Y}_{i} \rangle + \frac{\rho}{2} \|\mathcal{M}_{i} - \mathcal{X}\|_{F}^{2}.$$
(42)

According to the framework of ADMM, one can iteratively update \mathcal{M}_i s, \mathcal{X} , and \mathcal{Y}_i s as follows:

1.
$$\{\mathcal{M}_{1}^{k+1}, \dots, \mathcal{M}_{n}^{k+1}\} = arg\min_{\mathcal{M}_{1}, \dots, \mathcal{M}_{n}}$$
:
 $L_{\rho}(\mathcal{X}^{k}, \mathcal{M}_{1}, \dots, \mathcal{M}_{n}, \mathcal{Y}_{1}^{k+1}, \dots, \mathcal{Y}_{n}^{k+1})$.
2. $\mathcal{X}^{k+1} = arg\min_{\mathcal{X} \in \mathcal{Q}}$:
 $L_{\rho}(\mathcal{X}, \mathcal{M}_{1}^{k+1}, \dots, \mathcal{M}_{n}^{k+1}, \mathcal{Y}_{1}^{k}, \dots, \mathcal{Y}_{n}^{k+1})$.
3. $\mathcal{Y}_{i}^{k+1} = \mathcal{Y}_{i}^{k} - \rho(\mathcal{M}_{i}^{k+1} - \mathcal{X}^{k+1})$.

One can refer to (18) and (16) in the SiLRTC algorithm to obtain the closed form solutions for the first two steps. We summarize the HaLRTC algorithm in Algorithm 4. This algorithm can also be accelerated by adaptively changing ρ . An efficient strategy [28] is to let $\rho^0 = \rho$ (the input in Algorithm 4) and increase ρ^k iteratively by $\rho^{k+1} = t\rho^k$, where $t \in [1.1, 1.2]$.

Algorithm 4. HaLRTC: High Accuracy Low Rank Tensor Completion

Input:
$$\mathcal{X}$$
 with $\mathcal{X}_{\Omega} = \mathcal{T}_{\Omega}$, ρ , and K
Output: \mathcal{X}

1: Set $\mathcal{X}_{\Omega} = \mathcal{T}_{\Omega}$ and $\mathcal{X}_{\bar{\Omega}} = 0$.

2: for $k = 0$ to K do

3: for $i = 1$ to n do

4: $\mathcal{M}_i = \operatorname{fold}_i \left[\mathbf{D}_{\frac{\alpha_i}{\rho}} \left(\mathcal{X}_{(i)} + \frac{1}{\rho} \mathcal{Y}_{i(i)} \right) \right]$

5: end for

6: $\mathcal{X}_{\Omega} = \frac{1}{n} \left(\sum_{i=1}^{n} \mathcal{M}_i - \frac{1}{\rho} \mathcal{Y}_i \right)_{\bar{\Omega}}$

7: $\mathcal{Y}_i = \mathcal{Y}_i - \rho(\mathcal{M}_i - \mathcal{X})$

8: end for

Note that the proposed ADMM algorithm in this section aims to solve the tensor completion problem without observation noise, unlike the previous work in [38], [43], and [14]. Signoretto et al. [38] and Gandy et al. [14] also consider the noiseless case in (9). They relax the equality constraint in (9) into the noisy case in (4) and apply the ADMM framework to solve the relaxed problem with an increasing value of λ . However, our ADMM algorithm handles this equality constraint directly without using any relaxation technique. The comparison in Section 7 will show that our ADMM is more efficient than the ADM-TR Algorithm in [14].

Although the convergence of the general ADMM algorithm is guaranteed [28], the convergence rate may be slow. From the comparison in Section 7.3, we can observe that HaLRTC is comparable to FaLRTC and even more efficient to achieve a higher accuracy.

7 RESULTS

In this section, we first validate the tensor trace norm-based model in (9) by comparing to three heuristic models in Section 3.2 and the matrix completion model for tensor

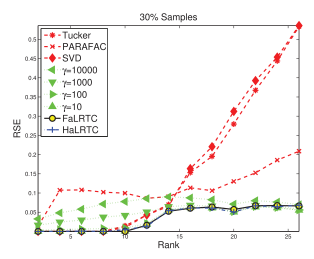


Fig. 2. The RSE comparison on the synthetic data. $I_1=I_2=I_3=50$. The ranks of the tensor are given by $r_1=r_2=r_3=r$, where r is equivalent to $2,4,6,8,\ldots,26$, respectively. Tucker: Tucker model heuristic algorithm; Parafac: Parafac model-based heuristic algorithm; SVD: the heuristic algorithm based on the SVD; $\gamma=10,000,\ \gamma=1,000,\ \gamma=100,$ and $\gamma=10$ denote the proposed SiLRTC algorithm with $\beta_i=1$ and $\gamma_i=\gamma$. We let $\mu=5$ in FaLRTC algorithm. The sample percentage is 30 percent.

completion. Then the efficiency comparison between SiLRTC, FaLRTC, and HaLRTC is reported. Several applications of tensor completion conclude this section. All experiments were implemented in Matlab (version 7.9.0) and all tests were performed on an Intel Core 2 2.67 GHz and 3 GB RAM computer.

In the following, we will use the 3-mode tensor $\mathcal{T} \in \mathbb{R}^{I_1 \times I_2 \times I_3}$ to explain how we generated the synthetic test data sets used in the following comparison. All synthetic data in Section 7 are generated in this manner. The tensor data follows the tucker model, i.e., $\mathcal{T} = \mathcal{C} \times_1 U_1 \times_2 U_2 \times_3 U_3$, where the core tensor \mathcal{C} is of size $r_1 \times r_2 \times r_3$ and r_i is of size $r_i \times r_i$. The entry of r_i is computed by

$$\mathcal{T}(i,j,k) = \sum_{1 \le m,n,l \le r} \mathcal{C}(m,n,l) U_1(i,m) U_2(j,n) U_3(k,l).$$
 (43)

The entries of U_i are random samples drawn from a uniform distribution in the range [-0.5, 0.5] and the entries of $\mathcal C$ are from a uniform distribution in the range [0,1]. Typically, the ranks of the tensor $\mathcal T$ unfolded respectively along three modes are $[r_1, r_2, r_3]$. The data is finally normalized such as $\|\mathcal T\|_F =$ the number of entries in $\mathcal T$.

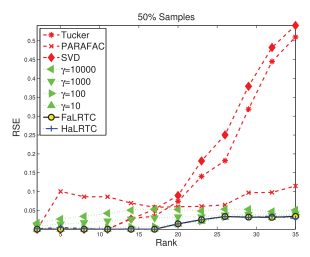


Fig. 3. See the caption of Fig. 2. The only difference is the sample percentage of 50 percent.

7.1 Model Comparison: Tensor Trace Norm versus Heuristic Models

We compare the proposed tensor completion model in (9) to three heuristic models in Section 3.2 on both synthetic and real-world data. We use SiLRTC, FaLRTC, and HaLRTC to solve (9) and compare their results to three heuristic methods. Since three heuristic models are nonconvex, multiple initial points are tested and the average performance is used for comparison. The initial value at each missing entry is generated from the Guassian distribution $N(\mu,1)$, where μ is the average of other observed entries. All experiments are repeated 10 times. The performance is measured by $RSE = \|\mathcal{X} - \mathcal{T}\|_F / \|\mathcal{T}\|_F$. One can easily verify its connection to the signal-to-noise ratio (SNR) by $SNR = -20\log_{10}RSE$.

In the SiLRTC algorithm, the value of α_i is set to 1/3 (3 is the mode number) and we only change the value of β_i . Let $\beta_i = \alpha_i/\gamma_i$. It is easy to see that when the γ_i s go to 0, the optimization problem in (15) converges to the original problem (7). In the FaLRTC algorithm, the parameters α_i s are set like in the SiLRTC algorithm and the parameters μ_i s are set as $\mu_i = \mu \frac{\alpha_i}{\sqrt{r_i}}$. Typically, μ is in the range [1, 10], if the data is normalized as above. The rank parameters of th three heuristic algorithms follow the ranks of the tensor $\mathcal T$ unfolded along each mode.

We choose the percentage of randomly sampled elements as 30 and 50 percent, respectively, and present the comparison in Figs. 2 and 3.

The brain MRI data is of size $181 \times 217 \times 181$. Although its ranks unfolded along three modes are [164, 198, 165],

TABLE 1
The RSE Comparison on the Brain MRI Data

RSE Comparison (10^{-4})

| Samples | Tucker | Parafac | SVD | SiLRTC $\gamma = 10^4$ | SiLRTC $\gamma = 10^3$ | SiLRTC $\gamma = 10^2$ | SiLRTC $\gamma = 10$ | FaLRTC | HaLRTC |
|---------|--------|---------|-----|------------------------|------------------------|------------------------|----------------------|--------|--------|
| 20% | 371 | 234 | 274 | 309 | 47 | 22 | 21 | 17 | 16 |
| 50% | 65 | 58 | 62 | 101 | 12 | 2 | 0 | 0 | 0 |
| 80% | 21 | 45 | 16 | 40 | 4 | 1 | 0 | 0 | 0 |

Tucker: Tucker model heuristic algorithm; Parafac: Parafac model-based heuristic algorithm; SVD: the heuristic algorithm based on the SVD; $\gamma=10,000,\,\gamma=1,000,\,\gamma=100,$ and $\gamma=10$ correspond to the proposed SiLRTC algorithm with $\beta_i=1$ and $\gamma_i=\gamma$. In the FaLRTC algorithm, μ is fixed as 5. We try different parameter values for three heuristic algorithms and report the best performance we obtain. The top, middle, and bottom parts of the table correspond to the sample percentage: 20, 50, and 80 percent, respectively.

TABLE 2 The RSE Comparison on the Synthetic Data with $r_i=2$

| RSE Comparison (10^{-4}), Size $20 \times 20 \times 20$ | | | | | | | | |
|---|------|------|------|----|--|--|--|--|
| Samples | MC1 | MC2 | MC3 | TC | | | | |
| 25% | 1663 | 1782 | 1685 | 34 | | | | |
| 40% | 247 | 258 | 241 | 2 | | | | |
| 60% | 0 | 0 | 0 | 0 | | | | |
| RSE Comparison (10 ⁻⁴). Size $20 \times 20 \times 20 \times 20$ | | | | | | | | |

| Samples | MC1 | MC2 | MC3 | MC4 | TC |
|---------|------|------|------|------|----|
| 20% | 1875 | 1763 | 2011 | 1804 | 3 |
| 40% | 92 | 102 | 97 | 88 | 0 |
| 60% | 0 | 1 | 0 | 0 | 0 |

RSE Comparison (10^{-4}), Size $20 \times 20 \times 20 \times 20 \times 20$ Samples II MC1 MC2 MC3 MC4 MC5 TC

| Samples | MC1 | MC2 | MC3 | MC4 | MC5 | TC |
|---------|------|------|------|------|------|----|
| 15% | 1874 | 1830 | 1663 | 1502 | 1688 | 50 |
| 40% | 125 | 119 | 131 | 114 | 136 | 0 |
| 60% | 0 | 0 | 0 | 0 | 0 | 0 |

MCk $(k=1,2,3,\dots)$ denotes the performance of the solution of the problem (44) with i=k. We use the FaLRTC algorithm to do tensor completion with $\mu=1$. The top, middle, and bottom parts of the table respond to different sizes of the synthetic data.

TABLE 3
The RSE Comparison on the MRI Brain Data

RSE Comparison (10^{-4})

| Sampl | es | MC1 | MC2 | MC3 | TC |
|-------|------|-----|-----|-----|----|
| 20% | · II | 670 | 781 | 862 | 17 |
| 50% | , | 27 | 36 | 75 | 0 |

 $\mu=5$ in the FaLRTC algorithm. MCk $(k=1,2,3,\ldots)$ denotes the performance of the solution of the problem (44) with i=k.

TABLE 4 The RSE Comparison on the Synthetic Data with the Size $60 \times 60 \times 60$

RSE Comparison (10^{-4}) , Samples percentage = 20%

| | S-MC1 | S-MC2 | S-MC3 | TC |
|-------------|-------|-------|-------|----|
| $r_i's = 2$ | 2224 | 2317 | 2365 | 0 |
| $r_i's = 4$ | 4887 | 4756 | 4441 | 1 |
| $r_i's = 6$ | 6479 | 6789 | 6247 | 1 |

S-MCk $(k=1,2,3,\dots)$ denotes the performance by applying matrix completion method on each slice. We use the FaLRTC algorithm to do tensor completion with $\mu=1$.

they can decrease to [35,42,36] if removing small singular values less than 0.01 percent of its Frobenius norm. Thus this is approximately an low rank data. We use the same normalization strategy above and report the comparison results in Table 1.

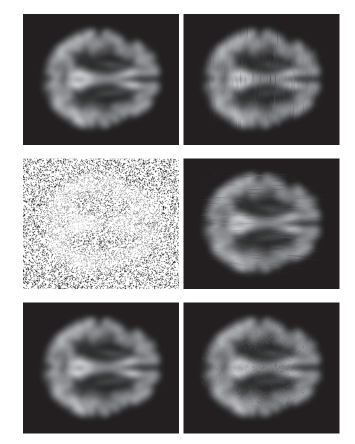


Fig. 5. The comparison of tensor completion and matrix completion. The left up image (one slice of the MRI) is the original; we randomly select pixels for removal, shown in white in the left middle image; the left bottom image is the reconstruction by the proposed FaLRTC algorithm with $\mu=5$; the right top, middle, and bottom images are, respectively, the results of matrix completion algorithms MC1, MC2, and MC3.

Results from Figs. 2 and 3 and Table 1 show that the proposed convex formulation outperforms the three heuristic algorithms. The performance of the three heuristic algorithms is poor for high rank problems. We can also observe that the proposed convex formulation is able to recover the missing information using a small number of samples. Next, we evaluate the SiLRTC algorithm using different parameters. We observe that the smaller the γ value, the closer the solution of (15) is to the original problem in (9). The performance of FaLRTC is similar to the SiLRTC algorithm with $\gamma=10$.

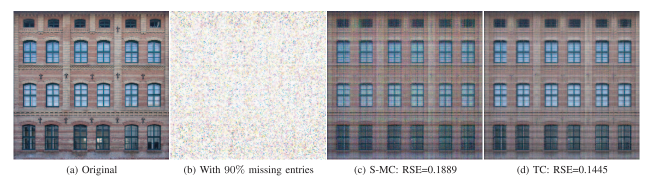


Fig. 4. The RSE comparison on the real image between tensor completion and S-MC. (a) The original image. (b) We randomly remove 90 percent entries from the original color image (3-mode tensor) and fill the value "255" on them. (c) The recovered result by S-MC which slices the image into red, green, and blue channels. (d) The recovered result by FaLRTC.

TABLE 5
The Efficiency Comparison on the Synthetic Data

| | Runnir | ig Time (sec) $I_1 =$ | $I_2 = I_3 = 100$ | $r_1 = r_2 = r_3 =$ | 2 Samples 20% | 6 | |
|--------------------------|-----------------|--------------------------|-------------------|---------------------|-----------------|------------------|------------------|
| RSE(SNR) | $10^{-1}(20db)$ | $3 \times 10^{-2}(30db)$ | $10^{-2}(40db)$ | $10^{-3}(60db)$ | $10^{-4}(80db)$ | $10^{-5}(100db)$ | $10^{-6}(120db)$ |
| SiLRTC: $\gamma = 10^4$ | 8.4674 | - | - | - | - | - | - |
| SiLRTC: $\gamma = 10^3$ | 83.7156 | 90.7864 | - | - | - | - | - |
| SiLRTC: $\gamma = 10^2$ | - | - | - | - | - | - | - |
| ADM-TR | 15.9333 | 28.7841 | 65.8791 | 94.1238 | - | - | - |
| FaLRTC: $\mu = 10^3$ | 2.8477 | 3.5597 | 8.5432 | 53.3950 | - | - | - |
| FaLRTC: $\mu = 10^2$ | 2.9097 | 3.6372 | 8.7292 | 37.4629 | 66.1966 | - | - |
| FaLRTC: $\mu = 10^1$ | 2.9532 | 3.6915 | 8.4905 | 36.9152 | 63.4942 | 68.6623 | - |
| FaLRTC: $\mu = 10^0$ | 3.0084 | 3.7605 | 8.6491 | 37.6047 | 64.3041 | 69.5687 | - |
| HaLRTC: $\rho = 10^{-8}$ | 9.2625 | 10.8970 | 11.9867 | 15.8007 | 19.6146 | 23.4286 | 27.2425 |
| HaLRTC: $\rho = 10^{-7}$ | 5.4485 | 7.3555 | 8.9900 | 11.9867 | 16.0731 | 19.8871 | 23.7010 |
| HaLRTC: $\rho = 10^{-6}$ | 3.5415 | 5.4485 | 7.3555 | 10.8970 | 14.7110 | 18.5249 | 22.3389 |
| Hal RTC: $a = 10^{-5}$ | _ | | | | _ | _ | _ |

Running Time (sec) $I_1 = I_2 = I_3 = I_4 = 50 \ r_1 = r_2 = r_3 = r_4 = 2 \ \text{Samples } 20\%$ $0^{-1}(20db) \quad 3 \times 10^{-2}(30db) \quad 10^{-2}(40db) \quad 10^{-3}(60db) \quad 10^{-4}(80db) \quad 10^{-4}(80db)$ 6(120db)RSE(SNR) $10^{-1}(20db)$ (100db)SiLRTC: $\gamma = 10^4$ 398.7844 SiLRTC: $\gamma = 10^3$ 554.4987 467.3572 SiLRTC: $\gamma = 10^2$ ADM-TR 198.9887226.7895 398.7835 588.9871 FaLRTC: $\mu = 10^3$ 25.359230.9946 76.0776 349.3935 509.7864 FaLRTC: $\mu = 10^2$ 25.9213 31.6816 77.7640 336.9775 514.9781 544.2382 FaLRTC: $\mu = 10^1$ 26.3341 32.1861 79.0023339.4174 FaLRTC: $\mu = 10^0$ HaLRTC: $\rho = 10^{-9}$ HaLRTC: $\rho = 10^{-8}$ 26.449632.3273 79.3488340.9059 517.2366 546.6251 82.5930 107.6211 112.6268 150.1690 185.2085 217.7451 252.7845 217.7451 52.559272.581777.5873 112.6268 147.6662 182.7056HaLRTC: $\rho = 10^{-7}$ 37.5423 $\overline{35.0394}$ 67.5761 167.6887 200.225492.6042 130.1465HaLRTC: $\rho = 10^{-6}$

We run the proposed SiLRTC algorithm with $\beta_i=1$ and three different values of γ_i (assume all γ_i s are identical to γ as before): 10^4 , 10^3 , and 10^2 . ADM-TR denotes the ADMM algorithm proposed in [14]. We report the best result we obtain by tuning the main parameters in this algorithm. The FaLRTC algorithm is run with four different values of μ (recall $\mu_i=\mu\frac{\alpha_i}{\sqrt{l_i}}$): 1,000, 100, 100, and 1. μ_i^k is updated iteratively based on the scheme in Section 5.3, i.e., $\mu_i^K=\mu\frac{\alpha_i}{\sqrt{l_i}}$, $\mu_i^0=0.4\alpha_i\|\mathcal{X}_{(i)}\|$, and $\mu_i^k=ak^{-p}+b$ with p=1.15 and K=200. We run the HaLRTC algorithm with five different inputs for ρ : 10^{-9} , 10^{-8} , 10^{-7} , 10^{-6} , and 10^{-5} . ρ is iteratively increased as discussed in Section 6, i.e., $\rho^{(0)}=\rho$, $\rho^{k+1}=t\rho^k$, where t=1.15. We use the accuracy measured by RSE from 10^{-1} to 10^{-6} as the stopping criteria and report the running time when the required accuracy is obtained. The top and bottom parts of the table record the running time on two different synthetic data respectively. "-" denotes the running time exceeds 100 seconds in the top table and 600 seconds in the bottom table.

7.2 Model Comparison: Tensor Trace Norm versus Matrix Trace Norm

In this section, we compare the behavior of matrix and tensor completion in practical applications. We have not obtained meaningful theoretical bounds for our tensor completion algorithms. Given a tensor \mathcal{T} with missing entries, we can unfold it along the ith mode into a matrix structure $\mathcal{T}_{(i)}$. Then, the missing value estimation problem can be formulated as a low rank matrix completion problem

$$\min_{X_{\Omega} = \mathcal{T}_{(i)\Omega}} : \|X\|_*. \tag{44}$$

We compare tensor completion and matrix completion on both the synthetic data and the real-world data and report results in Tables 2 and 3. We show an example slice of the MRI data in Fig. 5.

Results from these two tables indicate that the proposed low rank tensor completion algorithm outperforms the low rank matrix completion algorithm, especially when the number of missing entries is large.

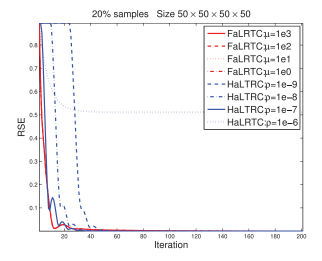
Next, we consider a natural idea for using the matrix completion technique for tensor completion. Without the unfolding operation, one can directly apply the matrix completion algorithm on each single slice of the tensor with missing entries. We call this as the slicewise matrix completion (S-MC). The comparisons on synthetic data and real data are reported in Table 4 and Fig. 4, which also show

the advantages of tensor completion. In particular, Table 4 indicates that tensor completion can almost perfectly recover the missing values while matrix completion performs rather poorly in this case.

The key reason why the proposed algorithm outperforms the matrix completion-based algorithms may be that tensor completion utilizes all information along all dimensions, while matrix completion only considers the constraints along two particular dimension.

7.3 Efficiency Comparison

This experiment concerns the efficiency of SiLRTC, FaLRTC, HaLRTC, and ADM-TR proposed in [14]. We compare them on the synthetic data and the results are summarized in Table 5. We report the running time of four algorithms when achieving the same recovery accuracy measured by RSE from 10^{-1} to 10^{-6} , respectively. We apply the efficient implementations of FaLRTC and HaLRTC by iteratively updating μ_i^k and ρ^k . We can observe from this table: 1) the larger the value of γ , the faster the SiLRTC algorithm, but a smaller value of γ can lead to a more accurate solution in the SiLRTC algorithm; 2) when three algorithms are configured to have the same recovery accuracy, the FaLRTC algorithm and the HaLRTC algorithm are much faster than the SiLRTC algorithm and the ADM-TR algorithm; 3) the FaLRTC algorithm is more efficient for obtaining the lower accuracy solutions while the HaLRTC algorithm is much more efficient for



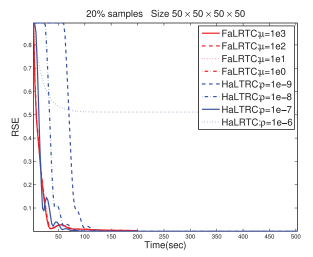


Fig. 6. The RSE comparison on the synthetic data. $I_1=I_2=I_3=I_4=50$ and $r_1=r_2=r_3=r_4=2$. All parameters are identical to the setting in the button part of Table 5.

obtaining higher accuracy solutions. Fig. 6 shows the RSE curves of the FaLRTC algorithm and the HaLRTC algorithm in terms of the number of iterations and the computation time. We can observe from this figure that the FaLRTC algorithm converges very fast at the beginning but takes some time to obtain a high accuracy. Thus, FaLRTC is a good choice if an accuracy lower than 10^{-2} is acceptable and the HaLRTC algorithm is recommended if a higher accuracy is desired. In practice, one might combine the

FaLRTC algorithm and the HaLRTC algorithm to improve the efficiency by running FaLRTC first and then HaLRTC.

7.4 Applications

In this section, we outline three potential application examples with three different types of data: images, videos, and reflectance data.

Images. Our algorithm can be used to estimate missing data in images and textures. For example, in Fig. 7, we show how missing pixels can be filled in a facade image. Note how our algorithm can propagate global structure even though a significant amount of information is missing.

Videos. The proposed algorithm may be used for video compression and video in-painting. The core idea of video compression is to remove individual pixels and to use tensor completion to recover the missing information. Similarly, a user can eliminate unwanted pixels in the data and use the proposed algorithm to compute alternative values for the removed pixels. See Fig. 8 for an example frame of a video.

Reflectance data. The BRDF is the "Bidirectional Reflectance Distribution Function." The BRDF specifies the reflectance of a target as a function of illumination direction and viewing direction and can be interpreted as a four mode tensor. BRDFs of real materials can be acquired by complex appearance acquisition systems that typically require taking photographs of the same object under different lighting conditions. As part of the acquisition process, data can be missing or be unreliable, such as in the MIT BRDF data set. We use tensor completion to estimate the missing entries in reflectance data. See Fig. 9 for an example. More results are shown in the video accompanying this paper.

8 Conclusion

In this paper, we extend low rank matrix completion to low rank tensor completion. We propose tensor completion based on a novel definition of the trace norm for tensors together with three convex optimization algorithms, SiLRTC, FaLRTC, and HaLRTC, to tackle the problem. The first algorithm, SiLRTC, is intuitive to understand and simple to implement. The latter two algorithms, FaLRTC and HaLRTC, are significantly faster than SiLRTC and use more advanced optimization techniques. Additionally, several heuristic algorithms are presented. The experiments show that the proposed algorithms are more stable and accurate in most cases, especially when the sample entries are very limited.

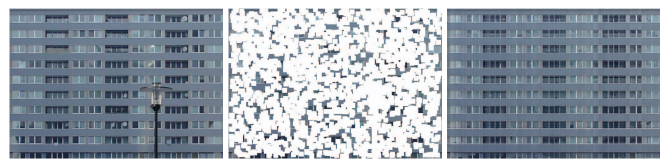


Fig. 7. Facade in-painting. The left image is the original image; we select the lamp and satellite dishes, together with a large set of randomly positioned squares, as the missing parts, shown in white in the middle image; the right image is the result of the proposed completion algorithm.







Fig. 8. Video completion. The left image (one frame of the video) is the original; we randomly select pixels for removal, shown in white in the middle image; the right image is the result of the proposed LTRC algorithm.







Fig. 9. The left image is a rendering of an original phong BRDF; we randomly select 90 percent of the pixels for removal, shown in white in the middle image; the right image is the result of the proposed SiLRTC algorithm.

Several application examples show the broad applicability of tensor completion in computer vision and graphics.

The proposed tensor completion algorithms assume that the data is of low rank. This may not be the case in certain applications. We plan to extend the theoretical results of Candès and Recht to the tensor case. We also plan to extend the proposed algorithms using techniques recently proposed in [8] and [48].

ACKNOWLEDGMENTS

This work was supported by US National Science Foundation (NSF) IIS-0812551, CCF-0811790, IIS-0953662, CCF-1025177, and NGA HM1582-08-1-0016. The authors thank the anonymous reviewers for pointing out ADMM and recent following work on tensor completion. They also thank S. Gandy for sharing the implementation of the ADM-TR algorithm.

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Ji Liu received the bachelor's degree in automation from the University of Science and Technology of China and the master's degree in computer science from Arizona State University, in 2005 and 2010, respectively. Currently, he is working toward the graduate degree in the Department of Computer Sciences at the University of Wisconsin–Madison. His research interests include optimization, machine learning, computer vision, and graphics. He won the KDD

best research paper award honorable mention in 2010.



Przemyslaw Musialski received the MSc degree from Bauhaus University Weimar, Germany, and the PhD degree from the Vienna University of Technology, Austria, in 2007 and 2010, respectively. From 2007 till 2010, he was with the VRVis Research Center in Vienna, where he was working on image processing and image-based urban modeling. In 2010, he continued this work at the Vienna University of Technology. Since 2011 he has been a post-

doctoral scholar at Arizona State University, where he is conducting research on image and texture processing.



Peter Wonka received the MS degree in urban planning and the doctorate degree in computer science from the Technical University of Vienna. Currently, he is with Arizona State University (ASU). Prior to coming to ASU, he was a postdoctoral researcher at the Georgia Institute of Technology for two years. His research interests include various topics in computer graphics, visualization, and image processing.



Jieping Ye received the PhD degree in computer science from the University of Minnesota, Twin Cities, in 2005. Currently, he is working as an associate professor in the Department of Computer Science and Engineering at Arizona State University (ASU). His research interests include machine learning, data mining, and biomedical informatics. He won the outstanding student paper award at ICML in 2004, the SCI Young Investigator of the Year Award at ASU in

2007, the SCI Researcher of the Year Award at ASU in 2009, the NSF CAREER Award in 2010, and the KDD best research paper award honorable mention in 2010 and 2011. He is a senior member of the IEEE.

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