**FLEXIFACT**: Scalable Flexible Factorization of Coupled Tensors on Hadoop

Alex Beutel*  
abeutel@cs.cmu.edu

Abhimanu Kumar *  
abhimank@cs.cmu.edu

Evangelos E. Papalexakis *  
epapalex@cs.cmu.edu

Partha Pratim Talukdar *  
ppt@cs.cmu.edu

Christos Faloutsos *  
christos@cs.cmu.edu

Eric P. Xing*  
epxing@cs.cmu.edu

Carnegie Mellon University, School of Computer Science

Abstract
Given multiple data sets of relational data that share a number of dimensions, how can we efficiently decompose our data into the latent factors? Factorization of a single matrix or tensor has attracted much attention, as, e.g., in the Netflix challenge, with users rating movies. However, we often have additional, side, information, like, e.g., demographic data about the users, in the Netflix example above. Incorporating the additional information leads to the coupled factorization problem. So far, it has been solved for relatively small datasets.

We provide a distributed, scalable method for decomposing matrices, tensors, and coupled data sets through stochastic gradient descent on a variety of objective functions. We offer the following contributions:

1. **Versatility**: Our algorithm can perform matrix, tensor, and coupled factorization, with flexible objective functions including the Frobenius norm, Frobenius norm with an $\ell_1$ induced sparsity, and non-negative factorization.
2. **Scalability**: FLEXIFACT scales to unprecedented sizes in both the data and model, with up to billions of parameters. FLEXIFACT runs on standard Hadoop.
3. **Convergence proofs**: We prove that FLEXIFACT converges, even with constraints like non-negativity. Moreover, we demonstrate this empirically.

In summary, our main contributions are:

1. **Versatility**: FLEXIFACT can operate under a wide spectrum of settings, including plain matrix factorization, tensor factorization, as well as coupled decompositions. Thus, FLEXIFACT includes several recent methods [9], [11], as special cases.
2. **Scalability**: FLEXIFACT scales very well both with the input size, as well as with the number of model parameters.
3. **Proof of convergence**: We prove that FLEXIFACT converges, even with constraints like non-negativity. Moreover, we demonstrate this empirically.
4. **Usability and Reproducibility**: Our implementation runs on stock Hadoop, as opposed to other recent methods [9]. We also open-source our code.
## Table 1: Feature Comparison of proposed FlexiFACT vs state of the art. (~ represents unknown or not directly applicable.) FlexiFACT contains existing state of the art as special cases.

| Data/Model                   | FlexiFACT | DSGD [9] | PSGD [21] | Matlab | GigaTensor [11] |
|------------------------------|-----------|---------|-----------|--------|-----------------|
| Matrix                       | ✓         | ✓       | ✓         | ~      | ✓               |
| Tensor                       | ✓         | ✓       | ✓         | ✓      | ✓               |
| Coupled Tensor/Matrix        | ✓         | ✓       | ✓         | ✓      | ✓               |

| Obj. Function                |           |         |           |        |                 |
|------------------------------|-----------|---------|-----------|--------|-----------------|
| Frobenius norm               | ✓         | ✓       | ✓         | ✓      | ✓               |
| Frobenius norm + ℓ₁ penalty | ✓         | ✓       | ✓         | ✓      | ✓               |
| Non-negativity constraints   | ✓         | ✓       | ✓         | ✓      | ✓               |
| Handles missing data         | ✓         | ✓       | ✓         | ✓      | ✓               |

| Scalability                  |           |         |           |        |                 |
|------------------------------|-----------|---------|-----------|--------|-----------------|
| in number of non-zeros       | ✓         | ~       | ✓         | ✓      |                 |
| in data dimensions           | ✓         | ~       | ✓         | ✓      |                 |
| in decomposition rank        | ✓         | ~       | ✓         | ✓      |                 |

| Proof of convergence         |           |         |           |        |                 |
|------------------------------|-----------|---------|-----------|--------|-----------------|
| Matrix Factorization         | ✓         | ✓       |           |        |                 |
| Tensor/Coupled Factorization | ✓         | ✓       |           |        |                 |
| Projections (ℓ₁ & non-negativity) | ✓         | ✓       |           |        |                 |

## Related Work and Background

### Matrix Factorization (MF): One of the most popular MF models is the Non-Negative Matrix Factorization [14]. Solutions range from alternating least squares (ALS) [20], to stochastic gradient descent (SGD) [7]. Gemulla et al. [9] made a breakthrough in scaling up MF: they propose a distributed version of Stochastic Gradient Descent (SGD) for MF which elegantly exploits the factorization structure of the problem, break the matrix into blocks and process them in parallel. Follow-up works [16, 19] have successfully extended this idea to the matrix completion problem using parallel stochastic updates. We build upon this insightful work, expanding on both the theory and implementation.

### Tensor Factorization. Tensors are multidimensional generalizations of matrices, widely applied in a variety of fields. See Kolda and Bader [12] for a comprehensive overview. The most popular Tensor Factorization is the so called Canonical Polyadic (CP) or PARAFAC decomposition [10]. PARAFAC with ℓ₁ constraints, which FlexiFACT is able to handle, is introduced in [15]. For in memory/single machine computations, the state of the art is the Tensor Toolbox [6], which provides an implementation of the ALS algorithm. Besides ALS there exist first order optimization techniques, such as [4]. In [15], the authors propose algorithms for incremental computation of the tensor decomposition, where the tensor is a stream. For large datasets residing on HDFS (Hadoop Distributed File System), the state of the art is GigaTensor [11] solves the PARAFAC decomposition using Alternating Least Squares (ALS).

### Coupled Factorizations. In this case, we have a tensor and a matrix, or two tensors, or two matrices, that share one dimension, as in the Facebook example of the introduction. Singh et al. [17] jointly factorize two related matrices for better decomposition and provide stochastic updates for this coupled MF optimization. For coupled matrix-tensor factorization, recent work [3] used first order optimization techniques.

## FlexiFACT Approach

### Table 2: Table of Symbols

| Symbol | Description |
|--------|-------------|
| X      | Data tensor |
| Y      | Data matrix |
| U, V, W| Factor matrices of X |
| U, A   | Factor matrices of Y |
| I × J × K| Dimensions of data tensor X |
| I × M  | Dimensions of data matrix Y |
| θ      | Any parameter in the objective |
| σ      | The parameter being updated |
| ∇      | Symbol for derivative |
| ηt     | Step size at iteration t |
| R      | Rank of decomposition |
| ◦      | Khatri Rao product |

As mentioned previously, we take on the problem of matrix, tensor and coupled factorization. In this section we will explain the variety of loss functions used in these tasks, the Stochastic Gradient Descent (SGD) update rules, and our partitioning scheme allowing for distribution of the SGD work. Although much of our description of the matrix factorization work is similar to [9], we will explain it here for completeness and clarity. Before we begin, it is important to clarify our notation. We will use capital boldface script characters.
like $\mathbf{X}$ to denote a tensor, capital boldface non-script characters e.g. $\mathbf{Y}$ to denote a matrix, and lowercase boldface character, e.g. $\mathbf{y}$, to denote a vector. $\mathbf{X}_{i,j,k}$ denotes the scalar in the $(i,j,k)$ position of the tensor $\mathbf{X}$, $\mathbf{Y}_{i,j}$ denotes the scalar in the $(i,j)$ position of matrix $\mathbf{Y}$, and $y_t$ denotes the scalar in the $i$th position of vector $\mathbf{y}$. We use $\mathbf{Y}_{i,*}$ to denote the vector of scalars $\mathbf{Y}_{i,j}$ for all $j$. Additionally, with a slight overloading of notation for simplicity and because our matrices and tensors may only have a small percentage of observed values, we say that $(i,j,k) \in \mathbf{X}$ if $\mathbf{X}_{i,j,k}$ is observed. A list of our commonly used symbols can be seen in Table 2.

3.1 Optimization Objectives We begin by explaining how we make use of objective functions for simpler cases like the Frobenius norm of matrices before expanding to more complex objectives.

Matrix Factorization For matrix factorization we would like to approximate our $I \times J$ data matrix $\mathbf{X}$ by $\mathbf{U}\mathbf{V}^T$, where $\mathbf{U}$ is of size $I \times R$ and $\mathbf{V}$ is of size $J \times R$. Therefore, we can have a loss function using the Frobenius norm as follows:

$$L(\mathbf{U}, \mathbf{V}) = \|\mathbf{X} - \mathbf{U}\mathbf{V}^T\|_F^2 = \sum_{i,j \in \mathbf{X}} L_{\mathbf{X}_{i,j}}(\mathbf{U}, \mathbf{V})$$

where $L_{\mathbf{X}_{i,j}}(\mathbf{U}, \mathbf{V}) = (\mathbf{X}_{i,j} - \sum_{r=1}^R \mathbf{U}_{i,r}\mathbf{V}_{j,r})^2$. As seen above, we divide our loss function into its component pieces $L_{\mathbf{X}_{i,j}}$ based on each observed point $\mathbf{X}_{i,j}$. This is necessary to use stochastic gradient descent.

Tensor Factorization For tensor factorization we would like to approximate our $I \times J \times K$ tensor $\mathbf{X}$ by a Khatri-Rao product $\sum_{r=1}^R \mathbf{U}_{i,r}\circ\mathbf{V}_{j,r}\circ\mathbf{W}_{k,r}$, where $\mathbf{U}$ is of size $I \times R$, $\mathbf{V}$ is of size $J \times R$ and $\mathbf{W}$ is of size $K \times R$ and we are performing an Khatri Rao product between these three matrices. We can analyze the loss in a few different ways. Following the standard Frobenius norm, as is common in PARAFAC, the loss is:

$$L(\mathbf{U}, \mathbf{V}, \mathbf{W}) = \|\mathbf{X} - \sum_{r=1}^R \mathbf{U}_{i,r}\circ\mathbf{V}_{j,r}\circ\mathbf{W}_{k,r}\|_F^2 = \sum_{(i,j,k) \in \mathbf{X}} L_{\mathbf{X}_{i,j,k}}(\mathbf{U}, \mathbf{V}, \mathbf{W})$$

where

$$L_{\mathbf{X}_{i,j,k}}(\mathbf{U}, \mathbf{V}, \mathbf{W}) = (\mathbf{X}_{i,j,k} - \sum_{r=1}^R \mathbf{U}_{i,r}\mathbf{V}_{j,r}\mathbf{W}_{k,r})^2.$$

Similarly we can induce sparsity in our parameter space with an $\ell_1$ penalty:

$$L(\mathbf{U}, \mathbf{V}, \mathbf{W}, \mathbf{A}) = \sum_{(i,j,k) \in \mathbf{X}} L_{\mathbf{X}_{i,j,k}}(\mathbf{U}, \mathbf{V}, \mathbf{W}) + \sum_{(i,j) \in \mathbf{Y}} L_{\mathbf{Y}_{i,j}}(\mathbf{U}, \mathbf{A})$$

Table 3 denotes the loss objectives for different coupled cases. For each of these we use SGD to minimize our loss and thus approximate our data.

3.2 SGD Updates For SGD we perform updates to our parameters $\mathbf{U}, \mathbf{V}, \mathbf{W}, \mathbf{A}$, which we will collectively refer to as $\Theta$ matrix whereas $\theta$ are the individual components of the matrix. This definition of $\Theta$ and $\theta$ will come in handy for parameter updates based on the gradient at individual data points. E.g. the update for tensor $\mathbf{X}$ are:

$$\theta^{(t+1)} = \theta^{(t)} - \eta_t \nabla L(\mathbf{X}_{i,j,k}^{(t)}(\theta^{(t)}))$$

For these update rules, we list below the differentials for each component $\sigma$ of $\theta$ where $(\nabla L_{\mathbf{X}_{i,j,k}}(\theta))_\sigma = \frac{\partial L_{\mathbf{X}_{i,j,k}}}{\partial \sigma}$:

$$(\nabla L_{\mathbf{X}_{i,j,k}}(\theta))_\sigma = \begin{cases} -2(\mathbf{X}_{i,j,k} - \sum_{r=1}^R \mathbf{U}_{i,r}\mathbf{V}_{j,r}\mathbf{W}_{k,r}) \mathbf{V}_{j,t}\mathbf{W}_{k,t} & \text{if } \sigma = \mathbf{V}_{j,t} \\ 0 & \text{if } \sigma = \mathbf{U}_{i,t}, \mathbf{V}_{j,t} \neq \mathbf{W}_{k,t} \end{cases}$$

similarly for $\sigma = \mathbf{V}_{j,t}$ or $\sigma = \mathbf{W}_{k,t}$. From this we observe that SGD update for $\mathbf{U}_{i,t}$ at a particular entry $\mathbf{X}_{i,j,k}$ (for a tensor $\mathbf{X}$) depends only on previous
Formulation

\[ \sum_{(i,j,k) \in X} \mathcal{L}_{X_{i,j,k}}(U, V, W) + \sum_{(i,j) \in Y} \mathcal{L}_{Y_{i,j}}(U, A) + \lambda(||U||_1 + ||V||_1 + ||W||_1 + ||A||_1) \]

Table 3: Table of Objective Functions. \( \Theta \) denotes any of the factors \( U, V, W \) or \( A \).

![Figure 1: Dividing the paired matrix and tensor into blocks such that no two of them share any row or a column or a third dimension in case of tensor.](image)

We first prove that two blocks in a stratum are independent. Figure 1 is a pictorial representation of the way we segment our simple matrix or a coupled tensor/matrix to enable parallelization. In order to run SGD on the blocks in parallel, we divide them such that no two blocks share common rows or columns. To be more precise, we say that a point \( x \in Z_b \) is the coordinates in the data, such as \( x = (x_i, x_j, x_k) \in X \). Two blocks \( Z_b \) and \( Z_{b'} \) are non-overlapping if for all \( x \in Z_b \) and \( x' \in Z_{b'} \), \( x_i \neq x'_i \) and \( x_j \neq x'_j \) and \( x_k \neq x'_k \). (We will prove later that this allows us to run the blocks in parallel.) We see that in the division shown in Figure 1 there are no two blocks share common rows or columns. More interestingly, we note that in Figure (c) blocks in the tensor \( X \) and the matrix \( Y \) share coordinates in the \( i \) dimension, and as a result, data points in the same \( i \) range must be in the same block across both data sets.

3.3 Blocking for Parallelization

Given this understanding of our optimization objective and SGD update rules, we would like to segment our data in such a way that certain blocks \( Z_b \) can be run in parallel, where we define \( Z_b \subseteq X \). Figure 1 is a pictorial representation of the way we segment our simple matrix or a coupled tensor/matrix to enable parallelization. In order to run SGD on the blocks in parallel, we divide them such that no two blocks share common rows or columns. To be more precise, we say that a point \( x \in Z_b \) is the coordinates in the data, such as \( x = (x_i, x_j, x_k) \in X \). Two blocks \( Z_b \) and \( Z_{b'} \) are non-overlapping if for all \( x \in Z_b \) and \( x' \in Z_{b'} \), \( x_i \neq x'_i \) and \( x_j \neq x'_j \) and \( x_k \neq x'_k \). (We will prove later that this allows us to run the blocks in parallel.) We see that in the division shown in Figure 1 there are no two blocks share common rows or columns. More interestingly, we note that in Figure (c) blocks in the tensor \( X \) and the matrix \( Y \) share coordinates in the \( i \) dimension, and as a result, data points in the same \( i \) range must be in the same block across both data sets.

Given this intuition, we provide a detailed description of our partition function. We call one set of independent blocks a stratum, and we denote the number of blocks in each stratum by \( d \). In order to cover all regions of \( X \), we need multiple strata. For a matrix we require \( d \) strata, and for tensors we require \( d^2 \) strata. For a stratum \( s \) we have blocks \( Z_i(s) \) for \( i = 0 \ldots d - 1 \). Each block \( Z = (b_i, b_j, b_k) \) where \( b_i, b_j, b_k \) are ranges in \( I, J, \) and \( K: b_i = (i[I/d], (i + 1)[I/d]), b_j = (j[J/d], (j + 1)[J/d]), b_k = (k[K/d], (k + 1)[K/d]) \). With this we define the blocks for stratum \( s \) as

\[ Z_i(s) = (b_i, b_{j,s}, b_{k,s,i}) \]

\[ j_{s,i} = (j + s) \mod d \]

\[ k_{s,i} = [(j + s)/d] \mod d \]

for \( i = 0 \ldots d - 1 \).

In our algorithm, we run the strata sequentially, but for each stratum we run SGD on the blocks in parallel. We consider running SGD on one stratum to be a subepoch in our algorithm, and running it on all strata an epoch. (Note, the order in which you run the strata does not matter, as long as they are each run once per epoch.) We can do this repeatedly, iteratively updating our parameters \( \theta \), until the algorithm converges. A more formal write up of the distributed stochastic gradient algorithm for a tensor (which can easily be generalized to matrices and coupled factorizations) is shown in Algorithm 1. We next offer a proof that this converges appropriately.

4 Proof of convergence with projections

The FlexiFACT approach is described in Algorithm 1. We first prove that two blocks in a stratum are interchangeable. We use this to prove that sequence of strata are a regenerative process, defined later in this section.
We use this to prove that our FLEXIFACT approach for Tensor and coupled case converges. Our generic constrained loss function for a tensor case is

\[
\mathcal{L} = L(U, V, W) + \lambda_u\|U\|_1 + \lambda_v\|V\|_1 + \lambda_w\|W\|_1
\]

(4.10) \[s.t.\] \(U_{i,r}, V_{j,r}, W_{k,r} \geq 0.\]

In the above projected loss equation 4.10 the parameter is always in a set, \(P\), constrained by the \(l\) and non-negativity constraints. The set \(P\) is a hyper-rectangle defined as \(\exists \, a_i < b_i, i = 1, \ldots, r\), such that \(P = \{\theta: a_i \leq \theta_i \leq b_i\}\) where \(a_i, b_i \in (-\infty, \infty)\). Here \(\theta\) is a parameter to be updated as defined in previous section (equation 3.3). The gradient based on equation 4.10 is:

\[
\nabla_{\theta} \mathcal{L} = \nabla_{\theta} L + p(\theta), \quad p(\theta) \in C(\theta)
\]

(4.11) where \(\theta\) is defined in Table 2, and \(L\) and \(\mathcal{L}\) are defined in equation 3.2. Function \(p()\) is the projection or constraint term of the gradient. The set \(C(\theta)\) is the union of the subgradients at \(\theta\). When \(\theta\) in interior of \(P, C(\theta)\) contains only the zero elements and contains the convex cone generated by the subgradients at \(\theta\) when \(\theta \in \partial P\), boundary of \(P\).

**Definition 4.1.** Two blocks \(Z_i\) and \(Z_i'\) in a given stratum are independent if for each \(x \in Z_i\) and \(x' \in Z_i'\) we have

\[
\nabla L_x(\theta) = \nabla L_x(\theta - \eta\nabla L_{x'}(\theta))
\]

and

\[
\nabla L_{x'}(\theta) = \nabla L_{x'}(\theta - \eta\nabla L_x(\theta))
\]

where \(\nabla L_x(\theta)\) is the partial differential of \(L_x\) w.r.t. \(\theta\) and \(\theta\) is the parameter we are updating.

**Theorem 4.1.** If blocks \(Z_i\) and \(Z_i'\) in a given stratum \(S_n\) are non-overlapping then they are independent (as defined previously in Definition 4.1).

**Proof.** For any two points \(x \in Z_i\) and \(x' \in Z_i'\) their rows or columns or any other coordinate do not overlap. From equation (3.4) we see that \(x\) does not modify \(\theta\) in positions for which \(i \neq i_1, j \neq j_1\) and \(k \neq k_1\). Therefore, because \(x\) and \(x'\) are not equal in any dimension, an update from \(\nabla L_{x'}\) will update different values than \(\nabla L_x\), where \(\nabla L_x\) is the gradient at point \(x\).

Additionally from equation (3.4) we see that any updates on \(\nabla L_{x'}\) only use values from \(U_{z_1, r}, A_{z_1}, \) and \(B_{z_2, r}\), and thus do not use any values that would be updated by \(\nabla L_x\). Because updates from \(x\) only effect parameters in \(x\)'s coordinates, updates from \(x\) are only based on parameters in \(x\)'s coordinates, and \(x\) and \(x'\) have no overlapping coordinates, we know that

\[
\nabla L_x(\theta) = \nabla L_x(\theta - \eta\nabla L_{x'}(\theta))
\]

and

\[
\nabla L_{x'}(\theta) = \nabla L_{x'}(\theta - \eta\nabla L_x(\theta))
\]

Therefore, \(Z_i\) and \(Z_i'\) are independent. By a similar argument

\[
\nabla \mathcal{L}_x(\theta) = \nabla \mathcal{L}_x(\theta - \eta\nabla \mathcal{L}_{x'}(\theta))
\]

and

\[
\nabla \mathcal{L}_{x'}(\theta) = \nabla \mathcal{L}_{x'}(\theta - \eta\nabla \mathcal{L}_x(\theta))
\]

\[\blacksquare\]

**Definition 4.2.** A process \(P(t), t \geq 0\) is regenerative if there exist time points \(0 \leq T_0 < T_1 < T_2 < \ldots\) such that the the remainder of the process after \(T_k\) \(P(T_k + t): t \geq 0\), for \(k \geq 1\) : (a) has the same distribution as the remainder of the process after \(T_0\), and (b) process \(P(T_0 + t): t \geq 0\) is independent of the process prior to \(T_k\) \(P(t): 0 \leq t < T_k\).

In other words a stochastic process with certain time points such that from a probabilistic view the process restarts itself at these time points is called a regenerative process. Intuitively this means a regenerative process can be split into i.i.d. cycles 35.

Based on equation 4.10 the projected sgd updates can be written as:

\[
\theta^{t+1} = \Pi_{\delta P}(\theta(t) + \eta \nabla L_x(\theta(t)))
\]

where \(\Pi_{\delta P}()\) is the projection of the updated gradient with respect to the original loss \(L(U,V,W)\). The projection step can be further broken up into

\[
\theta^{t+1} = \theta(t) + \eta \nabla L^*_x(\theta(t)) + \eta p(\theta(t)) \quad (using \ 4.11)
\]

\[
\theta(t) + \eta \nabla L^0(\theta(t)) + \eta \delta M + \eta \beta \eta \eta p(\theta(t))
\]

where \(L^*_x(\theta(t))\) is the loss function at stratum \(s_i\) at a point \(x\) in iteration \(t\) given parameter value in
previous iteration $\theta^{(t)}$. $\nabla L^0(\theta^{(t)})$ is the exact gradient in iteration $t$ given previous parameter value $\theta^{(t)}$. And

$$\delta M_t = \nabla L^0_x(\theta^{(t)}) - \nabla L^0(\theta^{(t)}) - \beta_t$$

where $\beta_t$ is the “error” before projection i.e. the error by which the update is outside $P$.

To prove the convergence of the method we define the following conditions, similar to the ones defined in [13, 9]:

**Condition 1.** $\nabla L^0(\theta)$ is continuous.

**Condition 2.** $\nabla L^0(\theta^{(t)})$ is bounded in second moment: $E[(\nabla L^0(\theta^{(t)}))^2] < \infty$ for all $\theta$.

**Condition 3.** The squared sum of the step sizes $\eta_t$ is bounded i.e. $\sum_t \eta_t^2 < \infty$.

**Condition 4.** The noise is martingale difference: $E[\delta M_{t+1} | \delta M_t, i \leq t] = \delta M_t$.

**Condition 5.** $E[\eta t \beta_i] < \infty$ with probability 1. Note that this is a condition on step-size. It implicitly says that the projection must not wander off infinitely outside the set $P$ over the iterations.

**Theorem 4.2.** The distributed SGD algorithm for tensor decomposition with projections, as presented in algorithm 2, converges.

**Proof.** The primary equations being updated each time in our iterations is equation 4.14. Rewriting it here we have:

$$\theta^{t+1} = \theta^{(t)} + \eta_t \nabla L^0(\theta^{(t)}) + \eta_t \delta M_t + \eta_t \beta_t$$

From theorem 4.1 we can see that the individual blocks in a given stratum are independent of each other’s updates and are interchangeable. We can also observe from Algorithm 1 that every stratum out of $d$ strata is picked exactly once in one cycle i.e. one epoch (outer while loop). Moreover two different cycles of strata i.e. iterations of the while loop are identical and independent. In other words the while loop forms an i.i.d cycle, and thus a regenerative process. The time-period of cycles is finite and bounded consequently that of the regenerative process too. Besides given all the conditions 1 to 5 as defined above, we have

$$\sum_{i=0}^{\kappa(1+t_s)-1} \left( \eta_i \delta M_i + \eta_i \beta_i \right) \rightarrow 0$$

for any arbitrary $\kappa$. The proof is similar to [13] and is valid due to the fact that noise is a martingale difference sequence and $\eta_i \delta M_i$ and $\eta_i \beta_i$ are an equicontinuous sequence ([13] Theorem 2.1, part 1, chapter 5; [9] follows a similar proof up to this point). We can now use this to analyze the updated with a projected loss. We find that equation 4.14 has the same set of stable points as

$$\theta^{t+1} = \theta^{(t)} + \eta_t \nabla L^0(\theta^{(t)}) + \eta_t \beta_t$$

Now we show that equation 4.18 converges. Through few algebraic manipulations it can be verified that the projection functions $p(\cdot)$ we have, $f_1$ soft threshold and non-negativity constraint project, are lipschitz continuous. Following the arguments of [13] (theorems 2.1, part 2) and with the assumption that updates $\theta^{(t)}$ are bounded (follow from the conditions 1 to 5 assumed earlier), equation 4.18 converges to a set of stationary points.

**5 MapReduce Implementation of FlexiFaCT**

Along with our theoretical analysis, we implemented our algorithm within the MapReduce framework [8]. To do this we used the open source Hadoop [11] version of MapReduce. The challenge is to turn the factorization problem into map() and reduce() functions, that Hadoop is designed to handle.

In our implementation we pass the data matrix or tensor as input to the mappers in the form $(i, j, k, X_{i,j,k})$. We also store our current parameters $\theta^{(s)}$, which could include $U$, $V$, $W$, and $A$ on the Hadoop File System (HDFS).

**FlexiFaCT Mapper:** Our Mapper function, is shown in Algorithm 2. It splits the data into the appropriate blocks and determines the order they should be processed in within each reducer. We also overload the default Hadoop partitioner, which typically just partitions on unique KEY values, and now partition only on $b_k$ so that each reducer represents a unique set of $i$ in $I$ or a unique set of rows in $U$. We additionally override the default Comparator, allowing us to sort our (KEY,VALUE) pairs within each reducer by the subepoch term calculated in the Mapper. We see here the Mapper is quite simple, the calculation of the blocks and the order within each reducer captures our partition function that allows us to perform SGD in this distributed fashion.

**Algorithm 2: FlexiFaCT Mapper (for tensor)**

**Input:** $I, J, K, d$

1: for all $(i,j,k, X_{i,j,k})$ do
2: $b_i = \lfloor \frac{i}{d} \rfloor$, $b_j = \lfloor \frac{i}{d} \rfloor$, $b_k = \lfloor \frac{k}{d} \rfloor$ {Get block index}
3: subepoch = $d \times ((b_k - b_i + d) \mod d) + ((b_j - b_i + d) \mod d)$
4: emit $((b_i, b_j, b_k, subepoch), (i, j, k, X_{i,j,k}))$
5: end for
θ values to disk (for another reducer to retrieve) and a new subepoch is reached, it must write its updated
only stores the components of θ it to HDFS. We then use the updated parameters
update the full parameter space
subsequent subepoch. (As before we use s to denote
the subepochs.) The reducer iterates over the points
in order, each time updating \( \theta^{(s)} \). Each reducer
only stores the components of \( \theta \) that correspond to its
current block in the current stratum. As such, when
a new subepoch is reached, it must write its updated
\( \theta \) values to disk (for another reducer to retrieve) and
read the most current \( \theta \) values for its next block in the
subsequent subepoch.

We run the MapReduce jobs iteratively. Each
MapReduce job is one epoch using all points in \( \mathbf{X} \) to
update the full parameter space \( \theta \) and ultimately to save
it to HDFS. We then use the updated parameters \( \theta \) in
the subsequent epoch (another run of the MapReduce
algorithm). We do this for a constant number of steps
or until the algorithm converges.

Reproducibility and Usability: This is a high
level overview of our implementation, but captures
the general method we use to both distribute our
work and optimize our speed within the MapReduce
framework. While this is not the typical way Hadoop
is programmed, it requires no modification of the Hadoop
framework and can be run on any standard Hadoop
cluster. Our code is open-sourced, and available at
http://alexbeutel.com/l/flexifact. It can run for
all of the data types and loss functions described in this
paper.

6 Experiments

6.1 Performance Evaluation In order to assess
how scalable and fast FlexiFaCT is, we conducted a
series of experiments in order to measure the running
time of FlexiFaCT with respect to 1) increasing
number of data points, 2) increasing dimensions of the
data and thus model, and 3) increasing rank of the
factorization. The first aspect has to do with scalability
in terms of data size, whereas the two latter aspects
refer to scalability with respect to parameter space size;
FlexiFaCT is able, as we demonstrate in the following
experiments, to scale easily in all three aspects. As a
baseline for tensor decomposition, we use GigaTensor
[11]. We also compared against PSGD [21], however, the
solutions obtained achieved much worse RMSE, and the
algorithm was not able to scale for very large number of
parameters (either rank or dimensions).

FlexiFaCT was implemented in Java, with
Hadoop 0.20.1 [8, 1]. We ran the experiments on the
OCC-Y cluster[4]. For the sake of experimentation, we
created a series of synthetic datasets wherein we were
able to control the three aspects we were testing: data
size, data dimensions, and rank. Additionally, we vali-
date our method’s correctness by presenting experimen-
tal evidence (on top of our theoretical guarantees), for
monotone convergence. In all cases, number of reducers
was constant and equal to 24.

Synthetic Data Generation To generate data we
first generate randomly matrix factors \( \mathbf{U}, \mathbf{V}, \mathbf{W} \) of
the specified dimension \( D \) (where \( I = J = K = D \))
and rank \( R = 30 \). We then randomly select data points
\((i, j, k)\) and add their value \( \mathbf{U}_{i,\ast} \odot \mathbf{V}_{j,\ast} \odot \mathbf{W}_{k,\ast} \) to the
data set. We do this until we have the desired number
of data points for each dataset. Unless otherwise stated,
we set \( D = 1 \) million and the number of data points to
10 million.

6.2 Scalability We now test FlexiFaCT on all
three types of scalability to demonstrate that it scales
in all dimensions and to unprecedented sizes.

Rank Scalability In testing the scalability with
respect to rank, we ran FlexiFaCT, GigaTensor, and
PSGD on a tensor and varied the rank from 25 up to
1000. Figure [2a] shows the running time for Flexi-
FACT, both for tensor and coupled factorizations, as
the rank (i.e. one of the parameter dimensions) in-
creases. We can see that FlexiFaCT scales linearly
as the rank of the factorization increases, having similar
timing behaviour both for plain tensor and coupled

\[\text{http://opencloudconsortium.org/tag/occ-y}\]
factorizations. GigaTensor, on the other hand, due to the fact that it is inverting an $R \times R$ matrix, locally, at every iteration slows down significantly for large ranks. We were unable to test GigaTensor on ranks larger than 100 due to the slow down. Although barely visible in the plot, we also ran PSGD on the data. However, it too couldn’t scale in rank due to the size of the parameter space. For rank above 50 the factor matrices could no longer fit in memory on each machine and thus PSGD could not run. This demonstrates FlexiFACT’s unique scalability in the parameter size. Additionally, we note that with $R = 1000$ and $D = 1$ million, the coupled FlexiFACT factorization scales to a total parameter space of 4 billion parameters.

**Data Dimensions Scalability** To test more directly the scalability as the dimension $D$ of the data tensor grows, we created a variety of tensors with varying dimension from 10,000 to 10 million. We decompose each tensor with $R = 50$. When testing the coupled FlexiFACT decomposition, we add an additional coupled matrix with 100,000 data points and the same dimensionality as the main tensor.

In Figure 2(b) we show how coupled factorization using FlexiFACT scales, as the dimensions of the data increase. We observe that FlexiFACT runs much faster than the baseline, GigaTensor. A likely explanation for the degree to which FlexiFACT is faster than GigaTensor is that FlexiFACT only focuses on the observed data points, where GigaTensor has to convert unobserved data points to zeros, thus slowing down the computation. We can see that FlexiFACT has a very smooth behaviour, scaling linearly with the dimension size. Again, when we are performing the coupled decomposition, we see that as the dimension scales our total parameter space reaches 2 billion parameters.

**Data Size Scalability** Last, for data scalability, we vary the number of observed data points from 1 million to 10 million. Figure 2(c) shows FlexiFACT’s running time as a function of the data size, i.e. the number of observations. We can see that FlexiFACT has, again, very smooth behaviour, and scales linearly with the number of observed elements. Again, we are significantly faster than GigaTensor, though the degree of difference is likely because it is must make unobserved points zeros for it to run.

**6.3 Correctness & Monotone Convergence** Besides speedup, we experimentally validate that FlexiFACT indeed decreases monotonically the objective function that it is minimizing. To test this we run on a small synthetic data set with $D = 10,000$ and 10 million data points, making the dataset very dense. We run a factorization with $R = 50$ using our implementation of PSGD and FlexiFACT with both $\ell_1$ sparsity and non-
negativity constraints. We then monitor the root mean squared error (RMSE).

Figure 5 shows that FLEXIFACT decreases the RMSE as expected, and at a much quicker pace than PSGD. It is important to note that the slow convergence of PSGD is because the problem (tensor factorization) is not convex, and thus Zinkevich et al. do not claim that their method works on such problems. However, we use PSGD as a comparison because it is not possible to track the RMSE with GigaTensor and thus PSGD is the closest competitor.

7 Conclusion

In this work we have introduced FLEXIFACT, a highly flexible, efficient, and scalable factorization tool. Our main contributions are

1. Versatility: FLEXIFACT, can operate under numerous factorization scenarios, including matrix, tensor, and coupled factorization as well as with non-negativity and sparsity constraints.

2. Scalability: FLEXIFACT scales very well with the input size, as well as with the number of parameters.

3. Proof of convergence: We provide theoretical guarantees for the convergence of FLEXIFACT, even in the presence of constraints.

4. Reproducibility and Usability: Our implementation works on stock Hadoop; furthermore, we ensure the reproducibility and outreach of FLEXIFACT by making our implementation publicly available.

Acknowledgements

Research was funded by grants NSF IIS-1247489, NSF IIS/HCC Grant #1302522, NSF Graduate Research Fellowship (Grant No. DGE-1252522), Defense Threat Reduction Agency under contract No. HDTRA1-10-1-0120, ARL under Cooperative Agreement Number W911NF-09-2-0053, and DARPA FA8750-13-20005. We also would like to thank the CMU Parallel Data Laboratory OpenCloud for providing infrastructure for experiments; this research was funded, in part, by the NSF under award CCF-1019104, and the Graduate Research Fellowship (Grant No. DGE-1252522), Defense Threat Reduction Agency under contract No. HDTRA1-10-1-0120, ARL under Cooperative Agreement Number W911NF-09-2-0053, and DARPA FA8750-13-20005. We also would like to thank the Open Cloud Consortium (OCC) and the Open Science Data Cloud (OSDC) for the use of resources on the OCC-Y Hadoop cluster, which was donated to the OCC by Yahoo! Any opinions, findings, and conclusions or recommendations expressed in this material are those of the author(s) and do not necessarily reflect the views of the funding parties. The U.S. Government is authorized to reproduce and distribute reprints for Government purposes notwithstanding any copyright notation here on.

References

[1] Apache Hadoop. http://hadoop.apache.org/ 2012.
[2] The matlab cmtf toolbox. http://www.models.life.ku.dk/joda/CMTF_Toolbox 2013.
[3] E. Acar, T. G. Kolda, and D. M. Dunlavy. All-at-once Optimization for Coupled Matrix and Tensor Factorizations. ArXiv e-prints, May 2011.
[4] Evrim Acar, Daniel M. Dunlavy, Tamara G. Kolda, and Morten Morup. Scalable tensor factorizations with missing data. In SDM, pages 701–712, 2010.
[5] S. Asmussen. Applied Probability and Queues. Wiley, 1987.
[6] B.W. Bader and T.G. Kolda. Matlab tensor toolbox version 2.2. Albuquerque, NM, USA: Sandia National Laboratories, 2007.
[7] Léon Bottou. Stochastic learning. In Olivier Bousquet and Ulrike von Luxburg, editors, Advanced Lectures on Machine Learning, Lecture Notes in Artificial Intelligence, LNAI 3176, pages 146–168. Springer Verlag, Berlin, 2004.
[8] Jeffrey Dean and Sanjay Ghemawat. Mapreduce: Simplified data processing on large clusters. OSDI, December 2004.
[9] Rainer Gemulla, Erik Nijkamp, Peter J. Haas, and Yannis Sismanis. Large-scale matrix factorization with distributed stochastic gradient descent. In ACM SIGKDD, pages 69–77, New York, NY, USA, 2011. ACM.
[10] R.A. Harshman. Foundations of the parafac procedure: Models and conditions for an “explanatory” multimodal factor analysis. 1970.
[11] Kang, E. Papalexakis, A. Harpale, and C. Faloutsos. Gigatensor: scaling tensor analysis up by 100 times-algorithms and discoveries. In ACM SIGKDD, pages 316–324. ACM, 2012.
[12] T.G. Kolda and B.W. Bader. From k-means to higher-way co-clustering: Multilinear decomposition with sparse latent factors. IEEE TSP, 61(2):493–506, 2013.
[13] B. W. Bader and T. G. Kolda. Matlab tensor toolbox version 2.2. Albuquerque, NM, USA: Sandia National Laboratories, 2007.
[14] E. Acar, T. G. Kolda, and D. M. Dunlavy. All-at-once Optimization for Coupled Matrix and Tensor Factorizations. ArXiv e-prints, May 2011.
[15] H. Kushner and G. Yin. Stochastic Approximation. SIAM review, 51(3), 2009.
[16] T.G. Kolda and B.W. Bader. From k-means to higher-way co-clustering: Multilinear decomposition with sparse latent factors. IEEE TSP, 61(2):493–506, 2013.
[17] A. Harpale, and C. Faloutsos. Gigatensor: scaling tensor analysis up by 100 times-algorithms and discoveries. In ACM SIGKDD, pages 69–77, New York, NY, USA, 2011. ACM.
[18] B. W. Bader and T. G. Kolda. Matlab tensor toolbox version 2.2. Albuquerque, NM, USA: Sandia National Laboratories, 2007.
[19] B. W. Bader and T. G. Kolda. Matlab tensor toolbox version 2.2. Albuquerque, NM, USA: Sandia National Laboratories, 2007.
[20] B. W. Bader and T. G. Kolda. Matlab tensor toolbox version 2.2. Albuquerque, NM, USA: Sandia National Laboratories, 2007.
[21] B. W. Bader and T. G. Kolda. Matlab tensor toolbox version 2.2. Albuquerque, NM, USA: Sandia National Laboratories, 2007.
[22] B. W. Bader and T. G. Kolda. Matlab tensor toolbox version 2.2. Albuquerque, NM, USA: Sandia National Laboratories, 2007.