Algorithms and Comparisons of Nonnegative Matrix Factorizations with Volume Regularization for Hyperspectral Unmixing

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Abstract—In this work, we consider nonnegative matrix factorization (NMF) with a regularization that promotes small volume of the convex hull spanned by the basis matrix. We present highly efficient algorithms for three different volume regularizers, and compare them on endmember recovery in hyperspectral unmixing. The NMF algorithms developed in this work are shown to outperform the state-of-the-art volume-regularized NMF methods, and produce meaningful decompositions on real-world hyperspectral images in situations where endmembers are highly mixed (no pure pixels). Furthermore, our extensive numerical experiments show that when the data is less separable and/or contains more endmembers, the regularizer based on the determinant of the Gramian performs the best results in most cases. For data that is less separable and/or contains more endmembers, the regularizer based on the determinant of the Gramian performs best in general.

Index Terms—nonnegative matrix factorization, volume regularization, hyperspectral unmixing, blind source separation

I. INTRODUCTION

Non-negative Matrix Factorization (NMF) is the following problem: given a matrix $X \in \mathbb{R}^{m \times n}$ and an integer $r$, find two matrices $W \in \mathbb{R}^{m \times r}$ and $H \in \mathbb{R}^{r \times n}$ such that

$$X \approx WH.$$ (1)

NMF has many applications; see, e.g., [4], [9], [6] and the references therein. In this work, we focus on blind hyperspectral unmixing (HU) [2] that aims at recovering the spectral signatures of the pure materials (called endmembers, represented as the columns of $W$) and their abundances in each pixel (represented as the columns of $H$) within a hyperspectral image $X$ where each column of $X$ is the spectral signature of a pixel; see Section II for more details. In order to tackle HU, we will use volume-regularized NMF (VRNMF) which can be formulated as follows

$$\min_{W,H} f(W,H;X) + \lambda V(W)$$

subject to $W \geq 0$, $H \geq 0$, $H^\top 1_r \leq 1_n$, (2)

where $1_n$ denotes the vector of ones of length $n$ and $W \geq 0$ indicates that $W$ is component-wise non-negative. The regularization parameter $\lambda \geq 0$ controls the trade off between the data fitting term $f(W,H;X)$ and the volume regularizer $V(W)$. The constraint $H^\top 1_r \leq 1_n$ is a relaxation of the sum-to-one constraint $H^\top 1_r = 1_n$ which requires that the abundances of the endmembers in each pixel sum to one. Because VRNMF minimizes the volume of $W$, the constraint $H^\top 1_r \leq 1_n$ will tend to be active for most pixels. It has been shown previously that this relaxation works better in practice as it allows to take into account for example different illumination conditions within the hyperspectral image; see for example [8]. The reason to consider VRNMF has a long history in blind HU and is motivated by geometric insights: the columns of $W$ (the endmembers) are the vertices of a convex hull that contains the data points. In the absence of pure pixels, that is, pixels containing a single endmember, minimizing the volume of the columns of $W$ allows to recover these endmembers under mild conditions; see Figure 1 for an illustration, and Section II for more details.

In this paper, we will consider the most widely used data fitting term, namely the least squares error $f(W,H;X) = \frac{1}{2} \|X - WH\|_F^2 = \frac{1}{2} \sum_{i,j}(X - WH)_{ij}^2$. For the volume regularizer, we will consider the following three functions

**Detminant:** $V_{\text{det}}(W) = \frac{1}{2} \det(W^\top W),$

**Log-det:** $V_{\log\det}(W) = \frac{1}{2} \log\det(W^\top W + \delta I_r),$

**Nuclear:** $V_{\text{nuc}}(W) = \|W\|_*,$

where $\det(A)$ is the determinant of matrix $A$, $I_r$ is identity matrix of size $r$, $\delta > 0$ is a constant to lower bound $V_{\log\det}$, and $\|A\|_*$ is the nuclear norm of $A$, that is, the sum of the singular values of $A$. VRNMF aims at fitting the data points within the convex hull of the columns of $W$ which should have a small volume. The reason to consider the functions $V_{\text{det}}$ and $V_{\log\det}$ is that $\sqrt{\det(W^\top W)/r!}$ is the volume of the convex hull of the columns of $W$ and the origin. Hence, $V_{\text{det}}$ is, up to a constant multiplicative factor, the square of that volume, while $V_{\log\det}$ is its logarithm. Let us write $V_{\text{det}}$ and $V_{\log\det}$ as functions of the singular values of $W$, denoted $\sigma_i(W)$ for $1 \leq i \leq r$:

$$V_{\text{det}}(W) = \frac{1}{2} \prod_{i=1}^r \sigma_i^2(W),$$

and

$$V_{\log\det}(W) = \sum_{i=1}^r \log(\sigma_i^2(W) + \delta).$$

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Both functions $V_{\det}$ and $V_{\logdet}$ are non-decreasing functions of the singular values of $W$. This motivates us to consider

$$V_*(W) = \|W\|_* = \sum_{i=1}^r \sigma_i(W).$$

The reason is that this regularizer is also a non-decreasing function of the singular values of $W$, and has been widely used in machine learning for several tasks such as matrix completion [20]. However, to the best of our knowledge, it has never been used in the context of VRNMF and it would be interesting to know how it compares to the standard choices $V_{\det}$ and $V_{\logdet}$.

We will answer the question “which volume function is better in which situations.” These are the main goals of this paper.

The contribution of this work is twofold: the first part of this work is algorithm design, in which we implement and enhance the algorithms to solve VRNMF by using block coordinate descent and optimal first-order methods. Experimental results will show that our algorithms perform better than the current state-of-the-art volume-regularization based method from [7]. The second part of this work is focused on model comparisons. We will answer the question “which volume function is better suited for VRNMF to tackle HU?” We do so by performing extensive numerical comparisons of VRNMF with different volume functions under various settings. The summary of the findings are as follows:

- For data that is highly separable, meaning that there exists data points close to the true endmembers, and in the presence of a few endmembers, VRNMF with $V_{\det}$ produces the best results in most cases.
- For data that is less separable or in the presence of a large number of endmembers, VRNMF with $V_{\logdet}$ performs best in general.

Finally, as a proof of concept, we showcase the ability of VRNMF to produce a meaningful unmixing on hyperspectral images using real-world data.

This work is the continuation of the conference paper [1]. The additional contributions of this extended version are the following:

- We base our numerical experiments only on real endmembers, as opposed to randomly generated ones in [1].
- We use a fine grid search by bisection to tune the regularization parameter $\lambda$.
- We implement VRNMF with the nuclear norm regularizer.
- We have improved our implementations; they are available from https://angms.science/research.html.
- We compare our implementation of VRNMF with the state-of-the-art volume-regularization algorithm RVolMin [7].

The remaining of this paper is organized as follows. In Section II, we give a more complete introduction to HU, followed by the discussion of the pure-pixel assumption also known as the separability assumption. This motivates the use of VRNMF when this assumption is violated. Section III gives the details about the enhancement and implementation of the algorithms for VRNMF. Section IV presents the experiments on VRNMF and Section V concludes the paper.

II. BRIEF REVIEW ON HU

We now give a brief review on HU; see [2], [14] and the references therein for more details. The goal of blind HU is to study the composition and the distribution of materials in a given scene being imaged. A scene usually consists of a few fundamental types of materials called endmembers, and the first goal of blind HU is to obtain the information of these endmembers from the observed hyperspectral image (HSI). HSI are images captured by sensors over different wavelengths in the electromagnetic spectrum. These images form a hyperspectral data cube of size $m \times \text{col} \times \text{row}$, where $m$ is the number of spectral bands, and “col” and “row” are the dimensions of the images, with $n = \text{col} \times \text{row}$. The $m$-by-$n$ data matrix $X$ is obtained by stacking the $m$-dimensional spectral signature of the pixels as the columns of $X$. Given the observed data matrix $X$, the goal of blind HU is to (i) identify the number of endmembers $r$, (ii) obtain the spectral signature of these endmembers, (iii) identify which pixel contains which endmember and in which proportion. This work does not consider problem (i) by assuming $r$ is known. In fact, model order selection is a topic of research on its own; see, e.g., [2] and the references therein. The focus of this work is (ii): recover the (ground truth) endmember spectral matrix, denoted by $W^{\text{true}}$, from the observed data $X$.

In fact, assuming (ii) is solved, (iii) can be tackled by solving a (convex) nonnegative least squares problem [2], [14]. The most widely used model for blind HU is the linear mixing model for which a hidden low-rank linear mixing structure for the data, namely $X = W^{\text{true}}H^{\text{true}} + N$. The data $X$ is hence generated by $W^{\text{true}}$ (the basis, where each column of $W^{\text{true}}$ is the spectral signature of an endmember), weighted by the matrix $H^{\text{true}}$ plus some noise $N$. The matrix $H$ in the HU literature is also called the abundance matrix and encodes how much each endmember (columns of $W^{\text{true}}$) is present in each pixel of the image: $H_{k,j}$ is the abundance of the $k$th endmember in the $j$th pixel. There are two physical constraints in HU: the non-negativity constraints $H \geq 0$ (spectral signatures are nonnegative), and the nonnegativity $H \geq 0$ and sum-to-one constraint $H^\top 1_r = 1_n$ (the abundances in each pixel are nonnegative and sum to one). In this paper, we consider a more general model, namely using $H^\top 1_r \leq 1_n$, that allows to take into account different intensities of illumination among the pixels of the image. Finally, NMF is the right model to perform blind HU under the linear mixing model, that is, to learn the endmember matrix $W$ and the abundance matrix $H$. 

from the data matrix $X$. However, NMF is a difficult problem in general [22].

A. Pure-pixel assumption

Separable NMF (SNMF) is able to solve blind HU when the data satisfies the separability condition, which is also known as the pure-pixel assumption in HU. It means the data $X$ contains at least $r$ pure pixels where each pure pixel contains only one endmember, and there is a one-to-one correspondence between the $r$ pure pixels and the $r$ endmembers. Mathematically, separability changes the NMF model (1) to

$$(SNMF): \quad X \cong W[,H]H\Pi_n = [X(:,A)\ X(:,A)H\Pi_n]$$

where $\Pi_n$ is a $n$-by-$n$ permutation matrix, and $H \in \mathbb{R}^{r \times (n-r)}$ has its columns with $l_1$ norm smaller than one. In SNMF, we have that $W = X(:,A)$, that is, the index set $A$ contains the indices of the pure pixels. Since all columns of $H$ have $l_1$ norm smaller than one, $X(:,A)$ and the origin are the $r$ extreme points of the data cloud $X$, that is, the convex hull of $X(:,A)$ and the origin encapsulates all the other points in $X$. Hence, SNMF is geometrically a vertex identification problem: given $X$, locate the extreme points $W = X(:,A)$ which will be exactly $W^{true}$ if the separability condition holds and no noise is present. Many algorithms exist to perform this task (referred to as pure-pixel search algorithms), e.g., vertex component analysis (VCA) [16] or the successive projection algorithm (SPA) [11]; see [2], [14] and the references therein for more algorithms and discussions. In this paper, we will compare VRNMF to SPA, as SPA is a state-of-the-art provably robust pixel search algorithm.

However, when the separability condition does not hold, pure-pixel search algorithms fail. In order to quantify how much separability is violated, we introduce the non-separability parameter $p \in (0,1]^r$. First, note that separability holds if and only if each row of $H$ contains at least one entry equal to one, that is, $\|h^j\|_\infty = 1$ for $j \in [r]$, where $h^j$ denotes the $j$th row of $H$ and $[r] = \{1,2,\ldots ,r\}$. Therefore, to break the separability condition, we need $\|h^j\| \leq p_j < 1$ for some $j$. Figure 1 gives an example with $r = 4$. Hence, having $\|h^j\| \leq p_j$ means that the maximum abundance of the $j$th endmember in all pixels is at most $p_j$, which sets a minimal amount of separation between the data points (the black dots in Figure 1) to the vertex $w_j$ (the black stars). In other words, $p_j$ controls the gap between $\{x_i\}_{i \in [n]}$ and $w_j$, where $x_i$ is the $i$th data point ($i$th column of $X$). Note that the entries of $p$ can be different in general meaning that we have asymmetric non-separability. For example, in Figure 1, data points are closer to vertex 1 than vertex 4. In the absence of pure pixels, that is, $p_j < 1$ for some $j$, it has been proved that, under mild conditions, minimizing the volume of the convex hull of the columns of $W$ allows to recover the endmembers. These conditions require that the data points are well spread within the convex hull spanned by the columns of $W$; see [6] for a recent survey on the topic.

Note that in Figure 1, the reconstructions given by SPA [11] (green vertices) and RVolMin [7] (deep blue vertices, a state-of-the-art minimum-volume NMF algorithm) are far away from the ground truth. VRNMF with $V_{det}$ (cyan vertices) produces a perfect recovery of $W^{true}$. The next section describes how we solve the VRNMF minimization problem to obtain such results.

III. SOLVING VRNMF

In this section, we describe how to solve (2). We use the framework of block coordinate descent (BCD) by solving the subproblems on $W$ and $H$ separately in an alternating scheme, as done in most NMF works [9]. Let us start with the subproblem for $H$.

A. Subproblem for $H$

Splitting $H$ into columns yields $n$ independent problems: for $1 \leq j \leq n$, solve

$$\min_{h_j \in \mathbb{R}^r_+} \|x_j - Wh_j\|_2^2, \quad \Delta = \{h \in \mathbb{R}^r_+: \|h\|_1 \leq 1 \}, \tag{3}$$

where $h_j$ is the $j$th column of $H$ and $\Delta$ is the $r$-dimensional unit simplex that encodes the non-negativity and sum-to-one constraints in (2). Assuming $\text{rank}(W) = r$ which is a standard assumption, this least squares problem over the unit simplex is a convex problem with strongly convex objective function. We use the accelerated projected gradient (APG) method from Nesterov [17] which requires $O\left(\sqrt{\kappa} \log \frac{1}{\epsilon}\right)$ iterations to reach an $\epsilon$-accurate solution, where $\kappa$ is the condition number of $W^\top W$ which is the Hessian of the objective function in (3). The convergence rate of APG is optimal as no other first-order method can have a faster convergence rate [17]. We defer the explanation of the acceleration scheme to section III-C.
compute the projection onto $\Delta$, we use the implementation from [8] requiring $O(r \log r)$ operations, and which uses the fact that the projection can be written as $P_{\Delta}(h) = |h - I|_{r+}$ where $I$ is a Lagrangian multiplier. Since $r$ is small (usually $r \leq 20$), this implementation is numerically as good as the optimal method [5] with complexity $O(r)$.

In summary, we solve (3) using an optimal first-order method. Note that (3) is parallelizable, we can solve the $n$ problems (3) in parallel.

### B. Subproblem for $W$ with $V_{\text{det}}$

We follow the idea from [24] and perform a block coordinate descent method on the columns $w_i$ of $W$ ($1 \leq i \leq r$). We have

$$||X - WH||_F^2 = ||h^T||_2^2 ||W_i||_2^2 - 2\langle X, h^TW_i \rangle + c,$$

(4)

$$\det(W^TW) = \gamma_i w^T_i Q_i Q_i^TW_i.$$  

(5)

In (4), $X_i \equiv X - \sum_{j \neq i} w_j h_j$ and $c$ is a constant independent of $w_i$. In (5), $\gamma_i \equiv \det(W^TW_i)$ and $Q_i$ is the orthonormal basis of the null space of $W_i^T$, where $W_i$ is $W$ without the column $w_i$; see [24, Appendix] for more details. Using (4) and (5), we obtain the following problem for each column of $W$:

$$\min_{W_i \succeq 0} \frac{1}{2} \gamma_i w^T_i (h^T_i I_m + \gamma_i Q_i Q_i^T) w_i - \langle X, h_i^T, w_i \rangle.$$  

Unlike the problem on $h$, here the objective function in $w_i$ depends on the other columns of $W$. We solve this quadratic program with nonnegativity constraints using APG, which is faster than the standard quadratic programming algorithms used in [24].

### C. Subproblem for $W$ with $V_{\text{logdet}}$

To solve for $V_{\text{logdet}}$, we use majorization minimization, similarly as in [7]. First we have

$$(\text{Lemma 2, [12]}) \quad \log \det(W^TW + \delta I_r) \leq ||W||_F^2 + c,$$

(6)

where $||W||_D = ||D^{1/2}W^T||_F$ is a weighted norm with $D = (Y^T Y + \delta I_m)^{-1} > 0$ for any matrix $Y$. This expression (6) comes from performing the first-order Taylor expansion of the concave function $\log \det(.)$. The inequality (6) holds when $Y = W$. In other words, we minimize the tight convex upper bound on the non-convex logdet function

$$\min_{W \succeq 0} \Phi(W) = \frac{1}{2} \langle W^T W, HH^T \rangle - \langle X, W^T H \rangle + \frac{1}{2} ||W||_D^2,$$

(7)

where $D = \left( W^{t-1} W^{t-1} + \delta I_r \right)^{-1}$. To solve (7), we again use APG. However, we embed the following two acceleration strategies to APG:

- The adaptive restart heuristic from [18]. The APG update of $W$ at iteration $t$ can be expressed as

$$W^{t+1} = \left[ W^t - \alpha_t \nabla \Phi(W^t + \beta_t \Delta_t) + \beta_t \Delta_t \right]_+,$$

where $\alpha_t$ is the step size, $\nabla \Phi$ is the gradient of the objective function and $[\cdot]_+$ is the projection onto the nonnegative orthant. Here $\beta_t \Delta_t$ is the momentum term added in Nesterov’s acceleration that extrapolates $W^t$ along the direction $\Delta_t = W^t - W^{t-1}$ and $\beta_t \in [0, 1]$ is a parameter with $\beta_0 = 0$. The extrapolation may increase the value of $\Phi$, and when this happens, we “restart” the APG scheme by reinitializing $\beta_t$, which reduces the APG back to a standard projected gradient step that decreases the value of $\Phi$.

- The general acceleration framework for NMF algorithms from [10]. There are several terms independent of $W$, in particular $HH^T$ and $XH^T$, in the gradient term and are the main computational cost. The idea has two components: (i) pre-compute the terms independent of $W$ outside the update to avoid repeated computation of the same terms, and (ii) perform the update on $W$ multiple times to reuse these precomputed terms (in most standard NMF algorithms, $W$ is only updated once before the update of $H$).

Finally, due to space limit, we do not present another majorization for logdet which is based on eigenvalue inequality proposed in the conference work [1]. In short, that approach is another relaxation of (7) that unlocks the coupling between columns of $W$ so that column-wise update similar to the one mentioned in section III-B can be used.

### D. Subproblem for $W$ with $V_s$

The nuclear norm regularized VRNMF problem on $W$ is

$$\min_{W \succeq 0} \frac{1}{2} \langle W^T W, HH^T \rangle - \langle X, W H \rangle + \lambda \|W\|_*.$$  

(8)

Ignoring for now the non-negativity constraints, we solve the resulting problem by proximal gradient which updates $W$ as follows:

$$W^{t+\frac{2}{3}} = \text{prox}_{\theta f, \rho} \left\{ W^{t+\frac{1}{3}} \right\}, \quad W^{t+\frac{2}{3}} := W^{t} - \alpha \nabla f(W^t),$$

where $\alpha$ is step size and $\nabla f$ is the gradient of $\frac{1}{2} \langle W^T W, HH^T \rangle - \langle X, W H \rangle$. For the step size, we use $\alpha = 1/L$ where $L = \|HH^T\|_2$ is the Lipschitz constant of the gradient. The proximal operator of $\|\cdot\|_*$ with step size $\theta$ has the closed-form expression given by the singular value thresholding (SVT) operator [3], we have

$$W^{t+\frac{2}{3}} = \text{SVT}_\theta \left( W^{t+\frac{1}{3}} \right) := U[\Sigma - \theta I]_+ V^T,$$

where $UV^T$ is the SVD of $W^{t+\frac{1}{3}}$. Finally, to take the non-negativity constraint into account, we simply put a nonnegative projection after SVT and set $W^{t+1} = [W^{t+\frac{2}{3}}]_+$. 

### E. Summary of the algorithms

Algorithm 1 shows the general framework for solving VRNMF. Our implementations are faster than previous works because of the use of Nesterov’s acceleration [17], adaptive restart [18] and the use of the acceleration strategy for NMF from [10]. Moreover, our update of $W$ in VRNMF using $V_s$ is new.

We refer to the algorithm using $V_{\text{det}}$, $V_{\text{logdet}}$ and $V_s$ as $\text{Det}$, $\text{logdet}$ and Nuclear, respectively.
Algorithm 1 VRNMF

Input: $X \in \mathbb{R}^{m \times n}$, an integer $r$, $\lambda \geq 0$
Output: $W \in \mathbb{R}_+^{m \times r}$, $H \in \mathbb{R}_+^{r \times n}$ for problem (2).

Initialize $(W_0^0, H^0)$ by SPA [11]

for $t = 1, 2, \ldots$ do
  Update of $W$
  Compute and store $HH^\top$ and $XH^\top$
  if $V = V_{\text{det}}$ then
    Update $W$ as stated in section III-B.
  else if $V = V_{\text{logdet}}$ then
    Update $W$ as stated in section III-C.
  else if $V = V_{\text{Nuclear}}$ then
    Update $W$ as stated in section III-D.
  end if
  Update of $H$ by APG [8].
end for

We end this section by briefly mentioning the convergence of these algorithms. VRNMF is a non-convex problem and it can be shown that the sequence $\{W_t, H_t^t\}_{t \geq 1}$ produced by Algorithm 1 converges to a first-order stationary point: For $\text{Det}$, the convergence comes from the standard result of coordinate descent [23]. For $\text{logdet}$, which involves the use of the upper bound $\Phi'$, convergence comes from the theory of [19]. For $\text{Nuclear}$, convergence result of proximal gradient applies [3]. Figure 2 shows a typical convergence curve of the algorithms.

IV. EXPERIMENTS

In this section we compare the VRNMF models on solving HU. We first describe the general experimental setting in section IV-A and then report the experimental results on recovering the ground truth $W^{\text{true}}$ under different asymmetric non-separability and different noise levels in the subsequent sub-sections. Finally we present results on two real-world data sets. All the experiments are run with MATLAB (v.2015a) on a laptop with 2.4GHz CPU and 16GB RAM. The codes available from https://angms.science/research.html.

A. Settings

1) Data generation: For synthetic experiments, we generate the observed data matrix as $X = [X^{\text{clean}} + N]_+$ with white Gaussian noise $N \in \mathbb{R}^{m \times n}$ with zero mean and variance $\sigma \geq 0$. We generate $X^{\text{clean}} = W^{\text{true}}H^{\text{true}}$, where $W^{\text{true}}$ comes from several datasets available from http://lesun.weebly.com/hyperspectral-data-set.html [25] (unlike the conference version [1] that generated $W^{\text{true}}$ at random); see Figure 3. We generate each column of $H \in \mathbb{R}_+^{r \times n}$ using the Dirichlet distribution of parameter 0.1. If $p = 1_r$, that is, separability condition holds, we take $H$ as $H^{\text{true}}$. Otherwise, we remove the columns of $H$ with at least one element in the $j$th coordinate that exceeds the value $p_j$, and resample again until $H$ satisfies the condition $\|h_j\|_\infty \leq p_j$ for all $j$. In this paper, we will use $p_j \in (0.5, 0.99)$ for all $j$. In all the experiments, we set the number of data points $n = 1000$ and the maximum number of iterations to 300.

For experiments on real data, we take the real data as $X$, without any preliminary dimension reduction, and without any pre-processing to suppress noise or to remove outliers.

2) Parameters of the algorithms: We will compare our three proposed algorithms $\text{Det}$, $\text{logdet}$ and $\text{Nuclear}$ with SPA [11] and RVolMin which is a state-of-the-art volume-regularized method [7]. For RVolMin, we use the same data fitting term (namely, the Frobenius norm), the same number of iterations, the same parameter search scheme and the same initialization as for our algorithms.

Given an observed data matrix $X$, all VRNMF algorithms have two main parameters: $r$ and $\lambda$. They also require an initialization $(W^{\text{ini}}, H^{\text{ini}})$. We assume $r$ is known. We generate $W^{\text{ini}}$ from $X$ using SPA [11], and generate $H^{\text{ini}}$ using the method described in Section III-A. The regularization parameter $\lambda$ should usually be chosen small. In fact, a large $\lambda$ forces the vertices of the convex hull of $W$ to be very close to each other, making $W$ ill-conditioned and/or rank-deficient. In particular, for VRNMF with $V_c$, the SVT operator set singular values smaller than $\lambda$ to zero, so a large $\lambda$ makes $W$ rank-deficient. The following describes how we tune $\lambda$. The goal here is to tune $\lambda$ so that each algorithm performs as best as possible for the considered problems. To achieve this goal, we use the ground truth $W^{\text{true}}$. We set $\lambda = \frac{f(W^{\text{ini}}, H^{\text{ini}})}{|V(W^{\text{ini}})|}$, where $(W^{\text{ini}}, H^{\text{ini}})$ are the initial solutions obtained by SPA, and $\lambda$ is a tuning variable within the search interval $I_0 = [10^{-6}, 0.5]$. We perform grid search by bisection in a greedy way to tune $\lambda$:

- (step-1) Take $\lambda_a = 10^{-6}$, $\lambda_b = 0.5$ initially.
- (step 2) Run the algorithm with $\tilde{\lambda}_a$, $\tilde{\lambda}_b$ and $\tilde{\lambda}_c$, where $\tilde{\lambda}_c = 0.5(\lambda_a + \lambda_b)$. Denote the performance of the solution $W$ produced under a specific $\lambda$ as $\text{MRSA}(\lambda)$, where $\text{MRSA}$ will be defined in the next paragraph.
- (step-3) Split the search interval $I_0 = [\lambda_a, \lambda_b]$ into two intervals $I_0^1 = [\tilde{\lambda}_a, \tilde{\lambda}_c]$ and $I_0^2 = [\tilde{\lambda}_c, \lambda_b]$. For each interval there are two MRSA values, we let the MRSA value of an interval be the sum of these values.
- (step-4) We repeat step-1 to step-3 on the interval with the lowest MRSA value. That is, at iteration $k$, we shrink the
search interval by half by defining the new search interval $I_{k+1}$ as

$$I_{k+1} \leftarrow \min_z \{ \text{MRSA}(I^1_k), \text{MRSA}(I^2_k) \}.$$ 

- If a draw happens in step-4, we perform two bisections on each of the interval $I^1_k$ and $I^2_k$ and set the next interval to be the one with the smallest MRSA.
- We repeat this process (step-1 to step-4) at most 20 times, or if the improvement from one iteration to the next is negligible, namely if

$$| \text{MRSA} \left( \lambda_{k+1} \right) - \text{MRSA} \left( \lambda_k \right) | \leq 10^{-4}.$$ 

Note that such greedy bisection search does not guarantee that $\lambda_k$ will converge to the best value, that is, the value that corresponds to the lowest MRSA. However, we have observed in extensive numerical experiments the effectiveness of this scheme, which will be illustrated in IV-B.

c) Performance metric: We measure the quality of a solution $W$ produced by VRNMF algorithms using the mean removed spectral angle (MRSA) between $W$ and $W^{\text{true}}$. 

Fig. 3. Top row The datasets from [25]. Bottom subplots The endmembers $W^{\text{true}}$ of the datasets. Best viewed in color.
TABLE I
COMPARISON OF MRSA VALUES OF THE BISECTION AND THE GRID SEARCH TO OBTAIN $\lambda^b$ AND $\lambda^*$, RESPECTIVELY. THE SECOND COLUMN IS THE NUMBER OF ITERATIONS NEEDED FOR THE BISECTION SEARCH TO TERMINATE.

<table>
<thead>
<tr>
<th>$p$</th>
<th>$\frac{\text{MRSA}(\lambda^b) - \text{MRSA}(\lambda^<em>)}{\text{MRSA}(\lambda^</em>)}$</th>
<th># of iterations</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.93</td>
<td>-0.0016±0.0002</td>
<td>2±0.00</td>
</tr>
<tr>
<td>0.89</td>
<td>-0.0143±0.0023</td>
<td>6.4±0.52</td>
</tr>
<tr>
<td>0.86</td>
<td>-0.0235±0.0029</td>
<td>7.9±0.32</td>
</tr>
<tr>
<td>0.83</td>
<td>-0.0429±0.0032</td>
<td>9±0.00</td>
</tr>
<tr>
<td>0.79</td>
<td>-0.0815±0.0089</td>
<td>11.1±0.32</td>
</tr>
<tr>
<td>0.76</td>
<td>-0.0919±0.0092</td>
<td>10.5±0.53</td>
</tr>
</tbody>
</table>

MRSA between two vectors $x, y \in \mathbb{R}^m \setminus \{0_m\}$ is defined as

$$\frac{100}{\pi} \cos^{-1}\left( \frac{\langle x - \bar{x}, y - \bar{y} \rangle}{\|x - \bar{x}\|_2 \|y - \bar{y}\|_2} \right) \in [0, 100].$$

(9)

MRSA gives a better measurement than relative percentage error as it purely depends on the shapes of $x$ and $y$, where the effects of shifts and scaling are removed. A low MRSA value means a good matching between $x$ and $y$. We measure the performance of algorithms by calculating the MRSA between $\{w_{i,1}^{\text{true}}\}_{i \in [r]}$ and $\{\bar{w}_{i,1}^{\text{true}}\}_{i \in [r]}$, which is defined as the mean of MRSA between each vector pair $\{w_i, \bar{w}_i^{\text{true}}\}$, such value is within $[0, 100]$.

B. Effectiveness of the bisection search on $\lambda$

In this section, we illustrate the effectiveness of our tuning strategy for $\lambda$. We perform experiments on the synthetic dataset where $W_{\text{true}}$ comes from the Samson dataset with $r = 3$ as follows. We consider 6 different values of the symmetric non-separability vector $p = [p_1, p_2, p_3]$ where $p_1 = p_2 = p_3$ are selected from the set $\{0.93, 0.89, 0.86, 0.83, 0.79, 0.76\}$. We run $\text{Det}$ with two parameter tuning schemes: (1) the bisection search mentioned in IV-A, and (2) a brute-force grid search: we run $\text{Det}$ with all 100 equally spaced steps values of $\lambda$ in the interval $I_0 = [10^{-6}, 0.5]$.

For each value of $p$, 10 data sets are generated randomly. Let us denote $\lambda^b$ the value of $\lambda$ obtained by the bisection search and $\lambda^*$ by the grid search. Table I shows the results between comparing the bisection search and the grid search displayed as $\frac{\text{MRSA}(\lambda^b) - \text{MRSA}(\lambda^*)}{\text{MRSA}(\lambda^*)}$ in the format of (mean±std) and the average number of iterations performed by the bisection search to reach $\lambda^b$. From Table I, we make the following two interesting observations:

1) As the purity goes down, more bisections are need to identify the best $\lambda$. This was expected since, for a high purity, the initialization (SPA) provides a good initial solution.

2) In all 50 cases, the value of $\lambda^b$ lead to a slightly smaller MRSA value than $\lambda^*$ while requiring less computations. This illustrates the fact that the bisection is able to identify the right value of $\lambda$ and to refine the search around that value with more precision than an expensive exhaustive search.

C. Comparison with MVC-NMF

Minimum volume constrained nonnegative matrix factorization (MVC-NMF) is one of the first minimum-volume NMF algorithm\(^1\) [15]. Let us run a small experiment to show that MVC-NMF does not compete with $\text{Det}$, $\text{logdet}$ and Nuclear and RVolMin [7]. Similarly as in the previous section, we use synthetic datasets where $W_{\text{true}}$ comes from the Samson dataset with $r = 3$. However, we use more difficult scenarios using the non-separability vectors $p = [p_1, p_2, p_3]$ where $p_1$ is selected from $\{0.95, 0.9, 0.85, 0.8, 0.75\}$, $p_2 = 0.79$, and $p_3 = 0.69$. For each value of $p$, 10 data sets are generated randomly. All algorithms take the same initialization (SPA), the same number of iterations (100) and the regularization parameter $\lambda$ is tuned using the bisection search. Figure 4 shows that MVC-NMF produces significantly worse results than the four other algorithms, hence will not be considered in the extensive numerical experiments the subsequent comparisons. It is difficult to pinpoint the reasons of the poor results of MVC-NMF; we see at least two of them: (1) MVC-NMF uses another regularizer which is the squared volume of the convex hull of the columns of $W$ without the origin, and (2) it does not use optimal optimization methods (for example, it uses a fixed step size of 0.001).

D. Using the Samson dataset with $r = 3$

Now we run more comprehensive experiments on the synthetic dataset where $W_{\text{true}}$ comes from the Samson data set with $r = 3$ as follows. We perform $10 \times 10 \times 10 = 1000$ experiments where the data in each experiment is generated using the non-separability vector $p = [p_1, p_2, p_3]$, where $p_i$ is selected from the set $P = \{0.99, 0.96, 0.93, 0.89, 0.86, 0.83, 0.79, 0.76, 0.73, 0.69\}$. Figure 5 shows the results in form of MRSA cubes for the algorithms SPA, Det, logdet, Nuclear and RVolMin, where each pixel in the cube is the result (in MRSA) over one trial. We then construct the recovery curves by counting the number of cases (pixels) in the cube that the MRSA value

\(^1\)The code available from https://github.com/aiclp/MVCNMF
is less than a threshold; see Figure 6. In general, the results in Figures 5 and 6 show that

- All VRNMF algorithms perform better than the state-of-the-art SNMF algorithm SPA, as the MRSA cubes of VRNMF have a wider blue region (lower MRSA) and their recovery curves in Figure 6 dominates that of SPA.

- When the data is highly non-separable (low \( p \)'s), VRNMF algorithms perform worse than SPA. In fact, the region of the cube corresponding to highly non-separable data in SPA is not as red as for the VRNMF approaches. The reason is that SPA always extract points from the data could hence these points are never too far from the vertices. However, when the data is highly non-separable, VRNMF may generate points further away than the vertices.

- Compared with RVolMin, logdet and Nuclear are consistently better, while Det performs similarly.

- logdet performs better than Det and Nuclear for highly non-separable data, as its red region is much more concentrated around the highly non-separable corner of the cube.

In terms of computational time, Det takes \(2.1\pm0.2\) seconds, logdet takes \(1.2\pm0.1\) seconds, Nuclear takes \(1.3\pm0.2\) seconds, and RVolMin takes \(2.1\pm0.0\) seconds. As expected, due to the inner loop and the computation of \(Q_i, \text{Det} \) is slower than the other algorithms.

**E. Using the Jasper dataset with \(r = 4\)**

In Figure 7, we perform the same experiment as in the previous section on the dataset Jasper with \(r = 4\), where we fix \(p_4 = 0.75\). Furthermore, Table II shows the result in MRSA on the dataset Jasper across three sets of predefined \(p\) values (highly separable with \(p_{\text{high}} = [0.9, 0.8, 0.7, 0.6]\), less separable with \(p_{\text{mid}} = [0.8, 0.7, 0.6, 0.51]\), and even less separable with \(p_{\text{low}} = [0.7, 0.65, 0.55, 0.51]\)) and three noise levels (\(\sigma = 0.001, 0.005\) and 0.01). Each item in the table (in mean±std) is the average over 20 trials.

We observe the following

- Figure 7 show that VRNMF algorithms are competitive with the state-of-the-art minimum volume based method RVolMin, where Det performs slightly better than RVolMin while logdet is significantly better.

- Tables II shows that all the methods improve the fitting accuracy of SPA, with Det has the best performance in all cases and RVolMin has the worst performances in all cases.

In terms of computational time, Det takes \(6.62\pm0.5\) seconds, logdet takes \(2.22\pm0.2\) seconds, Nuclear takes \(1.45\pm0.0\) seconds, and RVolMin takes \(2.7\pm0.0\) seconds. Hence, for the same reasons as in the previous experiment, Det is slower.

**F. Using the Urban dataset with \(r = 6\)**

Table III shows the result of the same experiment as in the previous section performed with the Urban dataset. Here \(p_{\text{high}} = [0.9, 0.75, 0.7, 0.65, 0.6, 0.85]\),

![Figure 5. MRSA cubes for the different NMF algorithms viewed from two different angles, for the synthetic Samson data sets with \(r = 3\). All the cubes share the same color scheme. In general, MRSA cubes for VRNMF have a wider blue region than that of SPA. However, when the vector \(p\) has a low value, VRNMF gives less accurate fitting than SPA (the red regions). VRNMF with logdet provides the best results (widest dark blue region). Best viewed in color.](image)

\[p_{\text{mid}} = [0.8, 0.7, 0.65, 0.6, 0.75, 0.8], \quad \text{and} \quad p_{\text{low}} = [0.7, 0.6, 0.55, 0.51, 0.65, 0.7]. \]

The noise levels are \(\sigma = 0.001, 0.005\) and 0.01. The results show that

- All the methods improve the fitting accuracy of SPA.

- Det has the best performance in most cases, with logdet as the first-runner up.

- Det performs well for \(p_{\text{high}}\) and \(p_{\text{mid}}\). With \(p_{\text{low}}\), it is not as good as logdet.

- Nuclear and RVolMin have the worst performances in all cases.

- The computational times are: Det \(7.5\pm0.2\), logdet \(1.8\pm0.1\), Nuclear \(1.5\pm0.0\) and RVolMin \(2.3\pm0.0\).
recovery rate/%

Fig. 6. Curves of recovery corresponding to Figure 5 for the synthetic Samson data sets with \( r = 3 \).

Fig. 7. Curves of recovery corresponds for the dataset Jasper with \( r = 4 \). Same experimental set up as the one in Figure 6, but where \( p_4 \) fixed at 0.75.

**TABLE II**

<table>
<thead>
<tr>
<th>Method</th>
<th>( p_{\text{high}} )</th>
<th>( p_{\text{mid}} )</th>
<th>( p_{\text{low}} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>SPA</td>
<td>5.40±0.60</td>
<td>12.62±0.18</td>
<td>20.76±0.23</td>
</tr>
<tr>
<td>Det</td>
<td>0.41±0.08</td>
<td>0.40±0.06</td>
<td>10.99±1.68</td>
</tr>
<tr>
<td>logdet</td>
<td>0.48±0.54</td>
<td>3.03±0.28</td>
<td>12.57±1.49</td>
</tr>
<tr>
<td>Nuclear</td>
<td>0.64±0.07</td>
<td>2.12±0.13</td>
<td>19.90±1.40</td>
</tr>
<tr>
<td>RVolMin</td>
<td>1.04±0.38</td>
<td>4.99±0.22</td>
<td>13.67±2.99</td>
</tr>
</tbody>
</table>

Across different noise levels \( (p = p_{\text{high}}) \):

<table>
<thead>
<tr>
<th>Method</th>
<th>( \sigma = 0.001 )</th>
<th>( \sigma = 0.01 )</th>
<th>( \sigma = 0.05 )</th>
</tr>
</thead>
<tbody>
<tr>
<td>SPA</td>
<td>5.40±0.60</td>
<td>12.62±0.18</td>
<td>20.76±0.23</td>
</tr>
<tr>
<td>Det</td>
<td>0.41±0.08</td>
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<tr>
<td>RVolMin</td>
<td>1.04±0.38</td>
<td>4.99±0.22</td>
<td>13.67±2.99</td>
</tr>
</tbody>
</table>

G. On the Cuprite dataset with \( r = 12 \)

Table IV shows the result on the same experiments conducted with the Cuprite dataset. Here we concatenate the \( p \) vector used in Urban two times to form the \( p \) vector with length 12 for the data Cuprite. For example, for \( p_{\text{Capride}} \), we use \( p_{\text{high}}^{\text{Capride}} = [p_{\text{high}}^{\text{Urban}} \quad p_{\text{high}}^{\text{Urban}}] \).

The result from the two tables show that

- All the methods improve the fitting accuracy of SPA.
- logdet has the best performance in all situations, with Det and Nuclear as the first-runner ups.
- RVolMin has the worst performances in most cases.

In terms of computational time, Det takes 28.1±0.9 seconds, logdet takes 3.8±1.1 seconds, Nuclear takes 3.4±0.1 seconds, and RVolMin takes 3.2±0.0 seconds. As expected, when \( r \) increase, Det takes much more time and it is not recommended for large \( r \).

In general, the results show that the VRNMF algorithms with Det and logdet performs very well, better than RVolMin and Nuclear in terms of fitting accuracy. As RVolMin consistently produce inferior results, we do not include it in the subsequent sections.

H. On image segmentation on real data

In this section we run the algorithms Det, logdet and Nuclear on real HU data Samson and Jasper. As stated in section IV-A, no pre-processing is performed on the raw
data and we use the raw data directly. The following states the specifications of the datasets. For the dataset Samson, \((m, n, r) = (156, 95^2, 3)\). For the dataset Jasper, \((m, n, r) = (198, 100^2, 4)\). We have tuned \(\lambda\) in the same way as for the synthetic datasets using the endmembers \(W_{\text{Ref}}\) from [25]. Figure 8 shows the decompositions. In the same figure, we also show the references provided by [25], and we list the numerical results in Table V. In Table V, the MRSA is calculated with respect to the reference \(W_{\text{Ref}}\) of [25]. The results show that all three VRNMF algorithms produce meaningful decomposition.

**Remark** The reference [25] used a sparsity regularization on \(H\) and hence the abundance map of [25] looks cleaner. It is out of the scope of this paper to consider a sparsity regularization on \(H\) as we are focusing on the volume regularization on \(W\). This is a direction for further research.

**V. CONCLUSION**

In this paper, NMF models with different volume regularizations (VRNMF) were investigated. We have developed highly efficient algorithms for these VRNMF models. The VRNMF algorithms are shown to be able to outperform both the state-of-the-art separable NMF algorithm SPA, and the volume-based methods MVC-NMF [15] and RVolMin [7]. Furthermore, extensive experimental results using real hyperspectral data showed that, when the data has a small rank \(r\) \((r \leq 4)\) and is highly separable, the volume regularizer based on the determinant (\(\text{Det}\)) provides the best results, although the the regularizer based on the logarithm of the determinant (\(\log\text{det}\)) provides almost as good decompositions. When the rank increases and/or the data becomes less separable, \(\log\text{det}\) performs best in most cases, while being computationally faster than \(\text{Det}\). Therefore, in practice, we recommend the use of \(\log\text{det}\).

**REFERENCES**


[10] Gillis, N., Glineur, F.: Accelerated multiplicative updates and hier-
Fig. 9. The distribution maps of the material in the whole scene obtained from the VRNMF algorithms. In (a), the colour code is: white – rock, blue – water, and green – tree. In (b), the colour code is: green – tree, blue – water, yellow – dirt and white – road. The map is constructed by taking the largest component of $h_j$ as the material of that pixel. As the factorization rank $r$ here is small and these datasets are highly separable, Det produces the best segmentation results. For example, the road in Jasper produced by Det is better identified than for the other two. Best viewed in color.


