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Hyperspectral Unmixing With Spectral Variability Using Adaptive Bundles and Double Sparsity

Tatsumi Uezato¹, Mathieu Fauvel, *Senior Member, IEEE*, and Nicolas Dobigeon², *Senior Member, IEEE*

Abstract—Spectral variability is one of the major issues when conducting hyperspectral unmixing. Within a given image composed of some elementary materials (herein referred to as endmember classes), the spectral signatures characterizing these classes may spatially vary due to intrinsic component fluctuations or external factors (illumination). These redundant multiple endmember spectra within each class adversely affect the performance of unmixing methods. This paper proposes a mixing model that explicitly incorporates a hierarchical structure of redundant multiple spectra representing each class. The proposed method is designed to promote sparsity on the selection of both spectra and classes within each pixel. The resulting unmixing algorithm is able to adaptively recover several bundles of endmember spectra associated with each class and robustly estimate abundances. In addition, its flexibility allows a variable number of classes to be present within each pixel of the hyperspectral image to be unmixed. The proposed method is compared with other state-of-the-art unmixing methods that incorporate sparsity using both simulated and real hyperspectral data. The results show that the proposed method can successfully determine the variable number of classes present within each class and estimate the corresponding class abundances.

Index Terms—Endmember variability, hyperspectral imaging, sparse unmixing, spectral unmixing.

I. INTRODUCTION

HYPERSPECTRAL analysis has received an increasing attention because of its high spectral resolution, which enables a variety of objects to be identified and classified [1]. Mixed pixels caused by the presence of multiple objects within a single pixel adversely affect the performance of hyperspectral analysis [2]. To address this problem, a wide variety of spectral unmixing methods have been developed over the last decades [3]–[6]. Spectral unmixing methods aim at decomposing a mixed spectrum into a collection of reference spectra (known as endmembers) characterizing the macroscopic materials present in the scene and their respective

proportions (known as abundances) in each image pixel [2]. Despite the large number of developed spectral unmixing methods, there are still major challenges for accurate estimates of endmember signatures and abundances [3]. Among these challenges, endmember variability may lead to large amounts of errors in abundance estimates [7]. It results from the fact that each endmember can rarely be represented by a unique spectral signature. Conversely, it is subject to so-called spectral variability, e.g., caused by variations in the acquisition process, the intensity of illumination, or other physical characteristics of the materials [8], [9]. Taking this endmember variability into account during the spectral unmixing process is one of the keys for successful application of spectral unmixing [10].

The methods that incorporate endmember variability can be categorized into two main approaches (see [7], [10], [11] for recent overviews). The first approach relies on the definition of a set of multiple spectral signatures, referred to as endmember bundles, to characterize each endmember class. Endmember bundles can be collected from field campaign or can be extracted from data itself using endmember bundle extraction methods [9], [12], [13]. The advantage of this approach is the simple representation of the endmember variability neither assuming any particular distribution nor adding additional terms describing this variability. This can be achieved by simply selecting physically possible spectral signatures from prior information (i.e., endmember bundles). Endmember bundles can be validated by experts in order to provide accurate representation about endmember variability [12]. Although traditional methods incorporating endmember bundles (see [14]) are known to be computationally expensive, more efficient methods have been recently developed and have shown great potential [15]–[18]. However, as pointed out in [19], it is unlikely that the endmember bundles completely represent endmember variability present in an image. Such incomplete endmember bundles may lead to poor estimates of abundances. The second approach uses physical or statistical descriptions of the endmember variability. More precisely, these methods describe the endmember variability thanks to a statistical distribution [20] or by incorporating additional variability terms in the mixing model [21]–[23]. This approach can potentially incorporate endmember bundles as supervised methods (see [24], [25]). The advantage of this approach results from the adaptive learning of the endmember variability. Indeed, state-of-the-art methods such as those recently introduced in [21]–[23] enable endmember spectra to spatially vary within each pixel in order to describe

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endmember variability. This is important since the endmember spectra to be used for the abundance estimation can be different between pixels. However, estimating endmember variability is a challenging task, especially when large amounts of endmember variability are present in an image. In addition, the statistical distribution or additional terms used in these methods may be overly simplified to represent endmember variability.

Both approaches demonstrate benefits and drawbacks. A natural question arises: is it possible to combine the strong advantages of both approaches to robustly represent endmember variability? This paper addresses the question and introduces a novel spectral unmixing method that bridges the gap between the aforementioned two approaches with the help of a double sparsity-based method inspired by Rubinstein *et al.* [26]. Specifically, the proposed method aims at adaptively recovering endmember spectra within each pixel to describe endmember variability while preserving simple representation. The proposed method is closely related to the existing methods. Thus, the main contributions of this paper are threefold: 1) to propose a novel spectral unmixing method that incorporates adaptive endmember bundles through double sparsity representation; 2) to give a systematic review of related work and show the relationship between the proposed method and existing methods; and 3) to provide comparison between the proposed method and other sparsity-based methods.

This paper is organized as follows. Section II describes related works and existing methods, while highlighting their inherent drawbacks. In Section III, a novel mixing model that incorporates endmember variability is proposed and its relationships with existing methods are discussed. Section IV introduces an associated unmixing algorithm designed to recover the endmember classes, adaptive bundles, and abundances. Sections V and VI show experimental results obtained from simulated data and real hyperspectral images. Finally, the conclusion is drawn in Section VII.

II. RELATED WORKS AND ISSUES RAISED BY EXISTING METHODS

A. Conventional (Variability-Free) Linear Mixing Model

Let $\mathbf{y}_i \in \mathbb{R}^{L \times 1}$ denote the L -spectrum measured at the i th pixel of a hyperspectral image. According to the linear mixing model (LMM), the observed spectrum of the i th pixel \mathbf{y}_i is approximated by a weighted linear combination of endmember spectra and abundance fractions

$$\mathbf{y}_i = \mathbf{M}\mathbf{a}_i + \mathbf{n}_i \quad (1)$$

where $\mathbf{M} \in \mathbb{R}^{L \times K}$ is the matrix of the spectral signatures associated with the K endmember classes, $\mathbf{a}_i = [a_{1i}, \dots, a_{Ki}]^T \in \mathbb{R}^{K \times 1}$ is the abundance fractions of the pixel, and $\mathbf{n}_i \in \mathbb{R}^{L \times 1}$ represents noise and modeling error. LMM is generally accompanied by abundance nonnegativity constraint (ANC) and the abundance sum-to-one constraint (ASC)

$$\forall k, \forall i, \quad a_{ki} \geq 0, \quad \text{and} \quad \forall i, \quad \sum_{k=1}^K a_{ki} = 1 \quad (2)$$

where a_{ki} is the abundance fraction of the k th class in the i th pixel. This model implicitly relies on the following assumption: each endmember class in \mathbf{M} is described by a unique spectrum and commonly used to unmix all the pixels of a given image. In other words, LMM does not account for spectral variability. However, as discussed previously, this is likely unrealistic because spectral variability is naturally observed in hyperspectral images, e.g., because of variations in illumination or physical intrinsic characteristics of materials [19].

B. Linear Mixing Models Incorporating Endmember Bundles

LMMs that consider endmember bundles have been developed. Endmember bundles can be extracted manually using the method developed in [12] or the ENVI software as done in [19]. Several methods that automatically extract endmember bundles have also been developed [9], [13], [27]. The automated endmember bundle extraction method proposed in [13] sequentially generates a number of subsets representing randomly selected pixels and extracts endmembers from each subset. Then, bundles are generated by grouping the extracted multiple endmembers. More recently, endmember bundle extraction methods that consider both spatial and spectral information have been developed [9], [27]. Endmember bundles can be potentially estimated by LMMs incorporating spectral variability [21]–[23]. These methods explicitly add a term representing endmember variability and, thus, estimate variable endmember spectra within each class. The estimated endmember spectra of each class can be subsequently grouped into endmember bundles. Although endmember bundles can be extracted by the above methods, in this paper, we assume that endmember bundles are *a priori* available and have been previously identified. This prevents unmixing results to be affected by the performance of the endmember bundle extraction step and enables fair comparison.

Multiple endmember spectral mixture analysis (MESMA) [14] allows the variability of the endmember spectrum representative of each class and a varying number of endmember classes present within each pixel. Although MESMA has been widely used for a variety of applications [14], [28], it owns several major limitations.

- 1) It is highly computationally expensive because MESMA needs to test a large number of combinations of endmember spectra [29].
- 2) MESMA tends to select an overestimated number of endmember classes because it uses the reconstruction error to select the appropriate combination of endmember spectra [30].
- 3) The performance of MESMA may significantly decrease when endmember spectra (or bundles) within each class do not completely represent the spectral variability [19].

To overcome these limitations, recent works have proposed a new class of methods that incorporate all endmember bundles defined as [15], [16]

$$\mathbf{E} = [\mathbf{E}_1 | \mathbf{E}_2 | \dots | \mathbf{E}_K] \quad (3)$$

where $\mathbf{E}_k \in \mathbb{R}^{L \times N_k}$ represents a set of endmember spectra (i.e., bundle) characterizing the k th class, N_k is the number of

endmember spectra in the k th class, and N is the total number of endmember spectra of all classes with $N = \sum_{k=1}^K N_k$. Generalizing LMM in (1), those methods first model a given observed pixel spectrum with respect to (w.r.t.) all spectra in endmember bundles and corresponding multiple abundances

$$\mathbf{y}_i = \mathbf{E}\mathbf{r}_i + \mathbf{n}_i \quad (4)$$

where $\mathbf{r}_i \in \mathbb{R}^{N \times 1}$ is multiple abundance fractions corresponding to each spectrum of the endmember bundles \mathbf{E} . As for LMM, ASC or ANC can also be imposed to \mathbf{r}_i . As a second step, multiple abundance fractions \mathbf{r}_i are summed within each class to generate a single abundance fraction for each class

$$\mathbf{a}_i = \mathbf{G}^T \mathbf{r}_i \quad (5)$$

with

$$\mathbf{G} = \begin{bmatrix} \mathbf{1}_{N_1} & \mathbf{0}_{N_1} & \cdots & \mathbf{0}_{N_1} \\ \mathbf{0}_{N_2} & \mathbf{1}_{N_2} & \cdots & \mathbf{0}_{N_2} \\ \vdots & \vdots & \ddots & \vdots \\ \mathbf{0}_{N_K} & \mathbf{0}_{N_K} & \cdots & \mathbf{1}_{N_K} \end{bmatrix} \quad (6)$$

where $\mathbf{1}_{N_k} \in \mathbb{R}^{N_k \times 1}$ is a column vector of ones and $\mathbf{0}_{N_k} \in \mathbb{R}^{N_k \times 1}$ represent an N_k -dimensional vector whose components are zeros. While these two steps are conducted separately in [15] and [16], they can also be considered jointly within a multitask Gaussian process framework [17], [19]. Even if these methods have been shown to be effective, a large number of endmember spectra within each class may be redundant. In such case, following a model selection inspiration, Veganzones *et al.* [15] introduce a complementary sparsity regularization on the multiple abundance vectors

$$\begin{aligned} \min_{\mathbf{r}_i} \quad & \frac{1}{2} \|\mathbf{E}\mathbf{r}_i - \mathbf{y}_i\|_2^2 + \lambda_r \|\mathbf{r}_i\|_1 \\ \text{s.t. } \forall i, \quad & \mathbf{r}_i \succeq \end{aligned} \quad (7)$$

where \succeq represents the element-wise comparison, $\|\cdot\|_2$ is the ℓ_2 -norm, and $\|\cdot\|_1$ is the ℓ_1 -norm which is known to promote sparsity. Once the multiple abundance vector \mathbf{r}_i has been estimated, it is normalized in order to reduce the effects of multiplicative factors and satisfy ASC. Following the same approach, further sparsity can be imposed using ℓ_p -norm [31] or reweighted ℓ_1 -approaches [32], [33]. Overall, this sparsity property allows the selection of a smaller number of endmember spectra. However, it may not lead to the selection of a smaller number of endmember classes. Conversely, to promote sparsity on the number of endmember classes, one strategy consists in formulating the unmixing problem through a sparse group lasso [34]

$$\begin{aligned} \min_{\mathbf{r}_i} \quad & \left\{ \frac{1}{2} \left\| \sum_{k=1}^K \mathbf{E}_k (\mathbf{g}_k \odot \mathbf{r}_i) - \mathbf{y}_i \right\|_2^2 \right. \\ & \left. + \lambda_g \sum_{k=1}^K \|\mathbf{g}_k \odot \mathbf{r}_i\|_2 + \lambda_r \|\mathbf{r}_i\|_1 \right\} \\ \text{s.t. } \forall i, \quad & \mathbf{r}_i \succeq 0 \end{aligned} \quad (8)$$

where \mathbf{g}_k is the k th column of \mathbf{G} , \odot is the element-wise product, and thus, $\mathbf{g}_k \odot \mathbf{r}_i$ extracts the elements in \mathbf{r}_i belonging

to the k th class. This approach has the great advantage of promoting sparsity in both the number of endmember spectra and the number of endmember classes. Another strategy relies on the concept of ‘‘social sparsity’’ that can exploit the structure of endmember bundles more explicitly [18]. The method assumes that \mathbf{r}_i can be partitioned into K groups representing each endmember class, leading to the optimization problem

$$\begin{aligned} \min_{\mathbf{r}_i} \quad & \left\{ \frac{1}{2} \|\mathbf{E}\mathbf{r}_i - \mathbf{y}_i\|_2^2 + \lambda_r \left(\sum_{k=1}^K \|\mathbf{g}_k \odot \mathbf{r}_i\|_p^q \right)^{\frac{1}{q}} \right\} \\ \text{s.t. } \forall i, \quad & \forall n, \mathbf{r}_i \succeq 0, \quad \sum_{n=1}^N r_{ni} = 1 \end{aligned} \quad (9)$$

where $\|\cdot\|_p$ is the ℓ_p -norm and r_{ni} is the n th multiple abundance fraction of the i th pixel. Finally, abundances associated each endmember class can be obtained by summing the multiple abundances within each class as in (5). This method can be considered as a generalized model, since by adjusting the values of (p, q) , it boils down to the group lasso (2, 1), the elitist lasso (1, 2), or the fractional case (1, q) where $q \in \mathbb{N}$ and $0 < q < 1$ [18]. The fractional case promotes sparsity of the selection of both endmember classes and spectra within each class. The aforementioned methods that incorporate all endmember bundles can potentially recover pixel-wise endmember variability, at the price of a thorough postprocessing analysis. More precisely, the pixel-wise endmember variability can be estimated by a weighted combination of all possible endmember spectra within each class, e.g., as done in [35].

All of the aforementioned sparsity-based methods still suffer from the following limitations.

- 1) *Interpretation of the Models*: The models are designed to estimate multiple abundance fractions \mathbf{r} corresponding to each spectrum in endmember bundles. Although the models can implicitly consider abundances for each class or endmember spectra for each pixel, they cannot independently describe pixel-wise endmember spectra. In order to generate pixel-wise endmember spectra, a thorough postprocessing analysis is required using a normalization technique [35].
- 2) *Adaptability to Describe Endmember Variability*: ASC imposed on \mathbf{r}_i does not allow a consistent description of the endmembers within each pixel. In addition, it cannot explicitly capture adaptive and hierarchical structure of endmember spectra for each pixel.

To overcome these two shortcomings, this paper capitalizes on this abundant literature to design a new multiple endmember mixing model introduced in Section III.

III. MULTIPLE ENDMEMBER MIXING MODELS

A. MEMM

The proposed model relies on three main ingredients, namely, endmember bundles, bundling coefficients, and abundances. According to this model, each endmember bundle is mixed to provide a suitable and adaptive endmember spectrum used to unmix a given pixel. The proposed multiple endmember mixing model (MEMM) is defined as

$$\mathbf{y}_i = \mathbf{E}\mathbf{B}\mathbf{a}_i + \mathbf{n}_i \quad (10)$$

where $\mathbf{B}_i \in \mathbb{R}^{N \times K}$ gathers so-called bundling coefficients of the i th pixel which decompose the endmember signatures according to the endmember bundles for the considered pixel. To enforce the bundle structure, the bundling coefficients \mathbf{B}_i associated with the pixel is defined as the following block-diagonal matrix:

$$\mathbf{B}_i = \begin{bmatrix} \mathbf{b}_{1i} & \mathbf{0}_{N_1} & \cdots & \mathbf{0}_{N_1} \\ \mathbf{0}_{N_2} & \mathbf{b}_{2i} & \cdots & \mathbf{0}_{N_2} \\ \vdots & \vdots & \ddots & \vdots \\ \mathbf{0}_{N_K} & \mathbf{0}_{N_K} & \cdots & \mathbf{b}_{Ki} \end{bmatrix} \quad (11)$$

where $\mathbf{b}_{ki} \in \mathbb{R}^{N_k \times 1}$ is the bundling coefficients for the k th class at the i th pixel. Each bundling coefficient must be nonnegative, and the bundling vector \mathbf{b}_{ki} is expected to be sparse. Indeed, multiple endmember spectra within each class are usually redundant and only a few endmember spectra within each class should be enough to unmix a pixel. This property can be induced by considering the following bundling constraints:

$$\forall i, \mathbf{B}_i \succeq 0 \quad \text{and} \quad \|\mathbf{B}_i\|_0 = \sum_{k=1}^K \|\mathbf{b}_{ki}\|_0 \leq s \quad (12)$$

where $\|\cdot\|_0$ is the ℓ_0 -norm that counts the number of nonzero elements and s is the maximum number of nonzero elements in \mathbf{B}_i , i.e., the maximum number of endmembers to be used within each class to describe the pixel. The ANC and the ASC are usually imposed. In addition, in this paper, complementary sparsity is imposed on each abundance vector, i.e.,

$$\forall k, \forall i, a_{ki} \geq 0, \quad \text{and} \quad \forall i, \sum_{k=1}^K a_{ki} = 1, \quad \|\mathbf{a}_i\|_0 \leq v \quad (13)$$

where v is the number of endmember classes to be used to decompose the image pixel.

B. MEMMs

The sparsity constraint (12) applied to \mathbf{B} can be slightly modified to obtain another meaningful set of constraints

$$\forall i, \mathbf{B}_i \succeq 0 \quad \text{and} \quad \forall k, \forall i, \|\mathbf{b}_{ki}\|_0 \leq 1. \quad (14)$$

The resulting model, referred to as MEMM_s in what follows, is designed to generate at most one scaled endmember spectrum for each class. Details on the optimization of MEMM_s are available in [36].

C. Relationships Between MEMM and Existing Models

1) *MEMM_s and MESMA*: When the sparsity constraint on the abundances $\|\mathbf{a}_i\|_0 \leq v$ in (13) is not considered, the optimization problem associated with the MEMM_s model described in Section III-B is equivalent to MESMA and sparse MESMA [37]. Unlike MESMA that considers the reconstruction error to determine the optimal combination of endmember classes within each pixel, MEMM_s incorporates the sparsity constraint to select the optimal combination. This prevents a larger number of endmember classes to be selected for each pixel.

2) *MEMM and Pixel-Wise Endmember Variability Models*: By denoting $\tilde{\mathbf{M}}_i = \mathbf{E}\mathbf{B}_i$, the equivalent endmember matrix associated with the i th pixel, MEMM models the observed pixel spectra as

$$\mathbf{y}_i = \tilde{\mathbf{M}}_i \mathbf{a}_i + \mathbf{n}_i \quad (15)$$

where $\tilde{\mathbf{M}}_i$ can be interpreted as a set of K spatially varying endmember spectra. This approach has also been adopted in recent works to incorporate endmember variability as additive factors [22], multiplicative factors [21], [38], or a combination of additive and multiplicative factors [23]. In particular, when $N_1 = \dots, N_k = 1$ in (3), the endmember bundles $\mathbf{E}_1, \dots, \mathbf{E}_K$ are reduced to unique endmember spectra characterizing each class. The associated bundling coefficient matrix $\mathbf{B} = \text{diag}[b_1, \dots, b_K]$ is diagonal, where each coefficient b_k scales the corresponding endmember spectrum \mathbf{E}_k ($k = 1, \dots, K$). Thus, MEMM generalizes the recently introduced extended LMM [21].

However, MEMM is different from the aforementioned methods since it resorts to *a priori* information (i.e., endmember bundles) to model the endmember variability. More precisely, MEMM describes the admissible variability within an endmember class as the convex cone spanned by the corresponding bundles. As a consequence, *per se*, MEMM offers an adaptive description of the spectral variability even when predefined endmember bundles do not completely capture this variability within each class.

Finally, when the existing methods impose ASC onto \mathbf{a}_i , they assume that mixed spectra belong to the simplex spanned by the endmember bundles. In such case, pixel-wise endmember spectra can be generated as convex combinations of spectra within each bundle where the weight of each bundling coefficient corresponds to a normalized abundance of a spectrum in the bundle. However, this is a limited assumption since observed spectra may be outside the simplex, e.g., when affected by variations in illumination. MEMM imposes ASC only onto the abundances \mathbf{a}_i of endmember classes and enables the bundling coefficients \mathbf{B}_i to scale the endmember signature, e.g., to capture variability induced by varying illumination (see experiments in Section VII).

3) *MEMM and Sparsity-Based Unmixing Methods*: By setting $\mathbf{r}_i = \mathbf{B}_i \mathbf{a}_i$, MEMM in (10) can be rewritten as

$$\mathbf{y}_i = \mathbf{E} \mathbf{r}_i + \mathbf{n}_i \quad (16)$$

similar to the existing models discussed in Section II. The main difference is that MEMM enables the multiple abundances \mathbf{r}_i to be decomposed into bundling coefficients \mathbf{B}_i within each class and abundances \mathbf{a}_i , resulting into a bilayer description of the abundances. This hierarchical decomposition has been also adopted by unmixing methods based on multilayer nonnegative matrix factorization [39]. However, each layer induced by MEMM (i.e., bundling matrix and abundance vector) has a clear and meaningful role. In addition, MEMM complements this bilayer hierarchy with a twofold structured, physically-motivated sparsity imposed on the multiple abundance vector \mathbf{r}_i . This bilevel sparsity has the significant advantage of reducing overfitting, and one may expect a

significant improvement of stability and interpretability of the abundance estimates.

The intrinsic structure of MEMM is also similar to the recently developed methods based on robust constrained matrix factorization [40] and kernel archetypoid analysis [41]. These methods model a set $\mathbf{Y} = [\mathbf{y}_1, \dots, \mathbf{y}_P] \in \mathbb{R}^{L \times P}$ of P pixel spectra as

$$\mathbf{Y} = \mathbf{YCA} + \mathbf{N} \quad (17)$$

where $\mathbf{C} \in \mathbb{R}^{P \times K}$ is a matrix gathering a set of coefficients, $\mathbf{A} = [\mathbf{a}_1, \dots, \mathbf{a}_P] \in \mathbb{R}^{K \times P}$ is the abundance matrix and $\mathbf{N} = [\mathbf{n}_1, \dots, \mathbf{n}_P] \in \mathbb{R}^{L \times N}$ is the error and noise matrix. Sparsity (induced by ASC or the use of ℓ_0 -pseudonorm) and nonnegativity constraints are imposed onto each column of \mathbf{C} and \mathbf{YC} can be interpreted as synthetic endmember spectra. These methods use the subset of whole image pixels to generate synthetic endmember spectra that are fixed within each image. On the other hand, MEMM uses the subset of endmember bundles to generate synthetic endmember spectra that may be different for each pixel.

Finally, MEMM is closely related to the fractional lasso-based method that has been recently proposed in [35]. Both methods impose sparsity for the selection of both endmember classes and endmember spectra, in a similar manner proposed in [34]. In addition, they can impose ASC and also recover pixel-wise endmember spectra and abundances for each endmember class. However, there are differences between these two methods.

- 1) MEMM defines bundling coefficients and, thus, can explicitly express pixel-wise endmember spectra $\tilde{\mathbf{M}}_i$ and abundances \mathbf{a}_i for each endmember class in the unmixing model. On the other hand, the method proposed in [35] is designed to estimate multiple abundance fractions \mathbf{r}_i for each possible endmember spectrum among endmember bundles in the unmixing model. Abundances for each endmember class or pixel-wise endmember spectra are subsequently recovered by conducting postprocessing steps. In other words, MEMM estimates the bundling coefficients independently whereas in [35], they are computed from the multiple abundance fractions.
- 2) While the bundling coefficients are only required to be positive in MEMM, both positivity and sum-to-one constraints are imposed by the model introduced in [35]. This leads to a different geometrical interpretation of pixel-wise endmember spectra (see Fig. 1). Pixel-wise endmember spectra derived by MEMM are inside the conic hull defined by each endmember bundle. The conic hull can also be interpreted as scaling factors discussed in [15] and [21] which are important, e.g., to describe varying illumination. On the other hand, the method in [35] generates pixel-wise endmember spectra as the convex combination of endmember spectra within each endmember bundle. Note that, if MEMM considers a sum-to-one constraint for each column in \mathbf{B} , MEMM can also consider the convex hull described by each endmember bundle.

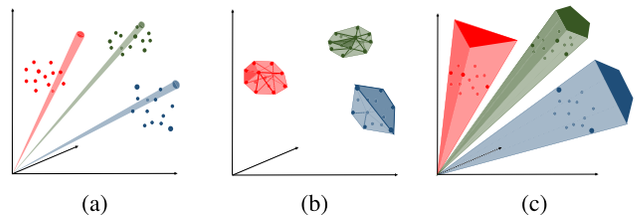


Fig. 1. Geometrical interpretation of potential endmember variability derived by different methods. Different colors represent different endmember classes. Points represent given endmember spectra within each class. Transparent colors represent potential endmember variability generated by the methods. (a) Potential endmember variability described by a line using MEMM_s or MESMA. (b) Potential endmember variability described by convex hull using the fractional lasso-based method or other methods that imposes ASC on \mathbf{r} (see [16]). (c) Potential endmember variability described by convex cone using the proposed MEMM.

IV. MEMM-BASED UNMIXING ALGORITHM

Unmixing according to the proposed MEMM can be formulated as the minimization problem

$$\begin{aligned} \min_{\mathbf{B}_i, \mathbf{a}_i} & \frac{1}{2} \|\mathbf{EB}_i \mathbf{a}_i - \mathbf{y}_i\|_2^2 \\ \text{s.t. } & \forall k, \forall i, \quad a_{ki} \geq 0, \quad \sum_{k=1}^K a_{ki} = 1, \quad \|\mathbf{a}_i\|_0 \leq v, \\ & \mathbf{B}_i \geq 0, \quad \|\mathbf{B}_i\|_0 \leq s. \end{aligned} \quad (18)$$

This minimization problem is similar to the double sparsity-inducing method proposed in [26]. Using an alternative formulation, the minimization problem can be written as the following nonconvex minimization problem:

$$\min_{\mathbf{B}_i, \mathbf{a}_i} \mathcal{J}(\mathbf{B}_i, \mathbf{a}_i) = \{f(\mathbf{B}_i, \mathbf{a}_i) + h(\mathbf{B}_i) + g(\mathbf{a}_i)\} \quad (19)$$

with

$$f(\mathbf{B}_i, \mathbf{a}_i) = \frac{1}{2} \|\mathbf{EB}_i \mathbf{a}_i - \mathbf{y}_i\|_2^2 \quad (20)$$

$$h(\mathbf{B}_i) = \iota_{\mathbb{R}_+}(\mathbf{B}_i) + \lambda_b \|\mathbf{B}_i\|_0 \quad (21)$$

$$g(\mathbf{a}_i) = \iota_{\mathbb{S}}(\mathbf{a}_i) + \lambda_a \|\mathbf{a}_i\|_0 \quad (22)$$

where λ_a and λ_b are parameters which control the balance between the data fitting term and the sparse regularizations, $\iota_C(\mathbf{x})$ is the indicator function on the set C [i.e., $\iota_C(\mathbf{x}) = 0$ when $\mathbf{x} \in C$ whereas $\iota_C(\mathbf{x}) = \infty$ when $\mathbf{x} \notin C$], and \mathbb{S} is the simplex defined by the ASC and ANC. Promoting the sparsity of the solution through penalized regularizations rather than a hard constraint allows the contribution of the sparsity term to be adjusted w.r.t. the data-fitting term rather than directly constraining the maximum number of classes. This flexibility allows different numbers of selected endmember spectra or classes to be selected for each pixel. Solving this optimization problem is challenging since the regularization functions h and g are nonconvex and nonsmooth. However, it can be tackled thanks to the proximal alternating linearized minimization (PALM) [42]. With guarantees to converge to a critical point, PALM iteratively updates the parameters \mathbf{a}_i and \mathbf{B}_i by alternatively minimizing the objective function w.r.t. these

Algorithm 1 Algorithm for MEMM-Based Unmixing

```

1: Input :  $\mathbf{y}_i, \mathbf{E}$ 
2: Initialization:  $\mathbf{a}_i^{(0)}$  and  $\mathbf{B}_i^{(0)}$ .
3: Set  $\mathbf{r}_i^{(0)}$  using an unmixing method (e.g. FCLS).
4:  $\mathbf{a}_i^{(0)} = \mathbf{G}^T \mathbf{r}_i^{(0)}$ 
5:  $\forall_k, \mathbf{b}_{ik}^{(0)} = (\mathbf{g}_k \odot \mathbf{r}_i^{(0)}) \oslash a_{ki}^{(0)}$ 
6: Main procedure:
7: while the stopping criterion is not satisfied do
8:    $\mathbf{b}_i^{(t+1)} \leftarrow \text{prox}_{c_t/\lambda_b}^h(\mathbf{b}_i^{(t)} - \frac{1}{c_t} \nabla_{\mathbf{b}_i} f(\mathbf{B}_i^{(t)}, \mathbf{a}_i^{(t)}))$ 
9:    $\mathbf{B}_i^{(t+1)} = \text{blkdiag}(\mathbf{b}_i^{(t+1)})$ 
10:   $\mathbf{a}_i^{(t+1)} \leftarrow \text{prox}_{d_t/\lambda_a}^g(\mathbf{a}_i^{(t)} - \frac{1}{d_t} \nabla_{\mathbf{a}_i} f(\mathbf{B}_i^{(t+1)}, \mathbf{a}_i^{(t)}))$ 
11: end while
12: Output :  $\mathbf{a}_i^{(t+1)}, \mathbf{B}_i^{(t+1)}$ 

```

parameters, i.e., by solving the following proximal problems:

$$\begin{aligned}
\mathbf{B}_i^{(t+1)} &\in \min_{\mathbf{B}_i} \left\{ h(\mathbf{B}_i) + \langle \mathbf{B}_i - \mathbf{B}_i^{(t)}, \nabla_{\mathbf{B}_i} f(\mathbf{B}_i^{(t)}, \mathbf{a}_i^{(t)}) \rangle \right. \\
&\quad \left. + \frac{c_t}{2} \|\mathbf{B}_i - \mathbf{B}_i^{(t)}\|_2^2 \right\} \\
\mathbf{a}_i^{(t+1)} &\in \min_{\mathbf{a}_i} \left\{ g(\mathbf{a}_i) + \langle \mathbf{a}_i - \mathbf{a}_i^{(t)}, \nabla_{\mathbf{a}_i} f(\mathbf{B}_i^{(t+1)}, \mathbf{a}_i^{(t)}) \rangle \right. \\
&\quad \left. + \frac{d_t}{2} \|\mathbf{a}_i - \mathbf{a}_i^{(t)}\|_2^2 \right\}. \tag{23}
\end{aligned}$$

The pseudocode for MEMM is shown in Algorithm 1, and these two steps are described in what follows.

A. Optimization w.r.t. \mathbf{B}_i

To optimize only w.r.t. the diagonal entries in \mathbf{B}_i , the objective function can be rewritten with the following decomposition:

$$\begin{aligned}
f(\mathbf{b}_i, \mathbf{a}_i) &= \frac{1}{2} \|\mathbf{U}_i \mathbf{b}_i - \mathbf{y}_i\|_2^2 \\
h(\mathbf{b}_i) &= \iota_{\mathbb{R}_+}(\mathbf{b}_i) + \lambda_b \|\mathbf{b}_i\|_0 \tag{24}
\end{aligned}$$

where

$$\begin{aligned}
\mathbf{U}_i &= [\mathbf{E}_1 a_{1i} | \cdots | \mathbf{E}_K a_{Ki}] \\
\mathbf{b}_i &= [\mathbf{b}_{1i}^T, \mathbf{b}_{2i}^T, \cdots, \mathbf{b}_{Ki}^T]^T.
\end{aligned}$$

This leads to the following updating rule:

$$\min_{\mathbf{b}_i} \left\{ h(\mathbf{b}_i) + \frac{c_t}{2} \left\| \mathbf{b}_i - \left(\mathbf{b}_i^{(t)} - \frac{1}{c_t} \nabla_{\mathbf{b}_i} f(\mathbf{B}_i^{(t)}, \mathbf{a}_i^{(t)}) \right) \right\|_2^2 \right\}$$

where $\nabla_{\mathbf{b}_i} f(\mathbf{B}_i^{(t)}, \mathbf{a}_i^{(t)}) = \mathbf{U}_i^T (\mathbf{U}_i \mathbf{b}_i - \mathbf{y}_i)$. Using similar computations as in [42], this can be conducted as

$$\mathbf{b}_i^{(t+1)} \in \text{prox}_{c_t/\lambda_b}^h \left(\mathbf{b}_i^{(t)} - \frac{1}{c_t} \nabla_{\mathbf{b}_i} f(\mathbf{B}_i^{(t)}, \mathbf{a}_i^{(t)}) \right) \tag{25}$$

where $c_t = \gamma_m \|\mathbf{U}_i^T \mathbf{U}_i\|_F$ represents a step size for each iteration. The proximal operator associated with f can be computed using the approach [42]. Finally, the bundling matrix \mathbf{B}_i can be reconstructed as $\mathbf{B}_i = \text{blkdiag}(\mathbf{b}_i)$, where $\text{blkdiag}(\cdot)$ generates the block diagonal matrix \mathbf{B}_i from the vector \mathbf{b}_i .

B. Optimization With Respect to \mathbf{a}_i

To optimize w.r.t. \mathbf{a}_i , the objective function can be rewritten using the decomposition

$$f(\mathbf{B}_i, \mathbf{a}_i) = \frac{1}{2} \|\tilde{\mathbf{M}}_i \mathbf{a}_i - \mathbf{y}_i\|_2^2 \tag{26}$$

$$g(\mathbf{a}_i) = \iota_{\mathbb{S}}(\mathbf{a}_i) + \lambda_a \|\mathbf{a}_i\|_0 \tag{27}$$

where $\tilde{\mathbf{M}}_i = \mathbf{E} \mathbf{B}_i$. Thus, updating the abundance vector can be formulated as

$$\min_{\mathbf{a}_i} \left\{ g(\mathbf{a}_i) + \frac{d_t}{2} \left\| \mathbf{a}_i - \left(\mathbf{a}_i^{(t)} - \frac{1}{d_t} \nabla_{\mathbf{a}_i} f(\mathbf{B}_i^{(t+1)}, \mathbf{a}_i^{(t)}) \right) \right\|_2^2 \right\}$$

where $\nabla_{\mathbf{a}_i} f(\mathbf{B}_i^{(t+1)}, \mathbf{a}_i^{(t)}) = \tilde{\mathbf{M}}_i^T (\tilde{\mathbf{M}}_i \mathbf{a}_i - \mathbf{y}_i)$. Using the proximal operator, this can be written as

$$\mathbf{a}_i^{(t+1)} \in \text{prox}_{d_t/\lambda_a}^g \left(\mathbf{a}_i^{(t)} - \frac{1}{d_t} \nabla_{\mathbf{a}_i} f(\mathbf{B}_i^{(t+1)}, \mathbf{a}_i^{(t)}) \right) \tag{28}$$

where $d_t = \gamma_a \|\tilde{\mathbf{M}}_i^T \tilde{\mathbf{M}}_i\|_F$ represents a step size for each iteration. Moreover the proximal mapping associated with g can be performed using the method developed in [43]. Details on the computation of these proximal operators are available in [36].

C. Initialization and Stopping Rule

MEMM requires initial estimates \mathbf{a}_i^0 and \mathbf{B}_i^0 of the abundance vector and bundling matrix, respectively. To do so, first, MEMM estimates a multiple abundance vector $\mathbf{r}_i^{(0)}$ using the state-of-the-art LMM-based unmixing method [e.g., fully constrained least squares (FCLS), see line 3]. Then, an initial estimate of the single abundance vector $\mathbf{a}_i^{(0)}$ is computed according to (5) (see line 4). Finally, the bundling matrix $\mathbf{B}_i^{(0)}$ is arbitrarily initialized as the corresponding scaling factor (see line 5, where \oslash stands for the element-wise division). Once initial estimates have been obtained, $\mathbf{a}_i^{(t+1)}$ and $\mathbf{b}^{(t+1)}$ are iteratively updated in lines 8–10. The algorithm stops when the difference between updated and previous values of the objective function $f(\mathbf{B}_i, \mathbf{a}_i)$ is smaller than a predetermined threshold.

V. EXPERIMENTS USING SIMULATED DATA

First, the relevance of the proposed MEMM and its variant MEMM_s has been evaluated, thanks to experiments conducted on simulated data sets.

A. Generation of Bundles

First, $K = 10$ spectra were selected from the USGS spectral library. The ten spectra were chosen so that the minimum angle between any two spectra was larger than 5° . This pruning prevented synthetic endmember bundles to overlap each other. Second, endmember bundles \mathbf{E}_k ($k = 1, \dots, K$) were designed by randomly generated $N_k = 30$ endmember spectra for each endmember bundle using the approach proposed in [44]. These bundles, depicted in Fig. 2, were used for generating the following two simulated data.

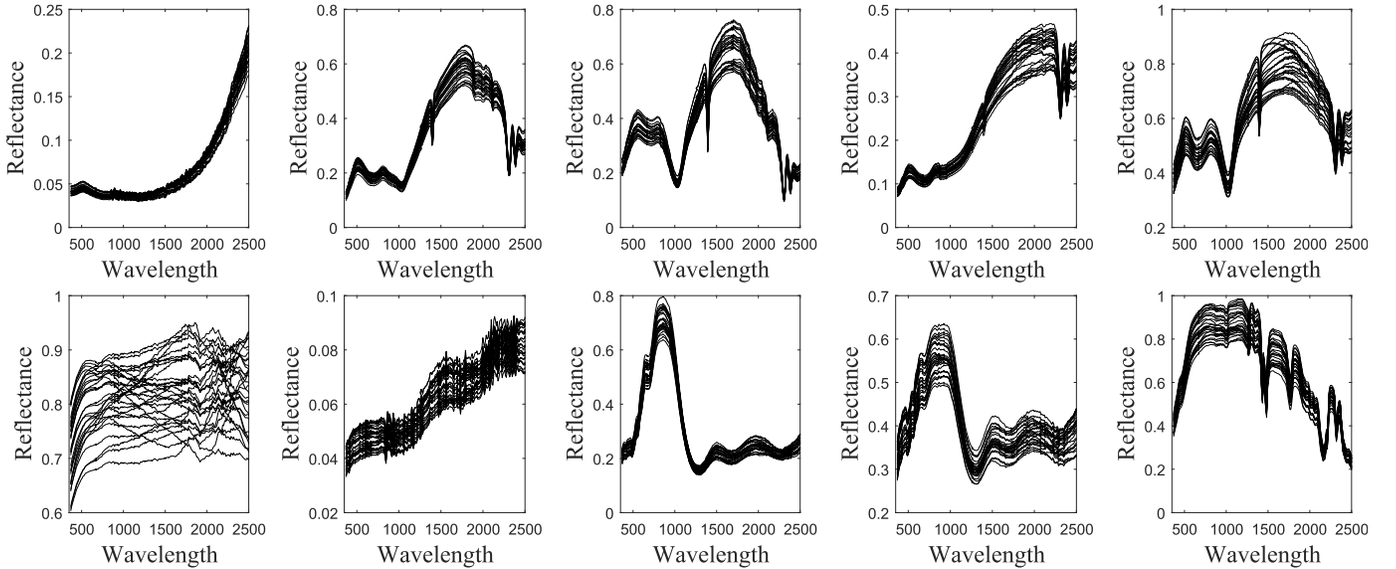


Fig. 2. Synthetically generated endmember bundles.

1) *Simulated Data Set 1 (SIM1)*: The first data set, referred to as SIM1 in what follows, was generated using MESMA. A mixed spectrum was generated using the following five steps. First, the number of endmember classes K was randomly determined in the set $(\{1, \dots, 5\})$. Using weighted random number, a smaller number is more likely to occur in the set. Second, a random combination of endmember classes was selected. Third, one spectrum within each selected endmember class was randomly chosen. Fourth, the abundances of the selected endmember classes were randomly generated using a Dirichlet distribution to jointly ensure ANC and ASC. Finally, a mixed spectrum was generated by a linear combination of endmember spectra of the selected endmember classes and the randomly generated abundances. A set of $P = 100$ mixed spectra were generated in this paper. Different amounts of additive Gaussian noise with corresponding signal-to-noise ratios (SNRs) of 50, 40, and 30 dB were considered to the mixed spectra. A complementary scenario with lower SNR (20 dB) is considered in [36].

2) *Simulated Data Set 2 (SIM2)*: SIM2 was generated using MEMM. A mixed spectrum was generated similarly as done in SIM1. The main difference is the bundling coefficients in MEMM. In order to generate the bundling coefficients, the number of spectra N_k was randomly chosen in the set $(\{1, \dots, 5\})$. The bundling coefficients of the randomly selected spectra were generated from a Dirichlet distribution. A mixed spectrum was generated by a linear combination of spectra, the bundling coefficients, and the abundances. As for SIM1, $P = 100$ mixed spectra were generated and different amounts of Gaussian noise were also added to SIM2.

3) *Simulated Data Set 3 (SIM3)*: For SIM3, each pixel is generated by selecting and then scaling a unique endmember spectrum belonging to a randomly chosen endmember class. More precisely, the observed pixel spectra \mathbf{y}_i of SIM 3 can be written as $\mathbf{y}_i = c_i \mathbf{e}_{k_i, n_i}$, where \mathbf{e}_{k_i, n_i} is the endmember spectrum defining the k th class and $c_i, k_i,$ and n_i are randomly

and uniformly drawn in the sets $(0.8, 1)$, $\{1, \dots, K\}$, and $\{1, \dots, N_k\}$, respectively. A set of $P = 100$ pixels were generated, and different levels of Gaussian noise were considered. These pure pixel spectra may be mistakenly unmixed by multiple endmember classes if endmember variability affected by the scaling factor is not correctly described by the considered model. To correctly select a single endmember, strong sparsity needs to be imposed while incorporating the scaling factor.

B. Compared Methods

MEMM and MEMM_s were compared with other five methods that incorporate endmember bundles and promote sparsity: FCLS [45], sparse unmixing by variable splitting and augmented Lagrangian (SUnSAL) [46], alternating angle minimization (AAM) [47], and methods based on group lasso, elitist lasso, and fractional lasso [18], [35]. Note that, FCLS and SUnSAL can incorporate endmember bundles by considering the model in (4). Abundances estimated by SUnSAL were normalized in order to be robust to the scaling factor caused by illumination variations as discussed in [15] and [21]. FCLS and SUnSAL were chosen for comparing the proposed methods with most widely used unmixing methods that promote sparsity. AAM was selected for comparison because it is a latest variant of MESMA and it is more computationally efficient than MESMA while achieving good performance. Group lasso, elitist lasso, and fractional lasso were also included for comparison because they can incorporate the structure of endmember bundles as well as MEMM. Note that, the fractional lasso is defined by using the $\ell_{1,q}$ -norm with $q = 0.1$ to impose greater sparsity for the selection of endmember classes, trying to mimic the $\ell_{1,0}$ -pseudo norm as discussed in [35]. MEMM, MEMM_s, and the methods based on group lasso, elitist lasso, and fractional lasso require initial estimates of abundances. In order to fairly compare methods, this paper used abundances estimated by FCLS for the initial estimates required for the methods.

SUnSAL, the methods based on group lasso, elitist lasso, and fractional lasso, and MEMM_s require a parameter λ_r or λ_a controlling sparsity regularization. MEMM requires two parameters λ_b and λ_a . In order to fairly compare the methods, these parameters were empirically determined in the set (0.0001, 0.001, 0.01, 0.1, 1, 5) for each simulated data so that the selected values produced the highest signal-to-reconstruction error (SRE_a). Parameter sensitivity of the proposed method is discussed in [36]. Finally, computation time was also discussed.

C. Performance Criteria

The main objective of this experiment was to assess the algorithm performance while selecting a combination of endmember classes and spectra, and estimating abundances corresponding to endmember classes and spectra. Three criteria were chosen for quantitative validation of the methods. To evaluate the quality of the reconstruction, one defines the SRE per endmember class and per endmember spectrum as [48]

$$\begin{aligned} \text{SRE}_a &\equiv \mathbb{E}[\|\mathbf{a}\|_2^2] / \mathbb{E}[\|\mathbf{a} - \hat{\mathbf{a}}\|_2^2] \\ \text{SRE}_r &\equiv \mathbb{E}[\|\mathbf{r}\|_2^2] / \mathbb{E}[\|\mathbf{r} - \hat{\mathbf{r}}\|_2^2] \end{aligned} \quad (29)$$

where \mathbf{a} and $\hat{\mathbf{a}}$ are the actual and estimated abundance vectors of all pixels, and \mathbf{r} and $\hat{\mathbf{r}}$ are the actual and estimated multiple abundance vectors of all pixels.

Second, the number of nonzero abundances were used to evaluate the sparsity level per endmember class and per endmember spectrum recovered by the methods (see [37], [48]). In this paper, normalized sparsity level (nSL) is defined as

$$\begin{aligned} \text{nSL}_a &\equiv \frac{1}{P} \sum_{i=1}^P \frac{\|\hat{\mathbf{a}}_i\|_0}{\|\mathbf{a}_i\|_0} \\ \text{nSL}_r &\equiv \frac{1}{P} \sum_{i=1}^P \frac{\|\hat{\mathbf{r}}_i\|_0}{\|\mathbf{r}_i\|_0}. \end{aligned} \quad (30)$$

As in [48], abundances smaller than 10^{-4} were considered as zero abundances. When the number of nonzero values of estimated abundances is equivalent to that of actual abundances, the nSL value becomes 1.

Finally, to validate the performance in selecting a relevant combination of endmember classes or spectra, one defines the distance between the two actual and estimated supports (DIST) [37], [49]

$$\begin{aligned} \text{DIST}_a &\equiv \frac{1}{P} \sum_{i=1}^P \frac{\max(|\mathcal{S}_i^a|, |\hat{\mathcal{S}}_i^a|) - |\mathcal{S}_i^a \cap \hat{\mathcal{S}}_i^a|}{\max(|\mathcal{S}_i^a|, |\hat{\mathcal{S}}_i^a|)} \\ \text{DIST}_r &\equiv \frac{1}{P} \sum_{i=1}^P \frac{\max(|\mathcal{S}_i^r|, |\hat{\mathcal{S}}_i^r|) - |\mathcal{S}_i^r \cap \hat{\mathcal{S}}_i^r|}{\max(|\mathcal{S}_i^r|, |\hat{\mathcal{S}}_i^r|)} \end{aligned} \quad (31)$$

where \mathcal{S} and $\hat{\mathcal{S}}$ are true and estimated support sets (i.e., indexes of nonzero values), $|\mathcal{S}|$ represents the total number of elements in the set \mathcal{S} , and \cap stands for the intersection operator. The figures of DIST_a and DIST_r evaluated the distance between two supports of endmember classes and the distance between two supports of endmember spectra, respectively.

TABLE I
SRE PER ENDMEMBER CLASS (SRE_a). THE SYMBOL (∞)
SHOWS PERFECT RECOVERY

	SNR	FCLS	AAM	SUnSAL	Group	Elitist	Fractional	MEMM _s	MEMM
SIM1	30dB	18.0507	18.6585	19.0019	19.4215	19.2333	19.4756	14.5739	18.6903
	40dB	22.9074	26.1146	22.9242	23.589	23.303	24.0219	14.8229	23.3532
	50dB	27.358	34.4346	27.5203	27.4184	26.898	27.6176	14.1579	28.0017
SIM2	30dB	16.1865	15.9426	16.6538	18.0736	16.4314	17.5376	12.1614	16.4785
	40dB	21.6942	20.2125	21.6339	22.0263	21.3809	24.128	13.6682	22.1381
	50dB	25.6617	22.2896	25.3859	25.9215	24.7702	26.4982	14.9405	26.5442
SIM3	30dB	14.5914	14.2696	14.5758	14.3931	14.3215	19.0764	14.1404	25.6203
	40dB	15.1873	14.6965	15.1079	14.8343	14.7525	16.9308	14.9971	∞
	50dB	15.7588	14.8567	15.6163	15.204	15.1532	17.2874	16.4725	∞

TABLE II
SRE PER ENDMEMBER SPECTRUM (SRE_r)

	SNR	FCLS	AAM	SUnSAL	Group	Elitist	Fractional	MEMM _s	MEMM
SIM1	30dB	1.1752	2.2584	1.7155	2.0398	1.5188	1.3071	-0.457	0.7697
	40dB	2.8611	5.0035	3.0215	3.0357	2.9957	2.9239	0.5171	2.7653
	50dB	3.6312	13.9842	3.823	3.7591	3.7576	3.649	0.17	3.5721
SIM2	30dB	1.0084	-0.1527	1.4259	1.8935	1.3043	1.1533	-2.0312	0.432
	40dB	2.3323	1.5198	2.4934	2.5741	2.48	2.4092	-2.1441	2.2462
	50dB	3.2175	3.4488	3.3204	3.3266	3.3666	3.2448	-3.0151	3.2304
SIM3	30dB	1.3047	0.7371	1.4879	1.5672	1.4417	1.0397	-1.8937	1.2851
	40dB	2.5007	1.2622	2.0889	2.5154	2.5172	2.3415	0.3956	2.5555
	50dB	3.1294	1.6732	3.215	3.095	3.1056	2.8264	-1.5401	3.3266

D. Results

SRE per class was calculated for each method and is reported in Table I. For SIM1 with 50 dB, AAM performed best among all methods. The performance of AAM, however, was degraded as SNR became lower. For data with 30 dB, the results derived from AAM were worse than those derived from other methods except MEMM_s. In addition, AAM performed poorly compared with other methods in SIM2. This showed that the MESMA-based approach (AAM) was less effective when given endmember bundles did not completely represent endmember variability present in the data and SNR of the data was low (<40 dB). This finding was also observed in [19]. MEMM produced better results for SIM2 with 50 dB than the other sparsity-methods and produced comparable results with 40 and 30 dB. MEMM_s performed poorly, compared with all methods. In SIM3, MEMM outperformed other methods as expected. This showed that MEMM could successfully select the correct support and describe the variability caused by scaling effects while other methods failed to describe the variability. SRE per spectrum was also calculated from each method and is reported in Table II. Compared with SRE per class, SRE per spectrum was very low for all methods. This showed that the exact recovery of multiple abundances \mathbf{r}_i was challenging under conditions where a large number of endmember spectra were present within each class.

The nSL in \mathbf{a}_i and \mathbf{r}_i is shown in Tables III and IV, respectively. For the number of nonzero abundances per class, the nSL_a values of MEMM were closer to 1 and performed best among all methods for SIM1, SIM2, and SIM3. This showed that the sparse constraint (the ℓ_0 norm) used in MEMM successfully imposed greater sparsity and led to the selection of smaller numbers of endmember classes. For the number of nonzero abundances per spectrum, AAM and MEMM_s produced the smaller number of nonzero abundances than other methods. This was because these methods selected

TABLE III
NSL PER ENDMEMBER CLASS (NSL_d)

	SNR	FCLS	AAM	SUnSAL	Group	Elitist	Fractional	MEMM _s	MEMM
SIM1	30dB	2.978	2.5045	1.8783	2.8432	4.6737	2.466	1.7205	0.9805
	40dB	2.8242	2.4623	2.247	2.6945	3.6512	2.0942	2.0705	1.1143
	50dB	2.5507	2.303	1.5393	2.6948	3.0268	1.772	2.0113	1.0185
SIM2	30dB	2.6933	2.3205	1.7102	2.5335	3.8612	2.0927	2.1673	0.9713
	40dB	2.7498	2.2693	2.0768	2.6227	3.073	1.7193	2.1525	1.1545
	50dB	2.619	2.1415	2.5373	2.458	3.0388	1.652	1.8788	1.0363
SIM3	30dB	4.44	3.42	2.85	3.67	7.2	5.58	2.91	1.02
	40dB	4.25	3.06	2.56	3.41	6.71	4.4	2.73	1
	50dB	3.9	2.91	2.55	5.39	6.53	4.13	1.92	1

TABLE IV
NSL PER ENDMEMBER SPECTRUM (NSL_r)

	SNR	FCLS	AAM	SUnSAL	Group	Elitist	Fractional	MEMM _s	MEMM
SIM1	30dB	6.527	2.5045	5.8797	22.1282	6.6413	14.545	1.7105	2.851
	40dB	8.3438	2.4623	7.0323	15.2065	9.0737	14.862	2.0052	4.622
	50dB	9.7863	2.303	7.6457	12.1585	11.1332	16.2827	1.9468	5.4307
SIM2	30dB	4.7619	1.7748	4.5617	20.7952	5.091	10.6147	1.6059	1.7862
	40dB	6.2133	1.6638	5.3686	13.4752	6.9975	11.7411	1.5778	3.4925
	50dB	7.5883	1.5571	7.3145	17.579	8.5828	12.8249	1.3488	5.1818
SIM3	30dB	10.79	3.42	9.88	38.16	11.7	43.24	2.86	2.26
	40dB	12.07	3.06	11.82	19.94	13.13	22.42	2.65	2.2
	50dB	14.26	2.91	11.33	15.7	14.95	21.71	1.86	7.88

TABLE V
DISTANCE BETWEEN ACTUAL AND ESTIMATED SUPPORTS
PER ENDMEMBER CLASS (DIST_d)

	SNR	FCLS	AAM	SUnSAL	Group	Elitist	Fractional	MEMM _s	MEMM
SIM1	30dB	0.5893	0.5465	0.3939	0.5866	0.6967	0.4816	0.3491	0.1195
	40dB	0.5689	0.5195	0.4616	0.5594	0.6368	0.3734	0.4498	0.1265
	50dB	0.5137	0.4804	0.268	0.5479	0.5795	0.3114	0.418	0.0758
SIM2	30dB	0.5725	0.5195	0.3775	0.5487	0.6538	0.4119	0.5025	0.1843
	40dB	0.5789	0.4797	0.4521	0.5534	0.5952	0.302	0.4938	0.162
	50dB	0.5345	0.4686	0.5247	0.5185	0.5825	0.2995	0.4179	0.1033
SIM3	30dB	0.7592	0.665	0.5783	0.6885	0.8457	0.7747	0.6	0.01
	40dB	0.7486	0.6205	0.537	0.672	0.8237	0.7198	0.5772	0
	50dB	0.7254	0.6253	0.553	0.7712	0.8149	0.6633	0.3767	0

TABLE VI
DISTANCE BETWEEN ACTUAL AND ESTIMATED SUPPORTS
PER ENDMEMBER SPECTRUM (DIST_r)

	SNR	FCLS	AAM	SUnSAL	Group	Elitist	Fractional	MEMM _s	MEMM
SIM1	30dB	0.9167	0.7651	0.853	0.94	0.8963	0.936	0.7493	0.8115
	40dB	0.8914	0.6721	0.8587	0.9137	0.8881	0.9215	0.7351	0.774
	50dB	0.8649	0.555	0.7831	0.8971	0.8911	0.9294	0.6586	0.7431
SIM2	30dB	0.9039	0.8389	0.8476	0.9332	0.8915	0.9094	0.8729	0.8142
	40dB	0.8791	0.7618	0.8525	0.9167	0.883	0.9088	0.8175	0.7993
	50dB	0.8738	0.6949	0.8696	0.9172	0.8816	0.9067	0.7277	0.8095
SIM3	30dB	0.9283	0.8332	0.9049	0.9668	0.9297	0.9649	0.851	0.7152
	40dB	0.9133	0.7922	0.8945	0.9377	0.9154	0.9488	0.785	0.5645
	50dB	0.918	0.7698	0.8901	0.9222	0.9191	0.9481	0.5783	0.6955

at most one spectrum within each class and enforced greater sparsity when selecting endmember spectra.

The errors in the support sets are shown in Tables V and VI. When estimating the support set per class, MEMM outperformed other methods for all data sets. Especially in SIM3 where a single class is present for each pixel, MEMM could perfectly recover the support in the data set for SNR levels of 50 and 40 dB. This showed that the double sparsity imposed by MEMM also led to the appropriate selection of the combination of endmember classes for each pixel. The performance of MEMM, however, was degraded when estimating the support set per spectrum. Other methods also performed poorly for the support set per spectrum. This shows

TABLE VII
COMPUTATION TIME FOR UNMIXING $P = 100$ PIXELS USING
ENDMEMBER BUNDLES OF $N = 300$ ENDMEMBER SPECTRA

Data	FCLS	AAM	SUNSAL	Group	Elitist	Fractional	MEMM _s	MEMM
SIM1	0.3006	517.7361	0.155	0.9517	1.2694	1.2801	0.4846	2.6625
SIM2	0.1966	549.1244	0.2616	0.8426	1.2122	1.2356	2.2311	2.5705
SIM3	0.1341	391.1602	0.1089	0.6244	0.8745	1.2769	1.8172	1.3913

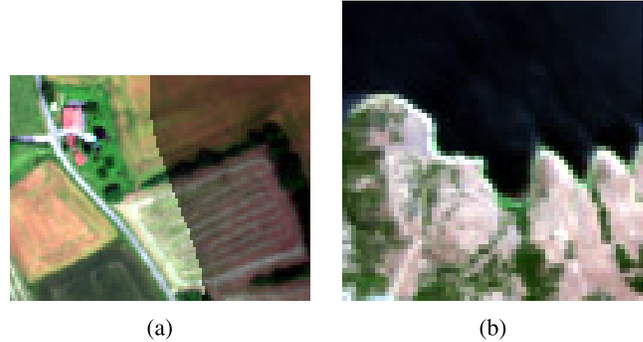


Fig. 3. Real hyperspectral images. (a) MUESLI image. (b) AVIRIS image.

that when multiple highly correlated endmember spectra are present within each class, the existing methods experience difficulty to select an optimal combination of endmember spectra.

These results also showed that better estimates in terms of extracted support (reflected by the DIST or SL values) did not necessarily lead to better SRE values. This is because even if the support and number of nonzero values are correctly recovered, estimated abundances in the correct support may be different from the actual ones. This effect may result from the auxiliary penalizations and constraints inherent to different methods (sum-to-one, nonnegativity, and various formulations of the structured sparsity).

Finally, the computation times of all methods are shown in Table VII. MEMM was more computationally expensive than FCLS, SUnSAL, group lasso, elitist lasso, and fractional lasso. The proposed methods, however, were computationally cheaper than AAM because they did not need to test a large number of combinations of endmember spectra.

VI. EXPERIMENTS USING REAL DATA

A. Description of Real Hyperspectral Images

The methods were finally compared on two real hyperspectral images. The first hyperspectral image was acquired in June 2016, over the city of Saint-André, France, during the MUESLI airborne acquisition campaign. The image was composed of 415 spectral bands. The spectral bands affected by noise (between 1.34–1.55 and 1.80–1.98 μm) were removed, leading to $L = 345$ spectral bands. In the image scene, spatially discrete objects were present. In this paper, each spatially discrete region was assumed to be composed of a single endmember class. Endmember bundles were extracted from each region. As a consequence, large amounts of endmember variability were expected to be present within each

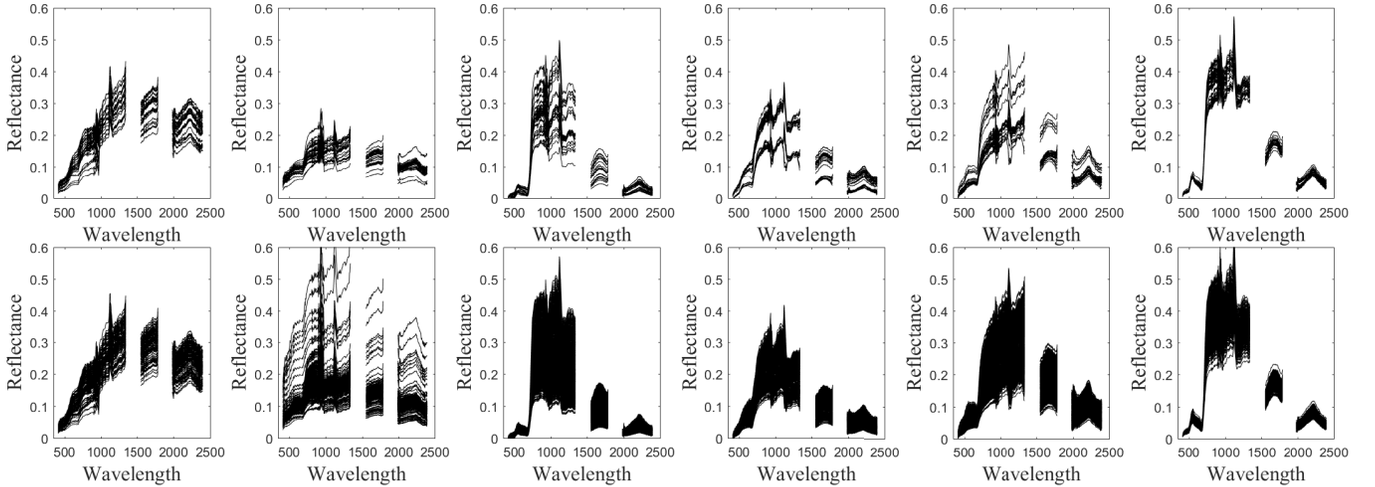


Fig. 4. (First row) Endmember bundles used for unmixing the MUESLI image. (Second row) Endmember bundles generated by MEMM.

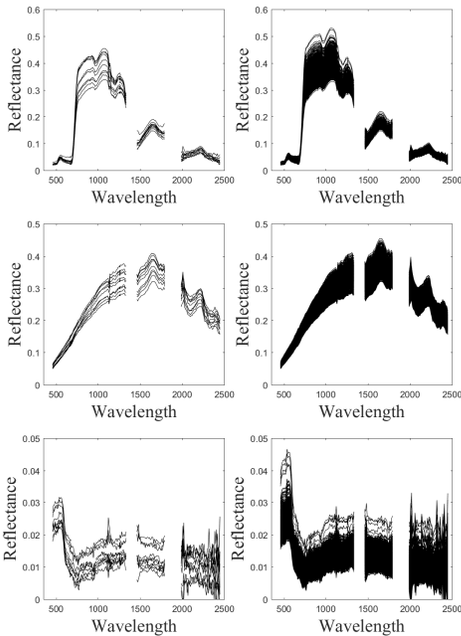


Fig. 5. (First column) Endmember bundles used for unmixing the AVIRIS image. (Second column) Endmember bundles generated by MEMM.

spatially discrete region associated with a particular class and mixed pixels were expected to be located in the boundary of these spatially discrete regions. Moreover, the scene of interest was composed of two flight lines under significantly different illumination conditions, as shown in Fig. 3(a). Thus, this image (referred to as MUESLI image) was used to evaluate whether the methods could accurately estimate abundances when large amounts of endmember variability were present. From this image, $K = 6$ endmember bundles composed of a total of $N = 180$ spectral signatures representing spatially discrete objects were extracted using the n -dimensional visualizer provided by the ENVI software. Endmember bundle extraction was done as follows. First, the N -dimensional visualizer was used to identify and extract pixels located near

the vertices of the point clouds, as also done by state-of-the-art methods [13], [27]. Multiple pixels near a particular vertex were considered as an endmember bundle. Finally, noisy pixels were removed from each endmember bundle. These bundles are represented in Fig. 4(top). Some endmember classes present in the studied area were affected by different illumination conditions. Unlike simulated data, ground truth was not available. Estimated abundances were qualitatively validated by visual inspection of the abundance maps.

The second image was acquired over Moffett Field, CA, USA, by the Airborne Visible/Infrared Imaging Spectrometer (AVIRIS). The image, depicted in Fig. 3(b), initially comprised 224 spectral bands. After the noisy spectral bands were removed, $L = 178$ spectral bands remained. The area of interest, composed of a lake and a vegetated coastal area, was considered in many previous studies, e.g., to assess the performance of unmixing methods. Thus, this second image (referred to as AVIRIS image) was used to test whether the proposed method could perform at least as well as the existing methods to analyze this widely used image scene. As for the AVIRIS image, $K = 3$ endmember bundles were extracted using the ENVI software and qualitative validation was conducted because of the lack of ground truth. The extracted bundles are depicted in Fig. 5(left).

B. Results

For both images, the proposed method was compared with FCLS, SUnSAL, AAM, and the methods based on group lasso, elitist lasso, and fractional lasso. The parameters (λ_r , λ_b and λ_a) required for the methods were empirically determined by qualitatively evaluating abundances derived from different values of the parameters. Finally, synthetic endmember bundles generated by MEMM were compared with endmember bundles initially used for unmixing.

Abundance maps estimated by the eight methods on the MUESLI image are depicted in Fig. 6. These maps show that the abundances estimated by MEMM and fractional lasso were more consistent at the boundary affected by different

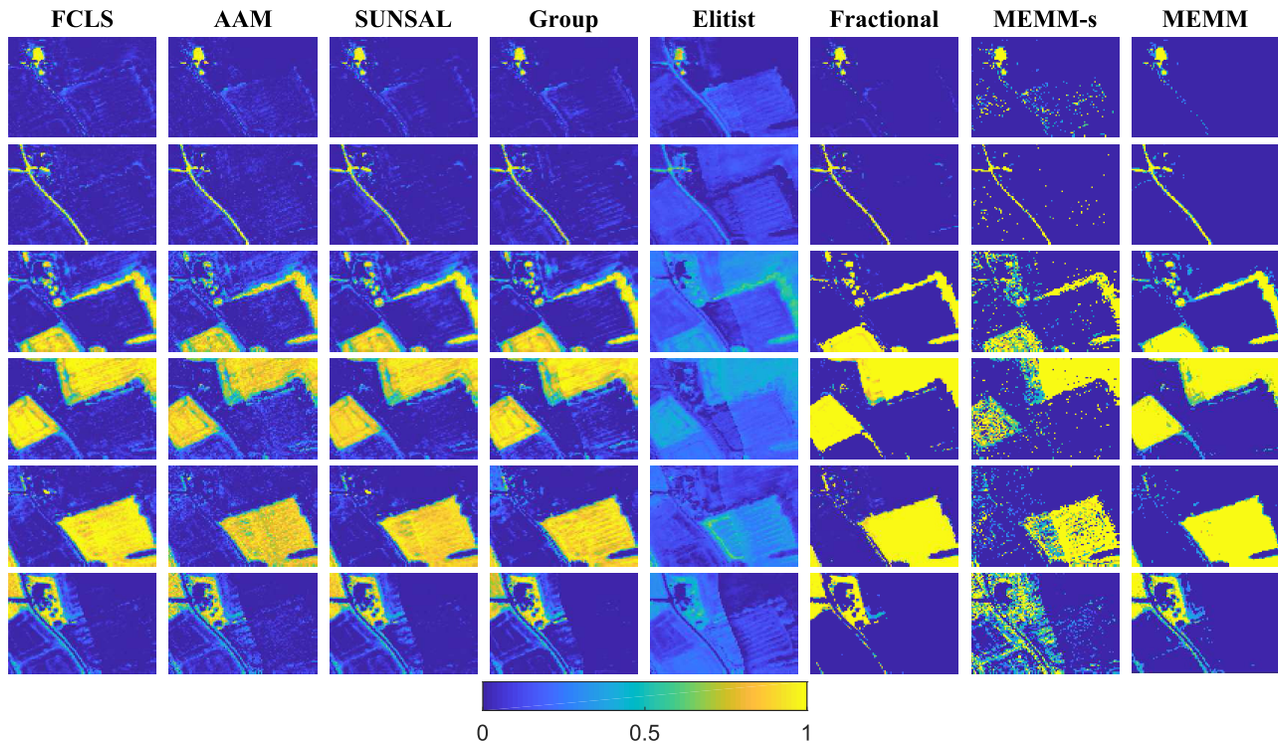


Fig. 6. MUESLI image: estimated abundance maps. (From top to bottom) Building, road, shrub, crop land 1, crop land 2, and grass.

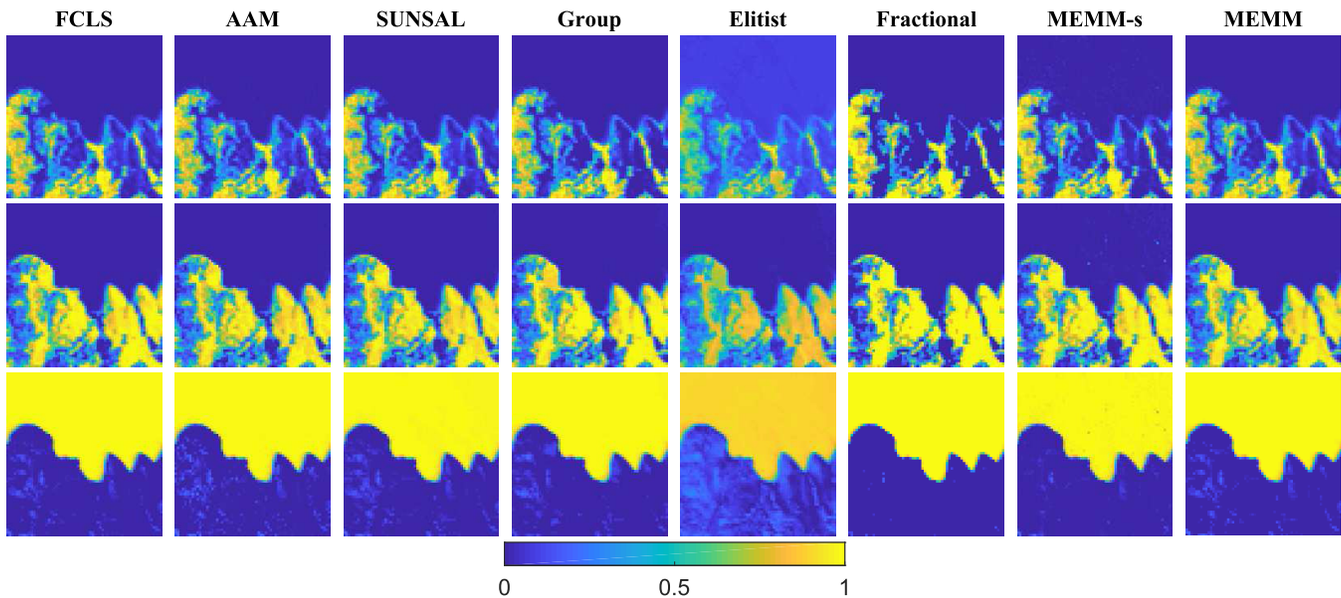


Fig. 7. AVIRIS image: abundance maps. (From top to bottom) Vegetation, soil, and water.

illumination conditions. This showed that MEMM and fractional lasso were more robust to different illumination conditions than concurrent methods. The abundances estimated by MEMM and fractional lasso were also high for each endmember class and showed less noisy. This suggested that MEMM and fractional lasso also promoted greater sparsity than other methods.

Fig. 7 shows the abundance maps estimated for the AVIRIS image. All methods except elitist lasso generated similar abundances. Abundances estimated by elitist lasso were different because it was designed to use a larger number

endmember classes for unmixing each pixel. MEMM produced similar abundances when compared with FCLS, group lasso. This showed that MEMM could perform at least as well as other sparsity-based methods to unmix this well-studied test site. MEMM_s, however, generated more noisy abundance maps. This showed that the initial estimates of abundances used in MEMM_s did not lead to an optimal combination of endmember spectra and the optimal abundances. Although MEMM_s may be more restrictive than MEMM, it may be important when at most one endmember spectrum from each bundle needs to be selected for each pixel. The performance

of MEMM_s, largely depends on initial estimates of multiple abundances \mathbf{r} . More accurate initial estimates are provided; better performance can be achieved. Therefore, MEMM_s may achieve comparable performance in much shorter computation time than the MESMA-based methods if accurate initial abundances \mathbf{r} can be estimated. This new insight is important because most existing methods focus on the development of efficient search of a combination of endmember spectra [29], [47] or the development of spectral library pruning methods to reduce computational complexity [50].

Finally, the endmember bundles recovered by MEMM were compared with endmember bundles initially used to unmix both hyperspectral images [see Figs. 4 and 5]. In Fig. 4, the synthetic endmember bundles filled the gaps that were present in the original endmember bundles. The extended endmember bundles showed more detailed spectral variability within each class in terms of both spectrum amplitudes and shapes also in Fig. 5. This enabled MEMM to generate adaptive endmember spectra within each pixel and estimate more accurate abundances even when initial endmember bundles did not completely represent endmember variability.

VII. CONCLUSION

This paper proposed a MEMM that bridges the gap between endmember bundle-based method and data driven-based methods. MEMM appeared to be superior to the existing methods as follows: 1) it had explicit physical meaning and independently describe pixel-wise endmember spectra and abundances of each class and 2) it independently imposed double sparsity for the selection of both endmember classes and endmember spectra rather than considering structured sparsity. MEMMs were tested and compared to the state-of-the-art methods using simulated data and real hyperspectral images. MEMM showed comparable results for estimating abundances while it outperformed other methods in terms of selecting a set of endmember classes within each pixel. This paper deeply focused on sparsity constraints for both bundling coefficients and abundances. However, other constraints (e.g., spatial constraints) can be easily incorporated in the proposed unmixing framework. Future works also include the Bayesian interpretation and identifiability of the proposed model. Indeed, as most of nonconvex problems, multiple solutions may arise from this multilayer matrix factorization task in the case of overlaps between convex cones spanned by the endmember classes.

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