Nonlinear unmixing of hyperspectral data with vector-valued kernel functions

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Abstract

This paper presents a kernel based nonlinear mixing model for hyperspectral data where the nonlinear function belongs to a Hilbert space of vector valued functions. The proposed model extends existing ones by accounting for band dependent and neighboring nonlinear contributions. The key idea is to work under the assumption that nonlinear contributions are dominant in some parts of the spectrum while they are less pronounced in other parts. In addition to this, we motivate the need for taking into account nonlinear contributions originating from the ground covers of neighboring pixels by practical considerations, precisely the adjacency effect. The relevance of the proposed model is that the nonlinear function is associated to a matrix valued kernel that allows to jointly model a wide range of nonlinearities and include prior information regarding band dependencies. Furthermore, the choice of the nonlinear function input allows to incorporate neighboring effects. The optimization problem is strictly convex and the corresponding iterative algorithm is based on the alternating direction method of multipliers (ADMM). Finally, experiments conducted using synthetic and real data demonstrate the effectiveness of the proposed approach.

Index Terms

Hyperspectral imaging, nonlinear mixing, vector-valued RKHS, ADMM.

I. INTRODUCTION

Hyperspectral images are a very powerful tool in remote sensing. They provide a spectrum for each pixel, which is a vector of reflectance values estimated over hundreds of contiguous spectral bands with high spectral resolution [1], [2]. The spatial resolution of hyperspectral images can vary from a few meters up to a hundreds of meters in the case of airborne remote sensing. No matter the spatial resolution in this range, the image possibly contains mixed pixels where the surface covered by the pixel contains more than one constituent material [3]. This is in contrast with a pure pixel where the corresponding surface only contains one constituent material also known as an endmember. According to the well-known linear mixing model (LMM), the spectrum associated with a mixed pixel is a linear combination of the endmembers spectra [4]–[6]. More formally:

$$s_n = \sum_{i=1}^{M} a_{i,n} \mathbf{r}_i + \mathbf{e}_n, \forall n = 1, \dots N,$$

$$\tag{1}$$

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where s_n is the L-dimensional spectrum of the n-th pixel, L is the number of frequency bands, M denotes the number of endmembers, $a_{i,n}$ is the abundance of the i-th endmember in the n-th pixel, r_i is the L-dimensional spectrum of the i-th endmember, e_n is a vector of Gaussian white noise, and N is the number of observations. All vectors are column vectors. The abundances, which represent the relative contributions of the endmembers [7], are positive and usually sum to one: $a_{i,n} \geq 0$ and $\sum_{i=1}^{M} a_{i,n} = 1$ respectively. The LMM (1) is a simplified spectral mixing model. It only considers light reaching the sensor that has interacted once with the imaged surface, and neglects complex interactions between light, the imaged surface, neighboring surfaces and the atmosphere.

More recently, there has been a considerable amount of studies devoted to nonlinear mixing models [8], [9]. In particular, bilinear models are among the most widely known to account for nonlinear effects [10]–[13]. The physical assumption underlying these models is that light beams go through multiple reflections before reaching the sensor, mainly due to the three dimensionality of real scenes and scattering in the atmosphere [14]–[16]. The mathematical expression established for multiple reflections is the term by term product of two reflectance vectors in the case of bilinear models, and more than two reflectance vectors in the case of multilinear models [17]. For example, the polynomial post nonlinear mixing model (PPNM) introduced in [14] considers bilinear contributions through the following formulation:

$$\boldsymbol{s}_{n} = \sum_{i=1}^{M} a_{i,n} \boldsymbol{r}_{i} + u_{n} \left(\sum_{i=1}^{M} a_{i,n} \boldsymbol{r}_{i} \right) \odot \left(\sum_{i=1}^{M} a_{i,n} \boldsymbol{r}_{i} \right) + \boldsymbol{e}_{n},$$
 (2)

where u_n is the nonlinearity parameter, and \odot denotes the Hadamard (element wise) product. On the right hand side of equation (2), the first term corresponds to the linear mixture and the second term corresponds to the nonlinear (bilinear) one. Another way for modelling the nonlinear mixtures of the endmembers is through the use of nonlinear (scalar-valued) functions in reproducing kernel Hilbert spaces [18], [19]. The advantage of kernel-based models over models similar to (2) is that they are non parametric which means that they do not impose a predetermined form for the nonlinearity. For example, the authors of [20] propose the following model:

$$s_n = \sum_{i=1}^{M} a_{i,n} r_i + \Psi_n(\mathbf{R}) + e_n, \tag{3}$$

where $\Psi_n(R) = [\Psi_n(r_{\lambda_1}) \dots \Psi_n(r_{\lambda_L})]^{\top}$, $\Psi_n(\cdot)$ is a nonlinear function in a reproducing kernel Hilbert space (RKHS), and r_{λ_i} denotes the *i*-th row in R. The authors of [20] show that model (3) is able to incorporate bilinear, multilinear as well as more complex nonlinear interactions between the endmembers depending on the kernel choice. The main drawback of both, bilinear and (scalar-valued) kernel based models, is that they impose the same function at all bands, which can be restrictive in practice. More precisely, bilinear models consider the same amount of bilinear contributions at all bands. In the case of the PPNM (2), the same weight $u_n a_{i,n} a_{j,n}$ is used to scale the bilinear contribution of $r_i \odot r_j$ at all bands. Similarly, the kernel-based model (3) considers the same scalar-valued nonlinear function $\Psi_n(\cdot)$ at all bands. In contrast with the aforementioned models, the authors of [21], [22] do not impose any analytical form for the nonlinear term. The nonlinear contribution is merely treated as a positive residual term that is sparsely (rarely) present among the observations [22]. Nevertheless, this model-free approach can be limiting itself since it does not control the nonlinear expression, hence it can prevent accurate estimations of the nonlinear contribution.

The proposed nonlinear mixing model circumvents the drawbacks of the previously cited models by assuming that the nonlinear function belongs to a reproducing kernel Hilbert space (RKHS) of vector-valued functions. This approach improves upon the case of RKHS of scalar-valued functions by allowing for variable nonlinear contributions at different

bands. The key idea is to work under the assumption that nonlinear contributions can be dominant in some parts of the spectrum and less significant in other parts [23]–[25]. Unlike the scalar valued case where the same function is considered at each band, the proposed model relaxes this constraint, and allows to take into account wavelength dependent nonlinear contributions. In particular, we focus on RKHS associated with a special type of kernels, namely separable kernels. This type of kernels jointly defines the form of the nonlinear contribution, and allows to include prior information regarding the similarity between the nonlinear contributions at different wavelength bands. Similarly to the PPNM model, the input of the nonlinear vector-valued function includes the linear mixture spectra present in the pixel. We go one step further by also including the spectra of the linear mixtures in neighboring pixels. This choice is motivated by the adjacency effect [24] which states that solar radiation reflected off adjacent surfaces can be scattered into the sensor's instantaneous field of view. Figure 1 shows two forms of the adjacency effect, and depicts how neighboring surfaces can nonlinearly contribute to the reflectance vector estimated for a pixel. The adjacency effects are usually removed in a pre-processing step known as atmospheric correction. There exists different empirical methods for atmospheric correction [24]–[26]. Nevertheless, the validity of some of these methods to correct the adjacency effect is still questionnable [27], and errors occurred by these methods can damage the quality of information extracted from remote sensing data [28]. As a result, accounting for potential adjacency effects through the input of the nonlinear function increases the mixing model accuracy.

The paper is organized as follows. Section II describes the nonlinear mixing model, and discusses approaches for constructing the matrix valued kernel, section III develops the optimization problem and the corresponding estimation algorithm. Finally, section IV validates the proposed mixing model using synthetic and real data.

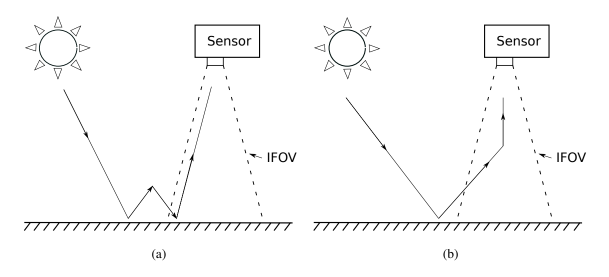


Fig. 1. Illustration of two forms of the adjacency effect resulting from: (a) multiple reflections involving the targeted surface and an adjacent surface, (b) reflection off an adjacent surface that is then scattered in the atmosphere into the sensor's instantaneous field of view (IFOV).

II. NONLINEAR MIXING MODEL

A. Model Description

First, we assume that the image is partitioned into patches or groups of pixels, and that the pixels in each patch are associated with a vector-valued nonlinear function. For ease of notations, assume that the available observations $S = [s_1, \ldots, s_N]$ belong to the same patch and that they are associated with the function f. The proposed nonlinear model decomposes the spectrum of a pixel into the sum of a linear and nonlinear part:

$$s_n = s_n^{\ln} + f(\tilde{v}_n) + e_n, \tag{4}$$

where $s_n^{\text{lin}} = \sum_{i=1}^M a_{i,n} r_i$, $\tilde{v}_n = \text{col}(\{s_i^{\text{lin}}\}_{i \in \mathcal{C}_n})$, $\text{col}(\cdot)$ is an operator that stacks its arguments on top of each other, and \mathcal{C}_n is a set of c pixels indices including n and c-1 of its neighboring pixels indices (for example $\mathcal{C}_n = \{n, n-1, n+1\}$). The nonlinear contribution in (4) is expressed in terms of the pixel and its neighbors linear mixtures. This is in accordance with several bilinear models such as the post polynomial nonlinear mixing (PPNM) model [13] which expresses the nonlinear contribution solely in terms of s_n^{lin} . The inconvenience of model (4) is that \tilde{v}_n depends on the unknown abundances. As a result, the corresponding optimization problem is not convex since the function f is unknown itself. In order to have a convex optimization problem, we propose to approximate s_i^{lin} by s_i , hence equation (4) becomes:

$$s_n = s_n^{\ln} + f(v_n) + e_n, \tag{5}$$

where $v_n = \operatorname{col}(\{s_i\}_{i \in \mathcal{C}_n})$, and it is assumed that $f(v_n)$ and e_n are small compared to s_n^{lin} . The latter assumption holds provided that the signal to noise ratio (SNR) is relatively high and that the linear part dominates the nonlinear one. The vector-valued approach offers an elegant and flexible way to jointly estimate multiple nonlinear functions at all bands since $f(v_n)$ implicitly corresponds to having different scalar-valued functions per band, i.e. $f(v_n) = [f_1(v_n) \dots f_L(v_n)]^{\top}$. As mentioned previously, the same nonlinear function is associated with all the pixels in the corresponding patch. It is important to note that the patch should contain enough pixels in order to have a good estimation of the nonlinear function. Moreover, it should be small enough to reflect the variability of the nonlinear function in the different regions of the image.

B. RKHS of vector-valued functions

The nonlinear function f used in (5) is a vector-valued function, its evaluation is a vector with L components representing the nonlinear contributions present in s_n at each band:

$$egin{aligned} f: & \mathbb{R}^{L\,c} &
ightarrow & \mathbb{R}^L \ & oldsymbol{v}_n &
ightarrow & oldsymbol{f}(oldsymbol{v}_n). \end{aligned}$$

As mentioned previously, we assume that f belongs to \mathcal{H}_k , a RKHS of vector-valued functions associated with the following kernel function:

$$\widetilde{k}: \mathbb{R}^{Lc} \times \mathbb{R}^{Lc} \to \mathbb{R}^{L\times L}
(\boldsymbol{v}_n, \boldsymbol{v}_{n'}) \to \widetilde{k}(\boldsymbol{v}_n, \boldsymbol{v}_{n'}).$$
(7)

Unlike the scalar-valued case, the kernel is a positive semi-definite matrix in $\mathbb{R}^{L\times L}$ rather than a positive scalar value. The overall Gram matrix \widetilde{K} associated with the function f is the matrix obtained from the evaluation of the kernel function (7) at all observation couples. It is a block matrix, such that the block indexed by (n, n') is given by:

$$\widetilde{\boldsymbol{K}}_{n,n'} = \widetilde{\boldsymbol{k}}(\boldsymbol{v}_n, \boldsymbol{v}_{n'}). \tag{8}$$

The Gram matrix consists of $N \times N$ blocks, where each block is an $L \times L$ matrix as defined in (8). As a result, \widetilde{K} is an $LN \times LN$ matrix. The representer theorem for vector-valued functions parallels the theorem in the scalar valued case. According to [29], $f(v_n)$ can be expressed as an expansion of the kernel function over all training points:

$$f(\mathbf{v}_n) = \sum_{n'=1}^{N} \widetilde{\mathbf{k}}(\mathbf{v}_n, \mathbf{v}_{n'}) \alpha_{n'}, \tag{9}$$

where $\alpha_{n'} \in \mathbb{R}^L$. As a result, estimating the nonlinear function reduces to estimating the coefficients $\{\alpha_{n'}\}_{n'=1}^N$. Finally, the norm of the function f in the RKHS $\widetilde{\mathcal{H}}_k$ can be written as:

$$\|\boldsymbol{f}\|_{\widetilde{\mathcal{H}}_{k}}^{2} = \sum_{n=1}^{N} \sum_{n'=1}^{N} \boldsymbol{\alpha}_{n}^{\top} \widetilde{\boldsymbol{k}}(\boldsymbol{v}_{n}, \boldsymbol{v}_{n'}) \boldsymbol{\alpha}_{n'}, \tag{10}$$

which gives a natural measure of the complexity of the function [29], [30].

C. Kernel design

The kernel function $k(v_n, v_{n'})$, as defined in equation (7), is an $L \times L$ matrix. The design of the kernel allows to jointly define the nonlinearity and include prior information regarding the similarity between the outputs of the nonlinear function at different bands. The authors of [29], [31] describe two possible kernel classes known as the transformable and the separable kernels. In what follows we describe these two classes of kernels, and explain their relevance in the nonlinear unmixing context.

1) Transformable and separable kernels: The first class of matrix valued kernels is known as transformable kernels. In the transformable case, the kernel $\tilde{k}(v_n, v_{n'})$ is defined in a component-wise fashion through a scalar valued kernel. Each component of the kernel is defined as:

$$\left[\widetilde{\boldsymbol{k}}(\boldsymbol{v}_{n},\boldsymbol{v}_{n'})\right]_{\ell,\ell'} = k(\boldsymbol{T}_{\ell}\boldsymbol{v}_{n},\boldsymbol{T}_{\ell'}\boldsymbol{v}_{n'}),\tag{11}$$

where k is a scalar valued kernel, and T_{ℓ} is an operator that extracts the c reflectance values in v_n corresponding to the ℓ -th band. More precisely, $T_{\ell}v_n = \operatorname{col}(\{s_{\ell,i}\}_{i\in\mathcal{C}_n})$ is a vector with c components. The scalar valued kernel acts jointly on pixels and bands indices, (n,n') and (ℓ,ℓ') respectively. The relevance of the transformable kernel is that the Gram matrix encodes the similarity between all pairs of pixels at all bands. Hence, it exploits all these correlations in order to jointly estimate the L components of the nonlinear function.

The second class of matrix valued kernels is known as the separable kernels. This class of kernels allows to incorporate prior information regarding the similarities between the different components of the vector valued function. The separable kernel is defined as the product between a scalar kernel acting on the input and an $L \times L$ positive semi-definite matrix encoding the similarities between the nonlinear contributions at different bands, i.e. between the different components of f. For this class, the kernel is defined as follows:

$$\widetilde{k}(\boldsymbol{v}_n, \boldsymbol{v}_{n'}) = k(\boldsymbol{v}_n, \boldsymbol{v}_{n'})\boldsymbol{\mathcal{E}},\tag{12}$$

where \mathcal{E} is an $L \times L$ positive semi-definite matrix. The norm of f gives further insight on how \mathcal{E} encodes the similarities between the nonlinear contributions at different bands. In fact, the norm of f in $\widetilde{\mathcal{H}}_k$ [32], [33] is given by:

$$\|\boldsymbol{f}\|_{\widetilde{\mathcal{H}}_{k}}^{2} = \sum_{\ell=1}^{N} \sum_{\ell'=1}^{N} \mathcal{E}_{\ell,\ell'}^{\dagger} \langle f_{\ell}, f_{\ell'} \rangle_{\mathcal{H}_{k}}, \tag{13}$$

where \mathcal{E}^{\dagger} is the pseudo inverse of \mathcal{E} , the scalar valued nonlinear functions f_1, \ldots, f_L belong to the RKHS \mathcal{H}_k associated with k, such that $\mathbf{f} = [f_1, \ldots, f_L]^{\top}$. Note that in the case of the separable kernel, given (8) and (12), the overall Gram matrix can be written in the following form:

$$\widetilde{K} = K \otimes \mathcal{E},\tag{14}$$

where \otimes is the kronecker product, and K is the $N \times N$ Gram matrix associated with the scalar valued kernel, namely $k_{n,n'} = k(\boldsymbol{v}_n, \boldsymbol{v}_{n'})$. Hereafter, we investigate a special structure for the matrix $\boldsymbol{\mathcal{E}}$. First, we assume that there exits prior information about the closeness between nonlinear contributions at different bands, i.e. between the functions f_1, \ldots, f_L . This prior can be modeled by a graph. We denote by $\boldsymbol{W} \in \mathbb{R}^{L \times L}$ the adjacency matrix of this graph [34]. More precisely, when two bands are likely to have similar nonlinear contributions, the corresponding nodes are connected by an edge and associated with a positive similarity weight $w_{\ell,\ell'} > 0$, otherwise $w_{\ell,\ell'}$ is set to zero. The authors of [32] show that when $\boldsymbol{\mathcal{E}}^{\dagger}$ is related to \boldsymbol{W} as follows:

$$\begin{cases}
\mathcal{E}_{\ell,\ell'}^{\dagger} = -w_{\ell,\ell'}, & \text{if } \ell \neq \ell', \\
\mathcal{E}_{\ell,\ell}^{\dagger} = \sum_{\ell'=1}^{N} w_{\ell,\ell'}, & \text{otherwise,}
\end{cases}$$
(15)

using (13), the norm of f in $\widetilde{\mathcal{H}}_{k}$ can be rewritten as:

$$\|f\|_{\widetilde{\mathcal{H}}_{k}}^{2} = \sum_{\ell=1}^{L} \|f_{\ell}\|_{\mathcal{H}_{k}}^{2} w_{\ell,\ell} + \frac{1}{2} \sum_{\ell=1}^{L} \sum_{\ell'=1}^{L} \|f_{\ell} - f_{\ell'}\|_{\mathcal{H}_{k}}^{2} w_{\ell,\ell'}.$$

$$(16)$$

From a regularization point of view, the norm of f as given by equation (16) is known as the graph regularizer. It penalizes the norms of the individual functions in addition to the differences between each pair of functions, hence forcing them to be similar. Moreover, the strength of the similarity between each pair of functions is determined by the corresponding weight. More precisely, a high value of $w_{\ell,\ell'}$ promotes a strong similarity between f_{ℓ} and $f_{\ell'}$, and conversely, a low value of $w_{\ell,\ell'}$ promotes a weak similarity between the two functions. In other words, the norm of f as given by equation (16) promotes similarity between the estimated nonlinearities at different bands in accordance with the prior information reflected through the graph. Finally, note that when $\mathcal{E} = I_L$, the norm of f reduces to the sum of the individual norms of its components $f_{\ell'}$. This corresponds to the case where there is no prior information between the nonlinearities at different bands.

In general, it is very likely that the nonlinear contributions at consecutive bands have smooth spectral variations. This prior information can be represented by a linear graph, where each node is connected to nodes at adjacent bands with unit weight. Nevertheless, more complex similarities can be incorporated using the graph structure. For example, in certain scenes nonlinear contributions can be dominant in certain spectral domains and less dominant in other spectral domains [23], [35]. This prior information can be represented by a clustered graph, where only nodes in spectral domains with similar nonlinear behavior are connected to each other.

2) Scalar kernel choice: The transformable and the separable kernels are both defined respectively in equations (11) and (12) using a scalar valued kernel k. Similarly to the case of functions in scalar valued RKHS, the choice of the kernel corresponds to a certain representation of the input data in a higher dimensional space known as the feature space [19]. Hence, looking at the feature space can provide guidelines for choosing an appropriate kernel.

We focus on the polynomial and Gaussian kernels due to their successful application to nonlinear unmixing in the scalar valued case [20], [36]. In particular, the second order homogeneous polynomial kernel:

$$k(\boldsymbol{v}_n, \boldsymbol{v}_{n'}) = (\boldsymbol{v}_n^{\top} \boldsymbol{v}_{n'})^2, \tag{17}$$

can be written as the inner product of the feature maps of v_n and $v_{n'}$, where the feature map is defined as follows:

$$\phi(\boldsymbol{v}_n) = [(v_{n,1})^2, \dots (v_{n,Lc})^2, \sqrt{2}(v_{n,1}v_{n,2})^2, \dots \sqrt{2}(v_{n,1}v_{n,Lc})^2, \sqrt{2}(v_{n,2}v_{n,3})^2, \sqrt{2}(v_{n,2}v_{n,Lc})^2, \dots \sqrt{2}(v_{n,Lc-1}v_{n,Lc})^2].$$

$$(18)$$

The feature map of the second order homogeneous polynomial kernel maps its input vector to all the possible pairwise products between its components. This can be seen as a representation of all possible second order interactions between the spectral values in the input vector. On the other hand, the Gaussian kernel:

$$k(\mathbf{v}_n, \mathbf{v}_{n'}) = \exp(-\frac{\|\mathbf{v}_n - \mathbf{v}_{n'}\|^2}{2\sigma^2}),$$
 (19)

can be expressed as an infinite series of higher order polynomial kernels:

$$k(\boldsymbol{v}_{n}, \boldsymbol{v}_{n'}) = \sum_{j=0}^{\infty} \frac{(\boldsymbol{v}_{n}^{\top} \boldsymbol{v}_{n'})^{j}}{\sigma^{2j} j!} \exp(-\frac{\|\boldsymbol{v}_{n}\|^{2}}{2\sigma^{2}}) \exp(-\frac{\|\boldsymbol{v}_{n'}\|^{2}}{2\sigma^{2}}).$$
(20)

Theoretically, the Gaussian kernel represents the case where an endless number of reflections occurs in the scene since it incorporates all higher order interactions between the input spectra. The drawback of the Gaussian kernel is that its feature map also contains a constant and a linear contribution (for j = 0 and j = 1 in equation (20)) which can hinder the estimation accuracy. Nevertheless, the Gaussian kernel shows satisfying results in practice as will be seen in the experiments.

III. ESTIMATION ALGORITHM

A. Optimization problem

In this section we derive the optimization problem aimed at estimating the abundances and the nonlinear function based on model (5). Assuming that the noise is white, Gaussian, with zero mean, and a possibly unknown variance, the maximum likelihood estimation leads to the least square (LS) optimization problem:

$$\min_{\boldsymbol{A}, \boldsymbol{f} \in \widetilde{\mathcal{H}}_{\boldsymbol{k}}} \quad \frac{1}{2} \sum_{n=1}^{N} \|\boldsymbol{s}_n - \boldsymbol{R} \boldsymbol{a}_n - \boldsymbol{f}(\boldsymbol{v}_n)\|^2, \tag{21}$$

where $\mathbf{R} = [\mathbf{r}_1, \dots, \mathbf{r}_M]$, and $\mathbf{a}_n = [a_{1,n}, \dots, a_{M,n}]^{\top}$. Problem (21) mainly ensures that the estimated model matches the observations. Nevertheless, the estimation of the abundances and the nonlinear function based on (21) is an underdetermined problem. As a result, it requires regularization and taking into account additional constraints on the abundances. For these reasons, we shall consider the following optimization problem:

$$\min_{\boldsymbol{A}, \boldsymbol{f} \in \widetilde{\mathcal{H}}_{\boldsymbol{k}}} \quad \frac{1}{2} \sum_{n=1}^{N} \|\boldsymbol{s}_{n} - \boldsymbol{R}\boldsymbol{a}_{n} - \boldsymbol{f}(\boldsymbol{v}_{n})\|^{2} + \frac{\lambda}{2} \|\boldsymbol{f}\|_{\widetilde{\mathcal{H}}_{\boldsymbol{k}}}^{2} + \mu \mathcal{J}(\boldsymbol{A})$$
subject to $a_{i,n} \succeq 0 \quad \forall i = 1, \dots, M, n = 1, \dots, N,$

$$\sum_{i=1}^{M} a_{i,n} = 1 \quad \forall n = 1, \dots, N,$$
(22)

where $A = [a_1, \dots, a_N]$, λ and μ are tuning parameters that control the tradeoff between the LS term and the two regularization terms. The first regularization, namely the ℓ_2 -norm of f in $\widetilde{\mathcal{H}}_k$, constrains the complexity of the estimated

function [30]. Furthermore, in the case of the separable kernel, it corresponds to the graph regularizer (16) which promotes certain similarities between the outputs of f. The second regularizer, namely $\mathcal{J}(A)$, aims at incorporating prior information about the abundances. For example, in the experiments we use the Frobenius norm of the abundances:

$$\mathcal{J}(\mathbf{A}) = \frac{1}{2} \|\mathbf{A}\|_{\mathsf{F}}^2,\tag{23}$$

known for promoting smoothness. As shown in the next section III-B, using another expression for $\mathcal{J}(A)$ is not cumbersome and affects one step in the iterative algorithm. Nevertheless, an advantage of using the ℓ_2 norm of the nonlinear function and the Frobenius norm of the abundances is that each regularization is strictly convex with respect to the corresponding unknown variable. Hence, the overall optimization problem is strictly convex with respect to all the unknown variables. Finally, the proposed optimization problem (22) imposes the positivity and sum-to-one constraints on the estimated abundances. Some of the nonlinear mixing models in the literature keep the sum-to-one constraint, as for example [13], [36]. It can be argued that this constraint should be relaxed to $\sum_{i=1}^{M} a_{i,n} \leq 1$ especially when dealing with real hyperspectral data. Even if this constraint is strictly enforced in the proposed optimization problem (22), it can be relaxed by introducing a shade endmember in the endmember matrix [37]. Furthermore, we show in the next section that dropping this constraint requires a simple modification of the iterative algorithm.

B. Iterative algorithm

In this section, we use the alternating direction method of multipliers (ADMM) [38] to solve the proposed optimization problem (22). The ADMM is a primal dual splitting method based on the augmented Lagrangian [39], [40]. Following the ADMM strategy, new variables and the corresponding consensus constraints are introduced in (22) in order to decouple the various terms in the objective function. We reformulate the optimization problem (22) in the following equivalent manner:

$$\min_{\boldsymbol{X},\boldsymbol{Z},\boldsymbol{f}\in\widetilde{\mathcal{H}}_{k}} \frac{1}{2} \sum_{n=1}^{N} \|\boldsymbol{s}_{n} - \boldsymbol{R}\boldsymbol{x}_{n} - \boldsymbol{f}(\boldsymbol{v}_{n})\|^{2} + \frac{\lambda}{2} \|\boldsymbol{f}\|_{\widetilde{\mathcal{H}}_{k}}^{2} + \mu \mathcal{J}(\boldsymbol{Z}) + \mathcal{I}_{\mathbb{R}_{+}^{M\times N}}(\boldsymbol{Z})$$
subject to $\mathcal{A}\boldsymbol{X} + \mathcal{B}\boldsymbol{Z} = \mathcal{C}$, (24)

with

$$\mathbf{A} = \begin{pmatrix} \mathbf{I}_{M \times M} \\ \mathbf{1}_{M}^{\top} \end{pmatrix}, \ \mathbf{B} = \begin{pmatrix} -\mathbf{I}_{M \times M} \\ \mathbf{0}_{M}^{\top} \end{pmatrix}, \ \mathbf{C} = \begin{pmatrix} \mathbf{0}_{M \times N} \\ \mathbf{1}_{N}^{\top} \end{pmatrix}$$
(25)

where \boldsymbol{X} and \boldsymbol{Z} are the ADMM variables, and $\mathcal{I}_{\mathbb{R}_{+}^{M\times N}}(\boldsymbol{Z})$ is the indicator of $\mathbb{R}_{+}^{M\times N}$ (i.e., $\mathcal{I}_{\mathbb{R}_{+}^{M\times N}}(\boldsymbol{Z})=0$ if $\boldsymbol{Z}\in\mathbb{R}_{+}^{M\times N}$ and $\mathcal{I}_{\mathbb{R}_{+}^{M\times N}}(\boldsymbol{Z})=\infty$ if $\boldsymbol{Z}\notin\mathbb{R}_{+}^{M\times N}$). Compared to problem (22), \boldsymbol{A} was substituted by \boldsymbol{X} and \boldsymbol{Z} , and a consensus constraint between the new variables was introduced. The positivity constraint was moved to the objective function through the indicator function, and the sum-to-one was incorporated within the equality constraint. As mentioned in the previous section, the sum-to-one constraint can be relaxed by adding a shade endmember to \boldsymbol{R} . Another alternative for relaxing the sum-to-one is by changing the definition of the matrices in (25) to:

$$\mathcal{A} = I_{M \times M}, \quad \mathcal{B} = -I_{M \times M}, \quad \mathcal{C} = \mathbf{0}_{M \times N}.$$
 (26)

The augmented Lagrangian associated with problem (24) is given by:

$$\mathcal{L}_{\rho}(\boldsymbol{X}, \boldsymbol{Z}, \boldsymbol{f}, \boldsymbol{\Lambda}_{\rho}) = \frac{1}{2} \sum_{n=1}^{N} \|\boldsymbol{s}_{n} - \boldsymbol{R}\boldsymbol{x}_{n} - \boldsymbol{f}(\boldsymbol{v}_{n})\|^{2} + \frac{\lambda}{2} \|\boldsymbol{f}\|_{\widetilde{\mathcal{H}}_{k}}^{2} + \mu \mathcal{J}(\boldsymbol{Z}) + \mathcal{I}_{\mathbb{R}_{+}^{M \times N}}(\boldsymbol{Z}) + \operatorname{tr}(\boldsymbol{\Lambda}_{\rho}^{\top}(\boldsymbol{A}\boldsymbol{X} + \boldsymbol{B}\boldsymbol{Z} - \boldsymbol{C})) + \frac{\rho}{2} \|\boldsymbol{A}\boldsymbol{X} + \boldsymbol{B}\boldsymbol{Z} - \boldsymbol{C}\|_{F}^{2},$$

$$(27)$$

where Λ_{ρ} is the matrix of Lagrange multipliers associated with the linear constraints in (24), and ρ is the penalty parameter. At each iteration, the ADMM algorithm consists of minimizing the augmented Lagrangian (27) sequentially. First, it is minimized with respect to the unknown variables $\{X, f\}$ and then with respect to Z while in each minimization keeping the other variables fixed to their previous estimate. Finally, it consists of updating the Lagrange multipliers matrix Λ_{ρ} associated with the linear constraints. This approach allows to break the optimization problem into a sequence of smaller and simpler sub-problems. The ADMM steps at each iteration, namely the $\{X, f\}$ minimization step, the Z minimization step, and the update of the Lagrange multipliers, are developed hereafter. To keep the notations simple, we drop the iteration index.

1) $\{X, f\}$ minimization step: This step consists of minimizing the augmented Lagrangian with respect to $\{X, f\}$. After discarding the terms independent of $\{X, f\}$ in (27), this step reduces to the following optimization problem:

$$\min_{\boldsymbol{X},\boldsymbol{f}\in\widetilde{\mathcal{H}}_{k}} \frac{1}{2} \sum_{n=1}^{N} \|\boldsymbol{s}_{n} - \boldsymbol{R}\boldsymbol{x}_{n} - \boldsymbol{f}(\boldsymbol{v}_{n})\|^{2} + \frac{\lambda}{2} \|\boldsymbol{f}\|_{\widetilde{\mathcal{H}}_{k}}^{2} \operatorname{tr}(\boldsymbol{\Lambda}_{\rho}^{\top} \boldsymbol{\mathcal{A}}\boldsymbol{X}) + \frac{\rho}{2} \|\boldsymbol{\mathcal{A}}\boldsymbol{X} + \boldsymbol{\mathcal{B}}\boldsymbol{Z} - \boldsymbol{\mathcal{C}}\|_{F}^{2}.$$
(28)

We rewrite (28) in the following equivalent form:

$$\min_{\boldsymbol{X},\boldsymbol{f}\in\widetilde{\mathcal{H}}_{k},\boldsymbol{E}} \quad \frac{1}{2} \sum_{n=1}^{N} \|\boldsymbol{e}_{n}\|^{2} + \frac{\lambda}{2} \|\boldsymbol{f}\|_{\widetilde{\mathcal{H}}_{k}}^{2} + \operatorname{tr}(\boldsymbol{\Lambda}_{\rho}^{\top} \boldsymbol{\mathcal{A}} \boldsymbol{X}) \frac{\rho}{2} \|\boldsymbol{\mathcal{A}} \boldsymbol{X} + \boldsymbol{\mathcal{B}} \boldsymbol{Z} - \boldsymbol{\mathcal{C}}\|_{F}^{2}$$
subject to
$$\boldsymbol{e}_{n} = \boldsymbol{s}_{n} - \boldsymbol{R} \boldsymbol{x}_{n} - \boldsymbol{f}(\boldsymbol{v}_{n}),$$

$$\forall n = 1, \dots, N,$$
(29)

where $E = [e_1, \dots, e_N]$, and solve its dual problem. The Lagrangian associated with problem (29) is given by:

$$\mathcal{L}(\boldsymbol{X}, \boldsymbol{f}, \boldsymbol{E}, \boldsymbol{\Lambda}) = \frac{1}{2} \sum_{n=1}^{N} \|\boldsymbol{e}_n\|^2 + \frac{\lambda}{2} \|\boldsymbol{f}\|_{\widetilde{\mathcal{H}}_{\boldsymbol{k}}}^2 + \operatorname{tr}(\boldsymbol{\Lambda}_{\rho}^{\top} \boldsymbol{\mathcal{A}} \boldsymbol{X}) + \sum_{n=1}^{N} \boldsymbol{\Lambda}_{n}^{\top} (\boldsymbol{s}_n - \boldsymbol{R} \boldsymbol{x}_n - \boldsymbol{f}(\boldsymbol{v}_n) - \boldsymbol{e}_n)$$

$$+ \frac{\rho}{2} \|\boldsymbol{\mathcal{A}} \boldsymbol{X} + \boldsymbol{\mathcal{B}} \boldsymbol{Z} - \boldsymbol{\mathcal{C}}\|_{\mathrm{F}}^2,$$
(30)

where $\Lambda = [\Lambda_1, \dots, \Lambda_N]$ is the matrix of Lagrange multipliers associated with the linear constraints in (29). The partial derivatives of the Lagrangian with respect to the primal variables, namely X, f and E, are:

$$\begin{cases} \frac{\partial \mathcal{L}}{\partial \mathbf{X}} &= \rho \mathbf{A}^{\top} \mathbf{A} \mathbf{X} - \mathbf{R}^{\top} \mathbf{\Lambda} + \mathbf{A}^{\top} \mathbf{\Lambda}_{\rho} + \rho \mathbf{A}^{\top} (\mathbf{B} \mathbf{Z} - \mathbf{C}), \\ \frac{\partial \mathcal{L}}{\partial \mathbf{f}} &= \lambda \mathbf{f}(\cdot) - \frac{1}{\lambda} \sum_{j=1}^{N} \widetilde{\mathbf{k}}(\cdot, \mathbf{v}_{j}) \mathbf{\Lambda}_{j}, \\ \frac{\partial \mathcal{L}}{\partial \mathbf{E}} &= \mathbf{E} - \mathbf{\Lambda}. \end{cases}$$
(31)

Setting the gradient of the partial derivatives in (31) to zero gives the primal variables as a function of the Lagrange multipliers:

$$\begin{cases}
\boldsymbol{X}^* &= \frac{(\boldsymbol{\mathcal{A}}^{\top} \boldsymbol{\mathcal{A}})^{-1}}{\rho} (\boldsymbol{R}^{\top} \boldsymbol{\Lambda}^* - \boldsymbol{\mathcal{A}}^{\top} \boldsymbol{\Lambda}_{\rho} - \rho \boldsymbol{\mathcal{A}}^{\top} (\boldsymbol{\mathcal{B}} \boldsymbol{Z} - \boldsymbol{\mathcal{C}})), \\
\boldsymbol{f}^*(\cdot) &= \frac{1}{\lambda} \sum_{j=1}^{N} \widetilde{\boldsymbol{k}} (\cdot, \boldsymbol{v}_j) \boldsymbol{\Lambda}_j^*, \\
\boldsymbol{E}^* &= \boldsymbol{\Lambda}^*.
\end{cases} (32)$$

To derive the Lagrange dual problem, the primal variables are substituted in (30) by their expressions from (32). This results in a quadratic form with respect to the Lagrange multipliers, and yields the following dual problem:

$$\max_{\mathbf{\Lambda}} \quad -\frac{1}{2} \operatorname{vec}(\mathbf{\Lambda})^{\top} Q \operatorname{vec}(\mathbf{\Lambda}) + \operatorname{vec}(\mathbf{\Lambda})^{\top} p, \tag{33}$$

with

$$\begin{cases}
Q = I_{L N \times L N} + \frac{1}{\lambda} \widetilde{K} + \frac{1}{\rho} I_{N \times N} \otimes D, \\
p = \text{vec}(S + \frac{1}{\rho} R(A^{\top} A)^{-1} A^{\top} (\Lambda_{\rho} + \rho(BZ - C))),
\end{cases}$$
(34)

where $D = R(A^{T}A)^{-1}R^{T}$, $\text{vec}(\cdot)$ is an operator that stacks the columns of a matrix on top of each other. As a result, the $\{X, f\}$ minimization step reduces to solving the following linear equation system:

$$Q\operatorname{vec}(\Lambda^*) = p, \tag{35}$$

with LN unknown variables. Once Λ^* is determined, it is substituted in (32) in order to evaluate the updated abundances. Note that the nonlinear function does not need to be evaluated at each iteration. It can be evaluated once, after the ADMM algorithm has converged.

2) Z minimization step: This step consists of minimizing the augmented Lagrangian with respect to Z. After discarding the terms independent of Z in (27) and accounting for the special structure of the matrices A, B, and C given in (25), problem (36) reduces to the following optimization problem:

$$\min_{\mathbf{Z}} \quad \frac{\rho}{2} \|\mathbf{Z} - \mathbf{X}\|_{\mathrm{F}}^{2} - \operatorname{tr}(\mathbf{\Lambda}_{\rho}^{\top} \mathbf{Z}) + \mu \mathcal{J}(\mathbf{Z}) + \mathcal{I}_{\mathbb{R}_{\perp}^{M \times N}}(\mathbf{Z}). \tag{36}$$

In particular, when $\mathcal{J}(\mathbf{Z})$ is the Frobenius norm (23), problem (36) reduces to the following positively constrained least squares problem:

$$\min_{\boldsymbol{Z}} \quad \frac{1}{2} \| \boldsymbol{Z} - \frac{\rho}{\rho + \mu} (\boldsymbol{X} + \frac{1}{\rho} \boldsymbol{\Lambda}_{\rho}) \|_{F}^{2} + \mathcal{I}_{\mathbb{R}_{\perp}^{M \times N}}(\boldsymbol{Z}). \tag{37}$$

The solution of problem (37) is obtained by a projection onto the positive orthant:

$$Z = \frac{\rho}{\rho + \mu} \left(X + \frac{1}{\rho} \Lambda_{\rho} \right)_{+}, \tag{38}$$

where $(\cdot)_+ = \max(0, \cdot)$ is applied component wise. As mentioned previously, $\mathcal{J}(\mathbf{Z})$ can be set to a regularization other than the Frobenius norm. For example, we demonstrate the case of the ℓ_1 norm known for promoting sparse abundances. In this case, problem (36) reduces to the following optimization problem:

$$\min_{\boldsymbol{Z}} \quad \frac{1}{2} \|\boldsymbol{Z} - (\boldsymbol{X} + \frac{1}{\rho} \boldsymbol{\Lambda}_{\rho})\|_{F}^{2} + \frac{\mu}{\rho} \|\boldsymbol{Z}\|_{1} + \mathcal{I}_{\mathbb{R}^{M \times N}}(\boldsymbol{Z}). \tag{39}$$

The solution of (39) is the well-known soft thresholding [41] applied to the projection onto the positive orthant:

$$Z = \operatorname{soft}_{\frac{\mu}{\rho}} ((X + \frac{1}{\rho} \Lambda_{\rho})_{+}), \tag{40}$$

where $\operatorname{soft}_{\frac{\mu}{\rho}}(\cdot) = \operatorname{sgn}(\cdot)(|\cdot| - \frac{\mu}{\rho})_+$, and the soft thresholding operator is applied component wise. The solution in equation (40) can be simplified to:

$$Z = (X + \frac{1}{\rho}\Lambda_{\rho} - \frac{\mu}{\rho})_{+}. \tag{41}$$

It is important to note that the sum to one constraint should be relaxed in (24) when the ℓ_1 norm is considered. Otherwise, $\mathcal{J}(\mathbf{Z})$ would be a constant in the feasible set, i.e. when the abundances are positive and sum to one [42].

3) **Update of the Lagrange multipliers:** The last step consists of updating the Lagrange multipliers according to the following rule,

$$\Lambda_{\rho}^{\to} = \Lambda_{\rho} + \rho (\mathcal{A}X + \mathcal{B}Z - \mathcal{C}), \tag{42}$$

where $\Lambda_{\rho}^{\rightarrow}$ denotes the updated matrix of Lagrange multipliers. This step can be seen as a gradient ascent of the augmented Lagrangian with respect to the Lagrange multiplier. Furthermore, it evaluates the running sum of the constraint residuals.

C. Implementation details

The ADMM steps described in the previous section are repeated until convergence. As suggested in [38], a reasonable stopping criteria is that the primal and dual residuals must be smaller than some tolerance thresholds, namely,

$$\|\mathcal{A}X + \mathcal{B}Z - \mathcal{C}\|_{\mathsf{F}} \le \epsilon_{\mathsf{pri}},$$
 (43)

$$\|\rho \mathcal{A}^{\top} \mathcal{B}(Z^{\rightarrow} - Z)\|_{F} \le \epsilon_{\text{dual}},$$
 (44)

The pseudocode of the proposed algorithm is given in Algorithm 1. The most computationally expensive step in the iterative algorithm is the X minimization step which requires solving an $LN \times LN$ system of linear equations (35). Solving the system of linear equations would have a memory complexity $\mathcal{O}(L^2N^2)$ and a runtime complexity $\mathcal{O}(L^3N^3)$. In practice, we do not need to compute the exact solution. The ADMM algorithm will converge even if the X minimization step is carried out only approximately [38], [40]. This allows us to solve (35) using an iterative algorithm which can reduce the runtime complexity. In the case of the separable kernel, both the memory and the runtime complexity can be reduced furthermore by exploiting the fact that Q is the sum of kronecker products.

The Conjugate Gradient method (CG) [43] is one of the most widely used iterative techniques for solving a large linear system of equations where Q is a positive definite matrix as in (35). The CG can yield the exact solution after LN iterations, but in practice a good initialization yields faster convergence [44]. At each iteration of the CG, the dominating operation is a matrix vector multiplication involving Q. In general, the number of operations required for multiplying Q by a vector is $\mathcal{O}(L^2N^2)$. However, in the case of the separable kernel, Q is a sparse matrix and can be written as the sum of kronecker products:

$$Q = I_L \otimes I_N + \frac{1}{\lambda} K \otimes \mathcal{E} + \frac{1}{\rho} I_N \otimes D, \tag{45}$$

where we have replaced \widetilde{K} by its expression from (14). For an efficient implementation of the product between Q and some vector $\text{vec}(\mathcal{J})$ where \mathcal{J} is an $L \times N$ matrix, the following relationships can be used:

$$\begin{cases}
(K \otimes \mathcal{E}) \operatorname{vec}(\mathcal{J}) = \operatorname{vec}(\mathcal{E} \mathcal{J} K), \\
(I_N \otimes D) \operatorname{vec}(\mathcal{J}) = \operatorname{vec}(D \mathcal{J})
\end{cases}$$
(46)

Given (45) and (46), the overall product between Q and some vector $\text{vec}(\mathcal{J})$ is given by:

$$Q\text{vec}(\mathcal{J}) = \text{vec}(\mathcal{J}) + \frac{1}{\lambda}\text{vec}(\mathcal{E}\mathcal{J}K) + \frac{1}{\rho}\text{vec}(D\mathcal{J}). \tag{47}$$

Equation (47) is expressed in terms of ordinary matrix products, which means that we do not have to compute any kronecker products. Compared to the case where the kronecker product is evaluated, the memory complexity is reduced from $\mathcal{O}(L^2N^2)$ to $\mathcal{O}(L^2+N^2)$ and the number of operations at each iteration is reduced from $\mathcal{O}(L^2N^2)$ to $\mathcal{O}(\max(L^2N,LN^2))$.

Algorithm 1 : $[X, F] = NDU(S, R, \lambda, \mu, \rho)$

- 1: Precompute $\mathcal{A}, \mathcal{B}, \mathcal{C}, \widetilde{K}, Q$
- 2: Initialize $oldsymbol{Z}, oldsymbol{\Lambda}_{
 ho}$
- 3: **while** $\operatorname{res}_{\operatorname{pri}} > \epsilon_{\operatorname{pri}}$ or $\operatorname{res}_{\operatorname{dual}} > \epsilon_{\operatorname{dual}}$ **do**

4:
$$p = \text{vec}(S + \frac{1}{\rho}R(A^{\top}A)^{-1}A^{\top}(\Lambda_{\rho} + \rho(BZ - C)))$$

5:
$$\operatorname{vec}(\mathbf{\Lambda}) = \mathbf{Q}^{-1}\mathbf{p}$$
 % See section III-C

6:
$$X = \frac{1}{\rho} (\mathcal{A}^{\top} \mathcal{A})^{-1} (\mathbf{R}^{\top} \mathbf{\Lambda} - \mathcal{A}^{\top} \mathbf{\Lambda}_{\rho} - \rho \mathcal{A}^{\top} (\mathcal{B} \mathbf{Z} - \mathcal{C}))$$

7:
$$oldsymbol{Z}^{ ext{old}} = oldsymbol{Z}$$

8: if
$$\mathcal{J}(\boldsymbol{Z}) = \|\boldsymbol{Z}\|_{\mathrm{F}}^2$$
 then

9:
$$Z = \frac{\rho}{\rho + \mu} \left(X + \frac{1}{\rho} \Lambda_{\rho} \right)_+$$

10: else if
$$\mathcal{J}(\boldsymbol{Z}) = \|\boldsymbol{Z}\|_1$$
 then

11:
$$oldsymbol{Z} = (oldsymbol{X} + rac{1}{
ho} oldsymbol{\Lambda}_{
ho} - rac{\mu}{
ho})_+$$

12: end if

13:
$$\Lambda_{
ho} = \Lambda_{
ho} +
ho(\mathcal{A}X + \mathcal{B}Z - \mathcal{C})$$

14:
$$\operatorname{res}_{\operatorname{pri}} = \|\boldsymbol{\mathcal{A}}\boldsymbol{X} + \boldsymbol{\mathcal{B}}\boldsymbol{Z} - \boldsymbol{\mathcal{C}}\|_{\operatorname{F}}$$

15:
$$\operatorname{res}_{\operatorname{dual}} = \|\rho \mathbf{A}^{\top} \mathbf{B} (\mathbf{Z} - \mathbf{Z} \operatorname{old})\|_{\operatorname{F}}$$

16: end while

17:
$$oldsymbol{F} = rac{1}{\lambda} \widetilde{oldsymbol{K}} ext{vec}(oldsymbol{\Lambda})$$

18:
$$\mathbf{F} = \text{reshape}(\mathbf{F}, L, N)$$

IV. EXPERIMENTS

A. Synthetic data: Illustrative examples

- 1) Data generation: The proposed approach is first illustrated using synthetic data. Several patches were generated according to three mixing models that incorporate the main assumptions underlying the proposed nonlinear mixing model, i.e., bilinear contributions, adjacency effects, and band selectivity. The bilinear contributions are created by adding pairwise products of spectra, the adjacency effect is created by adding bilinear contributions from neighboring pixels, and band selectivity is created by assigning a different weight to the nonlinear contributions at different bands. The three mixing models (MM) are denoted as MM 1, MM 2, and MM 3, and they are defined as follows:
 - MM 1 (bilinear contributions):

$$s_n = s_n^{\ln} + u \, s_n^{\ln} \odot s_n^{\ln} + e_n \tag{48}$$

• MM 2 (bilinear contributions + adjacency effects):

$$\mathbf{s}_n = \mathbf{s}_n^{\text{lin}} + u \sum_{i=n-2}^{n+2} \gamma_{n,i} \, \mathbf{s}_i^{\text{lin}} \odot \mathbf{s}_i^{\text{lin}} + \mathbf{e}_n \tag{49}$$

• MM 3 (bilinear contributions + adjacency effects + band selectivity):

$$s_n = s_n^{\text{lin}} + u \sum_{i=n-2}^{n+2} \gamma_{n,i} s_i^{\text{lin}} \odot s_i^{\text{lin}} \odot h + e_n$$
 (50)

where u in equation (48) is an attenuation parameter set to 0.2 in all the simulations, the coefficients $\gamma_{n,i}$ in equation (49) assign a different weight to the bilinear contributions coming from neighbors, the coefficients were set to $\gamma_{n,n-2} = \gamma_{n,n+2} = 0.05$, $\gamma_{n,n-1} = \gamma_{n,n+1} = 0.3$ and $\gamma_{n,n} = 0.4$, and h in equation (50) is an L dimensional vector where each component assigns a different weight to the nonlinear contribution at the corresponding band. Figure 2 (a) shows the entries of h that were used for the experiments. In fact, h was chosen such that it favors nonlinear contributions at the center of the spectrum and attenuates nonlinear contributions at the extremities of the spectrum. Note that h is unknown by all the unmixing methods used in the experiments. Several patches were created with N = 100 pixels using different numbers of endmembers and different values of the SNR. The endmembers were selected from the USGS spectral library of minerals. Their frequency bands are in the range 400 - 2560 nm, and were decimated such as to have L = 20 bands. Figure 2 (b) shows five endmembers spectra used in the simulations. The abundances were generated using a beta distribution with a unit shape parameter.

2) Unmixing methods: We tested three nonlinear unmixing algorithms. The first algorithm is the extended endmember matrix method. It considers the linear mixing model where the endmember matrix is extended by adding the pairwise products of the endmembers [45]. This algorithm is denoted as Ext in the experiments, it consists of solving a positively constrained least squares problem and has no tuning parameters. The second algorithm is the one proposed in [20], it is denoted as khype and based on scalar valued RKHS. Khype was tested with a Gaussian (G) and a second order homogeneous polynomial (P) kernel. The third algorithm is the one proposed in this paper, it is denoted in what follows as NDU (Nonlinear neighbor and band Dependent Unmixing). Similarly to khype, NDU was tested with a Gaussian (G) and a second order homogeneous polynomial (P) kernel. Furthermore, given that the kernel in NDU is matrix valued, it was tested using a transformable (Tr.) and a separable (Sp.) structure. In the case of the separable kernel, the graph that represents the similarities between the different bands is linear, i.e. each band is connected to the previous and next band, with unit weights. In order to determine v_n , the neighborhood was set to $C_n = \{n, n-1, n+1\}$.

Note that NDU requires tuning two parameters λ and μ , whereas khype [20] uses the same parameter to penalize the norm of the abundances and the nonlinear function ($\lambda = \mu$). In order to have a fair comparison between the two algorithms, we modified khype such as to have two distinct parameters λ and μ . For both algorithms, the tuning parameters were tested in the range [10^{-4} 10^{-3} 10^{-2} 10^{-1} 1 10]. The standard deviations of the Gaussian kernels used with khype and NDU were chosen such that the resulting Gram matrices have their values in the same range. The polynomial Gram matrices were scaled in order to have their values in the range [0 1]. Figure 3 shows the Gram matrices used with khype, and NDU in different settings and obtained with M=3 and SNR=40 dB. The first column in Figure 3 shows the $L\times L$ Gram matrices used by khype. The second and third columns in Figure 3 show the Gram matrices used by NDU obtained with a transformable and separable structure respectively. Recall that the NDU Gram matrices are $N\times N$ block matrices, where each block is an $L\times L$ matrix. In the case of the transformable kernel, each block is an $L\times L$ Gram matrix itself. Figure 3 shows that a block or sub-Gram matrix in the transformable kernel is similar the corresponding khype Gram matrix even though it is calculated using the observations themselves rather than the endmember matrix as in khype. Whereas for the separable kernel, each block is equal to \mathcal{E}^{\dagger} multiplied by the corresponding scalar valued kernel.

3) Performance measures: The abundance estimation accuracy was evaluated using the root mean square error (RMSE) defined as:

$$RMSE_{\boldsymbol{X}} = \sqrt{\frac{1}{MN} \|\boldsymbol{X} - \boldsymbol{X}^*\|_{F}^2},$$
(51)

where X represents the true abundances matrix and X^* respresents the abundances estimated by an unmixing algorithm. In addition to the abundances, each one of the unmixing algorithms estimates the nonlinear contributions. Let F and F^* denote the true and estimated $L \times N$ matrices of nonlinear contributions. The estimation accuracy of F was also evaluated using the RMSE defined as:

$$RMSE_{\mathbf{F}} = \sqrt{\frac{1}{LN} \|\mathbf{F} - \mathbf{F}^{\star}\|_{F}^{2}}.$$
(52)

4) Simulation results: Tables I, II and III report the results obtained using MM 1, MM 2, and MM 3 respectively. For each case, we report the root mean square errors of the estimated abundances (first term in brackets) and the estimated nonlinear contributions (second term in brackets). Khype and NDU require tuning two parameters, namely λ and μ , the best parameters are reported in brackets below the RMSE results. More precisely, the first term and second term in brackets correspond to the best values of λ and μ respectively. The estimation accuracy of the three methods became worse when the noise level increased. The same performance was observed for different values of M. In general, Khype outperformed Ext, and NDU outperformed both methods. Note that Ext gave results worse than NDU and khype even when MM 1 was used. This is most probably due to the fact that the sum to one constraint was not incorporated in the implementation of Ext, unlike the case of khype and NDU. In the majority of the cases, NDU gave the best RMSE of the abundances and the nonlinear part when used with the separable and polynomial kernel (Sp. + P). The polynomial and the Gaussian kernels usually had comparable results in all scenarios. The transformable kernel and the separable kernel had comparable results only when MM 1 and 2 were used, i.e. in tables I and II. However, the separable kernel outperformed the transformable kernel when MM 3 was used, i.e. in table III.

Figure 4 shows the true and estimated nonlinear contributions for all pixels at all bands obtained with P=3 and SNR=40 dB. The first row in Figure 4 shows the true nonlinear contributions. Whereas the following rows show the nonlinear contributions estimated with Ext, khype, NDU with a transformable kernel, and NDU with a separable kernel.

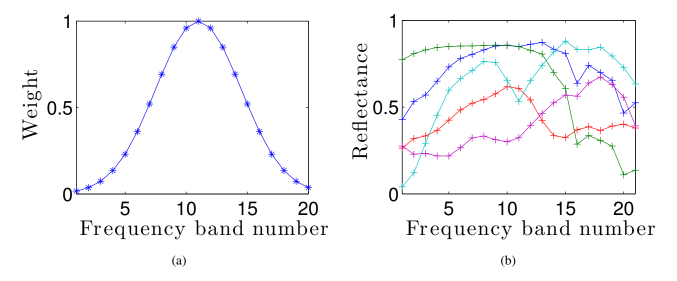


Fig. 2. Fom left to right: (a) The components of vector h corresponding to the weight of the nonlinear contribution at each band, (b) Endmembers spectra used to generate the illustrative examples.

For conciseness, we show the results with either the Gaussian or the polynomial kernel for Khype and NDU. In particular, the kernel that gave the best RMSE was chosen. Figure 4 mainly allows to visually compare the estimation of the nonlinear contribution obtained with the various methods and mixing models. The first column in Figure 4 corresponds to the results obtained with MM 1, i.e. with bilinear contributions. It can be seen that khype and NDU slightly outperformed Ext. The second column in Figure 4 corresponds to the results obtained with MM 2, i.e. with bilinear contributions and adjacency effects. The first figure in column 2 shows that the true nonlinear contributions at adjacent pixels (i.e. at adjacent columns in the image) have smooth variations. In this case, NDU with both the transformable and the separable kernel gave the smoothest results compared to Ext and khype. Recall that the kernels used by NDU account for adjacency effects through their input vector v_n . The third column in Figure 4 corresponds to the results obtained with MM 3, i.e. with bilinear contributions, adjacency effects, and band selectivity. In accordance with MM 3, the first figure in column 3 shows that the true nonlinear contributions are smooth, they are the most pronounced at the center of the spectrum and they are attenuated (almost zero) at the extremities of the spectrum. In this case, NDU with the separable kernel gave the best results. All the methods estimated the highest nonlinear contributions at the center of the spectrum. However, NDU with the separable kernel handled the attenuation of the bilinear contributions at the extremities of the spectrum better than the other methods. This is probably due to the fact that the prior information on the similarities between the nonlinear contributions at different bands is better in the case of the separable kernel than in the transformable kernel. More precisely, the separable kernel exploits the linear graph structure which promotes smooth variations at adjacent bands (i.e. adjacent rows in the image). Whereas, the transformable kernel exploits the correlations between all bands in order to estimate the nonlinear contributions.

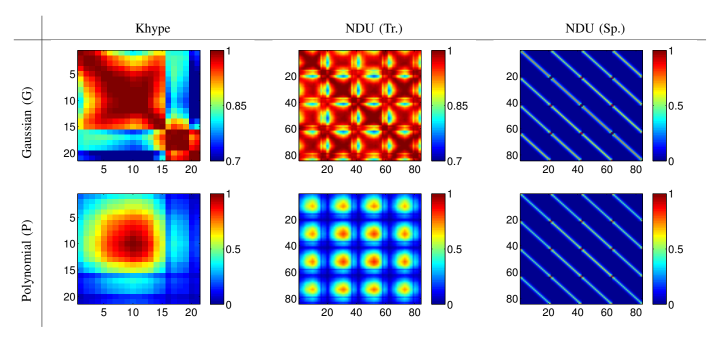
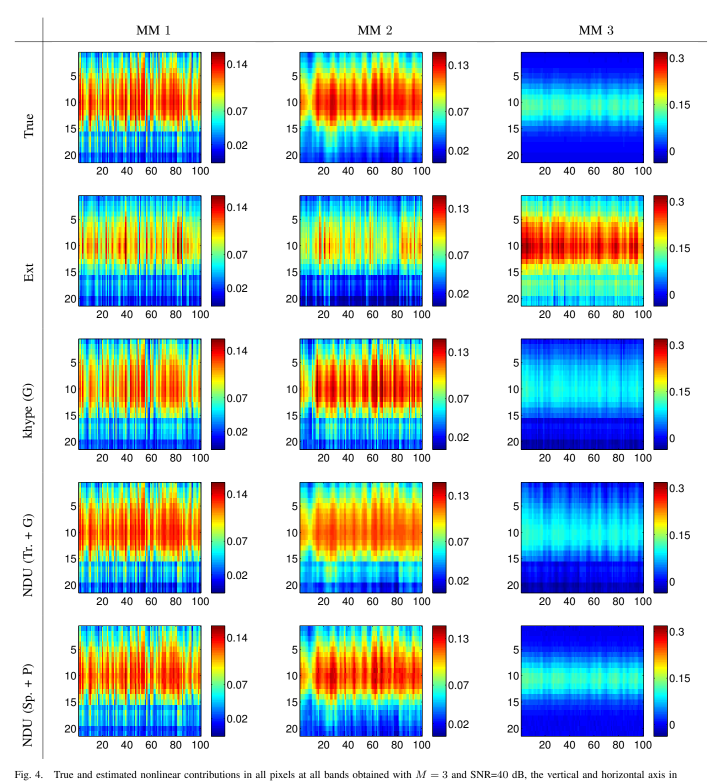


Fig. 3. Gram matrices obtained with M=3 and SNR = 40 dB. First column: Gram matrices used by khype, second column: left corners of the Gram matrices obtained using a transformable kernel, and third column: left corners of the Gram matrices obtained using a separable kernel. The first and the second row correspond to the Gaussian and polynomial kernels respectively.



each figure represent the frequency band and the pixel number respectively.

TABLE I ${\rm RMSE}\,(\times 10^{-2}) \ {\rm and} \ {\rm optimal} \ {\rm parameters} \ {\rm obtained} \ {\rm with} \ {\rm MM}\ 1. \ {\rm The} \ {\rm first} \ {\rm two} \ {\rm terms} \ {\rm in} \ {\rm brackets} \ {\rm are} \ {\rm the} \ {\rm RMSE} \ {\rm of} \ {\rm the} \ {\rm estimated}$ abundances (left term) and the nonlinear part (right term), and the second two terms in the lower brackets are the ${\rm optimal} \ {\rm values} \ {\rm for} \ \lambda \ ({\rm left} \ {\rm term}) \ {\rm and} \ \mu \ ({\rm right} \ {\rm term}).$

		SNR = 40			SNR = 30			SNR = 20	
	M = 3	M = 4	M = 5	M = 3	M = 4	M = 5	M = 3	M = 4	M = 5
Ext	(5.31, 2.35)	(4.98, 2.84)	(4.51, 2.63)	(7.38, 3.75)	(8.30, 5.84)	(7.19, 5.01)	(13.77, 8.06)	(15.28, 13.96)	(13.84, 11.22)
Khype	(2.01, 0.88)	(2.72, 1.14)	(2.60, 1.12)	(3.77, 1.55)	(4.68, 1.50)	(4.63, 1.84)	(9.52, 3.73)	(9.55, 3.12)	(9.33, 3.10)
(G)	(0.1, 0.01)	$(0.01, 10^{-3})$	$(0.01, 10^{-3})$	(0.1, 0.01)	(0.1, 0.01)	(0.1, 0.01)	(1, 0.1)	(1, 0.1)	(1, 0.1)
Khype	(2.79, 1.03)	(2.21, 1.20)	(2.08, 1.09)	(4.05, 1.55)	(4.21, 1.45)	(4.47, 1.74)	(8.73, 3.34)	(9.34, 2.91)	(9.01, 2.89)
(P)	$(0.01, 10^{-3})$	(0.1, 0.01)	(0.1, 0.01)	(0.1, 0.01)	(0.1, 0.01)	(0.1, 0.01)	(1, 0.1)	(1, 0.1)	(1, 0.1)
NDU.	(1.22, 0.54)	(1.50, 0.35)	(1.60, 0.29)	(2.82, 0.77)	(3.53, 0.95)	(3.65, 1.14)	(6.07, 2.26)	(7.79, 1.90)	(7.76, 2.11)
(Tr. + G)	(1, 0.01)	(1, 0.01)	(1, 0.01)	(10, 0.1)	$(1, 10^{-4})$	(10, 0.01)	$(10, 10^{-4})$	(10, 0.01)	(10, 0.01)
NDU	(1.43, 0.55)	(1.27, 0.26)	(1.35, 0.34)	(2.11, 0.76)	(3.12, 0.52)	(3.30, 0.67)	(5.55, 1.71)	(7.54, 1.66)	(7.87, 1.78)
(Tr.+P)	$(10^{-4}, 10^{-4})$	$(0.1, 10^{-4})$	$(0.1, 10^{-4})$	$(1, 10^{-4})$	$(1, 10^{-4})$	$(1, 10^{-4})$	$(10, 10^{-4})$	(10, 0.01)	(10, 0.01)
NFU	(1.79, 0.69)	(1.53, 0.52)	(1.98, 0.68)	(2.30, 1.36)	(3.81, 0.83)	(3.63, 0.89)	(6.17, 1.89)	(7.51, 2.73)	(7.71, 1.16)
(Sp+G)	$(0.01, 10^{-4})$	$(0.01, 10^{-3})$	$(0.01, 10^{-3})$	$(0.01, 10^{-4})$	(0.1, 0.01)	(0.1, 0.01)	(1, 0.01)	(0.1, 0.01)	(1, 0.1)
NDU	(1.17, 0.43)	(1.97, 0.72)	(2.15, 0.77)	(2.02, 0.85)	(3.73, 1.27)	(3.55, 0.98)	(5.79, 1.61)	(7.44, 1.65)	(7.61, 1.39)
(Sp+P)	$(10^{-3}, 10^{-4})$	$(10^{-3}, 10^{-4})$	$(10^{-3}, 10^{-4})$	$(0.01, 10^{-3})$	(0.1, 0.01)	$(0.01, 10^{-3})$	(1, 0.1)	(1, 0.1)	(1, 0.1)

TABLE II ${\rm RMSE}\,(\times 10^{-2}) \ {\rm and} \ {\rm optimal} \ {\rm parameters} \ {\rm obtained} \ {\rm with} \ {\rm MM}\ 2. \ {\rm The} \ {\rm first} \ {\rm two} \ {\rm terms} \ {\rm in} \ {\rm brackets} \ {\rm are} \ {\rm the} \ {\rm RMSE} \ {\rm of} \ {\rm the} \ {\rm estimated}$ abundances (left term) and the nonlinear part (right term), and the second two terms in the lower brackets are the ${\rm optimal} \ {\rm values} \ {\rm for} \ \lambda \ ({\rm left} \ {\rm term}) \ {\rm and} \ \mu \ ({\rm right} \ {\rm term}).$

		SNR = 40			SNR = 30			SNR = 20	
	M = 3	M = 4	M = 5	M = 3	M = 4	M = 5	M = 3	M = 4	M = 5
Ext	(4.27, 2.78)	(4.49, 3.32)	(3.80, 2.85)	(6.72, 3.44)	(8.03, 5.19)	(7.25, 4.65)	(12.98, 6.97)	(16.11, 12.57)	(13.57, 10.93)
Khype	(2.55, 0.99)	(2.33, 0.79)	(2.18, 0.88)	(4.37, 1.67)	(4.75, 1.34)	(5.07, 1.65)	(10.37, 4.18)	(9.41, 3.02)	(10.12, 2.93)
(G)	$(0.01, 10^{-3})$	$(0.01, 10^{-3})$	$(0.01, 10^{-3})$	$(0.1, 10^{-3})$	(0.1, 0.01)	(0.1, 0.01)	(1, 0.01)	(1, 0.1)	(1, 0.1)
Khype	(2.61, 1.05)	(2.26, 0.99)	(2.35, 1.14)	(3.76, 1.43)	(4.42, 1.27)	(5.06, 1.68)	(9.10, 3.30)	(9.16, 2.85)	(10.10, 2.85)
(P)	(0.1, 0.01)	$(0.1, 10^{-3})$	$(0.01, 10^{-3})$	(0.1, 0.01)	(0.1, 0.01)	(0.1, 0.01)	(0.1, 0.01)	(1, 0.1)	(1, 0.1)
NDU	(0.92, 0.64)	(1.54, 0.38)	(1.39, 0.33)	(1.97, 0.63)	(3.51, 0.60)	(3.93, 0.79)	(6.98, 2.53)	(8.51, 2.70)	(9.05, 2.71)
(Tr.+G)	(10, 0.01)	$(1, 10^{-4})$	$(1, 10^{-4})$	$(10, 10^{-3})$	$(10, 10^{-3})$	(10, 0.01)	$(10, 10^{-4})$	(10, 0.1)	(10, 0.01)
NDU	(1.75, 0.74)	(1.73, 0.43)	(1.62, 0.38)	(2.76, 1.04)	(3.42, 0.60)	(4.08, 0.73)	(5.82, 1.72)	(8.12, 2.10)	(9.14, 2.52)
(Tr+P)	(1, 0.01)	$(1, 10^{-4})$	$(1, 10^{-4})$	(10, 0.01)	$(10, 10^{-4})$	(10, 0.01)	$(10, 10^{-4})$	(10, 0.1)	(10, 0.01)
NDU	(1.92, 0.67)	(2.35, 0.84)	(1.87, 0.73)	(2.46, 1.40)	(3.71, 0.92)	(4.04, 1.03)	(5.67, 1.56)	(7.60, 1.38)	(8.93, 1.32)
(Sp.+G)	$(0.01, 10^{-4})$	$(0.01, 10^{-4})$	$(0.01, 10^{-4})$	$(0.01, 10^{-4})$	$(0.1, 10^{-3})$	$(0.1, 10^{-3})$	(1, 0.01)	(1, 0.1)	(1, 0.1)
NDU	(1.14, 0.39)	(2.21, 0.80)	(1.70, 0.61)	(2.12, 1.42)	(3.45, 0.83)	(3.91, 0.78)	(5.49, 1.85)	(7.36, 1.13)	(8.80, 1.03)
(Sp.+P)	$(10^{-3}, 10^{-4})$	$(0.01, 10^{-4})$	$(0.01, 10^{-3})$	$(10^{-3}, 10^{-4})$	$(0.1, 10^{-3})$	$(0.01, 10^{-3})$	$(0.1, 10^{-3})$	(1, 0.1)	(1, 0.1)

TABLE III ${\rm RMSE}\,(\times 10^{-2}) \ {\rm and\ optimal\ parameters\ obtained\ with\ MM\ 3}. \ {\rm The\ first\ two\ terms\ in\ brackets\ are\ the\ RMSE\ of\ the\ estimated\ abundances\ (left\ term)\ and\ the\ nonlinear\ part\ (right\ term),\ and\ the\ second\ two\ terms\ in\ the\ lower\ brackets\ are\ the\ optimal\ values\ for\ λ\ (left\ term)\ and\ μ\ (right\ term).$

		SNR = 40			SNR = 30			SNR = 20	
	M=3	M = 4	M = 5	M = 3	M = 4	M = 5	M = 3	M = 4	M=5
Ext	(9.21, 12.8)	(12.1, 15.7)	(8.97, 12.6)	(10.7, 13.0)	(12.7, 15.6)	(11.3, 15.5)	(17.7, 16.8)	(18.1, 21.0)	(15.9, 19.3)
Khype	(4.33, 2.18)	(3.55, 1.99)	(3.47, 1.91)	(5.80, 2.60)	(5.28, 2.20)	(5.48, 2.19)	(10.56, 5.21)	(9.59, 3.08)	(8.90, 3.53)
(G)	$(0.1, 10^{-3})$	$(0.1, 10^{-3})$	$(0.1, 10^{-3})$	$(0.1, 10^{-3})$	(0.1, 0.01)	(0.1, 0.01)	(10, 0.1)	(1, 0.01)	(10, 0.1)
Khype	(1.24, 2.63)	(4.34, 2.15)	(3.85, 2.30)	(2.56, 2.71)	(5.26, 2.24)	(4.85, 2.28)	(7.66, 3.31)	(9.44, 2.86)	(8.74, 3.33)
(P)	$(1, 10^{-3})$	(1, 0.01)	$(1, 10^{-3})$	(1, 0.01)	$(1, 10^{-3})$	(1, 0.01)	(1, 0.01)	(1, 0.1)	(10, 0.1)
NDU	(4.09, 1.95)	(3.20, 1.68)	(3.17, 1.68)	(4.39, 1.97)	(4.33, 1.78)	(4.63, 1.72)	(8.72, 3.25)	(10.06, 3.22)	(8.84, 3.15)
(Tr.+G)	$(10, 10^{-4})$	$(10, 10^{-4})$	$(10, 10^{-3})$	$(10, 10^{-4})$	$(10, 10^{-4})$	$(10, 10^{-4})$	$(10, 10^{-4})$	(10, 0.01)	(10, 0.01)
NDU	(5.96, 2.77)	(5.62, 2.30)	(5.32, 2.21)	(6.33, 2.80)	(6.62, 2.41)	(5.83, 2.31)	(10.15, 3.72)	(10.31, 3.38)	(8.81, 2.93)
(Tr+P)	(10, 0.1)	(10, 0.01)	$(10, 10^{-4})$	(10, 0.1)	(10, 0.01)	(10, 0.01)	(10, 0.1)	(10, 0.1)	(10, 0.1)
NDU	(1.29, 0.50)	(2.62, 1.03)	(1.75, 0.72)	(1.98, 0.69)	(3.40, 0.66)	(3.46, 0.68)	(6.29, 1.53)	(8.43, 1.62)	(7.99, 1.95)
(Sp+G)	$(0.1, 10^{-4})$	$(0.1, 10^{-3})$	$(0.1, 10^{-4})$	$(0.1, 10^{-4})$	$(0.1, 10^{-4})$	(1, 0.01)	(1, 0.01)	(10, 0.1)	(10, 0.1)
NDU	(0.95, 0.34)	(1.56, 0.42)	(1.51, 0.34)	(1.96, 0.41)	(3.25, 0.50)	(3.46, 0.48)	(6.15, 1.77)	(8.15, 0.98)	(7.64, 1.07)
(Sp+P)	$(0.01, 10^{-4})$	$(0.01, 10^{-3})$	$(0.01, 10^{-4})$	$(0.1, 10^{-3})$	$(0.1, 10^{-4})$	$(0.1, 10^{-3})$	(0.1, 0.01)	(1, 0.1)	(1, 0.1)

B. Real data: Gulf of Lion

1) Data set description and ground truth: The second set of experiments considers real data estimated by the Meris spectrometer and captured over the gulf of Lion in the south east of France. The image has 280×330 pixels, 13 spectral bands in the range 400 - 800 nm, and a spatial resolution of 300 m. This data set will be referred to as the "Meris" image in the following and is depicted in Figure 5 (a). Furthermore, Figure 5 (b) shows the corresponding classification map provided by Corine Land Cover (CLC) database [46]. Note that the two images were coregistered, and the classification map was chosen as close as possible to the date of the Meris image in order to have a consistent comparison. The classification map will be used for visual evaluation, in order to better evaluate and interpret the unmixing results provided by the various algorithms.

The CLC classification map has a spatial resolution approximately 10 times greater than the Meris data set's spatial resolution. Therefore, it was downscaled in order to obtain spatial abundance maps for the Meris image [47]. The spatial abundance of a certain class/endmember in a pixel in the estimated low resolution image is the average of the corresponding class occurrences in the corresponding window in the high resolution image. These maps are regarded as a potential visual ground truth that allow to better evaluate and interpret the unmixing results. The first row in Figure 7 shows the ground truth obtained from the CLC classification map, which corresponds to the proportions of three classes: water, agricultural areas, and forests and semi natural areas.

2) Unmixing results: We extracted 3 and 4 endmembers using virtual component analysis VCA [48]. We noticed that each time one of the endmembers extracted by VCA was not meaningful in the sense that the corresponding abundance map was not spatially coherent. Nevertheless, in the case with 4 extracted endmembers the three abundance maps corresponding to the meaningful endmembers were relatively in accordance with the estimated ground truth. In what follows we only show the results obtained with four endmembers. Figure 6 shows the estimated endmembers spectra, note that endmember 4 corresponds to the outlier. The abundance maps for each endmember were estimated using the fully constrained least squares approach (FCLS), Ext, khype and NDU. The separable kernel was used with NDU with a linear graph as in the experiments with synthetic data. Both khype and NDU were tested using a Gaussian and a second order homogeneous polynomial kernel. As in the previous section, v_n was defined using the pixels and its neighboring pixels spectra in particular the left and right neighbors were chosen. The tuning parameters λ and μ were set to 10 and 10^{-3} respectively for both khype and NDU. Unlike khype and Ext that were applied on each pixel separately, the image was divided into 10×10 patches and NDU was applied on each patch.

Table IV reports the root mean square error (RMSE) and the average spectral angle (ASA) of the reconstructed image using each algorithm. The RMSE is computed for all the observations in the image, and the ASA is given in radian. Table IV shows that NDU scored the best results in terms of both the RMSE and the ASA. The results obtained with Ext slightly improved the ones obtained with FCLS. Khype outperformed both methods, Ext and FCLS, and had results very close to the ones obtained with NDU. Figure 7 shows that the abundance maps estimated by the various algorithms are rather similar. As mentioned previously, the fourth endmember corresponds to noise hence its abundance maps are not shown in Figure 7. Figure 8 shows the nonlinear part estimated by NDU and khype at band 10. In fact, most of the areas where nonlinear contributions appear are mainly located on the boundaries of agricultural areas (endmember 2) surrounded by forests and semi-natural areas (endmember 3). Note that the nonlinear contributions estimated by NDU are relatively spatially smoother than the ones estimated by khype. However, NDU results exhibit some artifacts due to the fact that the

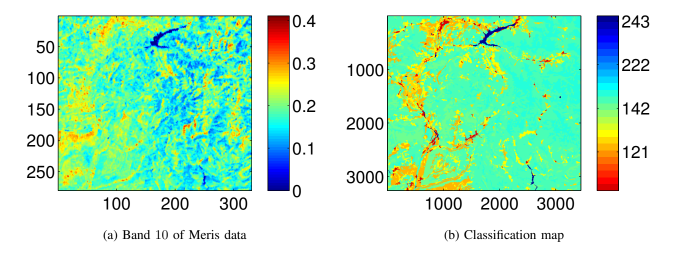


Fig. 5. Fom left to right: (a) band 10 of the Meris data set, (b) corresponding CLC classification map (See classes names in Appendix A).

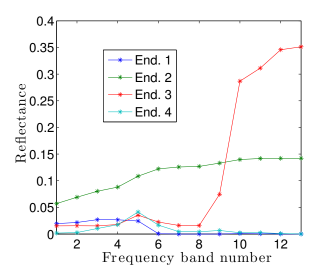


Fig. 6. Endmembers spectra estimated by VCA for the Meris data set.

image was partitioned into square patches.

Finally, Figure 9 compares the nonlinear contribution estimated by NDU and khype at all bands for the pixels delimited by rows 171 and 180 and columns 231 and 240. Both algorithms estimated the highest nonlinear contributions for this particular region. However, the nonlinear contributions have different variations throughout the spectral bands. NDU estimated the highest nonlinear contributions at the higher frequency bands whereas khype estimated almost the same level of nonlinearity at all frequency bands. Furthermore, NDU estimated smooth nonlinear contributions at adjacent pixels compared to khype. It can be concluded that NDU captures more spectral variability throughout the spectral bands and that it provides smooth nonlinear contributions at adjacent pixels.

TABLE IV ROOT MEAN SQUARE ERROR RMSE $(\times 10^{-2})$ and average spectral angle (ASA) in radian of the reconstructed spectra obtained with the Meris data set.

	RMSE	ASA
FCLS	1.14	0.0393
Ext	1.13	0.0343
Khype (G)	0.66	0.0338
Khype (P)	1.04	0.0381
Sep. (G)	0.41	0.0153
Sep. (P)	0.46	0.0204

V. CONCLUSION

This paper proposed a new kernel based nonlinear mixing model for hyperspectral data. The proposed vector-valued function is able to account for band dependent and neighboring nonlinear contributions. The proposed framework has several characteristics. It allows to handle in a unified framework several types of nonlinearities depending on the choice of the kernel function. Unlike nonlinear models proposed in the literature, it considers a different nonlinear function for each spectral band. The fact that the nonlinear function acts on the reflectance vectors observed in the corresponding pixel and its neighbors is intended to account for nonlinearities originating from the ground cover of the pixel and its neighbors. Furthermore, the separable kernel design can be used to incorporate prior information regarding the similarities between nonlinear contributions at different bands. In particular, a linear graph was proposed to promote smooth nonlinear variations between adjacent bands. The performance of the proposed approach was validated on synthetic and real data estimated by the Meris spectrometer and captured over the gulf of Lion in the south east of France. Finally, note that the proposed approach requires partitioning the image into patches which can result in artifacts in the estimated nonlinear part. Future work should aim at attenuating those artifacts through an extension of the vector-valued approach. More precisely, the vector-valued framework can be adapted such as to promote smoothness between the estimated nonlinear contributions at adjacent bands and between the nonlinear functions at adjacent patches simultaneously.

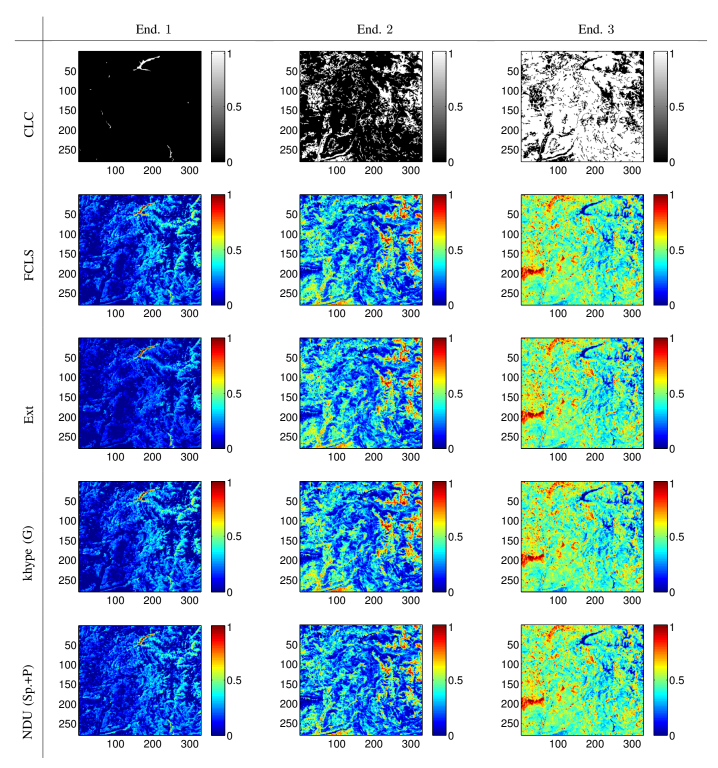


Fig. 7. Abundance maps of the first three endmembers obtained with VCA and corresponding to the Meris real data set. The abundance maps of End. 1, 2, and 3 correspond to water, agricultural areas, and forests and semi natural areas respectively.

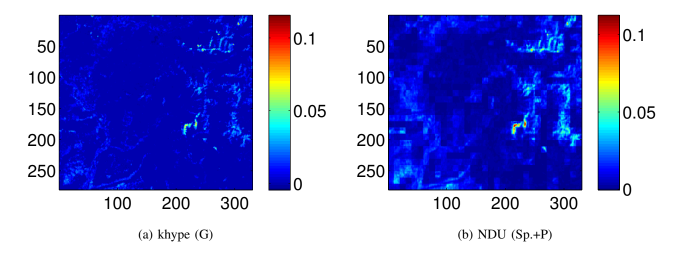


Fig. 8. Nonlinear contributions at all pixels at band 10 obtained with: (a) khype used with a Gaussian kernel (G) and (b) NDU used with a separable and polynomial kernel (Sp.+P).

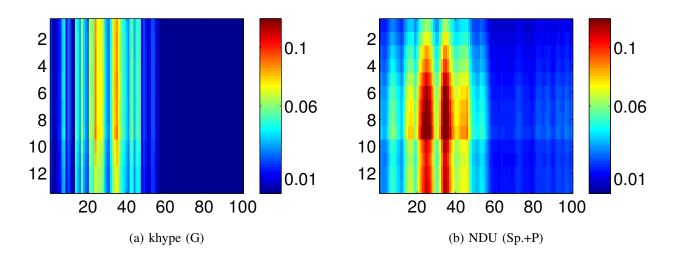


Fig. 9. Nonlinear contributions at all bands in some of the pixels in the Meris data set. The horizontal and vertical axis correspond to the frequency and pixel index respectively.

 $\label{eq:table v} TABLE\ V$ Classes in classification map of Meris data set.

Code	Color	description		
1		Artificial surfaces		
1.1		Urban fabric		
1.1.1		Continuous urban fabric		
1.1.2		Discontinuous urban fabric		
1.2		Industrial, commercial and transport units		
1.2.1		Industrial or commercial units		
1.2.2		Road and rail networks and associated land		
1.2.4		Airports		
1.3		Mine, dump and construction sites		
1.3.1		Mineral extraction sites		
1.4		Artificial, non-agricultural vegetated areas		
1.4.2		Sport and leisure facilities Not translated		
2		Agricultural areas		
2.1		Arable land		
2.1.1		Non-irrigated arable land		
2.2		Permanent crops		
2.2.1		Vineyards		
2.2.2		Fruit trees and berry plantations		
2.3		Pastures		
2.3.1		Pastures		
2.4		Heterogeneous agricultural areas		
2.4.2		Complex cultivation patterns		
2.4.3		Land principally occupied by agriculture		
2.4.4		Agro-forestry areas		
3		Forest and seminatural areas		
3.1		Forests		
3.1.1		Broad-leaved forest		
3.1.2		Coniferous forest		
3.1.3		Mixed forest		
3.2		and/or herbaceous vegetation associations		
3.2.1		Natural grasslands		
3.2.2		Moors and heathland		
3.2.3		Sclerophyllous vegetation		
3.2.4		Transitional woodland-shrub		
3.3		Open spaces with little or no vegetation		
3.3.1		Beaches, dunes, sands		
3.3.2		Bare rocks		
3.3.3		Sparsely vegetated areas		
3.3.5		Glaciers and perpetual snow		
5		Water bodies		
5.1		Inland waters		
5.1.1		Water courses		
5.1.2		Water bodies		

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