Abstract—A Metropolis-within-Gibbs sampler for piecewise convex hyperspectral unmixing and endmember extraction is presented. The standard linear mixing model used for hyperspectral unmixing assumes that hyperspectral data reside in a single convex region. However, hyperspectral data are often nonconvex. Furthermore, in standard endmember extraction and unmixing methods, endmembers are generally represented as a single point in the high-dimensional space. However, the spectral signature for a material varies as a function of the inherent variability of the material and environmental conditions. Therefore, it is more appropriate to represent each endmember as a full distribution and use this information during spectral unmixing. The proposed method searches for several sets of endmember distributions. By using several sets of endmember distributions, a piecewise convex mixing model is applied, and given this model, the proposed method performs spectral unmixing and endmember estimation given this nonlinear representation of the data. Each set represents a random simplex. The vertices of the random simplex are modeled by the endmember distributions. The hyperspectral data are partitioned into sets associated with each of the extracted sets of endmember distributions using a Dirichlet process prior. The Dirichlet process prior also estimates the number of sets. Thus, the Metropolis-within-Gibbs sampler partitions the data into convex regions, estimates the required number of convex regions, and estimates endmember distributions and abundance values for all convex regions. Results are presented on real hyperspectral and simulated data that indicate the ability of the method to effectively estimate endmember distributions and the number of sets of endmember distributions.

Index Terms—Endmember, hyperspectral, Markov chain Monte Carlo (MCMC), piecewise convex, sampling, spectral variation, unmixing.

I. INTRODUCTION

THE linear mixing model (LMM) represents the spectral signatures in a hyperspectral scene as convex combinations of endmembers. As stated in [1], endmembers are often defined as the spectral signatures of the distinct substances in a hyperspectral data set. The equation and constraints defining the LMM are as follows:

\[ x_j = \sum_{k=1}^{M} p_{jk} e_k + \epsilon_j. \] (1)

Here, \( N \) is the number of pixels, \( M \) is the number of endmembers, \( \epsilon_j \) is an error term, \( p_{jk} \) is the proportion of endmember \( k \) in pixel \( j \), and \( e_k \) is the \( k \)th endmember. The proportions satisfy the following constraints:

\[ p_{jk} \geq 0 \quad \forall k = 1, \ldots, M \] (2)

\[ \sum_{k=1}^{M} p_{jk} = 1. \] (3)

Mathematically, the LMM assumes that endmembers are vertices of a simplex that approximately encloses the spectra, or data points, present in an image. The approximation is represented by the error term.

Several methods have been developed for unmixing based on the LMM. These include methods, such as vertex component analysis (VCA) [2], that rely on the pixel purity assumption and assume that the endmembers can be found within the data set [2]–[5]. Methods have also been developed based on nonnegative matrix factorization [6]–[11], independent component analysis [12]–[14], and others [15]–[18]. All these methods search for a single set of endmembers and, therefore, a single convex region to describe a hyperspectral scene. Since these algorithms assume a single convex region, they cannot find appropriate endmembers for nonconvex data sets.

If a scene contains multiple distinct regions that do not share common materials and if each region contains linear mixtures of materials, then the set of all image spectra will consist of a union of simplices. A single simplex is convex, but the union of simplices is unlikely to be convex. Therefore, a piecewise convex model is a more appropriate model than a single convex region. Moreover, the extremal points of the individual convex sets may appear to be interior points in the convex hull of all image pixels. Hyperspectral images often exhibit these characteristics.

Consider the image shown in Fig. 1. This real hyperspectral data set is nonconvex and would be better represented with a piecewise convex representation of the data. By examining these nonconvex sets of spectra from hyperspectral images, endmembers may appear within the convex hull defined by the other endmembers in the scene. These interior endmembers cannot be recovered using methods based on the standard
Furthermore, the proposed method estimates the number of random simplices needed to describe the data set.

The S-PCUE method is a fully stochastic unmixing algorithm based on a piecewise convex model. The use of a piecewise convex representation was first presented in [21], [22], and [23]. The algorithms represented there will be referred to as piecewise convex endmember (PCE). The S-PCUE method differs from PCE because S-PCUE fully uses a complete Bayesian inference strategy whereas PCE does not. Although PCE uses the Dirichlet process for partitioning the data points into endmember sets, the endmembers and proportions were estimated by maximizing an objective function. Therefore, PCE is a stochastic expectation-maximization (EM) algorithm. The S-PCUE algorithm provides a fully stochastic extension of these methods by using a Gibbs sampling approach to sample all desired parameters. The S-PCUE algorithm estimates several sets of endmembers, the abundances for each data point, and the number of endmembers sets needed to represent a hyperspectral image. The proposed S-PCUE algorithm improves on the stochastic EM-type algorithms in both time complexity and convergence guarantees. Since the S-PCUE algorithm does not require a maximization step each iteration, the time per iteration is drastically reduced. Furthermore, since the proposed algorithm satisfies Markov chain Monte Carlo (MCMC) convergence properties, the proposed method inherits MCMC convergence guarantees.

II. S-PCUE METHOD

The S-PCUE method represents endmembers as random vectors and uses a Metropolis-within-Gibbs sampling technique to extract their values. Representing endmembers as random vectors provides the capability of representing the spectral variability associated with a particular material or environmental condition.

A. Model Description

Each set of endmember distributions represents the materials that mix together in some subset of the image scene. Each pixel in the scene is assumed to be a mixture of endmembers from one and only one set of endmember distributions. Therefore, the set of sets of endmember distributions defines a partition of the image pixels; that is, $X$ is a disjoint union $X = \bigcup_{r=1}^{R} \Gamma_r$. Each subset $\Gamma_r$ is assumed to satisfy the LMM with endmembers that are distributions. We therefore refer to them as convex sets. The number of endmember distribution sets is estimated by sampling from a Dirichlet process. In the current implementation, the endmembers are modeled using Gaussian distributions with a fixed isotropic diagonal covariance $e_{m,r} \sim N(e_{m,r}, S_{m,r})$, where $e_{m,r}$ is the mean value for the $m$th endmember distribution in the $r$th endmember distribution set and $S_{m,r}$ is the covariance for the $m$th endmember distribution in the $r$th convex set. In the current implementation, all endmember distributions are given the same fixed isotropic diagonal covariance $S$. Therefore, each of these covariance matrices is proportional to the identity matrix.

It is assumed that the LMM holds for the set of data points represented by each set of endmember distributions. Since
the endmembers are modeled using Gaussian distributions, the current implementation of S-PCUE follows the normal compositional model [20] in which endmembers are random vectors represented by Gaussian distributions and hyperspectral data points are random vectors distributed according to a convex combination of the associated Gaussian endmember distributions. The identity of the endmember set for each hyperspectral data point is unknown and is represented by a latent variable. Thus, the likelihood for a data point assigned to a single convex region is given by

\[
x_j \mid \mathbf{E}_r, \mathbf{p}_j \sim f(x_j \mid \mathbf{E}_r, \mathbf{p}_j) = N \left( \mathbf{p}_j \mathbf{E}_r, \sum_{m=1}^{M} p_{jm}^2 \mathbf{S}_{m,r} \right)
\]  

(4)

where \( x_j \) is the \( j \)th data point, \( r \) is the indicator variable for the \( r \)th set of endmembers, \( \mathbf{E}_r \) is a matrix whose rows are the endmember means for the \( r \)th set of endmembers, and \( \mathbf{p}_j \) is the vector of proportion values associated with the \( j \)th data point where \( p_{jm} \) is the \( m \)th element of this proportion vector and \( M \) is the number of endmember distributions in the \( r \)th partition. A somewhat subtle point that is noted here is that there is assumed to be only one true set of proportions for each data point. However, there are \( M \) estimates of the set of proportions, one for each partition. Let \( z_j \) denote the latent variable representing the partition to which the data point \( x_j \) is assigned. Then, the overall likelihood can be written as

\[
\prod_{r=1}^{R} \prod_{j \in I_r} f(x_j \mid \mathbf{E}_r, \mathbf{p}_j)
\]

where \( R \) is the number of convex sets, \( I_r = \{ j \mid z_j = r \} \subset \{1, \ldots, N\} \) denotes the set of indices of the data points that are assigned to the \( r \)th convex set, and \( \mathbf{E} = \{ \mathbf{E}_1, \ldots, \mathbf{E}_R \} \) denotes the set of endmember mean matrices.

For each partition, the means of the associated endmember distributions are assumed to share a Gaussian prior distribution

\[
e_{m,r} \sim N(\mu_{r}, \mathbf{C}_r), \quad m = 1, \ldots, M
\]  

(5)

where \( \mu_{r} \) and \( \mathbf{C}_r \) are the mean vector and full covariance hyperparameters, respectively, in the Gaussian prior distribution governing the \( r \)th set of endmember distribution means. These hyperparameters are also estimated within the S-PCUE algorithm. This prior for the mean of the endmember distributions of a partition was chosen for a number of reasons. First, it encourages the endmember distributions for each partition to have a smaller enclosed volume. In other words, the mean endmembers for each endmember set share a prior distribution that encourages the endmember distributions to have a tight fit around the data. Second, estimating a full covariance hyperparameter for each partition allows for a tight fit around data whose collective shape and size differ from partition to partition. Finally, this distribution is a conjugate prior to endmember distributions allowing for an efficient Gibbs sampling step.

The prior on all of the means over all sets \( \mu \) is also given by a Gaussian distribution whose mean is fixed at the mean of the input hyperspectral data and whose covariance is \( \mathbf{C}_\mu = \mathbf{I} \sigma_\mu^2 \), where \( \sigma_\mu \) is fixed to a large value relative to the spread of the entire input data set. This practice, of using marginal means to estimate hyperparameters, is in the spirit of empirical Bayes analysis [27]

\[
\mu_{r} \sim N \left( \frac{1}{N} \sum_{j=1}^{N} x_j, \mathbf{C}_\mu \right).
\]  

(6)

The prior used for the \( \mathbf{C}_r \) covariance matrices is the inverse-Wishart distribution. This prior distribution was chosen as it is the conjugate prior of the Gaussian distribution over a full covariance matrix.

The proportion values for all the data points in the image are given a Dirichlet prior

\[
p_{j} \mid z_j = r \sim D_M(\alpha_{1,r}, \ldots, \alpha_{M,r})
\]  

(7)

where \( D_M(\cdot) \) denotes the \( M \)-factor Dirichlet distribution whose density function is given by

\[
D_M(p_j \mid z_j = r) = \frac{\Gamma \left( \sum_{m=1}^{M} \alpha_{m,r} \right)}{\prod_{m=1}^{M} \Gamma(\alpha_{m,r})} \prod_{m=1}^{M} p_{jm}^{\alpha_{m,r} - 1}.
\]

By fixing the alpha values to one, the endmembers are encouraged to have a tight fit around the data [17]. In the current implementation, all alpha values are fixed to one. However, one could choose to use other values for each \( \alpha \). In the current implementation, the number of endmembers \( M \) is a set parameter fixed to a constant value across all partitions.

### B. Sampling Method

The S-PCUE algorithm iteratively samples the parameters and hyperparameters of interest using a Metropolis-within-Gibbs sampling method. The algorithm is outlined in the pseudocode shown in Algorithm 1. In the following, each step of the sampling algorithm, initialization, and the methods used for setting algorithm parameters are described.

**Algorithm 1—S-PCUE:** A Metropolis-within-Gibbs sampler partitions the data into convex regions, estimates the required number of convex regions, and estimates endmember distributions and abundance values for all convex regions. The method used for each step is described in the section shown in parentheses

1: Set Parameter Values and Initialize Partition (Section II-C)
2: for \( r \leftarrow 1 \) to \( R_{\text{initial}} \) Convex sets do
3: Initialize \( \mathbf{E}_r, \mathbf{P}_r, \) and \( \mathbf{C}_r \) (Section II-C)
4: end for
5: for \( k \leftarrow 1 \) to number of total iterations do
6: for \( r \leftarrow 1 \) to number of convex sets do
7: for \( j \leftarrow 1 \) to number of data points do
8: Sample proportions \( p_{j} \) for \( x_j \) for each set of endmembers (Section II-B1)
9: end for
10: for \( k \leftarrow 1 \) to number of endmembers in convex set \( r \) do

11: Sample \(e_{k,r}\) in convex set \(r\) (Section II-B2)
12: end for
13: Sample \(\mu_r\) (Section II-B3)
14: Sample \(C_r\) (Section II-B4)
15: end for
16: for \(k \leftarrow 1\) to \(K\) do
17: Sample new \(E_k^r\) and \(P_k^r\) matrices (Section II-B5)
18: end for
19: for \(j \leftarrow 1\) to number of data points do
20: Remove \(x_j\) from its current convex set
21: Compute DP partition probabilities for \(x_j\) using (12) and (13).
22: Sample a convex set for \(x_j\) based on the DP partition probabilities (Section II-B6)
23: if A new convex set is sampled then
24: Add the new endmember distribution set to \(E\) and assign \(x_j\) to this set
25: else
26: Update the label of \(x_j\) to the sampled endmember distribution set
27: end if
28: end for
29: end for

1) Sample Proportion Values: The proportion vectors for each data point and each set of endmembers are sampled in S-PCUE using a Metropolis--Hastings step. For implementation, a set of proportions is sampled for each data point for each set of endmember distributions. This is done to be able to compute likelihood values using appropriate proportion vectors for each set of endmembers. The Dirichlet prior shown in (7) is used as the proposal distribution. This results in the acceptance ratio shown in (8) used to accept or reject new proportion vector samples for each data point in each partition

\[
\begin{align*}
a &= \frac{\prod (p_j^{\text{new}} | X, E, z_j = r) D_M (p_j^{\text{old}} | z_j = r)}{\prod (p_j^{\text{old}} | X, E, z_j = r)} \\
&= \frac{f (x_j | X, p_j^{\text{new}}, z_j = r)}{f (x_j | X, p_j^{\text{old}}, z_j = r)}
\end{align*}
\]

where

\[
\prod (p_j | X, E, z_j = r) \propto f (x_j | X, p_j, z_j = r) D_M (p_j | z_j = r).
\]

The final equality is found since the proposal distribution is also the prior on the proportion vectors. In summary, the new proportion sample is found to be \(p_j^{\text{new}} = p_j^{\text{new}} \gamma + p_j^{\text{old}} (1 - \gamma)\), where \(\gamma = I(u < \min \{f (x_j | X, p_j^{\text{new}}, z_j = r) / f (x_j | X, p_j^{\text{old}}, z_j = r), 1\})\). Here, \(I(m) = 1\) when \(m\) is true and zero when \(m\) is false and \(u\) is randomly drawn from the uniform distribution over \([0, 1]\).

2) Sample Endmember Distribution Values: A Metropolis--Hastings step is also used to sample the means of each endmember distribution. The proposal distribution is a Gaussian mixture centered on the previous mean endmember value. The first Gaussian in the mixture has a diagonal covariance with small values, and the second Gaussian has a diagonal covariance with large values

\[
g (e_{m,r}^{\text{new}}, e_{m,r}^{\text{old}}) = w_n N (e_{m,r}^{\text{new}}, e_{m,r}^{\text{old}}, C_n) + (1 - w_n) N (e_{m,r}^{\text{new}}, e_{m,r}^{\text{old}}, C_w)
\]

where \(w_n\) is a fixed parameter used to determine the relative frequency sampling from a Gaussian with diagonal covariance whose diagonal covariances are either small or large. The covariance matrices \(C_n\) and \(C_w\) are fixed covariances used to generate the endmember samples. In all experimental results shown here, both \(C_n\) and \(C_w\) are fixed parameters set to isotropic diagonal covariance matrices.

The acceptance ratio will be

\[
a = \frac{\prod (e_{m,r}^{\text{new}} | X, P, z) g (e_{m,r}^{\text{old}} | e_{m,r}^{\text{new}})}{\prod (e_{m,r}^{\text{old}} | X, P, z) g (e_{m,r}^{\text{new}} | e_{m,r}^{\text{old}})}
\]

with \(f (e_{m,r} | \mu_r, C_r)\) as the distribution in (5).

As stated in Section II-A, the covariance for each endmember distribution is fixed to a diagonal isotropic matrix in the current implementation.

3) Sample Endmember Set Means: The hyperparameters \(\mu_r\), which represent the mean of the Gaussian prior for a set of endmembers, are sampled using a Metropolis--Hastings set. A Gaussian mixture is used as the proposal distribution. This step mimics the previous step described in Section II-B2 for sampling endmember distribution means. In the current implementation, the same Gaussian mixture used to generate new endmember samples in the previous step is used to generate the \(\mu_r\) samples. Similarly, the mixture is centered on the previous \(\mu_r\) value.

4) Sample Endmember Set Covariance: The covariance for each endmember set \(C_r\) is assumed to have an inverse-Wishart prior \(C_r \sim \text{InvWishart}(\Psi, t)\). Given that the inverse-Wishart is the conjugate prior to the Gaussian distribution with a full covariance, the covariance for each set is sampled directly using a Gibbs step from the posterior inverse-Wishart distribution resulting from the product of the likelihood of all of the endmember distribution means for a partition \(\prod_{m=1}^M N (e_{m,r} | \mu_r, C_r)\) and the \(\text{InvWishart}(\Psi, t)\) prior.

5) Sample Endmembers and Proportion for Potential New Partitions: Prior to sampling a partition label for each data point given updated endmember and proportion values, \(K\) new sets of endmember distributions and associated proportion values are sampled. These new sets allow for the addition of new convex sets during the update of the partition labels. This method of sampling \(K\) new endmember distribution sets follows from the method described in [28]. The number of new sets considered \(K\) is a fixed parameter in the implementation.

The proportion values for each data point and each of the \(K\) new endmember sets are drawn from the Dirichlet prior for the proportion values as defined in (7). The \(K\) new sets
of endmembers are drawn following the method defined in Algorithm 2.

Algorithm 2—Sampling New Endmember Sets

1: for $k \leftarrow 1$ to $K$ do
2: Sample $\mu_{R+k}$ from the Gaussian defined in (6)
3: Sample $C_{R+k}$ from the inverse-Wishart prior
4: for $m \leftarrow 1$ to $M$ do
5: Sample $e_{m,R+k}$ from the Gaussian centered at $\mu_{R+k}$ with covariance $C_{R+k}$
6: end for
7: end for

6) Sample Partition Labels: The labels $r$ are distributed according to a Dirichlet process. These labels determine the number of endmember distribution sets needed to describe an input hyperspectral data set as well as the partitioning of the data points into the various endmember distribution sets. The likelihood of a data point belonging to a convex region is computed for each existing $E$ and $P$ set and all potential new endmember sets that were sampled in the previous step as shown in

$$P(z_i = z_j | \mathbf{z}_{-i}, \mathbf{x}_i) = C \frac{n_{-i,j}}{\alpha + N - 1} f(\mathbf{x}_i | \mathbf{p}_r, \mathbf{E}_r, z_j = r) f(\mathbf{E}_r) f(\mathbf{p}_i | z_i = r),$$

$$j = 1, \ldots, R, j \neq i$$

$$P(z_i = R + 1 | \mathbf{z}_{-i}, \mathbf{x}_i) = C \frac{\alpha}{\alpha + N - 1} f(\mathbf{x}_i | \mathbf{p}_r^*, \mathbf{E}_r^*) f(\mathbf{E}_r^*) f(\mathbf{p}_i^*)$$

where $z_i$ is the indicator variable for the current data point $\mathbf{x}_i$, $C$ is a normalization constant, $n_{-i,j}$ is the number of data points excluding $\mathbf{x}_i$ in convex set $z_j$, $N$ is the total number of data points, $K$ is the number of new endmember distribution sets sampled, and $\alpha$ is the innovation parameter for the Dirichlet process.

C. Initialization and Parameter Settings

S-PCUE requires several parameters to be set prior to running the algorithm. For all experimental results shown, initialization for the algorithm and parameters are determined using the following methods.

1) Initialization: The initial convex sets and labels for each data point are determined by fitting the data to a Gaussian mixture using EM with random initialization. During each iteration of the EM algorithm, Gaussian components are checked to ensure that the covariance matrix for the corresponding Gaussian is not becoming singular. If that is the case, the Gaussian is removed, and the associated points are reassigned to the remaining Gaussians. Following the application of the EM algorithm, endmembers are initialized for each convex set using the VCA algorithm [2]. Finally, proportions for each data point are initialized randomly by drawing from a uniform Dirichlet distribution.

2) Parameter Settings: As described in Section II-A, the parameters for the Gaussian prior on the mean of the endmember distributions are to set to the mean of the data and variance of the data $\sigma_\mu = \text{var}(\mathbf{X})$.

Section II-B2 states that the means of the endmember distributions are sampled from a mixture of two Gaussians, one with a small covariance and another with a large covariance. Both Gaussians are centered on the previous endmember mean value. The covariances are set using an order weighted average. To compute these values, the pairwise Euclidean distances between each pair of input data points are computed and then sorted. The small covariance $C_n$ is set to the mean of the smallest pairwise distance for each data point. The large covariance $C_w$ is set to the mean of the median pairwise distance for each data point. For all runs of the algorithm, $w_n$ was set to 0.9 such that the majority of new means are drawn close to the previous value.

As described in Section II-B6, the sample labels are drawn from a Dirichlet process. The parameters required for this step are the number of new partitions considered $K$ and the innovation parameter $\alpha$. The primary consideration in setting the $K$ parameter is the time complexity per iteration. Larger values of $K$ require longer running time per iteration. $K = 5$ was found to be a good balance of running time, and the number of new partitions is considered. The innovation parameter is set to $\alpha = K/N$, where $N$ is the number of input data points.

The covariance governing each partition $C_r$ is sampled from an inverse-Wishart distribution as described in Section II-B4.

Fig. 2. (a) Results found using VCA algorithm on simulated data. (b) Results found using SPICE algorithm on simulated data.
The parameters for the inverse Wishart are set to $\mathbf{\Psi} = \mathbf{I}$, and $w$ is set to a scaled version of the mean of variances of the initial partitions.

III. RESULTS

The sample PCUE algorithm was run on simulated data and the Airborne Visible Infrared Imaging Spectrometer (AVIRIS) Indian Pines and Cuprite scenes. Results are shown on these data sets and discussed in the following sections.

A. Simulated Data

The S-PCUE algorithm was run on simulated data generated from three sets of two endmember distributions. The purpose of this simulated data set was to show that the proposed method is able to recover interior endmembers that are not found using conventional single partition unmixing algorithms. Each data point in this simulated set was generated as a convex combination of two endmembers with a total of three sets of endmember pairs. Simulated data generated in this fashion were selected to simulate the pairwise mixing seen in hyperspectral imagery such as that shown in [29]. Fig. 3(a) shows the simulated data set. The results found using the parameters settings described in Section II-C are shown in Fig. 3(b). As can be seen, the data are appropriately partitioned, and correct endmembers are estimated. For comparison, the sparsity promoting iterated constrained endmembers (SPICE) [30] and VCA [2] algorithms were also run on this data set. Both methods did not find the endmember located at (2, 3, 2).

The algorithm was run for 50,000 iterations and, from the 50,000 samples generated, the majority of samples partitioned the data into three convex regions, as shown in Fig. 3(c). The result shown in Fig. 3(b) was selected by, first, determining the number of convex sets found by the majority of the samples and, then, returning the result that had the largest likelihood and the appropriate number of convex sets. This method for selecting the final result was used in all of the runs of S-PCUE.

B. AVIRIS Indian Pines

S-PCUE was applied to the June 1992 AVIRIS Indian Pines data set as well, shown in Fig. 1. This data set, collected over an agricultural area in northern Indiana, was collected during very early growth stages. The data in this image are highly mixed resulting from the fields having very low crop cover and varying levels of residue from the previous year’s crops. The S-PCUE algorithm was applied to a subset of the points selected from the image (every tenth labeled pixel, resulting in 1031 pixels) for 50,000 iterations using the parameter settings described in Section II-C. As shown in the results in Fig. 4, the data are appropriately partitioned into three convex regions: (partition 1) woods, grass, and trees; (partition 2) hay, corn, and soybean; and (partition 3) stone Steel towers. For example, the algorithm pulls out a distinct hay endmember, whose proportion map is shown in Fig. 5(b). The soybean and corn fields, which are primarily soil and crop residue, are shown in the second two endmembers in Fig. 5(b).

For comparison, the SPICE and VCA algorithms were applied to the AVIRIS Indian Pines data set as well. For SPICE, the parameters used were $\mu = 0.001$ and $\gamma = 1$. SPICE resulted in estimating seven endmembers after initializing with 20 endmembers. The resulting proportion maps estimated by SPICE are shown in Fig. 7. VCA was used to estimate nine endmembers from the Indian Pines data set. VCA was set to estimate nine endmembers since this was the number of endmembers found by the proposed S-PCUE algorithm. The
Fig. 4. Scatter plot of results found on the AVIRIS Indian Pines data set after PCA dimensionality reduction from 220 to 3 dimensions. The large points correspond to endmembers in the scene.

Fig. 5. Proportion maps found after unmixing using the endmembers estimated from the S-PCUE algorithm. (a) Proportion maps corresponding to endmembers found for partition 1. This partition corresponds primarily to woods, grass, and trees. (b) Proportion maps corresponding to endmembers found for partition 2. This partition corresponds primarily to hay (endmember 1), corn, and soybean (endmembers 2 and 3, respectively). (c) Proportion maps corresponding to endmembers found for partition 3. This partition corresponds primarily to stone-steel towers.

Fig. 6. Color bar corresponding to all resulting proportion maps. Proportion values of zero correspond to dark blue, and proportion values of one correspond to dark red.

Fig. 7. Proportion maps found after unmixing using the endmembers estimated from the SPICE algorithm.

proportion maps corresponding to the endmembers estimated by VCA are shown in Fig. 8. Neither the SPICE nor VCA algorithm was able to estimate the strong hay endmember found by the S-PCUE algorithm. However, both SPICE and VCA were able to estimate a single stone-steel tower endmember which the S-PCUE assigned to a separate convex region (and, thus, three endmembers). As discussed in the following, future work will include estimating the number of endmembers needed per convex region. In the case of the convex region associated with stone-steel towers here, only a single endmember may be needed. The color bar used for all proportion maps is shown in Fig. 6.

C. AVIRIS Cuprite

S-PCUE was also applied to the AVIRIS Cuprite data set collected over Cuprite, NV. This area has little vegetation and visible mineral coverage. A mineral map of the area can be found in [31]. The S-PCUE algorithm was applied to a subset of the points selected from the image (every 101st pixel, resulting in 3113 pixels) for 50,000 iterations using the parameter settings described in Section II-C. Fig. 9 shows the 3-D plot of the data and the endmembers found following principal component
analysis (PCA) dimensionality reduction. Fig. 10 shows the proportion maps associated with each of the estimated endmembers over the full data set.

The endmembers found by S-PCUE were compared to spectra in the United States Geological Survey (USGS) spectral library. The spectra with the smallest spectral angle distance to the estimated endmembers were identified. These spectra are shown in Fig. 11. The average spectral angle distance between the estimated endmembers and identified spectra in the USGS spectral library was 0.037.

For comparison, the SPICE and VCA algorithms were run on the same AVIRIS Cuprite data. The VCA algorithm was set to estimate nine endmembers since S-PCUE found nine endmembers on this data. The SPICE parameters used were $\mu = 0.001$ and $\gamma = 1$. SPICE found nine endmembers after
Fig. 11. (Red) Estimated endmembers found using S-PCUE on the AVIRIS Cuprite data and (black) USGS spectral library spectra. The USGS spectra were selected by having the smallest spectral angle distance to the estimated endmembers. The USGS spectra selected for each endmember were (a) muscovite-medhi-Al Cu91-252d.26143, (b) buddingtonite cu93-260b.24428, (c) alunite na dickite_mv99-6-260.23945, (d) hematite_gds27.29282, (e) hematite_coat_br93-25a.26778, (f) calc_mont_amx6.24507, (g) buddingtonite_cu93-260b.24428, (h) alunite_gds82.1019, and (i) hematite_coat_br93-25a.26778.

Fig. 12. (Red) Estimated endmembers found using SPICE on the AVIRIS Cuprite data and (black) USGS spectral library spectra. The USGS spectra were selected by having the smallest spectral angle distance to the estimated endmembers. The USGS spectra selected for each endmember were (a) calcite_gds304.4124, (b) chlorite_mixture_cu93-65a.24855, (c) nontronite_sw1.16168, (d) montmorillonite_stx1.14644, (e) halloysite_kaolinite_cm29.25328, (f) montmorillonite-Na_cu93-52.25995, (g) buddingtonite_gds85.3924, (h) jarosite_gds98.11375, and (i) alunite_cu91-217g1.23924.

pruning 11 of the 20 initial endmembers. Figs. 12 and 13 show the endmembers found and their closest matching spectra in the USGS spectral library. The average spectral angle distance between the SPICE and VCA endmembers and spectral library spectra were 0.136 and 0.071, respectively. Thus, the proposed method estimated endmembers that have a smaller spectral angle distance between minerals matched in the USGS spectral library.

IV. CONCLUSION AND FUTURE WORK

The S-PCUE algorithm presented here uses a Gibbs sampling approach to estimate sets of endmember distributions, proportion values, and the number of endmember distribution sets needed to describe a nonconvex hyperspectral image. Thus, the proposed method provides an endmember estimation and spectral unmixing algorithm given a nonlinear representation of the data. Future work to further develop this method will include extending the algorithm to estimating covariance parameters for each endmember distribution and the number of endmember distributions per set. For example, consider the stone-steel tower partition shown in Fig. 5(c). This partition may be best represented with one or two endmembers; however, the current algorithm is restricted to the same number of endmembers per convex region. Currently, the method also does not restrict the endmembers to be nonnegative which is generally the case for hyperspectral imagery. Since, after applying dimensionality reduction, such as PCA or maximum noise fraction (MNF), the resulting data often have negative components, this restriction was not included. However, future work can include exploring
the use of truncated Gaussians for endmember proposal and prior distributions to restrict the endmembers to be nonnegative. Also, incorporating spatial correlations is an area of future work [32].

The proposed method applies a fully stochastic MCMC algorithm to a high-dimensional problem. The advantage of this is that the algorithm is guaranteed to converge to the joint distribution over the parameters. The implication is that one can truly expect to find globally optimal solutions by examining these distributions. Therefore, MCMC algorithms will almost certainly eventually find better solutions than algorithms that find local optima. Given these excellent convergence properties, it seems that MCMC algorithms should always be used. The difficulty is that the speed of the algorithm is slow compared to many existing unmixing algorithms in the literature. Furthermore, it is not obvious how to speed up such an algorithm since they appear to be inherently sequential. Generally, one cannot run Markov chains in parallel and take the union of the results while maintaining the convergence properties because they are different chains. However, Gopal [33] has made a tremendous theoretical breakthrough in the area of parallelization of MCMC algorithms. Roughly speaking, he has devised and demonstrated a method for running Markov chains in parallel in a way that maintains the convergence properties. Thus, MCMC methods can now be implemented correctly on clusters of computers consisting of tens, hundreds, or thousands of processors. Currently, their methods require some analysis of each specific problem. It seems clear that, once these methods are further developed, MCMC methods will become extremely attractive and potentially be the standard method for parameter estimation for difficult high-dimensional problems such as hyperspectral image analysis. The work described here, together with the research of Dobigeon et al. [17, 34, 35] and others [22], [36], represents progress in that direction. Hyperspectral unmixing is a very complex problem in general. In addition to piecewise convexity, there are a number of sources of nonlinearity that are only beginning to be considered for large-scale data sets. Complete analyses of hyperspectral data with multiple scattering, intimate mixtures, and multiple image components will certainly benefit from the type of research described in this paper.

REFERENCES


