Mixture of Latent Variable Models for Remotely Sensed Image Processing

by

Linlin Xu

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in
Geography

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Author’s Declaration

I hereby declare that I am the sole author of this thesis. This is a true copy of the thesis, including any required final revisions, as accepted by my examiners.

I understand that my thesis may be made electronically available to the public.
Preface Statement

The doctoral dissertation is compiled under the manuscript option, following the guidelines provided by the joint Waterloo-Laurier Graduate Program in Geography. Three manuscripts published in refereed journals, as listed below, are presented in Chapters 3 to 5 respectively, where the manuscripts are minorly changed for consistence formatting.


In all three manuscripts, I am the first author and my supervisor Prof. Dr. Jonathan Li is corresponding author. They are dominated by my intellectual effort. The roles of coauthors are explained in detail below.

The ideas in the first manuscript [1] were conceived by me. And I carried out the work, including experiments design and implementation, manuscript writing and revision. Dr. Li actively participated in the discussion of the experiment results and reviewing of the manuscript. In [2], I conceived the key ideas, conducted the experiments, wrote the manuscript and performed the revisions. The other coauthors participated in the reviewing of the manuscript. Dr. Wong provided critical suggestions on experimental design and reference choice. In [3], I initiated the research on PCA-based SAR image denoising, derived the equations in the method, designed and conducted the experiments, wrote and revised the manuscript. The other coauthors actively participated in discussion of the results and reviewing of the manuscript.

Signatures of coauthors indicate they are in agreement with the statement above.

Alexander Wong___________________________ Jonathan Li_______________________

Junhuan Peng_____________________________ Yuanming Shu____________________
Abstract

The processing of remotely sensed data is innately an inverse problem where properties of spatial processes are inferred from the observations based on a generative model. Meaningful data inversion relies on well-defined generative models that capture key factors in the relationship between the underlying physical process and the measurements.

Unfortunately, as two mainstream data processing techniques, both mixture models and latent variables models (LVM) are inadequate in describing the complex relationship between the spatial process and the remote sensing data. Consequently, mixture models, such as K-Means, Gaussian Mixture Model (GMM), Linear Discriminant Analysis (LDA) and Quadratic Discriminant Analysis (QDA), characterize a class by statistics in the original space, ignoring the fact that a class can be better represented by discriminative signals in the hidden/latent feature space, while LVMs, such as Principal Component Analysis (PCA), Independent Component Analysis (ICA) and Sparse Representation (SR), seek representational signals in the whole image scene that involves multiple spatial processes, neglecting the fact that signal discovery for individual processes is more efficient.

Although the combined use of mixture model and LVMs is required for remote sensing data analysis, there is still a lack of systematic exploration on this important topic in remote sensing literature. Driven by the above considerations, this thesis therefore introduces a mixture of LVM (MLVM) framework for combining the mixture models and LVMs, under which three models are developed in order to address different aspects of remote sensing data processing: (1) a mixture of probabilistic SR (MPSR) is proposed for supervised classification of hyperspectral remote sensing imagery, considering that SR is an emerging and powerful technique for feature extraction and data representation; (2) a mixture model of K “Purified” means (K-P-Means) is proposed for addressing the spectral endmember estimation, which is a fundamental issue in remote sensing data analysis; (3) and a clustering-based PCA model is introduced for SAR image denoising. Under a unified optimization scheme, all models are solved via Expectation and Maximization (EM) algorithm, by iteratively estimating the two groups of parameters, i.e., the labels of pixels and the latent variables. Experiments on simulated data and real remote sensing data demonstrate the advantages of the proposed models in the respective applications.
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List of Symbols

\( x_i^k \) \hspace{1em} p \times 1 \text{ dimensional observation at site } i \text{ in class } k

\( A^k \) \hspace{1em} p \times m \text{ dimensional linear transformation matrix in class } k

\( s_i^k \) \hspace{1em} m \times 1 \text{ dimensional latent variable at site } i \text{ in class } k

\( n^k \) \hspace{1em} p \times 1 \text{ dimensional noise variable in class } k

\( a_j^k \) \hspace{1em} the jth column vector in \( A^k \)

\( \Lambda_k \) \hspace{1em} covariance matrix of \( n^k \)

\( m^k \) \hspace{1em} p \times 1 \text{ dimensional mean vector of class } k

\( l_i \) \hspace{1em} label of observation at site \( i \)

\( I \) \hspace{1em} unit matrix

\( K \) \hspace{1em} number of classes

\( n \) \hspace{1em} number of observations
# List of Abbreviations

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<tr>
<th>Abbreviation</th>
<th>Full Form</th>
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<tr>
<td>EM</td>
<td>Expectation and Maximization algorithm</td>
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<tr>
<td>GMM</td>
<td>Gaussian Mixture Model</td>
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<td>ICA</td>
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<td>K-P-Means</td>
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<td>MLE</td>
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<td>MLL</td>
<td>Multi-level Logistic prior</td>
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<td>MPPCA</td>
<td>Mixture of Probabilistic Principal Component Analyzer</td>
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<td>MPSR</td>
<td>Mixture of Probabilistic Sparse Representation</td>
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<td>MRF</td>
<td>Markov Random Field</td>
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<td>PCA</td>
<td>Principal Component Analysis</td>
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<td>QDA</td>
<td>Quadratic Discriminant Analysis</td>
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<td>SAR</td>
<td>Synthetic Aperture Radar</td>
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<td>SR</td>
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Chapter 1
Introduction

1.1 Background

Remote sensing is the science of acquiring information about earth surface from a distance, using sensors typically onboard aircrafts or satellites (Lillesand et al., 2008). Remote sensors can be either active or passive. Synthetic aperture radar (SAR), as a typical active sensor, is capable of illuminating earth surface by microwave and collecting the backscattered waves from earth surface (Oliver and Quegan, 1998; Mott, 2007; Wang, 2008). Due to its ability to work irrespective of weather conditions or sun-light illumination, SAR has been widely used in remote sensing applications. Passive sensors, such as multispectral or hyperspectral sensors, on the other hand, capture the natural electromagnetic radiation that is reflected or emitted by earth surface. Since they obtain full spectral information with narrow spectral bands, hyperspectral sensors are good at discriminating different materials, and have been used in various applications including mineralogy, defense and environmental measurements (Richards and Jia, 1999; Shaw and Manolakis, 2002; Liang, 2004; Ustin, 2004; Lillesand et al., 2008; Bioucas-Dias et al., 2013).

While the advancement in remote sensing platforms provides great opportunities for a broad range of disciplines, the large and ever-increasing data volume demands efficient data processing and analysis techniques. The remote sensing data are usually provided as digital raster images. Therefore, image processing techniques are required to address many different tasks, such as image denoising, classification and spectral unmixing (Camps-Valls et al., 2011).

Image denoising aims to remove the undesirable information that contaminates the image. Noise in remote sensing images could be caused by many factors, depending on how the image was created. In particular, SAR sensor, as a coherent system, inherently produces speckle noise, which has salt-and-pepper appearance, and greatly impedes SAR image interpretation (Xie, et al., 2002). Noise reduction therefore always serves as a preprocessing step to enhance image quality (Buades et al., 2005).

Remote sensing image classification intends to infer the label/identity information of image pixels based on the spectral or spatial measurements (Lu and Weng, 2007; Mountrakis et al., 2011; Mulder, et al., 2011; Bioucas-Dias et al., 2013; Camps-Valls et al., 2014). Both supervised and unsupervised
techniques can achieve this purpose. Before performing classification, supervised classifiers are firstly trained on training samples with known labels, in order to learn the relationship between observations and labels. Unsupervised classifiers, on the other hand, do not need to be trained, and cluster the observations based on their internal structures.

Spectral unmixing task aims to estimate for each pixel the fractional abundances of endmembers, which are the spectra of pure materials (Plaza et al., 2009; Camps-Valls et al., 2011; Bioucas et al., 2012; Bioucas-Dias et al., 2013). The endmembers are assumed to be the underlying factors, which are responsible for generating the spectral pixels in multispectral or hyperspectral images. The estimation of endmembers as well as their abundances is a fundamental issue for remote sensing image analysis.

Remote sensing image processing is essentially an inverse problem, in which the observations are used to infer the properties of underlying geospatial processes that contribute to data generation (Wang, 2010). Therefore, knowing the data generating mechanism is crucial for solving inverse problems. If the function describing the relationship between the measurements and the underlying quantities is provided, data inversion can be solved by inverting the function. Unfortunately, in remote sensing, a function of explicit and exact form is usually unknown.

In order to achieve meaningful data inversion, prior information concerning data generation has to be used as guidance and regulation. In practice, statistical generative models are usually employed to describe the relationship between underlying quantities and measured ones, considering that stochastic generative models allow explicitly modeling the hidden variables associated with underlying generative mechanism, while in the meantime accommodating the noise in observations and uncertainties in human knowledge.

Efficient remote sensing data processing therefore relies on well-defined generative models that capture key factors in the relationship between the underlying physical process and the observations.

1.2 Motivation and Objectives

In remote sensing, three factors concerning the relationship between the observations and underlying spatial processes are of fundamental importance.

(1) Multiple spatial processes, instead of single one, contribute to generation the remote sensing images, given the complexity of the ground target. Consequently, observed image pixels of different sources tend to assume different spectral or spatial patterns. For example, an urban image usually
involves multiple ground cover types, which admit different textural structures in spatial domain, and varying spectral patterns in spectral space. Such source heterogeneity phenomenon is also witnessed at sub-pixel level. For example, an image pixel always involves the spectral contributions of multiple materials, whose spectra are called endmembers.

(2) Informative signals lie in latent space, instead of the original spectral/spatial space, due to noise and other uncertainties in remote sensing system. The unobserved variables in latent space, also called latent variables, may provide informative representation of the remote sensing data. For example, textual patterns of ground targets, as linear or nonlinear arrangements of pixels values, may serve as signatures of different land cover types. In addition, the latent variables may offer explanations of the data generation mechanism. For example, the abundances of endmembers reveal the material composition of a mixed pixel. Moreover, the latent variables can also help to reduce the dimensionality of high-dimensional measurements, which are not rare in remote sensing.

(3) Different spatial processes tend to associate with different groups of latent variables, instead of by the same group. For example, different ground cover types tend to admit different spectral signatures in latent spectral domain, and assume varying types of texture patterns in latent spatial space.

Due to the co-occurrence of above three factors, efficient data analysis therefore relies on well-defined generative models that are capable of accounting for both source heterogeneity effect and hidden variable effect, as well as their relations. Unfortunately, as two mainstream data analysis techniques, mixture models and latent variables models (LVM) are inadequate in addressing these important issues.

On the one hand, mixture models, such as K-Means, Gaussian Mixture Model (GMM), Linear Discriminant Analysis (LDA) and Quadratic Discriminant Analysis (QDA), although being capable of accounting for the effects caused by different sources, fail to address the latent variable effects. Consequently, the learning of mixture components will be rendered inefficient, due to the failure in addressing their association with latent variables. For example, since GMM characterizes a class by Gaussian distribution in the original space, and ignores the fact that classes could be better represented by discriminative signals in the hidden/latent feature space, it is difficult for GMM models to strike a good balance between model bias and model variance.
On the other hand, LVMs, such as Principal Component Analysis (PCA), Independent Component Analysis (ICA) and Sparse Representation (SR), explain only the latent variable effects, but fail to account for the source heterogeneity issue. As a result, the learning of latent variables will be affected and disturbed by the existence of mixture effect, due to the failure to explicitly model such effect. For example, because PCA seeks representational signals in the whole image scene that involves a mixture of sources, and neglects the fact that signal discovery for individual sources is more efficient, in image denoising problems, global PCA learnt for all classes is less efficient than local PCAs learnt for individual classes. In order to avoid confusion, it is worthwhile to mention that LVM here refers to continuous latent variable models.

Driven by the above considerations, this thesis therefore intends to explore mixture of LVM (MLVM) that is capable of accounting for both mixture effects and latent variables, in order to achieve efficient remote sensing data processing techniques. Although some MLVM models, such as mixture of probabilistic PCA (MPPCA, Tipping and Bishop, 1999) and mixture of factor analyzer (MFA, Ghahramani and Hinton, 1996; Fokoue and Titterington, 2003) have been developed in the statistical literature, no efforts have been conducted towards a systematic investigation, in the context of remote sensing data processing. Four main research questions or gaps remain unaddressed, which motivate the studies conducted in this thesis.

1. There is still a lack of a general framework that is capable of providing principles and guidelines for building MLVMs that suit a variety of remote sensing data processing tasks.

2. MLVM has not been developed for SR, which is emerging and powerful technique for feature extraction and data representation.

3. Since the pixel values in remote sensing images are nonnegative, the latent variables are also required to be nonnegative in some cases, e.g. spectral unmixing. Therefore, new MLVMs have to be developed to address this particularity of remote sensing data.

4. The diversity of remote sensing data type and applications requires new MLVMs that support different remote sensing data processing tasks, e.g. denoising, classification, spectral unmixing.

1.3 Thesis Structure

This thesis proposes to study the modeling and analysis of remotely sensed imagery from a probabilistic generative perspective. Simultaneous modeling of both the underlying spatial processes and hidden signals is achieved by MLVMs, where mixture components distinguish between different
spatial processes, and latent dimensions account for hidden signals in each component. The contribution of this thesis lies in the following aspects:

Chapter 2 introduces a probabilistic framework, enabling a principled way of modeling and estimating both source heterogeneity effect and hidden signal effect, under which three MLVMs are developed, and successfully applied to a variety of remote sensing applications in terms of the image processing tasks and the sensor types.

Chapter 3 describes a novel mixture of probabilistic SR (MPSR) model, to be incorporated with Markov random field (MRF) for supervised classification of hyperspectral remote sensing imagery, considering that SR is an emerging and powerful technique for feature extraction and data representation.

Chapter 4 presents a novel mixture of K Purified means (K-P-Means) model, for spectral endmember estimation, which is a fundamental issue in remote sensing data processing.

Chapter 5 presents a clustering-based PCA algorithm in Chapter 5, for state-of-the-art SAR image denoising.

Finally, Chapter 6 concludes the thesis and suggests future research directions.
Chapter 2
Mixture of Latent Variable Models

This Chapter starts with an overview of the mixture model and LVM, followed by the introduction to the framework of MLVM, and the descriptions of three variants of MLVM.

2.1 Mixture Model

Since multiple spatial processes are responsible for remote sensing data generation, mixture models, which account for this source heterogeneity effect, are essential for pattern discovery and prediction (McLachlan and Peel, 2000). In mixture models, the \( p \times 1 \) dimensional observation at site \( i \) in class \( k \), denoted by \( x_i^k \), can be expressed as a linear combination of the mean vector of a class \( \mathbf{m}^k \), plus the class-dependent noise \( \mathbf{n}^k \):

\[
x_i^k = \mathbf{m}^k + \mathbf{n}^k \quad (i = 1, 2, ..., n; k = 1, 2, ..., K)
\]

(2.1)

Mixture models differ on noise distributions (McLachlan and Peel, 2000). In particular, the GMMs are widely used for the tasks of clustering and classification of remote sensing data (e.g. Ju et al., 2003; Clark et al., 2005; Amato et al., 2008; Thessler et al., 2008; Brenning, 2009; Pu and Landry, 2012; Chen et al., 2013), where \( \mathbf{n}^k \) is Gaussian noise with zero mean and covariance matrix \( \Lambda_k \).

Accordingly,

\[
p(x_i^k) = \frac{1}{(2\pi)^{p/2}|\Lambda_k|^{1/2}} \exp\left\{-\frac{1}{2}(x_i^k - \mathbf{m}^k)^T \Lambda_k^{-1}(x_i^k - \mathbf{m}^k)\right\}
\]

(2.2)

Based on Eqs. (2.1) and (2.2), GMM infers the membership of \( x_i \), by MLE or its variants, such as EM algorithm (Bailey and Elkan, 1994; McLachlan and Peel, 2000).

Popular clustering or classification methods are variants of model defined by Eqs. (2.1) and (2.2). For example, K-Means assumes \( \Lambda_1 = \Lambda_2 = \cdots = \Lambda_K = \mathbf{I} \) with \( \mathbf{I} \) being unit matrix; LDA assumes \( \Lambda_1 = \Lambda_2 = \cdots = \Lambda_K = \mathbf{D} \), with \( \mathbf{D} \) being diagonal matrix; QDA allows \( \Lambda_k \) being different for different classes.

The mixture models as formulated by Eq. (2.1), where a class is characterized by a certain parametric distribution in original feature space, assume some limitations.

- Characterizing a cluster/class using the mean vector \( \mathbf{m}^k \) and covariance matrix \( \Lambda^k \) is difficult to strike a good balance between model bias and model variance. For example, in
QDA, the number of unknown parameters in $\Lambda^k$ will grow quadratically with the increase of data dimensionality. Consequently, given high dimensional remote sensing data, mixtures models will easily be overfitted, leading to poor generalization capability. Methods with constrained covariance structure, such as LDA and K-Means, on the other hand, provide compromised model flexibility, leading to large model bias. In contrast, MLVMs are capable of characterizing a class by latent bases, which contain less number of unknown parameters, and providing great model flexibility in the meantime (Tipping and Bishop, 1999). Therefore, it is worthwhile to explore the use of MLVM for remote sensing data clustering and classification.

- Characterizing a cluster/class using a certain parametric probabilistic distribution in original domain is problematic when $x_i^k$ does not assume that distribution. In contrast, MLVMs offer flexibility by representing a class by several latent bases, which are free of explicit statistical distributions. Moreover, since Gaussian distribution only captures second-order variance, how to characterize high-order within-class variance is essential when the Gaussian assumption is validated (Camps-Valls et al., 2011). Fortunately, LVMs, such as the SR that represents a class by non-orthogonal bases, or ICA that represents a class by independent bases, are capable of capturing higher-order correlations. Therefore, it is desirable to explore MLVMs for clustering or classification, where inner class variation is characterized by various latent bases, instead of a parametric distribution in original domain.

- Since mixture models do not address the latent variable effect, they are unable to uncover the hidden signals that associated with the underlying and unobservable physical processes, nor can they provide a quantitative explanation of the data generation mechanism.

### 2.2 Latent Variable Model (LVM)

Since remote sensing observations are always of high-dimensionality, with noise and outliers, LVMs that seek low-dimensional, noiseless, and meaningful structures in transformed space are crucial for inverse problems in remote sensing. Typical LVMs, such as PCA, ICA, FA, SR and nonnegative matrix factorization (NMF), have been widely used in remote sensing data processing for various purposes, including dimension reduction, feature extraction, and signal discovery (Kondratyev and
Pokrovsky, 1979; Huete, 1986; Miao and Qi, 2007; Amato et al., 2008; Ozdogan, 2010; Chen et al., 2011; Viscarra Rossel and Chen, 2011; Frappart et al., 2011; Small, 2012; Li et al., 2012).

In order to reduce confusion, it is important to point out that the term LVM here refers continuous latent variables model (Bishop, 2006). In a probabilistic formulation of LVM, $x_i$, i.e. the $p \times 1$ dimensional observation at site $i$, is expressed as a linear transformation $A$ of $m \times 1$ dimensional unknown latent variables $s_i$ with additive noise $n$ (Bell and Sejnowski, 1995; Tipping and Bishop, 1999; Lewicki and Olshausen, 1999; Aharon et al., 2006).

$$x_i = As_i + n \quad (i = 1,2,\ldots,n) \quad (2.3)$$

As we can see, the general term $m^k$ in Eq. (2.1) is expressed more specifically by $As_i$. Therefore, comparing with Eq. (2.1) that considers the overall effect of a physical process, Eq. (2.3) probes into the sources of the physical process that contribute to the observations. Nevertheless, Eq. (2.3) does not involve the label information, therefore ignores the effect caused by different physical sources. There are two essential limitations about LVMs.

- LVMs are inefficient in addressing label-related tasks, e.g. clustering and classification. The main reason is probably because the columns in $A$ are indiscriminative to different sources. Therefore the label information of observation $x_i$ could not be inferred from the representational relationship between $A$ and $x_i$. Consequently, the key issue in adapting LVM for the clustering or classification is to explicitly learn different $A$ for different classes, as is conducted in MLVM.

- Except from low efficiency in label-learning tasks such as clustering and classification, the above-mentioned LVMs are inadequate in discovering informative signals for some other image processing tasks, such as denoising. It is mainly due to the difficulties in capturing nonlinear and local structures in feature space when signal discovery is performed on the whole dataset, which assumes enormous complexity due to the source heterogeneity effect. On the other hand, it has proved more efficient to learn representational signals for individual sources separately (e.g. Tipping and Bishop, 1999). Therefore, it is desirable to explore mixture of LVMs where a LVM is built upon one component of the mixture, instead of all components.
2.3 Mixture of LVMS (MLVM)

2.3.1 Model Formulation

Given the limitations of mixture models and LVMs, this thesis therefore focuses on MLVMs, in order that the mixture models and LVMs can be mutually complementary and beneficial. In MLVM, $x_i^k$, i.e. the $p \times 1$ dimensional observation variable in class $k$, is expressed as a class-dependent linear transformation $A^k$ of $m \times 1$ dimensional class-dependent unknown latent variables $s_i^k$ with additive noise $n$.

$$x_i^k = A^k s_i^k + n \quad (i = 1, 2, ..., n; k = 1, 2, ..., K) \tag{2.4}$$

Therefore, MLVM models and learns both label information \{l_i\}, with $l_i$ being class label of $x_i$, and latent model information, i.e. \{A^k\} and \{s_i^k\}, as opposed to mixture model that addresses only label information, and LVM that considers only latent model information.

The essence of MLVM is to model simultaneously two key factors in remote sensing data generation, i.e. multiple spatial processes and hidden signals, using the mixture components to discriminate different spatial processes, and LVM to account for hidden signals in each component.

In terms of latent variables learning, MLVM is capable of providing latent variables of strong representation power, due to its capability to capture local structures in feature space. Moreover, learning latent variables for individual sources separately, instead of for all sources simultaneously, may lead to latent variables, not only of strong representational power, but also of strong discriminative or explanatory power.

In terms of label learning, MLVM is supposed to be more capable of strike a good balance between model bias and model variance, considering both the model flexibility due to factors, such as the adaptability of latent bases and the capability of latent variables to capture higher-order inner-class correlation, and the model rigidity due to factors, such as the less number of parameters required to character a class and the constraint imposed on latent variables and latent bases.

Due to these advantages, MLVM benefits both signal-discovery-related tasks (e.g. data representation, compression, denoising and spectral source separation) and label-learning tasks (e.g. clustering, classification and). In statistical literature, some models, such as mixture of PCA (Tipping and Bishop, 1999) and mixture of factor analysis (MFA, Ghahramani and Hinton, 1996; Fokoue and Titterington, 2003) have been developed, and successfully used in a variety of applications (Frey et
Nevertheless, these techniques only constitute limited examples of MLVM. There is still a lack of a general MLVM framework, providing principles and guidelines for building task-dependent MLVMs. Moreover, no explicit MLVMs have been used or developed for addressing the particularities of remote sensing applications.

2.3.2 Optimization scheme

There unknown parameters in Eq. (2.4) can be represented by $\gamma = \{\{A_k\}, \{s_k^n\}, \theta\}$, where $\theta$ parameterizes noise distribution. Although the maximum likelihood estimation (MLE) is usually used for estimating parameters of generative models, it fails the task here due to the existence of unknown label variables $\{l_i\}$. Nevertheless, the Expectation and Maximization (EM) algorithm can be employed to approximate MLE by treating $\{l_i\}$ as unobservable or missing information. The EM algorithm is capable of estimating both $\gamma$ and $\{l_i\}$ iteratively by treating one of them being known (Bailey and Elkan, 1994; Dempster et al., 1977). Therefore, the EM solution is obtained by alternating the E- and M-steps:

1. Firstly, initialize parameters $\gamma$;

2. E-step: estimate $\{l_i\}$ based on $\gamma$. In a probabilistic context, $\{l_i\}$ can be estimated by maximizing a posterior (MAP) distribution of $l_i$ given $x_i$.

$$l_i = \arg\max_{l_i} \{p(l_i|x_i)\} \quad (2.5)$$

$$p(l_i|x_i) \propto p(x_i|l_i)p(l_i) \quad (2.6)$$

where $p(x_i|l_i)$ denotes the class-dependent likelihood of $x_i$, which allows the modeling of spectral information, and $p(l_i)$ is the prior probability of labels, which allows the modeling of spatial information.

3. M-step: update $\gamma$ based on $\{l_i\}$. In this step, the essence is to learn latent variables in each class separately, using the observations in the associated class. In a probabilistic approaches, e.g. the probabilistic PCA (Tipping and Bishop, 1999) and probabilistic SR (Lewicki and Olshausen, 1999), $\gamma$ is estimated by firstly integrating out the latent variable $s$, then maximizing the ML of $x$ with respect to $A$ and $\theta$, finally estimating $s$ by maximizing its posterior distribution. Without considering the statistical distributions, $\gamma$ can be obtained efficiently by some matrix
decomposition and machine learning techniques, e.g. singular values decomposition (SVD) for learning PCA parameters, and K-SVD technique for learning SR parameters (Aharon et al., 2006).

(4) Repeat E- and M-step until the parameters stabilize or a certain number of iterations have been reached.

The EM algorithm is famous for its capability of increasing the likelihood of observations in each iteration. Nevertheless, there is no guarantee that it will converge to the global maximum of the likelihood function (Wu, 1983). In practice, considering the sensitivity to the initial values, EM algorithm can be performed multiple times using different initial values, in order to increase the chance of finding the optimum solution.

2.3.3 Model Specifications and Variations
Since the framework defined in Sections 2.3.1 and 2.3.2 is very flexible, model assumptions and optimization scheme can be further specified, in order to account for the particularities of different applications. Since different combinations of the specifications may lead to different variants of MLVM, principles and guidelines can therefore be provided for building task-dependent models. In chapter 2.4, three models are developed by adopting different model constraints and regulations.

2.3.3.1 Assumptions on $A^k$ and $s^k$
Different assumptions on $A^k$ and $s^k$ lead to different LVMs. The columns in $A^k$ define the projection directions that are capable of revealing “interesting” patterns. In a probability framework, $A^k$ is always assumed non-random, and the varying “interestingness” of $A^k$ is defined by different prior distributions of $s^k$. For example, to achieve uncorrelated projection directions, PCA assumes $s$ being Gaussian distributed with zero mean and identity covariance matrix (Tipping and Bishop, 1999); ICA achieves independent directions by assuming $s$ being super-Gaussian or sub-Gaussian distributed (Bell and Sejnowski, 1995), and SR obtains sparse signal by assuming $s$ admitting Laplacian or Cauchy distribution (Lewicki and Olshausen, 1999).

The number of columns in $A^k$ can be arbitrary. It can be bigger than the dimensionality of observations, e.g. in SR, or be equal to dimensionality of observations, e.g. in ICA, or be equal to the number of classes, e.g. in the proposed K-P-Means model. Generally speaking, larger number of latent bases enables better representation of inner-class variation, but in the meantime, increase model complexity.
Since the remote sensing spectral values are nonnegative, in order to achieve meaningful interpretation, the values of elements in $A^k$ and $s^k$ are required to be nonnegative in some circumstances, e.g. when learning spectral endmembers for spectral source separation.

Sometimes, it is not necessary to explicitly impose label constraint $A$ and $s$. Nevertheless, at least one of them has to be discriminative to different classes, in order that the other one can be class-dependent as well. For example, in the proposed K-P-Means model, although latent bases in $A$ are not explicitly labeled, their association with different classes are achieved by imposing class-discriminative constraints on $s$.

2.3.3.2 Assumptions on $n$

Different assumptions on $n$ lead to different mixture models. Although $n$ is normally assumed to follow a Gaussian distribution, it sometimes is assigned to other distributions in order to address the particularities of remote sensing dataset, e.g. $n$ follows Gamma distribution in the proposed clustering-based PCA model to accommodate the distinct statistical properties of SAR speckle noise.

Whether noise $n$ of different mixture components should follow the same distribution, depends on the capability of LVMs in representing class-discriminative information. While $n$ in Eq. (2.4) is assumed being the same for different classes, class-dependent noise, symbolized by $n^k$, will be used instead of $n$, in order to allow different noise distributions for different classes, if the class-dependent information cannot be totally explained by $A^k s^k$.

The complexity of the covariance matrix of $n$ depends on the representational capability of LVMs in capturing the correlation among multivariate variables. The covariance matrix of $n$ will be a full matrix, if the correlation effect among variables cannot be fully captured by $A^k s^k$. The covariance matrix of $n$ will be a diagonal matrix, if the correlation effect among variables can be effectively captured by $A^k s^k$. Moreover, the covariance matrix of $n$ will be isotropic matrix (whose off-diagonal elements are zeros, and diagonal elements have equal values), if variance heterogeneity effect among variables can be captured by $A^k s^k$.

The existence of $n$ allows the modeling of stochastic nature of remote sensing observations or the uncertainties in human prior knowledge concerning the data generating mechanism. However, if $n = 0$, then the model defined by Eq. (2.4) amounts to a deterministic model, which is impractical for remote sensing data modeling due to significant uncertainties in remote sensing system. Therefore,
even using a deterministic model, the noise in latent space still need to be estimated and separated in most applications, e.g. denosing, dimension reduction and feature extraction.

2.3.3.3 Classification vs. Clustering

For label learning tasks that aim to learn class labels of remote sensing observations, classification and clustering can be distinguished, based on whether $\gamma$ is known.

In classification, since $\gamma$ has been learnt from training samples, M-step in EM iteration can be avoided, and the estimation of labels $\{l_i\}$ requires performing E-step only once.

In clustering, however, the learning of $\{l_i\}$ has to be achieved iteratively by alternating the E- and M-steps until convergence.

2.3.3.4 Supervised vs. Unsupervised Latent Variable Learning

For latent variable learning tasks that intend to learn latent bases and latent variables, depending on whether $\{l_i\}$ are known, the tasks can be categorized into supervised and unsupervised ones.

In a supervised case, since the labels of observations $\{l_i\}$ are known, E-step can be avoided and latent variable learning can be achieved by performing M-step only once. In this case, the MLVM will degrade into $K$ LVMs, where $K$ denotes the number of classes. In unsupervised case, the learning of latent variables has to be performed iteratively by alternating the E- and M-steps until convergence.

2.3.3.5 Label Prior

In Eq. (2.6) the label prior $p(l_i)$ is used to model the spatial correlation effect among labels. In remote sensing observations, spatially-close pixels tend to be caused by the same spatial process. Therefore, they tend to admit the same label. The Markov random field (MRF) is a popular technique for modeling the spatial correlation effect in labels. It assumes that two pixels are correlated if only they are neighbors in spatial domain. If the label prior is adopted, in E-step, the estimation of labels requires solving a MAP problem, i.e. $l_i = \text{argmax}_{l_i} p(l_i | x_i)$. Otherwise, it degrades to a ML problem, i.e. $l_i = \text{argmax}_{l_i} p(x_i | l_i)$. 
2.4 Models Developed under MLVM Framework

Based on the framework defined by Eq. (2.4), three MLVMs are achieved by adopting different constraints and model specifications, in order to address different aspects of remote sensing data analysis.

2.4.1 Mixture of Probabilistic Sparse Representation (Abbreviation MPSR)

A mixture of probabilistic SR (MPSR) is proposed in Chapter 3 for supervised hyperspectral classification, considering the gap that while SR is an emerging and powerful technique for hyperspectral image representation, there is still a lack of a mixture of probabilistic approach for it. This Section starts with the model definition and optimization, followed by the discussion of the model characteristics.

2.4.1.1 Model Definition and Optimization

The generative model of MPSR is similar to Eq. (2.4), except that $A^k$ ($k = 1, 2, ..., K$) is assumed being known, and that $x_i^k$ is assumed being sparsely representable by only a few columns (also called atoms) in $A^k$. Accordingly, the class conditional distribution of $x_i^k$ is expressed as:

$$p(x_i^k) = \frac{1}{(2\pi)^{p/2}|\Lambda|^{1/2}} \exp\left(-\frac{1}{2} (x_i - A^k s_i^k)^T \Lambda^{-1} (x_i - A^k s_i^k)\right)$$

$$\Lambda = \begin{bmatrix}
\sigma_1 & 0 & 0 \\
0 & \ddots & 0 \\
0 & 0 & \sigma_p
\end{bmatrix}$$

(2.7)
(2.8)

Therefore, the unknown parameters include $y = \{s_i^k, A\}$ and $\{l_i\}$. Following the optimization scheme in Section 2.3.2, this model can be solved by EM algorithm which alternates two main steps: E-step: estimating $\{l_i\}$ given $y$, and M-step: updating $y$ given $\{l_i\}$. In order to address the spatial correlation effect, the E-step solves a MAP problem, where the label prior is modeled by MRF.

2.4.1.2 Model Characteristics

The benefits of MPSR can be summarized into the following aspects:

- Instead of characterizing the within-class variation by a covariance matrix in Eq. (2.2), MPSR captures the variation by the variability of bases in $A^k$. Note that the number of columns in $A^k$ (i.e. $m$) is allowed to be bigger than the dimensionality of spectral vector (i.e. $p$), and that the latent bases in $A^k$ are allowed to assume arbitrary distributions and
correlations. Due to these factors, $A^k$ can even be implemented by substituting its columns for training samples in class $k$, in a nonparametric manner. Therefore, $A^k$ provide flexibility and adaptability in capturing complex inner-class data structure, as opposed to the covariance matrix approach that is limited to explaining second-order correlation.

- Because of the great representational capability of $A^k s_i^k$, it is reasonable to assume that the noise $n$ is class-independent and admits a diagonal covariance matrix. Therefore, the number of parameters in the distribution of $n$ is greatly reduced, thus the risk of overfitting.

- In an unsupervised scenario, considering that learning latent bases (i.e. dictionary) for each class in MPSR, is more capable of capturing the complex data structure than learning latent bases for the whole dataset consisting of multiple classes, it is worthwhile to mention that assuming \{A$^k$\} to be unknown variables and learning \{A$^k$\} in MPSR may increase the representational capability of SR-based approaches for low-level tasks, such as image denoising and compression.

### 2.4.2 K-P-Means Model

The K-P-Means approach is proposed in Chapter 4, for spectral endmember estimation, which is a fundamental issue in remote sensing data processing. It is proved in this thesis that the combination of latent model and mixture model, as conducted in K-P-Means algorithm, is capable of providing a new route for spectral unmixing. This Section starts with the model definition of K-P-Means and the optimization method, followed by the discussion of the model characteristics.

#### 2.4.2.1 Model Definition and Optimization

The generative model of K-P-Means is the same to Eq. (2.4), except that $A^k (k=1,2,\ldots,K) = A$, and the label constraint on $A$ is achieved by imposing constraints on $s$, i.e., the elements in $s$ should be nonnegative, and in the $k$th class, the $k$th element should be bigger than the rest. Accordingly, the model can be formulated as:

$$x_i^k = \sum_{j=1}^{K} s_{ij} a_j + n, \text{ where } s_{ik} > \{s_{ij\neq k}\} \geq 0 \quad (2.9)$$

where $n$ is independently and identically (i.i.d.) white noise. Therefore, comparing with MPSR that imposes the sparsity constraint on $s$, K-P-Means imposes the constraint of $s_{ik} > \{s_{ij\neq k}\} \geq 0$ on $s$. Accordingly, Eq. (2.9) can be reformulated as:
\[
\mathbf{y}^k_i = (x^k_i - \sum_{j\neq k}^K s_{ij} a_j)/s_{ik} = a_k + \mathbf{n}, \text{where } s_{ik} > \{s_{ij \neq k}\} \geq 0
\] (2.10)

where \(\mathbf{y}^k_i\) is called the “purified” pixels, because it removes the contribution of less significant atoms/endmembers \(\{a_{j \neq k}\}\) associated with smaller coefficients \(\{s_{ij \neq k}\}\).

Following the optimization scheme in Section 2.3.2, the unknown parameters in K-P-Means, which include \(\mathbf{y} = \{\{s_{ij}\}, \{a_j\}\}\) and \(\{l_i\}\), are estimated by EM algorithm, which treats \(\{l_i\}\) as missing observations, and repeats the two steps until convergence: estimating labels \(\{l_i\}\) given \(\mathbf{y}\), and updating \(\mathbf{y}\) based on label information.

K-P-Means is designed for addressing a linear spectral unmixing problem, where a spectral pixel \(x_i\) can be expressed as a linear combination of spectral endmembers \(\{a_j\}\). The essence of K-P-Means is to separate the individual contributions of endmembers, and label a pixel by identifying the endmember that dominates this pixel. While K-P-Means are used here for spectral unmixing, it may be applicable to other clustering or signal discovery problems where the observations are a nonnegative linear combination of nonnegative signals.

2.4.2.2 Model Characteristics

The benefits of K-P-Means can be summarized into the following aspects:

- Comparing with GMM, the general term \(\mathbf{m}^k\) defined by Eq. (2.1) is expressed more specifically by \(\sum_{j=1}^K s_{ij} a_j\) in Eq. (2.9). Accordingly, as opposed to GMM, or mixture model in general, that consider the overall effect of a physical process, K-P-Means probes into the sources of the physical process that contribute to the observations. This property of K-P-Means allows it to separate the independent contribution of spectral endmembers (defined as the spectra of “pure” materials) in mixed pixels.

- Moreover, since K-P-Means characterizes a class by a number of \(K\) latent bases \(\{a_k\}\) which are more capable of capturing inner-class variance than single mean vector \(\mathbf{m}^k\) in Eq. (2.1), it is reasonable to assume that \(\mathbf{n}\) in K-P-Means admits less-complex covariance structure than in GMM. In the scenario where GMM characterizes a class by the mean vector and a full covariance matrix, and where K-P-Means characterizes a class by \(K\) latent bases \(\{a_k\}\) and an isotropic variance matrix of \(\mathbf{n}\), GMM will require \(K(p + p(p + 1)/2)\) parameters for characterizing all classes, while K-P-Means require only \(K(p + 1)\)
parameters for characterizing all classes. Since K-P-Means is capable of providing a parsimonious parameterization of clusters, it is less prone to overfitting. Moreover, there are no restrict assumptions on the distribution and number of \( \{a^k\} \), which gives the K-P-Means some flexibility to characterize the data variance.

- Comparing with LVMs defined by Eq. (2.3), where the mixed pixels \( \{x_i\} \), regardless of their label information, are used for learning latent bases \( A \), K-P-Means accounts for the label information by separating the individual contributions of different endmembers, and learns latent bases \( \{a_k\} \) based on the associated “purified” pixels \( \{y^k\} \). Therefore, by considering the label information, K-P-Means constitutes a powerful nonnegative matrix factorization technique.

### 2.4.3 Clustering-based Principal Component Analysis

The Clustering-based PCA model is proposed in Chapter 5, for addressing the SAR image denoising problem, which is fundamental for SAR image processing and interpretation. It is proved in this thesis the state-of-the-art SAR image denoising techniques can be achieved by performing PCA-based denoising for individual clusters, as conducted in clustering-based PCA. This Section starts with the model definition of clustering-based PCA and the optimization method, followed by the discussion of the characteristics of this model.

#### 2.4.3.1 Model Definition and Optimization

The generative model of clustering-based PCA is the same to Eq. (2.4), except that \( \{A^k\} \) are PCA bases, and that \( n \) is additive signal dependent noise (ASDN) that assumes zero-mean i.i.d. Gamma distribution.

\[
x^k_i = A^k \hat{s}^k_{y_i} + n
\]

where \( \hat{s}^k_{y_i} \) represents the noise-free latent variables, which is estimated by LMMSE in PCA domain. The task of denoising is achieved by estimating \( \hat{s}^k_{y_i} \) and reconstructing SAR image using \( y^k_i = A^k \hat{s}^k_{y_i} \).

Following the optimization scheme in Section 2.3.2, the unknown parameters in clustering-based PCA, which include \( \gamma = \{\{\hat{s}^k_{y_i}\},\{A^k\}\} \) and \( \{l_i\} \), are estimated by EM algorithm, which assumes the labels \( \{l_i\} \) as missing observation and repeats the two steps: E-step: estimating \( \{l_i\} \) given \( \gamma \), and M-step: updating \( \gamma \) given the label information. In E-step, label learning is achieved by performing
clustering in PCA domain. To reduce dimensionality and resist the influence of noise, several leading principal components (PCs), identified by the Minimum Description Length (MDL) criterion are used to feed the K-means clustering algorithm. In M-step, \( \{A^k\} \), after being learnt for different classes, are used to estimate \( \{s^k_{yi}\} \) via a LMMSE approach, in order to reconstruct the clean SAR image.

2.4.3.2 Model Characteristics

Clustering-based PCA algorithm assumes the following characteristics:

- Clustering-based PCA can be treated as an adaptation of MPPCA (Tipping and Bishop, 1999) for addressing the SAR image denoising problem. It assumes the main advantages of MPPCA model, i.e. learning PCA for individual classes is more efficient than learning PCA simultaneously for all classes. Nevertheless, it differs from MPPCA in terms of the implementations of EM steps, in order to fit into the SAR image denoising scenario.

- Although it is general practice to perform image denoising in latent space, it is not until recent years that it is recognized that image denoising is more efficient when latent models are learnt for individual classes. The effectiveness of denoising in latent domain depends highly on whether the latent variables can sparsely represent the scene signal. And the sparsity can be achieved by performing analysis on observations in the same class, which assume similar spectral or spatial patterns.

2.5 Chapter Summary

In this Chapter, a framework of MLVM was introduced, from a comparative perspective with the mixture model and LVM. Three variants of MLVM were described in terms of model assumptions and optimization scheme. The characteristics and advantages of these models relative to LVM and mixture model were discussed. It was demonstrated theoretically that the proposed MLVM models (i.e. MPSR, K-P- Means and clustering-based PCA) assume theoretical advantages over either LVM or mixture model. In the following Chapters 3, 4 and 5, the proposed models will be introduced in detail.
Chapter 3
MPSR for Bayesian Classification of Hyperspectral Imagery

This chapter presents a Bayesian method for hyperspectral image classification based on Sparse Representation (SR) of spectral information and Markov Random Filed (MRF) modeling of spatial information. We introduce a mixture of probabilistic SR (MPSR) approach to estimate the class conditional distribution, which proven to be a powerful feature extraction technique to be combined with labels prior distribution in a Bayesian framework. The resulting Maximum a Priori (MAP) problem is estimated by a graph cut $\alpha$-expansion technique. The capabilities of the proposed method are proven in several benchmark hyperspectral images of both agricultural and urban areas. © [2014] IEEE. Reprinted, with permission, from [Xu Linlin, and Li J., Bayesian classification of hyperspectral imagery based on probabilistic sparse representation and Markov random field, IEEE Geoscience and Remote Sensing Letters, 04/2014].

3.1 Introduction

The classification of hyperspectral remotely sensed imagery constitutes a challenging data-mining and machine learning problem due to not only the high dimensionality of various spectral bands, but also the ambiguity in spectral signatures of different classes caused by the existence of mixed pixels (Li et al., 2012). In light of these difficulties, one essential issue is how to extract the most compact and discriminative features from the high dimensional hyperspectral bands. Among many recent studies (Camps-Valls et al., 2010; Chen et al., 2011; Li et al., 2012; Chen et al., 2013; Xia et al., 2013), the Sparse Representation (SR) approach has proven to be an extremely powerful tool for hyperspectral image classification (Chen et al., 2011; Chen et al., 2013). It assumes that the high dimensional spectral vector can be sparsely represented by a few atoms in a dictionary consisting of training samples. Therefore, forcing sparsity, the training samples in all classes will compete for their involvement in representing the spectral vector. The most relevant class will eventually win large shares, resulting in small representational residual, while the wrong or less-relevant classes will have no or little involvement, leading to high representational residual. Therefore the label of a pixel can
be determined by selecting the minimum residuals among all classes. While this approach has proven its capability in revealing the most discriminative information hidden in high dimensional spectral vector, there is still a lack of probabilistic mixture approach which provides the probability features rather than residuals. A probabilistic mixture approach is especially important considering the facts that integrating contexture/spatial information is an essential issue for hyperspectral image classification (Camps-Valls et al., 2010; Chen et al., 2011; Li et al., 2012; Chen et al., 2013), and employing Markov Random Fields (MRF) method, a classic and powerful method for modeling spatial information, requires conditional probability in a Bayesian framework (Geman & Geman, 1984; Li, 2001; Deng & Clausi, 2005; Li et al., 2012).

In this chapter, we proposed a mixture of probabilistic SR (MPSR) approach to be integrated with MRF technique in Bayesian framework. Instead of using a unified dictionary consisting training samples from all classes, we design one dictionary for each class. And we therefore derive a conditional probability for spectral vector by sparsely representing it over the class-dependent dictionaries. While this probabilistic formulation of SR is used with MRF for hyperspectral data classification, it may also help other statistical methods in other applications. The rest of the chapter is organized as follows. Section 3.2 discusses the proposed MPSR method and its integration with MRF technique. In Section 3.3, experiments are designed to examine the performance of the proposed method. Section 3.4 concludes this study.

3.2 Proposed Approach

3.2.1 Problem Formulation

In this chapter, we denote the discrete lattice spanned by hyperspectral imagery by \( T \), and a site in the lattice by \( i \in T \). We represent the observation at site \( i \) by \( x_i \), a \( p \)-dimensional random vector taking on values of various spectral bands, and the label of site \( i \) by \( l_i \), a random variable taking on a class \( \{1, \ldots, K\} \). Then a hyperspectral image can be denoted as \( X = \{x_i | i \in T\} \), and the labels of this image as \( l = \{l_i | i \in T\} \). In the classification problem, we are trying to infer \( l \) based on \( X \), which in the Bayesian framework, can be achieved by maximizing the posterior distribution of \( l \) given \( X \),

\[
p(l|X) \propto p(X|l)p(l)
\]  

(3.1)
where \( p(x|l) \) denotes the probability distribution of spectral vector \( x \) conditioned on \( l \), which allows the modeling of spectral information; \( p(l) \) is the priori probability of labels, which allows the modeling of spatial information.

In this chapter, \( p(x|l) \) is approached by a novel MPSR approach to mine the most discriminative information hidden in spectral bands, while \( p(l) \) is implemented by the MRF-based Multi-level Logistic (MLL) prior to constrain regional smoothness. The MAP problem is solved by the graph cut \( \alpha \)-expansion algorithm.

### 3.2.2 Mixture of Probabilistic Sparse Representation

In this chapter, we assume that a spectral vector in a class can be sparse represented by the training samples in the same class. Therefore, as opposed to classic SR approach that adopts a unified dictionary for all classes (Chen et al., 2011; Li et al., 2012), we adopt separate dictionaries for different classes. We express the observed signal variable at site \( i \) that belongs to class \( k \) as:

\[
x_i^k = A_k s_i^k + n
\] (3.2)

where \( A_k = \{a_1^k, a_2^k, ..., a_{M_k}^k\} \) is the dictionary consisting of training samples in class \( k \); \( s_i^k \) is the sparse vector corresponding to class \( k \) whose non-zero elements define which columns in \( A_k \) will be used; and \( n \) is the class-independent zero-mean Gaussian noise with diagonal covariance matrix \( \Lambda \). Although it’s reasonable to assume different \( n \) for different classes, it would increase the number of unknown parameters, consequently the risk of overfitting. In our formulation, we assume that \( A_k s_i^k \) is capable of capturing the discriminative information in \( x_i^k \), thus the random noise \( n \) is class-independent. We treat \( A_k s_i^k \) as fixed effect; hence the class conditional likelihood of spectral vector \( x_i \) can be expressed as:

\[
p(x_i^k) = \frac{1}{(2\pi)^{n/2}|\Lambda|^{1/2}} \exp\left\{ -\frac{1}{2} (x_i - A_k s_i^k)^T \Lambda^{-1} (x_i - A_k s_i^k) \right\}
\] (3.3)

\[
\Lambda = \begin{bmatrix}
\sigma_l & 0 & 0 \\
0 & \ddots & 0 \\
0 & 0 & \sigma_p
\end{bmatrix}
\] (3.4)

The matrix \( A_k \) can be implemented as a dictionary storing training samples in class \( k \). Given the dictionary \( A_k \) the unknown sparse vectors \( s_i^k \) can be estimated by solving the following optimization problem.

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\[ \hat{s}_i^k = \arg \min ||A^k s_i^k - x_i||_2 \text{ subject to } ||s_i^k||_0 \leq \tau \] (3.5)

The \(l_0\) norm \(||.||_0\) will simply count the nonzero items in \(s_i^k\). So the optimal \(\hat{s}_i^k\) is estimated by minimizing the representation error with constraint on sparsity level. This NP-hard optimization problem can be solved by some greedy pursuit algorithms, such as Orthogonal Matching Pursuit (OMP) or Subspace Pursuit (SP). Interested readers are referred to Tropp & Gilbert (2007) and Dai & Milenkovic (2009) for further information. The estimation of the second unknown parameter \(\Lambda\) relies on the label information. This issue can be solved by Expectation Maximization (EM) algorithm by treating the label \(l\) as missing information (Deng & Clausi, 2005). Therefore \(\Lambda\) is estimated from representation residuals in an iterative manner (see Algorithm 1).

This MPSR leads naturally to a discriminative model. Assuming the labels of different sites are independent, according to the Bayes rule, the posterior probability of \(l_i\):

\[ p(l_i|x_i) \propto p(x_i|l_i)p(l_i) \] (3.6)

Assuming the classes are equally likely, then \(p(l_i|x_i) \propto p(x_i|l_i)\). Therefore, according to the MAP criterion, we can estimate \(l_i\) by maximizing \(p(x_i|l_i)\) over different classes. We refer to our classifier as MPSR, whose detailed implementation is summarized in Algorithm 1.

**Algorithm 1: MPSR**

**Input:** training dictionaries for all classes \(\{A^1, ..., A^K\}\), data matrix \(X=\{x_i| i \in T\}\)

**Output:** class labels \(I = \{l_i| i \in T\}\)

**Initialization:** \(\tilde{\Lambda}^{(0)} = I; t = 1; s_i^k = OMP(A^k, x_i, \tau)\) for \(k = 1, 2, ..., K\) and \(i \in T\)

while \(t \leq \text{iters or sum(diag}[\tilde{\Lambda}^{(t)} - \tilde{\Lambda}^{(t-1)}]) > s\) do

\(\tilde{l}_i = \arg \min_{l_i} \{-\log(p(x_i|l_i))\}\)

\(\tilde{\Lambda}^{(t)} = \text{var}([x_i - A^k \hat{s}_i^k]|i \in \text{test set})\)

end while

3.2.3 MRF-Based MLL Prior

Although MPSR itself constitutes a classifier, it ignores the contextual information which is of great importance for hyperspectral data classification. We therefore further incorporate the spatial information by using the MRF-based MLL prior. The MRF is a classical method for modeling
contextual information (Geman & Geman, 1984). It promotes identical class label for spatially close pixels. The MRF-based approach is often implemented by the MLL model, which can be expressed as (Li, 2001):

$$p(l) = \frac{1}{Z} \exp \left( -\sum_{i \in T} \sum_{u \in N_i} \delta(l_i, l_u) \right)$$  \hspace{1cm} (3.7)

where $N_i$ denotes the neighborhood centered at site $i$; and $\delta(l_i, l_u) = -1$ if $l_i = l_u$, while $\delta(l_i, l_u) = 1$ if $l_i \neq l_u$.

### 3.2.4 Complete Algorithm

The MPSR and MLL in Section 3.2.2 and 3.2.3 are incorporated into a Bayesian framework and solved by the MAP criterion. The optimal labeling $\hat{l}$ can be obtained according to MAP criterion:

$$\hat{l} = \text{arg min}_l \left\{ \sum_{i \in T} \left[ -\log p(x_i | l_i, \Lambda, s_i^l) - \gamma \sum_{u \in N_i} \delta(l_i, l_u) \right] \right\}$$  \hspace{1cm} (3.8)

where $\gamma$ is the weighting parameter that determines the relative contribution of the two components.

This combinational optimization problem of estimating $y$ given $\Lambda$ and $s_i^l$ is solved in this chapter by the graph-cut-based $\alpha$-expansion algorithm which proved being capable of providing efficient and effective approximation to the MAP segmentation in computer vision (Boykov et al., 2001; Bagon, 2006). We refer to the complete algorithm in this Section as MPSRMLL, whose detailed implementation is summarized in Algorithm 2. The time complexity of MPSRMLL is largely determined by the complexity of OMP algorithm: $O(\tau p M)$ with $M$ being the number of atoms in dictionary, and the complexity of the $\alpha$-expansion algorithm: $O(T)$ with $T$ being the number of pixels.

---

**Algorithm 2: MPSRMLL**

| Input: training dictionaries for all classes $\{A^1, ..., A^K\}$, data matrix $X=\{x_i| i \in T\}$ |
| Output: class labels $l = \{l_i| i \in T\}$ |
| Initialization: $\tilde{\Lambda}^{(0)} = I$; $t = 1$; $s_i^k = OMP(A^k, x_i, \tau)$ for $k = 1, 2, ..., K$ and $i \in T$ |
| while $t \leq \text{iters}$ or $\text{sum(diag} | \tilde{\Lambda}^{(t)} - \tilde{\Lambda}^{(t-1)} |) > s$ do |
| \hspace{1cm} $P = \{p(x_i| l_i = k)\} \text{ for } k = 1, 2, ..., K \text{ and } i \in T$ |
| \hspace{1cm} $\tilde{l}_i = \alpha - \text{expansion}(P, \gamma)$ |
| \hspace{1cm} $\tilde{\Lambda}^{(t)} = \text{var}(\{x_i - \tilde{\Lambda}_i s_i^l | i \in \text{test set}\})$ |
| end while |

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3.3 Experiments

We adopt three benchmark hyperspectral images: AVIRIS Indian Pines, University of Pavia and the Center of Pavia (referred to Hyperspectral Remote Sensing Scenes (2013) for detailed information) to test the proposed algorithms. The first image was captured by Airborne Visible/Infrared Imaging Spectrometer (AVIRIS) over a vegetation area in Northwestern Indiana, USA with spatial resolution of 20m, consisting of 145 × 145 pixels of 16 classes and 200 spectral reflectance bands after removing 20 water absorption bands (104–108, 150–163, and 220). The other two hyperspectral images are urban images acquired by the Reflective Optics System Imaging Spectrometer (ROSIS) with spatial resolution of 1.3m, consisting of 103 spectral bands after removing 12 noisy bands. The Pavia University scene is centered at the University of Pavia, consisting of 610×340 pixels, while the Pavia Center scene is at the center of the Pavia city, consisting of 1096×492 pixels. Both images have 9 ground-truth classes.

3.3.1 Design of Experiments

We implemented Algorithms 1 and 2 in Sections 3.2.2 and 3.2.4, which are referred as MPSR2 and MPSR2MLL. To examine the influence of $\Lambda$, we forced $\hat{\Lambda}$ in MPSR2 and MPSR2MLL to be unit matrix. And the resulting algorithms are referred to as MPSR1 and MPSR1MLL, respectively. We experimentally set $\text{iter} = 20$ and $s = 0.1$ for MPSR2 and MPSR2MLL, and $\gamma = 20$, $\tau = 5$ for all proposed algorithms. In Section 3.2.4, we explored the sensitivity of these parameters. We also implemented the OMP algorithm in Chen et al. (2011), and adopted the residuals in OMP as data cost to feed $\alpha$-expansion algorithm (referred to as OMPMLL). Moreover, since the MLRsubMLL approach in Li et al. (2012) is also MRF-based approach, we included this algorithm along with the MLRsub for comparison study. The smooth cost in MLRsubMLL was set to be 2 for optimal performance, while all other parameters followed Li et al. (2012).

For the labeled pixels in these datasets, we randomly select a certain number of pixels from each class as training samples, while the rest labeled pixels are used as test set. For Indian Pines dataset, training samples in each class constitute 10% of the total samples in that class. For the other two datasets, we adopt a popular approach, and the number of training samples in each class is the same as that in Chen et al. (2011). For further details the reader is referred to Chen et al. (2011).

To be consistent with the other researchers, we adopt three numerical measures, overall accuracy (OA), average accuracy (AA), and the $\kappa$ coefficient for evaluation purpose (Bagon, 2006). To account...
for the possible bias produced by random sampling, each experiment is performed 10 times on different sampling results. The numerical values in Table 3.1 are the average of the 10 realizations. But the maps in Figure 3.1 are from one realization.

3.3.2 Numerical Comparison

Table 3.1 provides the statistics of different algorithms on three benchmark dataset. Overall, MPSR2MLL greatly outperformed the other approaches on most datasets, achieving OA of 97.8%, 99.1% and 99.4% respectively.

Comparing with MPSR1 and MPSR2, the OAs of MPSR1MLL and MPSR2MLL increased on average 25%, 21%, and 6% on respectively the three datasets, indicating the importance and benefit of integrating SR-based classifier with MRF to utilize both spectral and spatial information for hyperspectral image classification. MLRsubMLL also increased significantly the performance of MLRsub. However, nearly no performance increase of OMPMLL over OMP was observed. It is mostly because OMP is hard-classifier which produces residual features rather than probability features.

Comparing with MPSR1 and MPSR1MLL, MPSR2 and MPSR2MLL achieved higher OA on Indian Pines, slightly higher values on Pavia U, and comparable values on Pavia C. These results justify the idea of accounting for the variance heterogeneity across different spectral bands. Moreover, they may also indicate that addressing variance inhomogeneity is more beneficial when the quality of training samples is low, considering that the Indian Pines dataset, on which the MPSR2 and MPSR2MLL achieve higher performance-increase than on the other two datasets, assumes higher dimensionality due to more spectral bands, heavier mixed pixel effect caused by lower spatial resolution, and smaller number of training samples in most classes than Pavia U and especially Pavia C.

It is desirable to compare MPSR2MLL and MLRsubMLL, since both approaches are MRF-based generative models for MAP classification. MPSR2 slightly outperformed MLRsub on Indian Pines and Pavia U, while MLRsub achieves better results on Pavia C. Nevertheless, the adoption of MLL prior enabled MPSR2MLL to achieve higher OA and \( \kappa \) values on all datasets.
Table 3.1: Overall accuracy, average accuracy, and κ statistic obtained by different methods

The best results are highlighted in bold typeface

<table>
<thead>
<tr>
<th>Classifies</th>
<th>Indian Pines</th>
<th></th>
<th></th>
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<td>AA(%)</td>
<td>κ</td>
<td>OA(%)</td>
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<td>κ</td>
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<td>80.4</td>
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<td>0.738</td>
<td>96.2</td>
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<tr>
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<td>64.7</td>
<td>0.632</td>
<td>80.4</td>
<td>83.1</td>
<td>0.738</td>
<td>96.3</td>
</tr>
<tr>
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<td>68.5</td>
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<td>76.2</td>
<td>77.9</td>
<td>0.701</td>
<td>94.6</td>
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<td>96.1</td>
<td>95.2</td>
<td>0.953</td>
<td>98.3</td>
</tr>
<tr>
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<td>98.8</td>
<td>0.987</td>
<td>99.4</td>
</tr>
</tbody>
</table>

![Image of classification results](image-url)
Figure 3.1: Classification maps obtained by different methods on AVIRIS Indian Pines dataset (overall accuracy are reported in the parentheses).

### 3.3.3 Visual Comparison

Figure 3.1 shows the classification maps by different algorithms on the Indian Pines image. Generally speaking, it indicates consistent results with the numerical measures. As we can see, algorithms without MLL prior, i.e. OMP, MPSR1, MPSR2 and MLRsub produced intense artifacts in the classification map due to the existence of mixed pixels in the image. Although all four algorithms performed seemingly well, careful inspection indicates that MPSR2 yields fewer artifacts than the others in certain classes, e.g. Grass/Pasture, Building-Grass-Tree-Drives and Soybeans-min. By combining with MLL prior, MPSR1MLL, MPSR2MLL and MLRsubMLL produced very smooth results, although there still exists misclassified patches in classes such as Soybeans-min and Building-Grass-Tree-Drives. Nevertheless, some small classes such as oats were totally misclassified, because of the lack of enough training samples for small classes. We also noticed that there are not much difference between the map of OMPMLL and OMP.

### 3.3.4 Sensitivity of Parameters

The Section explored the sensitivity of two important parameters, i.e. sparsity level and smooth cost for SR-based algorithms. Figure 3.2 plots the error bar of OA as a function of sparsity $\tau$ and smooth cost $\gamma$ based on the AVIRIS Indian Pines dataset.

Figure 3.2(a) indicates that MPSR-based algorithms achieved the highest performance when sparse level was 3. And from sparsity level of 3, the performance of MPSR-based algorithms reduced quite sharply. This is not surprising because increased sparsity level allows the wrong class to represent the test sample equally well as the true class, consequently leads to the loss of discriminative power. MPSR2MLL achieved higher OA than MPSR1MLL, and both MPSR1MLL and MPSR2MLL outperform OMPMLL when sparsity level is lower than 30. OMP achieved stable results and OMPMLL demonstrated slightly increased performance on high sparsity level.

In Figure 3.2(b), the increase in smooth cost increased the performance of MPSR1MLL and MPSR2MLL to a stable level, but did not indicate noticeable influence on OMPMLL. Moreover, MPSR2MLL achieved higher accuracy but lower variance than MPSR1MLL across most smooth-cost levels, indicating the worth of accounting for the variance heterogeneity in MPSR.
Figure 3.2: The error bar of OA as a function of sparsity $\tau$ (a) and smooth cost $\gamma$ (b).

### 3.4 Conclusion

In this chapter, we have proposed a mixture of probabilistic sparse representation approach to be integrated with MRF in Bayesian framework for hyperspectral image classification. We assume that the spectral vector in a class can be sparsely represented by the training samples in the same class. Moreover, the representation error is assumed being class-independent, with zero mean and diagonal covariance matrix. Based on these assumptions, we have derived the class conditional distribution of spectral vector, which is used with MRF labels prior distribution to form a MAP problem. The proposed approach is solved by graph cut $\alpha$-expansion techniques. On benchmark hyperspectral images, the proposed algorithm achieved new state-of-the-art performance.
Chapter 4
K-P-Means for Spectral Endmember Estimation

This chapter presents K-P-Means, a novel approach for spectral endmember estimation. Spectral unmixing is formulated as a clustering problem, with the goal of K-P-Means to obtain a set of "purified" spectral pixels to estimate endmembers. The K-P-Means algorithm alternates iteratively between two main steps (abundance estimation and endmember update) until convergence to yield final endmember estimates. Experiments using both simulated and real spectral images show that the proposed K-P-Means method provides strong endmember and abundance estimation results compared to existing approaches. © [2014] IEEE. Reprinted, with permission, from [Xu Linlin, Li J., Wong, A., and Peng, J., K-P-Means: a clustering algorithm of K “purified” means for spectral endmember estimation, IEEE Geoscience and Remote Sensing Letters, 03/2014].

4.1 Introduction

Accurate estimation of the spectra of pure materials called endmembers is essential to spectral unmixing that aims at estimating for each pixel the fractional abundances of endmembers. Current methods for endmember estimation can be categorized as geometric, statistical and sparse coding approaches (Bioucas et al., 2012). Although all these approaches have their own respective advantages, it is undeniable that endmembers extraction would be more straightforward if we have “pure” pixels that are due to individual endmembers, rather than multiple endmembers, for a number of reasons. First of all, classical geometric approaches that rely on the presence of pure pixels, such as vertex component analysis (VCA) (Nascimento & Bioucas-Dias, 2005) would achieve optimal performance. More intuitively, if we know the group of pixels that are due to a particular endmember, we can just use the mean value of pixels as an estimate of the endmember. Nevertheless, pure pixels are rare to obtain directly from the hyperspectral images due to factors such as low spatial resolution or the complexity of ground targets.

Given these considerations, this chapter therefore intends to explore the feasibility of obtaining “purified” pixels from mixed pixels in order to achieve simplified yet efficient endmember estimation. A “purified” pixel is defined as the residual of mixed pixel after removing the contribution of all endmembers except the one that dominates the pixel. We estimate “purified” pixels in two steps based on the abundance information of the hyperspectral image. First, we partition all pixels into several groups that are dominated by different endmembers. Second, for pixels in each group, we
remove the contributions due to non-dominant endmembers in that group. In the first step, since a cluster is defined by predominant endmembers, our approach differs from other label-utilizing approaches (Zare & Gader, 2010; Castrodad et al. 2011; Martin & Plaza, 2011) in spectral unmixing literature where a cluster may involve multiple significant endmembers. We treat the purified pixels in each group as realizations of endmember subject to random noise, and thereby use the expected value of the pixels as the endmember estimate. The resulting algorithm, which we will refer to as K-P-Means algorithm alternates iteratively between two main steps (abundance estimation and endmember update) until convergence to yield final endmember estimates. The capability of K-P-Means is proved by experiments on both simulated and real hyperspectral images.

4.2 K-P-Means

4.2.1 Problem Formulation and Motivations

This chapter addresses a linear spectral unmixing model where the observed spectral pixels stack $X$ is represented by endmember matrix $A$ and abundance matrix $S$ with independently identically distributed (i.i.d.) Gaussian noise $N$:

$$X = SA^T + N$$  \hspace{1cm} (4.1)

$$
\begin{bmatrix}
  x_1^T \\
  x_2^T \\
  \vdots \\
  x_n^T
\end{bmatrix} =
\begin{bmatrix}
  s_1^T \\
  s_2^T \\
  \vdots \\
  s_n^T
\end{bmatrix} (a_1, a_2, ..., a_K)^T +
\begin{bmatrix}
  n_1^T \\
  n_2^T \\
  \vdots \\
  n_n^T
\end{bmatrix}$$  \hspace{1cm} (4.2)

where $s_i$ is a $K \times 1$ nonnegative abundance vector, that measures the contribution of endmembers $a_j$ ($j = 1, 2, ..., K$) to $p \times 1$ dimensional spectral pixel $x_i$:

$$x_i = \sum_{j=1}^{K} s_{ij} a_j + n$$  \hspace{1cm} (4.3)

In most cases, the endmember collection $\{a_j\}$ contribute unequally to $x_i$, and the group of pixels dominated by $a_j$ is denoted by $G_j$. Therefore, the image can be partitioned into $K$ sets $G_j$ ($j = 1, 2, ..., K$). In order to reduce the coupling effect among endmembers, it is reasonable to infer $a_j$ ($j = 1, 2, ..., K$) separately from pixels in $G_j$. Nevertheless, mixed pixels in the same class may still admit multiple endmembers. In order to further remove the influence of less-dominant endmembers, it is desirable to use the proportion of $x_i$ that is solely due to the contribution of dominant endmember
to estimate $a_j$, as opposed to using $x_i$ wholly. We refer to $x_i$ after removing the contribution of less-dominant endmembers as “purified” pixel.

Not only good abundance information can be utilized to obtain “purified” pixels for enhanced endmember estimation, but accurate endmember estimates can in turn boost the accuracy of abundance estimation. Consequently, spectral unmixing can be treated as an iterative optimization issue by taking advantage of the label information from the abundance. We therefore present in the following Sections a K-P-Means clustering algorithm which intends to enhance endmember estimation based on the “purified” pixels by explicitly utilizing the label information.

### 4.2.2 K-P-Means Model

This Section formulates K-P-Means from a comparative perspective with the classical K-Means algorithm. In K-Means, the spectral vector in class $k$ can be expressed as:

$$x_i^k = m^k + n$$  \hspace{1cm} (4.4)

where $m^k$ is the mean vector of class $k$ and $n$ is class-independent white noise. Based on the following objective function:

$$\{m^k, l\} = min_{i,m} \sum_{k=1}^{K} \sum_{l_i=k} ||x_i - m^k||_2$$  \hspace{1cm} (4.5)

where $l = \{l_i | i = 1,2,...,n\}$ are the labels of pixels, K-Means algorithm iterates two steps: estimating $l$ given $\{m^k\}$, and estimating $\{m^k\}$ based on $l$.

Similarly, the generative model of K-P-Means is formulated as:

$$x_i^k = \sum_{j=1}^{K} s_{ij} a_j + n, \text{where } s_{ik} = \{s_{ij} \neq k\} \geq 0$$  \hspace{1cm} (4.6)

where the general term $m^k$ in K-Means is expressed more specifically by $\sum_{j=1}^{K} s_{ij} a_j$. It means that K-Means characterize a class by the mean vector $m^k$, while K-P-Means defines the class by the dominant endmember $a_k$, whose abundance $s_{ik}$ is the biggest. Therefore, comparing with K-Means that considers the overall effect of a physical process, K-P-Means probes into the sources of the physical process that contribute to the observations. The object function of K-P-Means can be expressed as:

$$\{a_k, l\} = min_{l,a} \sum_{k=1}^{K} \sum_{l_i=k} ||y_i - a_k||_2$$  \hspace{1cm} (4.7)

where $x_i$ in objective function of K-Means is substituted by:
Therefore, as opposed to K-Means that adopts mixed pixels \( \{x^k_i\} \) in class \( k \) for estimating the mean vector \( m^k \), K-P-Means excludes the contribution of less significant endmembers from estimating dominant endmember \( a_k \). Accordingly, \( a_k \) in the proposed algorithm can be treated as the mean vector of “purified” spectral pixels \( \{y^k_i\} \). That’s why our algorithm is termed K-P-Means. Based on above described model, K-P-Means iterates abundance estimation and endmember estimation, just as the two steps in K-Means, which are introduced in Section 4.2.3 and 4.2.4 respectively.

### 4.2.3 Abundance Estimation

Following Eq. (4.7), given \( \{a_j\} \), pixel labeling requires solving the following optimization issue:

\[
l_i = \arg\min_k \|x_i - \sum_{j \neq k}^K s_{ij} a_j \|_2 / s_{ik} \quad \text{s.t.} \quad \{s_{ij}\} \geq 0 \quad \text{and} \quad s_{ik} > \{s_{ijk}\} \tag{4.9}
\]

It means that \( x_i \) is associated with the \( k \)th endmember \( a_k \) which will take the largest coefficient \( s_{ik} \) when representation error is minimized. Suppose \( \{a_k\} \) are of similar scale, this optimization issue is equivalent to firstly estimating \( \{s_{ij}\} \) by solving:

\[
\arg\min_{\{s_{ij}\}} \|x_i - \sum_{j}^K s_{ij} a_j \|_2 \quad \text{s.t.} \quad \{s_{ij}\} \geq 0 \tag{4.10}
\]

then determining \( l_i \) by:

\[
l_i = \arg\max_k s_{ik} \tag{4.11}
\]

As we can see, the estimation of abundance in Eq. (4.10) is essentially a non-negative least square (NNLS) issue which can be efficiently solved by method in Lawson & Hanson (1974). Note that the sum-to-one constraint is not necessary, since we only need the relative magnitudes of abundances to determine dominant endmember. Therefore both K-Means and K-P-Means measure the “relevance” of a pixel to different clusters in order to determine its label. Nevertheless for K-Means the “relevance” is measured by the geometric “closeness” from the pixel to class centers, while for K-P-Means, it is measured by the magnitude of nonnegative contribution of endmembers to the representation of the pixel in a least squares sense.

### 4.2.4 Endmember Estimation

Following Eq. (4.7), given \( \{y^k_i\} \), K-P-Means update \( a_k \) based on the following generative model:
\[ y_i^k = a_k + n \] (4.12)

Since \( n \) is i.i.d. zero-mean Gaussian noise, the maximum likelihood estimation (MLE) of \( a_k \) is the expected value of \( \{y_i^k\} \). Note that it is possible to apply other endmember extraction techniques, such as VCA on \( y_i^k \) to produce candidates of \( a_k \), it however will introduce extra problems, such as the difficulty to determine the most relevant one.

4.2.5 Complete Algorithm

Assembling abundance estimation in Section 4.2.3 and endmember update in Section 4.2.4 into the iterative optimization framework, leads to the complete algorithm of K-P-Means, which is detailed in Algorithm 1. In endmember update step, in order to speed up convergence, the update of an endmember is allowed to utilize the endmembers that have been updated. The iteration of the two steps will stop if either the spectral angle difference (SAD, see Section 4.3) between endmember estimates in two continuous iterations is smaller than a given value (i.e. \( \tau \)), or a predefined maximum number of iteration (i.e. \( \text{iters} \)) is reached.

Algorithm 1: K-P-Means

- **Input**: spectral stack \( X \), number of clusters \( K \) and iteration threshold \( \tau \);
- **Output**: endmember \( \hat{A} \) and abundance \( \hat{S} \);
- **Initialization**: \( t := 1, \hat{A}^{(0)} = \text{VCA}(X) \); or \( \hat{A}^{(0)} = \text{randomly selected pixels \{x_r\}} \);
- **while** \( t \leq \text{iters} \) or \( \text{SAD}(\hat{A}^{(t)}, \hat{A}^{(t-1)}) \geq \tau \) **do**
  - (1) \( \hat{S} = \text{NNLS}(\hat{A}^{(t)}, X), \hat{l} \leftarrow \text{max}(\hat{S}) \);
  - (2) **for** \( k = 1, 2, ..., K \)
    - \( y_i^k = (x_i^k - \sum_{j \neq k}^K S_{ij} \hat{a}_j^{(t)}/s_{ik} \)
    - \( \hat{a}_k^{(t)} = \text{mean}(\{y_i^k\}) \)
  - **end for**
- **end while**

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4.3 Experiments

4.3.1 Simulated Study

A 64×64 sized image with mixed pixels of 4 endmembers randomly selected from the USGS digital spectral library (Clark et al., 1993) are simulated, following the procedure reported in Miao & Qi (2007). Using the 4 endmembers, mixed pixels are created by firstly dividing the entire image into 8×8 sized homogeneous blocks of one of the 4 endmembers, then degrading the blocks by applying a spatial low pass filter of 7×7. To further increase mixing degree, the remaining relatively “pure” pixels with 80% or larger single abundance are forced to take equal abundances on all endmembers. Zero-mean i.i.d. Gaussian noise is added to further degrade the image. The resulting image therefore resembles a highly mixed hyperspectral image with measurement errors or sensor noise, which is very challenging for spectral unmixing algorithms.

Two techniques, VCA [2] and MVC-NMF (Miao & Qi, 2007) are implemented using the code provided by their authors. VCA represents classical techniques that rely on the existence of pure pixels. Since VCA only extract endmembers, we estimate abundance using NNLS (Lawson & Hanson, 1974). The comparison with MVC-NMF is desirable since both K-P-Means and MVC-NMF deal with highly mixed pixels. MVC-NMF used as initial parameters the endmember estimated by VCA, and 150 iterations in maximum.

Moreover, three variants of K-P-Means are implemented. K-P-Means used as initial parameters both endmembers produced by VCA and pixels selected randomly from dataset, in order to explore the sensitivity of K-P-Means to initial parameters. The resulting algorithms are referred to as K-P-Means-VCA and K-P-Means-Random respectively. In K-P-Means-Random, 5 replicates are performed, each with a new set of initial endmembers, to obtain the solution with smallest residual. In order to prove the effectiveness of using “purified” pixels in K-P-Means, we introduce for comparison the “non-purified” approach (i.e. K-nonP-Means), where \( \hat{x}_i^k \) instead of \( y_i^k \) is used in Section 4.2.4 to update endmembers. All variants are implemented with \( \text{iters}=50 \) and \( \tau=0.01 \) without explicit explanation.

The consistency between estimated endmember \( \hat{a} \) and true endmember \( a \) is measured by the widely used spectral angle distance (SAD), defined as: \( \text{SAD} = \cos^{-1}(\langle a^T \hat{a} \rangle / (\|\hat{a}\|\|a\|)) \), and the spectral information divergence (SID), expressed as \( \text{SID} = D(a/\hat{a}) + D(\hat{a}/a) \), where \( D(x/y) \) measures the relative entropy between \( x \) and \( y \) (Chang & Heinz, 2000). The numerical measures for abundance \( s \)
are achieved by replacing \( a \) with \( s \) in SAD and SID. The resulting measures are called AAD and AID respectively.

The five methods are performed on simulated image to produce numerical measures. In order to investigate the noise robustness of different methods, they are tested on images with different noise levels measured by signal-to-noise ratio (SNR) (Miao & Qi, 2007). For each noise level, 20 images with independent noise realizations are processed to obtain statistics of numerical measures, as reported in Figure 4.1.

![Figure 4.1](image)

Figure 4.1: Performance comparison at different noise levels in terms of (a) SAD, (b) SID, (c) AAD and (b) AID. In these four statistics, smaller value means better result.

Overall, K-P-Means-VCA achieved much smaller SAD and SID values than VCA, and close results to MVC-NMF across all noise levels, indicating that K-P-Means is capable of extracting accurately the endmembers in highly mixed and noisy circumstance. Moreover, the endmember estimation of K-P-Means-VCA measured by SAD and SID seemed to be robust to noise level. As we can see, SAD and SID remained at very low values with SNR decreasing from 45 to 20, although
from 20 to 10 there was large increase in SAD and SID. When SNR=10, we noticed that K-P-Means-VCA achieved smaller SAD and SID than MVC-NMF.

In terms of abundance estimation, K-P-Means-VCA outperformed VCA according to the mean AAD and AID values across all noise levels. The variances of AAD and AID are also smaller in K-P-Means-VCA than in VCA. MVC-NMF achieved lower AAD and AID values than K-P-Means-VCA. But this advantage is less significant at low noise level. Overall, these results demonstrate that K-P-Means-VCA can achieve fairly accurate abundance estimation, although it is designed primarily for enhanced endmember extraction.

The observation that K-nonP- Means performed worse than K-P-Means-VCA and K-P-Means-Random demonstrates the importance and benefits of using “purified” pixels instead of the original pixels for endmember estimation. K-P-Means-Random outperformed VCA in terms of all measures across all noise levels, indicating K-P-Means is capable of achieving acceptable performance with random initializations. It is not surprising that K-P-Means-VCA performed better than K-P-Means-Random, considering the fact that good initial parameters can optimize the convergence properties of ill-posed optimization problems.

Endmember estimation by VCA was insensitive to the noise level change. The SAD and SID stayed almost unchanged with decreasing of SNR from 45 to 20. MVC-NMF performed better than the rest techniques in most cases, although its performance of endmember estimation decreased very fast from SNR = 20 to 10. We noticed that MVC-NMF performed very well when SNR=10 in Miao & Qi (2007). This inconsistency is probably because we used different endmember for simulation.

Table 4.1: Performance of K-P-Means-VCA and VCA, measured by mean SID and AID, over different image size and varying number of endmembers.

<table>
<thead>
<tr>
<th>Image size</th>
<th>Number of endmembers</th>
<th>64</th>
<th>128</th>
<th>256</th>
<th>512</th>
<th>4</th>
<th>6</th>
<th>8</th>
<th>12</th>
<th>15</th>
</tr>
</thead>
<tbody>
<tr>
<td>SID*1000</td>
<td>K-P-Means</td>
<td>1</td>
<td>0.5</td>
<td>0.3</td>
<td>0.6</td>
<td>1</td>
<td>2</td>
<td>14</td>
<td>21</td>
<td>50</td>
</tr>
<tr>
<td></td>
<td>VCA</td>
<td>7.5</td>
<td>5.1</td>
<td>5.6</td>
<td>5.6</td>
<td>6</td>
<td>10</td>
<td>59</td>
<td>57</td>
<td>100</td>
</tr>
<tr>
<td>AID</td>
<td>K-P-Means</td>
<td>1.0</td>
<td>0.9</td>
<td>0.8</td>
<td>1.0</td>
<td>0.9</td>
<td>2.0</td>
<td>3.4</td>
<td>6.1</td>
<td>6.9</td>
</tr>
<tr>
<td></td>
<td>VCA</td>
<td>2.6</td>
<td>1.5</td>
<td>1.5</td>
<td>1.7</td>
<td>1.8</td>
<td>4.9</td>
<td>6.8</td>
<td>8.9</td>
<td>12.9</td>
</tr>
</tbody>
</table>

In order to explore the sensitivity of K-P-Means to image size and number of endmembers, Table 4.1 presents the performances of VCA and K-P-Means-VCA, measured by mean SID and AID, over increasing image sizes from 64×64 to 512×512 and the numbers of endmembers from 4 to 15.
Generally speaking, K-P-Means is not sensitive to the increasing of image size, and the mean SID and AID values that achieved by K-P-Means-VCA are respectively around 10% and 50% of those achieved by VCA. However, the performances of both VCA and K-P-Means-VCA deteriorated with the increase of the number of endmembers. Nevertheless, the SID and AID values achieved by K-P-means are respectively 25% and 50% of the statistics achieved by VCA on average.

All algorithms were implemented under the MATLAB platform. On average, it took 0.04, 6.51, and 26.24 seconds, respectively by VCA, K-P-Means-VCA and MVC-NMF, to process images with 64×64 pixels, on a PC with a Pentium(R) 2.30GHZ Quad-Core processor.

4.3.2 Test on Real Hyperspectral Images

The Indian Pines image, captured by Airborne Visible/Infrared Imaging Spectrometer (AVIRIS) over a vegetation area in Northwestern Indiana, USA is used to test the proposed algorithms. The image has spatial resolution of 20m and contains 200 spectral reflectance bands after removing 20 water absorption bands (104–108, 150–163, and 220). The image consists of 145 × 145 pixels belonging to 16 different land cover types, as shown in Figure 4.2.

![Figure 4.2: The ground-truth map of 16 classes in AVIRIS Indian Pines image.](image-url)
In this experiment, K-P- Means- Random with \( iters = 50 \) and \( \tau = 0.01 \) extracted a number of 20 endmembers from pixels covered by ground truth classes. The abundance maps of eight selected endmembers are shown in Figure 4.3. As we can see, the maps from left to right, top to bottom correspond respectively to Grass/Trees, Hay-windrowed, Grass/Pasture, Soybeans-min, Corn-notill, Wheat, Wood, and Stone-steel Towers. These correspondences between abundance maps and ground truth classes may indicate that K-P- Means identified accurately the endmembers in the image, considering that different endmembers tend to dominate different classes. Nevertheless, the bright areas of most abundance maps do not match very well the ground truth, except Wheat and Stone-steel Towers. It is not surprising considering the gap that while K-P- Means is designed to identify
individual endmembers, the pixels in the same ground truth class may actually assume multiple significant endmembers, due to the complexity of ground targets in Indian Pines image.

Figure 4.4 shows the maps of the eight corresponding endmembers achieved by MVC-NMF, for comparison purpose. As we can see, most endmember maps achieved by MVC-NMF do not match the ground truth as well as the maps achieved by K-P-means, except the two maps correspond to Wheat and Stone-steel Towers.

4.4 Conclusion

This chapter has presented a K-P-Means algorithm for spectral endmember extraction. Based on abundance information, we proposed to obtain the “purified” pixels from the original mixed pixels for enhanced endmember estimation, which can in turn aid abundance estimation. Therefore, we interpreted spectral unmixing as an iterative optimization problem, and designed the K-P-Means algorithm which alternates iteratively between two main steps (abundance estimation and endmember update) until convergence to yield final endmember estimates. Experiments on both simulated and real hyperspectral images proved that K-P-Means is capable of estimating accurately both the endmembers and abundance.
Chapter 5
Clustering-based PCA for SAR Image Denoising

The combination of nonlocal grouping and transformed domain filtering has led to the state-of-the-art denoising techniques. In this chapter, we extend this line of study to the denoising of Synthetic Aperture Radar (SAR) images based on clustering the noisy image into disjoint local regions with similar spatial structure and denoising each region by the Linear Minimum Mean-Square Error (LMMSE) filtering in Principal Component Analysis (PCA) domain. Both clustering and denoising are performed on image patches. For clustering, to reduce dimensionality and resist the influence of noise, several leading principal components (PCs), identified by the Minimum Description Length (MDL) criterion are used to feed the K-means clustering algorithm. For denoising, to avoid the limitations of the homomorphic approach, we build our denoising scheme on additive signal-dependent noise (ASDN) model and derive a PCA-based LMMSE denoising model for multiplicative noise. Denoised patches of all clusters are finally used to reconstruct the noise-free image. The experiments demonstrate that the proposed algorithm achieved better performance than the referenced state-of-the-art methods in terms of both noise reduction and image details preservation. © [2014] IEEE. Reprinted, with permission, from [Xu Linlin, Li J., Shu, Y., and Peng, J., SAR image denoising via clustering-based principal component analysis, IEEE Transactions on Geoscience and Remote Sensing, 03/2014].

5.1 Introduction
Synthetic Aperture Radar (SAR), as a coherent imaging system is inherently suffering from the speckle noise, which has granular appearance and greatly impedes the automatic image processing and visual interpretation. Although multilook averaging is a common way to suppress speckle noise at the cost of reduced spatial resolution, it is more favorable to develop suitable filtering techniques. Classical filters, such as Lee filter (Lee, 1980), Frost filter (Frost et al., 1982) and Kuan filter (Kuan et al., 1985) that denoise SAR images in spatial domain by recalculating the center pixels of the filtering windows based on the local scene heterogeneity, although work well in stationary image area, they tend to either preserve speckle noise or erase weak scene signal at heterogeneous areas, e.g. texture area, boundary, line or point targets. In order to better preserve image edges, Yu & Acton (2002) designed a speckle reduction anisotropic diffusion (SRAD) method which can be treated as an edge-sensitive version of the classical filters. The performance of the Gamma MAP filter (Lopes et al.,
which denoises SAR image via maximum a posteriori criteria, depends highly on whether the imposed Gamma distribution can accurately describe SAR image.

Instead of denoising in spatial domain, it has been proved more efficient to perform the task in transformed domain where signal and noise are easier to separate. The wavelet techniques assume that noise mainly exists on the high frequency wavelet components and thus can be removed by filtering the wavelet coefficients in transformed domain. This idea has proved great success to denoise additive white Gaussian noise (AWGN). To adapt wavelet for SAR denoising, many techniques adopted the homomorphic approach where speckle noise subject to log-transformation is treated as AWGN and denoised in wavelet domain by thresholding (Gagnon & Jouan, 1997; Guo et al., 1994) or modeling (Achim et al., 2003; Solbø & Eltoft, 2004; Bhuiyan et al., 2007) the wavelet coefficients. However, since the performance of denoising is very sensitive to logarithmic operation that tends to distort the radiometric properties of SAR image, techniques based on additive signal-dependent noise (ASDN) model were developed in (Argenti & Alparone, 2002; Xie et al., 2003; Argenti et al., 2006; Argenti et al., 2008).

Although wavelet-based denoising methods have proved better efficiency than classical filters, limitations reside in the inadequate representation of various local spatial structures in images using the fixed wavelet bases (Muresan & Park, 2003; Zhang et al., 2010; He et al., 2011). On the other hand, locally learnt Principal Component Analysis (PCA) bases, a series of mutually orthogonal directions with sequentially largest variances, have shown better capability of representing structural features, e.g. image edges and texture. In PCA domain, the scene signal is mostly captured by several leading Principal Components (PCs), while the last few components with low variances are mainly due to noise. The denoising of AWGN has been achieved by filtering the PCs through linear minimum mean-square error (LMMSE). Examples include the adaptive PCA denoising scheme proposed by Muresan and Parks (2003) and local pixel grouping PCA (LPG-PCA) algorithm proposed by Zhang et al. (2010). Both methods have proved to be more effective than the conventional wavelet-based denoising methods. However, no efforts have been made to adapt PCA-based denoising to SAR images. Since SAR images assume signal-dependent noise, a new denoising model that takes into account this particularity is required.

A different line of research initiated in Buades et al. (2005) approaches image denoising as a nonlocal means (NLM) problem, where ‘nonlocal’ pixels whose neighborhoods share similar spatial structure, rather than ‘local’ pixels that are just geometrically near to each other are used to perform
weighted average with the weights measured by the Euclidean distances between the referenced image patch and the other candidate patches. The NLM method has been adapted for SAR image denoising by adjusting the similarity measure to the multiplicative nature of speckle noise (Coup et al., 2008; Zhong et al., 2009; Deledalle et al., 2009). Particularly, the probabilistic patch-based (PPB) algorithm in Deledalle et al. (2009) proved to achieve the state-of-the-art performance for SAR image denoising. Moreover, the idea of NLM has been extended to combine with the transformed domain denoising approaches, leading to the state-of-the-art image denoising techniques, e.g. the block-matching 3D filtering (BM3D) (Dabov et al., 2007), LPG-PCA (Zhang et al., 2010) and SAR-BM3D (Parrilli et al., 2012) algorithms. All methods take advantage of the enhanced sparsity in transformed domain when denoising is performed on image patches with similar structure. In these methods, block-matching approach was adopted to find for each patch in the image a group of similar patches. However, this approach faces the difficulty to define the threshold as to how “similar” to the reference patch is acceptable. It also has high computational cost.

In this study, we extend this line of study to denoise SAR images by explicitly addressing two issues. First, we build a new denoising model based on PCA technique to account for the multiplicative nature of speckle noise. Based on ASDN model, we derive a LMMSE approach for solving PCA-based denoising problem. Our approach is the first to build the PCA-based denoising method on the ASDN model for SAR image denoising. Besides SAR images, it is also applicable to other signal-dependent noise. Second, instead of using block-matching approach, we employ a clustering approach. We propose to use the combination of log-transformation, PCA and K-means methods for finding similar patches. Based on the statistical property of speckle noise, we proved the compatibility between PCA features and the K-means model. This clustering approach proved to be an competitive alternative to the block-matching approach adopted in Zhang et al. (2010), Deledalle et al. (2009) and Parrilli et al. (2012).

The rest of the chapter is organized as follows. Section 5.2 discusses data formation and PCA analysis. Section 5.3 derives the LMMSE filtering of speckle noise in PCA domain. Section 5.4 details the clustering-based scheme for SAR image denoising. In Section 5.5, the complete procedure of the proposed strategy that involves a second stage is discussed. In Section 5.6, experiments are designed to compare the proposed method with other popular denoising techniques. Results obtained using both simulated and real SAR images are presented and discussed. Section 5.7 concludes this study.
5.2 Data Formation & PCA Analysis

The SAR image as a collection of all the image patches is represented by a data matrix

\[ X = [x_1, x_2, \ldots, x_n]^T \]  \hspace{1cm} (5.1)

where \( n \) denotes the number of pixels, and \( x_i \) (\( i = 1, 2, \ldots, n \)) is a \( p \times 1 \) vector, representing the \( i \)th patch which is a small square window centered at the \( i \)th pixel.

![SAR image with patch variable](image)

**Figure 5.1:** Illustration of the acquisition of a patch in SAR image.

The PCA can be achieved by performing Singular Value Decomposition (SVD) on the covariance matrix of \( X \):

\[ \Sigma_x = \begin{pmatrix} C_{aa} & C_{ab} & \cdots & C_{ai} \\ C_{ba} & C_{bb} & \cdots & C_{bi} \\ \vdots & \vdots & \ddots & \vdots \\ C_{ia} & C_{ib} & \cdots & C_{ii} \end{pmatrix} = \begin{pmatrix} a_1^T \\ a_2^T \\ \vdots \\ a_p^T \end{pmatrix} \begin{pmatrix} \delta_1 & 0 & \cdots & 0 \\ 0 & \delta_2 & \cdots & 0 \\ \vdots & \vdots & \ddots & \vdots \\ 0 & 0 & \cdots & \delta_p \end{pmatrix} \begin{pmatrix} a_1^T \\ a_2^T \\ \vdots \\ a_p^T \end{pmatrix} \]  \hspace{1cm} (5.2)

where element \( C_{AB} \) in \( \Sigma_x \) represents the covariance between the two pixels at position \( A \) and \( B \) across the image. So \( \Sigma_x \) provides a statistical description of the relationship among pixels in SAR image. Pixels that do not belong to the same patch are considered uncorrelated. Thus the size of the patch determines the scale of spatial patterns that can be captured. Generally speaking, bigger sized patch considers larger range correlations and hence is more capable of capturing larger-scale repeated patterns in SAR image. \( a_i \) (\( i = 1, \ldots, p \)), \( p \times 1 \) vectors, denote the sequence of mutually orthogonal PCA bases onto which the projection of patches stack \( X \) produces the PCs with sequentially largest variances represented by \( \delta_i \) (\( i = 1, \ldots, p \)).

In PCA domain, several leading PCs capture most of the scene signal in image patches, while the last few components are mostly due to noise. In this chapter, we use PCA for both denoising and feature extraction. In Section 5.3, we develop a LMMSE criterion based on ASDN model to shrink the PCs. Thus denoising can be achieved by reconstructing SAR image using the processed PCs. In
Section 5.4, we use several leading PCs in logarithmic space identified by the Minimum Description Length (MDL) criterion to feed the K-means algorithm. This not only reduces the dimensionality and de-correlates the spatial variables, but also suppresses the noise contained in image patches.

5.3 SAR Image Denoising in PCA Domain

In Muresan & Parks (2003) and Zhang et al. (2010), the LMMSE shrinkage was conducted in PCA domain to remove AGWN. For SAR speckle noise, we can certainly adopt the homomorphic approach and apply the same methods in Muresan & Parks (2003) and Zhang et al. (2010). However since the performance of denoising is sensitive to log-transformation that tends to distort the radiometric dynamics of the SAR data, it is more appropriate to perform denoising in original space instead of logarithmic space. In this Section, we derive a new LMMSE shrinkage approach under the ASDN model. We assume the speckle noise is fully developed, thus a SAR image pixel can be modeled as:

\[ x = y \varepsilon \quad (5.3) \]

where \( y, \varepsilon, \) and \( x \) represent respectively unobserved scene signal, speckle noise and the observed signal. Based on Eq. (5.3), we get the ASDN model as:

\[ x = y + n \quad (5.4) \]

where \( n = y(\varepsilon - 1) \). Because \( \varepsilon \) has unit mean, thus \( n \) is a zero-mean signal-dependent noise. Hence the patch variable in SAR image can be described by:

\[ x = y + n \quad (5.5) \]

where \( x = [x_1, x_2, ..., x_p]^T \), \( y = [y_1, y_2, ..., y_p]^T \), \( n = [n_1, n_2, ..., n_p]^T \). Denote the covariance between \( x_i \) and \( x_j \) by \( \sigma_{x}^{ij} \), and the mean of \( x_i \) by \( \mu_{x}^{i} \), we can get:

\[
\sigma_{x}^{ij} = E[(x_i - \mu_{x}^{i})(x_j - \mu_{x}^{j})]
= E[(y_i\varepsilon_i - \mu_{x}^{i})(y_j\varepsilon_j - \mu_{x}^{j})] \quad (5.6)
\]

For fully developed speckle noise, \( x \) and \( s \) are uncorrelated, so we get \( E(y\varepsilon) = E(y)E(\varepsilon) \). Because \( E(\varepsilon) = 1 \), we get \( E(x) = E(y)E(\varepsilon) = E(y) \). So

\[
\sigma_{x}^{ij} = E(y_iy_j)E(\varepsilon_i\varepsilon_j) - \mu_{x}^{i}\mu_{x}^{j}
\]
We assume speckle noise is spatially uncorrelated, i.e. \( \sigma^{ij}_\varepsilon = 0 \) for \( i \neq j \). Thus we have:

\[
\sigma^{ij}_\varepsilon = \sigma^{ij}_y \quad \text{(for } i \neq j) 
\]

In the following analysis, we represent the empirical mean of the patches in \( \mathbf{X} \) by \( \bar{X} \) and we assume that the patch variable \( \mathbf{x} \) has been centralized. Denote the covariance of \( \mathbf{x} \) by \( \Sigma_x \), the PCA bases can be obtained by performing SVD on \( \Sigma_x \):

\[
\Sigma_x = \mathbf{A} \Delta \mathbf{A}^T
\]

where the column vectors in \( \mathbf{A} \) represent the PCA bases with sequentially largest variances, \( \Delta \) is the diagonal matrix of the variances of PCs, which are the projection of patch variables onto PCA bases:

\[
\mathbf{s} = \mathbf{A}^T \mathbf{x} = \mathbf{A}^T \mathbf{y} + \mathbf{A}^T \mathbf{n} = \mathbf{s}_y + \mathbf{s}_n
\]

where \( \mathbf{s}_y = \mathbf{A}^T \mathbf{y} \) and \( \mathbf{s}_n = \mathbf{A}^T \mathbf{n} \) stand respectively for the signal and noise parts in the projection. If \( \mathbf{s}_y \) is known, denoising can be achieved by performing inverse PCA transformation using \( \mathbf{s}_y \). In this research, \( \mathbf{s}_y \) is estimated by LMMSE criterion.

\[
\hat{\mathbf{s}}_y = E(\mathbf{s}_y) + \text{Cov}(\mathbf{s}_y, \mathbf{s}) \Sigma_x^{-1}(\mathbf{s} - E(\mathbf{s})) = 0 + \Sigma_{s_y} \Sigma_x^{-1}(\mathbf{s} - \mathbf{0}) = \Sigma_{s_y} \Delta^{-1} \mathbf{s}
\]

The \( ij \)th element of \( \Sigma_{s_y} \) can be estimated through the following equation:

\[
\sigma^{ij}_{s_y} = E \left( s_y(i)s_y(j) \right) - \mu_{s_y}^{i} \mu_{s_y}^{j} \\
= E \left[ [A^T(i,:)] [A^T(j,:)] \right] - \mu_{x}^{i} \mu_{y}^{j} \\
= E \left\{ \sum_k A(k,i)y_k \sum_t A(t,j)y_t \right\} - \sum_k A(k,i)\mu_{y}^{k} \sum_t A(t,j)\mu_{y}^{t} \\
= E \left\{ \sum_k \sum_t A(k,i)A(t,j)y_ky_t \right\} - \sum_k \sum_t A(k,i)A(t,j)\mu_{y}^{k} \mu_{y}^{t}
\]
\[
\begin{align*}
&= \sum_k \sum_t A(k, i) A(t, j) [E(y_k y_t) - \mu^k \mu^t] \\
&= \sum_k \sum_t A(k, i) A(t, j) \sigma^k \sigma^t \\
&= \Sigma_k A(k, i) A(k, j) \sigma^k + \Sigma_k \Sigma_{t, t\neq k} A(k, i) A(t, j) \sigma^k \sigma^t \\
&= \Sigma_k A(k, i) A(k, j) \sigma^k + \Sigma_k \Sigma_{t, t\neq k} A(k, i) A(t, j) \sigma^k \sigma^t
\end{align*}
\] (5.12)

Denote \( u_j = \varepsilon_j - 1 \). Because \( x_j \) and \( u_j \) are uncorrelated for fully developed speckle noise, we can get

\[
E(x_j^2) = E[(y_j + y_j u_j)^2] = E(y_j^2) + E(y_j^2 u_j^2) = E(y_j^2) (1 + \sigma^u)
\] (5.13)

From Eq. (5.13) and Eq. (5.7), we get \( \sigma^k = \sigma^x - \frac{\sigma^u}{\sigma^u + 1} E(x_k^2) \), so

\[
\sigma^x_y = \Sigma_k A(k, i) A(k, j) \left[ \sigma^x - \frac{\sigma^u}{\sigma^u + 1} E(x_k^2) \right] + \Sigma_k \Sigma_{t, t\neq k} A(k, i) A(t, j) \sigma^k \sigma^t
\] (5.14)

From Eq. (5.8), we get \( \sigma^k \sigma^t = \sigma^k \sigma^t \) for \( t \neq k \), so

\[
\sigma^x_y = \Sigma_k A(k, i) A(k, j) \left[ \sigma^x - \frac{\sigma^u}{\sigma^u + 1} E(x_k^2) \right] + \Sigma_k \Sigma_{t, t\neq k} A(k, i) A(t, j) \sigma^k \sigma^t
\] (5.15)

In Eq. (5.15), \( \sigma^u \) can be calculated from the theoretical distribution of speckle noise, e.g. for Gamma distribution \( \sigma^u = 1/L \) (Xie et al., 2002), where \( L \) stands for the ENL. \( A(i, j) \) is the \( i \)th element of \( A \) in Eq. (5.9). \( E(x_k^2) \) and \( \sigma^k \sigma^t \) are estimated by the respective empirical values: \( E(x_k^2) = \Sigma (k, k) + \Sigma^2(k) \), \( \sigma^k \sigma^t = \Sigma (k, t) \), where \( \Sigma (i, j) \) stands for the \( i \)th element of \( \Sigma_x \) and \( \Sigma^2(i) \) stands for the \( i \)th element of \( \Sigma_x \).

Given \( \Sigma_x \) in Eq. (5.11), the noise-free image patch can be obtained by performing inverse PCA transformation:

\[
\hat{\Sigma_x} = \Sigma_x + A \hat{\Sigma_y}
\] (5.16)

The denoised patches will finally be used to construct the noise-free SAR image.

**5.4 Clustering Scheme**

The effectiveness of denoising in PCA domain depends highly on whether the PCs can sparsely represent the scene signal. And the sparsity can be achieved by performing analysis on patches with similar spatial structure. There are two approaches for finding similar patches, block-matching and
clustering. Block-matching is a supervised approach, which finds, for each pixel on the image, a group of patches that are “similar” to the reference patch. This approach has high computational cost. Since it is hard to define the “threshold” as to how “similar” to the reference patch is acceptable, the researchers always turn to guarantee a minimum number of similar patches. However, it may render some of the selected patches less relevant to the referenced patch. As opposed to the block-matching approach adopted in BM3D and NLM, the clustering approach involved in the proposed approach finds similar patches in an unsupervised manner by adaptively partitioning the image into disjoint areas. It requires less computation. Moreover, since the “threshold” in clustering approach is adaptively determined by comparing the proximities of the candidate patch to different cluster centers, rather than being pre-defined, the clustering approach is supposed to be more capable of finding relevant patches than block-matching approach.

In this study, we adopt K-means algorithm (Lloyd, 1982) proposed by Lloyd considering its simplicity and speed. And we use the Euclidean distance to measure similarity in feature space. Performing K-means clustering on image patches also faces problems, such as high dimensionality, high correlation among features, and intense iterations due to poor initial parameter values. In this study, we adopt log-transformation and PCA to extract compact features to feed the K-means algorithm.

This Section is organized as follows: we start with the illustration of feature extraction techniques; we then prove the compatibility of the extracted features and K-means algorithm; lastly, we discuss parameter tuning and efficient realizations of the clustering algorithm.

5.4.1 Feature Extraction

Before extracting features for clustering, we apply log-transformation on original SAR image as a pre-processing step. It has been common practice to aid clustering by preprocessing heavy tail distributed variables using log-transformation (Liu et al., 2003; Liu et al., 2007). In particular, Liu in Liu et al. (2003) indicated that log-transformation significantly improved the clustering result, and Liu et al. (2007) demonstrated a 10% increase in clustering accuracy after applying the log-transformation. The speckle noise in SAR image follows Gamma distribution that is long-tailed. Moreover, speckle noise is signal-dependent which means it has bigger variance on brighter image areas. This unstable nature would produce large between-cluster overlapping. So the log-transformation is used here to de-skew the dataset and to stabilize the variance. The log-transformed data tends to be symmetrically
distributed with constant variance, thus is more desirable for statistical methods, such as PCA and K-means algorithms.

Although clustering can be performed directly on image patches, it always suffers from high dimensionality e.g. 5×5 patch produces 25 variables, and the intense speckle noise contained in the patch. In order to solve these problems, we adopt PCA as a feature extraction technique. Only K leading PCs that are mostly due to signal are used as features for clustering. The accurate estimation of K is important in the sense that the underestimation would lose useful information but overestimation would introduce noise and unnecessary computation cost. One popular approach determines K by setting a threshold to the percentage of variation explained by signal components. This approach is simple but rather subjective. In this study, we estimate K by MDL criterion which was proposed by Rissanen (1978) and has been used to determine the number of signals in Wax & Kailath (1985). The p × 1 dimensional image patch variable \( \mathbf{x} \) subject to log-transformation can be represented by the following equation:

\[
\mathbf{x} = \sum_{i=1}^{K} \mathbf{A}(:,i) \mathbf{s}_i + \mathbf{n}
\]  

where \( \mathbf{A} \) is the eigenvector matrix, whose ith column \( \mathbf{A}( :, i ) \) denotes the ith PCA bases, and \( \mathbf{n} \) denotes the log-transformed speckle noise that roughly satisfies Gaussian distribution with zero mean and diagonal covariance matrix \( I_p \sigma_n^2 \). We assume that the scene signal \( \mathbf{y} \) can be reconstructed by the first K PCs

\[
\mathbf{y} = \sum_{i=1}^{K} \mathbf{A}( :, i ) \mathbf{s}_i
\]  

where \( \mathbf{s}_i = \mathbf{y}^T \mathbf{A}( :, i ) \) stands for the ith PC. We can see that Eq. (5.17) is same with Eq. (1) in Wax and Kailath (1985), where the authors estimated the number of signals by

\[
K_{MDL} = \arg \min_k (p - k) \log \frac{\Sigma_{k+1}^p \delta_j}{p-k} - \log \prod_{k+1}^p \delta_j + \frac{k(2p-k)}{2n} \log n
\]  

where \( \delta_j \) stands for the jth biggest eigenvalue of \( \Sigma_{\mathbf{y}} \), n denotes the number of observations. \( K_{MDL} \) can be easily determined by comparing all the \( p - 1 \) solutions.

5.4.2 The Compatibility of PCA Features and K-means algorithm

K-means algorithm that relies on Euclidean distance implicitly assumes that the noise of input features satisfies independent and identically Gaussian distribution. In the following, we prove that PCA features described above satisfies this assumption. We reformulated Eq. (5.17) as:
\[ \bar{x} = \bar{y} + \bar{n} \] (5.20)

Since \( \bar{y} \) and \( \bar{n} \) are independent for fully developed speckle noise, we can get:

\[ \Sigma_{\bar{x}} = \Sigma_{\bar{y}} + \sigma_{\bar{n}} I_p \] (5.21)

where \( \Sigma_{\bar{x}} \) and \( \Sigma_{\bar{y}} \) denotes respectively the covariance matrix of \( \bar{x} \) and \( \bar{y} \). The PCA analysis can be achieved by performing SVD on \( \Sigma_{\bar{y}} \):

\[ \Sigma_{\bar{y}} = \bar{A}\bar{\Delta}\bar{A}^T \] (5.22)

where the column vectors in \( \bar{A} \) represent the PCA bases, \( \bar{\Delta} = \text{diag}(\delta_1, \ldots, \delta_p) \) is eigenvalue matrix.

Then, we have:

\[ \Sigma_{\bar{x}} = A\bar{\Delta}\bar{A}^T + \sigma_{\bar{n}} A\bar{\Delta}\bar{A}^T = \bar{A} \begin{bmatrix} \delta_1 + \sigma_{\bar{n}} & \cdots & 0 \\ \vdots & \ddots & \vdots \\ 0 & \cdots & \delta_p + \sigma_{\bar{n}} \end{bmatrix} \bar{A}^T \] (5.23)

So we can see that \( \Sigma_{\bar{y}} \) and \( \Sigma_{\bar{x}} \) share the same PCA bases. As in Eq. (5.10), the PCA features can be obtained by projecting image patch onto PCA bases:

\[ \bar{s} = \bar{A}^T \bar{x} = \bar{A}^T \bar{y} + \bar{A}^T \bar{n} = \bar{s}_y + \bar{s}_n \] (5.24)

where \( \bar{s}_y = \bar{A}^T \bar{y} \) and \( \bar{s}_n = \bar{A}^T \bar{n} \) stand respectively for the signal and noise parts in PCA feature. Denote the variance matrix of \( \bar{s} \) by \( \Sigma_{\bar{s}} \):

\[ \Sigma_{\bar{s}} = \Sigma_{\bar{s}_y} + \Sigma_{\bar{s}_n} = \begin{bmatrix} \delta_1 & \cdots & 0 \\ \vdots & \ddots & \vdots \\ 0 & \cdots & \delta_p \end{bmatrix} + \begin{bmatrix} \sigma_{\bar{n}} & \cdots & 0 \\ \vdots & \ddots & \vdots \\ 0 & \cdots & \sigma_{\bar{n}} \end{bmatrix} \] (5.25)

Since \( \Sigma_{\bar{s}_n} = \sigma_{\bar{n}} I_p \), the assumption of K-means algorithm on noise distribution can be well satisfied. Although this property could not guarantee the convergence of K-means algorithm to global minimum, it provides theoretical assurance that K-means performance can be optimized.

### 5.4.3 Parameters tuning and efficient realization

**Number of clusters**: The number of clusters \( T \) in the image depends on the definition of what constitutes a cluster. This issue is mostly application-oriented, e.g. for background subtraction, background and foreground should be represented as two clusters, but in content-based image analysis, the number of cluster is mainly determined by the number of objects in the image. Here we
have no high-level requirements on the notion of cluster, but only loosely constraint that a cluster is a collection of image patches with similar spatial structure. Thus the number of clusters cannot be and does not need to be estimated very accurate. A loose cluster can be splitted into several compact ones, which does not have too much influence on the denoising results. Nevertheless, the rough estimation of the number of patterns exist in the image is still important because overly underestimation would reduce the sparsity in PCA domain and the opposite would increase computation burden and also preserve unnecessary artifact. So the number of cluster can be better determined based on the complexity of scene. More complex image should be assigned more clusters to fully capture image details. Since PCA detects statistically uncorrelated sources, a more complex image scene tends to have larger number of signal PCs. So we use the number of signal PCs $K_{MDL}$ in Section 5.4.1 as estimate of the number of clusters. Moreover, to prevent over-segmentation, we set an upper limit on $T$. In this study, we require:

$$T = \max(K_{MDL}, 15)$$ (5.26)

**Size of cluster:** The number of patches in each cluster should be big enough for efficient estimation of the covariance matrix $\Sigma_X$. In this study, we constrain that each cluster should have at least 50 members. A cluster smaller than this value will be deleted and its members will be dispersed into the other clusters based on the proximity in Euclidean space.

**Initial cluster centers:** K-means clustering is very sensitive to initial parameter values. Poor assignment of initial parameters may cause longer time to converge. Because the PCs provide a contiguous membership indicator for K-means clustering (Ding & He, 2004), we estimate the labels of image pixels by sorting the values of the first PC and then splitting them evenly into $T$ groups. Given the initial labels, we estimate the centroids for each group.

**Deal with large image:** SAR images always have big size. Clustering on them directly tends to ignore weak patterns that involve small number of pixels. Hence some image details would be erased during the denoising stage. So in this study, a large image is divided into several sub-images which are denoised separately. The final noise-free image is reconstructed by all the denoised sub-images. There are no universal standard for the size of the sub-image. It should be adjusted according to the complexity of the image scene. Small size should be preferred for image with complex scene. For SAR image, a size of 64×64 pixels can achieve a good result based on our experiments. To avoid the boundary artifacts between neighboring sub-images, we design the neighboring sub-images to be
slightly overlapping. Based on our experiment, an overlapping of 5 pixels would produce smooth boundaries.

**Size of patch:** As discussed in Section 5.3, patch with big size can capture large scale patterns but would also increase the computation cost. For SAR images that are without strong texture patterns, the size of 3×3 pixel is sufficient according to our experiments. But larger patch size, such as 5×5 is required for heavily textured images.

### 5.5 Complete Procedure of the Proposed Approach

The complete procedure involves two stages (Figure 5.2): the first stage produces a denoised image which is referenced as a clean image in the second stage to feed the clustering algorithm and to aid the LMMSE shrinkage. The detailed procedure is given below:

Stage 1: The original SAR image is split into \( N \) sub-images which are \( M \times M \) sized (\( M=64 \)). For each sub-image, we repeat the steps of clustering and denoising, until all sub-images have been processed. Finally, we aggregate the denoised patches to produce the denoised SAR image.

Clustering: This step intends to identify image pixels whose neighborhoods have similar spatial structure. The \( i \)th sub-image is firstly log-transformed. Then we extract all the \( S \times S \) sized patches (\( S=5 \)) to form a data matrix which is then transformed into the PCA domain. The first \( K_{MDL} \) PCs are used to feed the K-means algorithm, where \( K_{MDL} \) is given in Section 5.4.1. The number of cluster is determined by Eq. (5.26). Other parameters, i.e. the size of cluster and the initial cluster centers are given in Section 5.4.3. The products of this step are the labels of all pixels in the \( i \)th sub-image.

Denoising: Given the labels, this step aims to denoise the \( i \)th sub-image. Image patches in each cluster are denoised separately. Note that the image patches are extracted from the original SAR image. For each cluster, patches of pixels belong to this cluster are extracted to form a data matrix. We calculate the empirical mean \( \bar{X} \) and variance matrix \( \Sigma_x \). Then the patches are denoised by the following operations: obtaining PCA bases (Eq. (5.9)), projecting onto PCA bases (Eq. (5.10)), shrinking PCs in PCA domain (Eq. (5.11), (5.15)), and transforming back into patch domain (Eq. (5.16)). This step does not stop until all the clusters in the \( i \)th sub-image have been denoised. The final product of this step is a collection of denoised patches in the \( i \)th sub-image.
Aggregation: In this step, the denoised patches are used to construct the noise-free image. Because the patches are overlapping, so each pixel in the image has many denoised values. The final value is estimated as their average. The final product of this step is a denoised image.

Stage 2: This stage goes through the same operations as stage 1, except that we use the denoised image in stage 1 to feed the clustering step and to aid the LMMSE shrinkage in the denoising step.

Figure 5.2: The flowchart of the proposed algorithm (left part: stage 1, right part: stage 2).
Clustering: Instead of the original SAR image, the denoised image produced in stage 1 is used for clustering to get the labels. Moreover, the log-transformation is avoided. Other operations are the same with stage 1.

Denoising: The denoising procedures on this stage are the same with stage 1, except that we use the denoised image in stage 1 to estimate $\Sigma_{sy}$ in Eq. (5.11). Given labels, we extract two sets of patches for each cluster. One set is from the original SAR image. This set is to be denoised. Another set is from the denoised image produced by stage 1. This set is treated as a collection of signal patches. Hence the covariance matrix of signal patches $\hat{\Sigma}_y$ can be estimated as the sample covariance matrix. The shrinkage of the first set of patches in PCA domain requires the estimation of $\Sigma_{sy}$. Here, instead of using Eq. (5.15), we use:

$$\hat{\Sigma}_{sy} = \Lambda^T \hat{\Sigma}_y \Lambda$$  \hspace{1cm} (27)

Aggregation: The denoised patches are used to estimate the final noise-free image. The aggregation procedures are the same with stage 1.

Stage 2 is basically a repetition of stage 1, except that we used the denoised image in stage 1 to perform clustering and to estimate $\Sigma_{sy}$ in Eq. (5.11). These modifications are motivated by the fact that the first stage can significantly suppress SAR speckle noise and achieve a cleaner image. Hence using the denoised image, instead of the noisy image can achieve more accurate clustering results. Moreover, treating the denoised image as clean image to estimate $\Sigma_{sy}$ is more efficient than performing shrinking on the noisy image. A second stage in the BM3D algorithm (Dabov et al., 2007) was motivated by similar considerations.

5.6 Results and Discussion

In this study, both simulated and real SAR images are used to test the proposed SAR denoising method. In order to achieve a quantitative evaluation, clean images are degraded by adding multiplicative noise. Thus we can treat the clean image as the true values and use numerical measures to evaluate the performance. Although the true values of real SAR images are unknown, we can achieve qualitative assessments based on visual interpretation. In this experiment, three other methods (i.e. PPB (Deledalle et al., 2009), LPG-PCA (Zhang et al., 2010) and SAR-BM3D algorithm (Parrilli et al., 2012)) are selected to compare with the proposed method. The selection of these methods is based on the considerations of both the availability of the codes and their relevance to our work. The
LPG-PCA represents the state-of-the-art denoising techniques for images with additive noise, while PPB and SAR-BM3D are the state-of-the-art methods for SAR image. Because LPG-PCA was designed to deal with additive noise, to adapt it to SAR image, we transform the speckle noise into additive noise by logarithmic operation before performing it on noisy SAR images. The biased means caused by log-transformation are also corrected. In all experiments, without explicit indication, the parameters of the above algorithms are set as suggested in the referenced papers, and our method is implemented by setting the patch size to be 5×5 pixels and the sub-images to be 64×64 pixels with 5 pixels overlapping with their neighbors. All the other parameters in our method are determined by the methods in Section 5.4.3.

5.6.1 Test with Simulated Images

A variety of image sources are considered in this experiment, including the benchmark test image, i.e. the Barbara (Figure 5.3a) in the image denoising literature, and the high-resolution optical satellite (i.e. IKONOS) image whose scene structure is similar to real SAR image (Figure 5.3b). An ideal SAR denoising method is required to be capable of removing speckle effectively, while in the meantime preserving image details (e.g. texture, edge and line target) that constitute the desired features for further analysis. So in order to fully examine the abilities of details preservation, an image comprising two texture parts with a smooth boundary is designed to be used in this experiment. As shown in Figure 5.3(c), the left part of the image is weakly textured with a wave-like appearance while the right part is with a strong mesh texture. Thus the performance of denoising methods on image with changing scene complexities can be investigated. Simulated SAR image are obtained by multiplying speckle noise with these clean images. In this experiment, we use speckle noise in amplitude format which satisfies a squared-root Gamma distribution (Xie et al., 2002). All images are degraded with four different levels of speckle noise, i.e. the ENL (denoted by L) equals to 1, 2, 4 and 16, respectively. To avoid randomness, 20 noisy images for each clean image are produced by multiplying different noise realizations. All noisy images are processed and the numerical evaluation is based on the average of the results.

In this study, two statistics (i.e. Signal-to-Mean-Square-Error Ratio (S/MSE) and β) are used to evaluate these denoising methods. S/MSE corresponding to SNR in case of additive noise is a very effective measure of noise suppression in multiplicative case (Gagnon & Jouan, 1997). On the other hand, to measure image detail preservation we employ β originally defined in Sattar & Floreby (1997). β should be equal to unity for an ideal detail preservation.
The zoom of denoised images by different methods when $L = 1$ are shown in Figure 5.4-5.6, and the values of the two statistics for $L = 1, 2, 4$ and 16 are summarized in Table 5.1, in which the best value in each unit is bold. Overall, it shows in Table 5.1 that the proposed method outperformed the other referenced algorithms in terms of both measures. This demonstrates that our method is good both at speckle noise suppression and image detail preservation. The row ‘prop.stage1’ in Table 5.1 was achieved by the first stage of our method. Comparing with row ‘prop.’, we can see that the second stage involved in our method can significantly improve the results. The row ‘prop.global’ in Table 5.1 was achieved by performing PCA denoising on the sub-images without the clustering step. The observation that values in ‘prop.global’ are lower than values in ‘prop.’ justifies the clustering approach in the proposed method.

Both LPG-PCA and the proposed method denoise SAR images in PCA domain. However, LPG-PCA works on AWGN obtained by performing log-transformation on SAR image, while the proposed method takes into account the multiplicative nature of speckle noise by building the denoising approach on ASDN model. The observation that the proposed method greatly outperformed LPG-PCA on most noise levels justifies the proposed denoising model for ASDN. We also observed that the performance of LPG-PCA is very sensitive to noise levels variation in logarithmic space. As we can see in Table 5.1, when $L=1$, LPG-PCA achieved lower statistics than the proposed method. But with the increasing of $L$, LPG-PCA tends to achieve comparable results with our method in terms of $\beta$. LPG-PCA even achieved higher $\beta$ on image Barbara when $L = 4$ and 16. This is reasonable because LPG-PCA was designed for AWGN. When $L$ is big, the speckle noise subject to logarithmic operation is very close to Gaussian white noise. Therefore, the method can achieve good results. However, when $L$ is small, speckle noise begins to deviate from Gaussian distribution and its mean value is no more zero. This discrepancy between empirical data and the model assumption may reduce the efficiency of LPG-PCA. As we can see, the images in Figs. 5.4-5.6 denoised by LPG-PCA show many small artifacts, while images by the proposed method have little artifacts but plenty image details.

It is noticed that our method was especially better at denoising the synthesized texture image (Figure 5.3c). In Figure 5.6, the image produced by the proposed method is the most similar to clean image. The images denoised by PPB and SAR-BM3D are blurred and the holes in boundary area are erased. The image produced by LPG-PCA has clear textural patterns, but assumes many artifacts. The statistics in Table 5.1 indicates consistent results. The proposed method achieved high values in both
measures, indicating good performance on both noise removal and detail preservation. In contrast, LPG-PCA achieved small S/MSE values, while PPB and SAR-BM3D achieved small β values. The clustering approach involved in the proposed method may have contributed to the superiority of the proposed method in dealing with textual images. In texture image, the increased scene complexity renders it difficult to find similar patches. Given the difficulty, the clustering approach might find more relevant patches than block-matching approach, leading to better preservation of texture patterns. On less-textured image, i.e. IKONOS, the proposed approach also achieved higher β values, and preserved more image details than the other methods. The observation that LPG-PCA and the proposed method outperformed SAR-BM3D in terms of detail preservation in highly textured image (i.e. Figure 5.6) may suggest that the PCA-based denoising approach is more efficient at dealing textural structures.

A good denosing method should be capable of removing speckle noise without sacrificing image details. PPB tended to erase image details too much. In Figs. 5.4-5.6, we see that the denoised images by PPB have very smooth appearances but also blurred boundaries and reduced details information. On the IKONOS image, LPG-PCA achieved higher S/MSE but lower β values than SAR-BM3D, while on the Syntexture image, LPG-PCA achieved lower S/MSE but higher β values. Our method achieved very high S/MSE and β values on most images.

The SAR-BM3D and PPB algorithms were implemented using the C language, while the other algorithms were implemented under the MATLAB platform. All the computations were conducted on a PC with a Pentium(R) 2.30GHZ Quad-Core processor. On average, it took 36.8, 53.1, 34.7 and 23.5 seconds, respectively by PPB, LPG-PCA, SAR-BM3D and the proposed method to process a 256×256 pixels simulated image. It is fair to compare the time efficiency of the proposed algorithm and LPG-PCA, because both methods are PCA-based and implemented in MATLAB language. The observation that the proposed algorithm consumes less than half of the time of LPG-PCA demonstrates the efficiency of clustering algorithm than block-matching approach. This conclusion is also supported by the shorter processing time of our algorithm than PPB and SAR-BM3D, especially considering the fact that C language is more efficient than MATLAB.
Figure 5.3: Clean images used in this study, (a) Barbara, (b) Optical satellite image (IKONOS), (c) Synthesized texture image. All images are 256×256 pixels big.

Table 5.1: Results (s/mse and $\beta$) on three images with different noise levels

<table>
<thead>
<tr>
<th>Noise Level</th>
<th>IKONOS</th>
<th>Barbara</th>
<th>Syntexture</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$L=1$</td>
<td>$L=2$</td>
<td>$L=4$</td>
</tr>
<tr>
<td>PPB</td>
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<td>16.56</td>
<td>17.97</td>
</tr>
<tr>
<td>Prop.</td>
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</tr>
<tr>
<td>$\beta$</td>
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<td>0.246</td>
<td>0.333</td>
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<td></td>
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Figure 5.4: Zoom of Barbara image degraded by single look speckle noise, (a) clean image, (b) noisy image, (c) PPB, (d) LPG-PCA, (e) SAR-BM3D, (f) the proposed method.

Figure 5.5: Zoom of IKONOS image degraded by single look speckle noise, (a) clean image, (b) noisy image, (c) PPB, (d) LPG-PCA, (e) SAR-BM3D, (f) the proposed method.
5.6.2 Test with real SAR images

The real SAR images used for testing different denoising methods are two TerraSAR-X sample imageries provided on the Astrium GeoInformation Services website. Both images are located at Canada Toronto, taken in December 2007 under the spotlight mode with 1m spatial resolution and incidence angle of 48.8°. But one is the Single Look Slant Range Complex (SSC) image, while the other one is the Spatially Enhanced (SE) Multi-look Ground-range Detected (MGD) with L=2. From the SSC image, we obtain two smaller images, and from the MGD image, we obtain one. The three images that comprise parking lots, roads and buildings are supposed to capture the major types of urban targets.

In this experiment, in addition to the denoising algorithms in the simulated study, we also tested the SRAD method in Yu & Acton (2002). We adopted the default patch size parameters for the referenced methods but a smaller size of 3×3 for the proposed method, because they experimentally allowed the respective best trade-offs between noise removal and detail preservation. The zooms of
these images denoised by different techniques are shown in Figs. 5.7-5.9. The results are basically consistent with the simulated study. The proposed method not only greatly suppressed speckle noise, e.g. all three denoised image are very smooth, but also preserved image details very well, e.g. the eight bright spots in Figure 5.7 were kept very well and the roads in Figure 5.8 were delineated very clearly. The SAR-BM3D also achieved good balance between noise removal and detail preservation. The PPB method achieved very clean images, but some image details were also smoothed out. The LPG-PCA, because it was not specifically designed for SAR speckle noise, produced many dark artifacts in Figs. 5.7 and 5.8 where noise level is high, but achieved smoother results in Figure 5.9 where less noise exists. Generally speaking, SRAD preserved point targets very well, but also produced undesirable artifacts.

![Images of denoised images](image)

Figure 5.7: Zoom of TerraSAR-X SSC image (112x95 pixels) of the parking lot located at the NE of the Macdonald-Cartier Freeway/Allen Road interchange, Toronto, Canada, with L=1, (a) SRAD, (b) original image, (c) PPB, (d) LPG-PCA, (e) SAR-BM3D, (f) the proposed method.
Figure 5.8: Zoom of TerraSAR-X SSC image (126×116 pixels) of the roads located at the SE of the Macdonald-Cartier Freeway/Allen Road interchange, Toronto, Canada, with L=1, (a) SRAD, (b) original image, (c) PPB, (d) LPG-PCA, (e) SAR-BM3D, (f) the proposed method.
5.7 Conclusion

In this chapter, we have presented a SAR images denoising scheme based on clustering the noisy image into disjoint local regions and denoising each region by LMMSE filtering in PCA domain. In the clustering step, in order to reduce dimensionality and resist the influence of noise, we identified several leading PCs in logarithmic domain by MDL criterion to feed the K-means algorithm. This clustering approach can be treated as the unsupervised counterpart of the commonly adopted block-matching approach. It requires less computation. Moreover, it is capable of adaptively identifying “similar” patches by considering the closeness to different clusters centers. In the denoising stage, in order to avoid the limitations of the homomorphic approach, we built our denoising scheme on ASDN and derived a PCA-based LMMSE denoising model for multiplicative noise. Our approach is the first to build the PCA-based denoising method on the ASDN model for SAR image denoising. Besides SAR images, it is also applicable to other signal-dependent noise. The denoised patches of all clusters were finally used to reconstruct the noise-free image. We tested our denoising scheme in both real and simulated SAR image with several other state-of-the-art methods. The results suggested that our method compared favorably w.r.t. the referenced methods in terms of both image detail preservation and speckle noise reduction.
Chapter 6
Conclusions and Recommendations

6.1 Summary and Contribution

This thesis systematically studies MLVM for modeling and analysis of remote sensing data. First of all, a framework is introduced to provide guidelines for building MLVMs, from a comparative perspective with mixture models and LVMs. The essence of MLVMs is to model simultaneously two key factors in remote sensing data generation, i.e. the source heterogeneity effect and the hidden signals effect, with mixture components discriminating different spatial processes, and latent dimensions addressing hidden signals in each component. The optimization scheme based on EM algorithm is described, and specifications of the framework for building task-dependent MLVM variants are discussed.

Secondly, a MPSR model is proposed for supervised hyperspectral classification, considering the gap that while SR is an emerging and powerful technique for hyperspectral feature extraction, there is still a lack of a probabilistic mixture model for it. In order to account for the spatial correlation effect in labels, a MRF label prior is adopted to be combined with MPSR in a Bayesian framework. The image classification based on MPSR model is achieved by solving a MAP problem in E-step of EM iteration. The capabilities of the proposed method are proven in several benchmark hyperspectral images of both agricultural and urban areas, by comparing with the advanced hyperspectral image classification method. Besides hyperspectral image classification, MPSR may also be applicable to other spatial classification issue involving high-dimensional features.

Thirdly, a K-P-Means model is proposed for addressing spectral unmixing issue. K-P-Means aims to learn nonnegative latent variables in nonnegative latent space, in order to account for the particularities of remote sensing spectral observations. By imposing label information on latent variables, K-P-Means model offers a novel interpretation of spectral unmixing. The optimization of K-P-Means model is achieved by alternating the E- and M-step until convergence. Experiments using both simulated and real spectral images show that the proposed K-P-Means model provides strong endmember and abundance estimation results compared to existing approaches. Besides its application in remote sensing, K-P-Means may also feasible for addressing other nonnegative data factorization issues or clustering issues in other fields.
Lastly, a clustering-based PCA model is designed for advanced SAR image denoising. In this model, the noise satisfies a Gamma distribution, in order to account for the particularities of SAR speckle noise. The optimization of this model involves two EM iterations. When learning labels in E-step, to reduce dimensionality and resist the influence of noise, several leading PCs in logarithmic space, identified by the MDL criterion are used to feed the K-means clustering algorithm. In M-step, after learning PCA for each cluster, denoising is achieved by performing LMMSE in PCA domain based on ASDN model. The experiments demonstrate that the proposed algorithm achieved better performance than the referenced state-of-the-art methods in terms of both noise reduction and image details preservation. Besides its application in SAR image denoising, the clustering-based PCA model may also provide a strong image representation scheme.

The main findings of this thesis are summarized into the following aspects:

(1) MLVM benefits remote sensing data processing, in terms of providing a sound data modeling framework, which is capable of capture key characteristics of remote sensing data, i.e. the underlying spatial processes and hidden signals. The theoretical benefits of MLVM are proved by the successfully application of three MLVMs in addressing different aspects of remote sensing data processing tasks.

(2) The MLVM is capable of achieving better remote sensing image classification than mixture models, considering that representing classes by discriminative signals in latent space, is more effective than characterizing classes by a certain parametric probabilistic distribution in original space. This conclusion is well justified by the higher classification accuracy achieved by MPSR in hyperspectral image classification, than an advanced method called MLRsubMLL, which has been demonstrated higher accuracy than mixture models such as QDA and LDA (Li et al., 2012).

(3) The MLVM is capable of addressing efficiently the spectral source separation issue, which is of fundamental importance in remote sensing data analysis, by simply adopting the idea of associating each hidden variable with a mixture component. This capability is well supported by the simplicity, higher accuracy and lower computational cost achieved by K-P-Means in spectral unmixing.

(4) As far as learning latent variables for image representation is concerned, MLVM also constitutes powerful technique, due to its ability to capture local structures in feature space. For example, in representational tasks, i.e. denoising, clustering-based PCA, which follows the key ideas of MLVM, achieves the state-of-the-art SAR denoising performance, mainly due to the fact that local
PCA learnt for individual classes is more capable of achieving noise-discriminative latent variable, than global PCA learnt for all classes.

In summary, the contributions of this thesis include:

(1) Introduce a probabilistic framework for modeling and estimating both source heterogeneity effect and hidden signals in remote sensing data. Under this framework, three MLVMs are successfully developed and applied to different remote sensing applications.

(2) Introduce a novel mixture of probabilistic sparse representation (MPSR) model and successfully integrate it with MRF label prior in Bayesian framework for hyperspectral image classification.

(3) Introduce the concept of “K purified means”, based on which a novel model called K-P-Means is developed for spectral endmember extraction.

(4) Introduce a clustering-based PCA model for achieving state-of-the-art SAR images denoising method.

(5) Demonstrate in experiments the effectiveness of the proposed models when comparing with the state-of-the-art methods in their respective applications.

(6) The proposed models are not limited to the demonstrated applications, but can possibly applied to applications in other disciplines.

6.2 Recommendations for Future Research

6.2.1 Incorporating Label Prior

In remote sensing data, the spatial correlation effect is substantial. Pixels that are spatially close to each other tend to assume similar labels. Therefore, it is very important to account for this effect in model building and optimization stage. Nevertheless, of the three proposed models, only MPSR adopts MRF label prior, in order to remove undesirable artifacts and achieve semantically meaningful classification results. The reason for not employing label prior in K-P-Means and clustering-based PCA is mainly because they are designed for learning representative latent variables, instead of for learning semantically meaningful labels. Nevertheless, it is still worthwhile to explore the outcome of the two models if the label prior was incorporated, considering that the spatial correlation effect may have an indirect influence on the latent variable learning.
6.2.2 Estimating Hyperparameters

Since MLVM involves both mixture model and LVM, it assumes more hyperparameters than both sub-models. The hyperparameters in MLVM include the number of classes, the number of latent bases, the number of active latent variables (e.g. the sparsity in SR) and the structure of the covariance matrix of noise. In supervised classification or latent variable learning, these hyperparameters determines directly the model generalization capability. The tuning of hyperparameters in supervised scenario can be achieved based on posterior information, by cross-validation techniques (Hastie et al., 2001). In unsupervised cases, although models are not designed for prediction, it is still crucial to estimate the hyperparameters, in order to achieve meaningful pattern discovery. The adjusting of hyperparameters can be achieved in Bayesian framework, by assuming prior distributions for hyperparameters. The resulting model can be solved by certain optimization techniques, such as variational Bayesian methods (Attias, 1999).

6.2.3 Unsupervised MPSR for Clustering and Latent Variable Learning

In this thesis, although MPSR is used for supervised spectral classification, it also has potential to be used for unsupervised classification and latent variable learning. Therefore, it is meaningful to explore the performance of MPSR on spectral clustering, by adopting some dictionary learning techniques, such as K-SVD, to learn dictionaries for different classes in M-step of EM iterations. It is also of great interests of to investigate on the capability of MPSR to learn representative latent signals for some other tasks, e.g. denoising. Note that clustering and latent variable learning can be achieved simultaneously by solving an unsupervised MPSR model, in which the labels and latent variables will be estimated iteratively by alternating the E- and M-step until convergence.

6.2.4 K-P-means for clustering

Although K-P-Means is mainly used in this thesis for learning latent variables, i.e. the endmembers, it is innately a clustering algorithm, where a cluster is represented by an endmember that admits dominant abundances on the associated pixel in that cluster. Although K-P-Means is capable of providing a map of dominant endmembers, it could not provide maps of different land cover types, because a land cover type always involves multiple endmembers. Therefore, in order to adapt K-P-Means for spectral image segmentation, the concept of cluster in K-P-Means has to be modified to allow the representation of a class by multiple endmembers, as opposed to by just one. Another necessary modification is to prescribe the class membership of endmembers. Accordingly, in E-step,
the representation error over different groups of endmembers will be used to infer the label of spectral pixel; in M-step, some techniques that allow the learning of multiple endmembers, such as VCA, can be used to update the endmembers for each class based on the label information of pixels.

6.2.5 K-P-Means for non-negative matrix factorization

By learning nonnegative latent variables in nonnegative latent space, K-P-Means amount to a nonnegative matrix factorization techniques, where a data matrix is decomposed into two nonnegative matrices, in order to learn meaningful patterns. Therefore, it is meaningful to explore the feasibility of using K-P-Means for addressing some nonnegative data representation issue, in a comparative perspective with typical nonnegative matrix factorization techniques.
References


Appendix A

List of Publications during PhD Thesis Work


Appendix B
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