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Hyperspectral data classification improved by minimum spanning forests

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Abstract. Remote sensing technology has applications in various knowledge domains, such as agriculture, meteorology, land use, environmental monitoring, military surveillance, and mineral exploration. The increasing advances in image acquisition techniques have allowed the generation of large volumes of data at high spectral resolution with several spectral bands representing images collected simultaneously. We propose and evaluate a supervised classification method composed of three stages. Initially, hyperspectral values and entropy information are employed by support vector machines to produce an initial classification. Then, the *K*-nearest neighbor technique searches for pixels with high probability of being correctly classified. Finally, minimum spanning forests are applied to these pixels to reclassify the image taking spatial restrictions into consideration. Experiments on several hyperspectral images are conducted to show the effectiveness of the proposed method. © 2016 Society of Photo-Optical Instrumentation Engineers (SPIE) [DOI: 10.1117/1.JRS.10.025007]

Keywords: hyperspectral images; supervised classification; minimum spanning forests; image analysis.

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1 Introduction

The technology of remote sensing^{1–4} has applications in several fields, including agriculture, forestry, hydrology, meteorology, mapping, environmental monitoring, land use, mineral exploration, and military surveillance.

Airborne and satellite sensors are typically used to measure the electromagnetic radiation reflected from the Earth's surface. Different types of surface materials, such as soil, water, vegetation, and minerals, can be distinguished from spectral reflectance signatures.

Advances in image acquisition techniques have allowed the extraction of large volumes of data with high spectral resolution. Such data, called hyperspectral, refer to the collection of information consisting of various spectral bands, which are defined in a wavelength range.

Hyperspectral image analysis^{5–11} is a challenging task and has received increasing interest in the community. Some inherent difficulties in the analysis of such data include its high dimensionality and noise contamination.

Although the spectral bands provide valuable information with high discriminative power that allows the identification of distinct classes of objects, some initiatives for combination of spatial and spectral information have been also presented in the literature.^{12–15}

This work investigates the use of spectral and spatial information to classify hyperspectral data. A three-stage supervised classifier is employed to determine the class for each pixel. An initial classification is performed by support vector machines (SVMs), whose result is used as input to K-nearest neighbor (K-NN) that searches for correctly classified pixels. Such pixels are used as seeds for a minimum spanning forests (MSF) algorithm to generate a final spatial-based classification. The proposed method is applied to a set of hyperspectral images, and its

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performance is compared to other classification approaches, including two which are also based on MSF.^{13,15}

The remainder of the paper is organized as follows. Section 2 briefly describes concepts related to the topic under investigation. The proposed method for hyperspectral data is presented in Sec. 3. Experimental results are described and discussed in Sec. 4. Section 5 presents the conclusions and directions for future work.

2 Background

The remote sensing area^{1–3,16} has employed aircraft and satellites to acquire information about the Earth's surface. In addition to traditional aerial photographs operating in the visible portion of the electromagnetic spectrum, continuous technological development has allowed the acquisition of information in other portions, such as microwave and thermal infrared.

Hyperspectral sensors, also known as imaging spectrometers, are instruments that capture images at several contiguous bands of spectral radiance with wavelengths typically varying between 0.4 and 2.5 μ m. Therefore, each pixel in the image contains a continuous spectrum that can be used to characterize objects in the scene.

Several hyperspectral sensors have been used to acquire data for a number of different applications, such as thematic mapping, geology, hydrology, agricultural assessment, mineral exploration, and study of global environmental and climate changes. Some important sensors include the hyperspectral digital imagery collection experiment,¹⁷ airborne visible/infrared imaging spectrometer,¹⁸ reflective optics system imaging spectrometer,¹⁹ compact airborne spectrographic imager,²⁰ airborne imaging spectrometer,²¹ airborne imaging spectrometer for different applications,²² geophysical and environmental research imaging spectrometer,²³ hyperspectral mapper,²⁴ aerosol mass spectrometer,²⁵ airborne prism experiment,²⁶ compact high-resolution imaging spectrometer,²⁷ digital airborne imaging spectrometer,²⁸ and thermal infrared multispectral scanner.²⁹

The analysis of hyperspectral data has received growing interest from the community, since they have spectral attributes with high discriminative power. Several classification methods have been proposed in the literature for labeling the image pixels according to specific classes, including Bayesian networks,³⁰ SVM,³¹ neural networks,³² nearest neighbors,³³ clustering algorithms,³⁴ and decision trees,³⁵ among others.

The process of classification consists in categorizing pixels or regions of the images into a set of different classes, such that pixels with similar attributes belong to the same class. Hyperspectral image classification methods can be generally divided into unsupervised and supervised techniques. Unsupervised techniques identify groups of pixels without prior knowledge of the data, whereas supervised techniques employ known labeled samples to assign pixels to one of the classes.

Many pixelwise classification approaches^{36,37} consider only spectral information of the pixels. Other methods take spatial information present in the image into account.^{12,38–40} Some strategies for including spatial information in the classification are watershed,⁴¹ minimum spanning forests,¹³ graph cuts,⁴² shortest paths,⁴³ random walks,⁴⁴ and Markov random fields.⁴⁵

The image analysis process typically uses a more succinct representation of the data or its components (objects) called the feature vector, which stores the most representative attributes of the regions of the image. The main features extracted from images to assist the analysis process are the color, texture, and shape of objects. The number of attributes or characteristics determines the dimension of the feature vector and usually depends on the application area and the properties that need to be discriminated.

Due to the high dimensionality⁴⁶ of the feature vector that represents the spectral information contained in hyperspectral images, feature selection techniques⁴⁷ are usually applied as preprocessing to the data classification. A widely used technique for data compression and dimensionality reduction is the principal component analysis (PCA).⁴⁸

3 Methodology

A general scheme for the proposed classification method is shown in Fig. 1. Given a set of hyperspectral bands and a set of classified samples (training data), the classification process

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Fig. 1 Overview of the proposed classification method. The first step produces an initial classification through SVM. The second step compares, for each pixel, the classification assigned by SVM and the classes of K-NN. If the classes agree, the pixel is selected as a seed. The final step uses the seeds in an MSF algorithm to produce a spatial classification.

of the hyperspectral image is initially obtained by using SVM. In a second step, a K-NN search is performed in order to choose the pixels that are most likely to be correctly classified by SVM. Such pixels are chosen as seeds for the MSF algorithm to produce a spatial-based classification. The details on each step are described as follows.

The hyperspectral image is a matrix of size $m \times n \times l$, so that each pixel is a vector $\mathbf{p}(x, y) = [p(x, y, 1), p(x, y, 2), \dots, p(x, y, l)]$. The set S_u of samples to be classified is the set of $m \times n$ pixels. For simplicity, a general pixel will be referred to as $\mathbf{p} = (p_1, p_2, \dots, p_l)$, dropping the two-dimensional indices. The classified samples form a set S_c of pairs $s_i = (\mathbf{x}_i, y_i)$ and $i = \{1, 2, \dots, j\}$, such that the *i*'th pair has feature vector $\mathbf{x}_i \in \mathbb{R}^p$ and belongs to class y_i .

Two measures are investigated as descriptors for the elements of both S_u and S_c . The first corresponds to the hyperspectral values. In such a case, the descriptor of a pixel **p** is the pixel itself, the vector $\mathbf{p} = (p_1, p_2, \dots, p_l)$. The second measure is the local entropy, which measures the intensity variation around a pixel. This information can be useful in distinguishing homogeneous regions (smaller values) from heterogeneous regions (larger values). The feature vector for a pixel **p** is a vector $\mathbf{e} = [e(p_1), e(p_2), \dots, e(p_l)]$, such that $e(p_i)$ is the entropy of p_i computed over a 9×9 neighborhood centered at p_i .

The hyperspectral and entropy descriptors for a sample $s_i \in S_c$ are given beforehand, so that the pair (\mathbf{p}_i, y_i) denotes s_i with hyperspectral descriptor and (\mathbf{e}_i, y_i) denotes s_i with entropy descriptor.

3.1 Support Vector Machines Classification

An SVM classifier receives as input the sets S_u and S_c . The PCA technique is applied to the entropy vectors **e** in order to obtain reduced vectors **e**'. This dimensionality reduction process is important to remove redundant or highly correlated features, which improves the performance of the classifier.

The SVM classifier is trained by using the pairs (\mathbf{e}'_i, y_i) of S_c . Given the trained model, SVM classifies the samples \mathbf{p} of S_u such that $L_1(\mathbf{p})$ is the assigned class. The assignment of all pixels produces an image L_1 of size $m \times n$.

3.2 Seed Identification

The SVM classification is checked against a K-NN search in order to identify pixels that have high probability of being correctly classified. The pixels identified in this step are called seeds. These pixels are considered in the next step as starting points for an MSF method which introduces spatial restrictions in the classification. All pixels that are not seeds will be reclassified by expanding the classification from the seeds. Algorithm 1 summarizes the process of seed identification.

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Input:
S_u set of pixels
S_c set of classified samples
L ₁ image of classes assigned by SVM
Output:
L ₂ image of seeds
C image of costs
1 for $\mathbf{p} \in S_u$ do
Find the pairs $\{(\mathbf{e}'_{c_1}, y_{c_1}), (\mathbf{e}'_{c_2}, y_{c_2}), \dots, (\mathbf{e}'_{c_k}, y_{c_k})\}$ from S_c which are the <i>k</i> nearest neighbors to the descriptor \mathbf{e}' of \mathbf{p}
3 if $L_1(\mathbf{p}) = y_{c_1} = y_{c_2} = \ldots = y_{c_k}$ then
4 Make $L_2(\mathbf{p})$ equal to $L_1(\mathbf{p})$
5 Make $C(\mathbf{p})$ equal to 0
6 else
7 Make $L_2(\mathbf{p})$ equal to -1
8 Make $C(\mathbf{p})$ equal to ∞
9 end
10 end

For each pixel $\mathbf{p} \in S_u$, a K-NN search is performed in the feature space to retrieve its k nearest neighbors among the set S_c . The entropy descriptors are also used in this step. Therefore, given the descriptor \mathbf{e}' of \mathbf{p} , the pairs of closest samples $\{(\mathbf{e}'_{c_1}, y_{c_1}), (\mathbf{e}'_{c_2}, y_{c_2}), \dots, (\mathbf{e}'_{c_k}, y_{c_k})\}$ from S_c are retrieved.

If the class of \mathbf{p} assigned by SVM is the same class of the pairs found with the K-NN search, $L_1(\mathbf{p}) = y_{c_1} = y_{c_2} = \ldots = y_{c_k}$, then \mathbf{p} is included as a seed pixel in the set L_2 , such that its class is the same as the one from SVM, $L_2(\mathbf{p}) = L_1(\mathbf{p})$. In addition to the class labeling, a cost of zero is also assigned to a seed, that is, $C(\mathbf{p}) = 0$. L_2 and C are images of size $m \times n$. These two images are further used in the MSF classification, such that $L_2(\mathbf{p})$ will store the classification of \mathbf{p} given by an MSF, whereas C will store the cost of such assignment.

All the pixels **p** for which the class assigned by SVM differs from any of the K-NN will be considered in the next step of the classification. These pixels are assigned an invalid class $L_2(\mathbf{p}) = -1$ and a high cost, $C(\mathbf{p}) = \infty$.

3.3 Minimum Spanning Forests Classification

The final step of the classification uses an MSF approach so that the classification takes spatial restrictions into account. This step performs its computations over the hyperspectral feature space; that is, the similarity of pixels is computed using the pixel vectors of values $\mathbf{p} = (p_1, p_2, \dots, p_l)$.

The classification is stored in the image L_2 . The cost image C and a minimum priority queue Q are used as auxiliary data structures. The seeds found in the previous step are initially inserted into the priority queue Q. The priority queue Q is maintained according to the costs of C. The MSF computation is performed as shown in Algorithm 2.

A minimum cost pixel **p** is retrieved from Q, and its four-neighborhood is analyzed. Suppose **q** is a neighbor pixel of **p** and $d(\mathbf{p}, \mathbf{q})$ is the Euclidean distance of their feature vectors

Algorithm 2 Minimum spanni	ng forests.
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Input:
S_u set of pixels
L ₂ image of classes
C image of costs
Q auxiliary minimum priority queue
Output:
L ₂ classified image
1 Insert into Q all $\mathbf{p} \in S_u$ such that $C(\mathbf{p}) = 0$
2 while Q is not empty do
3 Remove p from Q
4 for $\mathbf{q} \in S_u$ in the 4-neighborhood of \mathbf{p} do
5 if $C(\mathbf{q}) > d(\mathbf{p}, \mathbf{q})$ then
6 Make $L_2(\mathbf{q})$ equal to $L_2(\mathbf{p})$
7 Make $C(\mathbf{q})$ equal to $d(\mathbf{p}, \mathbf{q})$
8 if $q \in Q$ then
9 Update priority queue <i>Q</i>
10 else
11 Insert q into <i>Q</i>
12 end
13 end
14 end

 $\mathbf{p} = (p_1, p_2, \dots, p_l)$ and $\mathbf{q} = (q_1, q_2, \dots, q_l)$. If the cost $C(\mathbf{q})$ is greater than $d(\mathbf{p}, \mathbf{q})$, then \mathbf{q} is assigned to the class of \mathbf{p} ($L_2(\mathbf{q}) = L_2(\mathbf{p})$) and its cost is updated to the distance between the descriptors, $C(\mathbf{q}) = d(\mathbf{p}, \mathbf{q})$. In case \mathbf{q} is already in Q, the priority queue is updated; otherwise \mathbf{q} is inserted into Q.

The process just described is the basic iteration of the MSF algorithm. At each iteration, a pixel of minimum cost is retrieved from Q. Since the seeds are elements with cost zero, the classification will expand from them. The process is repeated until the priority queue is empty, such that all pixels have been classified.

4 Experimental Results

In this section, the proposed method is compared to four others: (i) SVM, (ii) SVM with PCA, (iii) the approach based on stochastic minimum spanning forest by Bernard et al.,¹³ referred here to as SMSF, and (iv) the approach based on probabilistic minimum spanning forest by Tarabalka et al.,¹⁵ referred here to as PMSF.

The SMSF and PMSF approaches are closely related to our proposed method, which are also based on the SVM classifier to obtain an initial classification and based on MSF for improving the results. On the other hand, the SMSF uses a stochastic scheme for the second step. A set of n maps is created and, for each one of these maps, a percentage of the pixels classified by SVM is randomly chosen to form the seeds for the MSF method. Then, MSF is applied to each map and the final classification is given by a voting scheme considering the results for all these maps.

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Dataset	Rows	Columns	Bands	Classes
Indian Pines	145	145	200	16
Pavia University	610	340	103	9
Pavia Centre	1096	715	102	9
Salinas-A	83	86	204	6
Salinas	512	217	204	16
Kennedy Space Center	512	614	176	13
Botswana	1476	256	145	14

Table 1 Summary of the main dataset characteristics.

The PMSF also differs in the second step, where the most reliable pixels classified by SVM are chosen as seeds for the MSF method.

Experiments are conducted on seven benchmark datasets.^{18,19,49} The main characteristics of each dataset (image dimensions, number of bands, and number of classes) are summarized in Table 1.

The algorithms were implemented in MATLAB[®].⁵⁰ The LIBSVM,⁵¹ with default parameters, was used to perform the classification based on SVM. The methods were evaluated through the overall and class accuracy rates.

Figure 2 shows an overview of the proposed classification method. Figures 2(a) and 2(b)depict one band of the Indian Pines dataset and its ground truth. The SVM classification is given in Fig. 2(c). The resulting seeds are shown in Fig. 2(d), whereas the final classification is shown in Fig. 2(e).

In the following sections, we initially tuned the parameters for SVM, our method, SMSF, and PMSF. Hyperspectral and entropy descriptors were compared. Entropy improved the results obtained with SVM when compared to the results through hyperspectral values. The same was not true for MSF classification, where the hyperspectral values tend to perform better than entropy. We presented some comparisons for SVM. For the MSF step, we used hyperspectral



(a) Indian Pines band



(b) Groundtruth



(c) SVM-PCA

(d) Seeds



Fig. 2 Example illustrating an overview of the proposed classification method.

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Fig. 3 Comparison of parameters for SVM classification. The lines are labeled as a pair defining which descriptor was used and whether PCA has been performed.

descriptors. The parameter k for the k-nearest neighbors used in our method was evaluated, as well as the parameters for SMSF and PMSF. After the tuning steps, we compared the methods.

4.1 Support Vector Machines and Descriptors

Figure 3 shows the behavior of the SVM classification according to the used descriptor, hyperspectral values, or entropy, as well as the use of PCA. Entropy provides a significant improvement on the classification results when compared to hyperspectral values, whereas the use of PCA helped to achieve better classification results relative to each descriptor.

4.2 K-Nearest Neighbor Setup

Figure 4 shows the impact on the classification according to the choice of the k value (of K-NN) for selecting the seeds in our method. The best results were obtained with the hyperspectral



Fig. 4 Impact of *k*-nearest neighbors for the selection of seed pixels. The lines are labeled as a pair defining which descriptor was used and whether PCA has been performed. The SVM result is depicted as a baseline for comparison.

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Fig. 5 Results obtained with the SMSF using different numbers of maps and percentages of pixels randomly selected as seeds. Each line reports the results for a certain percentage of seeds as the number of maps varies. The SVM result is depicted as a baseline for comparison.

descriptor. PCA did not provide any improvement. All the experiments were performed with 10% of samples for training the SVM. The SVM result is depicted as a baseline for comparison. The best result is achieved with k = 3.

4.3 SMSF Setup

The impact of the choice in terms of the number of maps and marks for the SMSF method is summarized in Fig. 5. All experiments used 10% of the pixels for the SVM classification. Even though the original paper¹³ does not use entropy and PCA for SVM, we have used these settings for a fair comparison of the MSF step; otherwise the results for SMSF would be degraded. Similarly to what occurred with our method, PCA and entropy do not improve the results of the MSF step. We suppress the results for such parameters and show the best results using



Fig. 6 Results obtained with the PMSF using different percentages of pixels selected as seeds and different reliability thresholds. Each line reports the results for a certain reliability threshold. The SVM result is depicted as a baseline for comparison.

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Class	SVM	SVM-PCA	SMSF	PMSF	SVM-MSF
Alfalfa	0.521	0.312	0.708	0.854	0.854
Corn-notill	0.798	0.827	0.869	0.891	0.850
Corn-min	0.781	0.817	0.884	0.893	0.905
Corn	0.657	0.738	0.752	0.786	0.862
Grass/pasture	0.792	0.828	0.846	0.906	0.937
Grass/trees	0.805	0.798	0.878	0.906	0.964
Pasture-mowed	0.739	0.000	0.000	0.000	0.000
Hay-windrowed	0.952	0.968	1.000	0.998	0.989
Oats	0.000	0.000	0.000	0.000	0.000
Soybeans-notill	0.889	0.907	0.946	0.929	0.938
Soybeans-min	0.943	0.954	0.977	0.965	0.978
Soybeans-clean	0.830	0.857	0.904	0.906	0.861
Wheat	0.753	0.858	1.000	0.926	0.984
Woods	0.932	0.930	0.967	0.942	0.972
Building-trees	0.711	0.737	0.851	0.713	0.661
Stone-steel	0.871	0.906	1.000	1.000	0.953
Overall	0.856	0.873	0.919	0.915	0.921

 Table 2
 Comparison for the Indian Pines dataset.

a feature descriptor based on hyperspectral values and no PCA step. This is also the setting used in the original paper. The best results were obtained with 20 maps (as in the original paper) and 10% of seeds randomly selected. In the original paper, the value is 3% for some datasets and 11% for other datasets. However, some of the datasets used here were not considered in the paper. Therefore, we choose 10% random samples for seed selection since this value achieves the best results for the datasets evaluated in our experiments.

Table 3 Comparison for the Pavia University data	aset
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Class	SVM	SVM-PCA	SMSF	PMSF	SVM-MSF
Asphalt	0.942	0.962	0.966	0.959	0.973
Meadows	0.971	0.978	0.984	0.973	0.989
Gravel	0.723	0.787	0.843	0.776	0.850
Trees	0.885	0.890	0.915	0.894	0.884
Painted metal sheets	0.973	0.988	0.999	0.999	0.995
Bare Soil	0.832	0.870	0.893	0.903	0.921
Bitumen	0.870	0.866	0.939	0.880	0.907
Self-blocking bricks	0.915	0.931	0.959	0.958	0.938
Shadows	0.842	0.864	0.836	0.845	0.887
Overall	0.921	0.938	0.953	0.941	0.955

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Class	SVM	SVM-PCA	SMSF	PMSF	SVM-MSF
Water	0.999	0.999	0.999	0.999	0.999
Trees	0.970	0.971	0.972	0.954	0.974
Asphalt	0.886	0.898	0.898	0.930	0.923
Self-blocking bricks	0.832	0.873	0.887	0.835	0.914
Bitumen	0.884	0.910	0.938	0.906	0.912
Tiles	0.961	0.964	0.963	0.952	0.965
Shadows	0.940	0.951	0.970	0.954	0.957
Meadows	0.990	0.992	0.991	0.995	0.995
Bare soil	0.611	0.626	0.677	0.589	0.694
Overall	0.971	0.975	0.979	0.974	0.979

 Table 4
 Comparison for the Pavia Centre dataset.

 Table 5
 Comparison for the Salinas-A dataset.

Class	SVM	SVM-PCA	SMSF	PMSF	SVM-MSF
Brocoli	1.000	0.986	0.997	0.929	1.000
Corn	1.000	1.000	1.000	1.000	0.997
Lettuce1	1.000	1.000	1.000	1.000	1.000
Lettuce2	1.000	1.000	1.000	1.000	1.000
Lettuce3	1.000	1.000	0.959	0.969	1.000
Lettuce4	1.000	1.000	0.996	0.983	0.999
Overall	1.000	0.999	0.994	0.988	0.999

 Table 6
 Comparison for the Salinas dataset.

Class	SVM	SVM-PCA	SMSF	PMSF	SVM-MSF
Brocoli1	0.999	0.999	1.000	0.999	0.998
Brocoli2	0.983	0.987	0.992	0.971	0.993
Fallow	0.998	0.999	0.997	0.995	0.999
Fallow rough	0.942	0.940	0.986	0.971	0.967
Fallow smooth	0.998	0.999	0.988	0.989	0.997
Stubble	0.999	0.997	0.995	0.991	1.000
Celery	0.999	0.998	1.000	0.998	0.999
Grapes	0.959	0.969	0.981	0.993	0.983
Soil	0.999	0.999	0.998	0.997	0.999
Corn	0.983	0.994	0.980	0.986	0.993
Lettuce1	0.985	0.976	0.968	0.958	0.978
Lettuce2	1.000	0.999	0.999	0.992	1.000
Lettuce3	0.987	0.971	0.922	0.916	0.990
Lettuce4	0.993	0.996	0.975	0.977	0.988
Vinyard1	0.925	0.944	0.952	0.991	0.962
Vinyard2	0.990	0.988	0.994	0.966	0.995
Overall	0.976	0.981	0.983	0.988	0.988

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Class	SVM	SVM-PCA	SMSF	PMSF	SVM-MSF
1	0.994	0.996	1.000	0.990	1.000
2	0.954	0.963	0.927	0.972	0.950
3	0.978	0.991	0.991	0.987	1.000
4	0.991	0.996	0.991	1.000	0.929
5	0.931	0.944	0.993	0.986	1.000
6	0.995	0.995	0.956	0.995	0.995
7	0.979	0.989	1.000	1.000	1.000
8	0.956	0.956	1.000	0.953	0.977
9	0.953	0.955	0.964	0.962	0.983
10	0.997	0.994	0.975	0.978	0.986
11	0.989	0.989	0.987	0.966	0.984
12	0.965	1.000	0.991	0.998	0.973
13	1.000	1.000	1.000	1.000	1.000
Overall	0.980	0.985	0.986	0.984	0.985

Table 7 Comparison for the Kennedy Space Center dataset.

4.4 PMSF Setup

These experiments were also conducted with 10% of the pixels for the SVM training. As in SMSF, the use of entropy and PCA for SVM was kept for a fair comparison of the MSF step. We only show the results for hyperspectral descriptors for the MSF step, once PCA and entropy did not result in any improvement.

Class	SVM	SVM-PCA	SMSF	PMSF	SVM-MSF
1	0.885	0.897	0.934	0.877	0.963
2	0.544	0.567	0.522	0.522	0.533
3	1.000	1.000	0.996	1.000	0.996
4	0.808	0.860	0.927	0.850	0.689
5	0.860	0.839	0.884	0.860	0.860
6	0.554	0.603	0.616	0.653	0.678
7	0.948	0.961	0.948	0.940	0.991
8	1.000	1.000	1.000	0.978	0.995
9	0.879	0.865	0.968	0.936	0.979
10	0.951	0.951	0.915	0.942	0.978
11	0.869	0.872	0.887	0.909	0.909
12	0.988	0.963	1.000	1.000	1.000
13	0.917	0.888	0.876	0.846	1.000
14	0.000	0.024	0.012	0.024	0.071
Overall	0.846	0.851	0.870	0.858	0.883

Table 8 Comparison for the Botswana dataset.

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Fig. 7 Post-hoc analysis of Friedman test for the different evaluated methods.

After the SVM step, PMSF computes the connected components on the resulting classification and the probability (reliability) of each classification being correct (for more details, refer to Ref. 15). These components and the following three parameters are used to select the seeds of the method:



Fig. 8 Visual comparison of the classification results given by SVM, SMSF, and SVM-MSF.

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- the way of selecting the seeds depends on whether a connected component is larger or smaller than a threshold *M*;
- if the connected component is larger than M, then a percentage P of its most reliable classifications is kept as seeds;
- otherwise, the pixels with reliability greater than a threshold S are kept as seeds.

The parameter S is the smaller probability value of the T% most reliable SVM classified pixels for the whole image. We will discuss the choice of T, as in the original paper. The parameter M was chosen to be 20, since it produced good results in our experiments, reflecting the choice in the original paper. The parameters P and T were set to ~5% as in the original paper; however, we did not obtain good results with such a choice. The differences may be due to the probability computations, even though we used LIBSVM,⁵¹ as in the original paper. The choice of these parameters for the PMSF method is summarized in Fig. 6. The overall accuracy stabilizes with P = 50% and T = 5%. The results get worse for T greater than 25%. The best results are obtained with T in the range (5%, 25%) (overlaid curves in Fig. 6).

4.5 Comparison

Tables 2–8 show the gradual improvement on the classification achieved by the proposed method when using SVM with PCA and then by adding the MSF step. The method also achieves



Fig. 9 Visual comparison of the classification results given by SVM, SMSF, and SVM-MSF.

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competing values of class accuracy and superior overall accuracy when compared to SMSF and PMSF. The experiments for these images were performed by using 10% of the pixels as training samples. The remaining pixels were used as test samples.

We performed the Friedman test and obtained a *p*-value of 0.0002, which suggests there is a significant difference between at least two methods. A post-hoc analysis based on the Tukey–Kramer test is shown in Fig. 7. The analysis shows that our method performs better than three methods (SVM, SVM-PCA, and PMSF) and has statistically equal results when compared against SMSF.

An interesting aspect of the proposed classification method is that, besides correcting erroneous classifications, the seed identification step also confirms many of the correct



Fig. 10 Visual comparison of the classification results given by SVM, SMSF, and SVM-MSF.

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classifications produced by the SVM classifier. The method preserves as many as possible of these correct classifications and works on the weak classified pixels, such as the ones close to spatial boundaries between classes.

Figures 8–10 show a visual comparison of the classification results produced by SVM, SMSF, and SVM-MSF. The images depict only the classes; that is, the respective background is removed and shown in black.

5 Conclusions

In this work, we described a method for hyperspectral data classification through the combination of spectral and spatial information. An SVM was employed to produce an initial classification, which was then refined by applying K-NN and MSF approaches, incorporating spatial information into the spectral classification process.

Experiments were conducted on several datasets, and the proposed approach was demonstrated to be very effective, providing results with high accuracy. Intermediate results also show that the entropy measure and PCA technique can be used to improve the classification obtained by the SVM classifier.

As directions for future work, we intend to investigate the combination of other complementary features, as well as the impact of using different parameters in the classification process.

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