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Abstract—Spectral-spatial processing has been increasingly explored in remote sensing hyperspectral image classification. While extensive studies have focused on developing methods to improve the classification accuracy, experimental setting and design for method evaluation have drawn little attention. In the scope of supervised classification, we find that traditional experimental designs for spectral processing are often improperly used in the spectral-spatial processing context, leading to unfair or biased performance evaluation. This is especially the case when training and testing samples are randomly drawn from the same image - a practice that has been commonly adopted in the experiments. Under such setting, the dependence caused by overlap between the training and testing samples may be artificially enhanced by some spatial information processing methods such as spatial filtering and morphological operation. Such enhancement of dependence in return amplifies the classification accuracy, leading to an improper evaluation of spectral-spatial classification techniques. Therefore, the widely adopted pixel-based random sampling strategy is not always suitable to evaluate spectral-spatial classification algorithms because it is difficult to determine whether the improvement of classification accuracy is caused by incorporating spatial information into classifier or by increasing the overlap between training and testing samples. To tackle this problem, we propose a novel controlled random sampling strategy for spectral-spatial methods. It can greatly reduce the overlap between training and testing samples and provides more objective and accurate evaluation.

Index Terms—Experimental setting, random sampling, spectral-spatial processing, data dependence, hyperspectral image classification, supervised learning

I. INTRODUCTION

Spectral-spatial processing have attracted increasing attentions during the past several years. Bringing spatial information into traditional single pixel based spectral analysis leads to better modelling of local structures in the image and facilitates more accurate land-cover and object classification. While a large portion of the hyperspectral remote sensing community have focused their research on improving classification accuracy by developing a variety of spectral-spatial methods [1], [2], [3], [4], few attention has been paid to experimental settings. Evaluation of hyperspectral image classification methods requires careful design of experiments such as appropriate benchmark data sets, sampling strategy to generate training and testing data, and appropriate and fair evaluation criteria [1], [5]. In the scope of supervised classification, we find that traditional experimental designs for spectral processing are often improperly used in the context of spectral-spatial processing, leading to unfair or biased performance evaluation. This is particularly the case when training and testing samples are randomly drawn from the same image/scene which is a common setting in the hyperspectral classification research due to limited availability of benchmark data and high cost of ground truth data collection.

Fig. 1 shows a typical spectral-spatial hyperspectral image classification system built on a supervised learning scheme. Training and testing samples are drawn from an image data set following a specific sampling strategy. After image preprocessing which may involve spectral-spatial operations, feature extraction step fuses the spectral and spatial information to explore the most discriminative feature for different classes. The extracted features are used to train a classifier that minimises the error on the training set. In the testing step, the learned classifier is used to predict the classes of testing samples based on the extracted features. The testing error is given by comparing the predicted labels with the ground truth, which can be used as a performance indicator for image preprocessing, feature extraction and classification methods.

In the experimental setting, the sampling strategy plays an important role in the classifier learning and evaluation. Given a dataset including a hyperspectral image and its land-cover classes or other ground truth data, in most cases, training and testing samples are not given in advance. A sampling strategy has to be employed to create the training and testing sets [6], [7], [8]. Random sampling is a natural choice since it treats all labeled data equally and each sample would be selected with the same probability. However, by this method, some classes with a small number of labeled samples may have much less selected samples than expectation. Therefore, a more sophisticated sampling method, stratified random sampling, is often used [7]. To guarantee each class having sufficient samples, it firstly groups those labeled samples into subsets based on their class labels, and then random sampling is carried out within each subset. In term of the number of training samples in each subset, it normally requires that proportion of each group should be the same as in the population. Then the rest of samples are employed as testing samples in the testing step. This method is very simple to implement, reproducible, and of statistical significance. To the best of our knowledge, a number of hyperspectral classification methods adopted this option in the experimental setting [2], [9], [3], [10], [11]. In the following sections, we refer to the stratified random sampling as random sampling.

Before proceeding to the issue of random sampling, we have to reaffirm some basic principles for supervised learning. Under statistical learning frame, a common assumption for inference purpose is that random variables are independent and identically distributed (i.i.d.). The identical condition implies that training and testing samples are generated from the same data distribution. The independent condition
In this paper, we study the relationship between sampling strategies and the spectral-spatial processing in hyperspectral image classification, when the same image is used for training and testing. We find that the experimental setting with random sampling makes data dependence on the whole image be increased by some spatial-spectral operations, and in turn increases the dependence between training and testing samples\(^1\). To address this problem, we propose an alternative controlled random sampling strategy to alleviate the side effect of traditional random sampling on the same hyperspectral image. This leads to a fairer way to evaluate the effectiveness of spectral-spatial methods for hyperspectral classification.

In summary, the contribution of this paper are in three aspects:

- We point out that the traditional random sampling from the same image experimental setting is not suitable for supervised spectral-spatial classification algorithms. This helps to re-examine the performance evaluation of various spectral-spatial classification methods.
- We find that under the random sampling setting, spectral-spatial methods can enhance the data dependence and improve the classification accuracy. We give a theoretical explanation for this phenomenon via computational learning theory.
- We propose a novel controlled random sampling strategy which can greatly reduce the overlap between training and testing samples caused by spatial processing, such that more objective and accurate evaluation can be achieved.

The rest of this paper is organized as follows. Section II reviews the spectral-spatial processing that have been commonly used for hyperspectral image classification. Section III provides an in-depth analysis on the dependency between training and testing samples. The spatial information embedded in the spectral-spatial processing under the experimental setting with random sampling is excavated and examined. Section IV analyzes the overlap between neighboring training and testing samples caused by spatial operations. Such overlap increases the dependence between training and testing samples, which may lead to mistakenly using of the testing data in the training process. Section V discusses the relationship among spectral-spatial processing, data dependence and classification accuracy via computational learning theory. A new sampling strategy is proposed in Section VI which reduces the influence of overlap between training and testing data. To prove its advantage over random sampling, a series of experiments are developed and results are presented in Section VII. At last, the conclusions are drawn in Section VIII.

II. SPECTRAL-SPATIAL PROCESSING IN HYPERSPECTRAL IMAGE CLASSIFICATION

The advantage of using hyperspectral data in land cover classification is that spectral responses reflect the properties of components on the ground surface [7]. Therefore, raw spectral responses can be used directly as the discriminative features of different land covers. At the same time, hyperspectral data also possesses the basic characteristic of the conventional images - the spatial information which corresponds to where a pixel locates in the image. The spatial information can be represented in different forms, such as structural information including the size and shape of objects, textures which describe the granularity and patterns, and contextual information which can express the inter-pixel dependency [3]. This assumption is also the foundation of development of most spectral-spatial methods for hyperspectral image classification.
In general, spectral-spatial information can contribute to hyperspectral image classification through three ways. Firstly, in image preprocessing, it can be used for image denoising, morphology, and segmentation. Image denoising enables the reduction of random noises introduced from sensor, photon effects, and calibration errors. Several approaches have been exploited for this purpose, for example, smoothing filters, anisotropic diffusion, multi-linear algebra, wavelet shrinkage, and sparse coding methods [13]. In most cases, denoising can be done by applying a local filter with designed or learned kernel across the whole image. In mathematical morphology, operations are performed to extract spatial structures of objects according to their spectral responses [14], [3]. Similar information is explored in image segmentation, which groups spatially neighboring pixels into clusters based on their spectral distribution [15], [9].

Secondly, common usage of joint spectral-spatial information lies in the feature extraction stage. While traditional spectral features are extracted as responses at a single pixel level in hyperspectral images, spectral-spatial feature extraction methods use spatial neighborhood to calculate features. Typical examples include texture features such as 3D discrete wavelet [10], 3D Gabor wavelet [16], 3D scattering wavelet[17], and local binary patterns [18]. Morphological profiles, alternatively, use closing, opening, and geodesic operators to enhance spatial structures of objects [19], [20], [21]. Other spectral-spatial features include spectral saliency [22], spherical harmonics [23], and affine invariant descriptors [24]. Heterogeneous features can be further fused using feature selection or reduction approaches [25].

Thirdly, some image classification approaches rely on spatial relation between pixels for model building. A direct way of doing so is calculating the similarity between a pixel and its surrounding pixels [26]. Markov random field, for example, treats hyperspectral image as dependent data and uses spectral information in the local neighborhood to help pixel class prediction [27], [9], [28]. Similar spatial structures are explored in conditional random fields [29], hypergraph modelling [30], and multi-scale analysis [11]. The spatial information can also be explored in constructing composite kernels in support vector machines [31]. While supervised learning approaches, such as K-nearest neighbors, linear discriminant analysis, Bayesian analysis, support vector machines, etc. are widely used in these classification tasks [32], [33], some approaches adopt semi-supervised or active learning strategies [34], [35].

### III. Spatial Information Embedded in Random Sampling

Random sampling makes the training and testing samples spread over the image, embedding plenty of underlying spatial information. In this section, we point out that the embedded spatial information will mistakenly influence the classifier learning and evaluation. We exploit this problem in a specific/extreme way, by which a hyperspectral classification task can even be done without spectral information.

In many benchmark hyperspectral datasets, pixels in the same class are not distributed randomly in the image. On the contrary, they tend to exist in continuous regions and follow a certain spatial distribution, especially when objects in the same materials present in the scene. Fig. 2 shows the false color composite and ground truth maps of five commonly used hyperspectral datasets, i.e., Botswana, Indian Pines (Indian), Kennedy Space Center (KSC), Pavia University (PaviaU), and Salinas scene (Salinas) [36]. In these images, there are strong dependencies between the spatial locations of pixels and land cover classes. This results in the potential using of the spatial structure and distribution of each single class. In most cases, if random sampling is used for selecting training and testing samples in the same image, the class label of a testing sample can be easily inferred only by its spatial relation with the training samples. This can be exemplified by Fig. 3, in which 5%, 10% and 25% of training data are sampled from the Indian Pines and Pavia University datasets. When it comes to 25% sampling rate, the spatial distribution of training samples (last column) is similar to the shape of the ground truth map (first column) in the spatial domain.

To show the extent that the classification accuracy is impacted by spatial information, we performed experiments on five benchmark datasets in Fig. 2. In the experiment, a nonlinear support vector machine (SVM) was employed because the land cover classes are not linearly separable in the spatial domain. The spatial coordinates were used as the spatial feature and no spectral information was included. The parameters of the SVM were learned via five-fold cross validation. Three sampling rates were explored, i.e. 5%, 10%, and 25% to generate the training data from all labeled samples, while the rest of labeled data served as the testing samples. In contrast to the spatial feature, the traditional spectral feature based methods were also implemented in which we followed the same setting as the spatial method.

Each test was repeated ten times in the experiment with random generation of training and testing samples. The overall classification accuracies (OA), average accuracies (AA) and Kappa Coefficient ($\kappa$) are shown in Table I for different methods. The comparison
between accuracies using spectral feature with SVM (Spe) and spatial feature with SVM (Spa1) shows some surprising results. Classification accuracy based on pure spatial feature has significantly outperformed the counterpart using pure spectral feature in all cases. In terms of overall accuracy, the spatial method achieves more than 93.8% accuracy on all datasets when only 5% of training samples are used, while the spectral method has only around 75.5% – 93.2% in accuracy. When the sampling rate becomes 25%, the accuracy almost reaches 100% for the spatial feature which agrees with the perceptual intuition in Fig. 3. Essentially, these phenomena are caused by the random sampling strategy on the same image. The results also show that higher sampling rate leads to increase of classification accuracy on all datasets.

In another point of view, the spatial classification can also be exploited in the local neighbourhood. Since the training samples spread uniformly in the image, it would be easy to find a nearest training sample for any testing samples that belong to the same class. An experiment was designed to test how the local information contributes the classification. We employed the K-nearest neighbor (KNN) classifier and set the parameter K to 1. The results are displayed in Table I under the columns of Spa2. It can be seen that the performance of Spa2 is comparable to the spatial method Spa1 on all datasets, which has significantly outperformed the spectral method on all datasets. It should be noted that in the KNN classification, predicting the label of testing samples is only based on the nearest training pixels in their spatial neighbourhood. This is similar to the mechanism of some spectral-spatial methods which also make use of the local spatial neighbourhood information but in a different way. This experiment further proves that the training data provide too much information on the spatial domain for the classification task. While classification based on spatial coordinates seems to perform better than the spectral information, it is infeasible in real applications in which unlabeled pixels are involved. Those unlabeled pixels are prone to be classified into its nearby class, thus producing a thematic map dramatically different from the reality. To exemplify this phenomenon, Fig. 4 shows the classification maps of the Indian Pines including the unlabeled pixels with 10% sampling rate. Although Sp1 and Sp2 achieve higher classification accuracy than Spe, their classification maps are far away from the ground truth map. Therefore this method is not acceptable in reality. In summary, these two experiments show that random sampling from the same image makes an underestimated amount of spatial information be embedded in the training set and the testing set. It is natural to raise the concern that they would interact with each other if spatial processing is applied to the image.

IV. OVERLAP BETWEEN TRAINING AND TESTING DATA FROM THE SAME IMAGE

The spectral-spatial methods make use of the spatial information in different forms and different ways as introduced in Section II. When it comes to the random sampling strategy, a more severe problem may happen in the spectral-spatial analysis, especially for the feature extraction stage. When only spectral responses are used, feature extraction is performed at single pixel, without exploring its
spatial neighborhood. Therefore, random sampling strategy provides a statistical solution for data splitting and there is no explicit overlap between training and testing samples. However, the spectral-spatial methods usually exploit information from neighborhood pixels. This is normally implemented by a sliding window with a specific size, for example, $3 \times 3$, $5 \times 5$ and so on. In each window, a kernel or filter is used to extract discriminative information. Since the training and testing samples are drawn from the same image, their features are almost certain to overlap in the spatial domain due to the shared source of information.

![Overlap between training and testing data on Indian Pines dataset under 5% sampling rate.](image)

Fig. 5. Overlap between training and testing data on Indian Pines dataset under 5% sampling rate.

![The regions for feature extraction from a training sample (O) and a testing sample (+) overlap with each other, as represented in gray color. The proportion of overlap is $\frac{3}{8}$ and $\frac{4}{5}$ for (a) $3 \times 3$ sliding window and (b) $5 \times 5$ sliding window, respectively.](image)

Fig. 6. The regions for feature extraction from a training sample (O) and a testing sample (+) overlap with each other, as represented in gray color. The proportion of overlap is $\frac{3}{8}$ and $\frac{4}{5}$ for (a) $3 \times 3$ sliding window and (b) $5 \times 5$ sliding window, respectively.

Fig. 5 shows the extent of overlap between training and testing data on the Indian Pines dataset. In the figure, the white dots show the locations of training samples, and the surrounding white squares cover a $3 \times 3$ region used for spectral-spatial feature extraction. The testing samples, however, may just lie in the the square and has its own surrounding regions. This brings about a shared region between features extracted from the training and testing data such that they interact with each other and lose the mutual independence. It is also evident that a larger filter leads to more overlap areas. An example is shown in Fig. 6 in which a $3 \times 3$ and $5 \times 5$ window will result in $\frac{3}{8}$ and $\frac{4}{5}$ of overlap for adjacent training and testing samples, respectively. Such overlap leads to using of the testing data for training purpose, and gives significant advantages to the spectral-spatial feature extraction approaches. This violates the basic principle of supervised learning that training and testing data shall not interact with each other. Depending on how feature is extracted, benefit of testing data may be explicit, for example when the spectral-spatial feature is extracted by concatenating the spectral responses of pixels in a neighborhood, or implicit, for example, by extracting texture features based on spatial frequency analysis such as discrete wavelet transform.

A. Experiment with a Mean Filter Based Spectral-spatial Method

In order to estimate how the overlap impacts the accuracy of spectral-spatial method with random sampling strategy, an experiment was carried out on the Indian Pines dataset. In this experiment, a linear SVM classifier was used to facilitate further comparison. The features were constructed by applying a mean filter to calculate the mean of the spectral responses in a neighborhood of the hyperspectral images, which was mathematically formulated as follows:

$$f(x, y) = \frac{1}{MN} \sum_{i=x-\frac{M}{2}}^{x+\frac{M}{2}} \sum_{j=y-\frac{N}{2}}^{y+\frac{N}{2}} S(i, j)$$  \hspace{1cm} (1)$$

where $M$ and $N$ are the width and height of neighborhood surrounding $(x, y)$. In the experiment, we set $M$ and $N$ both from 1 up to 27 with an interval of 2. $S(i, j)$ represents the spectral response at location $(i, j)$ and $f(x, y)$ is the feature extracted on location $(x, y)$ which contains both spectral and spatial information. This process can be considered as one of the simplest approaches to extract spectral-spatial features.

When the size of the neighborhood is $1 \times 1$, this reduces to extracting spectral feature only. Larger size of window results in more overlap. The calculated rate of testing samples covered by the neighborhood of training samples is shown in Fig. 7. When 5%
training data are sampled, 30.9% testing samples are covered by the 3×3 regions used to extract training features. When random sampling rate increases to 25%, the extent of overlap becomes 86.4%. The rise of sampling rate leads to rapid increase of overlap. Furthermore, when the size of filter grows, the overlap rate also increases rapidly. Eventually, when the overlap rate reaches 100%, all testing samples are used in the training process.

The experiment was repeated 10 times. In each time, the indices of the training and testing pixels were randomly generated. Features were generated using different settings of filter size and sampling rate. Under each setting, the same training and testing samples were used for fair comparison. The overall classification accuracies are shown in Fig. 8. Significant increase of the classification accuracy can be observed when spatial information is added to the spectral information. When the size of neighborhood increases, more testing data contribute to the training step, therefore the classification accuracy increases. It is also interesting to see that after the neighborhood increases to a specific size, the accuracy stops growing and tends to stable. This is probably because that when the neighborhood becomes too large, unlabeled data or samples from other classes are involved in the feature extraction, which neutralizes the benefits of overlap.

B. Non-overlap Measurement

Other than overlap, the increase of classification accuracy also owes to the better discriminative capability of spectral-spatial features. With larger filter size, the feature includes more spatial information. To demonstrate how the spatial neighborhood influences the effectiveness of spectral-spatial feature, we performed another experiment on those testing samples not overlapped with the training data.

Following the same setting as the previous experiment, we removed the testing samples that were covered by the training set and only test on the remaining samples. Table II shows the comparison of classification accuracy on all testing samples and non-overlap testing samples. The results show that when testing on non-overlap testing samples, the accuracy is improved when the neighborhood information is initially introduced by the 3×3 mean filter. However, when a larger size of filter is used, the accuracy of non-overlap testing samples does not increase and even decrease. The decrease could be caused by the fact that the non-overlap testing samples are easily influenced by the samples from other classes in the neighborhood. In contrast, the classification accuracy with overlapped testing samples has remarkable improvement when larger filter size is used.

Based on the above analysis, under the random sampling strategy, some filter-based spectral-spatial feature extraction methods would make the training and testing samples overlap and then interact with each other. Subsequently, in the training process, information from testing samples are included to train the classifier, which in return is used to classify the testing samples in the testing step. Although this kind of methods improves the classification results, they are not desired because they violate the basic assumption of supervised learning and their generalization is questionable. So far we have only analyzed a special case of spectral-spatial methods, it would be interesting to extend the analysis to a broader scope. Next, we try to discuss the data dependence and its impact on classification results by computational learning theory.

V. DATA DEPENDENCE AND CLASSIFICATION ACCURACY

Computational learning theory aims to analyze the computational complexity, feasibility of learning, and performance bound [37]. A widely known computational learning framework is the probably approximately correct (PAC) learning which estimates the sample complexity based on the required generalization error, probability of inference and complexity of a space of functions. Another classic theory is the Vapnik-Chervonenkis theory (VC theory). One of its functions is to bound the generalization ability of learning processes which is usually represented as the testing error $\hat{R}(h)$. 

Before introducing the computational learning theory, some basic learning concepts shall be firstly introduced in the scope of i.i.d. data. In computational learning, instead of considering the classification accuracy, a more general term, generalization error bound, is usually derived to describe the ability of learning algorithm to predict the unseen data. For a binary classification problem, given a hypothesis $h \in H$ where $H$ are all hypotheses, a target hypothesis $c$, and a sample set $S = (x_1, x_2, ..., x_m)$ following a distribution $D$, the empirical error (training error) $\hat{R}(h)$ and the generalization error (testing error) $R(h)$ can be defined as:

$$\hat{R}(h) = \frac{1}{m} \sum_{i=1}^{m} l(h(x_i), c(x_i))$$  \hspace{1cm} (2)

$$R(h) = \mathbb{E}_{x \in D} l(h(x), c(x))$$  \hspace{1cm} (3)

In Table II, the null values are due to the absence of non-overlapped testing samples.

<table>
<thead>
<tr>
<th>Filter Size</th>
<th>1</th>
<th>3</th>
<th>5</th>
<th>7</th>
<th>9</th>
<th>11</th>
</tr>
</thead>
<tbody>
<tr>
<td>All samples (5%)</td>
<td>72.1</td>
<td>86.1</td>
<td>90.2</td>
<td>91.4</td>
<td>92.1</td>
<td>92.3</td>
</tr>
<tr>
<td>Non-overlap (5%)</td>
<td>72.1</td>
<td>82.9</td>
<td>83.2</td>
<td>79.1</td>
<td>71.6</td>
<td>68.0</td>
</tr>
<tr>
<td>All samples (10%)</td>
<td>77.4</td>
<td>90.4</td>
<td>93.4</td>
<td>94.5</td>
<td>95.9</td>
<td>96.1</td>
</tr>
<tr>
<td>Non-overlap (10%)</td>
<td>77.4</td>
<td>86.2</td>
<td>84.8</td>
<td>77.9</td>
<td>65.3</td>
<td>NaN</td>
</tr>
<tr>
<td>All samples (25%)</td>
<td>82.4</td>
<td>94.6</td>
<td>97.5</td>
<td>98.2</td>
<td>98.5</td>
<td>98.7</td>
</tr>
<tr>
<td>Non-overlap (25%)</td>
<td>82.4</td>
<td>87.9</td>
<td>90.6</td>
<td>NaN</td>
<td>NaN</td>
<td>NaN</td>
</tr>
</tbody>
</table>
where $I$ is the error function and $E$ is the expectation.

Despite that the empirical error $\hat{R}(h)$ can be calculated once the training data $S$, its label $c(x_i)$ and the hypothesis $h$ are known, the generalization error cannot be estimated directly. In practice, simply decreasing $\hat{R}(h)$ by building complex classification model may not always minimize $\hat{R}(h)$ because it may lead to over-fitting. In order to bound $R(h)$, more factors have to be considered. Based on PAC learning, the generalization bound can be calculated as:

$$R(h) \leq \hat{R}(h) + \frac{1}{f} (log|F| + log\frac{1}{\delta})$$

which means that given training data of size $m$ and hypothesis complexity $|F|$, the inequality of generalization holds with probability no less than $1 - \delta$. This definition conforms to our understanding of learning that more training data leads to better learning outcome. Based on the inequality, the generalization bound can be tightened by increasing the training sample size $m$ or by decreasing the probability $1 - \delta$ which is equivalent to confidence of the inference. The complexity of hypothesis is determined by the learning models.

When the hypothesis sets are infinite, the above bound is uninformative. In order to impose generalization bound for infinite cases, the Redemacher complexity is introduced to measure the hypothesis complexity [38]. Specifically, it measures the variety of a set of functions by estimating the degree to which a hypothesis can fit random noise. The Rademacher complexity based generalization bound on i.i.d. data samples is defined as:

$$R(h) \leq \hat{R}(h) + \tilde{R}_n(H) + 3\sqrt{\frac{\log\frac{2}{\delta}}{2m}}$$

where $\tilde{R}_n(H)$ is the empirical Rademacher complexity. $1 - \delta$ is the probability of confidence and $m$ is the training sample size. $\tilde{R}_n(H)$ can be estimated by growth function or VC-dimension [37].

Even though these models provide generalization bounds for different learning algorithms, they are all based on the i.i.d. assumption. For non-i.i.d. data, the generalization bound has not been fully studied due to the lack of statistical model for dependent data. However, i.i.d. does not always hold in practice. In general, the samples in a hyperspectral image are not i.i.d., as the samples are spatially overlapping to each other in the image. The data dependence will inevitably happen no matter how carefully the sampling strategy is designed.

In recent years, researchers begin to develop new learning theories on this topic. Among all kinds of non-i.i.d. data, some data types possess the property of asymptotic independence, which is weaker than independence but stronger than dependence, for instances, time series model [39]. In order to define this kind of data, mixing condition is used to explicitly define the dependence of the future signal on the past signal based on decay. A commonly used model in non-i.i.d. scenario is the stationary $\beta$-mixing model [40]. Suppose events $A$ and $B$ are generated from a time sequence $\alpha \in (-\infty, +\infty)$ with an interval $k$, the definition of $\beta$-mixing coefficient is

$$\beta(k) = \sup_{m \in \alpha \in \alpha} \mathbb{E} \left[ \sup_{A \in \alpha_{m+k}} |P_r(A|B) - P_r(A)| \right]$$

This equation defines the dependence coefficient as the supremum of the difference between the conditional probability $P_r(A|B)$ and probability $P_r(A)$ when choosing arbitrary moment $m$ which separates event A and B. The sequence $\alpha$ is $\beta$-mixing if $\beta(k) \to 0$ when $k \to +\infty$. It implies that the dependence coefficient $\beta(k)$ decreases with the increase of interval $k$.

Several learning models have already been derived on stationary $\beta$-mixing data, such as VC-dimension bound[40], PAC learning [41] and Rademacher complexity [42]. In this work, the Rademacher complexity based generalization bound is employed since it associates the generalization bounds with $\beta$-mixing coefficient. It uses a technique to transferring the original dependent data to independent blocks. Let $2\mu$ be the number of blocks and each block contains $k$ consecutive points, then the size of sample $m = 2\mu k$. The original bound in Equation (5) is extended to $\beta$-mixing data as follows:

$$R(h) \leq \hat{R}(h) + \tilde{R}_n(H) + 3M \sqrt{\frac{\log\frac{2}{\delta}}{2\mu}}$$

where $M$ is the bound of a set of hypothesis $H$.

Compared with the i.i.d. case, this bound is not only related to the training error $\hat{R}(h)$, empirical Rademacher complexity $\tilde{R}_n(H)$, and probability $\delta$, but also relies on the $\beta$-mixing coefficient $\beta(k)$ which implies the degree of dependence among data. Considering the impact of $\beta$-mixing coefficient to the bound, this equation can be further simplified as:

$$R(h) \leq f(\beta(k)) + C$$

where $f(\beta(k))$ is a monotonicaly decreasing function. As a result, the generalization bound is tightened when the $\beta(k)$ increases, i.e. the dependence among data is enhanced.

Applying learning theory to hyperspectral image classification is challenging due to the complex statistical characteristic of hyperspectral images. To our knowledge, similar work is very rare. In the following experiments, we show that hyperspectral images share the same properties of $\beta$-mixing data.

Spectral feature extracted at image pixels often have strong dependence to their surrounding regions [4]. However, it is still questionable whether such dependence decreases with the increasing distance between the central pixel and its neighbouring pixels. In addition, since a hyperspectral image is a three-dimensional data, how the dependence is related to the spatial direction is still unknown. To check how the dependence varies with the distance, we performed a simple statistical analysis on the Indian Pines dataset. Here, the dependence between two pixels $X$ and $Y$ is approximated by the linear correlation coefficient of their spectral responses:

$$\rho_{X,Y} = \frac{cov(X,Y)}{\sigma_X \sigma_Y}$$

where $cov$ and $\sigma$ represent the covariance and standard deviation, respectively. A random location was firstly selected on this image, then the correlation coefficient $\rho$ was calculated between the pixel and its neighborhood pixels with different distances. We have calculated the result on a $9 \times 9$ patch. As expected, it does not show clear pattern at a single random pixel. Then we took account of pixels centered at all locations and calculated the mean of correlation coefficients in all patches based on the following:

$$c(x_i, y_i) = \frac{1}{M \times N} \sum_{x, y \in \mathbb{N}} \rho(S(x, y), S(x_i, y_i))$$

where $(x_i, y_i)$ is the neighbourhoud pixels of the central pixel $(x, y)$ in the patch. For a $9 \times 9$ patch, $x_i = x - 4, x - 3, ..., x + 4$ and $y_i = y - 4, y - 3,..., y + 4$. M and N are the width and height of the whole image. $S(x, y)$ represents the spectral response at location $(x, y)$. The statistical result is shown in Fig. 9. In the center of the patch, the intensity is one due to self-correlation. It clearly shows that with the increasing interval, the correlation coefficient gradually drops in all directions. This is consistent with the characteristic of $\beta$-mixing.

Now we can safely assume that hyperspectral images are $\beta$-mixing, and explore the relationship between the generalization bound with data dependence. Based on Equation (8), the bound is inversely related to $\beta$-mixing. As a consequence, the classification accuracy can be increased by enhancing the dependence between training and testing data. Recall that in the experiment with a mean filter based spectral-spatial method (Fig. 8), the accuracy increases with larger filters. It would be interesting to measure the pixel correlation to see how the mean filter influences the data dependence. Similarly, Equation 1 was used to calculate the mean spectral responses in a sliding window and the sizes of the windows were set to $1 \times 1, 3 \times 3, 5 \times 5, 7 \times 7, 9 \times 9, 11 \times 11$, respectively. Then the statistics on the
correlation coefficients on a $9 \times 9$ patch was calculated from the filtered images based on Equation 9 and Equation 10. Note when the $1 \times 1$ was used, this correspond to using the original image without filtering. The results are shown in Fig. 10 in which we only draw the correlation coefficients on a $9 \times 9$ patch was calculated from the projected illustration along X dimension from Fig. 9. From this new figure, two trends can be observed. Firstly, all curves drop continuously when the distance increases, which means that the processed data agree with the properties of $\beta$-mixing. Secondly, at the same distance, the larger the filter is, the stronger the dependence between the central pixel and its adjacent pixels becomes. Therefore, the overlap enhances the data dependence which tightens the error bound of the final classification results.

The theories presented above have explained why mean filter improves the classification accuracy, and they can be extended to other spectral-spatial operations that increase the data dependence. It should be noted that the above analysis is built on the assumption of random sampling for performance evaluation. Under such experimental setting, the improvement of classification accuracy comes from not only incorporating spatial information into classifier but also enhancing the dependence between training and testing data. While enhancement of such dependence in a proper way is needed to improve classification results, incorporating testing samples in the training procedure shall be avoided. Otherwise, it is difficult to determine whether the improvement of classification accuracy is caused by the former or latter. Therefore, to restrict the overlap rate between training and testing samples, we propose an alternative sampling strategy for evaluation of the spectral-spatial methods.

**Algorithm 1 Controlled Random Sampling Strategy**

**Require:** Hyperspectral Image $I$ and sampling rate $s$

**for** each class $c$ in $I$ **do**

Selects all unconnected partitions $P$ in the class $c$

**for** each partition $p$ in $P$ **do**

Calculate the number of training samples $n_t$ in the partition by $n_t = n_p \times s$

Randomly select a seed point $q$ in the partition

Applying the region-growing algorithm to extend $q$ to a region $r$ whose size is equal to $n_t$

**end for**

Combine these regions $r$ to form training samples $R_c$

**end for**

Combine the training samples $R_c$ and their corresponding class labels to get the whole training set $R$

**VI. A CONTROLLED RANDOM SAMPLING STRATEGY**

Following the discussion in previous sections, since random sampling from the same image is not suitable for evaluation the spectral-spatial methods, it is necessary to develop a new sampling strategy to separate the training and testing sets without overlap. It would be perfect if we could perform training and testing on two different images. Unfortunately, this is still infeasible in most cases due to the limited availability of benchmark datasets and high cost of ground truth data collection. Therefore, without changing much the current experimental setting, the goal is to significantly reduce the extent of data overlap and make the evaluation fair enough. Based on our analysis, the main problem of random sampling is that it makes the training and testing samples spatially adjacent to each other, leading to their overlap in the subsequent spatial operations. On the other hand, as a classical method, it has advantages such as simplicity, reproducibility, and statistical significance. As a result, the new sampling strategy should satisfies the following requirements. Firstly, it shall avoid selecting samples homogeneously over the whole image so that the overlap between training and testing set can be minimised. Secondly, those selected training samples should also be representative in the spectral domain, meaning that it shall adequately cover the spectral data variation in different classes. There is a paradox between these two properties, as the spatial distribution and the spectral distribution are coupling with each other. The first property tends to make the training samples clustered so that it generates less overlap between the training and testing data. However, the second property prefers training samples being spatially distributed as random sampling does, and covering the spectral variation in different regions of the image. Therefore, a good trade-off has to be achieved by the new sampling strategy. Thirdly, because there is no prior knowledge, we do not know which samples are more important than the others. Therefore the new method shall possess the property of randomness.

Here we propose a controlled random sampling strategy to achieve a compromise of the above considerations. Similar to random sampling, a pre-defined proportion of samples in each class is to be randomly selected as the training samples and the rest data serve as the testing samples. Those training samples shall be concentrated locally and dispersed globally. We borrow the idea of region growing to create region-shape training samples [43]. On the ground truth map, the seed points are randomly selected from different partitions of classes to make the training samples disperse globally and randomly. The partition referred here is a group of connected pixels with the same labels. For each class, there are usually several partitions distributed on the map, corresponding to the land cover of the same category in different locations. Then controlled random sampling proceeds with three steps. Firstly, it selects the unconnected partitions for the each class and counts the samples in each partition. This
step is to find the spatial distribution of each class and make sure that the selected training samples in the next step cover the spectral variance at the most extent. Secondly, for each partition, the training samples are generated by extending region from the seed pixel. In terms of region growing, it expands in all directions and takes account of 8-connected neighborhood pixels. All the adjacent pixels of seed pixels are examined and if they are within the same class, they work as the new seed points. This process is repeated until the amount of selected points reach a pre-defined number which is proportional to the number of pixels in the corresponding partition. This guarantees that the total number of training samples meet the pre-defined proportion of the whole data population. Thirdly, after the above steps are applied to all classes, those samples in the grown regions with their labels are chosen as the training samples and the rest of pixels work as the testing samples. In case when there are more partitions than the required training samples, partitions are again randomly sampled. A summary of this strategy is given in Algorithm 1.

In Fig. 11, we demonstrate different degrees of overlap between training and testing samples under random sampling and controlled random sampling strategies, after a Gaussian filter is applied. In the left column of the figure, the training and testing data are represented by colored dots and white regions in each partition. Applying the Gaussian filter creates the gray regions in the right column of the figure, representing the overlap between the training and testing data. It can be noticed that all the training samples are impacted by the testing data under random sampling. On the contrary, for controlled random sampling, only training samples at the edges of the training regions are influenced by the testing data. This figure clearly shows that the overlap from controlled random sampling is significantly less than that from the traditional random sampling.

To further illustrate how the controlled random sampling works with real datasets, examples on Indian Pines and Pavia University are given in Fig. 12 with 5%, 10% and 25% sampling rates. Compared with the random sampling strategy in Fig. 3, it can be observed that the spatial structure of each class can no longer be inferred from the training data as random sampling does. In the meantime, the training samples are still distributed across the whole image and a wide range of spectral variances are covered. Though this approach can not completely eliminate overlap between the training and testing data, the influence of testing data in the training stage can be greatly reduced to limited pixels at the boundaries of each training region. The experimental setting with the proposed sampling method can help us more accurately and objectively evaluate the performance of spectral-spatial methods.

VII. EXPERIMENTS

To prove the usability and advantage of the proposed controlled random sampling against random sampling, we have developed a series of experiments to test these two strategies when they are used to evaluate spectral-spatial operations in different stages of image classification. In the preprocessing step, we adopted a mean filter and a Gaussian filter as examples of smoothing and denoising operations. Then, we performed experiments with raw spectral feature to examine the effectiveness of the proposed sampling method when evaluating the spectral responses without spatial processing. Furthermore, two spectral-spatial feature extraction methods, i.e. 3D discrete wavelet and morphological profiles, were compared using two sampling methods. Finally, we compared the performance of two sampling strategies at varying sampling rates and discussed the limitation of the proposed method. In order to make the experiments more convincing, we adopted two widely used supervised classifiers, support vector machine (SVM) and random forest (RF) to validate our results. The SVM was implemented using the LIBSVM package [45], and the RF was implemented using the well-known Weka 3 data mining toolbox [46]. We present results on five benchmark datasets, i.e., Botswana, Indian Pines (Indian), Kennedy Space Center (KSC), Pavia University (PaviaU), and Salinas scene (Salinas).

A. Evaluation of Spectral-spatial Preprocessing Method

The spectral-spatial preprocessing step contributes to classification by improving the quality of hyperspectral images, reducing random noises, and enhancing specific features. By varying the parameters of mean filter and Gaussian filter, their influence to the classification accuracy under two sampling strategies can be analyzed. We undertook experiments on both Indian Pines and Pavia University datasets with SVM and RF, respectively. The results with mean filter are shown in Fig. 13. When traditional random sampling is used, the accuracy on the Indian Pines dataset increases with larger filter size when SVM and RF are adopted (Fig. 13(a) and (b)). For the Pavia University dataset (Fig. 13(c) and (d)), the accuracies also increase with larger filter size but decrease when the size reaches a specific value, which is slightly different from the results on the Indian Pines image. The reason may be that Pavia University has higher spatial resolution and interacts with filters in more complex way than the low spatial resolution Indian Pines data. The results confirm that using a mean filter with relative large size can increase the classification accuracy, up to 92.4% on Indian Pines and 98.0% on Pavia University. Essentially, it is mainly because larger filter leads to more overlap between the training and testing data. In contrast, when adopting the controlled random sampling strategy, the classification accuracy firstly improves marginally, but then dramatically drops with larger size filters. This is consistent with our expectation in evaluating the influence of spectral-spatial operations rather than the data dependence. Therefore, the proposed sampling method successfully avoids the problem of random sampling.

We then performed an experiment with Gaussian filter under the same setting to compare two sampling strategies. Among different denoising and smoothing approaches, Gaussian filter is a basic but effective tool to reduce the random noise in hyperspectral images. It works as a low-pass filter whose standard deviation controls the shape of filter and sets the threshold to remove the corresponding high frequency signal. The larger the stand deviation is, the lower frequency
the signal can be preserved and the image be more smoothed. We applied a Gaussian filter on each band of hyperspectral images with a range of standard deviations. The size of filters varies with the standard deviation so that the smoothing effect decays to nearly zero at the boundaries of filtering masks. Then the smoothed image was fed into the classifier. This experiment was repeated 10 times and the mean of overall accuracy was used as the evaluation criterion. The standard deviation ranged from $2^{-1}$ to $2^3$ with an interval of 0.5 on the exponential term. We plot the classification accuracy as a function of the standard deviation in Fig. 14 for random sampling and controlled sampling method, respectively. From Fig. 14 (a)-(d), we can see that the accuracy continuously increases until a specific point when Gaussian filter with larger standard deviation is used with random sampling strategy. This is consistent with the observation on the mean filter. We can assume that the Gaussian filter influences the data dependence to varying extents under different standard deviations, such that the classification accuracy is impacted by the filter parameter. This is also consistent with our earlier analysis that when data dependence is increased, the classification error bound will be tightened. However, this is not desired when evaluating a preprocessing method for image classification as we would like to know what is the actual contribution from the operation itself.

Compared with the random sampling, the controlled random sampling presents a different trend between the accuracy and standard deviation. The accuracy firstly improves marginally and then becomes stable or drops. This indicates that applying an appropriate Gaussian operator can reduce high-frequency signals in the data, and at the same time, local region is smoothed so that the differences between pixels in each region is reduced. These reduce the diversity of samples in each class to some extent, thus contributing to the final image classification. However, higher standard deviation of Gaussian may over-smooth the local samples, making the local samples highly similar. As a result, the classifier loses its generality to testing samples spatially far away from training samples. Under the new sampling strategy, Gaussian filter is able to improve the classification but not very significantly and the training and testing data dependence caused by overlap is no longer the dominant factor to the classification. Overall, these two experiments prove that the proposed sampling strategy is able to neutralize the improper benefit gained from enhancement of dependence between training and testing data.

B. Raw Spectral Feature

We then performed an experiment to compare two sampling strategies when raw spectral features were used on the benchmark datasets. The objective of this experiment is to examine the effectiveness and objectiveness of the proposed sampling method compared with random sampling. As mentioned in Section I, there is no issue with the experimental setting with random sampling when evaluating a pixel based spectral feature. But we still do not know whether the proposed sampling method is qualified in such a task.

In the experiment, only the raw spectral features were used without any spatial processing. Other settings were same as the previous experiment such as the classifiers, repetition of experimental runs, etc. The overall accuracy and standard deviation under random sampling and the controlled random sampling strategies(*) are reported in Tables III and IV for SVM and RF, respectively. Following observations can be made from the results. Firstly, higher sampling rate leads to increase of classification accuracy on all datasets. This is the same and expected for both sampling methods. Secondly, the standard deviation of the accuracy from the proposed sampling strategy is much higher than that of the random counterpart. This is due to the distinction of training data generated from the random seeds each time. Lastly, there is a reduction on the classification accuracy when the proposed sampling strategy is used. This is due to the fact that variations on the same class data in different regions are less sufficiently captured as some of them may not be included in the training samples when the proposed sampling strategy is used. The difference of accuracies is more evident on Indian Pines, Pavia University and Salinas datasets as these scenes include large blocks of regions in the same class, which leads to more benefits from spectral variation covered by random sampling strategy. For further illustrating this phenomenon, the classification maps on the Indian Pines and Pavia University under two sampling strategies are shown in Fig. 15. Compared with random sampling, those testing samples far away from the training regions are easily misclassified under controlled random sampling.

Despite the differences, this does not affect a fair evaluation of different algorithms with the proposed sampling strategy. In this experiment, assuming that the goal is to evaluate SVM and RF, it can be concluded from the results that SVM is a preferred classifier since it generates higher classification accuracy. Therefore, although the new sampling strategy has made the hyperspectral classification a more challenging problem and forces more rigorous evaluation to the feature extraction and classification approaches, it is still qualified in evaluating the algorithms in hyperspectral image classification.
C. Spectral-spatial Features

Now we turn our attention to test the proposed sampling strategy with two typical spectral-spatial feature extraction methods, i.e., 3D discrete wavelet transform (3D-DWT) and morphological profile. 3D-DWT is a typical example of filter-based methods. The morphological profile is a widely adopted spatial feature extraction method, including a number of variations for hyperspectral image classification.

1) 3D discrete wavelet transform: The discrete wavelet transform is derived from the wavelet transform which is a mathematical tool for signal analysis. Unlike Fourier transform, the advantage of wavelet transform is that the transformed signal provides time-frequency representation for the non-stationary signal, meaning that we can not only know whether a frequency component exists but also when it happens in a signal. The definition of continuous wavelet transform is shown as following:

$$
\Psi^x_{\psi}(\tau, s) = \int x(t) \cdot \psi_{\tau,s}(t) dt
$$

where $$\psi_{\tau,s}$$ is the basis functions (wavelet) with s and \(\tau\) that control the scale and translation, respectively. When it comes to discrete samples, DWT is implemented by a series of filters in the frequency domain. Since hyperspectral images consist of three dimensions, 3D-DWT exploits the correlation along the wavelength axis, as well as along the spatial axes, so that both spatial and spectral structures vary with the size of mean filter on the Indian Pines and Pavia University (PaviaU) datasets under random sampling (first row) and controlled random sampling (second row) strategies.

![Fig. 13](image13.png)

![Fig. 14](image14.png)
of hyperspectral images can be more adequately mapped into the extracted features.

In the implementation, we followed the multiple scale setting as described in [10], however, without the feature selection step. Firstly, the hyperspectral image was processed by a cascade of high pass filters and low pass filters. In each level, the data was decomposed into high frequency part and low frequency part. After three levels filters and low pass filters. In each level, the data was decomposed described in [10], however, without the feature selection step. Firstly, extracted features.

In order to keep the sub-cube and the original data cube at the same size, the filtered signals were not down-sampled as what the traditional DWT does. Then these sub-cubes were concatenated into the wavelet features. The multidimensional function was carried out along two spatial dimensions \(x\) and \(y\), as well as the spectral dimension \(\lambda\), respectively. The final concatenation worked as the feature for the whole data cube and can be represented as:

\[
\hat{C}_{\alpha}(x, y, \cdot) = \frac{1}{9} \sum_{i=x-1}^{x+1} \sum_{j=y-1}^{y+1} C(x, y, \cdot)
\]

(12)

In order to keep the sub-cube and the original data cube at the same size, the filtered signals were not down-sampled as what the traditional DWT does. Then these sub-cubes were concatenated into the wavelet features. The multidimensional function was carried out along two spatial dimensions \(x\) and \(y\), as well as the spectral dimension \(\lambda\), respectively. The final concatenation worked as the feature for the whole data cube and can be represented as:

\[
f(x, y) = (\hat{C}_{1}^x, \hat{C}_{2}^x, \hat{C}_{15}^x, \hat{C}_{1}^y, \hat{C}_{2}^y, \hat{C}_{15}^y, \hat{C}_{1}^\lambda, \hat{C}_{2}^\lambda, \hat{C}_{15}^\lambda)
\]

(13)

where \(f(x, y)\) is the 3D-DWT feature at location \((x,y)\).

The experimental results under random sampling strategy and controlled random sampling strategy (*) are shown in Table III and Table IV for SVM and RF, respectively. As expected, controlled random sampling strategy leads to lower accuracy compared with random sampling strategy on all datasets. Interesting observation can be obtained by comparing these results with the results on the raw spectral feature in Tables III and IV. On one hand, 3D-DWT performs better than raw spectral feature under both sampling methods. This indicates that the proposed method confirms that 3D-DWT is able to extract more discriminative information than raw spectral feature. On the other hand, under experimental setting with random sampling, 3D-DWT significantly improves the accuracy on all datasets over raw spectral feature. However, when testing it with the proposed controlled sampling strategy, the improvement can not reach the same level of significance, especially on Indian Pines, Pavia University, and Salinas datasets. It means that 3D-DWT does not perform that significantly better than the raw spectral features as expected, when eliminating the advantage of introducing information from the testing data into the training stage.

2) Morphological profile: To further analyze this issue, we undertook experiments on the mathematical morphology feature. Morphological operations employ the structuring elements in the image, making it possible to enhance or alleviate structures based on the specific requirements from users. The basic operators include erosion and dilation which expands and shrinks the structures, respectively. Combining them results in the opening (erosion-dilation) and closing (dilation-erosion) operations. These two processes can remove specific structures and noises without destroying the original primary structures in the image. The results of processing are called morphological profiles. Morphological profile based feature extraction method is able to explore the structures of objects based on the contrast and size of objects in the images, therefore, it has been widely studied for hyperspectral image classification [19], [21].

We followed a basic implementation of extended morphological profiles (EMP). The details of this method and its variation can be found in a survey paper from Fauvel et al [3]. The spatial feature was extracted as follows

\[
\Omega^{(n)}(I) = \left[\phi^{(n)}(I), ..., \phi^{(1)}(I), I, c^{(1)}(I), ..., c^{(n)}(I)\right]
\]

(14)

where \(\phi^{(n)}(I)\) and \(c^{(n)}(I)\) were the opening and closing operations with a disk-shape structural element of size \(n\), respectively. As different sizes of structuring elements were used, the morphological profile \(\Omega^{(n)}(I)\) was capable of integrating multi-scale information. Before the feature extraction, a principle component analysis (PCA) step was applied to hyperspectral images to reduce the dimension of the data. Then the morphological profiles were obtained on each of the \(m\) primary components:

\[
\Omega^{(n)}_{m}(I) = \left[\Omega_{1}^{(n)}(I), \Omega_{2}^{(n)}(I), ..., \Omega_{m}^{(n)}(I)\right]
\]

(15)

In the last step, the morphological profiles were stacked with the spectral response to form the spectral-spatial feature.

The classification results with two sampling strategies are shown in Table III and Table IV. Similar to the results on 3D-DWT, although...
The morphological profile feature has achieved better performance than the raw spectral method when tested with random sampling strategy, the improvement is not as significant when controlled random sampling is used. This is mainly because that spectral-spatial method does not take much advantage of the overlapped information between training and testing samples under the proposed method.

Directly comparing two completely different spectral-spatial methods may not make much sense since different features are more suitable to extract features on specific datasets or sensitive to specific classifiers. Here we analyze the results from another point of view, which may explain the advantage of the proposed sampling over random sampling. In Table III, 3D-DWT achieves higher accuracy than EMP on both Indian Pines and Pavia University datasets when random sampling is adopted. When adopting the new sampling strategy, 3D-DWT still performs slightly better than EMP on the Indian Pines, but EMP performs significantly better than 3D-DWT on the Pavia University. This is consistent with the fact that the morphology method is capable of extracting more spatial structures than 3D-DWT on the dataset with high spatial resolution [19].

Under the proposed sampling method, the properties of the spectral-spatial method can be more accurately reflected and evaluated in the experiments. This is impossible under random sampling because the classification result is strongly misled by the overlap between training and testing samples. Overall, the proposed sampling strategy reveals more real discriminative ability of the spectral-spatial methods, which is the purpose of the evaluation.

D. Relationship between Traditional Random Sampling and the Proposed Method

In previous experiments, we have used at least 5% of labeled data as training samples. In recent years, some researchers have used very few labeled samples to construct the training set. In such case, the potential overlap between training set and testing set would not be considered, which may explain the advantage of the proposed sampling over random sampling. In Table III, 3D-DWT achieves higher accuracy than EMP on both Indian Pines and Pavia University datasets when random sampling is adopted. When adopting the new sampling strategy, 3D-DWT still performs slightly better than EMP on the Indian Pines, but EMP performs significantly better than 3D-DWT on the Pavia University. This is consistent with the fact that the morphology method is capable of extracting more spatial structures than 3D-DWT on the dataset with high spatial resolution [19].

Under the proposed sampling method, the properties of the spectral-spatial method can be more accurately reflected and evaluated in the experiments. This is impossible under random sampling because the classification result is strongly misled by the overlap between training and testing samples. Overall, the proposed sampling strategy reveals more real discriminative ability of the spectral-spatial methods, which is the purpose of the evaluation.

3The percentage of overlap is defined as the percentage of the testing samples covered by the training features against all the testing samples, as described in Section IV.

Fig. 16. Overlap of training and testing data and classification accuracy on the Indian Pines dataset under random sampling and controlled random sampling strategies, when different percentage of labeled data were used as the training samples. The first row shows results of raw spectral responses and the second row shows the results of a $5 \times 5$ mean filter.
information based methods. This phenomenon is more obvious when a large amount of training data are used. An alternative controlled random sampling strategy is proposed to tackle these problems. It has been proved that the proposed method generates less overlap between training and testing samples for spectral-spatial methods compared with the random sampling. This advantage is even more obvious when the scale of training set is large. Therefore, this new strategy provides a proper way to evaluate the effectiveness of spectral-spatial operations and the corresponding classifiers.

Finally, it should be noted that the aim of this paper is not to criticize the spectral-spatial methods themselves or the exploration of spatial information. The concern is only on the widely adopted evaluation approach, or more strictly speaking, on the experimental setting. Under the experimental setting with random sampling, the performance evaluation may be not equally fair and unbiased for all spectral-spatial methods. This is especially the case for the practice that training and testing are performed on the same image. This problem is ultimately due to the lack of labeled hyperspectral data that are available for method evaluation. Therefore, a more urgent task for the research community is to build more benchmark datasets to facilitate future spectral-spatial hyperspectral image analysis research.

REFERENCE

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