Active Learning With Convolutional Neural Networks for Hyperspectral Image Classification Using a New Bayesian Approach

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I. INTRODUCTION

HSI (or imaging spectroscopy) [1] is based on the acquisition, measurement, analysis, and interpretation of spectra captured at different wavelength channels (throughout the visible and solar-reflected infrared spectrum) over an extensive observation area on the surface of the earth. A variety of imaging spectrometers are currently available, including airborne [e.g., the AVIRIS, the CASI, the ROSIS, and the Hypmap or the new hyperspectral missions based on HyperCam technology [2]–[9]] and spaceborne (e.g., the EO-1 Hyperion or the FTHSI on MightySat II [10]–[12]). Imaging spectrometers are also available on ground-based (stationary or handheld) platforms. These instruments allow for the acquisition of the solar-reflected spectrum in a large number of narrow and contiguous spectral bands (normally several hundreds) [13], creating data cubes in which each pixel contains a detailed contiguous spectral signature that can be used to characterize the objects in the scene with great precision and detail.

Several imaging spectrometers are currently operational, providing a large volume of hyperspectral data that can be used for a wide variety of applications, such as forestry, geology, precision agriculture, hydrology, ecological monitoring, scene recognition, military applications, and disaster monitoring [14]–[17]. For instance, we highlight the following spectrometers: AVIRIS [4], which measures the solar-reflected spectrum from 0.4 to 2.5 μm at the intervals of 0.01 μm creating hyperspectral images with 224 bands, EO-1 Hyperion, which also collects 242 bands in the range of 0.4–2.5 μm [16], [18], and ROSIS, which collects images with a spectral range from 0.43 to 0.96 μm [7], among others. Also, several new satellite missions will be soon operative and ready to collect data in a very similar spectral range. For instance, the imaging spectrometer included in the NASA HyspIRI [19] is expected to measure the visible-to-shortwave infrared in the range 0.38–2.5 μm or the German EnMAP [20] that is expected to collect data in the range 0.42–2.45 μm, as well as the Italian PRISMA Program [21].

The great amount of information that these spectrometers collect is very useful in pattern recognition, which has led to the development of multiple methods for the advanced classification of hyperspectral images [22], [23]. This includes unsupervised techniques (often called clustering methods) [24]–[28]. However, supervised classifiers are often preferred due to their capacity to provide high classification accuracy by considering class-specific information provided by labeled training samples [14].

In this sense, since their successful application in the field of pattern recognition in the 1990s [29], [30], ANNs have attracted the attention of a large number of researchers in the area of hyperspectral image classification [31], [32]. Their ability to learn by examples and to generalize, together with the following properties: 1) ANNs are nonparametric (i.e., they do not need prior knowledge of the statistical distribution of the classes) and 2) they offer multiple training techniques to deal with linearly nonseparable data [33], has made ANNs widely attractive for supervised classification of hyperspectral images compared with probabilistic methods.

In particular, DNNs [34], [35] have recently shown a great potential to yield high performance in image classification tasks [36]–[38]. DNNs are deep architectures (multilayer stack of simple modules) that have the capacity to learn more complex models than shallow ones [39], learning features at various levels of abstraction, i.e., the multilayer nonlinear transformations applied over ANNs architecture can adaptively extract more meaningful and discriminative features [40].

To date, four DNN models have been the mainstream DL architectures for the analysis of hyperspectral remote sensing images: DBNs, AEs, RNNs, and CNNs.

DL has emerged in part with DBN models [41], [42]. In [43], three DBNs to extract high-level features from hyperspectral data using spectral, spatial, and spectral–spatial information are introduced. In a similar way, [44] implements a DBN for feature extraction and classification, stacking spectral–spatial characteristics, while [45] investigates the hyperparameters used by the spectral and spectral–spatial DBNs in [43]. Another example is [46] in which a DBN is implemented introducing diversity promoting priors into the pretraining and fine-tuning phases in order to avoid the coadaption of latent factors.

On the other hand, the AE has been traditionally used as an unsupervised pixel-based method to learn useful features from data and perform dimensionality reduction. In the literature, we can find deep AE architectures, also called SAEs, for hyperspectral image classification, such as the SAE proposed in [47] that performs a two-step training strategy based on pixel spectrum, with an unsupervised representation learning and a supervised fine-tuning, before a final supervised classification step conducted by a logistic regression layer. In [48], an SAE is pretrained in unsupervised fashion with spectral data, and the features are extracted by a PCA+3D Gabor wavelet filter. In [49], three SAEs are introduced to generate high-level features from hyperspectral data using spectral, spatial, and spectral–spatial information with a logistic regression method performing the final classification. Following [49], in [50], two SAEs are proposed to extract spectral and spatial features that are stacked and embed into an SVM for classification.

SAEs and DBNs are successful DL methods for hyperspectral classification, improving their performance with the incorporation of spatial information in addition to the spectrum. However, both SAE and DBN models need to flatten the spectral–spatial features in 1-D vectors to satisfy their input accuracy by considering class-specific information provided by labeled training samples [14].
requirements, losing to a certain point the effectiveness of the spatial information [51] for characterization purposes.

Regarding RNN, it is a kind of network with loops in connections where node activations at each step depend on those of the previous step [52]. With the traditional pixel-based approach, the RNN exploits each hyperspectral pixel in the band-to-band fashion [52]. On the other hand, several CNN-based approaches (called CRNN [53]) have been implemented for hyperspectral classification. In [54], a 1-D CRNN is implemented, where spatial constraints are integrated by linear opinion pools. A similar model is used in [55], where a 1-D CRNN is trained in the semisupervised fashion with the labeled and unlabeled data (pixels) using pseudolabels. Again, RNNs and CRNNs can present the same problem as SAEs and DBNs: they need to adapt the spatial information in order to exploit it.

In this sense, we highlight CNNs [35] as a powerful tool for hyperspectral image classification [14] able to exploit both the spectral and the spatial information in an easy and natural way. CNNs successively apply convolution filters and pooling operations to the raw input data (which can be 1-D, 2-D, or 3-D), creating a hierarchy of layers whose outputs are increasingly complex feature vectors from the input data. In the literature, we can find multiple adaptations of these networks to hyperspectral analysis. Following a 1-D approach, [56] presents a five-layer 1-D CNN that receives \( n \times 1 \) input vectors, where \( n \) is the number of spectral bands, to classify hyperspectral images directly in the spectral domain. On the other hand, 2-D CNNs exploit the information from neighboring pixels in order to extract spatial features, whose input data are a patch of \( d \times d \) neighboring pixels [57], normally after applying PCA to extract the spectral features [58], [59]. Also, several approaches mix both 1-D and 2-D CNNs to extract spectral–space information, respectively [60], [61]. In contrast to these methods, several 3-D CNN models have been proposed that can learn both spatial and spectral features, taking as input data 3-D patches from the original hyperspectral data, processing each pixel by means of a 3-D convolution kernel in association with its spatial neighborhood and the corresponding spectral information [37], [62]–[64]. However, the application of CNNs to hyperspectral classification presents some issues, as they require a great amount of labeled data for fine-tuning the large number of training parameters that affect their generalization power, such as the number of hidden layers and their kernel size (which involves the number of weights, their biases, and the obtained feature maps), the pooling, padding, and stride sizes, the selected optimizer and its learning rate, and the batch size. These aspects make this kind of networks quickly overfit with small training sets which may lead to poor classification accuracy in the testing phase.

In general, the quality of ANN-based classification methods is strongly related to the quality and number of training samples available in advance [65]. In order to effectively learn the parameters of the classifier and to create a more robust and generalist model, a sufficient number of labeled samples are often required. However, in order to make the model as efficient as possible, the training set should be kept small and focused on the pixel samples that really help to improve the performance of the model. Moreover, the labeled samples are very difficult, expensive, and time consuming to collect in practice [66] and often only a few labeled samples are available in advance. This issue is particularly problematic in the hyperspectral image classification, since there is often an unbalance between the high dimensionality of the data and the limited number of training samples available [67], known as the curse of dimensionality or the Hughes effect [68]. As result, the ANN model may overfit the training data, which reduces its generalization capacity [69]. Some methods address this problem by using data augmentation techniques to generate additional training samples, performing basic transformations in the initial data set. Lu et al. [17] provide a method that offers robustness and flexibility in modeling scene images and report the improvements of the accuracy in scene recognition, constructing multiresolution features and modeling sparse features’ selection-based manifold regularization. In contrast, AL [70] has been used to facilitate the classification of hyperspectral image data sets, by including intelligently selected unlabeled samples from the original data set, i.e., the most informative samples, to the training set. This reduces the cost of acquiring large labeled training sets and the number of needed training samples [71]–[75].

However, the combination of AL with deep architectures, such CNNs, has been more difficult. This is because the goal of AL is to create a model composed by a predictor trained on a small set of well-chosen examples that can perform as efficiently as a predictor trained on a larger number of examples randomly chosen, while being computationally tractable, but CNNs often require large amounts of training data for training and are highly prone to overfitting when they are trained with small data sets. Also, similar to MLR, AL techniques rely on probabilistic functions, which indicate the probability of a sample to belong to the different existing categories, in order to create a model uncertainty but deep architectures normally do not represent model uncertainty, obtaining as final output the predicted class label instead the probability of each class. Moreover, conventional ANNs in general (and CNNs in particular) are based on the minimization of an error function [76], typically the least squared-error function between the desired class label and the obtained one in classification, and they cannot determine the level of uncertainty of their output results. In fact, the proposed model must be able to extract an output probability matrix from input data in order to apply the AL probabilistic function (called ranking function) and to extract those samples with more uncertainty, which will provide more information to the model.

To address this issue, in this paper, we consider BNNs [77], [78], a special kind of ANN that is robust to overfitting and is able to offer uncertainty estimates and a probabilistic interpretation of DL models by inferring distributions over the models’ weights, being able to learn from small data sets [79] and avoiding the tendency of conventional ANNs to make overconfident predictions in sparse data regions. In fact, we can consider BNNs as an extension of standard ANNs with posterior inference, adding a probability distribution on its weights [78], [80]. Recent works have showed that the Bayesian approach to CNNs (called hereinafter B-CNNs) can
offer robustness to overfitting on small data sets and improve their generalization capacity adding dropout at every weight layer (also called convolutional layer) of the CNNs, as a Bayesian approximation of the probabilistic model defined by the Gaussian process [81]–[83], which allows to represent the model uncertainty without introducing major changes to the network architecture. By taking advantage of BNNs (that offer good uncertainty estimates and are robust to overfitting) and following the methodology of [83], we propose, for the first time in the literature, an AL model with B-CNNs for hyperspectral remotely sensed data. The main innovative contributions of this paper can be summarized as follows:

1) the development (for the first time in the literature) of a dropout-based method (called B-CNN) to extract probabilistic information from 1-D, 2-D, and 3-D CNN models with the aim of performing accurate spectral–spatial feature-based classification of hyperspectral images using limited training data with different CNN architectures;

2) the development of a three-step-based training phase to perform AL over the proposed B-CNN for the first time in the hyperspectral image classification literature;

3) the exhaustive analysis and comparison of different acquisition functions to perform AL over the implemented B-CNN model, and a detailed comparison between the implemented B-CNNs and the standard 1-D, 2-D, and 3-D CNN classifiers for hyperspectral data in addition to other traditional hyperspectral data classifiers, such as RF, MLP, SVM, and MLR.

The remainder of this paper is organized as follows. Section II provides an overview of related works and presents the newly developed classifier model. Section III validates the proposed approach using three well-known hyperspectral data sets, highlighting the advantages of the newly proposed classifier. Finally, Section IV concludes this paper with some remarks and hints at plausible future research lines.

II. METHODOLOGY

A. Active Learning

AL has been adopted in remote sensing as an effective strategy to reduce the cost of acquiring large labeled training sets [75], and it is based on tree main aspects: 1) the availability of an initial training set; 2) the availability of a pool set; and 3) the use of an acquisition function. Let us denote by $D_{\text{train}} = \{X, Y\} = \{x_i, y_i\}_{i=1}^I$ a training set made up of $I$ labeled samples (where $x_i \in \mathbb{R}^d = [x_{i,1}, x_{i,2}, \ldots, x_{i,3}]$ is the input data, in our case a hyperspectral pixel vector, and $y_i = (1, 2, \ldots, C)$ is the corresponding label, with $C$ the number of different categories or classes) and $D_{\text{pool}} = [X] = \{x_i\}_{i=1}^{I+u} \in \mathbb{R}^d$ the pool of candidates, i.e., a set of $u$ unlabeled samples ($u \gg I$). The AL model is generally composed by a learner (trained with a few labeled samples, $D_{\text{train}}$) that iteratively selects new training examples from the pool of candidates ($D_{\text{pool}}$) that provide maximal information about the unlabeled data set and improve the model performance [74]. Algorithm 1 provides a general approximation of how AL works. As a result of the process illustrated in Algorithm 1, the classification accuracy given by the final selected training set is expected to be higher than the one obtained by using

![Fig. 1. Comparison between a conventional DNN (a multilayer perceptron or MLP) with three hidden fully connected layers and a CNN with three hidden CONV layers or kernels. The neurons in the CNN create 3-D blocks with sparse connectivity.](image-url)
randomly selected labeled samples. The acquisition function, in particular the user-defined heuristic, is a crucial point in AL. Tuia et al. [74] make a compilation of several heuristic methods, proposing a taxonomy of AL techniques. Here, we rely on posterior probability-based AL methods. These methods use the estimation of posterior probabilities of class membership, \( p(y|x) \), to rank the candidates in \( D_{\text{pool}} \). This kind of probability gives us an idea of the confidence of the class assignment, i.e., how good the classification is. However, DNNs, in general, and CNNs, in particular, normally do not calculate an uncertainty model that is needed for these AL methods. In Section II-B, we summarize how this problem is solved in [83].

### B. Bayesian-Convolutional Neural Networks

In contrast to conventional ANNs, the blocks or layers of neurons in CNNs operate such as kernels which are connected and applied over one region of the input image (also referred to as input volume hereinafter), i.e., layers are not fully connected to all neurons of the previous layer as in the standard multilayer perceptron or MLP (see Fig. 1). Each layer actually composes a feature extraction stage that can be of three kinds [64], [84].

1) **CONV Layer**: A layer where each node is in charge of computing the dot product (\( \cdot \)) between its own weights and a predefined region of the provided input volume to which it is connected. Actually, these layers work as kernels or filters where nodes share the same weights and bias, connecting the input volume to the output volume. Let us suppose a CONV layer that receives as input volume the data cube \( X \in \mathbb{R}^{d \times d \times n} \), where \( d \) represents the height and width and \( n \) represents the deep of the cube (also spectral bands). Each neuron in one filter that composes the CONV layer (\( k \) is the number of filters) will operate with a chunk of \( X \), in particular, with a \( l \times l \times q \) chunk (called filter bank, \( W^v \)). We can calculate the output of the neuron \((i, j, t)\) in the \( k \)th filter of the CONV layer as

\[
z_{i,j,t} = (X : W^v)_{i,j,t} = \sum_{i=0}^{l-1} \sum_{j=0}^{l-1} \sum_{t=0}^{q-1} x_{(i+s)i,(j+s)j,(t+s)t} \cdot w_{i,j,k} + b
\]

(1)

where \( z_{i,j,t} \) is the element \((i, j, t)\) in the \( k \)th feature map, \( x_{i,j,t} \) is an element of input data \( X \), \( w_{i,j,k} \) is a weight of the cube of weights \( W \), \( b \) is the bias, and \( s \) is the stride of the CONV layer. In fact, we can observe each filter as a window that moves itself on \( X \) in chunks of size \( l \times l \times q \) with a displacement dictated by \( s \). As a result, an output volume \( Z \) is obtained, which will be an array composed by \( k \) 1-, 2-, or 3-D feature maps depending on the kernel’s dimension.

2) **Nonlinearity Layer**: This layer is used to implement a nonlinear function (such as the sigmoid function or the ReLU [85]–[87]), which is then applied to each component of the obtained feature map to learn non-linear representations: \( A = f(Z) \).

3) **Pooling Layer**: Pooling layers are used to resume the output \( Z \) of several nodes in CONV layers using a pooling function. In addition, they also provide location invariant features. Normally, this layer executes a max operation (MAXPOOL layer) within a small region \( R \) defined by a kernel \( l \times l \times q \) over the resulting volume \( A \) after the nonlinearity layer, i.e., \( A \) is divided into several nonoverlapping maps, whose maximum values are mapped into the final output volume \( P = \max_{x \in R} A_x \).

Two characteristics make CNNs an ideal model for processing and classifying hyperspectral images: the sparse connectivity and shared weights. These features allow us to reduce the number of parameters to be learned by the network ensuring some degree of shift, scale, and distortion invariance. However, CNNs require huge amounts of data for regularization due to their model’s complexity and to the fact that they quickly overfit with small training sets. Such an overfitting problem makes that CNNs exhibit poor predictive performance in the testing phase. To avoid this problem, we adopt BNNs due to their robustness to overfitting and to their capacity to learn from small training sets.

Specifically, we use B-CNNs that combine the features of BNNs with the classification potential of CNNs. Another advantage of B-CNNs in our context is that they provide the uncertainty model that we need to apply AL techniques.

Given a training data set \( D_{\text{train}} = [X, Y] \) composed by inputs \( X = \{x_1, \ldots, x_i\} \) (where each \( x_i \in \mathbb{R}^n = [x_{i,1}, x_{i,2}, \ldots, x_{i,n}] \) and their corresponding outputs \( Y = \{y_1, \ldots, y_i\} \) (where each \( y_i \in \{1, 2, \ldots, c\} \)), the model posterior’ goal is to estimate a function \( y_i = f(x_i) \) as close as possible to the original function that has generated the outputs \( Y \). The Bayesian approach proposes to put some prior distribution over the space of functions \( p(f) \), so we can define a probability or likelihood on the output \( Y \) given the input \( X \) and a function \( f, p(Y|X, f) \). Therefore, the posterior distribution will be \( p(f|X, Y) = p(f|D_{\text{train}}) \) that captures the most likely functions given the observed data. In this way, the output \( y^* \) of a new input \( x^* \) can be predicted as the marginal likelihood

\[
p(y^*|x^*, D_{\text{train}}) = \int p(y^*|f^*) p(f^*|x^*, D_{\text{train}}) df^*.
\]

(2)

As (2) is normally intractable, we can approximate it adding a finite set of random variables \( \omega \) as follows:

\[
p(y^*|x^*, D_{\text{train}}) = \int p(y^*|f^*) p(f^*|x^*, \omega) p(\omega|D_{\text{train}}) df^* d\omega.
\]

(3)

In a BNN with weights \( W_i \) of size \( K_i \times K_{i-1} \) for each layer \( i \), the set of finite variables or parameters will be \( \omega = \{W_i\}_{i=1}^L \) (where \( L \) is the number of layers), and the posterior over \( \omega \) given \( X \) and \( Y \) will be \( p(\omega|D_{\text{train}}) \). However, the probability distribution \( p(\omega|D_{\text{train}}) \) is not tractable for a BNN. To infer the model posterior in a simple way, [83] proposes the use of variational inference as an approach based on Bernoulli approximation variational distributions (and relating this to dropout training) with the aim of not increasing the number of
parameters to be trained, as in other types of approaches such as the variational inference with Gaussian [88]. The first step is to define the approximating variational distribution \( q(W_i) \) for each BNN’s layer \( i (i = 1, \ldots, L) \) as

\[
W_i = M_i \cdot \text{diag}(\{z_{i,j}\}_{j=1}^{K_i})
\]

\[
i = 1, \ldots, L,
\]

\[
j = 1, \ldots, K_{i-1}
\]

where \text{diag} is the diagonal matrix with elements \( z_{i,j} \) that are Bernoulli distributed random variables with probabilities \( p_i \) and \( M_i \) are variational parameters to be optimized. Predictions follow (3). The change resides in replacing the intractable probability distribution \( p(\omega|\mathcal{D}_{\text{train}}) \) by the approximate distribution \( q(\omega) \) that belongs to a tractable family, which minimizes the Kullback–Leibler divergence, \( D_{KL}(q(\omega)||p(\omega|\mathcal{D}_{\text{train}})) = 0 \), a measure that returns the similarity between both the distributions

\[
q(y^*|x^*) = \int p(y^*|f^*)p(f^*|x^*, \omega)q(\omega)d\omega.
\] (4)

Using Monte Carlo integration, we can approximate the integral so that we can predict the probability that the output \( y^* \) corresponds to label \( c \) as follows:

\[
p(y^* = c|x^*, \mathcal{D}_{\text{train}}) = \int p(y^* = c|x^*, \omega)p(\omega|\mathcal{D}_{\text{train}})d\omega
\]

\[
\approx \int p(y^* = c|x^*, \omega)q(\omega)d\omega
\]

\[
\approx \frac{1}{T} \sum_{t=1}^{T} p(y^* = c|x^*, \hat{\omega}_t)
\]

being \( \hat{\omega}_t \sim q(\omega) \) called MC-dropout, while \( T \) are the stochastic forward passes. This Bernoulli approximation variational inference in BNNs can be implemented by adding dropout layers after certain weight layers in a network [83]. In the B-CNN model, this is the same than adding dropout to all CONV layers as well as inner-product layers.

C. AL Acquisition Function

The AL acquisition function \( a(x, \mathcal{M}) \) of a model \( \mathcal{M} \) with pool data \( \mathcal{D}_{\text{pool}} \) and inputs \( x \in \mathcal{D}_{\text{pool}} \in \mathbb{R}^d \) decides which data points \( x \) will be queried by an external oracle, which could be a human expert that performs the work of classifying the unlabeled data to be added to the training set \( \mathcal{D}_{\text{train}} \).

\[
S = \arg \max_{x \in \mathcal{D}_{\text{pool}}} a(x, \mathcal{M}).
\]

Gal et al. [89] make a review and a comparison between different acquisition functions. This paper performs a comparison with six different acquisition methods that have been adapted to AL methodology, taking into account different measurements, such as the entropy value and distances of the samples, among the random selection of samples.

1) Random Acquisition or Baseline: It chooses a point \( x_i \) following a uniformly random distribution from \( \mathcal{D}_{\text{pool}} \Rightarrow a(x_i) = \text{unif}() \), where \text{unif}() returns a draw from a uniform distribution over the interval \([0, 1]\).

2) Mean STD [90]: For each \( x_i \), it calculates \( \sigma(x_i) = (1/C) \sum c \sigma_c \), where \( C \) is the number of classes, \( c \) are the classes that \( x_i \) can take, and

\[
\sigma_c = \sqrt{\mathbb{E}_{q(\omega)}[p(y = c|x_i, \omega)^2] - \mathbb{E}_{q(\omega)}[p(y = c|x_i, \omega)]^2}.
\]

3) Maximum Entropy [91]: It chooses \( x_i \in \mathcal{D}_{\text{pool}} \) with the highest classification uncertainty, i.e., \( x_i \) that maximizes the predictive entropy

\[
\mathbb{H}[y_i|x_i, \mathcal{D}_{\text{train}}] := - \sum_c p(y_i = c|x_i, \mathcal{D}_{\text{train}}) \log p(y_i = c|x_i, \mathcal{D}_{\text{train}}).
\]

4) BALD [92]: This method chooses \( x_i \in \mathcal{D}_{\text{pool}} \) that are expected to maximize the mutual information between the predictions and the model posterior

\[
\mathbb{H}[y_i, \omega|x_i, \mathcal{D}_{\text{train}}] := \mathbb{H}[y_i|x_i, \mathcal{D}_{\text{train}}] - \mathbb{E}_{p(\omega|\mathcal{D}_{\text{train}})}[\mathbb{H}[y_i|x_i, \omega]]
\]

where \( \mathbb{H}[y_i|x_i, \mathcal{D}_{\text{train}}] \) is the entropy [91]. The selected points exhibit high variance in the input to the softmax layer.

5) Breaking Ties Criterion (BT-Criterion) [93], [94]: This method focuses on the boundary region between two classes with the aim of obtaining more diversity in the composition of the training set \( \mathcal{D}_{\text{train}} \). Sample \( x^{BT} \) is selected from \( \mathcal{D}_{\text{pool}} \) by

\[
x^{BT} = \arg \min_{x_i \in \mathcal{D}_{\text{pool}}} \left\{ \max_{c \in \mathcal{C}} \frac{p(y_i = c|x_i, \omega)}{\max_{c \in \mathcal{C}\{c^+\}} p(y_i = c|x_i, \omega)} \right\},
\]

where \( c^+ = \arg \max_{c \in \mathcal{C}} p(y_i = c|x_i, \omega) \) is the most probable label class for sample \( x_i \).

6) Mutual Information Criterion [70], [94]: It measures the mutual dependence between samples. In fact, this function selects the sample \( x^{MI} \) maximizing the MI between the obtained results and the class labels

\[
x^{MI} = \arg \max_{x_i \in \mathcal{D}_{\text{pool}}} I(\omega; y_i|x_i)
\]

where \( I(\omega; y_i|x_i) = (1/2) \log(|H^{MI}|/H) \) represents the MI between the obtained results and the class label \( y_i \), with \( H \) the posterior precision matrix and \( H^{MI} \) the posterior precision matrix after including the new sample \( x_i \).

In order to compare these acquisition functions, they have been adapted to be executed with AL methodology. For example, the BALD method has been approximated with \( q(\omega) \), as described by [89]

\[
\mathbb{H}[y_i, \omega|x_i, \mathcal{D}_{\text{train}}] := \mathbb{H}[y_i|x_i, \mathcal{D}_{\text{train}}] - \mathbb{E}_{p(\omega|\mathcal{D}_{\text{train}})}[\mathbb{H}[y_i|x_i, \omega]]
\]

\[
= - \sum_c p(y_i = c|x_i, \mathcal{D}_{\text{train}}) \log p(y_i = c|x_i, \mathcal{D}_{\text{train}})
\]

\[
+ \mathbb{E}_{p(\omega|\mathcal{D}_{\text{train}})} \left[ \sum_c p(y_i = c|x_i, \omega) \log p(y_i = c|x_i, \omega) \right].
\]
If we consider the BALD equation the identity
$$ p(y_i = c|x_i,D_{\text{train}}) = \int p(y_i = c|x_i,\omega)p(\omega|D_{\text{train}})d\omega, $$
we have
$$ \mathbb{E}[y_i,\omega|x_i, D_{\text{train}}] := -\sum_c \int p(y_i = c|x_i, \omega)p(\omega|D_{\text{train}})d\omega$$
$$ \cdot \log \int p(y_i = c|x_i, \omega)p(\omega|D_{\text{train}})d\omega$$
$$ + \mathbb{E}_p(\omega|D_{\text{train}}) \left[ \sum_c p(y_i = c|x_i, \omega) \log p(y_i = c|x_i, \omega) \right].$$

Now, we can apply Monte Carlo integration as follows:
$$ \mathbb{E}[y_i,\omega|x_i, D_{\text{train}}] := -\sum_c \int p(y_i = c|x_i, \omega)q(\omega)d\omega$$
$$ \cdot \log \int p(y_i = c|x_i, \omega)q(\omega)d\omega$$
$$ + \mathbb{E}_q(\omega) \left[ \sum_c p(y_i = c|x_i, \omega) \log p(y_i = c|x_i, \omega) \right]$$
$$ \approx -\sum_c \left( \frac{1}{T} \sum_{t=1}^T \hat{p}^t_c \right) \log \left( \frac{1}{T} \sum_{t=1}^T \hat{p}^t_c \right)$$
$$ + \frac{1}{T} \sum_{c,j} \hat{p}^t_{cj} \log \hat{p}^t_{cj}. $$

D. Proposed B-CNN Architecture for Active Learning

Finally, we present the new B-CNN architecture developed in this paper. It should be noted that the literature on CNNs applied to hyperspectral image classification shows different points of view on how the spatial and the spectral information in the original hyperspectral image can be used:

1) extracting only spectral information implementing a 1-D CNN architecture [14], [37], [56];
2) extracting only spatial information implementing a 2-D CNN architecture [57], [95]–[97];
3) extracting spectral–spatial information implementing a 3-D CNN architecture [37], [63].

In this regard, we emphasize that our B-CNN approach can be applied to 1-D, 2-D, and 3-D CNN architectures. This paper investigates the effects of applying the proposed Bayesian network to CNN models with the aim of performing hyperspectral classification based on spectral, spatial, and spectral–spatial features. In this sense, three B-CNN models have been implemented: 1-D, 2-D B-CNN, and 3-D B-CNN.

1) Spectral B-CNN Architecture: This model takes advantage of only the spectral information contained in the input hyperspectral image by developing a 1-D CNN architecture and performing a traditional pixelwise-based learning. Given the hyperspectral image $X \in \mathbb{R}^{h \times w \times n}$, where $h$ and $w$ are the height and width, respectively, and $n$ is the number of spectral bands, the 1-D B-CNN model will take as input data pixel vectors of the hyperspectral scene $X$, $x_i \in \mathbb{R} = [x_{i,1}, x_{i,2}, \ldots, x_{i,n}]$, where $i = 1, 2, \ldots, (h \cdot w)$. In this case, each input pixel vector $x_i$ is transformed through the net into feature maps, capturing the spectral information contained in $x_i$. Equation (1) can be rewritten as
$$ z_{rt} = (x \cdot W^r)_t = \sum_{i=0}^{q-1} x_{(t+1) \cdot s} \cdot w^*_i + b $$
where $q$ is the depth of the kernel, $z_{rt}$ is the $r$th neuron’s output in the $k$th filter, $x_i$ is one spectral band of the CONV layer’s input $x$, $W^r$ is the filter bank of the layer, characterized by $k$ kernels of size $1 \times q$, $w_i$ is a weight of vector $W$, $b$ is the bias of the layer, and $s$ is the stride.

In order to compare the implemented 1-D B-CNN model with the 1-D CNN baseline, the model’s architecture has been inspired by [56]. As we can see in Fig. 2, the proposed 1-D B-CNN model is composed by one input layer that receives the pixel vector with all its spectral bands. This input feeds one CONV layer, $c_1$, with $k_1^1$ kernels of size $1 \times q^1$, followed by the ReLU activation function and one maxpool layer, $mp_1$, whose kernel size is $1 \times m^1$. The output of $mp_1$ is reshaped into a vector in order to feed two fully connected layers at the end of the network. After the maxpool and first fully connected layers, dropout is implemented in order to perform the MC-dropout. Table I shows the details of the 1-D B-CNN implementation.

2) Spatial B-CNN Architecture: This model takes advantage of only the spatial information contained in the input image, reducing the number of spectral bands $n$ to 1 by applying PCA over original hyperspectral data sets. As result, a 2-D CNN architecture has been implemented, whose input is composed by patches of size $d \times d \times 1$ extracted from the hyperspectral scene. Normally, CNNs in general (and B-CNNs in particular) receive a completely normalized image prior to classification, i.e., a 3-D input array. However, in hyperspectral images, the classes are often mixed, so we feed the pixel (vectors of $1 \times n$) one by one to the B-CNN. This allows us to exploit the rich spectral information contained in the hyperspectral data in the case of the 1-D B-CNN, but we also need an additional mechanism in order to include also the spatial information in the 2-D model. In this case, we feed the network with the pixels that belong into a neighborhood window centered around each pixel under consideration. In this way, the input layer of the 2-D model accepts volumes of $d \times d \times 1$ [64], after processing the original scene with PCA. This requires a preprocessing stage in order to create patches of $d \times d \times 1$ for each pixel, where the desired label to be reached by the network will be the one owned by the central
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Fig. 3. Proposed spatial B-CNN (2-D B-CNN) architecture.

The pixel of the patch \([d/2 + 1, d/2 + 1, n]\) in this case, (1) can be rewritten as

\[
z_{i,j} = (X \cdot W^c)_{i,j} = \sum_{l=0}^{l-1} \sum_{j=0}^{l-1} x_{(i+s+l)_{i,j}+i,j} \cdot w_{i,j} + b
\]

where \(l\) is the height and width of the kernel, \(z_{i,j}\) is the output of the neuron \((i, j)\) in the \(k\)th filter, \(x_{i,j}\) is one pixel of the input patch \(X \in \mathbb{R}^{d \times d \times 1}\), \(W^c\) is the filter bank of the layer, characterized by \(k\) kernels of size \(l \times l\), \(w_{i,j}^{c}\) is a weight of matrix \(W\), and \(b\) is the bias of the layer and \(s\) the stride.

Fig. 3 shows the implemented 2-D B-CNN model. In this case, a deeper architecture has been selected. As we can observe, the input layer receives the hyperspectral patches of size \(d \times d \times 1\), which feeds the first CONV layer \(c1\). After that, three pairs of CONV+maxpool layers are implemented, \(c2\) and \(mp1\), \(c3\) and \(mp2\), and finally \(c4\) and \(mp3\). The network ends with two fully connected layers \(fc1\) and \(fc2\), where the last one performs the final classification. Dropout has been added at the end of certain layers in order to model the uncertainty of the network. Table I shows the details of the 2-D B-CNN implementation.

3) Spectral–Spatial B-CNN Architecture: This model takes advantage of both the spectral and the spatial information in the input hyperspectral image by developing a 3-D architecture that receives as input data patches of size \(d \times d \times n\), where \(n\) is the number of spectral bands. As in the 2-D B-CNN model, a preprocessing stage is required in order to create the input patches for each pixel [64], where the desired label to be reached by the network will be the central pixel of the patch \([d/2 + 1, d/2 + 1, n]\). In this case, each CONV layer performs (1).

As we can see in Fig. 4, the proposed spectral–spatial B-CNN consists of an input layer that receives the input patches, two CONV layers, \(c1\) and \(c2\) (with ReLU as a nonlinear activation function), two maxpool layers at the end of each CONV layer, \(mp1\) and \(mp2\), and two fully connected layers, \(fc1\) and \(fc2\). The last one is the output layer, which obtains the desired label for the input data. After each maxpool layer, a dropout layer is inserted in order to model the probability of the proposed network that will allow to obtain the uncertainty estimation. Table I provides additional details about the considered spectral–spatial B-CNN architecture.

The parameters of the considered 1-D, 2-D, and 3-D architectures, including the number and type of layers or kernel sizes, are one of the design choices of the proposed spectral, spatial, and spectral–spatial B-CNN models. In this sense, the architectures have been selected and defined in a way that is as general as possible to adapt them to different hyperspectral images [64]. In addition, our decision to use the same architectures for different data sets further illustrates that the proposed method can achieve good classification results on very different images, extracting the samples that maximize the information gained about the model, improving the model's performance.
the training, and showing its robustness, regardless of the fact that nonoptimal or customized topologies are adopted.

Also, we must remark that the proposed 1-D, 2-D, and 3-D B-CNN models have been developed as a computation graph using the library for machine intelligence Keras with TensorFlow back end over CUDA toolkit and the library of primitives for DNNs cuDNN. This computation graph is composed by connected nodes that represents operations (also called units of computation), while connections (or edges) represent the data consumed (input connections) and produced (output connections) in the unit. These connections allow us to represent the existing dependences between different operations, making it possible to identify those operations that can be executed in parallel in an easy way.

The process of our 1-D, 2-D, and 3-D B-CNNs follows two main steps. In a first step, the hyperspectral image is first loaded and a band-mean normalized version is calculated so that the values of the image are in the range $[0, 1]$. Then, the hyperspectral image’s ground truth is divided into two data sets: two randomly selected samples per class, $2 \cdot C$, will compose the initial training set, $\mathcal{D}^0_{\text{train}}$, and the remaining samples will compose the working set. From the working set, 50% of random selected pixels will compose the initial pool set, $\mathcal{D}^0_{\text{pool}}$, and the remaining 50% will be divided into testing samples (testing set with the 95% of samples) and validation samples (validation test with 5% of samples).

The next step is given by Algorithm 1. At this point, and with the aim of reducing the use of storage by the algorithm, the hyperspectral data are preprocessed in order to create the input samples that will feed the model (i.e., pixels vectors $1 \times n$, or patches $d \times d \times 1$ or $d \times d \times n$). In this sense, Algorithm 1 has been adapted to perform, for the first time in the literature, AL over the new B-CNN models for spectral, spatial, and spectral–spatial classification of hyperspectral data in an efficient way, both computationally and in terms of memory management. In $\epsilon = 0$, the training set $\mathcal{D}^0_{\text{train}}$ and the pool set $\mathcal{D}^0_{\text{pool}}$ are created as sets of $1 \times n$ pixel arrays.

Then, the B-CNN models are trained by a three-step process.

1) If we are working with 2-D or 3-D models, for each pixel $x_i \in \mathcal{D}^0_{\text{train}}$, we create a patch of size $d \times d \times 1$ or $d \times d \times n$, depending on the model’s dimension, centered on the pixel $x_i$, and assign the pixel’s label, $y_i$, to the patch. Once we have created the patches, these are sent to the network and the model is trained with MC-dropout in order to extract the labels $y_i'$, optimizing the cross-entropy function

$$H_s(y') = \sum_i y_i \log(y_i')$$

where $y_i$ is the original label of the $i$th sample and $y_i'$ is the predicted label obtained by the model.

2) Once a certain number of epochs have been executed and the weights and biases of the model have been adjusted, $\mathcal{D}^0_{\text{pool}}$ is sent as the test data to the network. MC-dropout is used to capture the confidence of the model in its predictions, calculating the probability of the output $y_i'$ for each $x_i$ in $\mathcal{D}^0_{\text{pool}}$, $P(y_i' | x_i, \mathcal{D}_{\text{train}})$. From an implementation point of view, $\mathcal{D}^0_{\text{pool}}$ passes through the network $T$ times, where $T$ is the number of stochastic forward passes. As a result, $T$ different outputs $y_i'$ have been obtained for each $x_i$ in $\mathcal{D}^0_{\text{pool}}$. To obtain the final probability, the average between all the outputs is calculated $y_i' = (1/T) \sum_{t=1}^{T} y_i'^{(t)}$, where $y_i'^{(t)}$ is the output of the model for $x_i$ in the $t$th stochastic forward pass [81].

3) The uncertainty over the model predictions, represented by the $T$ predicted probabilities, is used in the AL acquisition function in order to rank the unlabeled samples in $\mathcal{D}^0_{\text{pool}}$ according to their uncertainty. Then, those samples with higher score are selected, creating the $\mathcal{D}^0_{\text{selected}}$ set. With this process, those samples that provide more information and diversity to the network are considered to improve the final performance. To assign the
corresponding labels, each \( x_i \in \mathcal{D}_{\text{selected}}^0 \) is paired with its corresponding label \( y_i \). Finally, the selected pixels in \( \mathcal{D}_{\text{selected}}^0 \) are inserted into the training set as patches, each one with \( 1 \times n, d \times d \times 1 \), or \( d \times d \times n \), depending on the model’s dimension, creating the next \( \mathcal{D}_{\text{train}}^1 \). Also, the selected pixels in \( \mathcal{D}_{\text{selected}}^0 \) are deleted from the pool set, creating \( \mathcal{D}_{\text{pool}}^1 \).

After validating the model, the training process is repeated successively with each \( \mathcal{D}_{\text{train}}^i \) until a satisfactory result is achieved. We note that the aforementioned procedure allows us to avoid the calculation of the corresponding patch for all the pixels of the image, reducing the computation time and memory requirements of the algorithm.

### III. Experiments and Results

#### A. Experimental Configuration

In order to evaluate the performance of our newly developed approach, we use a hardware environment composed by a 6th Generation Intel Core i7-6700K processor with 8M of Cache and up to 4.20 GHz (four cores/eight-way multitask processing), 40 GB of DDR4 RAM with a serial speed of 2400 MHz, a GPU NVIDIA GeForce GTX 1080 with 8-GB GDDR5X of video memory and 10 Gb/s of memory frequency, a Toshiba DT01ACA HDD with 7200 r/min and 2 TB of capacity, and an ASUS Z170 pro-gaming motherboard. On the other hand, the used software environment is composed by Ubuntu 16.04.4 \( \times 64 \) as an operating system, CUDA 8 and cuDNN 5.1.5, and Python 2.7 as programming languages.

#### B. Hyperspectral Data Sets

In our experiments, three hyperspectral data sets have been used.

1. The first one is the well-known Indian Pines (IP) data set (Table II). This data set was gathered by AVIRIS [4]...
in 1992 over a set of agricultural fields with regular geometry and with a multiple crops and irregular patches of forest in Northwest Indiana. The IP scene has 145 × 145 pixels with 224 spectral bands in the range from 400 to 2500 nm, with 10-nm spectral resolution, 20-m spatial resolution, and 16-bit radiometric resolution. After an initial analysis, 4 zero bands and another 20 bands with lower SNR because of atmospheric absorption have been removed, retaining only 200 spectral channels. Moreover, about half of the pixels in the hyperspectral image (10 249 of 21 025) contain ground-truth information, which comes in the form of a single label assignment having a total of 16 ground-truth classes.

2) The second hyperspectral data set used in experiments was also collected by the AVIRIS instrument, in this case over Salinas Valley (SV), California (Table II). The covered area has 512 × 217 samples and the spatial resolution is 3.7 m/pixel; 204 out of the 224 bands are kept after 20 water absorption bands are removed. The ground truth is composed of 54 129 pixels and 16 land-cover classes, including vegetables, bare soils, and vineyard fields.

3) The third data set used in experiments is the Kennedy Space Center (KSC) (Table II), also collected by the AVIRIS instrument over Florida in 1996. Once noisy bands have been removed, the resulting image contains 176 bands with a 512 × 614 size, ranging from 400 to 2500 nm, and with 20-m spatial resolution. A total of 5122 pixels labeled in 13 classes, representing different land cover types, are considered for classification purposes.

C. Performance Evaluation

In order to test the proposed method, five different experiments have been carried out. In the first, second, third, and fourth experiments, the AL acquisition functions presented in Section II-C are tested considering an MLR classifier and the proposed 1-D, 2-D, and 3-D B-CNN models, respectively, with the aim of comparing the performance of each function over different classifiers, based on statistical or neural models, and using spectral, spatial, and spectral–spatial information. The fifth experiment makes a comparison between the AL methods (MLR: 1-D, 2-D, and 3-D B-CNN models adapted to AL) and the original ones (baseline MLR: 1-D, 2-D, and 3-D CNN models) with the aim of studying the impact of $D_{\text{train}}$ on the performance of both AL and traditional methods.

Also, we remark that each experiment uses the three considered hyperspectral data sets, running each model (with each acquisition function) over each scene 5 times, creating batches of 100 pixels, and working with the limited-memory Broyden–Fletcher–Goldfarb–Shanno optimizer [98], [99] in the case of the MLR, with $L_2$ as penalty and tolerance value fixed to 1e-18, being 1000 the number of maximum iterations, and with the Adam optimizer [100] for the 1-D, 2-D, and 3-D B-CNN models, with a learning rate of 0.001 and 100 epochs. Spatial and spectral–spatial patches have been created using a size of $d = 23$ for spatial patches and $d = 19$ for spectral–spatial patches with the aim of extracting enough spatial information from neighboring pixels. We have empirically observed that the value of $d$ should be large enough to characterize the spatial–contextual information around each pixel [64]. In this regard, $d = 23$ and $d = 19$ provide an appropriate compromise for the considered images (the selection of other close values of $d$ did not have a significant
impact on the final classification results). Finally, for each $c$, 10 unlabeled pixels have been chosen.

1) Experiment 1 (Performance of Different Acquisition Functions With the MLR): This experiment implements an AL-based MLR classifier. The output of the classifier directly gives the probability of each sample $x_i$ that belongs to the class $c$, i.e., $p(y_i = c|x_i, D_{\text{train}})$. This means that, to obtain the probabilities of $D_{\text{pool}}$, only one forward pass is needed, $T = 1$. Before running each implementation of AL-MLR, the testing set and the validation set are created. In this sense, $D_{\text{0train}}$ starts with $2\cdot C$ labeled samples. This means that IP and SV start with 32 labeled pixels and KSC starts with 26, while 50% of the remaining ground-truth samples is used to initialize the pool set $D_{\text{0pool}}$, i.e., IP’s $D_{\text{0pool}}$ starts with 5109 unlabeled pixels and SV with 27049. With the KSC data set, the percentage has been changed in order to reserve more data for the pool set, so that 85% of the remaining ground-truth samples have been selected to form $D_{\text{0pool}}$ with 4407 samples, instead of 2592. After $D_{\text{0train}}$ and $D_{\text{0pool}}$ are created, 5% of the remaining ground-truth data is used for validation and the remaining 95% for testing.

Table III shows the accuracy results of each acquisition function after 80 iterations, i.e., $\epsilon = 80$, with 10 acquisitions at each iteration. Focusing on the IP data set, the AL-MLR classifier with BT-criterion obtains the best overall accuracy (OA) when compared with the other functions, reaching 78.79% with only 8.12% of the ground truth, while the lowest OA is reached by the mean standard deviation (mean STD) with 73.90%. We can observe this behavior at different $D_{\text{train}}$ sizes in Fig. 5, which shows the evolution of the AL-MLR accuracy and execution times for each acquisition function. On the one hand, when more samples are added to $D_{\text{train}}$, higher accuracies are obtained until reaching a point in which adding more samples does not improve the classification accuracy, i.e., adding more samples does not add more information to the model. On the other hand, as more samples are added to $D_{\text{train}}$, higher execution times are needed by the model. Focusing on IP, BT-criterion’s accuracy stands out with 200 training samples, reaching the best classification results in less time than MI, BALD, and max-entropy. It is interesting to focus on the classification of Oats class (class 8) with only 20 pixels. We can observe how entropy-based...
methods, i.e., BALD and max-entropy, are able to reach better classification results for this particular class than distance-based methods, BT-criterion, and MI, while in classes with more samples (e.g., class 10, Soybeans-min), these methods reach very similar results. Also, MI, BALD, and max-entropy present very similar results in terms of both computation time and accuracy, being the random and mean STD the fastest methods but with the lowest accuracy results. In particular, we can observe that, although the random function reaches better OA than the mean STD, its average accuracy (AA) is slightly worse, i.e., the mean STD generalizes better than the random function, as it can be observed with Oats and Alfalfa (class 0 with 46 pixels), where the mean STD reaches better accuracy than the random function when less pixels are available.

A similar behavior can be observed for the SV data set, where distance-based methods reach better OA than random and entropy-based ones. In particular, the BT-criterion is able to reach 91.80% accuracy with only 1.53% of the ground truth, being max-entropy the function with the lowest OA, 86.25%. In general, entropy-based methods reach poorer OAs with the SV data set than the random function due to the spectral characteristics of the image and the number of pixels per class, which is quite balanced, being Lettuce-romaine-6wk, class 12 the one with less pixels, 916, where max-entropy and BALD reach the best OA with lower standard deviation than MI. Moreover, although BALD reaches a lower OA than the random function, it obtains better AA, so it can generalize better. Looking at Fig. 5, the second column shows the OAs and execution times of SV at different $D_{\text{train}}$ sizes. BT-criterion and MI stand out with 100 training samples, being the slowest methods, where random and mean STD are the fastest ones.

2) Experiment 2 (Performance of Different Acquisition Functions With the 1-D B-CNN): Our second experiment performs a comparison between all the acquisition functions for the proposed 1-D B-CNN model, whose architecture is described in Table I. In this case, 300 Monte Carlo iterations ($T = 300$) have been implemented, while $D_{\text{train}}$ and $D_{\text{pool}}$ follow the same initialization described in the MLR experiment. Table IV shows the obtained results in the 80th iteration of the model ($T = 80$) with 10 acquisitions at each iteration. Focusing on the IP scene, we can observe that the
distance-based BT-criterion reaches the highest OA value, 86.14% with 8.12% of the ground truth (i.e., 832 training samples), followed by the MI function, while the random function achieves the lower results with 81.83% of accuracy. As in the AL-MLR case, BALD and max-entropy reach very similar results with BALD exhibiting better generalization performance than the other tested acquisition functions. In Fig. 6, we can observe the performance of the proposed model. In the IP plot, the acquisition functions are very close one to each other at different $D_{\text{train}}$ sizes in terms of both OA values and execution times. In this case, the mean STD is the slowest method and the random function is the fastest one, while BT-criterion, MI, BALD, and max-entropy provide a very similar performance.

For the SV data set, the MI method reaches the best accuracy result, 97.27% with 1.54% of the ground truth (i.e., 832 training samples), demonstrating a high generalization power, which is overcome only by the BALD function. Observing Fig. 6, we can see that the BT-criterion has good OA between 100 and 300 training samples, being outperformed by MI with some variability. Again, the execution times are very similar for different functions, being mean STD the slowest one and random the fastest one, while BT-criterion, MI, BALD, and max-entropy are quite similar.

The results for the KSC are similar to those with the SV scene, with distance-based methods reaching the highest OA values, being the MI the best one: 93.57% OA with 15.85% of the ground truth. Also, the execution times are very similar with regard to those obtained for the SV data set.

With these data sets, we can also observe how 1-D B-CNN is able to scale logarithmically, rather than linearly, as in the case of the MLR. Also, we can see how the max-entropy and BALD functions suffer when a few samples are used. In this case, the model is not able to obtain good uncertainty values for these acquisition functions due to a poor dropout. To address this issue, we can add more uncertainty and variability to the model increasing the dropout values at the CONV and fully connected layers.

3) Experiment 3 (Performance of Different Acquisition Functions With the 2-D B-CNN): The third experiment performs a comparison of different acquisition function with the proposed 2-D B-CNN model, whose architecture is described in Table I. The parameter $T$ has been set to 300, and the initialization of $D_{\text{train}}$, $D_{\text{pool}}$, test and validation sets are the same as in our experiments with the 1-D B-CNN and MLR.

Table V shows the obtained results over the three considered hyperspectral data sets at iteration $\epsilon = 80$. Focusing on the IP data set, we can observe that distance-based methods are able to reach classification results over 99% accuracy, followed by entropy-based methods, with an OA around 98%. Fig. 7 shows the performance of the proposed spatial model with different sizes of $D_{\text{train}}$. MI and BT-criterion provide similar OA values from 600 to 800 training samples, while BALD remains close to max-entropy. On the other hand, although the random function is the fastest one, it reaches the lowest OA value, being BALD the slowest one, while BT-criterion, MI, and max-entropy exhibit similar execution times.

The SV data set provides very similar results, being the BT-criterion and MI the acquisition functions with better OA, followed quite closely by max-entropy. Again, in Fig. 7, we can observe how the BT-criterion and MI present very similar results, while max-entropy and BALD stay close one to each other, being mean STD and random the methods with the lowest OA results. The execution times are rather similar to the
Table VII

Classification Results Obtained by AL-Based Presented Methodologies in Comparison with Those Obtained with Traditional Hyperspectral Data Classifiers after 80 Iterations and 10 Acquisitions per Iteration

<table>
<thead>
<tr>
<th>Class</th>
<th>RF</th>
<th>MLP</th>
<th>SVM</th>
<th>1D-CNN</th>
<th>2D-CNN</th>
<th>1D-CNN</th>
<th>2D-CNN</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>25.43 (17.39)</td>
<td>56.52 (13.40)</td>
<td>57.17 (19.20)</td>
<td>23.48 (9.47)</td>
<td>55.8 (6.72)</td>
<td>94.86 (5.72)</td>
<td>97.33 (2.17)</td>
</tr>
<tr>
<td>0</td>
<td>62.67 (4.59)</td>
<td>76.27 (2.04)</td>
<td>76.90 (2.61)</td>
<td>74.24 (2.45)</td>
<td>82.05 (1.65)</td>
<td>94.40 (0.63)</td>
<td>96.29 (1.33)</td>
</tr>
<tr>
<td>0</td>
<td>27.5 (5.99)</td>
<td>63.26 (4.00)</td>
<td>64.51 (5.52)</td>
<td>72.37 (6.61)</td>
<td>92.98 (1.04)</td>
<td>95.21 (0.17)</td>
<td>97.24 (0.09)</td>
</tr>
<tr>
<td>0</td>
<td>34.09 (11.38)</td>
<td>57.85 (3.56)</td>
<td>65.82 (11.92)</td>
<td>19.44 (8.88)</td>
<td>63.9 (7.85)</td>
<td>97.47 (0.99)</td>
<td>99.16 (0.84)</td>
</tr>
<tr>
<td>0</td>
<td>81.04 (4.48)</td>
<td>83.79 (4.17)</td>
<td>89.01 (2.62)</td>
<td>74.95 (6.43)</td>
<td>92.82 (1.67)</td>
<td>97.09 (1.14)</td>
<td>98.17 (1.76)</td>
</tr>
<tr>
<td>0</td>
<td>94.88 (2.77)</td>
<td>94.47 (3.12)</td>
<td>95.70 (1.14)</td>
<td>84.77 (3.57)</td>
<td>99.25 (1.78)</td>
<td>99.77 (0.41)</td>
<td>100 (0.00)</td>
</tr>
<tr>
<td>0</td>
<td>31.07 (18.24)</td>
<td>72.14 (15.35)</td>
<td>81.07 (16.55)</td>
<td>60.71 (9.85)</td>
<td>73.78 (4.32)</td>
<td>94.63 (4.57)</td>
<td>97.10 (8.63)</td>
</tr>
<tr>
<td>0</td>
<td>98.62 (3.33)</td>
<td>96.92 (1.22)</td>
<td>97.89 (1.40)</td>
<td>97.07 (1.08)</td>
<td>96.88 (0.71)</td>
<td>100 (0.00)</td>
<td>96.99 (0.16)</td>
</tr>
<tr>
<td>0</td>
<td>28.01 (11.08)</td>
<td>60.00 (17.44)</td>
<td>70.00 (17.33)</td>
<td>41.00 (17.72)</td>
<td>66.67 (12.27)</td>
<td>100 (0.00)</td>
<td>85 (15.13)</td>
</tr>
<tr>
<td>0</td>
<td>66.01 (6.94)</td>
<td>74.50 (2.25)</td>
<td>73.30 (4.96)</td>
<td>67.47 (3.35)</td>
<td>84.43 (3.88)</td>
<td>89.87 (1.08)</td>
<td>96.14 (1.34)</td>
</tr>
<tr>
<td>0</td>
<td>85.56 (4.55)</td>
<td>83.15 (8.80)</td>
<td>83.79 (1.93)</td>
<td>73.78 (3.45)</td>
<td>94.78 (1.74)</td>
<td>97.13 (0.61)</td>
<td>98.23 (0.48)</td>
</tr>
<tr>
<td>0</td>
<td>97.68 (6.45)</td>
<td>78.51 (6.48)</td>
<td>78.24 (6.12)</td>
<td>92.91 (6.21)</td>
<td>99.25 (0.51)</td>
<td>99.77 (0.49)</td>
<td>100 (0.00)</td>
</tr>
<tr>
<td>0</td>
<td>92.20 (3.69)</td>
<td>97.56 (20.85)</td>
<td>89.83 (1.76)</td>
<td>98.54 (0.31)</td>
<td>98.21 (1.15)</td>
<td>99.76 (0.24)</td>
<td>99.51 (0.49)</td>
</tr>
<tr>
<td>0</td>
<td>93.06 (0.92)</td>
<td>94.06 (1.60)</td>
<td>94.22 (1.66)</td>
<td>94.21 (1.18)</td>
<td>96.28 (0.29)</td>
<td>99.29 (0.71)</td>
<td>99.48 (0.37)</td>
</tr>
<tr>
<td>0</td>
<td>41.50 (7.21)</td>
<td>63.34 (3.58)</td>
<td>54.27 (7.11)</td>
<td>58.55 (6.89)</td>
<td>53.89 (1.68)</td>
<td>98.45 (0.51)</td>
<td>95.73 (1.27)</td>
</tr>
<tr>
<td>0</td>
<td>83.35 (8.55)</td>
<td>86.24 (7.48)</td>
<td>90.00 (3.85)</td>
<td>85.18 (1.85)</td>
<td>90.32 (2.63)</td>
<td>94.09 (4.84)</td>
<td>97.06 (4.77)</td>
</tr>
</tbody>
</table>

IP data set, being the processing of SV slowest than IP. Again, BALD is the slowest method and random is the the fastest one, being BT criterion, max-entropy, and MI very similar. Finally, for the KSC data set, the best OA is reached with the BT criterion as an acquisition function with distance-based methods and entropy-based functions providing the highest overall values. Looking at Fig. 7, we can see in this case how BALD performs even worse than the random function until 600 training samples are reached. This is because the network has not achieved a sufficiently adjusted accuracy in the training phase, resulting from the great variability in the dropout. On the other hand, the execution times are very similar to those achieved in the experiments with the SV data set.

4) Experiment 4 (Performance of Different Acquisition Functions With the 3-D B-CNN): Our fourth experiment implements the proposed spectral–spatial B-CNN classifier using the six considered acquisition functions. Table VI and Fig. 8 show the obtained results.

With the IP data set, the best OA is reached by the BALD acquisition function, while max-entropy exhibits the best generalization power. Entropy-based methods are closely followed by distance-based methods, being random and mean STD the functions with the lowest OA. In Fig. 8, we can observe how the BT criterion stands out with 100 training samples, achieving good results with a very few samples, while BALD stands out with 600 training samples. However, BALD is the slowest method, being the random function the fastest one, while max-entropy, MI, and BT criterion exhibit similar execution times.

Focusing on SV, the best OA is reached by max-entropy, followed by MI and BALD. As we can see in Fig. 8,
Fig. 7. AL-based performance obtained by the 2-D B-CNN for different acquisition functions with different sizes of $D_{\text{train}}$. First column: results for the IP data set. Second column: results for the SV data set. Third column: results for the KSC data set.

Fig. 8. AL-based performance obtained by the 3-D B-CNN for different acquisition functions with different sizes of $D_{\text{train}}$. First column: results for the IP data set. Second column: results for the SV data set. Third column: results for the KSC data set.

BALD stands out with 100 training samples, until it is reached by MI and max-entropy. Again, BALD is the slowest method and random the fastest, while MI, max-entropy, and mean STD share the same computation time.

In the case of the KSC scene, all acquisition functions provide excellent classification performance, being random and mean STD the functions with the lowest OA (98.99% and 98.56%, respectively). In Fig. 8, we can
observe that the BT-criterion is able to reach high OA with a few training samples. Also, the BT-criterion, max-entropy, and MI exhibit similar execution times, being BALD and random the slowest and fastest acquisition functions, respectively.

5) Experiment 5 (Comparison With Other Traditional Classifiers): The fifth and final experiments perform a comparison between the AL implementations described in Sections III-C1–III-C4, with the best OA values for each hyperspectral data set, with traditional classifiers. For the IP data set, the AL-MLR and the 1-D B-CNN with the BT-criterion, 2-D B-CNN with the MI criterion, and 3-D B-CNN with the BALD criterion have been selected to be compared with the traditional RF, MLP, SVM, and MLR classifiers and also with the standard 1-D CNN, 2-D CNN, and 3-D CNN baselines, which have been implemented with the same parameters and architectures than the proposed AL approaches. In order to train the RF, MLP, SVM, and MLR, and 1-D CNN, 2-D CNN, and 3-D CNN baselines, only two pixels per class have been selected, while the remaining 800 pixels (the selected maximum size of $D_{train}$) have been randomly selected. This process has been repeated with the SV data set, selecting the AL-MLR and 2-D B-CNN with BT-criterion, 1-D B-CNN with MI, and 3-D B-CNN with max-
Focusing on the IP data set, we can observe the performance of pixelwise classifiers, being RF and the baseline MLR the classifiers that provide lower OA. We can observe how AL-MLR is better than the baseline MLR, but worse than SVM, while the spectral B-CNN model improves the classification results over the baseline 1-D CNN and the other pixelwise methods. Looking at spatial classifiers, the 2-D B-CNN is able to outperform the 2-D CNN results in 6.11 percentage points, improving also the generalization power. We can also observe this behavior with spectral–spatial classifiers, where the proposed B-CNN model outperforms the 3-D CNN baseline. Moreover, we can observe that, after adding spectral–spatial information, the classifier is able to improve its accuracy results. Fig. 9 presents these results in a graphical form, showing the classification maps obtained for each classifier. In Fig. 12, we can observe the performance of AL-MLR, 1-D B-CNN, 2-D B-CNN, and 3-D B-CNN with BT-criterion for the IP data set. We can see that the spectral–spatial B-CNN is able to reach a good classification accuracy with fewer training samples than the other AL-based classifiers, although it is the slowest method. Moreover, we can observe in Table VIII the number of training samples that each classifier needs to reach the accuracy percentage, being spectral–spatial B-CNN the one that needs...
The results for the SV data set are similar. In Table VII, we can see that the pixelwise classifiers based on AL outperform their baseline methods, being the AL-MLR better than MLR, while the spectral B-CNN is also better than the 1-D CNN baseline. Also, the spatial B-CNN outperforms the 2-D CNN baseline, being around 4.93 perceptual points better. Finally, the spectral–spatial B-CNN classifier is much better than the 3-D CNN baseline with 2.66 perceptual points better. In Fig. 10 shows the classification maps obtained by each classifier. Also, in Fig. 12, we can observe how B-CNNs are able to outperform the results of AL-MLR, standing out 150 training samples in the case of the 2-D B-CNN. In Table VIII, we can see that the spectral–spatial B-CNN needs less training data than the other classifier in order to reach 99% of accuracy.

The results obtained for the KSC data set are also quite similar to those obtained for the SV data set. In Table VII, we can see that the pixelwise classifiers based on AL outperform their respective baseline methods, as well as the RF, SVM, and MLP methods. Also, the spatial B-CNN model obtains better results than the baseline 2-D CNN, while the spectral–spatial B-CNN also outperforms the 3-D CNN baseline. These classification results can be observed in the graphical form in Fig. 11. In Fig. 12, we can observe in the third column the implemented AL-based methods with the BT-criterion as an acquisition function over the KSC data set. As we can see, the spectral–spatial B-CNN is able to reach good values with a few training samples, and in fact, this model can reach 99% accuracy with only 276 samples (i.e., 5.30% of the KSC’s ground truth), as shown in Table VIII.

### IV. Conclusion

In this paper, we have developed a new AL model with B-CNNs for hyperspectral image classification using spectral, spatial, and spectral–spatial features. The proposed approach offers robustness to overfitting on small labeled sets and improves the generalization capacity by including intelligently selected unlabeled training samples, integrating the spatial and the spectral information contained in the original hyperspectral image. To the best of our knowledge, this is the first time in the literature that AL is combined with CNNs (via BNNs) to perform robust hyperspectral image classification with very limited training sets. In this paper, we report very high classification accuracies using very limited labeled samples, avoiding the curse of dimensionality and the overfitting problems introduced by these kinds of networks. Our results also indicate that, by the proper selection of the acquisition function, AL offers a very good solution to avoid the aforementioned problems of overfitting with supervised deep networks. Future work will focus on improving the results obtained from the viewpoint of computational complexity, drawing additional comparisons with other established methods for spatial–spectral classification of remotely sensed hyperspectral and also validating the proposed techniques using multispectral data [101]. Finally, the inclusion of postprocessing methods, such as conditional random field [102], will also be studied in the future developments as a way to improve and smooth the classification maps obtained by the different tested methods.

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### References


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