A three-step classification framework to handle complex data distribution for radar UAV detection

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Abstract

Unmanned aerial vehicles (UAVs) have been used in a wide range of applications and become an increasingly important radar target. To better model radar data and to tackle the curse of dimensionality, a three-step classification framework is proposed for UAV detection. First we propose to utilize the greedy subspace clustering to handle potential outliers and the complex sample distribution of radar data. Parameters of the resulting multi-Gaussian model, especially the covariance matrices, could not be reliably estimated due to insufficient training samples and the high dimensionality. Thus, in the second step, a multi-Gaussian subspace reliability analysis is proposed to handle the unreliable feature dimensions of these covariance matrices. To address the challenges of classifying samples using the complex multi-Gaussian model and to fuse the distances of a sample to different clusters at different dimensionalities, a subspace-fusion scheme is proposed in the third step. The proposed approach is validated on a large benchmark dataset, which significantly outperforms the state-of-the-art approaches.

Keywords: radar UAV detection, micro-Doppler signature, greedy subspace
1. Introduction

Unmanned aerial vehicles have become an increasingly important radar target because of the low cost, wide applications and potential threats to public security. According to Grand View Research [1], the global market for commercial UAVs will grow by 17% every year. UAVs have been used for many different applications, e.g., package delivery, land surveillance, traffic monitoring and chasing birds in airport. However, UAVs may impose threats to public security, e.g., UAVs near airport may jeopardize the safety of airplanes [2], or UAVs may carry bombs or dangerous chemicals in a terrorist attack. Thus, it has become increasingly important to reliably detect UAVs using radars.

Early techniques using kinematic and radar cross-section characteristics [3] could not reliably differentiate UAVs from birds, as the both kinds are small and slow-moving targets. Many representations of micro-Doppler signature (mDS) have been explored, e.g., spectrogram [4–11], cepstrogram [12], cadence velocity diagram [13–15], others [16–19], and combinations of the aforementioned [20–22]. Particularly in [22], a rich source of features including spectrogram, cepstrogram and CVD are utilized. Most of the representations are closely related to spectrogram. mDS has been utilized in many radar-target-recognition tasks [23], e.g., airplane classification [24], ship detection [25], human detection [5], gait recognition [8, 9], action classification [10, 14] and vehicle classification [26]. Recently, Wi-Fi communication signals between UAVs and remote controllers have been utilized to detect UAVs [27]. However, when a UAV flies in an autonomous mode without Wi-Fi communication, such a technique will not work.

Machine-learning techniques have been utilized to automatically detect/classify UAVs using radars [6, 11, 15, 17–22, 28]. Artificial neural networks were applied on spectrum directly to classify different types of UAVs [28]. Support vector machine (SVM) and naive Bayes classifier were applied on the first five principal components extracted from spectrogram to differentiate UAVs from birds [6].
Huizing et al. employed Alexnet and LSTM-RNN on spectrograms to classify mini-UAVs [11], whereas Kim et al. utilized GoogLeNet on the image merged from spectrogram and CVD [20]. Similarly, SVM was applied on the feature vector obtained from spectrogram and CVD [21]. Patel et al. applied Alexnet on four time-frequency representations including spectrogram, cepstrogram and CVD for UAV classification [22]. Zhao and Su developed a cyclostationary analysis on the phase term of the radar signal to extract the mDS for UAV detection [18]. Very recently, empirical mode decomposition was employed to extract intrinsic mode functions for UAV classification [19]. Instead of detecting/classifying one UAV at a time, Zhang and Li detected multiple UAVs by using a k-means classifier on the mean CVD averaged along the Doppler frequency [15]. Most of these approaches utilized spectrogram or time-frequency representations that are derived from spectrogram, e.g., cepstrogram and CVD. Thus, the proposed approach also utilizes features derived from spectrogram. However, most approaches utilized the magnitude spectrogram only. As shown in [17], both phase and magnitude spectrograms are useful for classifying the radar signal.

The authors recently developed an automated UAV-detection system utilizing the regularized 2-D complex-log Fourier transform to extract spectrogram-like features and the subspace reliability analysis to remove unreliable feature dimensions [17]. Despite the success, three challenges remain. 1) The complex sample distribution of radar data. Subspace approaches utilizing up to the second-order statistics work well for Gaussianly distributed data [17, 29]. However, the high-dimensional mDS features deviate largely from Gaussian. 2) Outliers in radar data. Due to the poor signal-to-noise ratio of radar signal, it is error-prone for human to label the data, which leads to mislabeled data (outliers). The outliers are harmful for training classifiers. 3) The curse of dimensionality. It is difficult to robustly model the complex data distribution in a high-dimensional feature space.

In literature, these three challenges have been partially addressed. To model the complex distribution of radar data, Regev et al. utilized artificial neural
network to classify drones [28]. Zhao et al. utilized stacked auto-encoder and extreme learning machine for radar target recognition [30]. To be robust to the outliers in radar data, Dong et al. developed a joint sparse representation based on multi-task learning [31]. Many approaches have been devoted to address the curse of dimensionality. Specifically for radar target recognition, kernel joint discriminant analysis [32], sparse representation [31], subspace reliability analysis [17] and multiple kernel project subspace fusion [33] have been developed for dimensionality reduction. In this paper, an integrated three-step classification framework is proposed to address these three challenges.

In the first step, to handle the complex data distribution (Challenge 1) and the outliers (Challenge 2), the authors propose to utilize a greedy version of the sparse subspace clustering (SSC) algorithm [34, 35], the greedy subspace clustering (GSC) algorithm [36]. Gaussian mixture model (GMM) [37–39] is often used to model the complex data distribution, and the expectation-maximization (EM) algorithm [37] is often used to derive the mixture model. One critical challenge of the EM algorithm is that the GMM could not be reliably estimated due to insufficient training samples and the high feature dimensionality.

The sparse subspace clustering [34, 35] handles the complex distribution by clustering data according to the underlying subspace structure, which leads to a multi-Gaussian model if each cluster of samples follow the Gaussian distribution. The SSC algorithm is robust to outliers owing to the $l_1$ optimization when building the similarity matrix. As the SSC is slow, the authors propose to utilize the greedy subspace clustering [36]. Instead of the time-consuming $l_1$ optimization in the SSC, the GSC algorithm utilizes a nearest-subspace-neighbor algorithm to sequentially find the nearest neighbors to form linear subspaces. The neighborhood matrix is then used as the similarity matrix for subsequent spectral clustering. Similar outliers may form a cluster. Thus, a drop-off technique is proposed to remove samples in the smallest cluster as outliers.

In the second step, to tackle the curse of dimensionality (Challenge 3), a multi-Gaussian subspace reliability analysis (MGSRA) is proposed to remove the unreliable feature dimensions of the multi-Gaussian model derived in the first
step. The model cannot be reliably estimated due to insufficient samples in each
c cluster and the high dimensionality, especially the dimensions corresponding to
the small eigenvalues of covariance matrices. As the inverse of covariance matrix
is used to weigh the feature dimensions, those small eigenvalues will impose very
large and problematic weights to the corresponding dimensions [17, 29, 40].
Thus, the MGSRA algorithm is proposed to handle those unreliable feature
dimensions separately at different subspaces.

The proposed MGSRA is different from previous approaches [17, 41] in the
following aspects: 1) Most subspace approaches are designed based on a uni-
Gaussian model, whereas the MGSRA is built on a multi-Gaussian model, which
could better model the distribution of radar data. 2) Most subspace approaches
aim to find one linear subspace that meets a certain optimization criterion,
whereas the proposed MGSRA aims to find a set of linear subspaces separately
for each class. A problem thus arises naturally: how to optimally combine the
results from different subspaces?

In the third step, a subspace-fusion scheme is proposed to combine these
results. More specifically, the Mahalanobis distances of a sample to each cluster
center at a set of given feature dimensionalities are calculated. The rational of
choosing multiple dimensionalities is that it is difficult to determine the optimal
feature dimensionality for subspace approaches. Thus, a range of dimension-
alities covering the optimal one are sampled and the Mahalanobis distances at
these dimensionalities are evaluated. Then, the distances of a sample to different
cluster centers of different classes at different subspace dimensionalities are
treated as a feature vector, and a support vector machine is trained to combine
these distances. The proposed subspace fusion works better than traditional
approaches in which the distances are merged as a posterior probability, and
evaluated only at some fixed dimensionality for each class [37, 38].

The contributions of this study are summarized as follow: 1) Three chal-

lenes for radar UAV detection are identified: the complex data distribution,
the outliers and the curse of dimensionality. 2) A three-step classification frame-
work is proposed to address these challenges, i.e. a) the greedy subspace clus-
tering is utilized to handle the complex distribution and the outliers of radar data; b) a multi-Gaussian subspace reliability analysis is proposed to tackle the unreliable feature dimensions of the derived model; c) a subspace-fusion scheme is proposed to combine the subspace distances. 3) The proposed approach is systematically evaluated on a large benchmark dataset, and demonstrates a superior performance compared with the state-of-the-art approaches.

2. Proposed Three-Step Classification Framework

2.1. Challenges of UAV Detection

There are many challenges in detecting/classifying UAVs. Two of them, a robust feature representation and unreliable feature dimensions, were well addressed in the previous work [17]. Three others remain: the complex data distribution, the outliers and the curse of dimensionality.

2.1.1. Complex Sample Distribution of Radar Data

Subspace approaches often assume that data follow the Gaussian distribution [17, 29, 40–44], as the Gaussian model can be built using only mean and variance (covariance for multivariate Gaussian), which can be estimated easily from the data. However, in [29], Ren et al. showed that for visual recognition the histogram-like features do not follow the Gaussian distribution.

In this study, the authors find that it is insufficient to use a Gaussian distribution to model either the UAV class or the non-UAV class, as shown in Fig. 3 later in Section 7.2. This is primarily due to the following: 1) There are many different types of UAVs, e.g., helicopter, tricopter, quadcopter, hexacopter, octocopter and fixed-wing plane. One Gaussian distribution is not sufficient to model all these UAVs, especially the fixed-wing plane is significantly different from the rest. 2) The non-UAV class cannot be modeled as one Gaussian model either, as it consists of distinct background samples and bird samples. 3) Even for the same type of UAVs, data may not be Gaussianly distributed. All these lead to a complex data distribution.
2.1.2. Outliers in Radar Data

The radar micro-Doppler signatures are weak, much weaker than the main body Doppler. In addition, the thermal noise in a circuit and the noise/interference to radar receiver may contaminate radar signals. All these make it difficult to label radar targets. The labeling errors may come from different sources: 1) As the micro-Doppler signatures are weak, it is error-prone to manually label the data by analyzing the radar recordings; 2) The radar may capture the micro-motions of background objects, which will distort the radar signals of the target; 3) Due to the narrow radar beam, the target may fly in and out of the radar beam irregularly, without the notice of operators. 4) The mDS of fixed-wing plane is similar to that of gliding birds, which may be wrongly labeled as non-UAV if the UAV flies too far away and is not clearly visible in the video recordings. Some radar targets thus may be wrongly labeled, and known as outliers.

2.1.3. Curse of Dimensionality

The curse of dimensionality arises mainly due to high feature dimensionality and insufficient data. The feature representation used in the proposed approach, regularized 2-D complex-log Fourier transform [17], leads to a high-dimensional feature vector. (Refer to Table 7 for more information.) As a result, it is difficult to precisely model the complex data distribution using the limited number of samples in such a high-dimensional feature space.

More specifically, the high dimensionality leads to the following: 1) Traditional approaches such as the EM algorithm for GMM [37] do not work well here. The Gaussian mixture models are often either over-simplified or poorly estimated. The authors thus propose to utilize the greedy subspace clustering to find the underlying linear subspaces. 2) After clustering, it is still difficult to reliably estimate the multi-Gaussian model due to the high dimensionality, especially the dimensions corresponding to the small eigenvalues of the covariance matrices. Thus, a MGSRA algorithm is proposed to tackle these unreliable dimensions. 3) It is difficult to use the derived multi-Gaussian model to classify
samples in the high-dimensional feature space. Therefore, the authors propose to evaluate the Mahalanobis distances in the reduced subspaces at multiple dimensionalities and fuse them using the proposed subspace-fusion scheme.

2.2. Overview of the Proposed Approach

The block diagram of the proposed approach is shown in Fig. 1. The initial features are extracted by using the robust spectral analysis [17], and then the proposed three-step classification framework shown in Fig. 1 tackles the aforementioned three challenges. 1) To tackle the curse of dimensionality, among various clustering algorithms, the authors propose to utilize the greedy subspace clustering, as it could model the complex data distribution and handle the outliers at the same time. 2) The covariance matrices of the derived multi-Gaussian model are important but difficult to be reliably estimated. Thus, a multi-Gaussian subspace reliability analysis is proposed to tackle the unreliable feature dimensions of the covariance matrices. 3) Finally, a subspace-fusion scheme is proposed to evaluate the Mahalanobis distances of a sample to multiple cluster centers at different subspace dimensionalities. These distances are then fused by a support vector machine. In the following sections, the proposed approach will be illustrated in details.

3. Robust Spectral Analysis

The initial features are extracted using the regularized 2D complex-log-Fourier transform in [17]. The procedures are briefly summarized as follow:

Firstly, the time-series radar signal \( s(t) \) is segmented into \( I \) overlapping frames \( \{s_0, s_1, \ldots, s_{I-1}\} \), where \( s_i = \{s_i[n], n = 0, 1, \ldots, J-1\} \) is a vector of length \( J \). These \( I \) frames form a synthetic image \( S = [s_0, s_1, \ldots, s_{I-1}] \) of size \( I \times J \). The discrete Fourier transform \( f_i = [f_{i,0}, f_{i,1}, \ldots, f_{i,J-1}] \) of \( s_i \), \( f_i = \mathcal{F}\{s_i\} \), is computed as:

\[
f_{i,k} = \sum_{n=0}^{J-1} s_i[n] \exp(-j2\pi \frac{kn}{J}), \quad k = 0, 1, \ldots, J-1,
\]

(1)
where $\mathcal{F}\{\cdot\}$ denotes the discrete Fourier transform.

Secondly, the 2-D complex Fourier transform of $S$ is derived, which is equivalent to two 1-D spectral analysis on $S$:

$$
\mathcal{F}_{2D}\{S\} = \mathcal{F}_t\{\mathcal{F}\{S\}\},
$$

where $\mathcal{F}\{S\}$ is the Fourier transform on $S$, and $\mathcal{F}_t\{\cdot\}$ is the Fourier transform along the time axis. Previous approaches [12, 13] often utilize only the magnitude of $\mathcal{F}\{S\}$, whereas both phase and magnitude of $\mathcal{F}\{S\}$ are utilized in $\mathcal{F}_t\{\cdot\}$, because phase spectrums also carry important discriminant information for classification [17].

Thirdly, the weak micro-Doppler signatures are enhanced by taking the logarithm of the spectrum [17]. For $f_i = \mathcal{F}\{s_i\} = m_i \exp\{j\theta_i\}$,

$$
\log\{f_i\} = \log\{m_i\} + j\theta_i,
$$

where $m_i$ is the magnitude spectrum and $\theta_i$ is the phase spectrum. To balance the effects of $\log\{m_i\}$ and $\theta_i$, a weighting factor $w$ is introduced:

$$
\log\{f_i\} = \log\{m_i\} + jw\theta_i.
$$
$w$ is simply set as $w = 1/\pi$ so that the phase term is normalized to $[-1, 1]$.

Fourthly, a regularization term is introduced to Eqn. (4) to reduce the noise, because taking the logarithm not only enhances the weak micro-Doppler signature, but also enlarges the noise.

$$
\log \{ f_i \} = \log \{ m_i + c_i \} + jw\theta_i, \quad (5)
$$

where $c_i = \text{med} \{ m_i \}$, i.e. the median value of $m_i$. By adding such a regularization constant $c_i$, the logarithm of the strong frequency component will not be significantly altered, whereas the logarithm of the weak frequency component will be regularized close to $\log c_i$. The variations of noise frequency components are hence greatly reduced. Readers may refer to [17] for more details on the noise robustness of the robust spectral analysis.

Finally, the regularized 2-D complex-log-Fourier transform is derived as:

$$
F_R \{ S \} = F_t \{ \log \{ F \{ S \} \} \}, \quad (6)
$$

where $\log \{ F \{ S \} \}$ is calculated according to Eqn. (5).

4. Greedy Subspace Clustering

4.1. Limitations of EM Algorithm for Gaussian Mixture Model

The Gaussian mixture models [37, 38] have been widely used to handle complex data distributions. For $D$-dimensional feature $x \in \mathbb{R}^D$, the mixture probability density function (PDF) of the likelihood function is defined as:

$$
p(x|\Theta) = \sum_{i=1}^{M} \alpha_i p_i(x). \quad (7)
$$

This PDF is a weighted linear combination of $M$ Gaussian densities $p_i(x)$, each parameterized by a mean vector $\mu_i \in \mathbb{R}^D$ and a covariance matrix $\Sigma_i \in \mathbb{R}^{D \times D}$,

$$
p_i(x) = \frac{1}{(2\pi)^{D/2}|\Sigma_i|^{1/2}} \exp \left\{ -\frac{1}{2} (x - \mu_i)^T \Sigma_i^{-1} (x - \mu_i) \right\}. \quad (8)
$$

Collectively, the model is denoted as $\Theta = \{ \alpha_i, \mu_i, \Sigma_i \}$, for $i = 1, 2, \ldots, M$. 
The Expectation-Maximization algorithm [37] is often used to derive the mixture model. It starts with some initial estimation of \( \Theta \), and then updates \( \Theta \) by iteratively altering the following: 1) Estimate the membership weights for each sample; 2) Estimate the cluster weight, the mean and the covariance matrix of each cluster. Due to the curse of dimensionality, it is difficult to use the EM algorithm to build a precise mixture model for radar data. Hence the model is often simplified, e.g., by sharing the same covariance matrix among different classes and/or different mixture components, or by assuming that the covariance matrix is diagonal [37]. All these may oversimplify the model so that the discrimination power of the GMM is greatly reduced. To address this problem, the authors propose to utilize the greedy subspace clustering [36].

4.2. Motivations of Greedy Subspace Clustering

The greedy subspace clustering [36] is a greedy version of the sparse subspace clustering [35, 45]. The subspace clustering problem is formally defined as: Given data points \( \{ \mathbf{y}_i \in \mathbb{R}^D \}_{i=1}^N \) drawn from a union of independent linear subspaces \( \{ \mathcal{S}_i \}_{i=1}^n \), the target is to find dimensions \( \{ d_i \}_{i=1}^n \), subspace bases \( \{ \mathbf{A}_i \in \mathbb{R}^{D \times d_i} \}_{i=1}^n \) and permutation matrix \( \mathbf{P} \in \mathbb{R}^{N \times N} \) that segment the data,

\[
\mathbf{Y} = [\mathbf{y}_1, \mathbf{y}_2, \ldots, \mathbf{y}_N] = [\mathbf{Y}_1, \mathbf{Y}_2, \ldots, \mathbf{Y}_n] \mathbf{P},
\]

(9)

where \( \mathbf{Y}_i \in \mathbb{R}^{D \times N_i} \) are \( N_i \) data points drawn from \( \mathcal{S}_i \) and \( N = \sum_{i=1}^n N_i \). If \( \mathbf{y} \) is a new data point in \( \mathcal{S}_i \), it can be represented as a linear combination of the \( d_i \) points in the same subspace.

Let \( \hat{\mathbf{Y}}_i \in \mathbb{R}^{D \times (N-1)} \) denote the matrix obtained from \( \mathbf{Y} \) by removing the \( i \)-th column \( \mathbf{y}_i \), where \( \hat{i} \) means “not \( i \)”. \( \mathbf{c}_i \in \mathbb{R}^{N-1} \) derived by solving the \( l_1 \) optimization problem,

\[
\arg\min_{\mathbf{c}_i} \| \mathbf{c}_i \|_1 \text{ subject to } \mathbf{y}_i = \hat{\mathbf{Y}}_i \mathbf{c}_i,
\]

(10)

is a vector whose nonzero entries correspond to the points in \( \hat{\mathbf{Y}}_i \) lying in the same subspace as \( \mathbf{y}_i \). By inserting a zero entry at the \( i \)-th row of \( \mathbf{c}_i \), it becomes
an $N$-dimensional vector $\mathbf{c}_i \in \mathbb{R}^N$. The $l_1$ optimization is repeated for every $y_i$, $i = 1, 2, \ldots, N$. Then, the following coefficient matrix is obtained:

$$C = [\mathbf{c}_1, \mathbf{c}_2, \ldots, \mathbf{c}_N] \in \mathbb{R}^{N \times N},$$

(11)

which can be seen as the similarity matrix for $\mathbf{Y}$. Then, the spectral clustering algorithm [36] is applied on $C$ to segment the data.

The sparse subspace clustering is robust to noise and outliers owing to the $l_1$ optimization, but $l_1$ optimization is slow. On the collected UAV-detection dataset consisting of more than 10,000 training samples of 7236 dimensions, it takes more than 300 seconds for the $l_1$ optimization of one sample. The total execution time for all samples is about 35 days, which is too long. In addition, memory of a few gigabytes is required for each $l_1$ optimization, and hence parallel computing using a graphic card is not a feasible option. These are the motivations of using the greedy subspace clustering [36].

4.3. Nearest-Subspace-Neighbor Algorithm

The greedy subspace clustering [36] utilizes a nearest-subspace-neighbor (NSN) algorithm to sequentially find nearest neighbors to the subspace spanned by point $y_i$ and existing neighbors. The spectral clustering algorithm [36] is then applied on the neighborhood matrix for clustering. Formally, let $\mathcal{I}_i$ denote the set of neighbors for data point $y_i$, $[N]$ denote the set $\{1, 2, \ldots, N\}$, $\mathcal{U}$ denote the subspace spanned by the set of neighbors $\mathcal{I}_i$, $\mathcal{U}$ denote the set of orthonormal bases of subspace $\mathcal{U}$, and $\mathbf{I}\{\cdot\}$ denote an indicator function, which is one if the statement is true and zero otherwise. The NSN algorithm is summarized in Algorithm 1.

By initializing $\mathcal{I}_i \leftarrow \{i\}$, the first neighbor is chosen as $y_1$. The NSN algorithm then finds $K$ neighbors sequentially. At step $k$, a $k$-dimensional subspace $\mathcal{U}$ spanned by $y_i$ and the $k-1$ neighbors is constructed, and the point closest to the subspace is selected. After $k > k_{\text{max}}$, the subspace $\mathcal{U}$ constructed at step $k_{\text{max}}$ is used, and the points closest to the subspace $\mathcal{U}$ are chosen as neighbors for the rest of the process.
Algorithm 1 Nearest-subspace-neighbor algorithm

Input: A collection of data points \( \{y_i \in \mathbb{R}^D\}_{i=1}^N \), the number of expected neighbors \( K \), and maximum subspace dimension \( k_{\text{max}} \).

Output: A neighborhood matrix \( W \in \{0, 1\}^{N \times N} \).

\( y_i \leftarrow y_i / \|y_i\|_2, \forall i \in [N]. \quad \triangleright \text{Normalized to unit variance.} \)

\textbf{for} \( i = 1, 2, \ldots, N \) \textbf{do} \quad \triangleright \text{Run NSN for each data point.}

\( I_i \leftarrow \{i\}. \quad \triangleright \text{Initialize } I_i \text{ as } \{i\}. \)

\textbf{for} \( k = 1, 2, \ldots, K \) \textbf{do} \quad \triangleright \text{Iteratively add the closest point.}

\textbf{if} \( k \leq k_{\text{max}} \) \textbf{then}

\( U \leftarrow S\{y_j, j \in I_i\}. \quad \triangleright \text{Construct the subspace spanned by } y_j. \)

\textbf{end if}

\( j^* \leftarrow \arg \max_{j \in [N] \setminus I_i} \|U^T y_j\|_2. \quad \triangleright j^* \text{ is the nearest neighbor to } U. \)

\( I_i \leftarrow I_i \cup \{j^*\}. \quad \triangleright \text{Add } j^* \text{ to the set of nearest neighbors.} \)

\textbf{end for}

\( W_{ij} \leftarrow 1\{j \in I_i \text{ or } y_j \in U\}. \quad \triangleright \text{Construct the neighborhood matrix.} \)

\textbf{end for}
A brief time complexity analysis of the NSN algorithm is presented as follows: At step $k$ of the inner loop, the time complexity is $O(D^2 + k^3)$ to derive the spanned subspace $\mathcal{U}$ using Singular Value Decomposition. The time for $\arg\max_{j \in [N]} \|U^T y_j\|_2$ is $O(kDN)$. Thus, the time for the most inner loop in Algorithm 1 is $O(D^2 + k^3 + kDN)$. The time complexity for the whole algorithm is $O(N(KD^2 + \sum_{k=1}^K k^3 + \sum_{k=1}^K kDN)) = O(NKd^2 + NK^4 + NDK^2)$. Note that when $K \ll D$, the time complexity can be simplified as $O(NKD^2)$.

The GSC algorithm clusters data according to the underlying subspace structure. As a result, a multi-Gaussian model is derived for each class. The proposed MGSRA algorithm then removes the unreliable feature dimensions of the derived model, as illustrated in Section 5.

4.4. Outlier Removal by Cluster Drop-off

Although the GSC algorithm is robust to outliers to some extent, mislabeled data may be similar to each other and form a cluster, e.g., the UAV may fly out of the sight of a radar, but the radar recordings may be mislabeled as UAV samples, and these similar outliers may form a cluster. To tackle this problem, a simple heuristic is proposed to remove the outliers, i.e., the cluster with the smallest number of samples for each class is dropped off. In general, the mislabeled data only account for a small portion of the dataset. If they form a cluster, most likely they form the smallest cluster.

5. Multi-Gaussian Subspace Reliability Analysis

In [17], the subspace reliability analysis was utilized to remove the unreliable feature dimensions in the UAV and non-UAV classes separately in two different subspaces. The samples of each class are assumed to follow the Gaussian distribution. However, one Gaussian is not sufficient to model the complex data distribution. The authors thus propose to utilize the greedy subspace clustering to find the underlying subspace structure, as shown in the previous section, which naturally leads to a multi-Gaussian model.
The derived model could not be reliably estimated due to the curse of dimensionality. For the $i$-th cluster of the $j$-th class, the PDF of the Gaussian model is given as:

$$p_{ij}(x) = \frac{1}{(2\pi)^{D/2}|\Sigma_{ij}|^{1/2}} \exp \left\{ -\frac{1}{2}(x - \mu_{ij})^T \Sigma_{ij}^{-1}(x - \mu_{ij}) \right\},$$  \tag{12}

where $\mu_{ij} \in \mathbb{R}^D$ and $\Sigma_{ij} \in \mathbb{R}^{D \times D}$ are the mean vector and the covariance matrix for the $i$-th cluster of the $j$-th class, respectively. The key issue here is to reliably estimate the covariance matrices $\Sigma_{ij} \in \mathbb{R}^{D \times D}$ so that the Mahalanobis distance $(x - \mu_{ij})^T \Sigma_{ij}^{-1}(x - \mu_{ij})$ could be evaluated reliably. The small eigenvalues of the covariance matrices could not be reliably estimated. As the inverse of $\Sigma_{ij}$ is used to weigh the feature dimensions, those small eigenvalues are harmful for classification [40]. If the number of samples $N < D$, some eigenvalues of $\Sigma_{ij}$ will be zero and induce infinitely large weights. Even in the case that $\Sigma_{ij}$ has full rank, the small eigenvalues of $\Sigma_{ij}$ still cause trouble, as their inverses introduce problematic large weights to the feature dimensions. To tackle this problem, a multi-Gaussian subspace reliability analysis is proposed.

Denote the Mahalanobis distance of $x$ to the $i$-th cluster of the $j$-th class as

$$d_{ij}(x) = \frac{1}{2}(x - \mu_{ij})^T \Sigma_{ij}^{-1}(x - \mu_{ij}).$$  \tag{13}

The targets are to remove the small eigenvalues of $\Sigma_{ij}$ so that $d_{ij}(x)$ could be evaluated reliably, and to preserve the discriminant information among different classes, which mainly resides in the between-class scatter matrix

$$\Sigma_b = \sum_{j=1}^{c} (\mu_j - \mu)(\mu_j - \mu)^T,$$  \tag{14}

where $\mu_j$ is the mean vector for the $j$-th class, $\mu$ is the global mean and the number of class $c = 2$ for UAV detection. To remove the unreliable feature dimensions of $\Sigma_{ij}$, and preserve the discriminant information in $\Sigma_b$, the eigen-decomposition is applied on $S_{ij} = \Sigma_{ij} + \Sigma_b$ as:

$$S_{ij} = \Phi_{ij} \Lambda_{ij} \Phi_{ij}^T,$$  \tag{15}
where $\Phi_{ij}$ and $\Lambda_{ij}$ are the eigenvector and eigenvalue matrices of $S_{ij}$, respectively. Then, the eigenvectors are chosen corresponding to the leading $m$ eigenvalues of $S_{ij}$, i.e., $\Phi_{ijm}$, as the projection matrix. The Mahalanobis distance $d_{ij}(x)$ in the projected $m$-dimensional subspace becomes:

$$
d_{ijm}(x) = (x - \mu_{ij})^T \Phi_{ijm} (\Phi_{ijm}^T \Sigma_{ij} \Phi_{ijm})^{-1} \Phi_{ijm} (x - \mu_{ij}).$$  \hspace{1cm} (16)

The optimal feature dimensionality cannot be easily determined. Thus, many subspace approaches report the classification accuracies at different dimensionalities to show how the accuracies vary with the dimensionality, without determining the optimal dimensionality in advance. In the proposed approach, the distances are evaluated at a range of dimensionalities that probably will cover the optimal one. As these distances are evaluated in different subspaces, their scalings are different, and they should be properly weighted before fusion. Most importantly, a proper classification scheme needs to be developed for the derived multi-Gaussian model. To address these challenges, a subspace-fusion scheme is proposed as illustrated in the next section.

6. Proposed Subspace-Fusion Scheme

The proposed subspace-fusion scheme aims to combine the Mahalanobis distances defined in Eq. (16) and to build a classifier for the derived multi-Gaussian model. These Mahalanobis distances form a feature vector,

$$
d(x_k) = [d_{ijm}(x_k)],$$  \hspace{1cm} (17)

where $i = 1, 2, \ldots, M_j$ is the index of clusters, $j = 1, 2, \ldots, c$ is the index of classes, and $m$ is the index of subspace dimensionalities. Assume that all classes have the same number of clusters $M$, and the Mahalanobis distances are evaluated at $L$ different subspace dimensionalities for each cluster, the distance vector $d(x_k) \in \mathbb{R}^{M \times L}$. A support vector machine is trained using the derived feature vector $d(x_k), k = 1, 2, \ldots, N$. For a new testing sample $t$, $d(t)$ is evaluated and the class label is predicted using the trained SVM. The proposed subspace-fusion scheme has the following advantages:
1. For many subspace approaches [17, 29, 40–44], it is difficult to find the optimal feature dimensionality. In the proposed approach, the Mahalanobis distances are evaluated at a given set of dimensionalities, without the need of selecting the optimal dimensionality. It is much easier to choose a range of dimensionalities covering the optimal one, than to precisely determine it in advance.

2. The proposed scheme determines the optimal weights for the Mahalanobis distances through the trained SVM, which solves the problem of optimally combining these distances. It definitely outperforms other approaches in which the distances are evaluated at a single (optimal) dimensionality [37].

3. The proposed scheme addresses the challenge of developing a proper classifier for the multi-Gaussian model. Traditional maximum-a-posterior classifier for the GMM [37] cannot work properly here as the posterior probabilities cannot be reliably estimated due to the curse of dimensionality. The proposed approach tackles the problem by evaluating the distances in many reduced subspaces, and fusing them using a support vector machine.

7. Experimental Evaluation

7.1. Experimental Setup

The measurement data were collected by Thales using a low-power continuous-wave radar operating at X-band (9.7 GHz radio frequency). Some signals were sampled at 192 kHz and others at 96kHz. They are all normalized to 96kHz in the experiments. A horn antenna was manually adjusted toward the nearby target object. Bird samples were collected within the distance of 5-50 meters to the radar, and UAV samples were collected within the distance of 3-150 meters to the radar.

The dataset used in [17] consists of multiple radar recordings of UAVs and birds, varying in length. The total length of all recordings reaches 1058 seconds, including 854 seconds of UAV recordings and 204 seconds of non-UAV recordings. To better evaluate the performance of the proposed system, the dataset
is extended using additional data provided by Thales. The extended dataset consists of 48 radar recordings, including 2087-second recordings of UAVs and 322-second recordings of non-UAVs. The dataset covers a wide range of UAVs such as single-rotor, multi-rotor and fixed-wing types, and non-UAVs including background and targets most similar to UAVs such as birds. Hence the data distribution well represents the data population for UAV detection after filtering out targets obviously different from UAVs by other means. The model built on the dataset could be applied in practical scenarios. The sample spectrograms of UAVs and birds are shown in Fig. 2.

![Spectrograms](image)

(a) Spectrogram of Easystar glider  
(b) Spectrogram of Trex 450 helicopter  
(c) Spectrogram of a bird  
(d) Spectrogram of a group of birds  

Figure 2: Sample spectrograms of UAVs and birds.

As a longer duration is needed for the dynamic time warping (DTW) [4], the recordings are chopped into 1-second samples when evaluating DTW. For other approaches, the recordings are chopped into 50-ms samples, unless otherwise
stated. As a result, there are in total 2409 samples when evaluating DTW, and 48180 samples when evaluating others. This is a relatively large dataset for a two-class classification problem. Half of the dataset is randomly chosen as training samples, and the other half is chosen as testing samples. The experiments are repeated 10 times, and the average performance is reported.

Denote the number of UAV samples being correctly classified and falsely classified as $n_{TP}$ and $n_{FN}$, and the number of non-UAV samples being correctly classified and falsely classified as $n_{TN}$ and $n_{FP}$, respectively. The false acceptance rate (FAR) and the false rejection rate (FRR) are defined as follow:

\[
FAR = \frac{n_{FP}}{n_{FP} + n_{TN}},
\]

\[
FRR = \frac{n_{FN}}{n_{FN} + n_{TP}}.
\]

By varying the decision threshold, different combinations of FAR and FRR could be derived. When these two error rates are the same, it is defined as the equal error rate (EER). Three evaluation criteria are reported in this paper: EER, the FAR at the FRR of 1% (denoted as $FAR_{FRR=1\%}$) and the FAR at the FFR of 0.1% (denoted as $FAR_{FRR=0.1\%}$). These three criteria are chosen because: 1) The EER is commonly used in detection tasks. 2) To evaluate how the system performs at a low missing detection rate of UAVs (i.e., a low FRR), $FAR_{FRR=1\%}$ and $FAR_{FRR=0.1\%}$ are reported.

The regularized 2-D complex-log Fourier transform [17] is utilized as the initial feature representation, in which the spectrum utilizes 256 data points and the windows have 50% overlapping. After removing the clutter and some unreliable high-frequency components, the initial feature vectors have $201 \times 36 = 7236$ dimensions for 50-ms samples.

7.2. Analysis of Data Distribution

The distribution of the dataset is examined how far it deviates from the Gaussian. The feature vectors have 7236 dimensions. It is infeasible to visualize the data distribution in such a high-dimensional space. Thus, the first two principal components of the UAV/non-UAV class are extracted. Then,
The feature dimension is 7236. The first two principal components are extracted and used to plot the distribution of UAV/non-UAV samples in the subspace built from the first two principal components, and the clustering results with 3 and 4 clusters for UAV and non-UAV classes, respectively. The figure shows that the distributions of both UAV class and non-UAV class are far away from the Gaussian.

Clustering results using the GSC algorithm with 3 and 4 clusters for the UAV class and the non-UAV class are plotted respectively in the subspace built from the first two principal components, as shown in Fig. 3. Take note that the results are plotted in two different subspaces, as they utilize the first two principal components of UAV samples and non-UAV samples, respectively.

The following can be observed from the plots:

1. The distributions of both UAV and non-UAV classes are far away from
the Gaussian. This is consistent with the previous analysis that neither UAV nor non-UAV samples follow the Gaussian distribution.  

2. The data distribution of the UAV class is difficult to model, as there are many different kinds of UAVs. In addition, due to the pose variations, the micro-Doppler signatures of UAVs may appear very different.  

3. For the non-UAV class, there are roughly two clusters, which correspond to background samples and bird samples, two main types of non-UAVs in the current dataset.  

4. For both UAV and non-UAV samples, there are some outliers, which are far away from any cluster. Both the complex data distribution and the outliers are the motivations of using the greedy subspace clustering to handle these two challenges.  

7.3. **Comparison to State-of-the-Art Approaches**  

In literature, not many approaches are specifically designed for classifying UAVs from birds, except support vector machine on the integrated feature vector derived from spectrogram and cadence velocity diagram [21] and the authors’ previous approach [17]. The dynamic time warping [4] and the robust principal component analysis (PCA) [7] are two recent approaches published in reputable journals, but designed for other radar-target-recognition tasks. They are hence modified for UAV detection and compared with the proposed approach.  

7.3.1. **Classification Results Using Dynamic Time Warping**  

The dynamic time warping\(^1\) [4] is applied on the spectrogram to align the possible time variations. The recordings are chopped into 1-second samples due to computational complexity constraints. The optimal path derived by DTW is treated as the distance between two samples. The distances from one sample to all others are treated as the feature vector. A linear support vector machine with

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\(^1\)The matlab code of DTW can be downloaded from [http://labrosa.ee.columbia.edu/matlab/dtw/](http://labrosa.ee.columbia.edu/matlab/dtw/).
the cost parameter $C = 40$ is trained as the classifier. The average classification errors over 10 trials are shown in Table 1.

| EER    | $FAR_{FRR=1\%}$ | $FAR_{FRR=0.1\%}$ |
|--------|-----------------|-------------------|
| 18.69% | 92.93%          | 99.36%            |

Table 1: Classification errors using the dynamic time warping [4].

It can be seen that the dynamic time warping does not well solve the UAV-detection problem. The equal error rate is quite high. If the false rejection rate is expected to be low, the false acceptance rate is very high, as high as 92.93%.

7.3.2. Classification Results Using Robust PCA

The same procedures as in [7, 17] are utilized to implement the robust PCA. The feature vector is obtained by averaging the spectrogram over time. The minimum covariance determinant (MCD) estimator implemented using “rrcov” package in R programming is used to remove the outliers. PCA is then used to reduce the feature dimensionality. Finally, the feature vectors are normalized to zero mean with unit variance, and classified by a linear support vector machine with the cost parameter $C = 40$.

The error rates at various dimensionalities are shown in Fig. 4. These three figures follow the same trend, i.e., the error rates at very low dimensionality are high, drop with increasing dimensionality, and stabilize at high dimensionality. The lowest error rates in these three figures are achieved at 80 dimensions. The error rates at this optimal dimensionality are shown in Table 2. The robust PCA performs better than the DTW.

| EER    | $FAR_{FRR=1\%}$ | $FAR_{FRR=0.1\%}$ |
|--------|-----------------|-------------------|
| 10.15% | 53.41%          | 81.34%            |

Table 2: Classification performance using the robust PCA [7].
7.3.3. Support Vector Machine on Spectrogram and Cadence Velocity Diagram

A support vector machine was applied on spectrogram and cadence velocity diagram to differentiate UAVs from birds [21]. The implementation described in [21] is strictly followed. The feature vector consists of three parts: 1) The average spectrogram over time; 2) The first left singular vector of the spectrogram after the Singular Value Decomposition on the spectrogram; 3) The average cadence velocity diagram over cadence frequency. These three are then concatenated as the final feature vector and classified by a linear support vector machine with the cost parameter $C = 40$. This approach is named as SVM-S-CVD. The classification results are shown in Table 3. SVM-S-CVD performs better than DTW and RPCA, as it utilizes both spectrogram and CVD.

| Feature | EER | $\text{FAR}_{FRR}=1\%$ | $\text{FAR}_{FRR}=0.1\%$ |
|---------|-----|----------------------|----------------------|
| EER    | 7.46\% | 41.30\% | 90.24\% |

Table 3: The error rates for a support vector machine on the integrated feature vector extracted from spectrogram and cadence velocity diagram [21].

7.3.4. Performance Evaluation of Proposed Approach

Now the proposed approach (greedy subspace clustering, multi-Gaussian subspace reliability analysis and subspace-fusion scheme, denoted as GSC-MGSRA-SF) is compared with the subspace reliability analysis based on the uni-Gaussian model (denoted as SRA) [17]. Both utilize the robust spectral analysis in Sec-
tion 3 to extract the initial feature representation. In addition, GMM-MGSRA-SF is implemented to evaluate the performance gain over the GMM, where GSC is replaced by the Gaussian mixture model and the rest procedures are the same as the proposed approach.

**SRA:** The subspace reliability analysis is applied on two covariance matrices separately in two different subspaces, and a ratio test is employed to differentiate UAVs from non-UAVs. The dimensionality of these two subspaces is reduced using SRA to \{1, 2, \ldots, 10, 20, 30, \ldots, 200\}, respectively, and the performance at the optimal combination of these two dimensionalities is reported.

**Proposed GSC-MGSRA-SF:** As SRA is evaluated for at most 200 dimensions, the principal component analysis is applied to reduce the dimensionality from 7236 to 200 for a fair comparison. Then, the greedy subspace clustering is applied on the UAV class and the non-UAV class, respectively. To evaluate how the performance varies with the number of clusters, the number of clusters is explicitly chosen as 5, 10 and 20 for both classes. For the nearest-subspace-neighbor algorithm [36], the number of nearest neighbors is set as 40\(^2\) and the number of feature dimensions of the linear subspace is set to the default value 20. To remove the outliers that may form a cluster, the samples in the smallest cluster for each class are removed.

The Mahalanobis distances at dimensions of \{1, 2, \ldots, 10, 20, 30, \ldots, 200\} are evaluated for both UAV and non-UAV classes. These dimensionalities are well spread across the possible optimal dimensionality within 200. These distances are treated as the feature vector and classified by a linear support vector machine. The cost parameter for the SVM is explicitly set to 20. The proposed approach is denoted as GSC-MGSRA-SF-M, where \(M\) denotes the number of clusters used.

**GMM-MGSRA-SF:** To show the performance gain against the Gaussian mix-

\(^2\)The default value is 20. As there are thousands of samples, the similarity matrix is large. It will lead to a very sparse similarity matrix if utilizing only 20 nearest neighbors, and lead to numerical instability for the subsequent spectral clustering. Thus it increases to 40.
ture model, the GSC algorithm is replaced by GMM and the rest procedures are kept the same as the proposed approach. Due to the curse of dimensionality, the same diagonal matrix is used for all clusters of each class when building the mixture model using the EM algorithm. The number of mixture components is set to 5 for each class. The experimental results are summarized in Table 4.

| Method                  | EER  | FAR_{FRR=1\%} | FAR_{FRR=0.1\%} |
|-------------------------|------|---------------|-----------------|
| SRA [17]                | 5.56%| 25.20%        | 44.80%          |
| GMM-MGSRA-SF            | 4.76%| 16.15%        | 43.52%          |
| GSC-MGSRA-SF-5          | 3.95%| 14.50%        | 44.37%          |
| GSC-MGSRA-SF-10         | 3.13%| 7.91%         | 40.77%          |
| GSC-MGSRA-SF-20         | 3.05%| 7.55%         | 30.01%          |

Table 4: Comparison to SRA and GMM-MGSRA-SF, and evaluation of the proposed approach on different number of clusters used in the greedy subspace clustering algorithm.

The following can be observed from Table 4. The proposed GSC-MGSRA-SF outperforms SRA, which shows the advantages of modeling the complex data distribution as the multi-Gaussian model over the uni-Gaussian one. The proposed approach also outperforms GMM-MGSRA-SF, which shows the effectiveness of the GSC algorithm over the Gaussian mixture model. In general, the error rates decrease with the increase of the number of clusters used in GSC-MGSRA-SF. The multi-Gaussian model better models the complex data distribution by using more clusters. The performance gain becomes marginal when the number of clusters is large. When more clusters are used, the number of samples falling into each cluster becomes smaller. Thus, the distribution of each cluster may not be well estimated using a limited number of samples. As a result, the performance gain is small, or the performance may even drop if the number of clusters increases further. For the rest of experiments, the number of clusters is set to 20.

The performance comparisons to the state-of-the-art approaches are summarized in Table 5. The proposed approach significantly outperforms the others.
| Method                  | EER   | $FAR_{FRR=1\%}$ | $FAR_{FRR=0.1\%}$ |
|------------------------|-------|------------------|-------------------|
| DTW [4]                | 18.69%| 92.93%           | 99.36%            |
| RPCA [7]               | 10.15%| 53.41%           | 81.34%            |
| SVM-S-CVD [21]         | 7.46% | 41.30%           | 90.24%            |
| SRA [17]               | 5.56% | 25.20%           | 44.80%            |
| GMM-MGSRA-SF           | 4.76% | 16.15%           | 43.52%            |
| Proposed GSC-MGSRA-SF  | 3.05% | 7.55%            | 30.01%            |

Table 5: Summary of the comparisons to the state-of-the-art approaches.

### 7.4. Performance Evaluation on Noise Robustness

Noise is injected into the radar return signal to evaluate the noise robustness of the proposed approach. The signal-to-noise ratio (SNR) is calculated as:

$$SNR = 10 \log_{10} \left( \frac{P_x}{P_n} \right),$$  \hspace{1cm} (20)

where $P_n$ is the power of the injected Gaussian noise and $P_x$ is the power of the radar signal after removing the clutter. Gaussian noise is used as it is one of the most common noise types. The clutter is removed before injecting the noise as it is not relevant to the radar target but much stronger than the Doppler signatures. Note that the main body Doppler is much stronger than the micro-Doppler signatures. Thus, the actual SNR w.r.t. mDS is much lower than the reported SNR. The error rates for different SNRs are summarized in Table 6.

Table 6 shows that when the noise is small or even comparable to the micro-Doppler signatures, the proposed approach achieves a fairly good performance. The error rates do not change significantly when the noise level is low. The proposed approach is shown robust to noise. Even when the noise level is high, the error rates of the proposed approach remain at a reasonable level.

### 7.5. Performance Evaluation by Varying Observation Durations

Here, the proposed approach is evaluated for various observation durations. Intuitively, if there is a longer observation duration, more information about the
| SNR | EER  | $FAR_{FRR=1\%}$ | $FAR_{FRR=0.1\%}$ |
|-----|------|-----------------|-------------------|
| -10 | 12.79% | 39.12% | 66.35% |
| 0   | 5.87%  | 21.56% | 57.79% |
| 10  | 3.37%  | 8.39%  | 47.35% |
| 20  | 3.10%  | 8.10%  | 51.81% |
| clean | 3.05% | 7.55% | 30.01% |

Table 6: The classification errors vs. different SNRs. The proposed approach is robust to noise to a large extent.

radar target can be obtained, and hence a higher classification accuracy can be achieved, but the extracted feature vector will become larger.

In the previous experiments, the observation duration is set as 50 ms, as suggested by Thales. In this experiment, the system is evaluated for the observation durations of 10, 25, 50, 100 and 200 ms. Table 7 summarizes the initial feature dimensionality and the number of samples for different observation durations. If the number of samples is large, the NSN algorithm used in the greedy subspace clustering will take a long time to execute as it needs to loop through all samples to find the nearest subspace neighbors, and the subsequent spectral clustering requires a large amount of memory and a long execution time. On the other hand, if the initial feature dimensionality is large, the covariance matrix of the initial feature vector will be large and require a huge amount of memory.

The error rates and the average execution time of one sample vs. the observation durations are shown in Table 8. The proposed approach is trained and tested on a Dell PC with Intel Xeon Silver 4108 CPU @1.80 GHz. The experimental results are consistent with the previous analysis, i.e., longer the observation window, better the classification performance. When the duration is very short, e.g., 10 ms, the classification errors significantly increase because not even one full rotation cycle of the rotor blade of a UAV could be captured in such a short time. The intra-class variations of the UAV class greatly increase,
Observation duration (ms) | Initial feature dimensionality | The number of samples
---|---|---
10 | 1,206 | 240,900
25 | 3,417 | 96,360
50 | 7,236 | 48,180
100 | 14,874 | 24,090
200 | 29,949 | 12,045

Table 7: The initial feature dimensionality and the number of samples vs. the observation durations. If the duration is long, the feature dimensionality will be large. On the other hand, if the duration is short, the number of samples will be large.

and hence the error rates significantly increase. On the other hand, when the observation duration is sufficiently long, the error rates do not significantly decrease with a further increase of the duration, e.g., the performance gain from 100 ms to 200 ms is not as significant as others. In terms of execution time, although the proposed approach looks complicated, it could predict one sample in real time.

| Observation duration (ms) | EER | \(FAR_{FRR=1\%}\) | \(FAR_{FRR=0.1\%}\) | Execution time (ms) |
|---|---|---|---|---|
| 10 | 8.45% | 37.87% | 65.39% | 3.61 |
| 25 | 4.20% | 15.08% | 56.42% | 4.06 |
| 50 | 3.05% | 7.55% | 30.01% | 4.76 |
| 100 | 2.02% | 3.50% | 22.57% | 7.99 |
| 200 | **1.72\%** | **2.31\%** | **13.41\%** | **20.81** |

Table 8: Classification error rates and average prediction time of one test sample for different observation durations.

7.6. Demo of UAV Detection

A live demo for UAV detection is implemented using Matlab. The demo GUI is shown in Fig. 5. The sub-figure shown at the top is the spectrogram of
the radar signal. The sub-figure on the bottom-left is the video capture of the
target and the real-time classification result for the current sample of 50 ms is
shown on the right. The cumulative classification results as UAVs or non-UAVs,
and the classification time for the current sample are shown in bottom-middle.
Once the demo starts execution, the user will be prompted to choose a radar
recording for analysis. The overlapping ratio of 50% is preset in the demo, as
the model needs to be re-trained if the key parameters change. The overlapping
ratio controls the time resolution of the spectrogram. Larger overlapping ratio
means higher resolution and hence higher classification accuracy at a cost of
higher computational complexity. For a speed-accuracy trade-off, it is set to
50%. It takes about 15.33 ms to classify a sample using Matlab 2019a, on a
Dell PC with Intel Xeon Silver 4108 CPU @1.80 GHz. This demo shows that
the proposed system can detect UAVs reliably in real time.

![Spectrogram](image)

Figure 5: GUI of the UAV-detection demo. It shows that the proposed system could detect
UAVs reliably in real time.
8. Conclusion

In this paper, a three-step classification framework is proposed to address three challenges in radar UAV detection: outliers in the data, complex data distribution and the curse of dimensionality. In the first step, the authors propose to utilize the greedy subspace clustering to handle the outliers and model the complex data distribution. The expectation-maximization algorithm to derive the Gaussian mixture model could not well cluster the samples due to the curse of dimensionality. To circumvent this problem, a multi-Gaussian subspace reliability analysis is proposed in the second step to handle the unreliable feature dimensions of the derived multi-Gaussian model. In the third step, a subspace-fusion scheme is proposed to combine the distances of a sample to different clusters of different classes at different dimensionalities. The proposed system is compared with existing approaches on a large benchmark dataset, and significantly outperforms the state-of-the-art approaches.

The proposed three-step classification framework could well handle the complex distribution of radar data. However, a potential problem here is that the model in the early stage is optimized without considering the later ones. The future plan is to integrate these three steps as one unified algorithm, e.g., considering the reliability of newly added subspace during the GSC algorithm. The second potential research direction is to integrate the proposed subspace fusion with other subspace approaches, where the optimal dimension is difficult to determine or a single optimal dimension is not sufficient. Thirdly, it is still an open question how to optimally model the complex data distribution. The proposed framework demonstrates the effectiveness of the multi-Gaussian model. The plan is to explore other ways to construct the model, or extend this research to other pattern-recognition tasks, e.g., from UAV detection to UAV classification. Lastly, as a new dataset has been collected using SQUIRE radar (a FMCW radar) from Thales, the authors will explore the feasibility of not only detecting UAV, but also determining the direction and the distance of the UAV to the radar.
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