

Union and Intersection K-Fold Feature Selection

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Abstract: Feature selection (FS) is a vast research topic with many techniques proposed over the years. FS techniques may bring many benefits to machine learning algorithms. The combination of FS techniques usually improves the results as compared to the use of one single technique. Recently, the concepts of *explainability* and *interpretability* have been proposed in the *explainable artificial intelligence* (XAI) framework. The recently proposed *k-fold feature selection* (KFFS) algorithm provides dimensionality reduction and simultaneously yields an output suitable for explainability purposes. In this paper, we extend the KFFS algorithm by performing union and intersection of the individual feature subspaces of two and three feature selection filters. Our experiments performed on 20 datasets show that the union of the feature subsets typically attains better results than the use of individual filters. The intersection also attains adequate results, yielding human manageable (e.g., small) subsets of features, allowing for explainability and interpretability on medical domain data.

1 INTRODUCTION

The *machine learning* (ML) field is focused on learning from examples on a given dataset. The performance of ML techniques can be improved by reducing the dimensionality of the input data by keeping only the most relevant features, the key benefits are faster training and better generalization performance.


For dimensionality reduction, the use of *feature selection* (FS) techniques has been found appropriate. FS aims to identify the best performing set of features on a given task (Guyon et al., 2006; Guyon and Elisseeff, 2003; Bolon-Canedo et al., 2015). FS has a long research history and work towards improving FS techniques still continues (Alipoor et al., 2022; Chamlal et al., 2022; Huynh-Cam et al., 2022; Jeon and Hwang, 2023; Xu et al., 2022). FS techniques can be grouped into four categories: filters, wrappers, embedded, and hybrid (Guyon et al., 2006; Bolon-Canedo et al., 2015). In this paper, we use filter techniques, which assess the quality of subsets of features by using some metrics over the data, without resorting to any learning algorithm. In this sense, filter techniques


are referred to as *agnostic*. When dealing with high-dimensional data, we often find that filters are the only suitable approach, since the other techniques are too time-consuming and their use becomes computationally prohibitive (Hastie et al., 2009; Guyon et al., 2006; Escolano et al., 2009). For recent surveys on FS techniques, please see the publications by Remeseiro and Bolon-Canedo (2019), Pudjihartono et al. (2022a), and Dhal and Azad (2022).

In this work, we address the use of unsupervised and supervised FS filter techniques for different types of data. We propose to improve and extend the *k-fold feature selection* (KFFS) algorithm proposed by Ferreira and Figueiredo (2023), using combinations of heterogeneous filters. These combinations attain both adequate dimensionality reduction and improved performance. Moreover, the small dimensionality of reduced feature subspace allows for the human end user to focus on explainability and interpretability tasks.

1.1 Combination of Filters

We find the use of combination of filters in different applications. The problem of sleep disease diagnostic was addressed by Álvarez Estévez et al. (2011), with the monitoring of bio-physiological signals of

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patients during sleep, with polysomnography (PSG) data. A dataset with PSG of patients was used for the detection of arousals in sleep. From a set of 42 features extracted from biosignals methods to detect sleep events were developed. Using FS techniques the goal was to remove redundant features, identifying the best subset of features preserving classification accuracy. Wrapper and filter methods and combinations of these were considered, by union and intersection operations. Discarding the irrelevant features, a reduced dimensionality dataset was obtained, improving the accuracy of the classifiers.

The *heterogeneous ensemble feature selection* (HEFS) method proposed by Damte et al. (2023) fuses the output feature subsets of five FS filters with an union combination. It resorts to a merit-based evaluation to minimize redundancy of the obtained ensemble of features. In a multi-class intrusion detection dataset, HEFS leads to better performance than the individual FS methods.

Mochammad et al. (2022) proposed the *multi-filter clustering fusion* (MFCF) technique. A multi-filter method combining filter methods is applied as a first step for feature clustering; then, the key features are selected. The union of key features is used to find all potentially important features. An exhaustive search finds the best combination of selected features, to maximize the accuracy of the classification model. For rotating machinery problems, the fault classification models using MFCF yields good accuracy.

The intersection of common features selected by filter, wrapper, and embedded FS techniques was proposed by Bashir et al. (2022). A *support vector machines* (SVM) classifier is then trained on medical domain data, attaining better results as compared to the individual use of the FS methods.

Arya and Gupta (2023) introduced an ensemble filter-based FS approach combining ANOVA, Pearson correlation coefficient, mutual information, and Chi-square. The reduced feature sets are obtained with the union and intersection operations. Using decision tree, random forest, XGBoost, and CatBoost classifiers on the Edge-IIoT dataset (cyber-attack detection), we have 97.84% and 99.61% accuracy using the intersection and union feature sets, respectively.

An ensemble FS approach was proposed by Seijo-Pardo et al. (2017). The heterogeneous ensemble combines the result of different FS methods, with the same training data. The outputs of the base selectors are combined with different *aggregators* to obtain the resulting subset. On the experimental evaluation with the SVM classifier, ensemble results for seven datasets achieve comparable on better performance than the one attained by individual methods.

For reviews on ensemble FS methods and their combination, please see the publications by Bolón-Canedo and Alonso-Betanzos (2019) and Pudjihartono et al. (2022b).

1.2 Paper Organization

The remainder of this paper is organized as follows. In Section 2, we review some topics on feature selection. The proposed approach is detailed in Section 3. The experimental evaluation is reported in Section 4. Finally, Section 5 provides concluding remarks and directions of future work.

2 FEATURE SELECTION

We introduce notation and review some details of FS techniques in Section 2.1. An overview of the techniques considered in this work is presented in Section 2.2, including the *k-fold feature selection* (KFFS) algorithm, which we propose to extend in this work.

2.1 Notation

Regarding the notation followed in this paper, let $\mathbf{X} = \{\mathbf{x}_1, \dots, \mathbf{x}_n\}$ denote a dataset, represented as a $n \times d$ matrix (n instances on the rows and d features on the columns). Each instance \mathbf{x}_i is a d -dimensional vector, with $i \in \{1, \dots, n\}$. Each feature vector, a column of \mathbf{X} , is denoted as X_j , with $j \in \{1, \dots, d\}$. The number of classes is C , with $c_i \in \{1, \dots, C\}$ representing the class of instance \mathbf{x}_i . Finally, $\mathbf{y} = \{c_1, \dots, c_n\}$ represents the class labels for each instance, with $c_i \in \{1, \dots, C\}$.

In this work, we consider both unsupervised and supervised FS filters. The former do not use the class label vector \mathbf{y} , while the latter uses the label of each instance to perform the feature assessment. Some FS methods are based purely on the relevance of the features; they rank the features according to some criterion and then select the top-ranked ones. Other methods are based on the *relevance-redundancy* (RR) framework (Yu and Liu, 2004). In this case, the most relevant features are kept and a redundancy analysis is performed to remove redundant features.

2.2 Feature Selection Filters

We consider the three FS filters next described. The first technique is the *fast correlation-based filter* (FCBF) proposed by Yu and Liu (2003), based on the RR framework, computing the feature-class and feature-feature correlations. It starts by selecting a set

of features with correlation with the class label above some threshold (the predominant features). In the second step, redundancy analysis finds redundant features among the predominant ones. These redundant features are removed, keeping the ones that are the most relevant to the class. FCBF resorts to the *symmetrical uncertainty* (SU) (Yu and Liu, 2003) measure, defined as

$$SU(X_i, X_j) = \frac{2I(X_i; X_j)}{H(X_i) + H(X_j)}, \quad (1)$$

where $H(\cdot)$ denotes the Shannon entropy and $I(\cdot)$ denotes the *mutual information* (MI) (Cover and Thomas, 2006). The SU is zero for independent random variables and equal to one for deterministically dependent random variables, i.e., if one is a bijective function of the other.

The second FS technique is the Fisher ratio, a supervised relevance-only method. For the i -th feature, with $C = 2$, it computes the rank of the feature according to

$$FiR_i = \frac{|\bar{X}_i^{(-1)} - \bar{X}_i^{(1)}|}{\sqrt{\text{var}(X_i)^{(-1)} + \text{var}(X_i)^{(1)}}}, \quad (2)$$

where $\bar{X}_i^{(-1)}$, $\bar{X}_i^{(1)}$, $\text{var}(X_i)^{(-1)}$, and $\text{var}(X_i)^{(1)}$ are the sample means and variances of feature X_i , for the instances of both classes. It aims to measure how well each feature separates the two classes and is adequate as a relevance metric for FS purposes. For the multi-class case, $C > 2$, the FiR of feature X_i is given by (Duda et al., 2001; Zhao et al., 2010)

$$FiR_i = \frac{\sum_{j=1}^C n_j^{(y)} \left(\bar{X}_i^{(j)} - \bar{X}_i \right)^2}{\sum_{j=1}^C n_j^{(y)} \text{var} \left(X_i^{(j)} \right)}, \quad (3)$$

where $n_j^{(y)}$ is the number of occurrences of class j in the n -length class label vector y , and $\bar{X}_i^{(j)}$ is the sample mean of the values of X_i whose class label is j ; finally, \bar{X}_i is the sample mean of feature X_i .

The third FS filter is the relevance-only unsupervised *mean-median* (MM) criterion, which ranks features according to

$$MM_i = |\bar{X}_i - \text{median}(X_i)|. \quad (4)$$

The relevance of each feature is the absolute difference between the mean and median of X_i . This criterion is based on the idea that the most relevant features are the ones with more asymmetric distributions.

The k -fold *feature selection* (KFFS) filter, described in Algorithm 1, was proposed by Ferreira and

Figueiredo (2023) and it can work with any unsupervised or supervised FS filter.

KFFS follows the rationale that the importance of a feature is proportional to the number of times it is selected on the k -folds over the training data. It requires two parameters: the number of folds k to sample the training data and the threshold T_h to assess the percentage of choice of a feature by the filter on the k folds.

3 PROPOSED APPROACH

In Section 3.1, we present our key insights regarding the union and intersection of feature subspaces. The details of the proposed technique are presented in Section 3.2.

3.1 Union and Intersection

Our proposal is built upon the idea of the union and the intersection of feature subspaces, as depicted in Figure 2. Suppose that we have a feature space with d features and over that space we apply three different FS filters. These filters return feature subspaces with dimensionality m_1 , m_2 , and m_3 features, respectively. In Figure 2, we observe the union and the intersection among these feature subspaces, using an analogy with the additive RGB color scheme. The subspaces selected by FS methods 1, 2, and 3 are assigned to the primary R, G, and B colors, respectively. The intersection of the filter subspaces is represented by the corresponding results of the color addition on the RGB color space. To denote the number of features in common on the subspaces found by FS methods i and j , we use m_{ij} , with $i, j \in \{1, 2, 3\}$; on the case of three FS filters, we use the notation m_{123} .

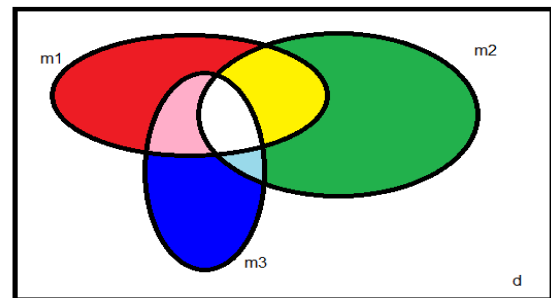


Figure 1: Feature subspace analysis for the case of three FS methods, on a d -dimensional space using a visual correspondence with the three primary colors.

Over these feature subspaces, we can compute statistics to assess the relation and (dis)similarities be-

Algorithm 1: k-Fold Feature Selection (KFFS) for filter FS by Ferreira and Figueiredo (2023).

Require: $X : n \times d$ matrix, n patterns of a d -dimensional dataset.
 @filter : a FS filter (unsupervised or supervised).
 k : an integer stating the number of folds ($k \in \{2, \dots, n\}$).
 T_h : a threshold (percentage) to chose the number of features.
 $y : n \times 1$ class label vector (necessary only in case of a FS supervised filter).
Ensure: idx : m -dimensional vector with the indexes of the selected features.

- 1: Allocate the *feature counter vector* (FCV), with dimensions $1 \times d$, such that each position refers to a specific feature.
 - 2: Initialize $FCV_i = 0$, with $i \in \{0, \dots, d-1\}$.
 - 3: Compute the k data folds in the dataset (different splits into training and test data).
 - 4: For each fold, apply @filter on the training data and update FCV_i with the number of times @filter selects feature i .
 - 5: After the k data folds are processed, convert FCV to percentage: $FCVP \leftarrow FCV/k$.
 - 6: Keep the indexes of the features that have been selected at least T_h times (expressed in percentage), $idx \leftarrow FCVP \geq T_h$.
 - 7: Return idx (the vector with the indexes of the selected features that have been selected at least T_h times).
-

tween them. The Jaccard index (JI) is one of such metrics, being defined as

$$JI(A, B) = \frac{|A \cap B|}{|A \cup B|}, \quad (5)$$

for sets A and B , where \cap denotes intersection, \cup denotes union, and $|\cdot|$ is the cardinality of the set. We have $0 \leq JI(A, B) \leq 1$. On the extreme cases, we have: if $A \cap B = \emptyset$, then $JI(A, B) = 0$; if $A \subseteq B$ or $B \subseteq A$ then $JI(A, B) = 1$. Other similar metrics are the Dice-Sorenson (DS) coefficient,

$$DS(A, B) = 2 \frac{|A \cap B|}{|A| + |B|}, \quad (6)$$

and the overlap coefficient or Szymkiewicz–Simpson (SS) coefficient,

$$SS(A, B) = \frac{|A \cap B|}{\min(|A|, |B|)}, \quad (7)$$

both ranging from 0 (maximally different) to 1 (maximally similar or one is a subset of the other).

3.2 Union and Intersection KFFS

Our proposal extends the KFFS algorithm as follows:

- Set the T_h and k parameters of KFFS to their values; by default, we set $k = 10$ and $T_h = 1$.
- Apply KFFS with two or three different FS filters, described in Section 2.2. We apply KFFS (@filter₁=FCBF), KFFS (@filter₂=FiR), and KFFS (@filter₃=MM) on the same k data folds, using the threshold T_h . Each filter will select different subsets of the input feature space, as depicted in Figure 2.
- Get the output indexes returned by each filter, idx_{fcfb} , idx_{fir} , and idx_{mm} .
- Combine the output indexes (idx_{fcfb} , idx_{fir} , and idx_{mm}) returned by the filters, with *union* and *intersection* of the indexes of the selected features.

- Return the two index vectors, given by

$$idx_{union} = idx_{fcfb} \cup idx_{fir} \cup idx_{mm};$$

$$idx_{intersection} = idx_{fcfb} \cap idx_{fir} \cap idx_{mm}.$$

The rationale is that by using and combining different filters, we are able to focus on different subsets of the original input feature space. We also expect that the combination of these feature subspaces will overcome the results of each individual FS method. The union of the feature subspaces will yield (much) larger subspaces than the intersection of these subspaces. In the intersection of the two or three subspaces, we will have a small number of features which are really relevant, since they are always selected regardless the FS filter.

The unsupervised MM filter and the supervised Fisher and FCBF FS filters were described in Section 2.2. The MM and Fisher filters are relevance-based methods, which select the top m most relevant features as follows:

- Compute the MM relevance by Equation (4) or Fisher ratio relevance by Equations (2) or (3), denoted as R_i , for each feature X_i , $i \in \{1, \dots, d\}$.
- Sort the relevance values by decreasing order.
- Compute the cumulative and normalized relevance values, leading to an increasing function whose values range to a maximum of 1.
- Keep the first top relevant m features, holding, say 90% of the accumulated relevance given by R_i .

The FCBF filter is a relevance-redundancy based method. We use its default parameter values.

4 EXPERIMENTAL EVALUATION

The proposed methods were evaluated with public domain datasets. Section 4.1 describes the datasets and

Table 1: Datasets with n instances, d features, and C classes.

Name	n	d	C	Problem/Task
Australian	690	14	2	Credit approval
Brain-Tumor-1	90	5920	5	Cancer detection
Brain-Tumor-2	50	10367	4	Cancer detection
Colon	62	2000	2	Cancer detection
Darwin	174	450	2	Alzheimer detection
Dermatology	366	34	6	Skin cancer detection
DLBCL	77	5469	2	B-cell malignancies
Drebin	15036	215	2	Malware detection
Heart	270	13	2	Heart disease
Hepatitis	155	19	2	Hepatitis survival
Ionosphere	351	34	2	Radar returns
Leukemia	72	7129	2	Leukemia detection
Leukemia-1	72	5328	3	Leukemia detection
Lymphoma	96	4026	9	Lymphoma detection
Prostate-Tumor	102	10509	2	Tumor detection
Sonar	208	60	2	Rock/Mine detection
Spambase	4601	57	2	Email spam
SRBCT	83	2308	4	Cancer detection
WDBC	569	30	2	Breast cancer
Wine	178	13	3	Wine cultivar

the evaluation metric. In Section 4.2, we report the experimental results for the individual filters, their union, and their intersection. In Section 4.3, we assess the effect of changing the threshold and the number of folds.

4.1 Datasets and Metrics

Table 1 describes the datasets used in our experiments. We have gathered 20 datasets with different types of data and problems, to assess the behavior of our proposed method in different classification task scenarios. The datasets are available from <https://csse.szu.edu.cn/staff/zhuzx/Datasets.html>, from the *Arizona State University* (ASU) repository (Zhao et al., 2010), from the UCI University of California at Irvine (UCI) repository (Dua and Graff, 2019), <https://archive.ics.uci.edu/ml/index.php>, from the *knowledge extraction evolutionary learning* (KEEL), <https://sci2s.ugr.es/keel/datasets.php> repository, and <https://jundongl.github.io/scikit-feature/datasets.html>.

The microarray datasets for cancer detection have “large d , small n ”, $d \gg n$. Other datasets are in the opposite situation, with $n \gg d$. We have also chosen both binary and multi-class datasets.

We use the FCBF and FiR implementation of the ASU repository. For the MM FS filter, we have our own implementation. We have considered the *naïve Bayes* (NB) classifier from *Waikato environment for knowledge analysis* (WEKA). NB classifier is sensitive to the presence of redundant features, suffering an increase in the test-set error rate in the presence of such features. Thus, it is useful to assess and compare the quality of the feature subspaces found by each method. Our key concern is to assess and compare the adequacy of the several feature subspaces and not to

find the best classifier. For comparison purposes, we have also used the *support vector machines* (SVM) classifier.

As evaluation metric, we consider the test-set error rate, with a 10-fold cross-validation procedure. We also analyze the size of the feature subsets.

4.2 Union and Intersection

Table 2 shows the average test set error rate (Err) and the average number of features m , over the ten folds, for the four combinations of unions among these subspaces.

In seven out of 20 datasets, the union of filters attains the best results. The best average global result is attained by KFFS(FCBF) closely followed by the union of the three filters. All FS filter lead to large reduction of the dimensionality of the data.

Table 3 reports the average test set error rate (Err) and the average number of features m , over the ten folds, for all the possible combinations of intersections among these subspaces. The results of the individual methods are the same as in Table 2.

For some cases, the intersection of the feature subspace is an empty set. In four out of the 20 datasets, the intersection of the filters attains better results than the use of individual filters. In generic terms, the intersection of filters also yields feature subspaces of reduced dimensionality.

4.3 Parameter Sensitivity

We analyze the effect of changing the threshold T_h in KFFS(FCBF), KFFS(FiR) and their union and intersection for the Prostate-Tumor dataset, in Figure 2. The FS approaches improve significantly the results of the baseline approach, with a consistent behavior. In KFFS, as we increase the threshold the dimensionality of the selected feature space decreases.

Table 4 reports the best threshold value for each dataset. We have made a grid search over all the possible threshold values from 0 to 100, and for each of the four filters KFFS(FCBF), KFFS(Fisher), and their union and intersection, we have recorded the highest threshold (fewer features) with the lowest test set error rate by the SVM classifier.

We now analyze the effect of changing the number of folds k in KFFS, for a fixed threshold T_h . The goal is to assess the sensitivity of our proposed method with the number of sampling folds on the training data. In Figure 3, we assess the test set error rate of the SVM classifier with 10-fold CV, on the DLBCL dataset, with ten different values of $k \in \{n/10, 2n/10, \dots, n\}$ and a fixed $T_h = 50$.

Table 2: Union evaluation. The average values of test set error rate (Err, in %) and the average number of features m for each individual FS filter and their union, on the ten folds of 10-fold CV, for all the benchmark datasets. We use KFFS with $k = 10$ and $T_h = 1$ with $@filter_1 = FCBF$, $@filter_2 = FiR$, and $@filter_3 = MM$. The lower Err is in boldface. Regarding the error rates, the Friedman test p-value is $p = 0.0010340240$ (≤ 0.05), thus having statistical significance.

Dataset	Baseline NB		Individual Filters						Union of Filters							
	Err	d	KFFS(FCBF)		KFFS(FiR)		KFFS(MM)		\cup_{12}		\cup_{13}		\cup_{23}		\cup_{123}	
			Err	m	Err	m	Err	m	Err	m	Err	m	Err	m	Err	m
Australian	23.48	14	24.06	8	23.48	9	24.93	9	23.33	10	23.91	11	23.62	12	23.62	12
Brain-Tumor-1	10.00	5920	12.22	473	18.89	205	43.33	1	12.22	617	12.22	474	18.89	206	12.22	618
Brain-Tumor-2	32.00	10367	28.00	359	24.00	205	34.00	121	22.00	535	24.00	474	32.00	303	20.00	630
Colon	40.48	2000	19.05	55	17.62	86	52.14	1	21.19	114	19.05	57	15.95	88	17.86	116
Darwin	12.68	450	12.06	112	13.20	57	14.38	62	11.50	139	14.41	151	12.65	104	11.50	174
Dermatology	2.80	34	3.63	19	26.53	8	2.80	29	3.63	23	2.80	31	2.80	30	2.80	31
DLBCL	18.21	5469	6.43	225	9.11	116	15.36	2	10.54	292	6.43	226	10.54	117	10.54	292
Drebin	16.64	215	8.73	17	19.89	28	21.56	48	18.24	42	20.12	62	20.69	52	19.14	66
Heart	15.56	13	16.67	8	15.19	10	17.78	10	15.56	11	15.19	11	15.19	12	15.19	12
Hepatitis	15.42	19	17.42	9	17.33	12	19.33	17	16.08	12	16.75	18	18.00	18	16.08	18
Ionosphere	18.81	34	8.83	15	18.25	16	19.38	23	16.25	23	17.10	27	19.10	23	16.81	27
Leukemia	1.43	7129	2.86	171	4.29	142	33.39	2	2.86	256	2.86	173	4.29	144	2.86	258
Leukemia-1	4.29	5327	5.71	204	4.29	149	54.46	2	4.29	301	5.71	207	4.29	152	4.29	304
Lymphoma	24.00	4026	23.11	848	18.00	128	15.56	112	19.89	904	22.00	921	12.67	218	20.89	964
Prostate-Tumor	37.09	10509	9.64	257	8.73	114	32.09	100	10.55	320	14.55	354	11.55	211	12.55	415
Sonar	32.74	60	34.67	18	34.67	20	31.64	14	35.62	25	30.81	31	33.21	32	32.74	36
Spambase	20.73	54	23.63	18	13.45	14	21.45	28	21.02	24	20.28	35	20.89	33	20.28	37
SRBCT	1.11	2308	0.00	203	1.11	145	6.11	118	1.11	267	0.00	297	1.11	239	1.11	351
WDBC	6.67	30	4.92	11	6.85	14	7.20	13	6.14	20	5.79	18	6.49	18	6.14	22
Wine	2.78	13	2.22	10	5.56	5	3.33	12	2.22	10	2.78	13	3.33	12	2.78	13

Table 3: Intersection evaluation. The average values of test set error rate (Err, in %) and the average number of features m for each individual FS filter and their intersection, on the ten folds of 10-fold CV, for all the benchmark datasets. We use KFFS with $k = 10$ and $T_h = 1$ with $@filter_1 = FCBF$, $@filter_2 = FiR$, and $@filter_3 = MM$. The lower Err is in boldface. Regarding the error rates, the Friedman test p-value is $p = 0.0010340240$ (≤ 0.05), thus having statistical significance.

Dataset	Baseline NB		Individual Filters						Intersection of Filters							
	Err	d	KFFS(FCBF)		KFFS(FiR)		KFFS(MM)		\cap_{12}		\cap_{13}		\cap_{23}		\cap_{123}	
			Err	m	Err	m	Err	m	Err	m	Err	m	Err	m	Err	m
Australian	23.48	14	24.06	8	23.48	9	24.93	9	24.06	7	25.65	6	24.78	6	25.80	5
Brain-Tumor-1	10.00	5920	12.22	473	18.89	205	43.33	1	20.00	62	-	0	-	0	-	0
Brain-Tumor-2	32.00	10367	28.00	359	24.00	205	34.00	121	30.00	30	34.00	6	38.00	23	46.00	3
Colon	40.48	2000	19.05	55	17.62	86	52.14	1	19.05	27	-	0	-	0	-	0
Darwin	12.68	450	12.06	112	13.20	57	14.38	62	12.58	30	13.73	23	13.17	15	16.50	11
Dermatology	2.80	34	3.63	19	26.53	8	2.80	29	27.09	5	5.03	17	30.16	7	30.71	4
DLBCL	18.21	5469	6.43	225	9.11	116	15.36	2	5.00	50	19.64	1	-	0	-	0
Drebin	16.64	215	8.73	17	19.89	28	21.56	48	11.34	3	11.79	2	21.14	24	11.79	2
Heart	15.56	13	16.67	8	15.19	10	17.78	10	16.67	8	17.41	7	18.52	9	18.15	7
Hepatitis	15.42	19	17.42	9	17.33	12	19.33	17	19.92	8	17.38	8	17.96	11	19.88	7
Ionosphere	18.81	34	8.83	15	18.25	16	19.38	23	12.83	8	11.12	10	19.10	15	13.69	7
Leukemia	1.43	7129	2.86	171	4.29	142	33.39	2	2.86	57	-	0	-	0	-	0
Leukemia-1	4.29	5327	5.71	204	4.29	149	54.46	2	4.11	52	-	0	-	0	-	0
Lymphoma	24.00	4026	23.11	848	18.00	128	15.56	112	19.11	72	13.67	39	16.67	22	20.89	9
Prostate-Tumor	37.09	10509	9.64	257	8.73	114	32.09	100	7.73	51	16.73	3	19.64	2	13.82	1
Sonar	32.74	60	34.67	18	34.67	20	31.64	14	35.60	14	40.90	2	40.86	2	41.86	1
Spambase	20.73	54	23.63	18	13.45	14	21.45	28	15.89	8	25.06	11	14.63	10	16.91	6
SRBCT	1.11	2308	0.00	203	1.11	145	6.11	118	1.25	81	4.86	24	3.61	24	7.22	14
WDBC	6.67	30	4.92	11	6.85	14	7.20	13	5.79	5	6.85	5	8.78	9	8.43	4
Wine	2.78	13	2.22	10	5.56	5	3.33	12	5.56	5	2.78	10	5.56	5	5.56	5

The number of folds k has a large impact on the end result for all filters. For lower values of k , we have a non-stationary behavior of the error rate curve. After a sufficiently large value of k , we observe a more stable behavior on the error rate. These results show that, for a specific dataset and problem, one should fine-tune both the T_h and k parameters to have better results.

5 CONCLUSIONS

In this paper, we have extended the KFFS filter algorithm by performing union and intersection of the individual feature subspaces of two and three heterogeneous FS filters. We have considered two supervised FS filters (FCBF and FiR) and one unsupervised filter (MM). Two of these filters are relevance based (FiR

Table 4: The best test set error rate (Err, in %), the corresponding average number of features m and Threshold, T_h , for KFFS(FCBF), KFFS(Fisher), and their union and intersection, for all the benchmark datasets. We use KFFS with $k = 10$ and the SVM classifier. The best result is in boldface.

Dataset	Baseline SVM		Individual Filters						Union \cup_{12}			Intersection \cap_{12}		
	Err	d	KFFS(FCBF)			KFFS(FiR)			Err	m	T_h	Err	m	T_h
			Err	m	T_h	Err	m	T_h						
Australian	14.49	14	14.49	4	91	14.49	7	81	14.49	8	81	14.49	3	91
Brain-Tumor-1	10.00	5920	10.00	75	31	10.00	5920	0	10.00	158	31	10.00	5920	0
Brain-Tumor-2	20.00	10367	20.00	10367	0	18.00	92	21	16.00	219	11	18.00	30	1
Colon	13.10	2000	11.43	17	21	13.10	17	91	11.43	117	1	13.10	2000	0
Darwin	17.12	450	16.60	35	31	14.38	37	11	16.57	70	21	17.12	450	0
Dermatology	3.36	34	3.08	20	1	3.36	34	0	2.79	17	51	3.36	34	0
DLBCL	2.50	5469	2.50	42	41	2.50	5469	0	2.50	79	41	2.50	25	11
Drebin	2.23	215	2.23	215	0	2.23	215	0	2.23	215	0	2.23	215	0
Heart	15.93	13	14.07	8	1	14.07	10	1	14.07	10	1	14.07	8	1
Hepatitis	23.29	19	19.38	5	61	18.17	10	21	17.46	10	61	18.79	7	11
Ionosphere	11.42	34	11.42	34	0	11.42	34	0	11.13	22	1	11.42	34	0
Leukemia	1.43	7129	1.43	7129	0	1.43	7129	0	1.43	7129	0	1.43	7129	0
Leukemia-1	1.43	5327	1.43	5327	0	1.43	5327	0	1.43	5327	0	1.43	5327	0
Lymphoma	4.33	4026	4.33	80	61	4.33	4026	0	4.33	93	71	4.33	4026	0
Prostate-Tumor	8.00	10509	6.00	24	61	6.00	54	51	5.00	65	61	4.00	48	1
Sonar	21.71	60	21.19	9	41	21.24	18	11	21.24	21	11	21.71	60	0
Spambase	10.06	54	10.06	54	0	10.06	54	0	10.06	54	0	10.06	54	0
SRBCT	0.00	2308	0.00	54	41	0.00	56	91	0.00	61	91	0.00	31	31
WDBC	2.28	30	2.28	30	0	2.28	30	0	1.93	17	21	2.28	30	0
Wine	0.56	13	0.56	8	91	0.56	13	0	0.56	9	81	0.56	13	0

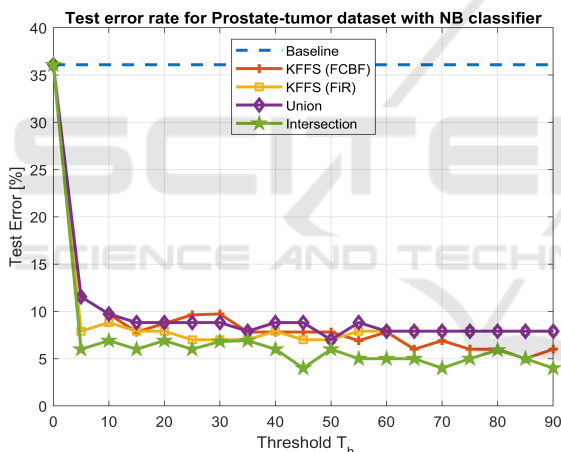


Figure 2: Test set error rate of the NB classifier with 10-fold CV, as a function of the threshold in KFFS, for KFFS(FCBF), KFFS(Fisher), and their Union and Intersection, with $k = 10$ on the Prostate-Tumor dataset.

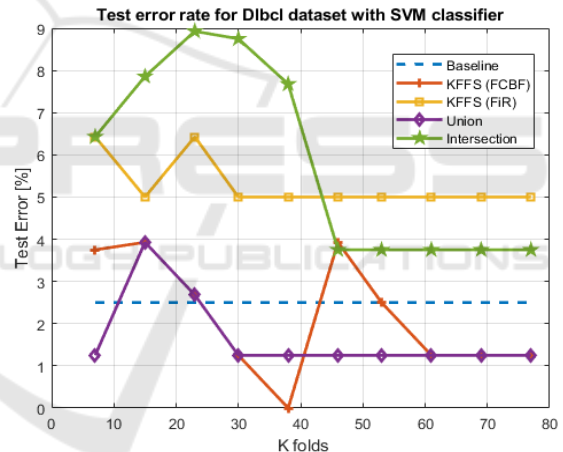


Figure 3: Test set error rate of the SVM classifier with 10-fold CV, as a function of the number of folds in KFFS, for KFFS(FCBF), KFFS(Fisher), and their Union and Intersection, with $T_h = 50$ on the DLBCL dataset.

and MM) while FCBF follows the RR framework. Our experiments on 20 datasets with diverse types of data and problems show that the union of the feature subsets typically attains better results than the individual filters. The intersection also attains adequate results, yielding human manageable subsets of features allowing for explainability and interpretability. By setting properly the threshold of the KFFS algorithm, we can control the dimensionality of the feature subspaces, reduced in such a way that allows for the domain expert (e.g., a medical doctor) to focus on the interpretation of the resulting variables.

However, in some cases, the subspace intersection is empty. The dimensionality of the subspace resulting from the intersection is typically much lower, as compared to the one from the union. When dealing with high-dimensional data, it is often the case that FS filters select different regions of the feature subspace. As future work, we aim to fine-tune the parameters of the method for each dataset or type of data/problem, individually. We will also explore the use of different thresholds per filter.

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