Unsupervised Feature Selection Using Feature Similarity

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Abstract—In this article, we describe an unsupervised feature selection algorithm suitable for data sets, large in both dimension and size. The method is based on measuring similarity between features whereby redundancy therein is removed. This does not need any search and, therefore, is fast. A feature similarity measure, called maximum information compression index, is introduced. The algorithm is generic in nature and has the capability of multiscale representation of data sets. The superiority of the algorithm, in terms of speed and performance, is established extensively over various real-life data sets of different sizes and dimensions. It is also demonstrated how redundancy and information loss in feature selection can be quantified with an entropy measure.

Index Terms—Data mining, pattern recognition, dimensionality reduction, feature clustering, multiscale representation, entropy measures.

1 Introduction

A important problem related to mining large data sets, both in dimension and size, is of selecting a subset of the original features [1]. Preprocessing the data to obtain a smaller set of representative features, retaining the optimal salient characteristics of the data, not only decreases the processing time but also leads to more compactness of the models learned and better generalization. When class labels of the data are available we use supervised feature selection, otherwise unsupervised feature selection is appropiate. In many data mining applications, class labels are unknown, thereby indicating the significance of unsupervised feature selection there.

Conventional methods of feature selection involve evaluating different feature subsets using some index and selecting the best among them. The index usually measures the capability of the respective subsets in classification or clustering depending on whether the selection process is supervised or unsupervised. A problem of these methods, when applied to large data sets, is the high-computational complexity involved in searching. The complexity is exponential in terms of the data dimension for an exhaustive search. Several heuristic techniques have been developed to circumvent this problem. Among them the branch and bound algorithm, suggested by Devijver and Kittler [2], obtains the optimal subset in expectedly less than exponential computations when the feature evaluation criterion used is monotonic in nature. Greedy algorithms like sequential forward and backward search [2] are also popular. These algorithms have quadratic complexity, but they perform poorly for nonmonotonic indices. In such cases, sequential floating

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searches [3] provide better results, though at the cost of a higher computational complexity. Beam search variants of the sequential algorithms [4] are also used to reduce computational complexity. Recently, robust methods for finding out the optimal subset for arbitrary evaluation indices are being developed using genetic algorithms (GAs) [5]. GA based feature selection methods [6] are usually found to perform better than other heuristic search methods for large and medium sized data sets; however, they also require considerable computation time for large data sets. Other attempts to decrease the computational time of feature selection include probabilistic search methods like random hill climbing [7], SCHEMATA+ [8], and Las Vegas Filter (LVF) approach [9]. Comparison and discussion of some of the above methods for many real-life data sets may be found in [6].

Since the interest of the article lies with unsupervised feature selection, we discuss here some of the existing methods which can be broadly classified into two categories. Methods in one such category involve maximization of clustering performance, as quantified by some index. These include sequential unsupervised feature selection algorithm [10], wrapper approach based on expectation maximization (EM) [11], maximum entropy based method [12], and the recently developed neuro-fuzzy approach [13]. The other category considers selection of features based on feature dependency and relevance. The principle is that any feature carrying little or no additional information beyond that subsumed by the remaining features, is redundant and should be eliminated. Various dependence measures like correlation coefficients [14], measures of statistical redundancy [15], or linear dependence [16], [17] have been used. Recently, the Relief algorithm [18] and its extensions [19] which identify statistically relevant features have been reported. Another algorithm based on conditional independence uses the concept of Markov blanket [20]. All these methods involve search and require significantly high computation time for large data sets. In [21], an algorithm which does not involve search and selects features by

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hierarchically merging similar feature pairs is described. However, the algorithm is crude in nature and performs poorly on real-life data sets. It may be noted that principal component analysis (PCA) [2] also performs unsupervised dimensionality reduction based on information content of features. However, PCA involves feature transformation and obtains a set of transformed features rather than a subset of the original features.

In the present article, we propose an unsupervised algorithm which uses feature dependency/similarity for redundancy reduction, but requiring no search. The method involves partitioning of the original feature set into some distinct subsets or clusters so that the features within a cluster are highly similar while those in different clusters are dissimilar. A single feature from each such cluster is then selected to constitute the resulting reduced subset. A new similarity measure, called maximal information compression index, is used in clustering. Its comparison with two other measures, namely correlation coefficent and leastsquare regression error is made. It is also demonstrated how "representation entropy" can be used for quantifying redundancy in a set.

The nature of both the proposed clustering algorithm and the newly introduced feature similarity measure is geared toward two goals—minimizing the information loss (in terms of second order statistics) incurred in the process of feature reduction, and minimizing the redundancy present in the reduced feature subset. The feature selection algorithm owes its low-computational complexity to two factors—1) unlike most conventional algorithms, search for the best subset (requiring multiple evaluation of indices) is not involved and 2) the new feature similarity measure can be computed in much less time compared to many indices used in other supervised and unsupervised feature selection methods. Since the method achieves dimensionality reduction through removal of redundant features, it is more related to feature selection for compression rather than for classification.

Superiority of the algorithm, over four related methods, namely branch and bound algorithm, sequential floating forward search, sequential forward search, and stepwise clustering, is demonstrated extensively on nine real-life data of both large and small sample sizes and dimensions ranging from 4 to 649. Comparison is made on the basis of both clustering/classification performance and redundancy reduction. Effectiveness of the maximal information compression index and the effect of scale parameter are also studied.

The organization of the article is as follows: In the next section, we describe measures of similarity between a pair of features. In Section 3, we present the proposed feature selection algorithm using the similarity measure and discuss some of its characterstics. In Section 5 we provide experimental results along with comparisons.

FEATURE SIMILARITY MEASURE

In this section, we discuss some criteria for measuring similarity between two random variables, based on linear dependency between them. In this context, we propose a new measure, called maximal information compression index, to be used for feature selection.

There are broadly two possible approaches for measuring similarity between two random variables. One is to nonparametrically test the closeness of probability distributions of the variables. Walds-Wolfowitch test and other run test [22] may be used for this purpose. However, these tests are sensitive to both location and dispersion of the distributions, hence not suited for the purpose of feature selection. Another approach is to measure the amount of functional (linear or higher) dependency between the variables. There are several benefits of choosing linear dependency as a feature similarity measure. It is known that if some of the features are linearly dependent on the others, and if the data is linearly separable in the original representation, the data is still linearly separable if all but one of the linearly dependent features are removed [16]. As far as the information content of the variables is concerned, second order statistics of the data is often the most important criterion after mean values [22]. All the linear dependency measures that we will discuss are related to the amount of error in terms of second order statistics, in predicting one of the variables using the other. We discuss below two existing [22] linear dependency measures before explaining the proposed maximal information compression index.

Correlation Coefficient(ρ). The most well-known measure of similarity between two random variables is the correlation coefficient. Correlation coefficient ρ between two random variables x and y is defined as

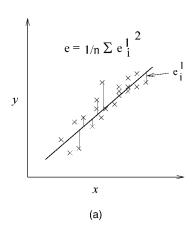
$$\rho(x, y) = \frac{\text{cov}(x, y)}{\sqrt{\text{var}(x)\text{var}(y)}},$$

where var() denotes the variance of a variable and cov() the covariance between two variables. If x and y are completely correlated, i.e., exact linear dependency exist, $\rho(x,y)$ is 1 or -1. If x and y are totally uncorrelated, $\rho(x,y)$ is 0. Hence, $1 - |\rho(x, y)|$ can be used as a measure of similarity between two variables x and y. The following can be stated about the measure:

- 1. $0 \le 1 |\rho(x, y)| \le 1$.
- 2. $1 |\rho(x, y)| = 0$ if and only if x and y are linearly
- 3. $1-|\rho(x,y)|=1-|\rho(y,x)|$ (symmetric). 4. If $u=\frac{x-a}{c}$ and $v=\frac{y-b}{d}$ for some constants a,b,c,d, then $1-|\rho(x,y)|=1-|\rho(u,v)|$ i.e., the measure is invariant to scaling and translation of the variables.
- The measure is sensitive to rotation of the scatter diagram in (x, y) plane.

Though correlation coefficient contains many desirable properties as a feature similarity measure, properties 4 and 5, mentioned above, make it somewhat unsuitable for feature selection. Since the measure is invariant to scaling, two pairs of variables having different variances may have the same value of the similarity measure, which is not desirable as variance has high information content. Sensitivity to rotation is also not desirable in many applications.

Least Square Regression Error (e). Another measure of the degree of linear dependency between two variables x and y



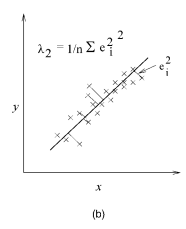


Fig. 1. Nature of errors in linear regression: (a) Least-square fit (e) and (b) least-square projection fit (λ_2).

is the error in predicting y from the linear model y = a + bx. a and b are the regression coefficients obtained by minimizing the mean square error

$$e(x,y)^{2} = \frac{1}{n} \sum_{i} (e(x,y)_{i})^{2},$$

 $e(x,y)_i=y_i-a-bx_i$. The coefficients are given by $a=\bar{y}$ and $b=\frac{\cos v}{(x,y)\mathrm{var}(x)}$ and the mean square error e(x,y) is given by $e(x,y)=\mathrm{var}(y)(1-\rho(x,y)^2)$. If y and x are linearly related e(x,y)=0, and if x and y are completely uncorrelated $e(x,y)=\mathrm{var}(y)$. The measure e^2 is also known as the residual variance. It is the amount of variance of y unexplained by the linear model. Some properties of the e are:

- 1. $0 \le e(x, y) \le \text{var}(y)$.
- 2. e(x, y) = 0 if and only if x and y are linearly related.
- 3. $e(x,y) \neq e(y,x)$ (unsymmetric).
- 4. If u = x/c and v = y/d for some constant a, b, c, d, then $e(x, y) = d^2e(u, v)$, i.e., the measure e is *sensitive* to scaling of the variables. It is also clear that e is invariant to translation of the variables.
- 5. The measure e is *sensitive to rotation* of the scatter diagram in x y plane.

Note that the measure e is not symmetric (property 3). Moreover, it is sensitive to rotation (property 5).

Now, we suggest a measure of linear dependency which has many desirable properties for feature selection not present in the above two measures.

Maximal Information Compression Index (λ_2) . Let Σ be the covariance matrix of random variables x and y. Define, maximal information compression index as $\lambda_2(x,y)=$ smallest eigenvalue of Σ , i.e.,

$$2\lambda_2(x,y) = (\operatorname{var}(x) + \operatorname{var}(y) - \sqrt{(\operatorname{var}(x) + \operatorname{var}(y))^2 - 4\operatorname{var}(x)\operatorname{var}(y)(1 - \rho(x,y)^2)}.$$

The value of λ_2 is zero when the features are linearly dependent and increases as the amount of dependency decreases. It may be noted that the measure λ_2 is nothing but the eigenvalue for the direction normal to the principle component direction of feature pair (x, y). It is shown in [2]

that maximum information compression is achieved if a multivariate (in this case bivariate) data is projected along its principal component direction. The corresponding loss of information in reconstruction of the pattern (in terms of second order statistics) is equal to the eigenvalue along the direction normal to the principal component. Hence, λ_2 is the amount of reconstruction error committed if the data is projected to a reduced (in this case reduced from two to one) dimension in the best possible way. Therefore, it is a measure of the *minimum amount of information loss* or the *maximum amount of information compression*, possible.

The significance of λ_2 can also be explained geometrically in terms of linear regression. It can be easily shown [22] that the value of λ_2 is equal to the sum of the squares of the perpendicular distances of the points (x,y) to the best fit line $y=\hat{a}+\hat{b}x$, obtained by minimizing the sum of the squared perpendicular distances. The coefficients of such a best fit line are given by $\hat{a}=\bar{x}cot\theta+\bar{y}$ and $\hat{b}=-cot\theta$, where

$$\theta = 2 \tan^{-1} \left(\frac{2 \operatorname{cov}(x, y)}{\operatorname{var}(x)^{2} - \operatorname{var}(y)^{2}} \right).$$

The nature of errors and the best fit lines for least-square regression and principal component analysis are illustrated in Fig. 1. λ_2 has the following properties:

- 1. $0 \le \lambda_2(x, y) \le 0.5(\text{var}(x) + \text{var}(y)).$
- 2. $\lambda_2(x,y) = 0$ if and only if x and y are linearly related.
- 3. $\lambda_2(x,y) = \lambda_2(y,x)$ (symmetric).
- 4. If $u = \frac{x}{c}$ and $v = \frac{y}{d}$ for some constant a, b, c, d, then $\lambda_2(x,y) \neq \lambda_2(u,v)$, i.e., the measure is *sensitive to scaling* of the variables. Since the expression of λ_2 does not contain mean, but only the variance and covariance terms, it is *invariant to translation* of the data set
- 5. λ_2 is *invariant to rotation* of the variables about the origin (this can be easily verified from the geometric interpretation of λ_2 considering the property that the perpendicular distance of a point to a line does not change with rotation of the axes).

The measure λ_2 possesses several desirable properties like symmetry (property 3), sensitivity to scaling (property 4), and invariance to rotation (property 5). It is a property of

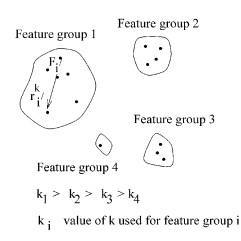


Fig. 2. Feature clusters.

the variable pair (x,y) reflecting the amount of error committed if maximal information compression is performed by reducing the variable pair to a single variable. Hence, it may be suitably used in redundancy reduction.

3 FEATURE SELECTION METHOD

The task of feature selection involves two steps, namely, partitioning the original feature set into a number of homogeneous subsets (clusters) and selecting a representative feature from each such cluster. Partitioning of the features is done based on the k-NN principle using one of the feature similarity measures described in Section 2. In doing so, we first compute the k nearest features of each feature. Among them the feature having the most compact subset (as determined by its distance to the farthest neighbor) is selected, and its k neighboring features are discarded. The process is repeated for the remaining features until all of them are either selected or discarded.

While determining the k nearest-neighbors of features, we assign a constant error threshold (ϵ) which is set equal to the distance of the kth nearest-neighbor of the feature selected in the first iteration. In subsequent iterations, we check the λ_2 value, corresponding to the subset of a feature, whether it is greater than ϵ or not. If yes, then we decrease the value of k. Therefore, k may be varying over iterations. The concept of clustering features into homogeneous groups of varying sizes is illustrated in Fig. 2. The algorithm may be stated as follows:

Algorithm:

Let the original number of features be D, and the original feature set be $O=\{F_i,i=1,\ldots,D\}$. Represent the dissimilarity between features F_i and F_j by $S(F_i,F_j)$. Higher the value of S is, the more dissimilar are the features. The measures of linear dependency (e.g., ρ,e,λ_2) described in Section 2 may be used in computing S. Let r_i^k represent the dissimilarity between feature F_i and its kth nearest-neighbor feature in R. Then

Step 1: Choose an initial value of $k \le D - 1$. Initialize the reduced feature subset R to the original feature set O,

i.e., $R \leftarrow O$.

Step 2: For each feature $F_i \in R$, compute r_i^k .

Step 3: Find feature F_{ii} for which r_{ii}^k is minimum.

Retain this feature in R and discard k nearest features of $F_{i'}$. (Note: $F_{i'}$ denotes the feature for which removing k nearest-neighbors will cause minimum error among all the features in R). Let $\epsilon = r_i^k$.

Step 4: If k > cardinality(R) - 1: k = cardinality(R) - 1.

Step 5: If k = 1: Go to Step 8.

Step 6: While $r_{i'}^k > \epsilon$ do:

(a) k = k - 1.

 $r_{i'}^k = \inf_{F_i \in R} r_i^k.$

("k" is decremented by 1, until the "kth nearest-neighbor" of at least one of the features in R is less than ϵ -dissimilar with the feature)

(b) If k = 1: Go to Step 8.

(if no feature in R has less than ϵ -dissimilar "nearest-neighbor" select all the remaining features in R)

End While

Step 7: Go to Step 2.

Step 8: Return feature set R as the reduced feature set.

3.1 Remarks

Computational Complexity. The algorithm has low-computational complexity with respect to both number of features and number of samples of the original data. With respect to the dimension (D), the method has complexity $\mathcal{O}(D^2)$. Among the existing search based schemes only sequential forward and backward search have complexity $\mathcal{O}(D^2)$, though each evaluation is more time consuming. Other algorithms like plus-l-take-r, sequential floating search and branch and bound algorithm [2] have complexity higher than quadratic. Most probabilistic search algorithms also require more than quadratic number of evaluations.

The second factor which contributes to the speedup achieved by the proposed algorithm is the low-computational complexity of evaluating the linear dependecy measures of feature similarity. If the data set contains lsamples, evaluation of the similarity measure for a feature pair is of complexity O(l). Thus, the feature selection scheme has overall complexity $\mathcal{O}(D^2l)$. Almost all other supervised and unsupervised feature evaluation indices (e.g., entropy, class seperability, K-NN classification accuracy) have at least $O(l^2)$ complexity of computation. Moreover, evaluation of the linear dependency measures involves computation using one-dimensional variables only, while the other measures often involve distance computations at higher dimensions. All these factors contribute to the large speedup achieved by the proposed algorithm compared to other feature selection schemes.

Notion of Scale in Feature Selection and Choice of k. In our algorithm k controls the size of the reduced set. Since k determines the error threshold (ϵ) , the representation of the data at different degrees of details is controlled by its choice. This characteristic is useful in data mining where multiscale representation of the data is often necessary. Note that the said property may not always be possessed by

Dataset	Method		Evaluation Criteria				
		Е	FFEI	S	KNNA (%)	BayesA (%)	Time (sec)
					Mean SD	Mean SD	
Isolet	SFS	0.52	0.41	1.09	95.02 0.89	92.03 0.52	14.01×10^4
d=310	SWC	0.71	0.55	2.70	72.01 0.71	68.01 0.44	431
D=617	Relief-F	0.70	0.52	2.24	95.81 0.81	95.52 0.47	5.03×10^3
k = 305	Proposed	0.50	0.40	1.07	96.00 0.78	95.01 0.52	440
Mult. Feat.	SFS	0.67	0.47	0.45	77.01 0.24	75.02 0.14	5.00×10^4
d=325	SWC	0.79	0.55	0.59	52.00 0.19	50.05 0.10	401
D=649	Relief-F	0.71	0.50	0.52	78.37 0.22	75.25 0.11	1.10×10^{3}
k = 322	Proposed	0.68	0.48	0.45	78.34 0.22	75.28 0.10	451
Arrhythmia	SFS	0.74	0.44	0.25	52.02 0.55	50.21 0.43	1511
d=100	SWC	0.82	0.59	0.41	40.01 0.52	38.45 0.38	70
D=195	Relief-F	0.78	0.55	0.27	56.04 0.54	54.55 0.40	404
k = 95	Proposed	0.72	0.40	0.17	58.93 0.54	56.00 0.41	74

TABLE 1
Comparison of Feature Selection Algorithms for Large-Dimensional Data Sets

E: Entropy, FFEI: Fuzzy Feature Evaluation Index, S: Class Separability, KNNA: k-NN classification accuracy, BayesA: naive Bayes classification accuracy, and SD: standard deviation. SFS: Sequential Forward Search and SWC: Stepwise Clustering. d: number of selected features, D: number of original features and k: parameter used by the proposed method.

other algorithms where the input is usually the desired size of the reduced feature set. The reason is that changing the size of the reduced set may not necessarily result in any change in the levels of details. In contrast, for the proposed algorithm, k acts as a scale parameter which controls the degree of details in a more direct manner.

Nonmetric Nature of Similarity Measure. The similarity measures used in the proposed algorithm need not be a metric. Unlike conventional agglomerative clustering algorithms, it does not utilize metric property of the similarity measures. Also, unlike the stepwise clustering method [21] used previously for feature selection, our clustering algorithm is partitional and nonhierarchical in nature.

4 FEATURE EVALUATION INDICES

Now, let us describe some indices that have been considered for evaluating the effectiveness of the selected feature subsets. The first three indices, namely class seperability, K-NN classification accuracy, and naive Bayes classification accuracy, do need class information of the samples while the remaining three namely, entropy, fuzzy feature evaluation index, and representation entropy, do not. Before we discuss them, we mention, for convenience, the following notations: Let l be the number of sample points in the data set, c be the number of classes present in the data set, d be the number of features in the original feature set d, d be the number of features in the reduced feature set d, d be the original feature space with

dimension D, and Ω_R be the transformed feature space with dimension d.

1. Class Separability [2]. Class seperability S of a data set is defined as $S = trace(S_b^{-1}S_w)$. S_w is the within class scatter matrix and S_b is the between class scatter matrix, defined as:

$$S_{w} = \sum_{j=1}^{c} \pi_{j} E\{(X - \mu_{j})(X - \mu_{j})^{T} | \omega_{j}\} = \sum_{j=1}^{c} \pi_{j} \Sigma_{j}$$

$$S_{b} = \sum_{j=1}^{c} (\mu_{j} - M_{o})(\mu_{j} - M_{o})^{T}$$

$$M_{o} = E\{X\} = \sum_{j=1}^{c} \pi_{j} \mu_{j},$$
(1)

where π_j is the a priori probability that a pattern belongs to class ω_j , X is the feature vector, μ_j is the sample mean vector of class ω_j , M_o is the sample mean vector for the entire data points, Σ_j is the sample covariance matrix of class ω_j , and $E\{.\}$ is the expectation operator. A lower value of the separability criteria S ensures that the classes are well separated by their scatter means.

 K-NN Classification Accuracy. Here, we have used the K-NN rule for evaluating the effectiveness of the reduced set for classification. Cross-validation is performed in the following manner—we randomly select 10 percent of the data as training set and

	T						T	
Dataset	Method		Evaluation Criteria					
		E	FFEI	S	KNNA (%)	BayesA (%)	Time (sec)	
					Mean SD	Mean SD		
	BB	0.50	0.30	0.28	90.01 0.71	88.17 0.55	1579	
Spambase	SFFS	0.50	0.30	0.28	90.01 0.72	88.17 0.55	1109	
	SFS	0.52	0.34	0.29	87.03 0.68	86.20 0.54	121.36	
d=29	SWC	0.59	0.37	0.41	82.04 0.68	79.10 0.55	11.02	
D=57	Relief-F	0.59	0.36	0.34	87.04 0.70	86.01 0.52	70.80	
k=27	Proposed	0.50	0.30	0.28	90.01 0.71	88.19 0.52	13.36	
	BB	0.67	0.47	0.29	78.02 0.47	62.27 0.41	1019	
Waveform	SFFS	0.68	0.48	0.31	77.55 0.45	62.22 0.41	627	
	SFS	0.69	0.49	0.37	74.37 0.44	59.01 0.42	71.53	
d=20	SWC	0.72	0.55	0.41	62.03 0.40	47.50 0.40	8.01	
D=40	Relief-F	0.73	0.54	0.38	74.88 0.41	62.88 0.40	50.22	
k = 17	Proposed	0.68	0.48	0.30	75.20 0.43	63.01 0.40	8.28	
	BB	0.65	0.44	0.07	75.96 0.35	65.10 0.28	150.11	
Ionosphere	SFFS	0.65	0.44	0.08	74.73 0.37	65.08 0.31	50.36	
	SFS	0.65	0.44	0.10	69.94 0.32	62.00 0.27	10.70	
d=16	SWC	0.66	0.47	0.22	62.03 0.32	59.02 0.25	1.04	
D=32	Relief-F	0.62	0.47	0.15	72.90 0.34	64.55 0.27	8.20	
k = 11	Proposed	0.64	0.43	0.10	78.77 0.35	65.92 0.28	1.07	

TABLE 2
Comparison of Feature Selection Algorithms for Medium-Dimensional Data Sets

BB: Branch and Bound, SFFS: Sequential Floating Forward Search

classify the remaining 90 percent points. Ten such independent runs are performed and the average classification accuracy on test set is used. The value of K, chosen for the K-NN rule, is the square root of the number of data points in the training set.

- 3. Naive Bayes Classification Accuracy. A Bayes maximum likelihood classifier [2], assuming normal distribution of classes, is also used for evaluating the classification performance. Mean and covariance of the classes are estimated from a randomly selected 10 percent training sample, and the remaining 90 percent of the points are used as test set. Ten such independent runs are performed and the average classification accuracy on test set is provided.
- 4. *Entropy* [10]. Let the distance between two data points p, q be

$$\mathcal{D}_{pq} = \left[\sum_{j=1}^{M} \left(rac{x_{p,j} - x_{q,j}}{max_j - min_j}
ight)^2
ight]^{1/2},$$

where $x_{p,j}$ denotes feature value for p along jth direction, and max_j, min_j are the maximum and minimum values computed over all the samples along

jth axis, M is the number of features. Similarity between p, q is given by $sim(p,q) = e^{-\alpha \mathcal{D}_{pq}}$, where α is a positive constant. A possible value of α is $\frac{-\ln 0.5}{\mathcal{D}}$. $\bar{\mathcal{D}}$ is the average distance between data points computed over the entire data set. Entropy is defined as:

$$E = -\sum_{p=1}^{l} \sum_{q=1}^{l} (\sin(p, q) \times \log \sin(p, q) + (1 - \sin(p, q)) \times \log (1 - \sin(p, q))).$$
(2)

If the data is uniformly distributed in the feature space, entropy is maximum. When the data has wellformed clusters uncertainty is low and so is entropy.

5. Fuzzy Feature Evaluation Index [13]. Fuzzy feature evaluation index (FFEI) is defined as:

$$\begin{split} FFEI &= \frac{2}{l(l-1)} \sum_{p} \sum_{q \neq p} \frac{1}{2} \\ & \left[\mu_{pq}^{R} (1 - \mu_{pq}^{O}) + \mu_{pq}^{O} (1 - \mu_{pq}^{R}) \right], \end{split} \tag{3}$$

where μ_{pq}^{O} and μ_{pq}^{R} are the degrees that both patterns p and q belong to the same cluster in the feature

Dataset	Method		Evaluation Criteria						
		Е	FFEI	S	KNNA (%)	BayesA (%)	Time (sec)		
					Mean SD	Mean SD			
	BB	0.65	0.40	0.90	64.03 0.41	63.55 0.40	4.01×10^4		
Forest	SFFS	0.64	0.39	0.81	67.75 0.43	66.22 0.41	3.02×10^4		
	SFS	0.64	0.41	0.98	62.03 0.41	61.09 0.40	7.00×10^{3}		
d=5	SWC	0.68	0.45	1.00	54.70 0.37	53.25 0.35	50.03		
D=10	Relief-F	0.65	0.40	0.90	64.03 0.41	63.55 0.40	2.80×10^4		
k=5	Proposed	0.65	0.40	0.90	64.03 0.41	63.55 0.40	55.50		
	BB	0.59	0.36	1.84	94.90 0.17	94.45 0.14	3.39		
Cancer	SFFS	0.59	0.36	1.84	94.90 0.17	94.45 0.14	6.82		
	SFS	0.61	0.37	2.68	92.20 0.17	91.05 0.15	1.16		
d=4	SWC	0.60	0.37	2.69	90.01 0.19	89.11 0.17	0.10		
D=9	Relief-F	0.59	0.36	1.84	94.90 0.17	94.25 0.17	0.91		
k=5	Proposed	0.56	0.34	1.70	95.56 0.17	94.88 0.17	0.10		
	BB	0.55	0.34	22.0	96.80 0.14	97.33 0.10	0.56		
Iris	SFFS	0.55	0.34	22.0	96.80 0.14	97.33 0.10	0.71		
	SFS	0.57	0.35	27.0	92.55 0.17	93.10 0.14	0.25		
d=2	SWC	0.60	0.37	29.2	92.19 0.19	93.02 0.17	0.01		
D=4	Relief-F	0.55	0.34	22.0	96.80 0.14	97.33 0.10	0.14		
k=2	Proposed	0.55	0.34	22.0	96.80 0.14	97.33 0.10	0.01		

TABLE 3
Comparison of Feature Selection Algorithms for Low-Dimensional Data Sets

spaces Ω_O and Ω_R , respectively. Membership function μ_{pq} may be defined as

$$\mu_{pq} = 1 - \frac{d_{pq}}{\mathcal{D}_m ax}$$
 if $d_{pq} \leq \mathcal{D}_{max}$
= 0, otherwise.

 d_{pq} is the distance between patterns p and q, and \mathcal{D}_{max} is the maximum separation between patterns in the respective feature spaces.

The value of FFEI decreases as the intercluster/intracluster distances increase/decrease. Hence, the lower the value of FFEI, the more crisp is the cluster structure. Note that the first two indices, class seperability and K-NN accuracy, measure the effectiveness of the feature subsets for classification, while the indices entropy and fuzzy feature evaluation index evaluate the clustering performance of the feature subsets. Let us now describe a quantitative index which measures the amount of redundancy present in the reduced subset.

6. Representation Entropy [2]. Let the eigenvalues of the $d \times d$ covariance matrix of a feature set of size d be $\lambda_j, j = 1, \ldots, d$. Let

$$\tilde{\lambda}_j = \frac{\lambda_j}{\sum_{i=1}^d \lambda_i}.$$

 $\tilde{\lambda_j}$ has similar properties like probability, namely, $0 \leq \tilde{\lambda_j} \leq 1$ and $\sum_{j=1}^d \lambda_j = 1$. Hence, an entropy function can be defined as

$$H_R = -\sum_{j=1}^d \tilde{\lambda}_j \log \tilde{\lambda}_j. \tag{4}$$

The function H_R attains a minimum value (zero) when all the eigenvalues except one are zero or, in other words, when all the information is present along a single coordinate direction. If all the eigenvalues are equal, i.e., information is equally distributed among all the features, H_R is maximum and so is the uncertainty involved in feature reduction. The above measure is known as representation entropy. It is a property of the data set as represented by a particular set of features, and is a measure of the amount of information compression possible by dimensionality reduction. This is equivalent to the amount of redundancy present in that particular representation of the data set. Since the

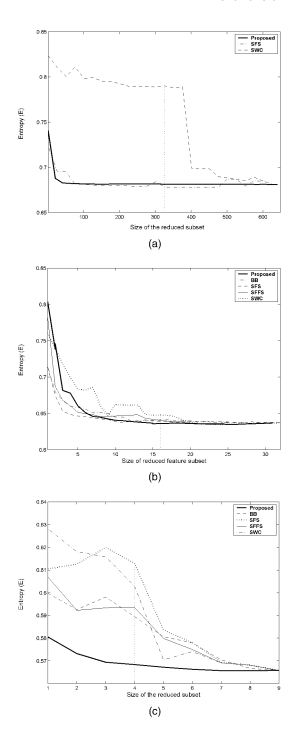


Fig. 3. Variation in classification accuracy with size of the reduced subset for—(a) Multiple features, (b) ionosphere, and (c) cancer data sets. The vertical dotted line marks the point for which results are reported in Tables 1, 2, and 3.

proposed algorithm involves partitioning of the original feature set into a number of homogeneous (highly compressible) clusters, it is expected that representation entropy of the individual clusters are as low as possible, while that of the final reduced set of features has low redundancy, i.e., a high value of representation entropy.

It may be noted that among all the *d*-dimensional subspaces of an original *D*-dimensional data set, the one corresponding to the Karhunen-Loeve coordinates [2] (for the first *d* eigenvalues) has the highest representation entropy, i.e., is least redundant. However, for large-dimensional data sets K-L transform directions are difficult to compute. Also, K-L transform results, in general, transformed variables and not exact subsets of the original features.

5 EXPERIMENTAL RESULTS AND COMPARISONS

Organization of the experimental results is as follows: First, the characteristics of the nine data sets used are discussed briefly. Then, performance of the proposed algorithm in terms of the feature evaluation indices, discussed in Section 4, is compared with five other feature selection schemes. Next, we have studied the redundancy reduction aspect of the algorithm quantitatively along with comparisons. Effect of varying the parameter k, used in feature clustering, is also studied.

Three categories of real-life public domain data sets are used: low-dimensional $(D \le 10)$, medium-dimensional $(10 < D \le 100)$, and high-dimensional (D > 100), containing both large and relatively smaller number of points. They are available from the UCI Machine Learning Repository [23]. Their characteristics are described below.

- 1. *Isolet*. The data consists of several spectral coefficients of utterances of English alphabets by 150 subjects. There are 617 features all real in the range [0, 1], 7,797 instances, and 26 classes.
- 2. *Multiple Features*. This data set consists of features of handwritten numerals ("0"-"9") extracted from a collection of Dutch utility maps. There are total 2,000 patterns, 649 features, and 10 classes.
- 3. Arrhythmia. It contains 452 samples, each having 279 attributes. Among the attributes 195 are real valued and are used for our experiments. The attributes represent parameters of ECG measurements and the task is to classify a patient into one of the 16 classes of cardiac Arrhythmia.
- 4. *Spambase.* The task is to classify an email into spam or nonspam category. There are 4,601 instances, 57 continuous valued attributes denoting word frequencies, and 2 classes.
- 5. Waveform. This consists of 5,000 instances having 40 attributes each. The attributes are continuous valued, and some of them are noise. The task is to classify an instance into one of the three categories of waves.
- 6. *Ionosphere*. The data represents autocorrelation functions of radar measurements. The task is to classify them into two classes denoting passage or obstruction in ionosphere. There are 351 instances and 34 attributes, all continuous.
- 7. Forest Cover Type. This is a GIS data set representing the forest covertype of a region. There are 54 attributes out of which we select 10 numeric valued attributes. There are 581,012 instances and eight classes.

Dataset	Method		CPU				
		FFEI	E	S	KNNA (%)	BayesA (%)	Time (sec)
					Mean SD	Mean SD	
Isolet	SFS	0.40	0.54	0.98	95.81 0.82	92.19 0.72	28.01×10^4
d=310, D=617	Proposed	0.40	0.50	1.07	96.00 0.78	95.01 0.52	440
Mult. Feat.	SFS	0.44	0.67	0.44	77.71 0.44	75.81 0.17	9.20×10^{4}
d=325, D=649	Proposed	0.48	0.68	0.45	78.34 0.22	75.28 0.10	451
Arrhythmia	SFS	0.40	0.77	0.21	53.22 0.59	52.25 0.44	2008
d=100, D=195	Proposed	0.40	0.72	0.17	58.93 0.54	56.00 0.41	74
	BB	0.40	0.65	0.90	64.03 0.41	63.55 0.40	9.21×10^{4}
Forest	SFFS	0.40	0.66	0.83	67.01 0.45	66.00 0.44	7.52×10^4
	SFS	0.43	0.66	1.01	61.41 0.44	60.01 0.41	17.19×10^3
d=5, D=10	Proposed	0.40	0.65	0.90	64.03 0.41	63.55 0.40	55.50

TABLE 4
Comparison of Feature Selection Algorithms for Large Data Sets when Search Algorithms use FFEI as the Selection Criterion

- 8. *Wisconsin Cancer*. The popular Wisconsin breast cancer data set contains nine features, 684 instances, and two classes.
- 9. *Iris.* The data set contains 150 instances, four features, and three classes of Iris flowers.

5.1 Comparison: Classification and Clustering Performance

Four indices, namely, entropy (2), fuzzy feature evaluation index (3), class seperability (1), and K-NN and naive Bayes classification accuracy are considered to demonstrate the efficacy of the proposed methodology and for comparing it with other methods. Four unsupervised feature selection schemes considered for comparison are:

- 1. Branch and Bound Algorithm (BB) [2]. A search method in which all possible subsets are implicitly inspected without exhaustive search. If the feature selection criterion is monotonic BB returns the optimal subset.
- 2. Sequential Forward Search (SFS) [2]. A suboptimal search procedure where one feature at a time is added to the current feature set. At each stage, the feature to be included in the feature set is selected from among the remaining available features so that the new enlarged feature set yields a maximum value of the criterion function used.
- 3. Sequential Floating Forward Search (SFFS) [3]. A near optimal search procedure with lower computational cost than BB. It performs sequential forward search with provision for backtracking.
- Stepwise Clustering (using correlation coefficient) (SWC) [21]. A nonsearch based scheme which obtains a reduced subset by discarding correlated features.

In our experiments, we have mainly used entropy (2) as the feature selection criterion with the first three search algorithms.

Comparisons in terms of five indices was made for different sizes of the reduced feature subsets. Tables 1, 2, and 3 provide such a comparative result corresponding to high, medium, and low-dimensional data sets when the size of the reduced feature subset is taken to be about half of the original size as an example. Comparison for other sizes of the reduced feature set is provided in Fig. 3 considering one data set from each of the three categories, namely, multiple features (high), ionosphere (medium), and cancer (low). The CPU time required by each of the algorithms on a Sun UltraSparc 350 MHz workstation are also reported in Tables 1, 2, and 3. Since the branch and bound (BB) and the sequential floating forward search (SFFS) algorithms require infeasibly high computation time for the large data sets, we could not provide the figures for them in Table 1. For the classification accuracies (using K-NN and Bayes), both mean and standard deviations (SD) computed for ten independent runs are presented.

Compared to the search-based algorithms (BB, SFFS, and SFS), the performance of the proposed scheme is comparable or slightly superior, while the computational time requirement is much less for the proposed scheme. On the other hand, compared to the similarity based SWC method the performance of the proposed algorithm is much superior, keeping the time requirement comparable. It is further noted that the superiority in terms of computational time increases as the dimensionality and sample size increase. For example, in the case of low-dimensional data sets the speedup factor of the proposed scheme compared to BB and SFFS algorithms is about 30-50, for Forest data

Dataset	ВВ	SFFS	SFS	SWC	Relief-F	Proposed
Isolet	-	-	2.91	2.87	2.89	3.50
Mult. Ftrs.	-	-	2.02	1.90	1.92	3.41
Arrhythmia	-	-	2.11	2.05	2.02	3.77
Spambase	2.02	1.90	1.70	1.44	1.72	2.71
Waveform	1.04	1.02	0.98	0.81	0.92	1.21
Ionosphere	1.71	1.71	1.70	0.91	1.52	1.81
Forest	0.91	0.82	0.82	0.77	0.91	0.91
Cancer	0.71	0.71	0.55	0.55	0.59	0.82
Iris	0.47	0.47	0.41	0.31	0.47	0.47

 ${\it TABLE 5} \\ {\it Representation Entropy H_R^s of Subsets Selected using Some Algorithms}$

which is low-dimensional but has large sample size the factor is about 100, for medium-dimensional data sets, BB and SFFS are about 100 times slower and SFS about ten times slower, while for the high-dimensional data sets SFS is about 100 times slower, and BB and SFFS could not be compared as they require infeasibly high run time.

It may be noted that the aforesaid unsupervised feature selection algorithms (namely, BB, SFFS, SFS) usually consider "entropy" as the selection criterion. Keeping this in mind detailed results are provided in Tables 1, 2, and 3. However, we have also run the experiments using another unsupervised measure, namely, fuzzy feature evaluation index (FFEI) (3). Table 4 shows, as an illustration, the results only for the four large data sets (Isolet, Multiple Features,

Arrhythmia, and Forest Covertype). These results corroborate the findings obtained using entropy.

In a part of the experiments, we compared the performance with a supervised method Relief-F, which is widely used. We have used 50 percent of the samples as design set for the Relief-F algorithm. Results are presented in Tables 1, 2, and 3. The Relief-F algorithm provides classification performance comparable to the proposed scheme inspite of using class label information. Moreover, it has much higher time requirement, specially for data sets with large number of samples, e.g., the Forest data. Its performance in terms of the unsupervised indices is also poor.

Statistical significance of the classification performance of the proposed method compared to those of the other algorithms is tested. Means and SD values of the accuracies,

TABLE 6	
Redundacy Reduction using Different Feature Similarity Mea	sures

Dataset	Similarity Measure: λ_2		Simila	rity Measure: e	Similarity Measure: ρ		
	H_R^g	H_R^s	H_R^g	H_R^s	H_R^g	H_R^s	
Isolet	0.001	3.50	0.007	3.01	0.003	3.41	
Mult. Ftrs.	0.002	3.41	0.008	2.95	0.007	3.01	
Arrhythmia	0.007	3.77	0.017	2.80	0.010	3.41	
Spambase	0.04	2.71	0.07	2.01	0.05	2.53	
Waveform	0.10	1.21	0.14	1.04	0.11	1.08	
Ionosphere	0.05	1.81	0.07	1.54	0.07	1.54	
Forest	0.10	0.91	0.17	0.82	0.11	0.91	
Cancer	0.19	0.82	0.22	0.71	0.19	0.82	
Iris	0.17	0.47	0.22	0.31	0.17	0.47	

 H_R^g : Average representation entropy of feature groups, H_R^s : representation entropy of selected subset, λ_2 : maximal information compression index, e: least-square regression error, and ρ : correlation coefficients.

computed over 10 independent runs, are used for this purpose. A generalized version of paired *t*-test suitable for both inequal means and variances is used. The above problem is the classical Behrens-Fisher problem in hypothesis testing, a suitable test statistic is described and tabled in [24] and [25], respectively. It is observed that the proposed method has significantly better performance compared to the SWC algorithm for all the data sets, and the SFS algorithm for most of the data sets. For the other algorithms, namely Relief-F, BB, and SFFS, the performance is comparable, i.e., the difference of the mean values of the classification scores is statistically insignificant.

5.2 Redundancy Reduction: Quantitative Study

As mentioned before, the proposed algorithm involves partitioning the original feature set into certain number of homogeneous groups and then replacing each group by a single feature, thereby resulting in the reduced feature set. Representation entropy (H_R) , defined in Section 4, is used to measure the redundancy in both the homogeneous clusters and the final selected feature subset. H_R when computed over the individual clusters should be as low as possible (indicating high redundancy among the features belonging to a single cluster), while giving as high value as possible for the selected subset (indicating minimum redundancy). Let us denote the average value of H_R computed over the homogeneous groups by H_R^g and the value of H_R for the final selected subset by H_R^g .

Table 5 shows the comparative results of the proposed method with other feature selection algorithms in terms of H_R^s . It is seen that the subset obtained by the proposed scheme is least redundant having the highest H_R^s values.

To demonstrate the superiority of the maximal information compression index λ_2 , compared to the other two feature similarity measures (ρ and e) used previously, we provide Table 6, where we have compared both H_R^s and H_R^g values obtained using each of the similarity measures, in our clustering algorithm. It is seen from Table 6 that, λ_2 has superior information compression capability compared to the other two measures as indicated by the lowest and highest values of H_R^g and H_R^s , respectively.

5.3 Effect of Parameter k

In our algorithm, the size of the reduced feature subset and hence, the scale of details of data representation is controlled by the parameter k. Fig. 4 illustrates such an effect for three data sets—multiple features, ionosphere, and cancer, considering one data from each of the high, medium, and low categories. As expected, the size of the reduced subset decreases overall with increase in k. However, for medium and particularly large-dimensional data (Fig. 4a), it is observed that for certain ranges of k at the

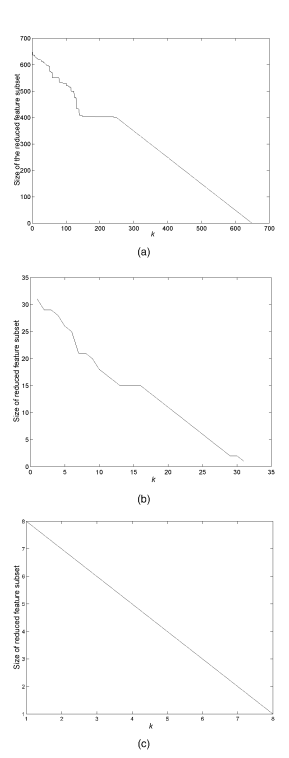


Fig. 4. Variation in size of the reduced subset with parameter k for—(a) multiple features, (b) ionosphere, and (c) cancer data.

lower side, there is no change in the size of the reduced subset, i.e., no reduction in dimension occurs. Another interesting fact observed in all the data sets considered is that, for all values of k in the case of small dimensional data sets, and for high values of k in the case of medium and large-dimensional data sets, the size of the selected subset varies linearly with k. Further, it is seen in those cases,

^{1.} The test statistic is of the form $v=\frac{\bar{x}_1-\bar{x}_2}{\sqrt{\lambda_1s_1^2+\lambda_2s_2^2}}$, where \bar{x}_1,\bar{x}_2 are the means, s_1,s_2 the standard deviations, and $\lambda_1=1/n_1,\lambda_2=1/n_2,\,n_1,n_2$ are the number of observations.

 $d + k \approx D$, where *d* is the size of the reduced subset and *D* is the size of the original feature set.

CONCLUSION AND DISCUSSIONS

An algorithm for unsupervised feature selection using feature similarity measures is described. The novelty of the scheme, as compared to other conventional feature selection algorithms, is the absence of search process which contributes to the high-computational time requirement of those feature selection algorithms. Our algorithm is based on pairwise feature similarity measures, which are fast to compute. It is found to require several orders less CPU time compared to other schemes. Unlike other approaches, which are based on optimizing either classification or clustering performance explicitly, here we determine a set of maximally independent features by discarding the redundant ones. This enhances the applicability of the resulting features to compression and other tasks like forecasting, summarization, and association mining in addition to classification/clustering. Another characterstics of the proposed algorithm is its capability for multiscale representation of data sets. The scale parameter k used for feature clustering efficiently parametrizes the tradeoff between representation accuracy and feature subset size. All these make it suitable for a wide variety of data mining tasks involving large (in terms of both dimension and size) data sets.

Besides formulating the novel clustering algorithm, we have defined a feature similarity measure called maximal information compression index. One may note that the definition of the said parameter is not new, it is its use in feature subset selection framework which is novel. The superiority of this measure for feature selection is established experimentally. It is also demonstrated through extensive experiments that representation entropy can be used as an index for quantifying both redundancy reduction and information loss in a feature selection method.

In the present article, we have measured the information loss in terms of second order statistics. The similarity measure used for the feature selection algorithm is selected/defined accordingly. One may modify these measures suitably in case even higher order statistics are used. In this regard, modifications of correlation indices [22] which measure higher order polynomial dependency between variables may be considered. Also, the similarity measure is valid only for numeric features; its extension to accomodate other kinds of variables (e.g., symbolic, categorical, hybrid) may also be investigated.

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