Gene Microarray Cancer Classification using Correlation Based Feature Selection Algorithm and Rules Classifiers

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Abstract—Gene microarray classification problems are considered a challenge task since the datasets contain few number of samples with high number of genes (features). The genes subset selection in microarray data play an important role for minimizing the computational load and solving classification problems. In this paper, the Correlation-based Feature Selection (CFS) algorithm is utilized in the feature selection process to reduce the dimensionality of data and finding a set of discriminatory genes. Then, the Decision Table, JRip, and OneR are employed for classification process. The proposed approach of gene selection and classification is tested on 11 microarray datasets and the performances of the filtered datasets are compared with the original datasets. The experimental results showed that CFS can effectively screen irrelevant, redundant, and noisy features. In addition, the results for all datasets proved that the proposed approach with a small number of genes can achieve high prediction accuracy and fast computational speed. Considering the average accuracy for all the analysis of microarray data, the JRip achieved the best result as compared to Decision Table, and OneR classifier. The proposed approach has a remarkable impact on the classification accuracy especially when the data is complicated with multiple classes and high number of genes.

Keywords—Feature selection, gene expression data, Correlation-based Feature Selection algorithm, Decision Table, JRip, and OneR.

1 Introduction

Cancer is considered as one of the dreadful diseases and diagnosis of cancer is very important in initial stage for its proper treatment [11]. Cancer data is a collection of thousands of genes and DNA microarray is used to determine the expression level of genes [21]. Microarray gene selection and classification is considered very challenges task since the datasets are large and abundant in noisy genes [12]. The main problem in microarray datasets arises from the fact that the genes greatly outnumber the sample observations [27]. Thus, feature selection methods is needed in microarray cancer datasets so as to select suitable feature set that makes the classifier more accurate and faster [7]. The main benefits of using feature selection is improved the performance of...
the classifier by minimizing redundancy, removing noisy data and maximizing the relevance [8]. The relevance and the redundancy level of a gene are often measured by correlation coefficients such as: Symmetrical Uncertainty [13], Mutual Information (MI) [32], Pearson’s correlation [6][28].

Problems in high-dimensional data analysis have motivated the researchers to search for possible solutions and propose viable algorithms [14]. Alternatively, different meta-heuristic algorithms have been adapted for feature selection issues [19][29]. Examples of these algorithms are Principle Component Analysis [34], Genetic Algorithm [3], Ant Colony Optimization [9], Simulated Annealing [16] and Particle Swarm Optimization [5][33].

Correlation-based Feature Selection (CFS) is a simple filter algorithm that ranks feature subsets according to a correlation-based heuristic evaluation function [38]. CFS is one of the most well-known feature selection algorithms that take advantage of a redundancy filter. The CFS evaluates subsets of features using the hypothesis “Good feature subset contain features that are highly correlated with the classification and yet uncorrelated to each other” [39].

CFS evaluates the worth of a subset of attributes by considering the individual predictive ability of each feature along with the degree of redundancy between them [19]. Correlation coefficients is used to estimate correlation between subset of attributes and class, as well as inter-correlations between the features. Relevance of a group of features grows with the correlation between features and classes, and decreases with growing inter-correlation. CFS is used to determine the best feature subset and is usually combined with search strategies such as forward selection, backward elimination, bi-directional search, best-first search and genetic search. In this paper, Greedy Stepwise is used as search method with CFS algorithm.

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Background

Nowadays, various kinds of machine learning and statistical approaches are used to classify tumour cells accurate such as support vector machines [15], k-nearest neighbor [30], and neural network techniques [24]. Also, several researchers thus hybridize the classification techniques with optimization algorithms for further enhancement of accuracy [1][22].

Two-phase hybrid model is suggested in [23] for cancer classification, integrating Correlation-based Feature Selection (CFS) with improved-Binary Particle Swarm Optimization (iBPSO) using Naive-Bayes classifier as the only classifier.

The Pearson’s Correlation Coefficient (PCC) [18] is used in combination with Binary Particle Swarm Optimization (BPSO) or Genetic Algorithm (GA) along with various classifiers for the selection and classification of high dimensional microarray data. It was noticed that the PCC filter showed a remarkable improvement in the classification accuracy when it was combined with BPSO or GA. Also, the results show that the BPSO is working faster and has better performance than GA.

Additionally, the Probabilistic Attribute-Value have used in [4] for Class Distinction (Pavicd) algorithm as a feature selection in microarray cancer datasets. The
Pavicd algorithm works on the space of feature values instead of the features’ space. Experiments show that Pavicd gets the best performance in terms of running time and classification accuracy when using Ripper-k and C4.5 as classifiers.

A Collaborative Representation (CR)-based classification with regularized least square was developed [31] to classify gene data. The CR codes a testing sample as a sparse linear combination of all training samples and then classifies the testing sample by evaluating which class leads to the minimum representation error. Experiments results on several diseases show that the CR-based algorithm achieves high classification accuracy and fast computational speed than the traditional classifiers, such as support vector machine algorithm.

The K-Nearest Neighbor (K-NN) classifier and feature selection using ANOVA test was developed based on MapReduce programming model [26]. The approach works in a distributed manner on scalable clusters. The algorithms are successfully implemented on Hadoop framework and comparative analysis is done using various microarray datasets.

A multi-test decision tree (MTDT) was applied for solving biological problems [10]. The application of several univariate tests in each non-terminal node of the decision tree is considered. Comparison results with eight classifiers show that MTDT has a statistically significantly higher accuracy than popular decision tree classifiers, and it was highly competitive with ensemble learning algorithms.

Moreover, the Support Vector Machine (SVM) classifier was applied on four microarray datasets [2]. The study analyzed two different kernels of SVM; radial kernel and linear kernels. The results showed that the SVM exceeded the performance and accuracy compared to K-nearest neighbor (KNN) and neural network (NN).

3 The Proposed Work

In this paper, 11 different high dimensional datasets are applied. The Correlation-based Feature Selection (CFS) with Greedy Stepwise search method is proposed for genes selection. Also, multiple classifiers are utilized to show the quality of each of them.

3.1 Datasets

The performance of three classifiers; Decision Table, JRip, and OneR are investigated using eleven (11) publicly available microarray datasets [37]. A brief overview of these datasets such as number of gene, number of instance, and number of class is summarized in Table 1.
Table 1. Gene microarray datasets

| Dataset       | Gene number | Instance number | Class number                                                                                     |
|---------------|-------------|-----------------|---------------------------------------------------------------------------------------------------|
| Breast Cancer | 24481       | 97              | 2 classes, 46 relapse, 51 non-relapse                                                             |
| CNS           | 7129        | 60              | 2 types, 21 survivors, 39 failures                                                                |
| Colon Tumor   | 2000        | 62              | 2 types, 40 Tumor, 22 Normal                                                                       |
| Leukemia      | 7129        | 72              | 2 types of acute leukemia, 47 Acute Lymphoblastic Leukemia (ALL), 25 Acute Myeloid Leukemia (AML) |
| Leukemia_3C   | 7129        | 72              | 3 types of acute leukemia, 38 B-cell ALL, 9 T-cell ALL, 25 AML                                     |
| Leukemia_4C   | 7129        | 72              | 4 types of acute leukemia, 38 B-cell, 9 T-cell, 21 BM AML, 4 PB AML                                 |
| Lung Cancer   | 12600       | 203             | 5 types, 139 adenocarcinoma (AD), 17 normal lung (NL), 6 small cell lung cancer (SMCL), 21 squamous cell carcinoma (SQ), 20 pulmonary carcinoid (COID). |
| Lymphoma      | 4026        | 66              | 3 different adult lymphoid malignancies, 46 diffuse large B-cell lymphoma (DLBCL), 9 Follicular Lymphoma (FL), 11 Chronic Lymphocytic Leukemia (CLL). |
| MLL           | 12582       | 72              | 3 types, 24 acute lymphoblastic leukemia (ALL), 20 Mixed- Lineage Leukemia (MLL), 28 acute myeloblastic leukemia (AML). |
| Ovarian Cancer| 15154       | 253             | 2 types, 162 Cancer, 91 Normal                                                                     |
| SRBCT         | 2308        | 83              | 4 different cases, 29 Ewing sarcoma (EWS), 11 Burkitt lymphoma (BL), 18 neuroblastoma (NB), 25 rhabdomyosarcoma (RMS). |

These datasets are related to studies of human cancer, including Breast Cancer, Central Nervous System (CNS), Colon Tumor, Leukemia, Leukemia_3C, Leukemia_4C, Lung Cancer, Lymphoma, Mixed Lineage Leukemia (MLL), Ovarian Cancer, and Small Round Blue-Cell Tumor (SRBCT).

### 3.2 Correlation based feature selection algorithm

A common procedure for choosing the foremost applicable characteristics within the dataset is to use correlation. Correlation is additional strictly remarked as Pearson’s coefficient of correlation in statistics. We are able to compute the correlation among every characteristic and also the output variable and choose solely those characteristics that have a moderate-to-high positive or correlational statistics (close to -1 or 1) and drop those characteristics with a coffee correlation (value near zero).

Correlation-based feature selection (CFS) ranks characteristics in keeping with a heuristic analysis operate supported correlations [17]. The operate gauges subsets product of characteristic vectors, that are correlative with the category label, however freelance of every alternative.

The CFS methodology assumes and accepts that impertinent options show a coffee correlation with the category and thus ought to be unheeded by the algorithmic rule. On the opposite hand, excess options ought to be examined, as they’re typically powerfully correlative with one or additional of the opposite attributes [36].
3.3 Classification model

In this paper, three classifiers are applied: Decision Table, JRip, and OneR. The choice of various classifiers is due to the fact that there is no any specific classifier to work perfectly for all datasets and not all classifiers work in the same way on a dataset.

- Decision Table constructs a decision table majority classifier [25]. It evaluates feature subsets using best-first search and can use cross-validation for evaluation.
- JRip implements RIPPER, including heuristic global optimization of the rule set [35].
- OneR is the 1R classifier with one parameter: the minimum bucket size for discretization [20].

4 Experimental Design and Results Discussion

Initially, the Decision Table, JRip, and OneR classifiers were applied on the original datasets. Then, the all eleven datasets were filtered using Correlation-based Feature Selection (CFS) algorithm. After that, the filtered datasets were tested against the applied classifiers. This was done in order to compare the classification accuracy of the dataset with the one before filtration. In each dataset, experiment was performed in full training method and 2-folds to 10-folds cross validation.

The features of the datasets were filtered and the number of selected genes using CFS is tabulated in Table 2. From the results, we can notice that CFS has selected fewer number of genes compared to that of original dataset. The results show that the number of selected genes for Breast Cancer is reduced from 24481 to 138, CNS from 7129 to 39, Colon Tumor from 2000 to 26, Leukemia from 7129 to 79, Leukemia_3C from 7129 to 104, Leukemia_4C from 7129 to 119, Lung Cancer from 12600 to 548, Lymphoma from 4026 to 175, MLL from 12582 to 142, Ovarian Cancer from 15154 to 35, and SRBCT from 2308 to 112 genes.

Table 2. Number of selected genes before/after applying CFS algorithm

| Dataset       | #Total Genes | #Gene After CFS |
|---------------|--------------|-----------------|
| Breast Cancer | 24481        | 138             |
| CNS           | 7129         | 39              |
| Colon Tumor   | 2000         | 26              |
| Leukaemia     | 7129         | 79              |
| Leukaemia-3C  | 7129         | 104             |
| Leukaemia-4C  | 7129         | 119             |
| Lung Cancer   | 12600        | 548             |
| Lymphoma      | 4026         | 175             |
| MLL           | 12582        | 142             |
| Ovarian Cancer| 15154        | 35              |
| SRBCT         | 2308         | 112             |
The accuracy of the classifiers applied on the original and filtered datasets was evaluated as shown in Table 3. Results in bold indicate the best performed classifier for each specific dataset. The results show that generally the accuracy of the classifiers on the filtered dataset performed better results when compared with those applied directly on the original datasets. However, there are some cases with few classifiers in which the accuracy on the original dataset is same as filtered dataset.

In addition, the results show that the Decision Table and JRip are the best classifiers for the datasets, while the OneR is the worst classifier. The Breast Cancer had the highest accuracy of 88.7% with JRip, CNS (90.0% with Decision Table), Colon (96.8% with Decision Table and JRip), Leukemia (98.6% with Decision Table), Leukemia_3C (100.0% with Decision Table), Leukemia_4C (98.6% with Decision Table and JRip), Lung (97.5% with JRip), Lymphoma (100.0% with JRip), MLL (95.8% with JRip), Ovarian (100.0% with Decision Table), and SRBCT (97.6% with JRip).

### Table 3. Accuracy for the original and filtered microarray datasets

| Datasets      | Classifier       | 2-Fold 5-Fold 7-Fold 9-Fold 10-Fold Average |
|---------------|------------------|-----------------------------------------------|
| **Breast Cancer** | Decision Table   | 79.4 64.9 50.5 57.7 64.9 50.5 63.9 60.8 55.7 61.9 69.0 |
|               | Decision Table-CFS | 85.6 68.0 66.0 59.8 75.3 73.2 67.0 62.9 72.2 67.0 69.7 |
|               | JRip              | 84.5 58.8 53.6 57.7 56.7 49.5 54.6 59.8 56.7 66.0 59.8 |
|               | Jrip-CFS          | 88.7 62.9 73.2 59.8 68.0 70.1 71.1 69.1 59.8 68.0 69.1 |
|               | OneR              | 82.5 55.7 46.4 45.4 49.5 52.6 54.6 51.5 42.3 56.7 53.7 |
|               | OneR-CFS          | 81.4 58.8 53.6 59.8 54.6 53.6 61.9 54.6 58.8 57.7 59.5 |
| **CNS**       | Decision Table   | 90.0 63.3 56.7 56.7 51.7 63.3 60.0 56.7 75.0 63.0  |
|               | Decision Table-CFS | 90.0 63.3 71.7 71.7 73.3 66.7 81.7 68.3 70.0 76.7 73.3 |
|               | JRip              | 75.0 61.7 68.3 40.0 68.3 51.7 56.7 60.0 60.0 58.3 60.0 |
|               | Jrip-CFS          | 88.3 71.7 80.0 63.3 68.3 70.0 68.3 71.7 68.3 65.0 71.5 |
|               | OneR              | 88.3 60.0 55.0 55.0 48.3 48.3 56.7 65.0 53.3 48.3 57.8 |
|               | OneR-CFS          | 83.3 53.3 76.7 63.3 70.0 58.3 66.7 65.0 58.3 65.0 66.0 |
| **Colon**     | Decision Table   | 96.8 62.9 72.6 71.0 69.4 72.6 75.8 72.6 72.6 77.4 74.4 |
|               | Decision Table-CFS | 96.8 67.7 85.5 80.6 80.6 83.9 80.6 88.7 83.9 82.9 92.9 |
|               | JRip              | 85.5 71.0 75.8 75.8 67.7 75.8 74.2 72.6 69.4 75.8 74.4 |
|               | Jrip-CFS          | 96.8 88.7 74.2 82.3 80.6 72.6 82.3 87.1 82.3 77.4 82.4 |
|               | OneR              | 88.7 51.6 75.8 74.2 71.0 66.1 79.0 71.0 62.9 69.4 71.0 |
|               | OneR-CFS          | 88.7 64.5 77.4 82.3 79.0 77.4 79.0 71.0 77.4 74.2 77.1 |
| **Leukemia**  | Decision Table   | 98.6 76.4 88.9 80.6 90.3 83.3 81.9 81.9 84.7 83.3 85.0 |
|               | Decision Table-CFS | 98.6 76.4 90.3 81.9 91.7 83.3 87.5 83.3 86.1 88.9 86.8 |
|               | JRip              | 95.8 76.4 87.5 80.6 91.7 83.3 84.7 84.7 87.5 87.5 86.0 |
|               | Jrip-CFS          | 95.8 76.4 90.3 84.7 95.8 83.3 90.3 84.7 87.5 91.7 88.1 |
|               | OneR              | 95.8 76.4 87.5 84.7 87.5 90.3 80.6 79.2 86.1 81.9 85.0 |
|               | OneR-CFS          | 95.8 76.4 87.5 84.7 87.5 90.3 80.6 77.8 86.1 83.3 85.0 |
| **Leukemia_3C** | Decision Table   | 100.0 72.2 69.4 76.4 81.9 90.3 70.8 86.1 84.7 84.7 81.7 |
|               | Decision Table-CFS | 98.6 81.9 72.2 79.2 81.9 90.3 84.7 84.7 81.9 81.9 83.7 |
|               | JRip              | 98.6 76.4 77.8 80.6 86.1 88.9 90.3 91.7 90.3 90.3 87.1 |
| Datasets | Classifier | Full Train | 2-Fold | 3-Fold | 4-Fold | 5-Fold | 6-Fold | 7-Fold | 8-Fold | 9-Fold | 10-Fold | Average |
|----------|------------|------------|--------|--------|--------|--------|--------|--------|--------|--------|--------|---------|
| Leukemia-4C | JRip-CFS | 98.6 | 79.2 | 83.3 | 83.3 | 91.7 | 91.7 | 94.4 | 94.4 | 94.4 | 94.4 | 90.6 |
|           | OneR-CFS  | 86.1 | 58.3 | 77.8 | 69.4 | 70.8 | 75.0 | 69.4 | 68.1 | 72.2 | 69.4 | 71.7 |
|           | Decision Table | 98.6 | 52.8 | 66.7 | 69.4 | 83.3 | 75.0 | 76.4 | 73.6 | 79.2 | 77.8 | 75.3 |
| SRBCT     | JRip-CFS  | 98.6 | 75.0 | 72.2 | 76.4 | 87.5 | 81.9 | 77.8 | 79.2 | 80.6 | 79.2 | 80.8 |
| Lung      | OneR-CFS  | 83.3 | 63.9 | 70.8 | 70.8 | 77.8 | 72.2 | 69.4 | 68.1 | 76.4 | 76.4 | 72.9 |
| Lymphoma  | JRip-CFS  | 98.6 | 72.2 | 83.3 | 87.5 | 80.6 | 83.3 | 87.5 | 90.3 | 90.3 | 90.3 | 95.3 |
|           | Decision Table | 98.6 | 50.0 | 51.8 | 53.3 | 64.4 | 61.5 | 62.5 | 64.8 | 67.2 | 64.8 | 61.1 |
| MLL       | OneR-CFS  | 83.3 | 65.3 | 70.8 | 70.8 | 77.8 | 72.2 | 69.4 | 68.1 | 76.4 | 76.4 | 72.9 |
| Ovarian   | JRip-CFS  | 98.6 | 72.2 | 83.3 | 87.5 | 80.6 | 83.3 | 87.5 | 90.3 | 90.3 | 90.3 | 95.3 |
| SRBCT     | Decision Table | 98.6 | 50.0 | 51.8 | 53.3 | 64.4 | 61.5 | 62.5 | 64.8 | 67.2 | 64.8 | 61.1 |

Furthermore, it is noticeable that the highest average accuracy is achieved using JRip-CFS as compared to other classifiers. For example, Lymphoma data had the highest accuracy of 93.8% with JRip-CFS, and 86.1% with JRip, while the accuracy
was 85.5% with Decision Table-CFS, 83.6% with Decision Table, 84.5% with OneR-CFS, and 82.6% with OneR.

Considering the datasets analysis in full training method and 2-folds to 10-folds cross validation. The results showed that the JRip exceeded the performance and accuracy compared to Decision Table and OneR. As example, Figure 1 shows the classification accuracy for the SRBCT dataset using the 3 classifiers in the ten different tests. The results in Figure 1 prove that JRip combined with CFS was the best functional method since it yielded better results than the other methods.

Moreover, considering the average accuracy for the 11 datasets as shown in Table 4. The average accuracy of all 11 data was 78.2% (Decision Table), 82.4% (Decision Table-CFS), 80.7% (JRip), 86.0% (Jrip-CFS), 73.1% (OneR), and 75.3% (OneR-CFS). It is clear that the classifier’s accuracy is improved after selection process using CFS. Also, we have clearly noticed that the average accuracy of JRip was better than Decision Table and OneR.

### Table 4. Average accuracy for the 11 microarray data

| Classifier          | Accuracy for the full training and cross validation method |
|---------------------|-----------------------------------------------------------|
|                     | Full Tran | 2-Fold | 3-Fold | 4-Fold | 5-Fold | 6-Fold | 7-Fold | 8-Fold | 9-Fold | 10-Fold | Average |
| Decision Table      | 61.0      | 63.0   | 74.4   | 85.0   | 81.7   | 75.3   | 83.7   | 83.6   | 83.9   | 96.8     | 72.2    |
| Decision Table-CFS  | 69.7      | 73.3   | 82.9   | 86.8   | 83.7   | 80.8   | 86.5   | 85.5   | 86.5   | 96.9     | 73.9    | 82.4    |
| JRip                | 59.8      | 60.0   | 74.4   | 86.0   | 87.1   | 81.0   | 88.3   | 86.1   | 82.2   | 97.2     | 85.7    | 80.7    |
| Jrip-CFS            | 69.1      | 71.5   | 82.4   | 88.1   | 90.6   | 85.3   | 91.6   | 93.8   | 87.1   | 97.3     | 89.5    | 86.0    |
| OneR                | 53.7      | 57.8   | 71.0   | 85.0   | 71.7   | 72.9   | 75.3   | 82.6   | 78.5   | 96.9     | 58.3    | 73.1    |
| OneR-CFS            | 59.5      | 66.0   | 77.1   | 85.0   | 74.4   | 73.1   | 76.1   | 84.5   | 79.0   | 96.9     | 56.7    | 75.3    |
5 Conclusion

Usually, microarray data is characterized by noisiness as well as increased dimensionality. Therefore, selecting relevant genes is an imperative in microarray data analysis. In this paper, CFS is proposed to select the relevant features. Also, Decision Table, JRip, and OneR classifiers are proposed to classify the microarray data. The comparative analysis proved that the accuracy of all classifiers is improved using filtered datasets compared with their accuracy on the original datasets. This indicates that the feature selection by CFS not only improved the efficiency of the classification process but also its accuracy is enhanced. Furthermore, it can be seen that JRip has presented the highest classification accuracy among all the other classifiers. Further, this paper can be extended by considering the applicability of another features selection techniques such as Genetic Algorithm, Principle Component Analysis, Simulated Annealing, Ant Colony Optimization, and Particle Swarm Optimization.

6 References

[1] Agrawal S, Agrawal J. Neural network techniques for cancer prediction: A survey. Procedia Computer Science. 2015; 60:769–774. https://doi.org/10.1016/j.procs.2015.08.234
[2] Almoeirifi, M, (2015). Classification Approaches for Microarray Gene Expression Data Analysis. Master thesis. Laurentian University, Sudbury, Ontario, Canada.
[3] Alshamlan, H.M., G.H. Badr and Y.A. Alohal, 2015. Genetic Bee Colony (GBC) algorithm: A new gene selection method for microarray cancer classification. Computational Biology Chemistry, 56: 49-60. https://doi.org/10.1016/j.compbiolchem.2015.03.001
[4] Angulo, A.P. Gene Selection for Microarray Cancer Data Classification by a Novel Rule-Based Algorithm. Information 2018, 9, 6; https://doi.org/10.3390/info9010006.
[5] Ardjani, F., K. Sadouni and M. Benyettou, 2010. Optimization of SVM multiclass by particle swarm (PSO-SVM). Proceedings of the 2nd International Workshop on Database Technology and Applications, Nov. 27-28, IEEE Xplore press, Wuhan, China. https://doi.org/10.1109/dbta.2010.5658994
[6] Bolón-Canedo, V., N. Sánchez-Marono, A. Alonso-Betanzos, J.M. Benítez and F. Herrera, 2014. A review of microarray datasets and applied feature selection methods. Information Sci., 282: 111-135. https://doi.org/10.1016/j.ins.2014.05.042
[7] Annavarapu, C.S., S. Dara, H. Banka, 2016. Cancer microarray data feature selection using multi-objective binary particle swarm optimization algorithm. EXCLI J., 1;15:460-473. DOI: 10.17179/excli2016-481
[8] Chen Y, Li Y, Narayan R, Subramanian A, Xie X. Gene expression inference with deep learning. Bioinformatics. 2016; 32(12):1832–1839. PMID: 26873929 https://doi.org/10.1093/bioinformatics/btw074
[9] Chen, Y., D. Miao and R. Wang, 2010. A rough set approach to feature selection based on ant colony optimization. Pattern Recognition Lett., 31: 226-233. https://doi.org/10.1016/j.patrec.2009.10.013
[10] Czajkowski M, Grzes M, Kretowski M. Multi-test decision tree and its application to microarray data classification. Artificial Intelligence in Medicine. 2014; 61(1):35–44. doi: https://doi.org/10.1016/j.artmed.2014.01.005
[11] Dash, R., 2018. An adaptive harmony search approach for gene selection and classification of high dimensional medical data. J. King Saud University-Computer and Information Sciences. https://doi.org/10.1016/j.ksuci.2018.02.013

[12] Dashbhan, M., M. Balafar and P. Suravajhala, 2018. Gene selection for tumor classification using a novel bio-inspired multi-objective approach. Genomics, 110: 10-17. https://doi.org/10.1016/j.ygeno.2017.07.010

[13] Fakhoury M. Autistic spectrum disorders: A review of clinical features, theories and diagnosis. International Journal of Developmental Neuroscience. 2015; 43:70–77. PMID: 25862937 https://doi.org/10.1016/j.ijdevneu.2015.04.003

[14] Ferreira, L.G.; dos Santos, R.N.; Oliva, G.; Andricopulo, A.D. Molecular Docking and Structure-Based Drug Design Strategies. Molecules 2015, 20, 13384–13421. https://doi.org/10.3390/molecules200713384

[15] Ghaddar, B. and J. Naoum-Sawaya, 2018. High dimensional data classification and feature selection using support vector machines. Eur. J. Operational Res., 265: 993-1004. https://doi.org/10.1016/j.ejor.2017.08.040

[16] Gonzalez-Navarro, F. and L.A. Belanche-Muñoz, 2014. Feature selection for microarray gene expression data using simulated annealing guided by the multivariate joint entropy. Inter. J. Comput. Sci. Applications. https://doi.org/10.13053/cys-18-2-1473

[17] Hall, A.M. Correlation-based feature selection for machine learning, Tech. Rep., Doctoral Dissertation, University Of Waikato, Department of Computer Science, 1999.

[18] Hameed, S.S., F.F. Muhammad, R. Hassan and F. Saeed, 2018. Gene Selection and Classification in Microarray Datasets using a Hybrid Approach of PCC-BPSO/GA with Multi Classifiers. Journal of Computer Science, Volume 14, Issue 6, Pages 868-880. https://doi.org/10.3844/jcssp.2018.868.880

[19] Hira, Z. M., and Gillies, D. F., A Review of Feature Selection and Feature Extraction Methods Applied on Microarray Data, Adv Bioinformatics. 2015: 198363. https://doi.org/10.1155/2015/198363.

[20] Holte, R.C. (1993). Very Simple Classification Rules Perform Well on Most Commonly Used Datasets, Mach Learn, 11, 63-91.

[21] Hong, H.J.; Koom, W.S.; Koh, W.-G. Cell Microarray Technologies for High-Throughput Cell-Based Biosensors. Sensors 2017, 17, 1293. https://doi.org/10.3390/s17061293

[22] Huang HH, Liu XY, Liang Y. Feature Selection and Cancer Classification via Sparse Logistic Regression with the Hybrid L1/2+2 Regularization. PloS one. 2016; 11(5):e0149675. https://doi.org/10.1371/journal.pone.0149675 PMID: 27136190 https://doi.org/10.1371/journal.pone.0149675

[23] Jain, I., V.K. Jain and R. Jain, 2018. Correlation feature selection based improved-Binary Particle Swarm Optimization for gene selection and cancer classification. Applied Soft Computing, 62: 203-215. https://doi.org/10.1016/j.asoc.2017.09.038

[24] Khan J, Wei JS, Ringner M, Saal LH, Ladanyi M, Westermann F, et al. Classification and diagnostic prediction of cancers using gene expression profiling and artificial neural networks. Nature medicine. 2001; 7(6):673. https://doi.org/10.1038/89044 PMID: 11385503

[25] Kohavi, R. (1995). The power of decision tables, Machine Learning: Ecml95, 912, 174-189.

[26] Kumara M, Rath NK, Swain A, Rath SK. Feature selection and classification of microarray data using MapReduce based ANOVA and KNearest neighbor. Procedia Comput Sci 2015; 54: 301-10. https://doi.org/10.1016/j.procs.2015.06.035

[27] Łatkowski, T. and S. Osovski, 2015. Data mining for feature selection in gene expression autism data. Expert Systems Applications, 42: 864-872. https://doi.org/10.1016/j.eswa.2014.08.043
[28] Miyahara, K.; Pazzani, M.J. Collaborative filtering with the simple bayesian classifier. In Proceedings of the Pacific Rim International Conference on Artificial Intelligence, Melbourne, Australia, 28 August–1 September 2000; pp. 679–689. https://doi.org/10.1007/3-540-44533-1_68

[29] Pashaei, E. and N. Aydin, 2017. Binary black hole algorithm for feature selection and classification on biological data. Applied Soft Computing, 56: 94-106. https://doi.org/10.1016/j.asoc.2017.03.002

[30] Ruskin, H.J. Computational Modeling and Analysis of Microarray Data: New Horizons. Microarrays 2016, 5, 26. https://doi.org/10.3390/microarrays5040026

[31] Shen L, Jiang H, He M, Liu G (2017) Collaborative representation-based classification of microarray gene expression data. PLoS ONE 12 (12): e0189533. https://doi.org/10.1371/journal.pone.0189533

[32] Torkkola, K. Feature extraction by non-parametric mutual information maximization. J. Mach. Learn. Res. 2003, 3, 1415–1438.

[33] Tran, B., B. Xue and M. Zhang, 2014. Improved PSO for Feature Selection on High-Dimensional Datasets. In: Simulated Evolution and Learning, Dick, G. (Ed.), Lecture Notes in Computer Science, Springer, Cham. https://doi.org/10.1007/978-3-319-13563-2_43

[34] Wang, A.; Gehan, E. Gene selection for microarray data analysis using principle component analysis. Stat. Med. 2005, 24, 2069–2087. https://doi.org/10.1002/sim.2082

[35] Witten, I.H., and Frank, E. (2005) Data Mining: Practical machine learning tools and techniques. 2nd Ed. Morgan Kaufmann.

[36] Wosiak, A. and D. Zakrzewska, Integrating Correlation-Based Feature Selection and Clustering for Improved Cardiovascular Disease Diagnosis, Complexity, vol. 2018, Article ID 2520706, 11 pages, 2018. https://doi.org/10.1155/2018/2520706.

[37] Zhu, Z., Y.S. Ong and M. Dash, 2007. Markov blanket embedded genetic algorithm for gene selection. Pattern Recognition, 40: 3236-3248. https://doi.org/10.1016/j.patcog.2007.02.007

[38] Yildirim, P., 2015. Filter Based Feature Selection Methods for Prediction of Risks in Hepatitis Disease, International Journal of Machine Learning and Computing, Vol. 5, No. 4. https://doi.org/10.7763/ijmlc.2015.v5.517

[39] Hall, A., Mark, and A. Smith, Lloyd., 1997. Feature Subset Selection: A Correlation based Filter Approach. International Conference on Neural Information Processing and Intelligent Information Systems.

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