Multiple similarly effective solutions exist for biomedical feature selection and classification problems

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Binary classification is a widely employed problem to facilitate the decisions on various biomedical big data questions, such as clinical drug trials between treated participants and controls, and genome-wide association studies (GWASs) between participants with or without a phenotype. A machine learning model is trained for this purpose by optimizing the power of discriminating samples from two groups. However, most of the classification algorithms tend to generate one locally optimal solution according to the input dataset and the mathematical presumptions of the dataset. Here we demonstrated from the aspects of both disease classification and feature selection that multiple different solutions may have similar classification performances. So the existing machine learning algorithms may have ignored a horde of fishes by catching only a good one. Since most of the existing machine learning algorithms generate a solution by optimizing a mathematical goal, it may be essential for understanding the biological mechanisms for the investigated classification question, by considering both the generated solution and the ignored ones.

This study focuses on the binary classification problem in the biomedical big data. A binary classification algorithm tries to tune the parameters of a machine learning model by optimizing the discrimination power of samples from two groups. This problem setting is widely employed to facilitate the clinical drug trials between treated participants and controls, the genome-wide association studies (GWASs) between participants with or without a phenotype, and the biomarker screening procedures between patients of a specific disease and healthy controls of similar baseline characteristics,

Most of the biomedical classification algorithms are deterministic computational algorithms, and the same single solution will be generated for the same input dataset. Modern biotechnologies may produce thousands or millions of data points, or features, for a single sample, and this renders finding the globally optimal solution impossible within a reasonable period of time. So almost all the existing biomedical classification algorithms are approximate algorithms and can only deliver locally optimal solutions. For example, a Support Vector Machine (SVM) builds a hyperplane that maximizes the margin to two groups of samples, and its classification performance heavily depends on the kernel function. Even a special case of an SVM was proven to be NP-hard, so the general SVM model is also NP-hard. A Naïve Bayes (NBayes) classifier is not NP-hard, but has a strong assumption the independence between features. Feature selection may improve the performance of the classification algorithms by reducing the feature dimensions.

A few challenges still remain to be resolved for biomedical classification problems. Firstly, it’s difficult to select the best solution from a few alternative ones with minor or even no performance differences. But most of the existing classification algorithms output only one solution, and discard all the other candidate solutions with similar classification performances, generated during the optimization procedure. Secondly, a classification algorithm usually has some strong assumptions on the dataset, and tries to optimize a mathematical function based on these assumptions. The complexity of biomedical big data does not necessarily follow these assumptions, e.g.

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set ALL2 in Table 2. Even the worst triplet achieved the overall accuracy 0.830 for the dataset ALL2, which was
features with ranks larger than 30 may work very well together with the other features, as in the case for the data-
evaluated by the individual t-test didn’t always appear in the good solutions of all the four datasets, and even the
than one feature selection solution achieved very good performances, as demonstrated in Table 2. The best feature
worse (0.4% in accuracy) than the best model generated by the feature selection algorithm CFS in9. But CFS rec-
larger than or equal to 0.80 for the dataset CNS, and even the worst feature triplet outperformed the previous best
Table 2 showed that there are 1,707 triplets achieved accuracy accuracies larger than the cutoff were listed in the last four rows.

| Dataset  | ALL2 | CNS  | Adeno | DLBCL |
|----------|------|------|-------|-------|
| ELMs in Top20Features | 3    | 1673 | 10000 | 859   |
| ELMs in AllFeatures  | 3    | 313  | 9994  | 174   |
| MinAcc(McTwo)        | 0.651| 0.643| 0.878 | 0.914 |
| MaxAcc(McTwo)        | 0.837| 0.843| 0.919 | 0.987 |
| Top20Features: MinAcc(ELM) | 0.833| 0.944| 1.000 | 1.000 |
| Top20Features: MaxAcc(ELM) | 0.833| 1.000| 1.000 | 1.000 |
| AllFeatures: MinAcc(ELM) | 0.833| 0.889| 1.000 | 0.958 |
| AllFeatures: MaxAcc(ELM) | 0.833| 1.000| 1.000 | 1.000 |

Table 1. Summary of the best ELM models trained on the four datasets. The models with accuracies larger than 0.800 were collected for the two difficult datasets ALL2 and CNS, and the accuracy cutoff 0.900 was used for the two easy datasets Adeno and DLBCL. Except the heading row, the first two rows are the number of ELM models using the training matrices Top20Features and AllFeatures, respectively. The next two rows “MinAcc(McTwo)” and “MaxAcc(McTwo)” gave the minimum and maximum binary classification accuracies of the models generated on the same datasets. The minimum and maximum accuracies of the ELM models with the accuracies larger than the cutoff were listed in the last four rows.

SVM’s kernel function and Naïve Bayes’s inter-feature independence. So even the mathematically global optimum of a classification algorithm may not be a biologically best solution.

So this study proposes the hypothesis that the current machine learning algorithms might have ignored a for-
est of similarly good trees by taking one of these trees as the final solution. We support this hypothesis with multi-
ple similarly good solutions for two binary classification problems, i.e. disease diagnosis and biomarker detection.
The existence of such similarly-well solutions suggests that picking one of them may not be a good choice, and the algorithm users may want to make the choices based on their own domain-specific expertises.

Results and Discussion

Multiple ELMs performed similarly well. We detected more than one ELM models with similarly good classification accuracies for both rounds of experiments, i.e. Top20Features and AllFeatures. Firstly, our ELM models outperformed all the models on the three datasets CNS, Adeno and DLBCL generated in9, and the maximum improvement 15.7% in accuracy was achieved on the dataset CNS. The ELM models performed slightly worse (0.4% in accuracy) than the best model generated by the feature selection algorithm CFS in9. But CFS recom-
ended 56 features, compared with the 20 features by the Top20Features ELM models.

Table 1 also shows the existences of more than one solution with similarly well classification performances. For the two easy datasets Adeno and DLBCL, hundreds or even more of ELM models with very good classification performances exist. For the difficult dataset CNS, even the worst ELM model performs better than the best model reported previously9. And there exist three ELM models with the same classification accuracy 83.3% for both the models of Top20Features and AllFeatures on the dataset ALL2.

Support Vector Machine (SVM) and Naïve Bayes (NBayes) are two widely used classifiers, and were also
evaluated on the T20Features experiment of the two difficult datasets ALL2 and CNS. They are deterministic
classification algorithms, and will generate the same models if the input datasets do not change. When the default
random seed 0 was used, neither SVM and NBayes achieved accuracies better than 0.8. So this study conducted
another randomized experiments for SVM and NBayes. The dataset was randomly split into 70% training and 30%
testing datasets, respectively. An accuracy was calculated for the testing dataset using the model trained over
the training dataset. 10,000 random runs were carried out, but no SVM or NBayes models achieved accuracies
better than 0.8 on the difficult dataset ALL2. For the other difficult dataset CNS, there were 471 SVM models with
accuracies better than 0.8, and NBayes established 1,874 models with accuracies better than 0.8. So in summary,
ELM outperformed SVM and NBayes on the two difficult datasets, and may be a good candidate algorithm to
investigate the hypothesis in this study.

Some of these accurate ELM models are significantly different to each other. An ELM model was represented by
the matrix of the internal parameters, as described above. A heatmap was generated for the difference matrix
between two ELM models, as shown in Fig. 1. A white color represents no difference between the corresponding
weights in the two ELM models, and red represents the maximum difference. For the two best ELM models on the
dataset ALL2, the maximum difference reaches 4.8 for the mode Top20Features, and a larger difference 6.4 for the
model AllFeatures. A significant difference was observed in both heatmaps on the dataset ALL2. Similar patterns
were observed for the other three datasets, as illustrated in Fig. 1.

There exist multiple feature selection solutions with best classification accuracies. Firstly, more
than one feature selection solution achieved very good performances, as demonstrated in Table 2. The best feature
evaluated by the individual t-test didn’t always appear in the good solutions of all the four datasets, and even the
features with ranks larger than 30 may work very well together with the other features, as in the case for the data-
set ALL2 in Table 2. Even the worst triplet achieved the overall accuracy 0.830 for the dataset ALL2, which was
similar to the best performance 0.837 in the previous study. And the best feature triplet outperformed the best
previous model by 0.030 in the overall accuracy. Table 2 showed that there are 1,707 triplets achieved accuracy
larger than or equal to 0.80 for the dataset CNS, and even the worst feature triplet outperformed the previous best
model by 0.101 in the overall accuracy. There are 19,040 and 6,803 feature triplets achieved 1.000 in accuracy for the two easy datasets Adeno and DLBCL, respectively.

The averaged ranks of the selected triplets demonstrated the existences of many features that were not high in ranks but constituted very good feature triplets. The minimum feature ranks of these triplets were larger than 1.000 for all the four datasets. So there exist triplets that didn’t consist of the feature with the smallest t-test P-value for all the four datasets.

Multiple good solutions were detected for all the four datasets, and they scattered across the 3-D space, since $x \leq y \leq z$. The scattering pattern was also observed for the best ten models, as shown in Fig. 2.

Figure 2 shows the scattering map of the selected feature triplets in the 3D plots in Fig. 2. The x, y, z coordinates of a triplet were the ranks of the three features in the top-50 p-value features of t-test. A red color in the heatmap indicates the accuracy of 1.00 and the blue indicates the accuracy of 0.72. For each dataset, the best 10 solutions and the solutions which reach the cutoff value are both shown in the 3D scattering map. For dataset CNS, the best 10 solutions have similar red colors, so they achieved similar classification accuracies. However, they scattered across the 3D space, and clearly demonstrated their differences on the feature compositions. Even more variations were observed for all the solutions with accuracy $\geq 0.8$. Similar patterns were also detected for all the other three datasets.

We further investigated whether a good triplet may consist of three features with low classification performances alone. One of the interesting examples is the triplet $\{D50683\_at, HG961-HT961\_at, Z35402\_rna1\_s\_at\}$ for the dataset DLBCL. This triplet achieved 95.8% in accuracy, but their individual performances were not very high. The feature $D50683\_at$ only achieved 66.7% in accuracy, while the other two features $HG961-HT961\_at$ and $Z35402\_rna1\_s\_at$ alone performed slightly better, with accuracies 83.3% and 75.0%, respectively. So it's

Table 2. Summary of the best ELM models trained on triplets of the four datasets. The models with accuracies larger than 0.800 were collected for the two difficult datasets ALL2 and CNS, and the accuracy cutoff 0.900 was used for the two easy datasets Adeno and DLBCL. Except the heading row, the first row is the number of ELM models using the training matrices with 3 features. The next two rows “MinAcc(McTwo)” and “MaxAcc(McTwo)” gave the minimum and maximum binary classification accuracies of the models generated on the same datasets. The minimum and maximum accuracies of the best ten ELM models with the accuracies larger than the cutoff were listed in the following two rows. And the last row shows the averaged rankings of the 10 triplets with the best accuracies.

| Dataset | ALL2 | CNS | Adeno | DLBCL |
|---------|------|-----|-------|-------|
| Triplets | 21 | 1707 | 19040 | 6803 |
| MinAcc(McTwo) | 0.651 | 0.643 | 0.878 | 0.914 |
| MaxAcc(McTwo) | 0.837 | 0.843 | 0.919 | 0.987 |
| MinAcc(ELM) | 0.830 | 0.944 | 1.000 | 1.000 |
| MaxAcc(ELM) | 0.867 | 1.000 | 1.000 | 1.000 |
| (Min, Max) ranks | 7,000, 32,000 | 4,000, 36,333 | 1,333, 29,000 | 12,667, 32,333 |
necessary to use more features for much better classification performances, and these three features may contribute complementary information for the classification modeling.

Validations of the above two observations. The above two observations were further validated on six independent binary classification datasets. The six datasets were described in the section Material and Methods, and the same experimental procedure was carried out on these datasets. The similar significant difference was detected between the best two ELM classification models of each of the six dataset, as shown in Fig. 3 and summarized in Table 3. All the six datasets have at least two ELM models with identical classification performances using the top 20 features ranked by t-test, and these two models were significantly different to each other, as demonstrated by the heatmaps in Fig. 3. Only two of the six datasets Gas1 and Gas2 have the best two ELM models with the same classification accuracies, and they are significantly different to each other, too. The classification accuracies of the best two ELM models of the other cases using all the features in Fig. 3 demonstrated slight differences, which may be due to that only 10,000 random runs of ELM optimizations were conducted in this study. The data also suggested that there exist many noisy features in the biomedical datasets and it's necessary to remove some before the training of a classification model. In either situation, there holds the existence of multiple similarly-well solutions for the investigated biomedical classification problems.

All the triplet feature subsets with Acc ≥ 0.8 for the six independent datasets were summarized in Fig. 4 and Table 4. We may observe that there are a large number of triplets with Acc ≥ 0.8 for each of the six datasets. The averaged rankings of the three features in the best 10 triplets were summarized in the last row of Table 4, and suggested that these features are different to each other. So the best 10 triplets used different features, but they achieved the same classification accuracies, as illustrated in Fig. 4. So there exist multiple feature selection solutions with similarly-well performances for the six independent datasets.

Like the 4 datasets argued above, the six more datasets investigated in this section have the same situation that features may be complementary to each other and their combination achieved much better. For example, the combination of the three features 208600_s_at, 202469_s_at and 212481_s_at achieved 93.2% in accuracy while they alone can only achieve 79.7% (208600_s_at), 70.4% (202469_s_at) and 86.3% (212481_s_at) in accuracies, respectively.

There exist multiple feature selection solutions with the best classification precisions or recalls. Besides the performance measurement accuracy, precision and recall were also used to evaluate the trained models. Precision was defined as TP/(TP + FP) and recall was defined as TP/(TP + FN), where TP, FP and FN were the numbers of true positives, false positives and false negatives, as defined in Eq. Similar patterns were observed using the two performance measurements precision and recall, as summarized in Table 5 and demonstrated in Figs 5 and 6.

All the best ten models achieved the same precision 1.000 for all the four datasets although the difficult dataset CNS has much fewer models than the other three datasets, as in Table 5 and Fig. 5. The topological distributions of the best 10 models do not form a tight cluster for all the four datasets, suggesting that these 10 triplets didn't use similar features. The data in Table 5 also supported this observation that the averaged rankings of three features in the best 10 triplets are different to each other. For example, the dataset Adeno has a triplet with the averaged ranking 1.333, while another of its best 10 triplets has the averaged ranking 24.000.
Significantly different features were used for the 10 triplets with the best recalls for the four datasets. But the numbers of good solutions varied for different performance measurements, suggesting that the solution optimization procedure heavily depends on the optimization goal.

Biological hints on multiple solutions with the same best classification performances. As shown in the above experiments, there exist multiple solutions with best classification performances, which were demonstrated in ten different datasets. We explored the biological functions of different triplets with the similarly best classification accuracies on the datasets ALL2, CNS, Adeno and DLBCL.

The dataset ALL2 has two groups of features \{1803\_at, 32783\_at, 39271\_at\} and \{36912\_at, 1599\_at, 1854\_at\}, and they achieved 0.867 and 0.833 in accuracies, respectively. It was demonstrated that X-ray irradiation may induce G1-arrest in the MOLT-4 lymphocytic leukemia cells, and this process was usually accompanied by...
reduced activity of CDK2 but increased activity of CDK1 (1803_at)
19. Another literature suggested that MYBL2 (1854_at) was over-expressed in some acute myeloid leukemias
20. So it's interesting to observe that these two leukemia-associated biomarker genes accompanied by two other features may accurately separate the ALL patients with relapse from those without.

The two triplets {L17131_rna1_at, M73547_at, D78012_at} and {U50136_rna1_at, J02611_at, D17793_at} generated 0.889 and 0.833 in accuracies for the binary classification problem CNS. The POLYPOSIS LOCUS PROTEIN 1 (M73547_at) has a known association with CNS and patients with familial polyposis have the increased risk of tumor development in extracolonic sites, including the central nervous system21. It was also proposed that the expression level of Apolipoprotein D (J02611_at) is correlated with the prognosis in several types of malignancy, including CNS astrocytomas and medulloblastomas22.

The dataset Adeno has two triplets {M77836, J02854, T64297} and {H06524, H43887, U37019} among the ten best solutions. These two triplets have not shared features, but both achieved the best classification accuracies.

**Figure 4.** Three-D dot plots for the triplet feature subsets of the six independent datasets using accuracy as the measurement. The axes x, y and z are the t-test ranks of three features in each solution, and $x \leq y \leq z$. The color of each point reflects the accuracy of the model it represents on the corresponding dataset.

| Dataset  | Leuk | Lym | Stroke | ALL1 | Gastric1 | Gastric2 |
|----------|------|-----|--------|------|----------|----------|
| Triplets | 14703| 13416| 8090   | 19600| 19600    | 19600    |
| MinAcc(ELM) | 1.000| 1.000| 1.000  | 1.000| 0.980    | 1.000    |
| MaxAcc(ELM) | 1.000| 1.000| 1.000  | 1.000| 0.980    | 1.000    |
| (Min, Max) | 3.667| 4.333| 10.667 | 11.000| 23.333   | 10.333   |
| ranks     | 17.667| 11.667| 12.667 | 6.000 | 23.667   | 19.667   |

**Table 4.** Summary of the best ELM models trained on the triplets of the six independent datasets. The meaning of each row is similar to Table 2.
1,000. M77836, also known as pyrroline-5-carboxylate reductase 1 (PYCRI) was experimentally observed as an upregulated protein in an in vitro progression model of the colorectal adenoma–to–carcinoma sequence using quantitative 2-DE and is complemented by Western blot validation. Another immunohistochemistry study confirmed that H06524 (Gelsolin precursor) is severely down-regulated in all adenocarcinomas tested while is expressed in normal cells of the colon. So both triplets have genes biologically associated with adenocarcinoma. The existence of these two triplets with the best accuracy 1.000 may represent two distinctive functional modules expressed in normal cells of the colon.

The easy dataset DLBCL has two groups of features \{X62078_at, L33842_rna1_at, J02645_at\} and \{X56494_at, M57710_at, U19495_s_at\}, both of which achieved 0.917 in accuracy. The gene Lectin, galactoside-binding, soluble, 3 (LGALS3, galectin 3, M57710_at) is a significantly upregulated gene in both the FL and DLBCL lymphoma samples when compared with the expression of the immune escape genes at the single gene level in control ble, 3 (LGALS3, galectin 3, M57710_at) is over-expressed in the REF/REL group of DLBCL patients and was suggested to be associated with the resistance to immunochemotherapy of the DLBCL patients.

**Conclusion**

Our experimental data demonstrated that more than one best solution exist for the disease classification and feature selection problems. Some essential information may be lost if we only focus on one solution for these two optimization problems. So we recommend that at least multiple solutions with very close optimization performances should be delivered by optimization algorithms like classification and feature selection.

**Material and Methods**

The python source code for testing ELM, SVM and NBayes on the two difficult datasets ALL2 and CNS may be freely accessed at http://www.healthinformaticslab.org/supp/.

**Proposed Methodology of Disease Classification.** Disease diagnosis is a binary classification problem that we determine whether a given sample has the investigated disease or not based on its data. There are two groups of samples in such a problem. There is usually one group of samples carrying a specific disease, and the other group of samples consists of the control samples. A highly accurate classification model may greatly help the clinical doctors improve the diagnosis sensitivity and reduce the false positives.

ELM (Extreme Learning Machine) is a machine learning algorithm that implements a feed-forward neural network with a single layer of hidden nodes for classification or regression. ELM features generating a very good classification model even with the utilization of a vector of randomly generated input weights. It's also very efficient on dealing with a huge amount of features.

So we hypothesize that there may be multiple ELM classification models with similar prediction accuracies. The low computational requirement of ELM makes it possible to run ELM multiple times within a reasonable period of time.

**Experimental settings of Disease Classification.** We chose four publicly available datasets, i.e., ALL2, CNS, Adeno and DLBCL. These four datasets are representative since the datasets ALL2 and CNS are difficult in classification while the other two datasets Adeno and DLBCL are easy to be separated, as shown in a recent work. All these datasets are binary classification problems.

Two experimental strategies were employed to find multiple optimized ELM models. Firstly, features in each dataset were evaluated for their discriminative abilities using t-test, and only 20 features with the smallest P-values

| Dataset     | ALL2 | CNS | Adeno | DLBCL |
|-------------|------|-----|-------|-------|
| Precision(Triplets) | 16535 | 732 | 17169 | 14625 |
| MinPrecision(ELM)    | 1.000 | 1.000 | 1.000 | 1.000 |
| MaxPrecision(ELM)    | 1.000 | 1.000 | 1.000 | 1.000 |
| (Min, Max) ranks of best-precision 10 models | 18.667 | 10.667 | 1.333 | 11.667 |
| Recall(Triplets)     | 4906  | 1369 | 17613 | 198   |
| MinRecall(ELM)       | 1.000 | 1.000 | 1.000 | 1.000 |
| MaxRecall(ELM)       | 1.000 | 1.000 | 1.000 | 1.000 |
| (Min, Max) ranks of best-Recall 10 models | 19.333 | 2.000 | 25.000 | 19.667 |

Table 5. Summary of the ELM models with the best performance measurements precision and recall trained on triplets of the four datasets. The numbers of models with precisions and recalls larger than CutOff were collected for the four datasets ALL2/CNS/Adeno/DLBCL in the row “Precision(Triplets)” and “Recall(Triplets)”, respectively. CutOff is 0.800 for the two difficult datasets ALL2 and CNS, and 0.900 for the two easy datasets Adeno and DLBCL. The minimum and maximum precisions of the best ten ELM models with the precisions larger than the cutoff were listed in the rows “MinPrecision(ELM)” and “MaxPrecision(ELM)”. And the next row gave the averaged rankings of the 10 triplets with the best precisions. The last three rows were defined similarly for the performance measurements Recall.

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were kept for ELM constructions. This round of experiments was denoted as Top20Features. Secondly, all the features in a dataset were used to train the ELM models, and this round is denoted as AllFeatures.

10,000 optimizations of ELM models using different randomly generated input weights were carried out for each round of experiments. Each dataset was split into two parts, i.e. 70% as the training and 30% as the test datasets. The binary classification accuracy on the test dataset was calculated for comparison, as defined in 9.

The ELM module Python-ELM version 0.3 of the Python programming language was used in this study 27. This module implements an MLP (Mult-Layer Perceptron) for the random input layer. For a dataset with m features, an \( n \times m \) weight matrix will be generated based on the training data, where \( n \) is the number of units in the layer,

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**Figure 5.** Three-D dot plots for the four datasets using precision as the measurement. The axises x, y and z are the ranks of each solution, and \( x \leq y \leq z \). The color of each point reflects the precision of the model on the corresponding datasets.

**Figure 6.** Three-D dot plots for the four datasets using recall as the measurement. The axises x, y and z are the ranks of each solution, and \( x \leq y \leq z \). The color of each point reflects the recall of the model on the corresponding datasets.
and n are 150 and 290 for the modes Top20Features and AllFeatures, respectively. This matrix was extracted to represent the trained ELM model, and a comparison was conducted between the weight matrices of two ELM models. A difference matrix of two weight matrices is defined to be the absolute value of the first matrix minus the second one. All the experiments in this study were carried out in an Inspur Gene Server G100, with 256GB memory, 28 Intel Xeon® CPU cores (2.4 GHz), and 30TB RISC1 disk space.

Proposed Methodology of Feature Selection. Besides the classification problem, we also investigated the existence of multiple similarly good feature subsets for the four datasets, i.e. ALL2, CNS, Adeno and DLBCL. Almost all the existing feature selection algorithms are deterministic, and they generate only one solution. So we chose the top 50 ranked features based on the t-test P-values for the four datasets, and conducted an exhaustive screening of all the triplets from these 50 features. Each dataset is randomly split into two parts: 80% as training dataset and 20% as test dataset. The accuracy of test dataset is used to represent the binary classification accuracy. For each of the four datasets, we screened 5000 triplets, and 100 triplets with the highest accuracy were saved. The accuracy cutoff 0.8 was chosen for the two difficult datasets ALL2 and CNS, and 0.9 for the two easy datasets Adeno and DLBCL. Besides accuracy, two other performance measurements precision and recall were used to test our hypothesis. The same cutoff values were used to screen the solutions, so that the solutions with top performances were highlighted.

An extended validation on six more datasets. An extended validation on six more binary-classification transcriptomic datasets were employed to further support our hypothesis. The widely used Leukaemia (Leuk) dataset has 47 acute lymphoblastic leukaemia (ALL) and 25 acute myeloid leukaemia (AML) samples. Another dataset Lymphoma (Lym) has 22 germlinecentre and 23 activated B-like DLBCL samples. The 40 samples of the dataset Stroke (accession: GSE22253) consists of 20 ischemic stroke patients and 20 controls. One more dataset ALL1 was chosen from the same database of ALL2, and it has 95 B-cell and 33 T-cell acute lymphoblastic leukemia (ALL) samples. The two gastric cancer datasets Gas1 and Gas2 were retrieved from the database Gene Expression Omnibus Gastric1/Gastric2 (accession: GSE29272). Gas1 consists of 72 non-cardia gastric cancer samples and 72 normal controls. Gas2 has 62 cardia gastric cancer samples and 62 normal controls.

References
1. Coppo, R. et al. Risk factors for progression in children and young adults with IgA nephropathy: an analysis of 261 cases from the VALIGA European cohort. Pediatr Nephrol 32, 139–150, https://doi.org/10.1007/s00467-016-3469-3 (2017).
2. Utkin, L. V., Chekh, A. I. & Zhuk, Y. A. Binary classification SVM-based algorithms with interval-valued training data using triangular and Epanechnikov kernels. Neural Netw 80, 53–66, https://doi.org/10.1016/j.neunet.2016.04.005 (2016).
3. Lopresti, A. L. & Drummond, P. F. Efficacy of curcumin, and a saffron/curcumin combination for the treatment of major depression: A randomised, double-blind, placebo-controlled study. J Affect Disord 207, 188–196, https://doi.org/10.1016/j.jad.2016.09.047 (2017).
4. Pantalio, G. F. et al. Genome wide association study (GWAS) for grain yield in rice cultivated under water deficit. Genetica 144, 651–664, https://doi.org/10.1007/s10709-016-9992-z (2016).
5. Zhang, W. et al. Genetic variants in the PIWI-piRNA pathway gene DCP1A predict melanoma disease-specific survival. Int J Cancer 139, 2730–2737, https://doi.org/10.1002/ijc.30409 (2016).
6. Urquidi, V. et al. A microRNA biomarker panel for the non-invasive detection of bladder cancer. Oncotarget, https://doi.org/10.1086/624782 (2016).
7. Vidmar, T. & Kanisch, G. A recursive deterministic algorithm for treatment of true coincidence summing effects in gamma-ray spectrometry. Appl Radiat Isot 70, 726–732, https://doi.org/10.1016/j.apradiso.2011.12.021 (2012).
8. Brown, W. M., Faulon, J. L. & Sale, K. A deterministic algorithm for constrained enumeration of transmembrane protein folds. Comput Biol Chem 29, 143–150, https://doi.org/10.1016/j.compbiolchem.2005.03.001 (2005).
9. Ge, R. et al. McTwo: a two-step feature selection algorithm based on maximal information coefficient. Bioinformatics 30, 142, https://doi.org/10.1093/bioinformatics/btw620 (2014).
10. Chiang, D. Y. et al. High-resolution mapping of copy-number alterations with massively parallel sequencing. Nat Methods 6, 99–103, https://doi.org/10.1038/nmeth.1276 (2009).
11. Long, T., Chen, M., Jiang, S. & Lu, W. Continuous leaf optimization for IMRT leaf sequencing. Med Phys 43, 5403, https://doi.org/10.1118/1.4962030 (2016).
12. Peng, R., Leung, C. S., Constantinides, A. G. & Zeng, W. I. Lagrange Programming Neural Network for Nondifferentiable Optimization Problems in Sparse Approximation. IEEE Trans Neural Netw Learn Syst, https://doi.org/10.1109/TNNLS.2016.2575860 (2016).
13. Amari, S.-i. & Wu, S. Improving support vector machine classifiers by modifying kernel functions. Neural Networks 12, 783–789 (1999).
14. Scholkopf, B. & Smola, A. J. Learning with kernels: support vector machines, regularization, optimization, and beyond. (MIT press, 2001).
15. Brooks, J. P. Support vector machines with the ramp loss and the hard margin loss. Operations research 59, 467–479 (2011).
16. Lewis, D. D. Naive (Bayes) at forty: The independence assumption in information retrieval. Neural Networks 17, 346–354, https://doi.org/10.1016/S0893-6080(00)00084-0 (2001).
17. Zou, Q., Zeng, J., Cao, L. & Ji, R. A novel features ranking metric with application to scalable visual and bioinformatics data classification. Neurocomputing 173, 346–354, https://doi.org/10.1016/j.neucom.2014.12.123 (2016).
18. Lin, C. et al. LibD3C: Ensemble classifiers with a clustering and dynamic selection strategy. Neurocomputing 123, 424–435, https://doi.org/10.1016/j.neucom.2013.08.004 (2014).
19. Wu, J. et al. Unscheduled CKD1 activity in G1 phase of the cell cycle triggers apoptosis in X-irradiated lymphocytic leukemia cells. Cellular & Molecular Life Sciences 63, 2538–2545 (2006).
20. Doli, S. et al. Study of the S427G polymorphism and of MYBL2 variants in patients with acute myeloid leukemia. Leukemia & Lymphoma 57, 1 (2015).
21. Williamson, P. M. & Smith, K. V. Association of central nervous system sarcoma with familial polyposis coli. Clinical & Experimental Neurology 15, 127–130 (1978).
22. Hunter, S. B. et al. Apolipoprotein D expression in primary brain tumors: analysis by quantitative RT-PCR in formalin-fixed, paraffin-embedded tissue. Journal of Histochemistry & Cytochemistry Official Journal of the Histochemistry Society 53, 963–969 (2005).
23. Roth, U. et al. Differential expression proteomics of human colorectal cancer based on a syngeneic cellular model for the progression of adenoma to carcinoma. Proteomics 10, 194 (2010).
24. Gay, F. et al. In colon carcinogenesis, the cytoskeletal protein gelsolin is down-regulated during the transition from adenoma to carcinoma. *Human Pathology* **39**, 1420–1430 (2008).

25. Tosolini, M., Algans, C., Pont, F., Ycart, B. & Fournié, J. J. Large-scale microarray profiling reveals four stages of immune escape in non-Hodgkin lymphomas. *Oncoimmunology* **5**, e118246 (2016).

26. Martin Stenson, S. B.-E. et al. Overexpression of Ribosomal Proteins Is Associated with Resistance to Immunochemotherapy in Diffuse Large B-Cell Lymphoma Patients. *Blood* **120**, 4121 (2016).

27. Ding, S., Zhao, H., Zhang, Y., Xu, X. & Nie, R. Extreme learning machine: algorithm, theory and applications. *Artificial Intelligence Review* **44**, 103–115 (2015).

28. Golub, T. R. et al. Molecular classification of cancer: class discovery and class prediction by gene expression monitoring. *Science* **286**, 531–537 (1999).

29. Alizadeh, A. A. et al. Distinct types of diffuse large B-cell lymphoma identified by gene expression profiling. *Nature* **403**, 503–511, [https://doi.org/10.1038/35000501](https://doi.org/10.1038/35000501) (2000).

30. Krug, T. et al. TTC7B emerges as a novel risk factor for ischemic stroke through the convergence of several genome-wide approaches. *J Cerebr Blood F Met* **32**, 1061–1072 (2012).

31. Chiaretti, S. et al. Gene expression profile of adult T-cell acute lymphocytic leukemia identifies distinct subsets of patients with different response to therapy and survival. *Blood* **103**, 2771–2778, [https://doi.org/10.1182/blood-2003-09-3243](https://doi.org/10.1182/blood-2003-09-3243) (2004).

32. Wang, G. S. et al. Comparison of Global Gene Expression of Gastric Cardia and Noncardia Cancers from a High-Risk Population in China. *Plos One* **8** (2013).

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Author Contributions

J.L. and C.X. worked on designing the experiment, writing the code and running most of the experiments. J.L. and C.X. drafted the initial version of the manuscript. W.Y. and Y.S. were involved in carrying out the experiments and result discussion. F.Z. conceived the project, designed the experiment and wrote the manuscript. W.Z. helped discuss, draft and polish the manuscript.

Additional Information

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