Prediction of Lung Cancer Based on Serum Biomarkers by Gene Expression Programming Methods

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Abstract

In diagnosis of lung cancer, rapid distinction between small cell lung cancer (SCLC) and non-small cell lung cancer (NSCLC) tumors is very important. Serum markers, including lactate dehydrogenase (LDH), C-reactive protein (CRP), carcino-embryonic antigen (CEA), neurone specific enolase (NSE) and Cyfra21-1, are reported to reflect lung cancer characteristics. In this study classification of lung tumors was made based on biomarkers (measured in 120 NSCLC and 60 SCLC patients) by setting up optimal biomarker joint models with a powerful computerized tool - gene expression programming (GEP). GEP is a learning algorithm that combines the advantages of genetic programming (GP) and genetic algorithms (GA). It specifically focuses on relationships between variables in sets of data and then builds models to explain these relationships, and has been successfully used in formula finding and function mining. As a basis for defining a GEP environment for SCLC and NSCLC prediction, three explicit predictive models were constructed. CEA and NSE are frequently-used lung cancer markers in clinical trials, CRP, LDH and Cyfra21-1 have significant meaning in lung cancer, based on CEA and NSE we set up three GEP models-GEP1 (CEA, NSE, Cyfra21-1), GEP2 (CEA, NSE, LDH), GEP3 (CEA, NSE, CRP). The best classification result of GEP gained when CEA, NSE and Cyfra21-1 were combined: 128 of 135 subjects in the training set and 40 of 45 subjects in the test set were classified correctly, the accuracy rate is 94.8% in training set; on collection of samples for testing, the accuracy rate is 88.9%. With GEP2, the accuracy was significantly decreased by 1.5% and 6.6% in training set and test set, in GEP3 was 0.82% and 4.45% respectively. Serum Cyfra21-1 is a useful and sensitive serum biomarker in discriminating between NSCLC and SCLC. GEP modeling is a promising and excellent tool in diagnosis of lung cancer.

Keywords: Lung cancer - diagnosis - gene expression programming - biomarker - cyfra21-1

Introduction

Lung cancer is one of the most frequent types of cancer in the world, both in terms of incidence and mortality, leading to three million deaths annually and resulting in an enormous global health problem (Cho, 2007). Known as diseases caused by uncontrolled cell growth in one or two lungs, its primary symptoms include chest pain, shortness of breath, cough and coughing up blood. Lung cancer may spreads to other organs and spreading to the brain can cause headache, vomiting, psychosis, unilateral limb paresthesia and visual impairment. In spite of the progress and efforts we made in treatment and diagnosis of lung cancer patients, the diagnosis of overall survival at five years is only 15%, and more than 75% patients at the time of diagnosis is already presenting with advanced stage of disease when therapeutic options are very limited (Patz et al., 2007).

The histological types of lung cancer are complicated because there are many types of normal airway epithelial cells. In the process of tumor genesis, pluripotent stem cells can differentiate into different directions, so that there is a significant heterogeneity of lung cancer in the histological types. Based on histopathological presentation, lung cancer is sub-divided into four major histological subtypes: small cell lung cancer (SCLC), squamous cell carcinoma (SCC), adenocarcinoma (ADC), and large cell carcinoma (LCC). The latter three, collectively referred as non-small cell lung cancer (NSCLC). So, from clinical point of view, domestic and overseas, these four completely different types of lung cancer are divided into two categories: small cell lung cancer (SCLC) (16.8%) and non-small cell lung cancer (NSCLC) (80.4%) (Travis et al., 1995). Since the biological behavior of small cell lung cancer and other types of lung cancer are significantly different-namely, the former is clinically characterized by...
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highly malignant, early widespread metastases, sensitivity to radiotherapy and chemotherapy, so the treatment of small cell lung cancer is different to other types of lung cancer. As different types of lung cancer have different treatments, the differential diagnosis in early stage is very important to improve the survival rate. Therefore, early diagnosis of lung cancer types may play a very important role in the improvement of therapeutic outcomes. Together with the advances in imaging studies (such as Computed Tomography, Whole-body positron-emission tomography, Chest Radiograph, Magnetic Resonance Imaging and Sputum Cytology) and endoscopic examinations (such as bronchoscope and mediastinoscope), the development of biomarkers useful for serum diagnosis is crucial for the early diagnosis of lung cancer.

Biomarkers are reflecting the chemical and biological substances presented in the tumor. They may not exist in normal adult tissue or found only in embryonic tissue. Their content in the tumor tissues could significantly exceed the content in normal organization. Their quantitative or qualitative change may indicate the nature of the tumor, so as to understand the tumor tissue, cell differentiation and cell function, to help the diagnosis, classification, prognosis judgment and treatment of tumor (Leidinger et al., 2010).

Detection of serum tumor marker levels becomes ways to improve the rate of early diagnosis of lung cancer (Zhang et al., 2013). Markers which have been widely used for diagnosing lung cancer include CEA (Yang et al., 2014), Cyfra21-1 (Tomita et al., 2010), NES (Chu et al., 2011), LDH (Ziaian et al., 2014), CRP (Onitilo et al., 2012) and other indexes. However, single marker of lung cancer shows low diagnostic specificity and sensitivity, having poor value for the diagnosis of lung cancer (Leidinger et al., 2010). Therefore, there is a consensus that a combination of markers can increase the diagnostic specificity and sensitivity than single marker (Schneider et al., 2002; Farlow et al., 2010; Wang et al., 2013; Li et al., 2014). For example, in diagnosing some malignant tumors, such as prostate or ovarian cancer, by combining multiple biomarkers data, the diagnostic performance was improved rapidly than individual serum biomarker (Armsellem-Ouazana et al., 2005; Kim et al., 2012). In recent years, people have done a lot of researches on statistical point of view, hoping to establish a kind of intelligent diagnosis model based on multiple tumor marker detection data, in order to overcome the influence of some subjective and individual factors, and improve the further tumor diagnosis.

Over the last decades, machine learning algorithms have shown to be a potential tool in medicine. The Naives Bayes algorithm and the support vector machine (SVM) (Gopinath and Shanthi, 2013), Fuzzy Logic and Artificial Neural Networks (ANN) (Biglarian et al., 2012) have been used as auxiliary tools in diagnosis and prognosis of cancer. These methods train as a classifier according to the features of essential biomarkers and other index. Then test the classifier in the same samples or the other samples. For example, Fuzzy logic-based tumor-marker profiles improved sensitivity in the diagnosis of lung cancer (Schneider et al., 2002), and ANN significantly improved the sensitivity of biomarkers in the Diagnosis of Lung Cancer (Feng et al., 2012).

As these machine learning methods are easily available as software packages and can achieve a highly accurate model of classification, they can be broadly adapted to predict lung cancer types by using input biomarker data. While, what is difficult for them is to explain how these data were used for classification. Moreover, little methods are able to select appropriate data automatically. So we can say the predictive results which without revealing the intrinsic link of each variable are “black box”. Researchers tend to analyze the relationship between biomarkers and the lung cancer with statistical methods. So we proposed a new evolutionary algorithm -Gene expression programming (GEP).

Gene expression programming (GEP) is proposed by a Portuguese scientist called Candida Ferreira in 2000 and it is a new type of adaptive evolutionary algorithm based on biological structure and function. What it learns specifically is about the relationships between variables in different sets of data, and it builds models to describe these relationships. GEP does not require an accurate formulation of physical relationships, and can find the precise expression for different markers. The markers that show negative correlation with lung cancer will not be selected by the GEP classifier.

GEP is originated in the field of biology and developed from the genetic algorithm (GAs) and genetic programming (GP). It inherits the traditional advantages of GAs and GP, based on these advantages, a genetic operation was developed which is specific to GEP. A large number of experiments prove that the GEP algorithm and a variety of improved algorithms have a very good performance in using simple coding to solve complex problems.

On the other hand, GEP, have played a significant role to predict essential proteins indispensable for cell survive (Zong et al., 2013). GEP has been used to predict adverse events of radical hysterectomy in cervical cancer patients (Kusy et al., 2013). Moreover, a GEP was applied to automatically detect the population with fatty liver in computed the teat date of serum glucose, the total cholesterol, low density lipoprotein cholesterol, high density lipoprotein cholesterol and triglyceride from 196 power plant workers. However, there is no relevant literature researched that GEP is effective in auxiliary diagnosis of lung cancer.

In this study we record the level of an assortment of biomarkers which previously proved to have prognostic or diagnostic value of lung cancer, as a first step in the effort to improve the diagnostic accuracy of biomarkers and establish a novel multi-analyze serum biomarker test for prediction of lung cancer through the use of Gene Expression Programming. In this way can we develop a best artificial calculation model that can be widely used in lung cancer types predicting.

Materials and Methods

Subjects

We selected 180 patients who were diagnosed with
biopsy-proven untreated pulmonary malignant disease. They all hospitalized in the Affiliated Hospital of Qingdao University from January 2006 to September 2013. The histological subtypes in the 180 patients with lung cancer were NSCLC in 120, SCLC in 60 patients. Approval has been obtained from the relevant ethics committee and all participants were given the written informed consents. Histological diagnosis of primary lung cancer was established according to the revised classification of lung tumors of the World Health Organization and the International Association for Lung Cancer Study. A summary of the clinical characteristics of the subjects, together with a breakdown of each group by age, gender and more detailed information is presented in Table 1.

Biomarker selection

Our GEP classifier is constructed to predict lung cancer types based on various biomarkers. Because CEA and NSE are the most widely used biomarkers among these patients, we implement experiments based on data of both of them. C-reactive protein (CRP) is an acute-phase protein produced mainly by hepatocytes in the presence of inflammation and serum CRP concentrations were significantly higher in NSCLC patients compared to the SCLC controls (Lee et al., 2009). Serum LDH levels have been correlated with poor prognosis and resistance to chemotherapy and radiotherapy in various neoplastic diseases (Zhao et al., 2013). Serum Cyfra21-1 is one of the most important serum markers in the diagnosis of non-small cell lung cancer (NSCLC), especially squamous-cell carcinoma (Ono et al., 2013; Wang et al., 2013). So, based on CEA and NSE, we set up three GEP models - GEP1 (CEA, NSE, Cyfra21-1), GEP2 (CEA, NSE, LDH), GEP3 (CEA, NSE, CRP).

Measurement of serum biomarker concentrations

Specimens: 10 ml blood was collected from each patient in the fasting state, and processed immediately by centrifugation at 3000 rpm at room temperature for 10 min in a centrifuge.

Kits and Instruments: Levels of CEA, NSE and Cyfra21-1 were measured by using the Electrochemiluminescence immunoassay instrument. Their kits are from Roche Germany. Test them according to the standard operating procedure for kits. LDH and CRP are measured by using Olympus AU2700 automatic biochemical analyzer and auxiliary reagents. All operations are enforced according to the instruction strictly.

Normal Serum Reference Values: LDH: <245 U/L; CEA: <5.0 ng/mL; Cyfra21-1: <3.3 ng/mL; NSE: 15.7-17.1 ng/mL; CRP: <8 ng/mL.

Statistical analysis of individual serum biomarkers

Statistical analyses were performed using SPSS19.0. All data were expressed as mean±standard, groups comparison was conducted using analysis of variance (ANOVA). P-values<0.05 were considered significant difference, P>0.05 showed no significant difference among two groups.

Gene expression programming

Data preparation of GEP: Gene encodes the two types of lung tumors randomly. We code small cell lung cancer as 0, non-small cell lung cancer as 1. All data was entered into the computer twice. Original data are divided into 4 equal datasets, and one fold is used to train the classifier and the remaining three folds are used for testing the predictive performance of GEP. Since the ratio of SCLC and NSCLC in original data is about 1.3 (SCLC: NSCLC=60:180), each fold data maintains the same ratio of SCLC and NSCLC in original data. So there are 45 patients (SCLC: n=15 NSCLC: n=30) in testing set and 135 patients (SCLC: n=45 NSCLC: n=90) in training set. As we established three GEP models, the process is repeated three times to generated three classifiers, with each of the three datasets used exactly once as testing data.

GEP theory: To build a classifier of predicting lung cancer types using GEP, the following major steps are needed: defining a chromosome using a function and terminal set, initializing a population and generating a group of chromosomes, defining a fitness function for evaluating chromosomes, selecting eugenic ones from populations, reproducing a group of chromosomes of the next generation, and deciding the termination of the model. Figure 1 illustrates the flowchart of building GEP classifier.

In this work, GEP is developed to predict two essential...
lungs cancer types. First of all, a set of functions and terminals is chosen to define chromosomes that will be expressed as nonlinear entities. The set of functions contains arithmetic operators and logic operators, such as +, -, *, +, min, max, equal, sqrt, log, exp, abs, while the set of terminals contains variables combination of serum biomarkers (for example: CEA+NSE+LDH and CEA+NSE+Cyfra21-1) and relevant coefficients. Then we build the chromosomal structure. For each chromosome the length of its head and the number of genes will be given.

The second step is to randomly initialize population. The input parameter indicates the size of population. For example, when GEP is used to deal with CEA+NSE+Cyfra21-1, we vary the whole number of populations from 100 to 180, and keep track of results each model produces. According to the results, the performance of models gradually increases as the number of populations rises. Thus, we choose the maximum number, 180, as the input populations.

The third step is to define a fitness function to evaluate the individual chromosome. In order to obtain an optimal output, we select SSPN as our fitness function which is described as the product of sensitivity (SN), specificity (SP), positive predictive value (PPV), and negative predictive value (NPV). The formula is as follows: SSPN=SNi x Pi x PPvi x NPvi. Where SN, SP, PPV, NPV are calculated respectively by using the following formulas for each chromosome: SNi=TPi/(TPi+FPi), SPi=TNi/(TNi+FNi), PPvi=TPi/(TPi+FPi), NPvi=TNi/(TNi+FNi).

The TNi, TPi, FNi and FPi are the numbers of true negatives, true positives, false negatives, and false positives, respectively. Given the content of 5 biomarkers, we use the fitness function to compute the scores of all chromosomes in population.

The forth step is to select the top 30% populations as the eugenic ones. Then the fifth step is that performing a set of genetic operations (including mutation, transposition and crossover) on eugenic ones reproduces chromosomes.

Finally, we choose the maximum as the number of generations to decide the termination of the model. In this study, the parameters used in our GEP classifier are listed in Table 2. We develop the program that predicts lung cancer types based on GEP in C++ Language.

### Results

**Test result of individual serum tumor markers**

SCLC and NSCLC groups are significantly different regarding to CEA, NSE, CPR, LDH and Cyfra21-1 concentrations in serum from 60 SCLC patients and 180 NSCLC subjects. The results of the measurements are as shown in Table 3.

**Classification and prediction**

Investigation of GEP models and their performance in diagnosing: We set up three GEP models. Observe their performances in diagnosing lung cancer types, then choose the one which obtains the highest prediction accuracy rate as the optimum GEP model.

The results of running the three GEP models were given in Table 4. It can be observed that in GEP3, 128 of 135 subjects in the training set as well as 41 of 60 subjects in the test set were classified correctly, its training set accuracy was 94.82% and the test set accuracy was 91.11%. By contrast, with the same function set, GEP1 represented the accuracy 94.07% and 77.78% while GEP2 represented 93.33% and 80.00% for training and testing set respectively. Compared with GEP1, the accuracy was significantly decreased by 1.5% and 6.6% in training set and test set in GEP2, in GEP3 was 0.82% and 4.45% respectively. It was concluded that the most accurate model was GEP3. The combination of CEA, NSE and Cyfra21-1 has the optimum predicting performance.

### Table 2. Parameters Used in Our GEP Method

| Parameter | Description of Parameter | setting of parameter |
|-----------|--------------------------|----------------------|
| P1        | Function set             | + - * / Exp Log Logi |
| P2        | Number of genes          | 5                    |
| P3        | Head Size                | 8                    |
| P4        | Linking Function         | Addition             |
| P5        | Number of chromosomes    | 50                   |
| P6        | Number of generation     | 200                  |
| P7        | Number of genes          | 5                    |
| P8        | Number of Tries          | 3                    |
| P9        | Max. Complexity           | 5                    |
| P10       | Mutation rate            | 0.044                |
| P11       | IS Transposition Rate    | 0.1                  |
| P12       | RIS Transposition Rate   | 0.1                  |
| P13       | Gene Transposition Rate  | 0.1                  |
| P14       | One-Point Recombination Rate | 0.3                 |
| P15       | Two-Point Recombination Rate | 0.3                 |
| P16       | Gene Recombination Rate  | 0.1                  |

### Table 3. Comparison of 5 Biomarkers Distribution in SCLC and NSCLC

| Biomarker | NSCLC(n=120) Mean ± Standard | SCLC(n=60) Mean ± Standard | P-value |
|-----------|------------------------------|---------------------------|---------|
| LDH       | 161.290±62.179               | 209.880±161.322           | >0.005  |
| CRP       | 25.079±24.817                | 14.935±21.078             | <0.001  |
| CEA       | 51.493±77.529                | 25.074±40.957             | <0.001  |
| NSE       | 13.638±5.571                 | 62.972±63.012             | <0.001  |
| CYFRA 21-1| 12.447±15.814                | 6.418±9.567               | <0.001  |

*Compare Small cell lung cancer group and non-small cell lung cancer group, the differences of CEA, NSE, CRP and Cyfra21-1 were statistically significant <0.01, the differences of LDH were not statistically significant: P>0.05.*
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Table 4. The Accuracy of GEP Models in Predicting Lung Cane Types

| GEP model       | Combination       | Training set (n=135) | Best fitness | Accuracy | Best fitness | Test set (n=45) | Accuracy |
|-----------------|-------------------|---------------------|--------------|----------|--------------|----------------|----------|
| GEP1            | 1.CEA+NSE+LDH     | 127                 | 94.07%       | 41       | 77.78%       |
| GEP2            | 2.CEA+NSE+CRP     | 126                 | 93.33%       | 36       | 80.00%       |
| GEP3            | 3.CEA+NSE+Cyfra21-1| 128                 | 94.82%       | 41       | 91.11%       |

**Table 5. Statistics of Test Set and Training Set**

| Statistics        | Training | Test |
|-------------------|----------|------|
| Calc. Errors:     | 0        | 0    |
| Accuracy:         | 0.9482   | 0.9111 |
| Error:            | 0.0518   | 0.0889 |
| Sensitivity:      | 1        | 1    |
| Specificity:      | 0.8444   | 0.7333 |
| PPV:              | 0.9278   | 0.88235 |
| NPV:              | 1        | 1    |
| Correlation Coefficient (CC): | 0.88 | 0.88 |
| Mean Squared Error (MSE): | 0.05 | 0.05 |
| Root Mean Squared Error (RMSE): | 0.23 | 0.23 |
| Relative Absolute Error (RAE): | 0.12 | 0.12 |
| Mean Absolute Error (MAE): | 0.05 | 0.05 |
| Relative Squared Error (RSE): | 0.23 | 0.23 |
| Root Relative Squared Error (RRSE): | 0.4 | 0.4 |

Variables and “+ * /Exp Log Logi” were used as function set. Therefore, a mathematical model of output variables was developed by using GEP. The program was coded in C++. The obtained formulations correspond to the following equations are: i) GEP1 (CEA+NSE+LDH) model is: \[ y = \text{apsLogid[1]+apsLogi[\log10 (d[1])]+apsLogid[\log10 (d[1])]+apsLogid[\log10 (d[1])]} \]. In this model, \( d[0], d[1], d[2] \) were LDH, CEA, NSE respectively; ii) GEP2 (CEA+NSE+CRP) model is: \[ y = \text{apsLogid[1]+apsLogi[\log10 (d[1])]+apsLogid[\log10 (d[1])]+apsLogid[\log10 (d[1])]} \]. In this model, \( d[0], d[1], d[2] \) were CRP, CEA, NSE.

The expression of GEP classifier with the best prediction performance, GEP2 (CEA+NSE+CRP) model was obtained as: \[ y = \text{apsLogid[1]+apsLogi[\log10 (d[1])]+apsLogid[\log10 (d[1])]} \]. In this model, \( d[0], d[1], d[2] \) were CRP, CEA, NSE, Cyfra21-1 respectively.

Analysis of the best GEP model

Expression tree: In order to achieve the geneticization of function expression, we need to use the K-expression (from Kava language) to represent the gene fragment. The process of expression is shown in Figure 2. Table V shoes the statistics of test set and training set.

Discussion

Because of its characteristics such as high degree of malignancy and difficulty to detect it early, lung cancer leads to unsatisfactory outcomes and 5-year survival rate. Roughly divided into two groups according to pathology: NSCLC (80.4%) and SCLC (16.8%), lung cancer becomes one of the major diseases which seriously threat human’s life and health (Travis et al., 1995; Welch et al., 2000). Patients with NSCLC are treated differently from those with SCLC. If the NSCLC can be diagnosed in early stage, it is possible that the patients would undergo surgery and may achieve healing, thereby reducing mortality. Therefore, the distinction between them is extremely important (Travis, 2011).

Many studies have considered lung cancer types (Li et al., 2005; Taguchi et al., 2007; Kligerman and Abbott, 2010; Wrona and Jassem, 2010; Nevins, 2011; Raj et al., 2011; Travis, 2011; Barash et al., 2012). For example, CT scanning, sputum cytology, LDCT, routine chest radiograph and, most recently, molecular biomarkers have evaluated various approaches for lung cancer classification. Based on chest X-Ray or sputum cytology, early screening tests of high risk individuals have not shown improvement in disease-specific survival. Chest LDCT scan, which recent NCCN guidelines endorsed as a screening tool for lung cancer, has been proven as an effective tool for the early detection of resectable disease. However, questions remain regarding the definition of lung cancer patients: how long and how frequent to screen; and high cost and potential toxicity from radiation exposure. The abnormality of the peripheral blood tumor markers is often earlier than radiographic abnormality; therefore for patients have no clinical symptoms, any abnormal markers could have prompted significant.

The expression of biomarkers in lung cancers is useful in the diagnosis and clinical management of patients with lung cancer. Biomarkers provide insight into histogenesis, interrelationships, and biological behavior of lung tumors. People have found a lot of tumor markers which were valuable for lung cancer diagnosis, efficacy detection and relapse diagnosis, such as AFP, PSA and CA125, have been proven respectively to be effective in the screening of liver, prostate and ovarian cancers (Perkins et al., 2003). Due to the complexity of tissue origin of lung cancer, involving the cancerous process of multiple genes and the heterogeneity of tumor antigens expression, a single tumor biomarker can not reflect lung cancer biological characteristics very well (Wang et al., 2013). Therefore, the sensitivity of a single tumor marker for diagnosis of lung cancer is relatively low (Tureci et al., 2006). Combining tumor marker examination can improve the lung cancer diagnostic rate, namely, combinatorial analysis of tumor markers is a new development direction of the experiment teaching as well as an active research topic recently. Regarding the importance and the need for finding a new simple effective method for lung cancer type detection, we proposed a GEP method to determine the biomarker combination that best discriminates NSCLC from SCLC subjects based on a variety of biomarkers obtained from the clinic and they are easily obtained in the impoverished region.
This chapter presents data on lung cancer detection, involving some of the most studied and interesting lung cancer biomarkers to date—Cyfra21-1, NSE, CEA, as well as markers in clinical application such as CRP, LDH. GEP1 trained with 3 biomarkers (CEA, NSE, Cyfra21-1), correctly classified 128 of 35 subjects in the training set as well as 40 of 45 subjects in the test set, acquiring the accuracy of 94.82% and 88.89% in the training set and test set. The accuracy of classification rate was slightly reduced in GEP2 (CEA, NSE, LDH) and GEP3 (CEA, NSE, CRP) since one biomarker was changed by 1.5% and 6.6%, 0.82% and 4.45% in the training set and test set, respectively. It is acknowledged that accuracy in GEP1 is higher than GEP2 and GEP3. GEP1 gained the best performance which trained with Cyfra21-1, NSE, CEA. Cyfra21-1 which showed the most powerful result compared to the other tumor markers, revealed a distinctive potential to differentiate patients with NSCLC from SCLC.

Cyfra21-1, which is a polypeptide tumor marker, is produced by almost all human cells and designated circulating cytokeratin-19 fragment (Wieskopf et al., 1995; Molina et al., 2003; Nakata et al., 2004). It is a unique epitope from a polypeptide and abundantly elaborated following cell death, whose diagnostic utility and prognostic relevance have been demonstrated in stomach cancer, colorectal cancer, breast cancer, and cervical cancer (Gaarenstroom et al., 1995; Lee, 2013; Wang et al., 2013; Gwak et al., 2014). Having been classified into 20 subtypes based on isoelectric point and differences in the molecular mass which was determined by 2-dimensional electrophoresis, Cyfra21-1 is a useful auxiliary biomarker in the diagnosis of NSCLC, and it has been reported that it has particularly high specificity for the diagnosis of lung squamous cell carcinoma (Ono et al., 2013). Therefore based on NSE, CEA and Cyfra21-1, the GEP1 accuracy is outstanding than GEP2 and GEP3. Moreover, the clinical information such as age, gender, smoking, nodules, hemoptysis and other index, may improve the GEP performance.

In summary, GEP is a nonlinear method of artificial intelligence with good reproducibility, which makes full use of the thoughts of biological replication expression. Based on GEP, we find a superior biomarker combination to distinguish lung cancer types. The auxiliary mode is convenient, economy and can be widely used in poverty-stricken areas. However, with the emergence of new predictive tumor markers, we are confined no longer to a single or several determinate tumor markers joint detection, but to large sample size, high amount of information and large scale on gene and protein levels.

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