Classifying Lung Adenocarcinoma and Squamous Cell Carcinoma using RNA-Seq Data

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ABSTRACT

Background: Lung adenocarcinoma (LUAD) and lung squamous cell carcinoma (LUSC) are two primary subtypes of non-small cell lung carcinoma (NSCLC). Currently, the most widely used method to discriminate between LUAD and LUSC is hematoxylin-eosin (HE) staining. However, this method sometimes is unable to make the precise diagnosis on LUAD or LUSC. More accurate diagnostic approaches are highly desired.

Methods: We propose to use gene expression profile to discriminate NSCLC patient’s subtype. We leveraged RNA-Seq data from The Cancer Genome Atlas (TCGA) and randomly split the data into training and testing subsets. To construct classifiers based on the training data, we considered three methods: logistic regression on principal components (PCR), logistic regression with LASSO shrinkage (LASSO), and kth nearest neighbors (KNN). Performances of classifiers were evaluated and compared based on the testing data.

Results: All gene expression-based classifiers show high accuracy in discriminating LUSC and LUAD. The classifier obtained by LASSO has the smallest overall misclassification rate of 3.42% (95% CI: 3.25%-3.60%) when using 0.5 as the cutoff value for the predicted probability of belonging to a subtype, followed by classifiers obtained by PCR (4.36%, 95% CI: 4.23%-4.49%) and KNN (8.70%, 95% CI: 8.57%-8.83%). The LASSO classifier also has the highest average area under the receiver operating characteristic curve (AUC) value of 0.993, compared to PCR (0.987) and KNN (0.965).

Conclusions: Our results suggest that mRNA expressions are highly informative for classifying NSCLC subtypes and may potentially be used to assist clinical diagnosis.

KEY WORDS: LUAD; LUSC; Principal Components; LASSO; Kth Nearest Neighbors.

ABBREVIATIONS: NSCLC: Non-Small Cell Lung Carcinoma; LUAD: Lung adenocarcinoma; LUSC: Lung squamous cell carcinoma; RNA-Seq: RNA sequencing; PCR: Principal Components Regression; LASSO: Logistic regression with lasso shrinkage; KNN: Kth nearest neighbors; ROC: Receiver Operating Characteristic; AUC: Area under the ROC curve.

INTRODUCTION

Lung cancer has the second highest estimated new cases rate and the highest estimated death rates for both male and female. According to American Cancer Society (ACS), the estimated new cases in 2015 are 115,610 and 105,590 which account for about 14% and 13% of all new cancer cases for male and female. The estimated deaths are 86,380 and 71,660 which account for about 28% and 26% of all deaths associated with cancer for male and female.1 Lung cancer can be classified as small cell lung carcinoma and non-small cell lung carcinoma (NSCLC). NSCLC weights more than 80% of all lung cancer. NSCLC can be sub-classified into Lung squamous cell carcinoma (LUSC), Lung adenocarcinoma (LUAD), and large cell carcinoma.2 Approximately 20% of all lung cancers are LUSC and it has the worst prognosis, and about 40% of all NSCLC cancers are LUAD.2
Despite the differences in prognosis, subtypes of NSCLC have been treated by similar strategies. The treatment effects vary in LUSC and LUAD patients. With the rapid development of the targeted therapies for NSCLC, more efficient treatments are available for the both NSCLC subtype. However, to choose treatments, especially targeted therapies and combination of interventions, we need accurate sub-typing between them.

Currently, the most widely used method to distinguish between LUAD and LUSC is hematoxylin-eosin (HE) staining of the tumor tissue sections observed under a light microscope. However, due to the reasons such as unclear structures in tumors and small biopsies with a limited number of tumor cells, by using only HE staining, it is difficult to make the precise diagnosis on LUAD and LUSC. Since the molecular profiling is different between LUAD and LUSC, Immunohistochemical (IHC) staining can help to diagnose between LUAD and LUSC. However, it needs knowledge of reliable IHC markers. Yu et al designed a fully automated informatics pipeline to extract quantitative image features and build classifiers to distinguish survival outcomes for lung cancer. They applied the classifiers to distinguish between LUAD and LUSC and obtained 0.75 as the highest area under the curve (AUC) value.

High-throughput data obtained from microarrays and RNA sequencing can be used to identify appropriate biomarkers for IHC staining. However, limited biomarkers are applied to IHC staining. By using all available high-throughput data, we may obtain better diagnostic outcomes. In our study, we applied three methods directly to public available RNA-Seq data released by the cancer genome atlas (TCGA) from National Cancer Institute (NCI).

METHODS

Data Source

Normalized level 3 RNA sequencing data of tumor samples were obtained from the R package RTCGARNA seq. The data includes 576 LUAD and 552 LUSC cases. The outcome to predict is LUAD versus LUSC, and the predictors are 20,259 gene expressions quantified using RNA-Seq by expectation maximization (RSEM) values. The original data includes 20531 genes; however, 272 have all zero values and thus were excluded from the analyses. In addition, we applied log transformation to achieve an approximately normal distribution for the data.

Overview of Classifier Construction and Evaluation

We applied the following three methods to predict lung adenocarcinoma (LUAD) and lung squamous cell carcinoma (LUSC): principal components regression (PCR), logistic regression with LASSO shrinkage (LASSO), and kth nearest neighbors (KNN). We randomly split the data into training and testing at a ratio of 3:1, where the training data were used to construct a classifier and the testing data were used to evaluate its performance. Details on the three methods are described in the next three sections.

After a classification model was constructed, we applied it to the testing data to obtain a predicted probability for each sample to be a certain subtype. We applied 0.5 as the cutoff point of the predicted probability to classify samples into the two subtypes. A set of statistics including overall misclassification rates, misclassification rates for LUAD and LUSC, and AUC values were calculated. In addition, we believed that genes used in the selected models were informative. Therefore, we tracked those genes from our selected models for PCR and LASSO. Due to method difference, KNN doesn’t return information on genes. For PCR, we selected 10 genes (5 with the largest positive and 5 with the largest negative loading values) from each of the first two principal components. For LASSO, we recorded all genes included in the selected model. The frequency of those genes will be reported in the Results section.

For each method, we repeated the process 500 times using 500 randomly generated seed numbers. Within each replication, we applied function “set.seed” with fixed seed number. In this way, we got same split datasets for all three methods to make the final results more comparable. All data processing and analyses were performed in R (version 3.3.2).

PCR: Logistic Regression on Principal Components

The “prcomp” function from R Stats package was used to obtain principal components from training data. We fitted the training data with a different number of principal components and recorded the AIC value for each model. The model with the smallest AIC value was identified as the best model.

LASSO: Logistic Regression with Lasso Shrinkage

The R package “glmnet” is used to fit the LASSO model. To select the shrinkage parameter lambda in the LASSO model, we further split the training dataset at a ratio of 2:1. We fit 100 models with different lambda values using the first portion of the data and calculate the overall misclassification rate of each model using the second portion of the data. The cutoff point to calculate the overall misclassification rate was set as 0.5. The best lambda value was chosen as the model that yields the smallest overall misclassification rate.

KNN: kth Nearest Neighbors

The R package “class” was used to build a KNN-based classifier. We applied the same strategy as what we did for LASSO to select the best number of nearest neighbors (K) in the KNN model. Specifically, the training data were split at a ratio of 2:1, where the first portion was used to fit 20 KNN models with K ranging from 1 to 20. Then the 20 models were applied to the second portion of the data to calculate the overall misclassification rate (cutoff point=0.5) for each model. The best model (best number for K) was selected as the one that achieved the smallest overall misclassification rate.
RESULTS

Table 1 shows the mean value and 95% confidence intervals for overall misclassification rate and misclassification rates for LUAD and LUSC. As we mentioned in Methods section, all misclassification rates were obtained by using 0.5 as the cut-off point. LASSO has the smallest overall misclassification rate 3.42% (95% CI: 3.25%-3.60%), followed by PCR 4.36% (95% CI: 4.23%-4.49%) and KNN 8.70% (95% CI: 8.57%-8.83%). Compared to PCR and KNN, LASSO also has the smallest misclassification rate for both LUAD (2.55%) and LUSC (4.36%).

Receiver operating characteristic (ROC) curves with area under the curve (AUC) values for all three methods are displayed in Figure 1. LASSO has the highest average AUC value (0.993), compared to PCR (0.987) and KNN (0.965).

Figure 2 shows the classification result from one replication. The test dataset for this replication included 145 LUAD and 137 LUSC cases. Among them, 7LUAD cases (red triangle) were classified into LUSC and 12LUSC cases (green plus) were classified into LUAD. The AUC value was 0.974 if only the first two principal components (PCs) were used. To explore whether increasing number of PCs in the model would help to reduce the misclassification rates and increase the AUC value, we considered a model including the first thirteen PCs. We got same overall misclassification rate and same misclassification rates for LUAD and LUSC from the selected model using the same cutoff point. The AUC value was 0.985, slightly higher than the AUC value from using only two PCs. Since by only using the first two PCs, LUAD and LUSC were well separated, we only recorded genes with large effects from the first two PCs.

Frequencies of genes identified by the first two principal components from PCR and by the selected model from LASSO are shown in Figure 3. There are 12 genes identified by the first principal component, 16 genes identified by the second one, and 408 genes identified by LASSO. We only display 20 genes with the highest frequency from LASSO. Among all identified genes, 21 genes have a 90% or higher frequency rate. Three genes, CALML3, DSG3, and KRT5 were identified by both PCR and LASSO.

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**Table 1:** Summary Statistics for PCR, LASSO, and KNN. Mean Misclassification Rates and 95% Confidence Intervals are shown in Percentages. All Statistics Presented in the Table are based on 500 Replications.

|          | PCR       | LASSO     | KNN       |
|----------|-----------|-----------|-----------|
| Misclassification Rate (%) | Mean (95% CI) | Mean (95% CI) | Mean (95% CI) |
| LUAD     | 4.36 (4.23-4.49) | 2.55 (2.41-2.69) | 7.25 (7.01-7.49) |
| LUSC     | 7.02 (6.84-7.20) | 4.36 (4.13-4.59) | 10.16 (9.93-10.39) |
| Overall  | 5.64 (5.51-5.76) | 3.42 (3.25-3.60) | 8.70 (8.57-8.83) |

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**Figure 1:** ROC Curves for PCR (plot A), LASSO (plot B), and KNN (plot C). Max AUC: Maximum AUC Value; Min AUC: Minimum AUC Value; Avg AUC: Mean AUC Value; 95% CI: 95% Confidence Interval for AUC Value. All Statistics Presented in the Figures are based on 500 Replications.
DISCUSSION

In this paper, we considered three different statistical methods to classify LUSC and LUAD patients based on their gene expression profiles. All the three methods have a low overall misclassification rate of less than 9% and high AUC of greater than 0.96. Our analyses demonstrate that gene expression data can accurately discriminate LUSC and LUAD samples, which does not specifically depend on the choice of statistical method.
Therefore, gene expression profile may potentially be used in clinic to enhance the diagnosis of LUSC and LUAD.

One issue we had is to standardize the data. Some genes have many zero values. After we split the whole data into subsets, the subsets may contain all zeros for certain genes, and those genes could be different in training and testing data, and they could also be different for each replication. To confirm the results that we got for all three methods, we removed all genes that have constant zeros for all subsets. This further reduced the number of genes included in the analysis. Then we applied three methods again on the standardized data. No significant changes were seen after the standardization. Therefore, we chose to process the data without standardization.

The model fitting based on the KNN method was sometimes unstable in our analysis. It might be caused by the high dimensions of the data. Different algorithms on distance metric can be used to improve the classification. Also, we only tested a number of neighbors (K) from 1 to 20. Larger K may improve the diagnostic performance. In contrast, PCR and LASSO were very stable. As for computational time, both PCR and LASSO were much less than it for KNN. On average, one replication took less than 1 minute and less than 10 seconds for PCR and LASSO, respectively. One KNN replication for K from 1 to 20 took more than 10 minutes. When we increased K, the computation time increased significantly. Since PCR and LASSO performed better on sub-typing LUAD and LUSC and had shorter computation time, we consider PCR and LASSO to be better methods than KNN.

Another advantage of PCR and LASSO is that we can get information on the contribution of each gene. Among those genes identified by our methods, SFTA3, DSG3, DSC3, and CALML3 were found to be useful to distinguish LUAD and LUSC from each other using earlier version TCGA data. In addition, DSG3, NAPSA, KRT3, KRT6A, KRT6B, and SFTA2 were identified as potential biomarkers for distinguishing between the two subtypes using different data sources. Unlike our study, those studies applied various differently expressed gene screening methods to identify potentially informative genes. Although, our main aim is to discriminate LUAD versus LUSC, we successfully identified many genes which were found in other studies. We believe that those high dimension reduction methods can help to discover potential biomarkers to distinguish between the two subtypes. Those methods also can be applied to other disease types known to have different molecular profiling.

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CONFLICTS OF INTEREST

The authors have no conflicts to declare.

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