Logistic Regression Analysis for LncRNA-Disease Association Prediction Based on Random Forest and Clinical Stage Data

BO WANG$^{1,3}$ AND JING ZHANG$^{2,1}$

$^1$College of Computer Science and Technology, Harbin Engineering University, Harbin 150001, China
$^2$School of Information Science and Engineering, University of Jinan, Jinan 250022, China
$^3$College of Computer and Control, Qiqihar University, Qiqihar 161006, China

Corresponding author: Jing Zhang (ise_zhangjing@ujn.edu.cn)

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ABSTRACT An increasing amount of studies have found that LncRNA plays an important role in various life processes of the body. In current prediction research on lncRNA-disease associations, correlation analysis of disease prognosis is overlooked. In this study, a logistic regression prediction model based on tumor clinical stage data and the expression quantity of lncRNA transcript is constructed. The proposed model is based on unknown human lncRNA-disease associations combining with the clinical stage data. Firstly, the importance of the characteristic variable is calculated by the proposed CVS$_g$C-RF algorithm. Secondly, 95 lncRNAs, which are most closely related to prostate cancer, are calculated from 480 alternative lncRNAs by CASO and CVS$_e$CS-CF. On the basis of the above 95 lncRNAs, the CSPA-PL algorithm is used to select a further 22 lncRNAs that are most closely related to the tumor clinical stage for prostate cancer. Finally, 22 lncRNAs are used to construct a logistic regression prediction model. Additionally, this method is applied to lung cancer data; 16 lncRNAs are selected to construct a logistic regression prediction model for lung cancer. Experimental results show that the best results for ROC Area, the accuracy and recall rate of the prediction model are achieved by the proposed method for prostate cancer and lung cancer, which provides a promising basis for subsequent prediction studies of lncRNA-disease associations.

INDEX TERMS LncRNA-disease association, random forest, logistic regression analysis, clinical stage data.

I. INTRODUCTION
Long non-coding RNA (IncRNA) is non-coding RNA with more than 200 nucleotides in length [1]. It has very important biological functions and is another important area in the bioinformatics field [2], [3]. Studies show that IncRNA is closely correlated with many diseases, such as lung cancer [4], [5], Alzheimer’s disease [6], osteosarcoma [7], breast cancer [8], gastric cancer [9], colon cancer [10], prostate cancer [11], cervical cancer [12], etc. At present, more and more researchers are engaged in research in this area, which is an important molecular target in the diagnosis and treatment of disease. It is extremely important to study the relationship between IncRNA and the prognosis of cancer patients by utilizing clinical data. Current research in IncRNA is in the initial stages; people still know little about the deep mechanisms in the occurrence and development of cancer. Therefore, it is important to study IncRNA, which has a significant impact on the prognosis of cancer patients by using bioinformatics combined with clinical data.

Relevant research results in recent years are broadly divided into three types, as follows.

The first type is machine-learning-based methods and known disease-related IncRNAs. For instance, Yu et al. [13] proposed a new method called CFNBC based on the Naïve Bayes classifier to predict IncRNA-disease association. The novelty of CFNBC lies in the introduction of the item-based collaborative filtering algorithm and Naïve Bayes classifier,
which guarantee that CFNBC can be applied to predict potential lncRNA-disease associations efficiently without entirely relying on known miRNA-disease associations. Cui et al. [14] developed a novel model called BLM-NPAI for predicting lncRNA-disease associations. The main advantage of BLM-NPAI was that it could also make predictions using nearest neighbors for some lncRNAs and diseases without any association. Chen and Yan [15] used semi-supervised learning to predict the potential associations between lncRNAs and diseases, and proposed the first lncRNA-disease association prediction model (LRLSLDA) on the premise that similar functions of lncRNA tended to result in similar diseases. However, the model was too complex and exhibited high computational complexity. Meanwhile many parameters need to be selected in the calculation process. Huang et al. [16] improved the calculation of disease similarity based on the framework of LRLSLDA to further improve the prediction results, and presented a new method, ILNC-SIM. This approach kept the general hierarchical structure information of disease DAGs and determined the disease similarity calculation based on an edge-based method. Finally, the prediction performance was improved to some extent, but there were still some limitations. For example, the similarity score in the model needs to be further optimized. The lack of unrecorded but real lncRNA-disease associations had a large impact on the model, and the integration of multiple types of data was lacking. Chen [17] built a new approach (KATZLDA) by integrating known lncRNA-disease associations, lncRNA expression profiles, lncRNA functional similarity, disease semantic similarity and Gaussian interaction profile kernel similarity to predict the potential lncRNA-disease associations. The biggest advantage of KATZLDA was that it could be effectively applied to new diseases and lncRNAs without any known associations. However, the learning network built by KATZLDA was based on the known correlation relationship, so this was limited by the known learning knowledge and had certain limitations in prediction. Zhao et al. [18] constructed a multi-source data set by integrating multidimensional data (genome, regulatory group and transcriptome), and proposed a Bayesian classification method using this multi-source data to predict the lncRNA–disease associations. Experimental results showed that this method successfully identified 707 lncRNAs related to human cancer. However, this method was a supervised classification algorithm which required a large number of negative cases, but these are difficult to obtain.

The second type is network-based methods. For instance, Li et al. [19] present a novel network consistency projection approach called NCPLDA for lncRNA-disease association prediction. The network was built by integrating the lncRNA-disease association probability matrix with the integrated disease similarity and lncRNA similarity. Zhou et al. [20] proposed a new method (RWRHLD) that built a heterogeneous network of lncRNA–disease associations, on which a random walk algorithm was executed. However, the limitation was that the incomplete coverage of the lncRNA crosstalk network and the lncRNA–disease associations could lead to inaccurate predictions. Liu et al. [21] established a bidirectional network of protein-coding genes (PCG) and lncRNA for the prostate cancer and protein interaction databases based on lncRNAs and PCG expression maps, and further realized lncRNA–disease association prediction based on this network. However, the method was limited by the incomplete protein interaction database, and its performance had some limitations.

The third type is RW-based methods (RW is short for random walk). For instance, Li et al. [22] proposed a prediction model called LRWHHLDA for inferring LncRNA-disease association. LRWHHLDA can be implemented in the case of lacking known lncRNA-disease associations by using an improved local random walk method. Yu et al. [23] used multidimensional heterogeneous data to construct lncRNA networks with similar functions and the disease ontology to construct disease networks. On this basis, BRWMLDA was proposed to predict the lncRNA-disease associations. BRWMLDA improved the random walk model and the prediction performance to some extent.

To summarize, the limitations of the current research were described by the review [24] and the aforementioned discussions. Current studies have ignored correlation analysis of clinical prognosis, concerns of the prediction model have been limited to a single lncRNA forecast. The clinical prognosis of the disease associated with lncRNA information is rarely involved, such as tumor clinical stage, tumor pathological stage, survival time, disease status, family history of genetic diseases, and so on.

In this study, a logistic regression prediction model of lncRNA-disease associations based on the tumor clinical stage data was constructed. Three kinds of circular allelism operations (\(\Gamma_{\text{center}}(\Theta_{\text{sub}}^{\text{tumor}}), \Gamma_{\text{X-axis}}(\Theta_{\text{sub}}^{\text{normal}}), \Gamma_{\text{Y-axis}}(\Theta_{\text{sub}}^{\text{normal}})\)) were proposed for the prediction model. The calculation of the significance of characteristic variables based on random forests was proposed, and the selection algorithm for the characteristic variables was given. Finally, the clinical stage prediction algorithm of cancer-associated lncRNA was implemented using the simplified characteristic variables. Experimental results showed that the proposed method had a higher predictive performance.

II. MATERIALS AND METHODS

A. LncRNA DATA

The lncRNA expression data for prostate cancer was obtained from the lncRNAAtlas database [25]. A total of 220 samples were obtained (denoted by \(S_{\text{normal,tumor}} = \{S_1, \ldots, S_{220}\}\)), including 44 normal samples (denoted by \(S_{\text{normal}} = \{S_1, \ldots, S_{44}\}\)) and 176 cancer samples (denoted by \(S_{\text{tumor}} = \{S_{45}, \ldots, S_{220}\}\)). Based on the differential expression P-value (\(P \leq 0.001\)) of lncRNA transcripts between \(S_{\text{normal}}\) and \(S_{\text{tumor}}\), 480 lncRNA transcripts with significant differences (denoted by \(L_{R1}, L_{R2}, \ldots, L_{R480}\)) in ascending order of P-value) were obtained. Of these, 480 lncRNA transcripts
were denoted by \( L_r = \{L_{r1}, L_{r2}, \ldots, L_{ri} \mid 1 \leq i \leq 480 \} \), \( L_{r\text{sub}} \) was the subset of \( L_r (L_{r\text{sub}} \subseteq L_r) \) and the expression of \( L_r \) on \( S_i \) was denoted by \( L_r^{S_i} = \{L_{r1}^{S_i}, L_{r2}^{S_i}, \ldots, L_{ri}^{S_i} \mid 1 \leq i \leq 480 \} \). Figure 1 shows the details of the top 20 IncRNA transcripts in the 480 transcripts, where each row represents one IncRNA transcript. The first column is the rank of the transcript, the second column is ensemble gene ID, the third column is the gene name, the fourth column is the ensemble transcript ID, the fifth column contains the P values of the differential expression between normal samples and cancer samples. Each column after the fifth column is the expression quantity of the transcript in the sample. These transcripts were the more pronounced differences between normal and cancer samples.

### B. CLINICAL DATA

Clinical data associated with \( S^{normal \cup tumor} \) were obtained from the TCGA database (https://cancergenome.nih.gov). The aliquot barcode of the clinical data in \( S^{normal} \) (size 44) and \( S^{tumor} \) (size 176) are presented in Table 1. Each \( S_i \) contains 70 clinical reference values. Some of these were retained as follows: barcode and sample type of \( S^{normal \cup tumor} \) (denoted by \( P^{normal \cup tumor} = \{P_1, \ldots, P_{220}\} \)), tumor clinical stage of \( S^{tumor} \) (denoted by \( C^{tumor} = \{CT_1, \ldots, CT_{176}\} \)). Of these, the barcode was used to correlate the clinical data with the IncRNA data, the sample type was used to select the characteristic variables of the IncRNA-disease associations, and the tumor clinical stage was used to predict IncRNA with significant impact on the prognosis of cancer patients combined with the clinical data. In this case, the clinical stage (CTNM) was performed using the TNM stage system, where T represents the tumor size, N represents lymph node metastasis, and M represents distant metastasis. The distribution of \( CT^{tumor} \) (size 176) associated with \( S^{tumor} \) is shown in Table 2. As can be seen from this table, 118 effective clinical stage data and 58 invalid clinical stage data were obtained. Variance analysis of \( \{T_1 \cup T_2\} \) and \( \{T_1 \cup T_2 \cup T_3\} \) showed that \( \{T_1 \cup T_2\} \) has a better-balanced distribution and is conducive to machine classification learning. Finally, 104 available data (denoted by \( CT^{\leq tumor} = CT^{tumor} \cap \{T_1 \cup T_2\} = \{CT_1^\leq, CT_2^\leq, \ldots, CT_{104}^\leq\} \) ) were selected from 118 valid clinical stage data.

The following two matrices were constructed by combining IncRNA data and clinical data for the prediction study in this paper.

(a) The matrix \( M^{CV} \) of the characteristic variables is shown in (1), which contains 480 columns of characteristic variables, 1 categorical variable column, and 220 rows of sample data. After the selection algorithm, the \( \lambda \) IncRNAs that were most closely related to prostate cancer were screened from 480 characteristic variables.

\[
M^{CV} = L_r^{S_tumor} \sim P^{normal \cup tumor} = \begin{bmatrix}
L_r^{S_1} & L_r^{S_2} & \cdots & L_r^{S_n} & P_1 \\
L_r^{S_1'} & L_r^{S_2'} & \cdots & L_r^{S_n'} & P_2 \\
\vdots & \vdots & \ddots & \vdots & \vdots \\
L_r^{S_m} & L_r^{S_m'} & \cdots & L_r^{S_m''} & P_m
\end{bmatrix} \\
(n = 480, m = 220)
\]

\[
P_{1 \leq i \leq 44} = [normal, tumor] \times \begin{bmatrix} 1 \\
0 \\
1 \end{bmatrix},
\]

\[
P_{45 \leq i \leq 220} = [normal, tumor] \times \begin{bmatrix} 0 \\
0 \\
1 \end{bmatrix} \quad (1)
\]

(b) The matrix \( M^{PP} \) of prognosis prediction is shown in (2), which contains \( \lambda \) columns of characteristic variables, 1 categorical variable column, and 104 rows of sample data.

\[
M^{PP} = L_r^{S_tumor} \sim CT^{\leq tumor} = \begin{bmatrix}
L_r^{S_1} & L_r^{S_2} & \cdots & L_r^{S_1} & CT_1^{\leq} \\
L_r^{S_1'} & L_r^{S_2'} & \cdots & L_r^{S_1'} & CT_2^{\leq} \\
\vdots & \vdots & \ddots & \vdots & \vdots \\
L_r^{S_m} & L_r^{S_m'} & \cdots & L_r^{S_m'} & CT_m
\end{bmatrix} \\
(L_r^{S_1} = L_r^{S(1-45)}, m = 104)
\]
C. CASO METHOD

The circular allelism subarea operation (abbreviated to CASO) for characteristic variable selection is described below. The 480 \( L_r \)s in \( M^{CV} \) formed a circular queue \( \Theta \). According to the importance of each \( L_r \), the \( L_r \) in descending order are evenly clockwise distributed on the ring \( \Theta \). \( \Theta \) contains 480 nodes in descending order of importance (denoted by \( \Theta = \{S_1, S_2, \cdots, S_{480}\} \)). The \( \text{Significance}(L_r) \) is the importance of \( L_r \). The descending rank of \( \text{Significance}(L_r) \) is \( j = \text{rank}(\text{Significance}(L_r)) \).

Here, the node \( S_j \) is \( L_r \), and the subset of \( \Theta \) is denoted by \( \Theta_{[a,b]} = \{S_a, \cdots, S_b\} (1 < a < b < 480) (\Theta_{[a,b]} \subset \Theta) \). The descending queue formed with \( L_r \) in descending order according to \( \text{Significance}(L_r) \) is \( Q_{\text{Dec}}(L_r) \).

The circular queue \( \Theta \) is formed as shown in Figure 2. The descending order from \( S_1 \) to \( S_{480} \) is evenly clockwise distributed on \( \Theta \). The center symmetric points of \( S_a \) and \( S_b \) are \( S_{a-\text{center}} \) and \( S_{b-\text{center}} \). The X-axis symmetric points of \( S_a \) and \( S_b \) are \( S_a-X \) and \( S_b-X \). The Y-axis symmetric points of \( S_a \) and \( S_b \) are \( S_a-Y \) and \( S_b-Y \).

Figure 2 illustrates the following. The red line, area and data represent the center allelism operation. The green line, area and data represents the X-axis allelism operation. The yellow line, area and data represents the Y-axis allelism operation.

The circular queue \( \Theta \) is divided into four areas (the first quartile area, the second quartile area, the third quartile area, and the fourth quartile area).

\textbf{Definition 1 (Center Allelism Operation)} \( \Gamma_{\text{center}}(\Theta_{[a,b]}) \):

\[ \Gamma_{\text{center}}(\Theta_{[a,b]}) \text{ is defined as } \Theta_{[a,b]} \cup \Theta_{[a-\text{center},b-\text{center}]} \] is the center allelism area of \( \Theta_{[a,b]} \). The set of central symmetric points of \( \Theta_{[a,b]} = \{S_a, \cdots, S_b\} \) is denoted by \( \Theta_{[a-\text{center},b-\text{center}]} = \{S_{a-\text{center}}, \cdots, S_{b-\text{center}}\} \).

\textbf{Definition 2 (X-Axis Allelism Operation)} \( \Gamma_{X-axis}(\Theta_{[a,b]}) \):

\[ \Gamma_{X-axis}(\Theta_{[a,b]}) \text{ is defined as } \Theta_{[a,b]} \cup \Theta_{[X-axis,b-\text{axis}]} \cdot \Theta_{[X-axis,a-\text{axis}]} \] is the X-axis allelism area of \( \Theta_{[a,b]} \). The set of X-axis symmetric points of \( \Theta_{[a,b]} = \{S_a, \cdots, S_b\} \) is denoted by \( \Theta_{X-axis} = \{S_a-X, \cdots, S_b-Y\} \).

\textbf{Definition 3 (Y-Axis Allelism Operation)} \( \Gamma_{Y-axis}(\Theta_{[a,b]}) \):

\[ \Gamma_{Y-axis}(\Theta_{[a,b]}) \text{ is defined as } \Theta_{[a,b]} \cup \Theta_{[Y-axis,b-\text{axis}]} \cdot \Theta_{[Y-axis,a-\text{axis}]} \] is the Y-axis allelism area of \( \Theta_{[a,b]} \). The set of Y-axis symmetric points of \( \Theta_{[a,b]} = \{S_a, \cdots, S_b\} \) is denoted by \( \Theta_{Y-axis} = \{S_a-Y, \cdots, S_b-Y\} \).

The red area in Figure 2 is \( \Gamma_{\text{center}}(\Theta_{[a,b]}) \), which is located in the first quartile area and the third quartile area. The green area in Figure 2 is \( \Gamma_{X-axis}(\Theta_{[a,b]}) \), which is located in the first quartile area and the second quartile area.

Additionally, \( \Theta_{X-axis} \) is closer to \( \Theta_{[a,b]} \) than \( \Theta_{[a-\text{center},b-\text{center}]} \) is centered on \( \Theta_{[a,b]} \) and \( \Theta_{Y-axis} \) is far from \( \Theta_{[a,b]} \), the above \( \Gamma_{\text{center}}(\Theta_{[a,b]}) \), \( \Gamma_{X-axis}(\Theta_{[a,b]}) \), and \( \Gamma_{Y-axis}(\Theta_{[a,b]}) \) constitute the circular subset \( \Theta \) required by the next algorithm.

D. RANDOM FOREST

Random forest (abbreviated to RF) is an enhanced classifier constructed by multiple decision trees. In the process of building the decision tree, it is necessary to order the importance of variables. Since a random forest has a large number of decision trees, the importance obtained from each decision tree could be integrated to obtain the final importance rank of the variables. The selection of characteristic variables is carried out according to the order of the variables, which is more stable and reliable than a single decision tree. In the selection of \( M^{CV} \) characteristic variables based on RF, RF contained \( \alpha \) trees (denoted by \( T = \{T_1, \cdots, T_i, \cdots, T_{\alpha}\} \)). 480 \( L_r \)s (denoted by \( L_r = \{L_{r1}, \cdots, L_{ri}, \cdots, L_{r480}\} \)) in \( M^{CV} \) are the characteristic variable set, and 220 \( Ps \) (denoted by \( P = \{P_1, \cdots, P_i, \cdots, P_{220}\} \)) in \( M^{CV} \) are the classified variable set.

E. CVS\( _G \)C-RF ALGORITHM

The significance computing of the characteristic variables based on RF (abbreviated to CVS\( _G \)C-RF) is given in Algorithm 1. Here, OOB means out-of-bag. Records are extracted from the original data to construct the training set for decision tree learning. Because this process uses sampling with replacement, some samples are not included, termed out-of-bag. On average, 37% of the data is not selected in each
sampling with replacement, which is often used to validate the constructed decision tree model. If the characteristic variable \( L_{R_1} \) upset on OOB has no effect on the result of the decision tree, then \( L_{R_1} \) is deemed to be not important. If the reverse is true, then \( L_{R_1} \) is very important.

**Algorithm 1 CVS_{sub}-C-RF(\( L_{R_{sub}}, \tau, \eta \))**

```plaintext
1: \( RF(\{L_{R_{sub}}\}, \tau, \eta) \);
2: \( \forall \in T_k \in \text{Do} \)
3: \( OOB(T_k) = \text{MeanDecreaseAccuracy}^{\text{O}}(\text{OOB} \in T_k) \);
4: \( \forall \in L_{R_1} \in \text{L}_{R_{sub}} \text{Do} \)
5: \( T_k^{L_{R_1}} \leftarrow \text{Randomupset}([L_{R_1}^{S_1}, L_{R_1}^{S_2}, \ldots, L_{R_1}^{S_{220}}]) \);
6: \( \text{MeanDecreaseAccuracy}^{\text{O}}(\text{OOB} \in T_k^{L_{R_1}}) \);
7: \( \text{Significance}(\tau) = \text{OOB}(T_k^{L_{R_1}}) - \text{OOB}(T) \);
8: \( \text{L}_{R_{sub}} \leftarrow \text{L}_{R_{sub}} - \text{L}_{R_1} \);
9: \( \text{end for} \)
10: \( T \leftarrow T - T_k \);
11: \( \text{end for} \)
12: \( \text{Significance}(\tau) = \sum_{i=1}^{\tau} \text{Significance}(\tau) \);
13: \( Q^{\text{Dec}}(\text{L}_{R_{sub}}) = \text{Sort}(\text{L}_{R_{sub}}) \text{Significance}(\tau) \);
14: \( \text{return} \; Q^{\text{Dec}}(\text{L}_{R_{sub}}) \);
```

The CVS_{sub}-C-RF algorithm is shown in Algorithm 1. In this algorithm, \( RF(\{L_{R_{sub}}\}, \tau, \eta) \) is a random forest containing \( \tau \) decision trees being trained on \( L_{R_{sub}} \); \( \tau \) is the number of decision trees contained in a random forest. \( \eta \) is the number of random characteristic variables contained in each partition \( \eta = \left\lfloor \sqrt{RF_{\text{cv-number}}(L_{R_{sub}}) + 0.5} \right\rfloor \).

\( RF_{\text{cv-number}}(L_{R_{sub}}) \) is the number of characteristic variables in \( L_{R_{sub}} \). The relationship between \( \tau \) and \( RF_{\text{cv-number}}(L_{R_{sub}}) \) is \( \tau \propto RF_{\text{cv-number}}(L_{R_{sub}}) \).

**MeanDecreaseAccuracy^{\text{O}}(\text{OOB})** is a standardized prediction error rate for OOB data. \( \text{Randomupset}([L_{R_1}^{S_1}, L_{R_1}^{S_2}, \ldots, L_{R_1}^{S_{220}}]) \) is the random upset operation on a characteristic variable \( L_{R_1} \). All characteristic variables in \( L_{R_{sub}} \) are arranged in descending order according to \( \text{Significance}(\tau) \) (denoted by \( \text{Sort}(\text{L}_{R_{sub}}) \text{Significance}(\tau) \)). Further, a queue \( Q^{\text{Dec}}(\text{L}_{R_{sub}}) \) is formed.

**F. DISCUSSION OF CVS_{E}**

A good characteristic variable selection (abbreviated to CVS_{E}) algorithm must possess both global selectivity and local stability. Global selectivity and local stability are mutually restricted. For example, the larger the selection range, the more complex the mutual relations among the characteristic variables, and bidirectional influence relations coexist. Local stability is needed for adjustment, but is limited by the selection range, which leads to insufficient coverage of the correlation between characteristic variables. At this point, an increase in global selectivity is required. Therefore, how to adjust the global selectivity and local stability of an algorithm is very important, and could affect the performance of CVS_{E}.

**G. CVS_{sub}-CS-CF ALGORITHM**

CVS_{E} combines CASO and CVS_{E}-C-RF (named CVS_{E}-CS-CF). CVS_{E}-CS-CF algorithm for \( M^{CV} \) is provided in Algorithm 2. The \( L_{R_1} \) in \( M^{CV} \) is selected by the CVS_{E}-CS-CF algorithm. The \( \lambda L_{R_1} \) most closely related to categorical variables is selected in 480 \( L_{R_1} \). In order to possess both global selectivity and local stability, the CVS_{E}-CS-CF algorithm is divided into a primary stage, a stable stage and a run-off stage. The primary stage can guarantee global selectivity, the stable stage guarantees local stability, and the run-off stage guarantees the final selection result by combining global selectivity and local stability.

**Algorithm 2 CVS_{sub}-CS-CF(\( L_{R_1}, \omega_{0-9}, \eta_{0-9}, \Delta, \Omega, \alpha, \beta)\)**

```plaintext
1: \( \Omega = CVS_{E} - RF(L_{R_1}, \omega_{0-9}, \eta_{0-9}) \);
2: \( \text{Setpenalty} \leftarrow \emptyset \), \( \text{Setshock} \leftarrow \emptyset \);
3: \( \text{while} distance([\Delta, \Omega]) \geq \Omega \; \text{do} \)
4: \( \text{if} \{ [\alpha, \beta] \} \Delta \text{dimidiate then} \)
5: \( Q^{++f} = CVS_{E} - RF(L_{R_1}, \omega_{0-9}, \eta_{0-9}) \);
6: \( Q^{++f} = CVS_{E} - RF(L_{R_1}, \omega_{0-9}, \eta_{0-9}) \);
7: \( Q^{++f} = CVS_{E} - RF(L_{R_1}, \omega_{0-9}, \eta_{0-9}) \);
8: \( \text{else} \)
9: \( \text{end if} \)
10: \( \text{end while} \)
11: \( \text{for each} \; Q_f \{ f \in [1, \max(f)] \} \; \text{do} \)
12: \( \text{for each} \; L_{R_1} \in Q_f \cup \{ \Theta_{[1,22]} \} \; \text{do} \)
13: \( \text{Setpenalty} \leftarrow L_{R_1} \);
14: \( \text{end if} \)
15: \( \text{end for} \)
16: \( \text{end if} \)
17: \( \text{if} \; L_{R_1} \in [\Theta_{[1,22]}] \) and \( L_{R_1} \neq Q_f \; \text{then} \)
18: \( \text{Setpenalty} \leftarrow L_{R_1} \);
19: \( \text{end if} \)
20: \( \text{if} \; L_{R_1} \in [\Theta_{[1,22]}] \) and \( L_{R_1} \in Q_f \; \text{then} \)
21: \( \text{Setshock} \leftarrow L_{R_1} \);
22: \( \text{end if} \)
23: \( \text{end for} \)
24: \( \text{return} \; \Theta_{\text{select}} \), \( \lambda \);```

**Primary Stage:** The 480 \( L_{R_1} \)s are ranked in descending order of importance, with the top 2\( \Omega \) \( L_{R_1} \)s entering the candidate area and the remaining 480 −2\( \Omega \) \( L_{R_1} \)s entering the observation area. Since the \( L_{R_1} \) entering the candidate area is selected in the global range of 480, global selectivity is obtained. However, the larger the scope, the more complex the relationship of \( L_{R_1} \) will become, and the bidirectional influence relationship exists. In the following stable stage,
and the three allelism areas of the above observation areas are rearranged in descending order of importance. (This is the CVS\textsubscript{g}-C-RF algorithm that was discussed previously.) Each \( L_r \) that is screened from the candidate area into the observation area is added to the penalty set. The top \( \Omega L_r/\Omega S \) that are screened from the observation area into the candidate area are added to the shock set. Each \( L_r \) in \( Set_{\text{penalty}} \) has a potential risk of poor stability. Each \( L_r \) in \( Set_{\text{shock}} \) has strong reactivation activity. The detailed process is described in steps 3-24 of Algorithm 2.

Run-Off Stage: \( Set_{\text{penalty}} \) and \( Set_{\text{shock}} \) are used to update each \( L_r \) in the candidate area. Remove \( L_r \) in \( Set_{\text{penalty}} \) from the candidate area and add it to \( Set_{\text{shock}} \) in the candidate area. The updated candidate area is the result of characteristic variable selection (denoted by \( \Theta_{\text{select}} \)). The number of characteristic variables in \( \Theta_{\text{select}} \) is \( \lambda \). The detailed process is described in steps 25-27 of Algorithm 2.

In Figure 3, \( \Omega \) is set to 60, the candidate area is \([S_1, S_{120}]\), and the observation area is \([S_{120}, S_{480}]\). \( \Delta_1, a \) and \( b \) are set to 120, 1, and 120 in Figure 3(a). The area where the CASO is performed is \([S_1, S_{120}] \) (shown in pink). The center allelism area is \( \Theta_{\text{sub-center}} = [S_{241}, S_{360}] \) (shown in red). The X-axis allelism area is \( \Theta_{\text{sub-X-axis}} = [S_{121}, S_{240}] \) (shown in green). The Y-axis allelism area is \( \Theta_{\text{sub-Y-axis}} = [S_{361}, S_{480}] \) (shown in yellow). Finally, the three allelism area combinations are \([S_1, S_{120}] \cup [S_{241}, S_{360}], S_{121} \cup [S_{121}, S_{240}], \) and \([S_1, S_{120}] \cup [S_{361}, S_{480}]\). \( \Delta_1, a \) and \( b \) are set to 60, 1, and 60 in Figure 3(b). The area where the CASO is performed is \([S_1, S_{60}] \) (shown in pink). The center allelism area is \( \Theta_{\text{sub-center}} = [S_{241}, S_{300}] \) (shown in red). The X-axis allelism area is \( \Theta_{\text{sub-X-axis}} = [S_{181}, S_{240}] \) (shown in green). The Y-axis allelism area is \( \Theta_{\text{sub-Y-axis}} = [S_{361}, S_{480}] \) (shown in yellow). Finally, the three allelism area combinations are \([S_1, S_{60}] \cup [S_{241}, S_{300}], S_{181} \cup [S_{181}, S_{240}], \) and \([S_{121}, S_{240}] \cup [S_{361}, S_{480}]\). \( \Delta_1, a \) and \( b \) are set to 60, 1, and 120 in Figure 3(c). The area where the CASO is performed is \([S_{61}, S_{120}] \) (shown in pink). The center allelism area is \( \Theta_{\text{sub-center}} = [S_{301}, S_{360}] \) (shown in red). The X-axis allelism area is \( \Theta_{\text{sub-X-axis}} = [S_{121}, S_{180}] \) (shown in green). The Y-axis allelism area is \( \Theta_{\text{sub-Y-axis}} = [S_{361}, S_{420}] \) (shown in yellow). Finally, the three allelism area combinations are \([S_{61}, S_{120}] \cup [S_{301}, S_{360}], S_{61} \cup [S_{121}, S_{180}], \) and \([S_{61}, S_{120}] \cup [S_{361}, S_{420}]\).

**FIGURE 3.** A CASO process of \( L_r \) in candidate area.

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**H. CSPA-PL**

The CVS\textsubscript{g}-CS-CF algorithm selected \( \lambda \) lncRNAs that were most closely related to prostate cancer. Next, \( \lambda \) lncRNAs are correlated with the prognosis data of the tumor clinical stage, and a logistic regression model is adopted to propose a clinical stage prediction algorithm for cancer-associated lncRNA (abbreviated to CSPA-PL). Concerning the related operations of \( \Theta_{\text{select}} \), CSPA-PL is divided into an inspection stage and an optimization stage. CSPA-PL is described in Algorithm 3.
Algorithm 3 CPSP-PL($\Theta_{select}, \lambda, M^{PP}$)

1: $\Theta_{select}_{sub-pre} = \Theta_{select}_{1, \frac{1}{2}1} [\frac{1}{2} + 1, \lambda]$; $\Theta_{select}_{sub-rear} = \Theta_{select}_{1, \frac{1}{2}1} [\frac{1}{2} + 1, \lambda]$;
2: $Z_{[1,n]}^{pre} \leftarrow \text{Logistic} - \text{Step}(\pi_{\Theta_{select}_{sub-pre}}(M^{PP}))$;
3: $\{\gamma^{pre}\} \leftarrow \min(\gamma_{[1,n]}^{pre}) \text{AIC};$
4: $Z_{[1,n]}^{rear} \leftarrow \text{Logistic} - \text{Step}(\pi_{\Theta_{select}_{sub-rear}}(M^{PP}))$
5: $\min(Z_{[1,n]}^{rear}) / \text{AIC};$
6: for each $L_{i}$ in minimum($Z_{[1,n]}^{rear}$) do $\gamma_{[1,n]}^{rear}$ do
7: if $\text{Value}_{\gamma^pre}(L_{i}) < 0.01$ then $\gamma_{[1,n]}^{rear}$ do
8: $\{\gamma^{rear}\} \leftarrow L_{i}$; $\gamma_{[1,n]}^{rear}$ do
9: end if $\gamma_{[1,n]}^{rear}$ do
10: end for $\gamma_{[1,n]}^{rear}$ do
11: $\{\text{Buffer} - \text{pool}\} = [\{\gamma^{pre}\} \cup \{\gamma^{pre}\}] \times [\{\gamma^{rear}\}]$;
12: for each $\text{Buffer} - \text{pool}$ in $\text{Buffer} - \text{pool}$ do $\text{Buffer} - \text{pool}$ do
13: $\Phi_{1} \leftarrow \text{Logistic}(\pi_{\text{Buffer} - \text{pool}}(M^{PP}))$; $\text{Buffer} - \text{pool}$ do
14: $\Phi_{optimal} \leftarrow \max(\Phi_{1})$ $\text{AccuracyRate}$;
15: end for $\text{Buffer} - \text{pool}$ do
16: $\text{return} \Phi_{optimal}$

Inspection Stage: Before being applied to the clinical stage prediction model, the $\lambda, L_{i}$s most closely associated with prostate cancer need to be inspected. The detailed process is described in steps 1-11 of Algorithm 3. In the inspection stage, $\Theta_{select}$ is divided into $\Theta_{select}_{1, \frac{1}{2}1}$ and $\Theta_{select}_{1, \frac{1}{2}1} [\frac{1}{2} + 1, \lambda]$, and then $\text{Logistic} - \text{Step}(\cdot)$ is performed on $\Theta_{select}_{1, \frac{1}{2}1}$ and $\Theta_{select}_{1, \frac{1}{2}1} [\frac{1}{2} + 1, \lambda]$. $\text{Logistic} - \text{Step}(\cdot)$ employs stepwise selection of variables for the logistic regression model. $\pi_{\Theta_{select}_{sub-pre}}(M^{PP})$ is the projection of the variable $\Theta_{select}_{sub-pre}$ on $M^{PP}$. $\pi_{\Theta_{select}_{sub-pre}}(M^{PP})$ is the projection of the variable $\Theta_{select}_{sub-rear}$ on $M^{PP}$. The process data of $\text{Logistic} - \text{Step}(\cdot)$ on $\pi_{\Theta_{select}_{sub-pre}}(M^{PP})$ is recorded in $\gamma_{[1,n]}^{pre}$. The process data of $\text{Logistic} - \text{Step}(\cdot)$ on $\pi_{\Theta_{select}_{sub-rear}}(M^{PP})$ is recorded in $Z_{[1,n]}^{rear}$. $\{\gamma^{pre}\} \leftarrow \min(\gamma_{[1,n]}^{pre}) \text{AIC}$ means that the set of $L_{i}$ with the smallest AIC value in $\gamma_{[1,n]}^{pre}$ is put into $\gamma_{[1,n]}^{pre}$. Steps 5-10 indicate that the set of $L_{i}$ with the smallest AIC values and significance less than 0.01 in $\gamma_{[1,n]}^{pre}$ is put into $\gamma_{[1,n]}^{rear}$. The Cartesian product of $\{\gamma^{pre}\}$ and $\{\gamma^{rear}\}$ is performed to form a buffer pool ($\text{Buffer} - \text{pool}$).

Optimization Stage (Steps 12-17 of Algorithm 3): Each element in $\text{Buffer} - \text{pool}$ constructs a logistic regression model, and the model with the highest accuracy (denoted by $\max(\Phi_{1})$ $\text{AccuracyRate}$) is selected as the optimal prediction model (denoted by $\Phi_{optimal}$).

III. RESULTS AND DISCUSSION

A. PERFORMANCE EVALUATION OF CVS$_{C}$-RF

When constructing RF in the CVS$_{C}$-RF algorithm, the number of decision trees $\tau$ in the random forest has a large impact on the performance and efficiency of the algorithm. In order to determine the optimal value $\tau_{optimal}$, three groups of experiments were carried out under the premise of $\tau \propto RF_{cv-number} (LR_{sub})$. Each group of experiments involved 10 randomized experiments for different $\tau$, and a comparative analysis was given using the lost count and stability. The calculation of the lost count is shown in (3). The loss count of the $j$-th $\tau$ is denoted $\text{Lost}(\tau_{j})$. The value of $j$ is an integer between 1 and $h$ (denoted by $Z_{[1,h]}$). $\sum_{j=1}^{10} \text{Lost}(\tau_{j})$ is the loss count of the $j$-th randomized experiment for $\tau_{j}$.

$$\text{Lost}(\tau_{j}) = \sum_{i=1}^{10} \text{Lost} (\tau_{j}) (j \in Z[1,h])$$

Stability was investigated from two aspects: internal stability and external stability.

The calculation of internal stability is shown in (4). The internal stability of $\tau_{j}$ in the top $d$ ranges is denoted $\text{Internal-stability}(\tau_{j} | pre - d)$ in (4). The union of the results for $\tau_{j}$ on 10 randomized experiments in the top $d$ ranges is denoted $\bigcup_{j=1}^{10} L_{i} (\tau_{j}^{pre-d})$. The occurrence counts of $L_{i}$ in the union are denoted $\text{count}(L_{i} | \bigcup_{j=1}^{10} L_{i} (\tau_{j}^{pre-d}))$.

$$\text{Internal-stability}(\tau_{j} | pre - d) = \sum_{i=1}^{d} \text{count}(L_{i} | \bigcup_{j=1}^{10} L_{i} (\tau_{j}^{pre-d}))$$

The calculation of external stability is shown in (5). The external stability of $\tau_{j}$ in the top $d$ ranges is denoted $\text{External-stability}(\tau_{j} | pre - d)$ in (5). The union of the results for $\tau_{j} \in [h, h]$ in the top $d$ ranges is denoted $\bigcup_{j=h}^{h} L_{i} (\tau_{j})$. The occurrence counts of $L_{i}$ in the union are denoted $\text{count}(L_{i} | \bigcup_{j=h}^{h} L_{i} (\tau_{j}))$.

$$\text{External-stability}(\tau_{j} | pre - d) = \sum_{i=1}^{d} \text{count}(L_{i} | \bigcup_{j=h}^{h} L_{i} (\tau_{j}))$$

In the first group of experiments, $RF_{cv-number} (LR_{sub})$ is set to 480, and 18 groups of data are taken in the interval $[3000, 11500] (h = 18, \bar{h} = 9)$. The experimental results are shown in Table 3 and Table 4. It can be seen from Table 3 that the lost count gradually approaches 0 from $\tau_{0} = 7000$, and there are two fluctuations of 1 loss in $\tau_{11} = 8000$ and $\tau_{12} = 8500$, and 0 loss is stable from $\tau_{13} = 9000$. It can be seen from Table 4 that from $\tau_{12} = 8500$, the internal stability of the top 60 and the top 80 both reached 100%, the top 100 reached more than 98.90%, and the top 120 reached more than 93.50%. From $\tau_{12} = 8500$, the external stability
TABLE 3. Loss count analysis of $RF_{cv-number}(Lr_{sub}) = 480$.

| $\tau$ | $h$ | serial number of 10 randomized experiments | lost count |
|-------|-----|------------------------------------------|------------|
|       |     | $T_1$ | $T_2$ | $T_3$ | $T_4$ | $T_5$ | $T_6$ | $T_7$ | $T_8$ | $T_9$ | $T_{10}$ |
| 3000  | 1   | 5     | 1     | 3     | 3     | 2     | 1     | 5     | 3     | 3     | 8       | 4       | 7       | 41       |
| 3500  | 2   | 3     | 4     | 5     | 3     | 1     | 1     | 4     | 3     | 3     | 3       | 3       | 3       | 29       |
| 4000  | 3   | 3     | 5     | 3     | 3     | 2     | 1     | 2     | 3     | 1     | 2       | 1       | 9       | 21       |
| 4500  | 4   | 1     | 1     | 1     | 1     | 1     | 1     | 1     | 1     | 1     | 1       | 1       | 1       | 1         |
| 5000  | 5   | 0     | 1     | 0     | 1     | 1     | 1     | 1     | 1     | 1     | 1       | 1       | 1       | 1         |
| 5500  | 6   | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0       | 0       | 0       | 0         |
| 6000  | 7   | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0       | 0       | 0       | 0         |
| 6500  | 8   | 0     | 1     | 0     | 2     | 2     | 0     | 0     | 0     | 1     | 0       | 0       | 0       | 0         |
| 7000  | 9   | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0       | 0       | 0       | 0         |
| 7500  | 10  | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0       | 0       | 0       | 0         |
| 8000  | 11  | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0       | 0       | 0       | 0         |
| 8500  | 12  | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0       | 0       | 0       | 0         |
| 9000  | 13  | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0       | 0       | 0       | 0         |
| 9500  | 14  | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0       | 0       | 0       | 0         |
| 10000 | 15  | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0       | 0       | 0       | 0         |
| 10500 | 16  | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0       | 0       | 0       | 0         |
| 11000 | 17  | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0       | 0       | 0       | 0         |
| 11500 | 18  | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0       | 0       | 0       | 0         |

FIGURE 4. Loss count curve of $RF_{cv-number}(Lr_{sub}) = 480$.

of the top 60 and the top 80 both reached 100%, the top 100 reached more than 99.40%, and the top 120 reached more than 95.75%. Figure 5 shows that both the internal stability and external stability had a relatively high stability trend from $\tau_{13}$. The left boundary ($\tau_{13} = 9000$) was moved two digits to the right. Finally, $\tau_{optimal}$ was set to $\tau_{15} = 10000$, and $\eta$ was set to $\left\lfloor \sqrt{480 + 0.5} \right\rfloor = 22$.

In the second group of experiments, $RF_{cv-number}(Lr_{sub})$ is set to 240, and 13 groups of data are taken in the interval [1000, 7000] ($h = 13, \bar{h} = 6$). The experimental results are shown in Table 5 and Table 6. It can be seen from Table 5 that the lost count gradually approaches 0 from $\tau_6 = 3500$, and there is a fluctuation of 1 loss in $\tau_7 = 4000$, and 0 loss is stable from $\tau_8 = 4500$.

It can be seen from Table 6 that from $\tau_6 = 3500$, the internal stability of the top 60 and the top 80 both reached 100%, the top 100 almost reached 100% (except a fluctuation of 99.80% for $\tau_{10} = 5500$), and the top 120 reached more than 95.33%. Figure 7 shows that both the internal stability and external stability had a relatively high stability trend from $\tau_{10}$ (except for a fluctuation in the top 100). The left boundary ($\tau_{8} = 4500$) was moved three digits to the right. Finally, $\tau_{optimal}$ was set to $\tau_{11} = 6000$, and $\eta$ was set to $\left\lfloor \sqrt{240 + 0.5} \right\rfloor = 15$.

In the third group of experiments, $RF_{cv-number}(Lr_{sub})$ is set to 120, and 9 groups of data are taken in the interval [500, 2500] ($h = 9, \bar{h} = 6$). The experimental results are shown in Table 7 and Table 8. It can be seen from Table 7 that 0 loss is stable from $\tau_6 = 1000$. It can be seen from Table 8 that from $\tau_6 = 1000$ the internal stability
TABLE 5. Loss count analysis of $RF_{cv-number}(L^{sub}) = 240$.

| $\tau$ | $h$ | $L_{1}$ | $L_{2}$ | $L_{3}$ | $L_{4}$ | $L_{5}$ | $L_{6}$ | $L_{7}$ | $L_{8}$ | $L_{9}$ | $L_{10}$ |
|-------|-----|--------|--------|--------|--------|--------|--------|--------|--------|--------|--------|
| 1000  | 1   | 7      | 9      | 11     | 11     | 10     | 12     | 3      | 6      | 12     | 87     |
| 1500  | 2   | 7      | 2      | 2      | 3      | 2      | 2      | 2      | 1      | 3      | 26     |
| 2000  | 3   | 4      | 1      | 3      | 0      | 1      | 1      | 2      | 1      | 2      | 17     |
| 2500  | 4   | 2      | 0      | 1      | 0      | 1      | 0      | 1      | 0      | 0      | 5      |
| 3000  | 5   | 1      | 0      | 0      | 0      | 0      | 0      | 1      | 0      | 0      | 2      |
| 3500  | 6   | 0      | 0      | 0      | 0      | 0      | 0      | 0      | 0      | 0      | 0      |
| 4000  | 7   | 0      | 0      | 0      | 0      | 0      | 0      | 0      | 0      | 0      | 1      |
| 4500  | 8   | 0      | 0      | 0      | 0      | 0      | 0      | 0      | 0      | 0      | 0      |
| 5000  | 9   | 0      | 0      | 0      | 0      | 0      | 0      | 0      | 0      | 0      | 0      |
| 5500  | 10  | 0      | 0      | 0      | 0      | 0      | 0      | 0      | 0      | 0      | 0      |
| 6000  | 11  | 0      | 0      | 0      | 0      | 0      | 0      | 0      | 0      | 0      | 0      |
| 6500  | 12  | 0      | 0      | 0      | 0      | 0      | 0      | 0      | 0      | 0      | 0      |
| 7000  | 13  | 0      | 0      | 0      | 0      | 0      | 0      | 0      | 0      | 0      | 0      |

TABLE 6. Stability analysis of $RF_{cv-number}(L^{sub}) = 240$.

| $\tau$ | internal stability(%) | external stability(%) |
|-------|------------------------|-----------------------|
|        | top 60 | top 80 | top 100 | top 120 | top 60 | top 80 | top 100 | top 120 |
| 3500   | +     | +     | +       | 95.67   | +     | +     | +       | 99.98   |
| 4000   | +     | +     | +       | 95.93   | +     | +     | +       | 96.67   |
| 4500   | +     | +     | +       | 95.67   | +     | +     | +       | 98.12   |
| 5000   | +     | +     | +       | 96.33   | +     | +     | +       | 97.19   |
| 5500   | +     | +     | 99.80   | 95.50   | +     | +     | +       | 97.98   |
| 6000   | +     | +     | +       | 96.75   | +     | +     | +       | 97.81   |
| 6500   | +     | +     | +       | 96.41   | +     | +     | +       | 97.92   |
| 7000   | +     | +     | +       | 96.83   | +     | +     | +       | 96.67   |

FIGURE 5. Stability distribution of $RF_{cv-number}(L^{sub}) = 480$.

FIGURE 6. Loss count curve of $RF_{cv-number}(L^{sub}) = 240$.

80 both reached 100%, and the top 100 reached more than 94%. Figure 9 shows that both internal and external stability had a relatively high stability trend from $\tau_8$. The left boundary ($\tau_6 = 1000$) was moved two digits to the right. Finally, $\tau_{optimal}$ was set to $\tau_8 = 2000$, and $\eta$ was set to $\lceil \sqrt{1200} + 0.5 \rceil = 11$.

Figure 4, Figure 6 and Figure 8 show the following: The lost count for $\tau_j$ was more in the early stage and decreased greatly in the middle stage, but was unstable. It decreased to 0 in the later stage and tended to be stable. Note: “+” in Tables 4, 6 and 8 represents 100.

B. PERFORMANCE EVALUATION OF CVS$_{E-CS-CF}$

The parameter settings for the CVS$_{E-CS-CF}$ algorithm are shown in Table 9. The $Set_{penalty}$ obtained by the CVS$_{E-CS-CF}$ algorithm is denoted $Set_{penalty}(i \in [1, 25])$. 

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TABLE 7. Loss count analysis of $RF_{cv-\text{number}}(Lr_{\text{sub}}) = 120$.

| $\tau$ | $h$ | $T_1$ | $T_2$ | $T_3$ | $T_4$ | $T_5$ | $T_6$ | $T_7$ | $T_8$ | $T_9$ | $T_{10}$ | Lost count |
|-------|-----|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|-----------|
| 500   | 1   | 2     | 3     | 2     | 1     | 1     | 2     | 4     | 2     | 2     | 3       | 22        |
| 600   | 2   | 2     | 0     | 0     | 1     | 0     | 0     | 1     | 1     | 0     | 2       | 7         |
| 700   | 3   | 1     | 2     | 2     | 1     | 0     | 1     | 0     | 0     | 1     | 0       | 8         |
| 800   | 4   | 0     | 0     | 0     | 0     | 1     | 1     | 1     | 1     | 1     | 0       | 5         |
| 900   | 5   | 1     | 1     | 0     | 1     | 0     | 0     | 0     | 0     | 0     | 0       | 4         |
| 1000  | 6   | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0       | 0         |
| 1500  | 7   | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0       | 0         |
| 2000  | 8   | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0       | 0         |
| 2500  | 9   | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0     | 0       | 0         |

FIGURE 7. Stability distribution of $RF_{cv-\text{number}}(Lr_{\text{sub}}) = 240$.

FIGURE 8. Loss count curve of $RF_{cv-\text{number}}(Lr_{\text{sub}}) = 120$.

(As shown in Table 10). The location of $Set_{\text{penalty}}^i$ in $\Theta$ is denoted $\text{position}(Set_{\text{penalty}}^i)^\Theta$. The relative position coefficient of $Set_{\text{penalty}}^i (RPC(Set_{\text{penalty}}^i))$ is calculated via (6). The $Set_{\text{shock}}$ containing 30 elements is shown in Table 11.

$$RPC(Set_{\text{penalty}}^i) = \frac{\text{position}(Set_{\text{penalty}}^i)^\Theta}{\Delta} \quad (6)$$

$RPC$ is a value between 0 and 1. The smaller it is, the more important the $Lr_i$ is. This indicates that the location of $Set_{\text{penalty}}^i$ in $\Theta$ is at the front. The larger the $RPC$ value, the less important the $Lr_i$ is. This indicates that the location of $Set_{\text{penalty}}^i$ in $\Theta$ is later in the queue. If $RPC(Set_{\text{penalty}}^i)$ is larger, $Set_{\text{penalty}}^i$ will be punished. It indicates that the algorithm could protect those $Lr_i$ with higher importance, so the algorithm has better stability. According to Figure 10, 96% of $RPC(Set_{\text{penalty}}^i)$ in $Set_{\text{penalty}}$ are above 0.49 (except 0.44), the mean of which is 0.79. This indicates that the top 60 $Lr_i$ in $\Theta$ are stably protected and the algorithm has good stability. Ultimately, $\lambda$ lncRNAs most closely related to prostate cancer are selected by the CVSSe-CS-CF algorithm from 480 lncRNAs ($\lambda = 95$). The results are shown in Table 12.

C. PERFORMANCE EVALUATION OF CSPA-PL

After implementing Logistic − Step($\pi_{\text{select}} M^{PP}$) in the inspection stage of the CSPA-PL algorithm, a total
TABLE 8. Stability analysis of $RF_{cv}$-number ($L_{\text{sub}}$) = 120.

| $\tau$ | internal stability(%) | external stability(%) |
|--------|------------------------|----------------------|
|        | top 60 | top 80 | top 100 | top 60 | top 80 | top 100 |
| 1000   | +      | 98.25  | 92.70   | +      | 99.06  | 94.50   |
| 1500   | +      | 98.88  | 92.60   | +      | 99.69  | 94.00   |
| 2000   | +      | 99.38  | 93.50   | +      | 95.00  |
| 2500   | +      | 99.88  | 94.00   | +      | 95.50  |

TABLE 9. Parameter setting of CVS e-CS-CF algorithm.

$\begin{array}{cccccccccc}
L_r & \tau_0 & \eta_0 & \tau_1 & \epsilon_1 & \tau_2 & \epsilon_2 & \Delta & \Omega & a & b & f \\
\hline
M^{CF} & 10000 & 22 & 6000 & 15 & 2000 & 11 & 120 & 60 & 1 & 121 & 0
\end{array}$

TABLE 10. Result of $Set_{penalty}$.

| $L_r$ | gene name | $L_r$ | gene name | $L_r$ | gene name |
|-------|------------|-------|------------|-------|-----------|
| 41    | FAM201A    | 99    | MATN1-AS1  | 193   | TSPAN10   |
| 43    | RP11-262H14.1 | 101  | RP11-244F12.3 | 204   | RP11-696N14.1 |
| 45    | RP11-713P17.3 | 104  | LINC0476   | 236   | AC139666.1 |
| 61    | AC093627.10 | 112  | CTC-308K20.1 | 239   | RP11-834C11.3 |
| 66    | CTC-504A5.1 | 119  | POLR2J4    | 251   | RP11-558F24.4 |
| 67    | CTC-774J1.2 | 120  | AMZ2P1     | 331   | MSRI      |
| 73    | RP11-573H1.2 | 122  | AM2-AS1    | 382   | RP11-446H18.3 |
| 90    | RP11-206L10.11 | 151  | LINC00086  |
| 97    | LINC00338  | 169  | CTBP1-AS1  |

TABLE 11. Result of $Set_{shock}$.

| $L_r$ | gene name | $L_r$ | gene name | $L_r$ | gene name |
|-------|------------|-------|------------|-------|-----------|
| 19    | AC030090.1 | 135   | KB-431C1.4 | 223   | PVT1      | 272   | PRKY     |
| 34    | RPL13P5    | 160   | UHRF1      | 225   | SNHG15    | 310   | ZNFX1-AS1 |
| 63    | RNF126P1   | 170   | RP11-72M17.1 | 230    | AP001372.2 | 369   | MTMR9LP  |
| 70    | TTTY15     | 185   | KTN1-AS1   | 234   | SNHG7     | 383   | RP11-488C13.5 |
| 87    | RHPN1-AS1  | 197   | RP11-29402.2 | 243    | NDUFB2-AS1 | 395   | LINC00402 |
| 102   | RPS-916L1.7 | 211   | DLX6-AS1   | 253   | CTD-2185M18.1 | 408  | MLLT4-AS1 |
| 114   | AC002505.4 | 214   | RP4-65936.2 | 258   | WHAMMP2   |
| 118   | ATG9B      | 219   | RP11-500G10.1 | 267   | FLNB-AS1  |

FIGURE 10. Comparison of Relative position coefficient distribution on $Set_{penalty}$.

of 21 groups ($Z_{pre}^{[1,21]}$) were obtained. The AIC distribution of $Z_{pre}^{[1,21]}$ is shown in Figure 11. This shows that the value of $Z_{pre}$ is the minimum (157.66). So, $[\tau_{pre}]=\{L_r(1)\}$ is obtained. After implementing Logistic $\rightarrow$ Step$\left(\tau_{select}(M^{PP})\right)$, a total of 39 groups ($Z_{pre}^{[1,39]}$) were obtained. In this, the value of $Z_{pre}^{[1,39]}$ is the minimum (172.29). There are three $L_r$s in $Z_{pre}^{[1,39]}$, the P-values of which were less than 0.01. So, $[\tau_{pre}]=\{L_r(1)\}$ is obtained. The values of $i(1)$ and $i(2)$ in the above $L_r(1)$ and $L_r(2)$ are shown in Table 13. There were three alternative sets in Buffer $\rightarrow$ pool, which were $\text{Buffer} \rightarrow \text{pool}_1 = \{\{\tau_{pre}\} \cup L_r(1)\}$, $\text{Buffer} \rightarrow \text{pool}_2 = \{\{\tau_{pre}\} \cup L_r(79)\}$ and $\text{Buffer} \rightarrow \text{pool}_3 = \{\{\tau_{pre}\} \cup L_r(110)\}$. Next, $\text{Buffer} \rightarrow \text{pool}_s$ was obtained by executing Logistic $\rightarrow$ Step$\left(\tau_{select}(M^{PP})\right)$ on $\text{Buffer} \rightarrow \text{pool}_i$, which produced $\text{Buffer} \rightarrow \text{pool}_s = \{\{\tau_{pre}\} \cup L_r(1)\}$, $\text{Buffer} \rightarrow \text{pool}_s = \{\{\tau_{pre}\} \cup L_r(11)\}$ and $\text{Buffer} \rightarrow \text{pool}_s = \{\{\tau_{pre}\} \cup L_r(79)\}$. The values of $j(1)$, $j(2)$ and $j(3)$ in the above $L_r(1)$, $L_r(11)$, and $L_r(79)$ are shown in Table 14. The accuracy comparison experiment for $\text{Buffer} \rightarrow \text{pool}_s$ for the logistic regression model shows that the accuracy rate of $\text{Buffer} \rightarrow \text{pool}_s = \{\{\tau_{pre}\} \cup L_r(79)\}$ is the highest. Finally, the optimal logistic regression model for tumor clinical stage (denoted by $\Phi_{optimal}$) is obtained, which contained 22 $L_r$s (as shown in Table 15).
TABLE 12. Result of $\lambda_{select}$ ($\lambda = 95$).

| $\lambda$ | $L_{r_1}$ | gene name   | $\lambda$ | $L_{r_1}$ | gene name   | $\lambda$ | $L_{r_1}$ | gene name   |
|-------|---------|-------------|-------|---------|-------------|-------|---------|-------------|
| 1     | 6       | LINC00665   | 25     | 62      | SRD5A2      | 49     | 79      | NEURL3      |
| 2     | 3       | RP11-342C23.4 | 26     | 9       | TAM222A-AS1 | 50     | 46      | RPS-1121A15.1 |
| 3     | 7       | RP11-368H7.2 | 27     | 27      | BOLA3-AS1  | 51     | 55      | RP1-166D19.1  |
| 4     | 30      | CTD-31962G.2 | 28     | 29      | ACT3526   | 52     | 140     | DANCR       |
| 5     | 20      | AP001626.1  | 29     | 25      | AC073543.13 | 53    | 186     | RP1-18920.4  |
| 6     | 15      | RP11-429716.7 | 30    | 21      | LINC00261  | 54     | 348     | AP000662.4  |
| 7     | 28      | CTG-497E21.4 | 31     | 23      | RP1-401F24.4 | 55    | 246     | LGA05842.1  |
| 8     | 12      | CTD-252T21.11 | 32    | 26      | MAGI2-AS3  | 56     | 54      | RP1-379F4.4  |
| 9     | 53      | RP3-467K16.2 | 33     | 47      | RP1-316P17.2 | 57    | 33      | BPX19-AS1  |
| 10    | 14      | RP1-279F6.1  | 34     | 5       | RP1-231P20.2 | 58    | 32      | BX82465.0  |
| 11    | 64      | SNHG16      | 35     | 86      | RP1-108M4.9 | 59     | 121     | HX22-AS2A  |
| 12    | 8       | MSL3P1      | 36     | 98      | PGM5-AS1   | 60     | 139     | MR222HG    |
| 13    | 51      | LINC00087   | 37     | 82      | ALO7862.14 | 61     | 116     | CD27-AS1   |
| 14    | 10      | PCA3        | 38     | 22      | AC073133.1 | 62     | 115     | C107076-S1  |
| 15    | 65      | LINC00992   | 39     | 137     | RP1-66B24.4 | 63    | 36      | ADAMTS9-AS2 |
| 16    | 40      | ADAMTS9-AS1 | 40     | 50      | SNHG3      | 64     | 38      | DNM1B-AS1  |
| 17    | 42      | RP4-647C14.2 | 41     | 17      | ERVH48.1   | 65     | 74      | WW2C-AS2   |
| 18    | 4       | LINC00675   | 42     | 105     | RP11-168G6.1 | 66    | 48      | LHA-F-AS1  |
| 19    | 198     | MIR203HG    | 43     | 156     | XBBAC-B135H16.15 | 67   | 59      | RP11-57A19.2 |
| 20    | 13      | RP11-627G23.1 | 44    | 31      | RP1-324H6.5 | 68     | 58      | RP1-17A19.1 |
| 21    | 16      | RP11-279F6.1 | 45     | 56      | SENP3-EFA4A | 69    | 117     | NPY6R      |
| 22    | 52      | MIR31HG     | 46     | 75      | RP1-24M17.7 | 70     | 427     | AC058791.2 |
| 23    | 84      | LINC00085   | 47     | 166     | LINC00605  | 71     | 72      | RP11-265M18.2 |
| 24    | 35      | RP11-412D9.4 | 48     | 91      | SERTAD4-AS1 | 72    | 106     | ZNF300P1   |

FIGURE 11. Contrastive distribution on AIC value of $Z_{pre}$ $\{1,21\}$.

TABLE 13. The $i(1)$ value of $L_{r_{(1)}}$ in $\{\gamma_{pre}\}$, the $i(2)$ value of $L_{r_{(2)}}$ in $\{\gamma_{rear}\}$.

| $L_{r_{(1)}}$ value of $i$ | $L_{r_{(2)}}$ value of $i$ |
|--------------------------|--------------------------|
| $L_{r_{(1)}}$ | 6,15,12,14,64,8,40,42,4,16,62,27,29,25,21,47,82,137,50, | 17,166,162 |
| $L_{r_{(2)}}$ | 6,15,12,14,64,8,40,42,4,16,62,27,29,25,21,47,137,50, | 17,166,79 |
| $L_{r_{(3)}}$ | 6,15,12,53,14,64,8,10,40,42,14,35,62,27,29,25,23,2 |
|                      | 6,47,5,82,137,50,17,166 |

It can be seen that the CSPA-PL algorithm could further select 22 $L_{r_{(1)}}$s from the 95 $L_{r_{(1)}}$s most closely related to prostate cancer, which are most closely related to the tumor clinical stage of prostate cancer.

In order to verify the universality of the work presented in this paper, we also chose the lung cancer data set in the lncRNA database and TCGA database for experimentation. A total of 290 samples were obtained, including 46 normal samples and 245 cancer samples. First, the importance of the characteristic variable was calculated using the CVS gC-RF algorithm. Second, 120 IncRNAs which were most closely related to prostate cancer, were calculated from the 480 alternative IncRNAs by CASO and CVS e-CS-CF. On the basis of the above 120 IncRNAs, the CSPA-PL algorithm was adopted to further select 16 IncRNAs that were most closely related to the tumor clinical stage of lung cancer. Finally, 16 IncRNAs were used to construct a logistic regression prediction model.

D. PREDICTION RESULT

Three state-of-the-art methods (MlrLDAcp [26], REP-Tree [27], NaïveBayes [28]) were selected to compare with CSPA-PL by 10-fold cross validation. The comparison experiments were carried out from three aspects: ROC area, prediction accuracy and recall rate. The results of ROC area for prostate cancer are shown in Figure 12. The mean ROC area of three compared methods for prostate cancer was 0.673, and the ROC area of the CSPA-PL method in this paper was 0.857, which was the largest and 1.27 times that of the other methods. The results of ROC area for lung cancer are shown in Figure 13. The mean ROC area of the compared methods
TABLE 15. Result of $L_r$ in $\Phi_{\text{optimal}}$

| $L_r$ | gene name    | $L_r$ | gene name    | $L_r$ | gene name    |
|-------|--------------|-------|--------------|-------|--------------|
| x6    | LINC00665    | x4    | LINC00675    | x47   | RP11-316P17.2|
| x15   | RP11-429J17.6| x16   | RP11-279F6.1 | x137  | RP11-66B24.4 |
| x12   | CTD-2527121.1| x35   | RP11-412D9.4 | x50   | SNHG3        |
| x14   | RP11-279F6.1 | x62   | SRD5A2       | x17   | ERVH48.1     |
| x64   | SNHG16       | x27   | BOLA3-AS1    | x166  | LINC00665    |
| x8    | MSL3P1       | x29   | C1orf126     | x79   | NEURL3       |
| x40   | ADAMTS9-AS1  | x25   | AC073343.13  |       |              |
| x42   | RP4-647C14.2 | x21   | RP11-279F6.1 |       |              |

**FIGURE 12.** ROC area comparison on prostate cancer.

**FIGURE 13.** ROC area comparison on lung cancer.

**FIGURE 14.** AV-PR comparison.

for lung cancer was 0.666, and the ROC area of the CSPA-PL method in this paper was 0.842, which was the largest and 1.26 times that of the other methods. In the comparison experiment for recall rate, the accuracy and recall rate were often mutually restricted and offset each other. Therefore, AV-PR was implemented and used in this paper as the mean of the prediction accuracy and recall rate. The results for AV-PR are shown in Figure 14 (The prostate cancer is shown by blue, the lung cancer is shown by red.). The mean AV-PR of the compared methods for prostate cancer was 0.723, and the AV-PR of CSPA-PL with the maximum was 0.889, which was about 1.231 times that of the other methods. The mean AV-PR of the compared methods for lung cancer was 0.784, and the AV-PR of CSPA-PL with the maximum was 0.896, which was about 1.142 times that of the other methods. These results indicate that the accuracy rate and ROC area for CSPA-PL were both good.

**IV. CONCLUSION AND DISCUSSION**

Although some methods have been applied to lncRNA-disease association prediction, clinical prognostic data were rarely involved. In this study, we constructed a clinical stage prediction algorithm for cancer-associated lncRNA (CSPA-PL), which utilized cancer clinical stage data. CSPA-PL was based on unknown human lncRNA-disease associations combining with the clinical stage data. The core modules of CSPA-PL included CASO, the CVSE-RF algorithm, and the CVSE-CS-CF algorithm. For CASO, a learning mode was formed in which the first quartile area was a defensive area and the other quartile areas were offensive. Three symmetric ideas were adopted, which were center allelism,
X-axis allelism and Y-axis allelism. The CVS₂-RF algorithm employed a variable selection algorithm based on random forests as the core to calculate the importance of the characteristic variables. This method exhibited good robustness. Experimental results showed that the proposed method in this study has good predictive performance.

The value of this model lies in the following:
(a) It provides a strong research foundation for the prediction of prognosis information for cancer patients by lncRNA-disease association.
(b) The simplified lncRNAs in the model were the closest to predicting the relationship between lncRNA and the disease, which provides a favorable research premise for subsequent studies of this association.

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