PLIT: An alignment-free computational tool for identification of long non-coding RNAs in plant transcriptomic datasets

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Abstract

Long non-coding RNAs (lncRNAs) are a class of non-coding RNAs which play a significant role in several biological processes. RNA-seq based transcriptome sequencing has been extensively used for identification of lncRNAs. However, accurate identification of lncRNAs in RNA-seq datasets is crucial for exploring their characteristic functions in the genome as most coding potential computation (CPC) tools fail to accurately identify them in transcriptomic data. Well-known CPC tools such as CPC2, IncScore, CPAT are primarily designed for prediction of IncRNAs based on the GENCODE, NONCODE and CANTATAdb databases. The prediction accuracy of these tools often drops when tested on transcriptomic datasets. This leads to higher false positive results and inaccuracy in the function annotation process. In this study, we present a novel tool, PLIT, for the identification of lncRNAs in plants RNA-seq datasets. PLIT implements a feature selection method based on L_1 regularization and iterative Random Forests (iRF) classification for selection of optimal features. Based on sequence and codon-bias features, it classifies the RNA-seq derived FASTA sequences into coding or long non-coding transcripts. Using L_1 regularization, 31 optimal features were obtained based on lncRNA and protein-coding transcripts from 8 plant species. The performance of the tool was evaluated on 7 plant RNA-seq datasets using 10-fold cross-validation. The analysis exhibited supe-

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rior accuracy when evaluated against currently available state-of-the-art CPC tools. *Keywords:* lncRNA; LASSO; iterative Random Forests; Random Forests; RNA-seq; Ensembl Plants; CANTATAdb

Recent advances in genome sequencing have led to the discovery of thousands

1. Introduction

of non-coding RNA transcripts. Using RNA Sequencing (RNA-Seq) and epigenome sequencing, a new class of RNA transcripts i.e. long non-coding RNAs (lncRNAs) is defined as those having transcript length > 200 nucleotides. Although this class of RNA lacks protein-coding ability, they have been found involved in the regulation of biological processes such as enzymatic activity regulation, genomic loci imprinting, transcription, translation, and cellular differentiation [1]. Several lncRNA databases such as GENCODE, NONCODE and CANTATAdb have been developed for storage of lncRNAs [2, 3, 4]. These databases provide valuable resources for further identification of novel lncRNAs from genomic sequences. Although many lncRNAs have been identified in plants and animals, accurate computational identification of these lncRNAs in RNA-seq datasets remains one of the major problems in plants.

Therefore, an efficient, accurate and robust computational algorithm is required to predict lncRNAs in plants to further investigate their potential roles. Computational prediction of lncRNAs has been viable for the past few years. These methods generally use machine learning approaches to classify RNAs into different classes. Several tools have been developed including: Coding Potential Calculator 2 (CPC2) [5], Coding-Non-Coding Index (CNCI) [6], Coding Potential Assessment Tool (CPAT) [7], and a predictor of long non-coding RNAs and messenger RNAs based on improved k-mer scheme (PLEK) [8]. CPC2 computes the coding probability of the sequence by computing its peptide length, isoelectric point, Fickett score [9] and ORF integrity. CPC2 employed SVM using RBF kernel for training 17984 protein-coding and 10452 non-coding transcripts from Refseq [10], Ensembl (v87), and EnsemblPlants (v32) databases [11]. Tools such as CPAT and IncScore [7, 12] classified protein-coding and non-coding transcripts based on logistic regression model

as machine learning classifier using sequence-based features such as open-reading frame (ORF) size, ORF length, ORF coverage, GC content, Fickett score and hexamer score; whereas others such as CNCI and LncRNA-MFDL [13] classified lncRNAs us-

- ing adjoining nucleotide triplets (ANT) features to identify most-like CDS (MLCDS) regions in each transcript. PLEK [8] uses calibrated k-mer frequencies of a sequence and sliding-window approach as features for classification using SVM classifier from LIBSVM package. Currently developed alignment-free tools such as CPC2, lncScore, CPAT and PLEK work well with FASTA sequences derived from the GENCODE, NON-
- CODE or CANTATAdb databases, but perform poorly on FASTA sequences derived from RNA-seq data. Thus, an accurate tool is required for prediction of lncRNAs in plants.

In this work, we have developed a new alignment-free tool named Plant LncRNA and Identification Tool (PLIT) which uses L_1 regularization for feature selection and a Random Forest classifier for classification of sequences. For lncRNA identification, PLIT implements 73 sequence and codon-bias based features. The framework implements an optimization module called LASSO iterative Random Forest-Feature Selection (LiRFFS) [14, 15] which selects an optimal feature set from training and validation set features. The selected feature set can be used for identification of lncRNAs directly from RNA-seq derived FASTA sequences. The optimal features were selected based on coding and long non-coding FASTA sequences from the Refseq database [10]. The prediction accuracy of the PLIT was benchmarked against other existing tools. In total, 31 features which included ORF length, ORF coverage, Hexamer Score, GC content, and codon-bias features such as Codon Usage Bias, Relative Codon Bias, and Relative Synonymous Codon Usage were selected. Compared with the existing tools, PLIT exhibited 15-30% increase in the prediction accuracy when evaluated with 10-fold Cross Validation and repeated 10-fold Cross Validation with data shuffling on different plant RNA-seq datasets. The availability of the RNA-seq based plant lncRNA prediction tool will provide a useful resource

for identification of novel lncRNAs in plants. PLIT is freely available on GitHub: https://github.com/deshpan4/PLIT.

2. Methods

2.1. Data description

For extracting optimal feature set from FASTA sequences, a random selection of protein-coding and lncRNA transcript sequences from eight plant species was obtained from Refseq Release 91 [10]. Transcript sequences for *Arabidopsis thaliana*, *Brassica rapa*, *Brassica napus*, *Brassica oleracea*, *Zea mays*, *Oryza sativa*, *Solanum tuberosum* and *Solanum lycopersicum* were downloaded. lncRNA sequences were filtered by applying a threshold cutoff of 200bp on non-coding RNA (ncRNA) FASTA files.

For performance evaluation, the lncRNA genomic coordinates for seven plant species (*Arabidopsis Thaliana, Glycine Max, Oryza Sativa, Solanum Lycopersicum, Sorghum Bicolor, Vitis Vinifera,* and *Zea Mays*) were downloaded from CANTATAdb v2.0 [4] as negative examples, and protein-coding coordinates were downloaded from Ensembl Plants Release 41 [11] as positive examples. The RNA-seq data for the seven plant species were obtained from the NCBI SRA database [16] with accession numbers PRJNA268115, PRJNA237837, PRJNA293380, PRJNA478448, PRJNA318972, PRJNA356948 and PRJNA484195. A description of the total number of lncRNA transcript sequences and number of annotated sequences has been provided in Table

Table 1: Summary of IncRNAs used in plant RNA-Seq datasets.

Data set	lncRNAs used	lncRNAs annotated	Read length	Coverage
A. thaliana	4373	1027	50-51	23x
G. max	2819	612	178-202	4x
O. sativa	2759	265	250	12x
S. lycopersicum	4308	549	250	6x
S. bicolor	2456	189	202	4x
V. vinifera	4019	1292	105	7x
Z. mays	10761	1029	202	1x

2.2. Data preprocessing

1.

The first 15 base pairs of the sequence reads are trimmed using Cutadapt [17] to remove adapter and low-quality sequences with Q-score less than or equal to

- 30. For *A. thaliana* reads, trimmed reads are aligned to the TAIR10 genome using the Tophat2 mapper [18] with custom parameter values (minimum intron length = 40, maximum intron length = 5000, segment length = 20, segment mismatches = 1, max multihits = 1). The trimmed sequence reads of *G. Max* were mapped to Glycine_Max_v2.0 genome with custom parameter values (minimum intron length = 30, maximum intron length = 15000, segment mismatches = 1, max multihits = 1).
- Trimmed reads for *Oryza Sativa* L. ssp. Japonica were mapped to IRGSP-1.0 genome with min intron length of 20, max intron length of 15000, segment mismatches of 1 and max multihits of 1. For *S. Lycopersicum*, the trimmed reads were aligned to the tomato genome (SL2.50) with min anchor length more than 8 nt, segment mismatches = 1 and max multihits = 1. The trimmed reads for *Sorghum Bicolor* were
- mapped to the Sorgum genome (Sorghum_Bicolor_NCBIv3) and trimmed reads of *Vitis Vinifera* were aligned to grape genome (IGGP_12x) with segment mismatches = 1 and max multihits = 1. For *Zea Mays*, the trimmed reads were aligned to the Maize genome (AGPv4) with the custom parameter values: min intron length = 5, max intron length = 60000, segment length = 25, segment mismatch = 1 and max multihits
- = 1. Remaining parameters were kept to default.

2.3. Feature extraction

For extraction of features from the RNA-Seq derived genomic sequences, the transcript sequences were first extracted from the Binary Alignment Map (BAM) file produced by the Tophat2 mapper [18]. Based on reference alignment of sample reads, a consensus FASTA sequence for each transcript coordinate was constructed by a two-step process: (1) SNP and INDEL calling of BAM file using SAM-tools mpileup [19] that generated a Variant Call Format (VCF) file, and (2) sequence extraction from the genome and consensus sequence generation using variants from VCF by the SAMtools *faidx* tool [19]. The mpileup function collects the information from the BAM file and computes the likelihood. This is stored in a Binary VCF (BCF) format. The Bcftools consensus function creates a consensus FASTA transcript sequence based on reference genome by applying the VCF variants. The sequence obtained can be used for extraction of features for lncRNA classification.

To construct a random forest model, 73 ORF-based and codon-bias features were extracted for each sequence in a given dataset. The features were selected based on the published results on sequence measures and codon bias measures [20, 21]. Features extracted from FASTA sequences can be categorized into either ORF-based features or codon bias features. These features constitute a feature set $F = f_1, f_2, f_3, \ldots, f_{73}$, where f_i denotes the i^{th} feature. The features are discussed below.

115 2.3.1. ORF and Sequence based features

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We extracted three ORF-based features: maximum ORF length (f_1) , ORF coverage (f_2) and mean ORF coverage (f_3) and four sequence-based features: transcript length (f_4) , GC content (f_5) , Fickett score (f_6) and Hexamer score (f_7) .

- 1. ORF length (f_1) : f_1 is one of the most fundamental features used to distinguish lncRNA from mRNA as majority of protein-coding genes have ORFs greater than 100 amino acids [22].
- 2. ORF coverage (f_2): f_2 is the ORF coverage defined as length of the longest ORF divided by transcript length. This feature has also been shown to produce good classification performance when compared to ORF length [7, 12].
- 3. Overall ORF coverage (f_3) : f_3 is the overall ORF coverage defined as the average of total ORF lengths divided by transcript length for the sequence.
 - 4. Transcript length (f_4) : f_4 is the total length of each transcript sequence.
 - 5. GC content (f_5): f_5 is the GC content, which is also a common measure to differentiate lncRNA from protein-coding transcripts, as coding sequences have been reported to have higher GC content in exons over introns [23]. The GC content was calculated by counting the frequency of GC motifs for each sequence.
 - 6. Fickett score (f_6): f_6 is the Fickett score [9] which was obtained by calculating four base pair position values in the transcript sequence. f_6 is calculated as follows. Let

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A_1 = Number of A's in positions 1, 4, 7, 10, ...,
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 A_2 = Number of A's in positions 2, 5, 8, 11, ..., and

 A_3 = Number of A's in positions 3, 6, 9, 12,

Then $A_{position}$ is defined as,

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$$A_{position} = \frac{MAX(A_1, A_2, A_3)}{MIN(A_1, A_2, A_3) + 1} \tag{1}$$

and $T_{position}$, $G_{position}$ and $G_{position}$ are calculated similarly. In a similar manner, $A_{content}$, $T_{content}$, $G_{content}$ and $G_{content}$ of the sequence are determined by calculating percentage composition of each base in the sequence. These eight values are then converted to a probability value (p) using a lookup table [9] and multiplied by a weight (w) for each base. The Fickett score f_6 is then determined as:

$$f_6 = \sum_{i=1}^8 p_i w_i. (2)$$

7. Hexamer score (f_7) : f_7 is the Hexamer score which is computed by making a hexamer table of 4096 (64×64) k-mers using a reference set of coding and non-coding sequences. The Hexamer score is calculated by first measuring frequencies of hexamers in the test set sequences. The logarithmic ratio of coding and non-coding sequences is then computed for each hexamer having non-zero frequency in the test set. Positive f_7 indicates higher probability of protein-coding sequence whereas a negative score indicates a higher probability of non-coding RNA sequence. The in-frame hexamer frequency of protein-coding sequences is given by $F(h_i)$ where i = 1, 2, ..., 4096 and inframe hexamer frequency of lncRNA sequences is given by $F'(h_i)$ where i = 1, 2, ..., 4096. Therefore, for each hexamer sequence, $H_1, H_1, H_1, ..., H_m$, where m is observed in the test sequence. f_7 is given by:

$$f_7 = \frac{1}{m} \sum_{i=1}^{m} \log \left(\frac{F(h_i)}{F'(h_i)} \right)$$
 (3)

2.3.2. Codon Bias features

In protein-coding genes, the translational mapping process of codons (or nucleotide triplets) to amino acids involve usage of synonymous codons which code the same amino acids that are non-distinguishable at protein level. However, it has been reported that there exists a non-uniform codon usage in most genes i.e. codon bias [24, 25]. Many indices have been proposed for measuring codon bias, among which we carefully selected six codon-bias measures which are important in distinguishing lncRNAs from mRNAs.

1. Frequency of the optimal codons (f_8): This feature is calculated as ratio of the total number of optimal codons to the total number of synonymous codons. The Frequency of the optimal codons (Fop) was also one of the measures proposed by Ikemura [25]. The number of codons of optimal codons is calculated as: $O_{opt} = \sum_{c \in C_{opt}} O_c$, where C_{opt} is defined as subset of optimal codons from all codons C and O_{tot} is the total number of codons in the sequence. Therefore, f_8 is calculated as: $f_8 = \frac{O_{opt}}{O_{tot}}$.

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- 2. Codon Usage Bias (f_9) : The Codon Usage Bias (CUB) which assesses codon bias in test set sequence relative to reference set of sequences based on weighted sum of distances of relative codon usage frequencies between the reference set and test set sequences [26]. The reference set is used as standard to which other sequences can be compared. f_9 is defined as: $f_9 = \sum_{a \in A} F_a d(f_a, f_a^{ref})$, where F_a is frequency of amino acid a in the test set sequence; f_a and f_a^{ref} are codon frequencies for amino acid a in test and reference sets, respectively; and f_a^{ref} and f_a^{ref} . These are calculated as: f_a^{ref} is the frequency of codon f_a^{ref} and f_a^{ref} is the frequency of amino acid f_a^{ref} in test set sequences and f_a^{ref} is the frequency of amino acid f_a^{ref} in reference set sequences.
- 3. Relative Codon Bias (f_{10}) : The Relative Codon Bias (RCB) [27] is a measure that defines the contribution of codons as: $w_c^{RCB} = \frac{(O_c E[O_c])}{E[O_c]}$, where $E[O_c]$ is the expected number of codon occurrences in three codon positions. Once w_c^{RCB} is determined, f_{10} is calculated by the following formula for each sequence: $f_{10} = exp\left(\frac{1}{O_{tot}}\sum_{C \in C}\log w_c^{RCB}\right) 1.$
- 4. Weighted sum of relative entropy (f_{11}) : This measures the degree of deviation from equal codon usage [28]. Therefore, f_{11} is defined as sum of relative

entropy of each amino acid weighted by its relative frequency in the test sequence which is given by: $f_{11} = \sum_{a \in A} F_a E_a$. Here F_a is the relative frequency of amino acid a in the test sequence and E_a is computed as: $E_a = \left(\frac{H_a}{\log_2 k_a}\right)$, where k_a number of synonymous codons observed in the test sequence and H_a is the entropy which measures uncertainty of codon usage in the test sequence for amino acid a computed as: $H_a = \sum_{c \in C_a} f_{ac} \log_2 f_{ac}$.

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- 5. Synonymous Codon Usage Order (f_{12}): This is also an entropy-based codon bias measure and is similar to f_{11} which differs only by the way entropy is calculated for each amino acid [29]. Instead of calculating the relative entropy, the normalized difference between maximum and observed entropy is computed as $E_a = \frac{\log_2 k_a H_a}{\log_2 k_a}$, and then f_{12} is computed as $f_{12} = \sum_{a \in A} F_a E_a$.
- 6. Relative Synonymous Codon Usage (RSCU): This measure defines the relationship between observed codon frequencies and the number of times codon is observed when synonymous codon usage is random with no codon bias [30]. This is calculated as: $RSCU_{ac} = \frac{O_{ac}}{\frac{1}{k_a}\sum_{c\in C_a}O_{ac}}$ where O_{ac} is the frequency of codon c for amino acid c and is computed for 61 codons individually by the above equation. Methionine (M), Tryptophan (W) and stop codons were excluded from the analysis as M and W do not have any synonymous codons and stop codons do not contribute any information. Therefore, in total RSCU provided 61 features for the classification: f_{13}, \ldots, f_{73}

The random forest model for unified 6-plant species was constructed by building a training set of 22471 balanced coding (positive) and non-coding (negative) sequences, whereas the test set sequences consisted of 7529 sequences. The training and test set sequences in plant RNA-Seq datasets were divided into a proportion containing 75% and 25% of the total coding and non-coding sequences. Training and test datasets were constructed as randomized set of 8962 (Training) and 994 (Test) sequences for *A. thaliana*, 5050 (Training) and 562 (Test) sequences for *G. max*, 4968 (Training) and 550 (Test) sequences for *O. sativa*, 7756 (Training) and 860 (Test) sequences for *S. lycopersicum*, 4422 (Training) and 490 (Test) sequences for *S.*

bicolor, 7236 (Training) and 802 (Test) sequences for *V. vinifera*, 16902 (Training) and 1878 (Test) sequences for *Z. mays*. For obtaining an optimal set of features, a unified set of 6 plant species, namely, *A. thaliana*, *Z. mays*, *O. sativa*, *B. napus*, *B. rapa* and *B. oleracea*, has been constructed. The dataset has been divided into training and validation sets which were used by LiRFFS algorithm for obtaining an optimal feature set.

2.4. Feature selection

The selection of optimal features is an important optimization for classification. Wrapper-based Feature Selection (FS) methods such as Sequential Forward Selection (SFS) [31] or SVM-recursive feature elimination (SVM-RFE) [32] are computationally inefficient and can fail to identify optimal feature subsets. Whereas filterbased FS methods such as mRMR [33], Chi-square [34] and Information Gain [35], assign relevance score or rank to each feature by considering each feature separately and ignoring dependencies between features which leads to worse classification performance. Regression based approaches, such as least-squares estimate methods, often produce larger variance during model fitting which leads to over-fitting and poor generalization. Least Absolute Shrinkage and Selection Operator (LASSO) is a feature selection method which combines least-square loss with L1 norm constraint and produces sparse features by shrinking coefficients to zero. Other approaches such as ridge regression [14, 36] use L2 norm due to which it produces non-zero coefficients and therefore becomes inefficient for feature selection. Usage of Lq norm (with q < 1 or q > 1) approaches for optimization are generally nonconvex and make the minimization computationally challenging.

The PLIT framework implements LASSO and an iterative Random Forest Feature Selection (LiRFFS) algorithm (Algorithm 1) for identifying the principal set of collective features yielding the highest accuracy. It works by iterative selection of features based on varying the value of LASSO parameter lambda (λ) (Figure 1).

As λ changes, non-zero beta coefficients are generated which corresponds to the selection of features using L1-regularized optimization of LASSO [14]. The β coefficients are calculated on training set features for each λ using the following

equation:

$$\beta^{LASSO} = \underset{r}{\operatorname{argmin}} \frac{1}{2n} \| X\beta - y \|_{2}^{2} + \lambda \| \beta \|_{1}$$
 (4)

where, $\lambda \geq 0$, $\|X\beta - y\|_2^2$ is the loss function (i.e. sum of squares), $\|\beta\|_1$ is the penalty term and λ is the tuning parameter which controls the strength of the penalty. Features extracted from the coding and noncoding sequences are divided into training and validation sets. β coefficients are calculated on each λ value. The selected features for each λ are iteratively applied on the validation set to obtain the accuracy vector. The optimal feature set is obtained by selecting the feature set that produces the prediction accuracy between the tolerance accuracy value and the maximum prediction accuracy value. The optimal feature set can be used for building the model for classification of test set transcript sequences.

The algorithm (Algorithm 1) takes λ_{lower} , λ_{upper} and $\lambda_{step-size}$ as lower limit, upper limit and step-size input values for creation of λ values. The default values of λ_{lower} , λ_{upper} and $\lambda_{step-size}$ are chosen as 0.00001, 0.1 and 0.00001, respectively. ntrees is defined as number of trees required for generating random forests (ntrees=400 (default)). Tolerance is the maximum threshold value allowed (tolerance=0.5). X_{train} , X_{val} , Y_{train} and Y_{val} are the features and class values of training and validation sets, respectively. The function estimateBetaLASSO is used for calculation of β^{LASSO} values which is computed on the value of λ . The β^{LASSO} values are calculated by: $\frac{1}{2n} \| X\beta - y \|_2^2 + \lambda \| \beta \|_1$. Coordinate Descent (CD) minimization algorithm is implemented for minimizing the objective function with respect to each of its coordinate directions. Results of the β coefficient values for each λ value is stored in the betaE list. p contains the number of features from the training set. sel-Prob contains the selection probabilities of the features for the classification using iterative Random Forests. maxAccIndex contains the maximum prediction accuracy among n iterations of validation set sequences for particular value of λ , and accRfPred contains the prediction accuracies of all λ values. The algorithm returns a feature list of minimal and maximal optimal features.

Algorithm 1 LiRFFS algorithm

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Input: \lambda_{lower}, \lambda_{upper}, \lambda_{stepsize}, \beta, n, tolerance, ntrees, X_{train}, X_{val}, Y_{train}, Y_{val}
Output: Minimal and maximal optimal feature lists
 1: \lambda = Create list based on \lambda_{lower}, \lambda_{upper} and \lambda_{step-size} values
 2: betaLASSO = function estimateBetaLASSO(X_{train}, Y_{train})
 3: for i = 0 to length(\lambda) do
      betaE = minimize the betaLASSO using CD minimization
      if (values in betaE < tolerance) then
 5:
         Set values in betaE = 0
 6:
         betaENonZero = length(values in betaE = 0)
 7:
 8:
      end if
      betaEArray = betaE
 9:
      if (betaENonZero < length(betaEArray−1)) then
10:
         for j = 1 to betaENonZero do
11:
            [X_{trainF}, X_{valF}, Y_{trainF}, Y_{valF}] = [X_{train}[j], X_{val}[j], Y_{train}[j], X_{val}[j]]
12:
         end for
13:
         selProb = replicate the values of \frac{1}{n}
14:
         Initialise rf as list
15:
         for iter = 1 to n do
16:
            rf[iter] = RandomForest(X_{trainF}, Y_{trainF}, X_{valF}, Y_{valF}, selProb, ntrees)
17:
            selProb = giniImportance(rf[iter])
18:
19:
         maxAccIndex = extract index values of the \lambda value having
20:
         the maximum prediction accuracy stored in r f[iter]
         Store the maxAccIndex value in accRfPred list
21:
22:
      end if
23: end for
24: for i = \max AccIndex to 0 do
      diffArrNeg = accRfPred[i] - accRfPred[i-1]
26: end for
27: for i =maxAccIndex to length(accRfPred) do
      diffArrPos = accRfPred[i] - accRfPred[i+1]
29: end for
30: Extract index values of diffArrNeg and diffArrPos values
    and store in thresArrNeg and thresArrPos lists
31: Extract Last elements from thresArrNeg and thresArrPos lists
32: return Optimal feature list based on thresArrPos and thresArrNeg lists
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2.5. 10-fold cross validation and repeated 10-fold cross validation with data shuffling

For evaluating the prediction accuracy of PLIT against CPC2, CPAT, IncScore and PLEK tools, a 10-fold Cross Validation (CV) and repeated 10-fold CV with data shuffling benchmarking was performed on the coding and non-coding sequences extracted from the RNA-seq datasets. From the complete sequence set, 10% were selected as test set and 90% as training set in each fold consisting of balanced IncRNA and protein-coding sequences. For repeated 10-fold CV, five repetitions were performed with shuffling of sequences in each iteration followed by 10-Fold CV in each repetition.

2.6. Performance evaluation criteria

To assess classification performance of lncRNAs and mRNA transcripts, Accuracy (ACC), Sensitivity (SENS), Specificity (SPEC), F1-Score (F1), Negative Predictive Value (NPV) and Matthews Correlation Coefficient (MCC) metrics were used which are defined as follows.

- Accuracy =
$$\frac{TP+TN}{TP+FP+FN+TN}$$

- Sensitivity or Recall =
$$\frac{TP}{TP+FN}$$

- Specificity =
$$\frac{TN}{FP+TN}$$

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- Positive Predictive Value or Precision = $\frac{TP}{TP+FP}$

$$- F1\text{-Score} = \frac{2*(Precision*Recall)}{Precision+Recall}$$

- Negative Predictive Value or NPV = $\frac{TN}{TN+FN}$

- Matthews Correlation Coefficient or MCC =
$$\frac{(TP*TN)-(FP*FN)}{\sqrt{(TP+FP)(FN+TN)(FP+TN)(TP+FN)}}$$

In all the above, TP = True Positive, TN = True Negative, FP = False Positive and FN = False Negative.

3. Results

3.1. Results of optimal feature selection

The selection of optimal features was performed on a unified dataset of 6 plant species (A. thaliana, Z. mays, O. sativa, B. napus, B. rapa and B. oleracea). The dataset consisted of 22,468 (lncRNA and mRNA) transcript sequences selected as training set and 7,532 sequences selected as validation set. An optimal feature set was selected based on λ values ranging from 0.1 to 1.0×10^{-5} . Figure 2 shows the prediction accuracies of lncRNA and mRNA sequences in the validation set over the range of λ values. Based on tolerance cutoff value of 0.5, two feature sets, namely, the 7 feature set (7F) and the 31 feature set (31F) were selected having minimal and maximal optimal features. 7F is selected based on the least number of features producing higher prediction accuracy having accuracy within the tolerance threshold value from the maximum prediction accuracy λ value. Whereas 31F is selected based on the maximum number of features having prediction accuracy within the tolerance threshold value from the maximum prediction accuracy λ value. Prediction of test set sequences was performed based on the optimal feature sets obtained from LASSO-iRF computation.

The performance of the 7F, 31F and 73F feature sets were compared on *Arabidopsis thaliana, Zea mays, Sorghum bicolor* and *Vitis vinifera* RNA-seq test sets containing lncRNA and mRNA transcripts. The AUC plots (Figure 3) shows that performance of 31F was similar to 73F, whereas 7F showed slight decrease in the AUC value as compared to 31F and 73F. A difference of 0.02, 0.0085 and 0.02 in the AUC values for 7F can be observed in *Arabidopsis thaliana, Sorghum bicolor* and *Vitis vinifera* test set sequences (Figure 3(a), (c) and (d)). Although the difference is minor, prediction of lncRNAs by the 7F feature set increases the False Positive Rate as compared to the 31F feature set.

3.2. Performance of PLIT feature groups on plant RNA-seq datasets

To evaluate performance of PLIT, the tool was evaluated on coding and noncoding sequences obtained from seven plant RNA-seq species. Individual performance of PLIT in different species (Table 2) shows that on an average, a higher accuracy and specificity values were obtained for *A. thaliana, G. max*, *O. sativa, S. lycopersicum, V. vinifera* and *Z. mays* datasets. High specificity value indicate accuracy in identification of lncRNA sequences in the RNA-seq dataset. An evaluation of F1 and accuracy values displayed precision in detecting lncRNA sequences. A measure of NPV also demonstrates accuracy in determining the true positives and true negatives from the test set.

Table 2: Performance (percentage accuracy) of PLIT on different plant RNA-seq datasets.

Data set	ACC	SENS	SPEC	F1	NPV	MCC
Arabidopsis thaliana	77.16	76.05	78.27	77.16	76.57	0.54
Glycine max	84.16	82.92	85.41	84.16	83.33	0.68
Oryza sativa	87.09	86.18	88.00	87.09	86.42	0.74
Solanum lycopersicum	85.35	83.95	86.74	85.34	84.39	0.70
Sorghum bicolor	88.57	88.98	88.16	88.57	88.88	0.77
Vitis vinifera	81.04	75.81	86.28	81.00	78.10	0.62
Zea mays	94.94	94.25	95.63	94.94	94.32	0.89

An area under receiver operating characteristic (AUC) curve gives better insight about the ability of a classifier to separate two classes. From the RNA-seq datasets, a higher AUC was observed for *S. lycopersicum*, G. max, *O. sativa*, *V. vinifera*, and *Z. mays* with an average AUC of 0.933 (Figure 4). A slightly lower AUC was observed for *S. bicolor* and *A. thaliana* having AUC of 0.75 and 0.85 respectively. The AUC and the evaluation metrics clearly indicates that PLIT predict the lncRNA sequences in plants without overfitting the training data.

The performance of PLIT was benchmarked against the four popular coding potential computation tools: CPAT, CPC2, PLEK and IncScore, using the plant RNA-seq test datasets. An initial benchmarking analysis based on the prediction accuracy (Table 3) shows that PLIT exhibited much higher prediction accuracies in all the plant RNA-seq test set sequences. The accuracy of PLIT ranged from 76.5% to 96.7%, while only IncScore achieved the accuracy >90% among other tools. The accuracy for IncScore ranged between 62.7% to 92.6%, whereas CPC2, PLEK and CPAT demonstrated accuracies between 52.1% and 89.07%. Apart from accuracy, it was noticed that PLIT achieved higher sensitivity and specificity in all the test sets which

indicates that PLIT is higher quality classifier for plants. CPAT demonstrated comparable sensitivity but demonstrated much lower specificity. On the other hand, CPC2, PLEK and IncScore achieved higher specificity but much lower sensitivity. Lower sensitivity implies producing higher false negatives i.e. classifying coding sequences as long non-coding transcripts, whereas lower specificity implies increase in false positive results i.e. classifying long non-coding as coding transcripts (Table 4 and Table 5). Table 4 and 5 demonstrates that CPC2 and IncScore shows lower Sensitivity and higher Specificity values for all the plant species, whereas CPAT and PLEK produces lower Specificity and higher Sensitivity values. Overall, the results demonstrated PLIT as more accurate and robust tool for differentiating IncRNA and protein-coding transcripts in plants exhibiting consistently greater accuracy, Sensitivity and Specificity metrics in all plant species.

Table 3: Performance comparison (percentage accuracy) of PLIT with existing tools on different plant RNA-seq datasets.

Data set	CPAT	CPC2	PLEK	lncScore	PLIT
Arabidopsis thaliana	53.01	50.40	64.98	67.70	76.05
Glycine max	65.42	57.98	67.90	73.22	86.88
Oryza sativa	64.49	66.30	69.38	78.44	86.59
Solanum lycopersicum	55.58	58.48	63.02	66.74	86.04
Sorghum bicolor	66.93	60.41	64.69	71.22	87.96
Vitis vinifera	52.24	58.97	65.21	63.34	78.30
Zea mays	87.91	59.53	79.39	92.33	96.43

Table 4: Sensitivity comparison of PLIT with existing tools on different plant RNA-seq datasets.

Data set	CPAT	CPC2	PLEK	IncScore	PLIT
Arabidopsis thaliana	65.30	32	67.75	57.96	76.25
Glycine max	82.70	32.62	57.76	66.06	86.17
Oryza sativa	78.17	47.10	67.27	77.81	88.40
Solanum lycopersicum	72.22	27.44	56.65	63.19	84.18
Sorghum bicolor	81.68	48.16	73.06	68.16	88.98
Vitis vinifera	80	35.91	61.30	56.53	73.81
Zea mays	91.39	44.51	87.64	87.22	96,16

3.3. Results of 10-fold Cross Validation (CV) performance benchmarking

To assess the performance of PLIT in plant RNA-seq datasets, a 10-fold CV performance benchmarking was performed. The prediction accuracy of PLIT against

Table 5: Specificity comparison of PLIT with existing tools on different plant RNA-seq datasets.

Data set	CPAT	CPC2	PLEK	lncScore	PLIT
Arabidopsis thaliana	51.67	68.81	63.18	78.27	75.85
Glycine max	60.09	83.33	79.07	81.56	87.58
Oryza sativa	59.75	85.50	71.74	79.34	84.78
Solanum lycopersicum	53.19	89.53	71.62	72.79	87.90
Sorghum bicolor	61.56	72.65	52.65	77.55	86.93
Vitis vinifera	51.16	82.04	69.57	70.57	82.79
Zea mays	84.97	74.54	71.14	97.44	96.70

CPAT, CPC2, PLEK and lncScore was tested on each fold. To evaluate the classification performance of PLIT, 31F was used for comparing the prediction accuracies of test sets.

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Transcript length distribution of TAIR10-annotated and EST-derived lncRNA transcripts demonstrates the degree of sequence length variation in lncRNA transcripts (Figure 5). Sequences derived from the TAIR10 annotation data ranges between 200 bp and 8000 bp whereas sequences derived from EST analysis ranges widely between 200 bp and 7.8×10^5 bp. Additionally, ORF count of EST-lncRNA sequences reveals counts greater than 700 ORFs per frame. Such extremely long lncRNA sequences are generally mis-classified as protein-coding transcripts, due to which the overall prediction accuracy decreases.

The 10-fold CV benchmarking on seven plant RNA-seq test set sequences (Figure 6) demonstrates PLIT's superior performance exhibiting significant improvement in the prediction accuracy across all the folds. Among all the tools, CPC2 demonstrated poor accuracy in predicting the lncRNAs in the *A. thaliana*, *G. max*, *O. sativa*, *S. bicolor* and *Z. mays* datasets. The prediction accuracies ranged between 49.7% and 53% for *A. thaliana*, 57% to 63% for *G. max*, 63% to 69% for *O. sativa*, 56% to 66% for *S. bicolor* and 50% to 60% for *Z. mays*. Similar to CPC2, CPAT also generated lower prediction accuracies with accuracies ranging between 49% to 54% for *A. thaliana* and *V. vinifera* species. CPAT generated comparatively higher prediction accuracy against PLEK and CPC2 on *S. bicolor* dataset. PLEK and lncScore exhibited similar performance for *A. thaliana*-EST derived sequences, *G. max* and *V. vinifera* displaying accuracies in the range of 60% to 76%. lncScore, on the other

hand, demonstrated higher performance among CPAT, PLEK and CPC2 tools with accuracies ranging between 65% and 70% for *A. thaliana* TAIR10 annotated lncRNA sequences and *S. lycopersicum*, 74% to 80% for *O. sativa*, and 70% to 78% for *S. bicolor*. The difference in the accuracies between PLIT and other tools ranges between 9% to 29% in *A. thaliana*, 11% to 27% for *G. max*, 7% to 20% for *O. sativa*, 16% to 29% for *S. lycopersicum*, 12% to 25% for *S. bicolor*, 13% to 27% for *V. vinifera*, and 3% to 46% for *Z. mays*. However, for the *Z. mays* test set transcript sequences, the highest difference was produced by CPC2 alone. CPAT and PLEK generated accuracy difference ranging between 3 to 8%, whereas PLEK exhibited a difference of 16% against PLIT.

3.4. Results of repeated 10-Fold Cross Validation performance benchmarking with data shuffling

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To further evaluate the efficiency and robustness of PLIT tool, a repeated 10-fold CV benchmarking was performed by repeatedly shuffling the coding and long non-coding sequences in each iteration. The prediction accuracies in each iteration were averaged to calculate a mean accuracy value along with standard error around the mean value (Figure 7). PLIT generated a mean accuracy 78.07% with SE of 1.2, whereas lncScore, CPC2, PLEK and CPAT displayed mean accuracies of 68.41%, 50.7%, 53.5% and 63.15% along with SE ranging between 0.7 to 1.3 in *A. thaliana* (Figure 7a and b). PLIT showed mean prediction accuracy range of 84.5 - 86.6% for *G. max, O. sativa, S. lycopersicum* and *S. bicolor* datasets (Figure 7c, d, e and f), whereas for *V. vinifera* and *Z. mays* (Figure 7g and h), mean values of 78% and 96.2% were obtained respectively. The SE value ranged between 1.2 and 1.7 for all the plant species except *Z. mays* which displayed a SE of 0.45.

Prediction accuracies of CPC2, CPAT and lncScore over different test set sequences in several plant species displayed an average SE of 1.8 for *G. max, O. sativa* and *S. bicolor.* In *S. lycopersicum* dataset, an average SE of 1.38 was observed. For *V. vinifera*, CPC2, PLEK and lncScore generated SE of 1.5, whereas PLEK produced a lower SE value of 0.5. The mean accuracy and SE plots demonstrates the consistency of accuracy values of PLIT across several folds and repetitions when tested against currently

known and popular tools. As mentioned previously in Section 3.3, the difference in the accuracy remained consistently similar with slight deviation along the mean value.

3.5. Comparison of PLIT-LiRFFS against mRMR feature selection method

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The results of LiRFFS-based feature selection implemented in PLIT were compared against the feature set obtained from the mRMR (minimum-redundancy maximum relevancy) method. Using Mutual Information Difference (MID) in mRMR, 31 feature set was obtained on the unified dataset of 6 plant species. mRMR selected transcript length among ORF and sequence-based features. The remaining 30 RSCU features were selected by mRMR-based feature selection. Whereas LiRFFS selected 7 ORF and sequence-based features and 5 codon-biased and 19 RSCU-based codonbiased features were selected. The results of the feature selection from LiRFFS and mRMR approaches were applied and tested on five species: G. max, O. sativa, S. lycopersicum, S. bicolor, and v. vinifera. Tables 6, 7 and 8 demonstrate comparison of prediction accuracy, sensitivity and specificity on different plant RNA-Seq datasets. The results clearly demonstrate a significant increase in prediction accuracy of 6.19% in O. sativa, 6.4% in S. lycopersicum and 4.7% in S. bicolor datasets, whereas a slightly higher increase of 0.36% and 1.74% for G. max and V. vinifera datasets. Sensitivity and specificity analysis results exhibits higher metric values in all the species except G. max and V. vinifera where a minor increase of 1.06% in Sensitivity and 0.25% was observed.

 $Table \ 6: \ Performance \ comparison \ (percentage \ accuracy) \ of \ LiRFFS \ algorithm \ against \ mRMR-based \ selected \ features \ implemented \ on \ different \ plant \ RNA-seq \ datasets.$

Data set	LiRFFS Accuracy	mRMR Accuracy
Glycine max	83.27	82.91
Oryza sativa	86.91	80.72
Solanum lycopersicum	84.30	77.90
Sorghum bicolor	85.92	81.22
Vitis vinifera	79.42	77.68

Table 7: Sensitivity comparison of LiRFFS algorithm against mRMR-based selected features implemented on different plant RNA-seq datasets.

Data set	LiRFFS Sensitivity	mRMR Sensitivity	
Glycine max	81.85	82.91	
Oryza sativa	86.91	82.54	
Solanum lycopersicum	83.25	76.74	
Sorghum bicolor	81.22	81.63	
Vitis vinifera	75.06	71.32	

Table 8: Specificity comparison of LiRFFS algorithm against mRMR-based selected features implemented on different plant RNA-seq datasets.

Data set	LiRFFS Specificity	mRMR Specificity	
Glycine max	84.69	82.91	
Oryza sativa	86.91	78.91	
Solanum lycopersicum	85.34	79.07	
Sorghum bicolor	90.61	80.81	
Vitis vinifera	83.79	84.04	

4. Discussion

As RNA-Seq technology has been widely developed for identification of novel lncRNAs, recent advances in computer science has enabled computational prediction of lncRNAs solely from genomic sequence. Many tools developed for identification of lncRNAs paid less attention for their identification of the sequences derived from RNA-sequencing studies in plants. Therefore, it is necessary to develop a tool which could identify these long non-coding sequences with higher accuracy. In this study, we developed a new tool, called PLIT for accurately identifying the lncRNA transcript sequences from RNA-seq datasets. PLIT constructs a set of sequencebased and codon-bias features from the target FASTA sequence derived from RNAseq data. The tool additionally implements and provides a feature selection-based approach for identifying a set of optimal features using LASSO and iRF classifier methods. The implementation of the LiRFFS algorithm in PLIT selected 24 codonbias and 7 sequence features. The selected features shows that identification of lncRNA sequences primarily depends on open reading-frames, length of transcript, coverage of the ORFs in different sequences, frequency of hexamers, base pair positioning in the reading frames, GC content, contribution of frequency of base pairs in the codons and selection of synonymous codons in the transcript sequences. Furthermore, some of the extracted features such as GC content, sequence length as well as the ORF length which are implemented in PLIT have been previously administered in PredCircRNA tool for classification of circular RNAs from other lncRNA sequences [37]. Another study conducted by Hu et al. (2015) developed RNAfeature for characterizing novel ncRNAs by finding significant features shared by various ncRNA sequences [38]. The study determined 10 essential features including structures, sequences, expression profiles and histone modification signals. When compared with RNAfeature, the common feature implemented in both PLIT, PredCircRNA and RNAfeature tools is the GC content. This feature signifies its importance in lncRNA identification. Apart from GC content, RNAfeature used DNA and protein sequence conservation features, RNA secondary structure stability, homologs, conservation features and ORF property feature which is constructed from multiple sequence alignment method.

In this study, a set of distinct 8 plant species from the Refseq database have been used for individually testing the prediction performance of the feature set implemented in PLIT tool. Furthermore, a unified set of 6 plant species from the Refseq database was constructed containing model and non-model organisms which were used for obtaining optimal set of features. A trimmed (minimal) set of features from the feature selection results can be further used for obtaining accurately identifying lncRNA sequences in plant RNA-Seq datasets. Therefore, current study focuses on the accurate prediction and benchmarking against other widely known tools. lncRNA sequences deposited in CANTATAdb database have been used as negative samples whereas protein-coding genes deposited in Ensembl database have been used as positive samples for constructing training and test set sequences for benchmarking the prediction accuracy of PLIT tool. The optimal feature set is obtained by iteratively constructing a training model from the training sequence set and testing the model on the validation set of coding and non-coding sequences. Since, Refseq and Ensembl databases contains lncRNA and protein-coding sequences for model and non-model species, a supervised classification approach can be implemented on the reference set of sequences which can potentially generate a set of optimal features. This reduced set of features can be used as input in further downstream

analysis in wide variety of RNA-Seq plant species for the accurate identification of lncRNA sequences from protein-coding genes.

Identification of lncRNA sequences primarily exhibits their selection based on the ORFs on the DNA strand and position of purines and pyramidines in the reading frame. Additionally, the effect of increased GC content in lncRNA prediction is linked to the codon usage pattern where higher GC content indicates heterogeneity [39]. A study conducted by Zhou et al. (2014) confirms the existence of linear relationship between amino acid usage and genomic GC content [39]. Another study conducted by Biro et al. (2008) discusses about the correlations between individual codons as well as codon residues at different codon positions [40]. Furthermore, non-randomness of synonymous codon usage is highly affected by tRNA pool size having a primary role in the reading frame and imposing constraints on the synonymous codon usage [41, 42].

These results provide insights into the preferential selection of synonymous codons in the classification process. LiRFFS produced a minimal and maximal set of optimal feature sets from the training and validation datasets constructed from six plant species. The AUC profiles of the 31F optimal feature set on plant RNA-seq datasets demonstrates comparatively higher performance when compared against the 7F set. The similarity of the ROC curves for 31F and 73F sets indicate better selection of features represented by greater prediction performance. Test set sequences used in RNA-Seq datasets were used to demonstrate prediction accuracy of PLIT tool against other existing tools and its application for identifying novel lncRNAs based on the optimal feature set.

When comparing against currently popular state of the art alignment-free tools such as CPAT, CPC2, PLEK and lncScore [5, 7, 8, 12], PLIT generated much better prediction accuracy values when tested on several plant datasets. From 10-fold CV and repeated 10-fold CV analyses, it was found that the prediction accuracies of other tools were comparitively lower with average differences ranging between 9 to 30%. The sensitivity and specificity values obtained from other tools displayed a biased prediction thereby generating greater false negatives or false positives. Prediction performance of PLIT showed accuracies >80% on most plant species whereas other

tools performed inconsistently with accuracies varying significantly across different species.

Compared with other tools, PLIT provides unique set of features for accurate identification of lncRNAs transcripts which provides several advantages over other tools. First, apart from commonly known distinguishing sequence-based features such as ORF length, ORF coverage, GC content, Fickett score and Hexamer score, it takes advantage of codon-biased features to increase its discriminative power. Second, PLIT implements a powerful semi-supervised optimization approach for selection of principal features which can be applied on plants and vertebrates. Third, the implemented LiRFFS and random forest classifier for feature selection and prediction of lncRNAs offers robustness, efficiency and suitability on multiple model as well as non-model plant species. The results of PLIT revealed higher accuracy with balanced sensitivity and specificity on all test sets. This implies that the models generated by PLIT did not overfit the training data.

5. Conclusions

In this work, we developed a novel tool, PLIT, for accurate identification and discovery of lncRNA sequences particularly well-suited for RNA-seq data from plants. PLIT exceeds the prediction performance over other tools on various parameters.

The ability to identify and differentiate various lncRNA transcripts was demonstrated with several cross-validation tests on different RNA-seq datasets. Using LiRFFS, optimal features were identified based on the FASTA sequences from Refseq database. Evaluation with K-fold and repeated K-fold CV demonstrated consistent and superior performance of PLIT on all plant datasets against other tools. Thus, PLIT is a stable, robust and accurate tool for distinguishing long non-coding and protein-coding transcripts from plant RNA-seq data.

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550 References

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- [1] X. Liu, L. Hao, D. Li, L. Zhu, S. Hu, Long Non-coding RNAs and Their Biological Roles in Plants, Genomics, Proteomics & Bioinformatics 13 (3) (2015) 137–147. doi:10.1016/j.gpb.2015.02.003.
- [2] J. Harrow, A. Frankish, J. M. Gonzalez, E. Tapanari, M. Diekhans, F. Kokocinski, B. L. Aken, D. Barrell, A. Zadissa, S. Searle, I. Barnes, A. Bignell, V. Boychenko, T. Hunt, M. Kay, G. Mukherjee, J. Rajan, G. Despacio-Reyes, G. Saunders, C. Steward, R. Harte, M. Lin, C. Howald, A. Tanzer, T. Derrien, J. Chrast, N. Walters, S. Balasubramanian, B. Pei, M. Tress, J. M. Rodriguez, I. Ezkurdia, J. Van Baren, M. Brent, D. Haussler, M. Kellis, A. Valencia, A. Reymond, M. Gerstein, R. Guigó, T. J. Hubbard, GENCODE: The reference human genome annotation for the ENCODE project, Genome Research 22 (9) (2012) 1760–1774. doi:10.1101/gr.135350.111.
 - [3] Y. Zhao, H. Li, S. Fang, Y. Kang, W. Wu, Y. Hao, Z. Li, D. Bu, N. Sun, M. Q. Zhang, R. Chen, NONCODE 2016: An informative and valuable data source of long non-coding RNAs, Nucleic Acids Research 44 (D1) (2016) D203–D208. doi: 10.1093/nar/gkv1252.
 - [4] M. W. Szcześniak, W. Rosikiewicz, I. Makałowska, CANTATAdb: A collection of plant long non-coding RNAs, Plant and Cell Physiologydoi:10.1093/pcp/ pcv201.
- [5] Y. J. Kang, D. C. Yang, L. Kong, M. Hou, Y. Q. Meng, L. Wei, G. Gao, CPC2: A fast and accurate coding potential calculator based on sequence intrinsic features, Nucleic Acids Researchdoi:10.1093/nar/gkx428.
 - [6] L. Sun, H. Luo, D. Bu, G. Zhao, K. Yu, C. Zhang, Y. Liu, R. Chen, Y. Zhao, Utilizing sequence intrinsic composition to classify protein-coding and long non-coding transcripts, Nucleic Acids Research 41 (17). doi:10.1093/nar/ gkt646.

- [7] L. Wang, H. J. Park, S. Dasari, S. Wang, J. P. Kocher, W. Li, CPAT: Coding-potential assessment tool using an alignment-free logistic regression model, Nucleic Acids Research 41 (6). doi:10.1093/nar/gkt006.
- [8] A. Li, J. Zhang, Z. Zhou, PLEK: a tool for predicting long non-coding RNAs and messenger RNAs based on an improved k-mer scheme, BMC Bioinformatics 15 (1) (2014) 311. doi:10.1186/1471-2105-15-311.
 - [9] J. W. Fickett, Recognition of protein coding regions in DNA sequences, Nucleic Acids Research 10 (17) (1982) 5303-5318. doi:10.1093/nar/10.17.5303.
- [10] N. A. O'Leary, M. W. Wright, J. R. Brister, S. Ciufo, D. Haddad, R. McVeigh, B. Rajput, B. Robbertse, B. Smith-White, D. Ako-Adjei, A. Astashyn, A. Badretdin, Y. Bao, O. Blinkova, V. Brover, V. Chetvernin, J. Choi, E. Cox, O. Ermolaeva, C. M. Farrell, T. Goldfarb, T. Gupta, D. Haft, E. Hatcher, W. Hlavina, V. S. Joardar, V. K. Kodali, W. Li, D. Maglott, P. Masterson, K. M. McGarvey, M. R. Murphy, K. O'Neill, S. Pujar, S. H. Rangwala, D. Rausch, L. D. Riddick, C. Schoch, A. Shkeda, S. S. Storz, H. Sun, F. Thibaud-Nissen, I. Tolstoy, R. E. Tully, A. R. Vatsan, C. Wallin, D. Webb, W. Wu, M. J. Landrum, A. Kimchi, T. Tatusova, M. DiCuccio, P. Kitts, T. D. Murphy, K. D. Pruitt, Reference sequence (RefSeq) database at NCBI: Current status, taxonomic expansion, and functional annotation, Nucleic Acids Research 44 (D1) (2016) D733–D745. doi:10.1093/nar/gkv1189.
- [11] D. R. Zerbino, P. Achuthan, W. Akanni, M. R. Amode, D. Barrell, J. Bhai, K. Billis, C. Cummins, A. Gall, C. G. Girón, L. Gil, L. Gordon, L. Haggerty, E. Haskell, T. Hourlier, O. G. Izuogu, S. H. Janacek, T. Juettemann, J. K. To, M. R. Laird,
 I. Lavidas, Z. Liu, J. E. Loveland, T. Maurel, W. McLaren, B. Moore, J. Mudge, D. N. Murphy, V. Newman, M. Nuhn, D. Ogeh, C. K. Ong, A. Parker, M. Patricio, H. S. Riat, H. Schuilenburg, D. Sheppard, H. Sparrow, K. Taylor, A. Thormann, A. Vullo, B. Walts, A. Zadissa, A. Frankish, S. E. Hunt, M. Kostadima, N. Langridge, F. J. Martin, M. Muffato, E. Perry, M. Ruffier, D. M. Staines, S. J. Trevan-

- ion, B. L. Aken, F. Cunningham, A. Yates, P. Flicek, Ensembl 2018, Nucleic Acids Researchdoi: 10.1093/nar/gkx1098.
 - [12] J. Zhao, X. Song, K. Wang, IncScore: alignment-free identification of long noncoding\nRNA from assembled novel transcripts, Sci. Rep. 6 (2016) 34838. doi:10.1038/srep34838.
- [13] X. N. Fan, S. W. Zhang, lncRNA-MFDL: identification of human long non-coding RNAs by fusing multiple features and using deep learning, Molecular Biosystems 11 (3) (2015) 892–897. doi:10.1039/c4mb00650j.
 - [14] R. Tibshirani, Regression Selection and Shrinkage via the Lasso (1996). arXiv: 11/73273, doi:10.2307/2346178.
- [15] S. Basu, K. Kumbier, J. B. Brown, B. Yu, Iterative random forests to discover predictive and stable high-order interactions., Proceedings of the National Academy of Sciences of the United States of America 115 (8) (2018) 1943–1948. doi:10.1073/pnas.1711236115.
- [16] R. Leinonen, H. Sugawara, M. Shumway, The sequence read archive, Nucleic Acids Researchdoi:10.1093/nar/gkq1019.
 - [17] M. Martin, Cutadapt removes adapter sequences from high-throughput sequencing reads, EMBnet.journal 17 (1) (2011) 10. doi:10.14806/ej.17.1. 200.
- [18] D. Kim, G. Pertea, C. Trapnell, H. Pimentel, R. Kelley, S. L. Salzberg, TopHat2: accurate alignment of transcriptomes in the presence of insertions, deletions and gene fusions., Genome biology 14 (4) (2013) R36. doi:10.1186/gb-2013-14-4-r36.
 - [19] H. Li, B. Handsaker, A. Wysoker, T. Fennell, J. Ruan, N. Homer, G. Marth, G. Abecasis, R. Durbin, The Sequence Alignment/Map format and SAMtools, Bioinformatics 25 (16) (2009) 2078–2079. arXiv:1006.1266v2, doi:10.1093/bioinformatics/btp352.

- [20] J. W. Fickett, C. S. Tung, Assessment of protein coding measures., Nucleic acids research 20 (24) (1992) 6441–6450. doi:10.1093/nar/20.24.6441.
- [21] A. Roth, M. Anisimova, G. M. Cannarozzi, Measuring codon usage bias, in:

 Codon Evolution: Mechanisms and Models, 2012. doi:10.1093/acprof:

 osobl/9780199601165.003.0013.
 - [22] M. C. Frith, A. R. Forrest, E. Nourbakhsh, K. C. Pang, C. Kai, J. Kawai, P. Carninci, Y. Hayashizaki, T. L. Bailey, S. M. Grimmond, The abundance of short proteins in the mammalian proteome, PLoS Genetics 2 (4) (2006) 515–528. doi:10.1371/journal.pgen.0020052.

640

645

- [23] M. Amit, M. Donyo, D. Hollander, A. Goren, E. Kim, S. Gelfman, G. Lev-Maor, D. Burstein, S. Schwartz, B. Postolsky, T. Pupko, G. Ast, Differential GC Content between Exons and Introns Establishes Distinct Strategies of Splice-Site Recognition, Cell Reports 1 (5) (2012) 543–556. doi:10.1016/j.celrep.2012.03.013.
- [24] B. Clarke, Darwinian evolution of proteins, Science 168 (190) 1009–1011. doi: 10.1126/science.168.3934.1009.
- [25] T. Ikemura, Correlation between the abundance of yeast transfer RNAs and the occurrence of the respective codons in protein genes, Journal of Molecular Biology 158 (4) (1982) 573–597. doi:10.1016/0022-2836(82)90250-9.
- [26] S. Karlin, J. Mrázek, What drives codon choices in human genes?, Journal of molecular biology 262 (4) (1996) 459–72. doi:10.1006/jmbi.1996.0528.
- [27] U. Roymondal, S. Das, S. Sahoo, Predicting gene expression level from relative codon usage bias: An application to escherichia coli genome, DNA Research

 16 (1) (2009) 13–30. doi:10.1093/dnares/dsn029.
 - [28] H. Suzuki, R. Saito, M. Tomita, The 'weighted sum of relative entropy': A new index for synonymous codon usage bias, Gene 335 (1-2) (2004) 19–23. doi: 10.1016/j.gene.2004.03.001.

- [29] X.-F. Wan, D. Xu, A. Kleinhofs, J. Zhou, Quantitative relationship between synonymous codon usage bias and GC composition across unicellular genomes.,

 BMC evolutionary biology 4 (2004) 19. doi:10.1186/1471-2148-4-19.
 - [30] P. M. Sharp, T. M. F. Tuohy, K. R. Mosurski, Codon usage in yeast: Cluster analysis clearly differentiates highly and lowly expressed genes, Nucleic Acids Research 14 (13) (1986) 5125–5143. doi:10.1093/nar/14.13.5125.
- [31] P. Pudil, J. Novovičová, J. Kittler, Floating search methods in feature selection, Pattern Recognition Letters 15 (11) (1994) 1119–1125. doi:10.1016/0167-8655(94)90127-9.
 - [32] M. L. Huang, Y. H. Hung, W. M. Lee, R. K. Li, B. R. Jiang, SVM-RFE based feature selection and taguchi parameters optimization for multiclass SVM Classifier, Scientific World Journal 2014. doi:10.1155/2014/795624.

- [33] H. Peng, F. Long, C. Ding, Feature selection based on mutual information: Criteria of Max-Dependency, Max-Relevance, and Min-Redundancy, IEEE Trans. on Pattern Analysis and Machine Intelligence 27 (8) (2005) 1226–1238. doi: 10.1109/TPAMI.2005.159.
- 675 [34] Y. T. Chen, M. C. Chen, Using chi-square statistics to measure similarities for text categorization, Expert Systems with Applications 38 (4) (2011) 3085–3090. doi:10.1016/j.eswa.2010.08.100.
 - [35] C. Lee, G. G. Lee, Information gain and divergence-based feature selection for machine learning-based text categorization (2006). doi:10.1016/j.ipm. 2004.08.006.
 - [36] D. W. Marquardt, Generalized inverses, ridge regression, biased linear estimation, and nonlinear estimation, Technometrics 12 (3) (1970) 591–612. doi: 10.1080/00401706.1970.10488699.
- [37] X. Pan, K. Xiong, PredcircRNA: computational classification of circular RNA from other long non-coding RNA using hybrid features, Mol. BioSyst. 11 (8)

(2015) 2219-2226. doi:10.1039/C5MB00214A.

URL http://xlink.rsc.org/?DOI=C5MB00214A

690

- [38] L. Hu, C. Di, M. Kai, Y. C. T. Yang, Y. Li, Y. Qiu, X. Hu, K. Y. Yip, M. Q. Zhang, Z. J. Lu, A common set of distinct features that characterize noncoding RNAs across multiple species, Nucleic Acids Research 43 (1) (2015) 104–114. doi: 10.1093/nar/gku1316.
- [39] H. Q. Zhou, L. W. Ning, H. X. Zhang, F. B. Guo, Analysis of the relationship between genomic GC content and patterns of base usage, codon usage and amino acid usage in prokaryotes: Similar GC content adopts similar compositional frequencies regardless of the phylogenetic lineages, PLoS ONEdoi: 10.1371/journal.pone.0107319.
- [40] J. C. Biro, Does codon bias have an evolutionary origin?, Theoretical Biology and Medical Modellingdoi:10.1186/1742-4682-5-16.
- [41] M. A. Antezana, M. Kreitman, The nonrandom location of synonymous codons suggests that reading frame- independent forces have patterned codon preferences, Journal of Molecular Evolutiondoi:10.1007/PL00006532.
 - [42] D. J. Lipman, W. J. Wilbur, Contextual constraints on synonymous codon choice, Journal of Molecular Biologydoi:10.1016/0022-2836(83)90063-3.

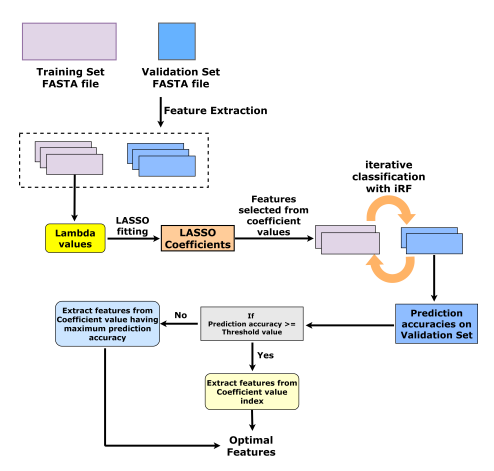


Figure 1: LiRFFS workflow. Sequence and codon-bias features from the training and validation lncRNA and protein-coding sequences are extracted. LASSO coefficients are generated from the training set and iteratively applied on the validation set using an iRF classifier to generate the prediction accuracy at each λ value. Prediction accuracies are compared against the minimum threshold tolerance value. If the prediction accuracy produced by a particular λ value is \geq minimum threshold tolerance value, the optimal features are selected from the filtered coefficient set. If accuracy is lower then the λ value, the optimal features are selected from the λ value producing the maximum prediction accuracy.

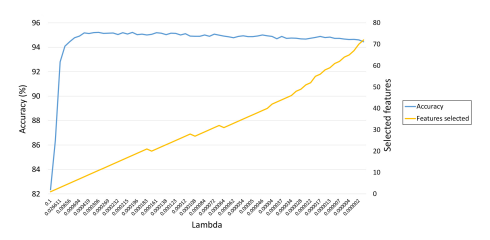
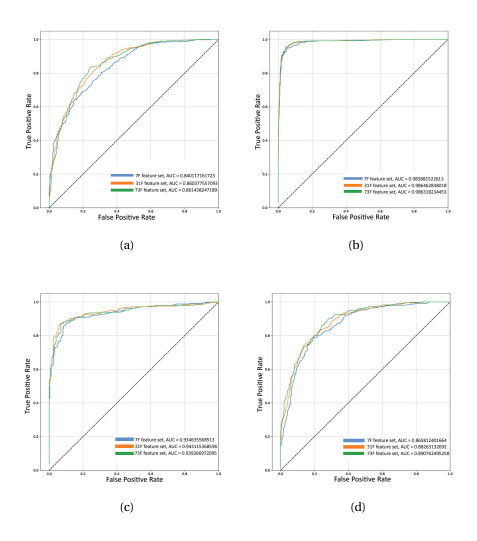


Figure 2: Feature selection bar plot. The bar plot shows prediction accuracy of the features selected at different λ values performed on validation set lncRNA and protein-coding transcripts on six plant species. X-axis shows range of λ values ranging from 0.1 to 1×10^{-6} . Primary y-axis shows the prediction accuracy and secondary y-axis shows the number of features selected at each λ value.



 $Figure \ 3: \ ROC \ plots \ of \ (a) \ \textit{Arabidopsis thaliana}, \ (b) \ \textit{Zea mays}, \ (c) \ \textit{Sorghum bicolor}, \ and \ (d) \ \textit{Vitis vinifera} \\ RNA-seq \ test \ datasets, \ showing \ comparison \ of \ AUC \ values \ for \ 7F, \ 31F \ and \ 73F \ feature \ sets.$

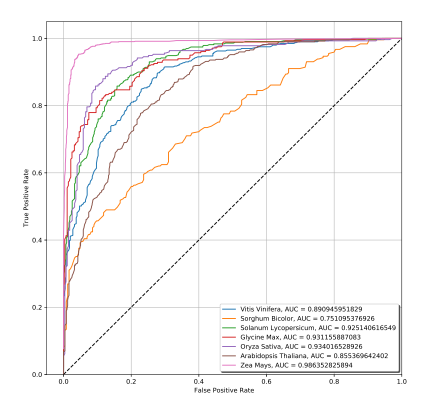
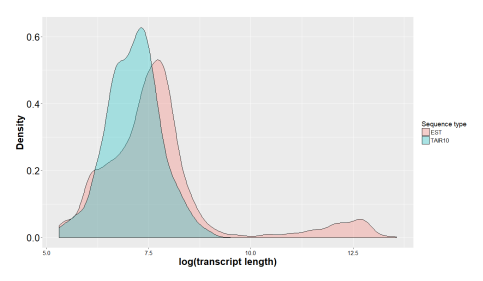


Figure 4: ROC curves and AUC values of PLIT for different RNA-seq plant data sets.



 $Figure \ 5: \ Density \ distribution \ of \ transcript \ lengths \ of \ lncRNA \ sequences \ in \ ATH \ TAIR10-annotated \ and \ EST-predicted \ results. \ X-axis \ is \ log \ of \ transcript \ lengths \ and \ y-axis \ is \ density.$

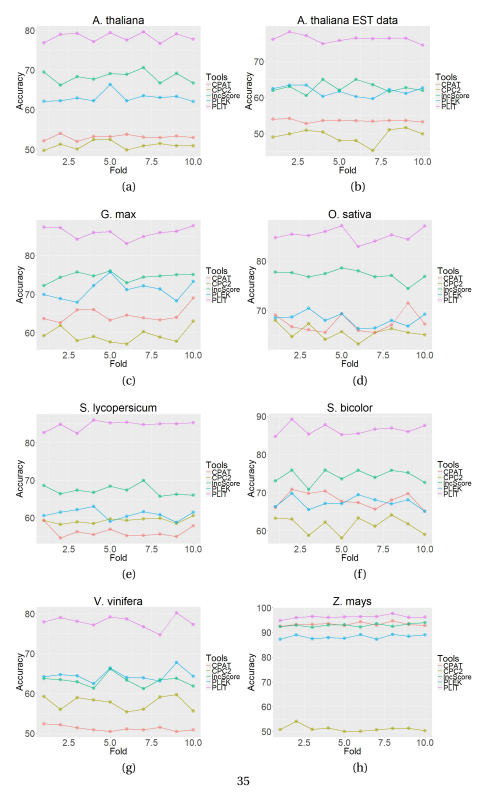


Figure 6: Plots illustrating performance of PLIT against other existing tools based on 10-fold Cross Validation benchmarking analysis for (a) *A. thaliana*, (b) *A. thaliana*-EST derived lncRNA sequences, (c) *G. max*, (d) *O. sativa*, (e) *S. lycopersicum*, (f) *S. bicolor*, (g) *V. vinifera*, and (h) *Z. mays*. X-axis represents folds whereas y-axis represents percentage accuracy.

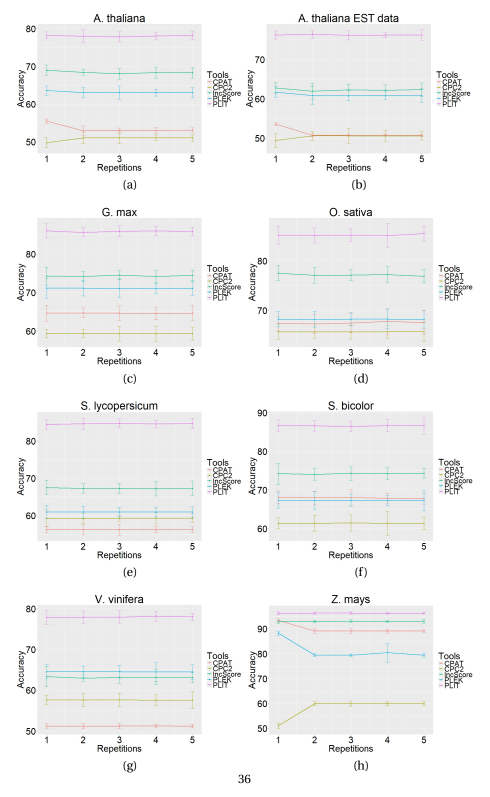


Figure 7: Plots illustrating repeated 10-fold Cross Validation benchmarking analysis with data shuffling. Five repetitions have been performed on each RNA-seq dataset. Prediction accuracy of PLIT has been bechmarked against CPAT, CPC2, PLEK and IncScore tools by averaging the accuracies over the folds and calculating the mean value on each repetition. The error bars indicate the Standard Error around the mean value. The following RNA-seq datasets have been used for repeated 10-fold CV analysis: (a) *A. thaliana*, (b) *A. thaliana*-EST derived IncRNA sequences, (c) *G. max*, (d) *O. sativa*, (e) *S. lycopersicum*, (f) *S. bicolor*, (g) *V. vinifera*, and (h) *Z. mays*. X-axis represents repetitions whereas y-axis represents percentage accuracy.