funbarRF: DNA barcode-based fungal species prediction using multiclass Random Forest supervised learning model

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**Abstract**

**Background:** Identification of unknown fungal species aids to the conservation of fungal diversity. As many fungal species cannot be cultured, morphological identification of those species is almost impossible. But, DNA barcoding technique can be employed for identification of such species. For fungal taxonomy prediction, the ITS (internal transcribed spacer) region of rDNA (ribosomal DNA) is used as barcode. Though the computational prediction of fungal species has become feasible with the availability of huge volume of barcode sequences in public domain, prediction of fungal species is challenging due to high degree of variability among ITS regions within species.

**Results:** A Random Forest (RF)-based predictor was built for identification of unknown fungal species. The reference and query sequences were mapped onto numeric features based on gapped base pair compositions, and then used as training and test sets respectively for prediction of fungal species using RF. More than 85% accuracy was found when 4 sequences per species in the reference set were utilized; whereas it was seen to be stabilized at ~88% if ≥7 sequence per species in the reference set were used for training of the model. The proposed model achieved comparable accuracy, while evaluated against existing methods through cross-validation procedure. The proposed model also outperformed several existing models used for identification of different species other than fungi.

**Conclusions:** An online prediction server "funbarRF" is established at http://cabgrid.res.in:8080/funbarrf/ for fungal species identification. Besides, an R-package funbarRF (https://cran.r-project.org/web/packages/funbarRF/) is also available for prediction using high throughput sequence data. The effort put in this work will certainly supplement the future endeavors in the direction of fungal taxonomy assignments based on DNA barcode.

**Keywords:** BOLD systems, DNA barcode, ITS, Fungal taxonomy, CBOL

**Background**

In meta-genomic studies, taxonomy classification is crucial for characterizing microbial communities [1]. In particular, prediction of unknown fungal specimens and conservation of their genomic resources are vital for studying and preserving fungal diversity [2]. However, identification of specimens that lacked morphological character is often difficult [3]. In this direction, molecular technique like DNA barcoding [4] has been successfully employed in the recent years for species identification [5–7]. In this technique, a standard genomic region is used to distinguish species based on barcode-gap [8]. The COI (cytochrome c oxidase subunit I) gene of mitochondrial DNA was first accepted as the barcode by the CBOL (consortium for barcode of life) [9] for prediction of animal species [3]. Later on, the matK and rbcL genes of chloroplast region were adopted by CBOL as barcodes for identification of plant species [10]. As far as fungus is concerned, the ITS of rDNA that includes ITS1 and ITS2 separated by 5.8S genic region (Fig. 1a), has been accepted by almost all the mycologists as the molecular region for species identification [11–13]. Considering the importance of barcoding in the preservation of species diversity as well as for other applications, the CBOL has been continuously emphasizing on...
the development of new approach(s) for identification of unknown species based on its barcode sequence [14, 15]. However, reference datasets having barcode sequences with known species labels are essential for the prediction of unknown species. For fungal species identification, two important ITS reference databases namely UNITE [13] and Warcup [16] have been developed. Besides, the BOLD (barcode of life data) [9] system also provides taxonomic information for fungal species identification. As far as prediction of fungal species is concerned, few computational approaches namely RDP classifier [16, 17], SINTAX [1], Mycofier [18] and MOTHUR [19] were proposed in the past. The RDP classifier employed naïve Bayes algorithm for taxonomy assignment, based on k-mer (k=8) similarity features [17]. Similar k-mer (k=8) features were also utilized in the SINTAX algorithm for taxonomy prediction by using a non-Bayesian classifier [1]. The k-nearest neighbor (kNN) algorithm was implemented in MOTHUR for taxonomy classification, based on k-mer (k=8) similarity measures [19].
Feature generation is a crucial step in computational predictions using biological sequences [24]. Since the biological sequences are the strings of alphabets, they should be transformed to numeric vectors before being employed as input in supervised learning-based predictors [25]. As far as barcode-based species identification using machine learning predictors is concerned, sparse encoding technique was adopted by Weitschek et al. [15]. In another study, Meher et al. [26] encoded the barcode sequences based on the composition of contiguous k-mer, for species identification using RF [22] machine learning technique. Specific to fungal species, k-mer features [26] were employed in RDP classifier, SINTAX algorithm and Mycofier for encoding barcode sequences into numeric vectors. Recently, Brinda et al. [27] shown that the spaced k-mer [21] provides significantly higher accuracy as compared to the contiguous k-mer. Therefore, in the present study, the g-spaced base pair features [21] were used to encode the barcode sequences into numeric feature vectors. Five kinds of g-spaced features namely 1-spaced (g=1), 2-spaced (g=2), 3-spaced (g=3), 4-spaced (g=4) and 5-spaced (g=5) were computed. This is similar to the di-nucleotide compositions with skips of 1, 2, 3, 4 and 5 nucleotides respectively [21]. For any nucleotide sequence of length N, each g-spaced feature set results in 16 descriptors. The frequency of the di-nucleotide s and t with g-gap (g-spaced feature value) is given by $D_g(s, t)/(N-1)$, where $s, t = A, T,$

**Table 1** Distribution of collected fungal barcode sequences over different genomic regions. It can be seen that >56000 sequences out of 59847 sequences are from ITS (including ITS1 and ITS2) region. These 59847 barcode sequences are belonged to 3770 species, where at least 3 sequences are present for each species.

| Genomic region | 18S | 28S | 5.8S | AOX-fmt | atp6 | COI-5P | COI | COXIII | ITS | ITS1 | ITS2 |
|----------------|-----|-----|------|---------|-----|-------|-----|--------|-----|------|------|
| # Sequences    | 5   | 6   | 2418 | 79      | 3   | 595   | 3   | 3      | 51886| 2428 | 2421 |

Mycofier, naïve Bayes classifier coupled with k-mer ($k=5$) features was adopted for identification of fungi at genus label [18].

Though concerted efforts have been put for the development of above mentioned tools and techniques that have advanced our knowledge for species identification using DNA barcode, still there is a room for further improvement. The 8-mer similarities have been adopted in RDP classifier, SINTAX and MOTHUR, where the number of features are large (i.e., $4^8$). So, prediction with same accuracy using less number of features is one of the aims of this study. Further, tool like PROTAX [7] depends upon the output of third party software BLAST [20], which itself takes longer time for performing sequence alignment for larger size dataset. Thus, the other aim of this work is to develop an alignment free tool for prediction of fungal species. Furthermore, the supervised machine learning techniques such as naïve Bayes classifier, kNN, Bayesian regression model have been successfully employed for taxonomy assignments of fungal species, as evidenced from the above mentioned studies. Keeping above in mind, we have proposed a supervised learning-based prediction model for identification of fungal species, by analyzing their barcode sequences. In the proposed model, gapped base-pair compositions [21] were used as features and Random Forest (RF) [22] methodology as predictor. The performance of the developed model was not only evaluated with fungal species but also for the prediction of other species as well. We believe that the developed approach will supplement the existing tools and techniques for species identification using DNA barcode.

**Methods**

**Barcode sequences of fungal species**

The Warcup dataset (17878 sequences belonging to 8551 species) was used to test the predictive ability of the RDP classifier [16] and SINTAX algorithm [1]. Besides, the RDP classifier was also evaluated with UNITE dataset (145019 sequences belonging to 10297 species). Further, performance of another machine learning-based classifier i.e., Mycofier [18] was tested on fungal ITS sequences from the NCBI GenBank (https://www.ncbi.nlm.nih.gov/). None of the above studies have used fungal barcode sequences of BOLD systems (http://www.boldsystems.org/), which is one of the most widespread endeavor in the field of barcode-based species identification [23]. Therefore, we preferred the BOLD database for collecting the fungal ITS sequences for our study. At first, 68565 barcode sequences belonging to 4182 species (at least 3 sequences per species), across all the 7 phyla of fungal kingdom were collected. Excluding sequences with non-standard nucleotide bases, 60348 sequences confined to 4100 species were obtained. Further excluding 330 species with 1 or 2 sequences, 3770 species with 59847 barcode sequences were retained for the analysis. Among 59847 sequences, more than 56000 are from ITS regions and rests are from other genomic portions (Table 1). Out of 59847 sequences, 1485 (2.481% of 59847) and 17549 (29.32% of 59847) sequences are found common with the Warcup and UNITE datasets respectively (Fig. 1b). So, the prepared dataset consists of ~70% non-redundant (with Warcup and UNITE) sequences (excluding the 18038 common sequences present in Warcup and UNITE datasets, which is 30.14% of 59847).
Supervised learning technique

Supervised learning methods are promising for DNA barcode-based species identification [15]. For instance, supervised learning techniques namely SVM (with sequential minimal optimization) [29], C4.5 (J48) [30], RIPPER [31] and Naïve Bayes [32] were employed by Weitschek et al. [15] for species identification based on DNA barcode. In SPIDBAR [26], RF supervised learning technique was applied for prediction of species using barcode sequences. Specific to the fungal species identification, Naïve Bayes algorithm was employed in RDP [16], SINTAX [1] and Mycofier [18], whereas kNN was used in MOTHUR [19].

Motivated by the successful application of machine learning techniques in earlier studies, we preferred to use RF supervised learning model for identification of fungal species in the present study. Here, the class labels are the species names of fungi and the number of classes is same as the number of distinct species present in the dataset. Also, there are other advantages of using RF i.e., it is non-parametric (independent of the probability distribution of the dataset), robust to noise and can handle large datasets [27]. Since there were more than two species of fungus, a multi-class RF [33] model was built for prediction of species.

Random Forest (RF)

RF [22] is an ensemble learning method, consisting of several classification trees [34], where each classifier (classification tree) is constructed on a bootstrap resample of the learning dataset. Since each classifier is built upon a bootstrap sample, on an average 36.8% of observations do not play any role in the construction of each classification tree and are called Out-Of-Bag (OOB) instances [35]. In other words, each classifier in RF is built on 2/3rd of the learning data and tested on the 1/3rd OOB sample. These OOB samples are the source of data for measuring the prediction error of RF. More clearly, the error for each classifier in RF is measured based on its OOB samples (called as OOB error) and these OOB errors are averaged over all the decision trees to compute the OOB error of the forest. As far as prediction of test instance is concerned, each classifier of RF votes each test instances to one of the pre-defined K classes and the test instance is predicted by the label of winning class [35]. There are two important parameters in RF i.e., mtry (number of variables to choose at each node for splitting) and ntree (number of decision trees to construct in the forest), tuning of which is required to achieve maximum prediction accuracy. For tuning of ntree, the RF was trained by using the feature set g=1, g=1+2, g=1+2+3, g=1+2+3+4 and g=1+2+3+4+5 with varying number of decision tress (1 to 500) and default mtry (\( \sqrt{\text{no.of variables}} = \sqrt{p} \)). The number of trees after which the OOB-error rate got stabilized was considered as the optimal ntree. With the optimum ntree, RF was again trained with the same datasets with varying mtry values (1, \( \sqrt{p} \), \( \sqrt{p^2} \), \( 2\sqrt{p} \), \( 3\sqrt{p} \), \( 5\sqrt{p} \), p). The mtry that generated the lowest OOB-error rate was considered as the optimal mtry. A flow chart describing the process involved in prediction using RF method is shown in Fig. 1d. For implementing RF methodology, the function randomForest available in the R-package “randomForest” [36] was used.

Training and validation

At least four sequences per species (class) are required to train the supervised learning classifier for species identification using DNA barcode [15]. However, we have considered those species for which at least three sequences were also available. Here, seven different datasets were prepared with 3, 4, 5, 6, 7, 8 and 9 sequences per species respectively. The sequences in these datasets were randomly drawn from the original dataset. Number of sequences and species for each category are given in Table 2. For the dataset with k sequences per species, a k-fold CV procedure [37] was employed to evaluate the species identification success rate (SISR) of the proposed model. For k-fold CV, k subsets were prepared by randomly splitting the whole dataset in such a manner that one sequence of each species was present in each subset. In the k-fold CV procedure, k-1 subsets were utilized for training of the model and the rest one subset was utilized for validating the

Table 2 Number of sequences, species, sequences/species for the considered seven categories of datasets. For instance, in the first category there are 3770 species with 11210 sequences, where each species has 3 sequences. Further, in the category with k sequences per species, a k-fold cross validation was adopted where k-1 sequences per species were used to train the model and rest one sequence was used to assess the model accuracy.

| #Sequence/Species | 3   | 4   | 5   | 6   | 7   | 8   | 9   |
|-------------------|-----|-----|-----|-----|-----|-----|-----|
| #Species          | 3770| 3461| 2777| 2328| 1998| 1773| 1498|
| #Sequence         | 11210| 13844| 13885| 13968| 13986| 14184| 13482|
corresponding trained model in each fold. In this procedure, all the \( k \) subsets were provided equal opportunity to be used as validation set, where the accuracy was measured in terms of SISR averaged over \( k \) folds of the CV. The SISR is defined as follows:

Let \( N_h \) be the number of query sequences belong to the \( h^{th} \) species (class) and \( n_h \) be the number of query instances correctly classified into \( h^{th} \) class, where \( h=1, 2, \ldots, H \). Then the SISR can be computed as

\[
\text{SISR} = \frac{\sum_{h=1}^{H} n_h}{\sum_{h=1}^{H} N_h}.
\]

Prediction for other species

To check the suitability of the proposed approach for the prediction of other species (other than fungi), its performance was assessed on five different taxonomical entities namely Inga, Drosophila, Cypraeidae, Fish and Bat. The barcode sequences for these entities were retrieved from http://dmb.iasi.cnr.it/blog.php, which have also been utilized in earlier developed species identification methods [15, 38]. The numbers of sequences for the reference and query datasets for these entities are given in Table 3.

Table 3 Summary of the training and test datasets for five different taxonomical entities.

| Dataset          | Taxonomical entity | Drosophila | Inga  | Fish  | Bat   | Cypraeidae |
|------------------|--------------------|------------|-------|-------|-------|------------|
| #Train (reference) | 419                | 791        | 515   | 682   | 1656  |
| #Test (query)    | 116                | 122        | 111   | 144   | 352   |

#Train: Number of sequences in the training set
#Test: Number of sequences in the test set

Comparison with existing fungal taxonomy prediction method

The proposed computational model was further compared with the existing fungal species identification methods namely RDP classifier, SINTAX and MOTHUR. We used the executable code of the MOTHUR (https://github.com/mothur/mothur/releases/tag/v1.40.5), RDP classifier (https://sourceforge.net/projects/rdp-classifier/) and SINTAX (http://www.drive5.com/usearch/manual/cmd_sintax.html) for implementing the corresponding algorithms in our fungal datasets. The performances of the methods were evaluated with a dataset of 1363 species (10 sequences per species). Accuracies were computed over 10-fold CV, where one sequence of each species was present in each fold. We preferred to use 10 sequences per species, because the datasets up to 9 sequences/species were utilized for assessing the SISR of the proposed computational model (see subsection Training and validation).

Results

Parameter optimization analysis

In all the five model representations (\( g=1, g=1+2, g=1+2+3, g=1+2+3+4+5 \)) the OOB-error rates are seen to be stabilized after \( ntree=400 \) (Fig. 2a), for all the seven datasets (3-9 fold). It can also be seen that the OOB-errors are lower for the dataset with larger number of sequences per species. For instance, OOB-error rates are lower for all the model representation with 9 sequences per species than that of others (Fig. 2a). It is further observed that the OOB-error is lowest for \( g=1+2+3+4+5 \), as compared to the other model representation (Fig. 2b). Though, the errors are getting stabilized around \( ntree=400 \) (Fig. 2a), the optimum value of \( ntree \) was kept as 500 anticipating further improvement. With the optimum value of \( ntree (=500) \), it is further observed
that OOB-errors are minimum for the dataset with 9 sequences per species for all the seven mtry values and five model representations (Fig. 2c). Further among the five model representations, OOB-error is seen to be lowest for $g=1+2+3+4+5$ and that is with $mtry=\sqrt{p}$, which is the default mtry value (9 in the present study) in RF (Fig. 2d). Thus $g=1+2+3+4+5$ is the best model representation with lowest OOB-error, and the optimum values of RF parameters ntree and mtry are 500 and 9 respectively.
Analysis of g-spaced base pair features
Although the OOB error rate is found to be lowest for the model representation \( g=1+2+3+4+5 \), cross validation analysis was performed in all the five model representations (feature sets) to have a comprehensive comparative analysis. The SISRs for different number of sequences per species and for different combinations of \( g \) (i.e., model representations) are shown in Fig. 3. The SISRs are observed to be gradually increased while numbers of sequences per species are increased, for all the combinations of \( g \) (Fig. 3v). In particular, SISR reached 80%, when 4 sequences per species are used to train the model (Fig. 3a). The success rates are observed to be higher for \( g=1+2+3+4+5 \) as compared to \( g=1, g=1+2, g=1+2+3+4, g=1+2+3 \). Also, it is seen that the SISRs are \( \geq 80\% \) for all the model representations, when \( \geq 5 \) sequences per species are used for training (Fig. 3b). Though \( \geq 80\% \) success is achieved even for 4 sequences per species in the training dataset, that is only for \( g=1+2+3+4 \) and \( 1+2+3+4+5 \). Further, SISRs are increased upto 7 sequences per species in the training set, and almost stabilized thereafter (Fig. 3a). The success rates are also found to be more stable, when the prediction model is trained with a large number of sequences (Fig. 3a). The SISRs are further observed to be more stable, when more combinations of g-spaced base-pair features are used in the prediction model (Fig. 3b).

Performance analysis based on k-mer features
In one of our recent studies [26], RF classifier along with k-mer feature was found performing better than the existing machine learning and rule-based approaches [15] for species identification, other than fungi. Thus, we compared the performance between g-spaced and k-mer features. Four different compositions of k-mer \((k=1, 2, 3, \text{ and } 4)\) features were employed here for fungal species identification using RF classifier. Since SISR reached \(~80\%\) when 4 sequences per species were used to train the prediction model (Fig. 3a), the datasets with 5 and 6 sequences per species were only used to compare the SISR of k-mer feature vector with that of model representation \( g=1+2+3+4+5 \). The results of the comparison in terms of SISRs are given in Table 4. The SISRs are observed to be higher for larger combinations of k-mer features. At the same time, the accuracies were also found to be more stable both for k-mer and g-spaced features, when large number of sequence per species were included for training. Though the accuracies for k-mer and g-spaced feature sets are observed at par, the number of features for k-mer are larger than that of g-spaced feature sets. For instance, the number of features for k-mer \( 1+2+3+4 \) is 340 which is much larger than that of \( g=1+2+3+4+5 \) feature set (Table 4). Thus, it may be said that g-spaced features are more efficient in capturing the variability of the nucleotide distribution present in the barcode sequences of fungal species.

Performance analysis in other species
The SISRs of the proposed approach (RF with feature set \( g=1+2+3+4+5 \)) are shown in Fig. 3c. From the figure, it can be seen that the SISRs for other species are much higher (>92%) as compared to that of fungi (<90%). It is further observed that the SISR is low in plant (Inga) than that of others, and this may be due to the fact that except Inga, others are from animal kingdom [45]. It is further noticed that the SISRs in animal and plant species are higher than that of fungi and this may be due to the fact that in fungi ITS regions are used as barcodes which are not highly conserved as that of COI or trnTD [12]. Nevertheless, the SISRs are observed between 92-99%, and thus the proposed approach may be efficiently employed for identification of species other than fungi based on DNA barcode.

Performance analysis using simulated datasets
With the feature set \( g=1+2+3+4+5 \) and RF classifier \((ntree=500, mtry=9)\), the median of the prediction accuracies are observed to be >96% for the effective population sizes 1000 and 10000, whereas it is ~94% for 50000 (Fig. 3d). Further, the prediction accuracies are seen to be declined with increase in the effective population sizes (Fig. 3e). Nonetheless, >90% accuracy are observed in each set for all the three simulated datasets (Fig. 3e).

Comparative analysis for prediction of species other than fungi
The SISRs of the developed model (RF classifier with \( g=1+2+3+4+5 \) features) are ~10% higher as compared to that of similarity-based approaches (Fig. 4a). Further, diagnostic-based method outperformed the similarity- and tree-based approaches, which is corroborated with the results of Weitschek et al. [15]. Though the success rate for the diagnostic-based approach is >90%, it is ~5% less than that of proposed approach (Fig. 4a). Thus, it is inferred that the proposed approach can also achieve higher SISR than that of other ad-hoc methods for prediction of other species.

Comparative analysis for prediction of fungal species
The accuracies of funbarRF and MOTHUR are observed ~89%, which is 2% higher than that of RDP and SINTAX algorithms (Fig. 4b). Further, the stability of the accuracy is found to be highest for RDP and lowest for funbarRF algorithm. It is also seen that 10650 correctly predicted sequences (out of 13630) are common to all the four methods (Fig. 4c). Though the SISRs are seen at par for funbarRF and MOTHUR, number of sequences predicted by funbarRF (370) are distinct from the other
classifiers are higher than that of MOTHUR (141) (Fig. 4c). This implies that the sequences that are not correctly predicted by MOTHUR are also correctly predicted by funbarRF. Thus, the funbarRF can be more efficient than MOTHUR for fungal species identification. Furthermore, ≥99% of the sequences predicted by RDP and SINTAX algorithms are found to be predicted either by MOTHUR or funbarRF or both (Fig. 4c).
Prediction software

Software development is an integral part, as far as the research in the field of computational biology is concerned. Here also, we have established a prediction server “funbarRF” (http://cabgrid.res.in:8080/funbarrf/) for fungal species identification. A snapshot of the server page is shown in Fig. 5a. The user interface of the server was designed using HTML, where the PHP and R-programs were implemented at the back end for execution of the proposed approach. The user has to submit both reference and query sequences in FASTA, with the sequence identifiers in BOLD format. Two result files are generated pertaining to the reference (training) and query (test) sets (Fig. 5b). Number of instances observed and correctly predicted for each reference species are given in training-result-file, whereas the predicted labels

| Feature-type | Feature combination | #Features | #Sequences/Species |
|--------------|---------------------|-----------|--------------------|
| k-mer        | 1+2                 | 20        | 76.37±4.91 79.61±3.33 |
|              | 1+2+3               | 84        | 79.21±4.71 82.72±2.81 |
|              | 1+2+3+4             | 340       | 80.61±4.03 83.68±2.85 |
| g-spaced     | g=1+2+3+4+5         | 96        | 81.74±2.72 83.49±2.36 |

Table 4 Species identification success rates for different combinations of k-mer and g-spaced feature sets, where 4 and 5 sequences per species were used to train the prediction model. It can be seen that though the species identification success rates for both feature sets are at par, number of k-mer features used are larger than that of g-spaced features.

Fig. 4 (a) The SISRs of the proposed model, similarity-, tree- and diagnostic-based methods for taxonomy prediction of Drosophila, Inga and Cypraeidae. b Accuracy of different taxonomy prediction method for prediction of fungal species using DNA barcode. c Number of correctly predicted fungal species that are common in different taxonomy prediction methods.
Fig. 5 (a) Snapshot of the server page of the funbarRF and (b) result page after execution of an example dataset.
for query sequences are shown in test-result-file (Fig. 5b). To facilitate prediction using high throughput sequences, an R-package named as “funbarRF” (https://cran.r-project.org/web/packages/funbarRF/) has also been developed.

**Discussion**

New species identification (taxonomically) is an integral part of biodiversity surveys that are essential for formulating policies to conserve endangered species [38]. DNA barcoding provides an alternative for molecular identification of those micro-organisms for which morphology-based species identification is often difficult [47–49]. In DNA barcoding, one of the fundamental issues is how best one can assign a correct taxonomy to an unknown specimen based on the known taxonomy of the sequences of reference library [15, 46, 50, 51]. Further, commonly used rule-based methods are dependent upon the alignment of the barcode sequences [50]. Though the alignment for coding region like COI is easier, it may not be that much easier for ITS non-coding region due to larger variability in length and indels [51]. This study presents a new computational approach that involves the feature generation based on g-spaced nucleotide base pairs and application of RF for identifying species using DNA barcode, with an emphasis on fungi.

The developed model was evaluated on 3770 fungal species, where the performance was analyzed based on cross validation technique. Though the identity between any two nucleotide sequences in a dataset are generally kept <80% to avoid over estimation while performing classification using machine learning techniques, this pre-processing step is mostly feasible in classification where large numbers of sequences are present in different classes. However, this pre processing step may not be feasible in the present context, because the numbers of sequences in each class (species) are very small and the numbers of classes are also larger (1498 to 3770). In other words, if such (similar) sequences are excluded, the size of the dataset will be reduced further by which the model may not be able to capture the variation present in different classes (species). We also found the similarities between sequences of different classes (species) at threshold 0.8 (results not reported), when the similarity check was performed using CD-HIT program [52]. Thus, we feel that there is a less probability of over-estimation. To the best of our knowledge, we have also not found any earlier studies [15, 16, 18, 19, 26] reporting such pre-processing step, as far as species identification using DNA barcode is concerned.

Five different combinations of g-spaced base pair features were used to encode the barcode sequences that were subsequently used as input in RF classifier for species identification. Higher SISR was found for the training dataset with higher number of sequences per species. This may be due to the fact that with increase in the number of sequences per species, variability present between the species in terms of nucleotide distribution was captured more accurately.

Performances based on g-spaced base pair features were further compared with that of contiguous k-mer features, where the accuracies corresponding to 80 g-spaced base pair features were found similar with that of 340 k-mer features. This implies that, higher number of k-mer features may be required as compared to that of g-spaced features to achieve a certain level of accuracy. Though more features usually lead to a better performance, redundant features often causes misclassification and thereby reduction in classification accuracy [53]. So, one of the probable reasons for the relatively poor performance of $k=1+2+3+4$ as compared to that of $g=1+2+3+4+5$ may be that the k-mer features may have induced more redundancy, which may not be the case in g-spaced base-pair features.

We could not evaluate the proposed model on Warcup and UNITE datasets due to the constraint of computational power. However, the developed model was compared against those which were evaluated on these datasets, and found comparable accuracy for fungal species identification. Thus, the proposed model will certainly complement the prevailing efforts for prediction of fungal species. The developed method was not compared against the Mycofier, as it has been developed for prediction of fungi at genus label. We also did not evaluate the accuracy of the developed model against PROTAX, because we found it difficult to identify the exact feature sets the PROTAX require. Moreover, the PROTAX depends upon the result of multiple sequence alignment of barcode sequences which itself takes longer time.

The developed approach was also assessed for prediction of other species. While evaluated with 5 different taxonomical entites, the proposed model achieved >90% accuracy. Besides, the proposed approach achieved >95% SISR in three diverged taxonomical entities i.e., *Drosophila*, *Inga* and *Cypraidae*, and the same was found much higher than that of rule-based approaches. Furthermore, the proposed method confirmed >90% accuracy with the simulated datasets. Therefore, it may be inferred that the developed technique is not only capable for predicting the fungal species, but also other species as well.

**Conclusion**

This study presents a computational model for prediction of fungal species based on DNA barcode. The developed web server and R-package “funbarRF” will provide a platform for identification of fungi at species label. Besides, it can also be useful for identification of other species. So far so good, the proposed computational model is believed to be helpful for the taxonomists working on fungal species.
Abbreviations
BLAST: Basic local alignment search tool; BOLD: Barcode of life data; CBOL: Consortium for barcode of life; COI: Cytochrome c oxidase subunit I; CV: Cross validation; ITS: Internal transcribed spacer; kNN: k-nearest neighbor; NJ: Neighbor joining; NN: Nearest neighbor; OOB: Out-of-bag; PAR: Parsimony; rDNA: ribosomal DNA; RF: Random forest; SSR: Species identification success rate

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Availability of data and materials
All the datasets used in this study are available at http://cagbgridres.in:8080/funbarf/dataset/

Authors’ contributions
PKM and ARR formulated the problem; SG and RT collected and processed the dataset; PKM, TKS, SG and RT analyzed the dataset; PKM developed the prediction method and wrote the R-codes; TKS and PKM designed the server; PKM, SG and RT, and ARR revised the manuscript; All authors read and approved the final version.

Ethics approval and consent to participate
Not applicable.

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Not applicable.

Competing interests
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References
1. Edgar RC. SINTAX: a simple non-Bayesian taxonomy classifier for 16S and ITS sequences. In: bioRxiv; 2016. https://doi.org/10.1101/074161.
2. Hawkins DL. Fungal diversity and its implications for genetic resource collections. Studies in Mycology. 2004;50:9–18.
3. Roe AD, Rice AV, Bromilow SE, Cooke JE, Spelling FA. Multilocus species identification and fungal DNA barcoding: insights from blue stain fungal symbionts of the mountain pine beetle. Molecular Ecology Resources. 2010; 10(6):946–59.
4. Hebert PD, Cywinska A, Ball SL, deWaal JR. Biological identifications through DNA barcodes. Proceedings of the Royal Society B: Biological Sciences. 2003;270(1512):313–21.
5. Shenoy BD, Jeewon R, Hydev KD. Impact of DNA sequence-data on the taxonomy of anamorphic fungi. Fungal Diversity. 2007;26(1):1–54.
6. Giraud T, Refrégier G, Le Gac M, de Vienne DM, Hood ME. Speciation in fungi. Fungal Genetics and Biology. 2008;45(6):791–802.
7. Somervuo P, Koskela S, Pennanen J, Henriik Nilsson R, Ovaskainen O. Unbiased probabilistic taxonomic classification for DNA barcoding. Bioinformatics. 2016;32(19):2920–7.
8. Das S, Deb B. DNA barcoding of fungi using Ribosomal ITS Marker for genetic diversity analysis: A Review. International Journal of Pure & Applied Bioscience. 2015;3(3):160–7.
9. Ratnasingham S, Hebert PDR. BOLD: The Barcode of life data system available from http://www.barcodinglife.org. Molecular Ecology Notes. 2007; 7:335–64.
10. Hollingsworth PM, Forrest LL, Spouge JL, Hailabaei M, Ratnasingham S, van der Bank M, Chase MW, Cowan RS, Erickson DL, Fazekas AJ. A DNA barcode for land plants. Proceedings of the National Academy of Sciences of USA 2009, 106(31): 12794-12797.
11. Seifert KA. Progress towards DNA barcoding of fungi. Molecular Ecology Resources. 2009;9:82–9.
12. Schoch CL, Seifert KA, Huhndorf S, Robert V, Spouge JL, Levesque CA, Chen W. Fungal Barcoding Consortium: Nuclear ribosomal internal transcribed spacer (ITS) region as a universal DNA barcode marker for Fungi. Proceedings of the National Academy of Sciences of the United States of America. 2012;109(16):6241–6.
13. Killigil U, Nilsson RH, Abaarenk K, Tedesoro L, Taylor AF, Bahram M, Bates ST, Bruns TD, Bengtsson-Palme J, Callaghan TM, et al. Towards a unified paradigm for sequence-based identification of fungi. Mol Ecol. 2013;22(21): 5271–7.
14. Bertolazzi P, Felici G, Weitsche E. Learning to classify species with barcodes. BMC Bioinformatics. 2009:1457.
15. Weitsche E, Fison G, Felici G. Supervised DNA barcodes species classification: analysis, comparisons and results. BioData Mining. 2014;7(1):4.
16. Deshpande V, Wang Q, Greenfield P, Charleston M, Ponnas-Affaro A, Kuke CR, Cole JR, Midgley DJ, Tran-Dinh N. Fungal identification using a Bayesian classifier and the Warcup training set of internal transcribed spacer sequences. Mycologia. 2016;108(1):1–5.
17. Wang Q, Gartty GM, Tiedje JM, Cole JR. Naive Bayesian classifier for rapid assignment of rRNA sequences into the new bacterial taxonomy. Appl Environ Microbiol. 2007;73(16):5261–7.
18. Delgado-Serrano L, Restrepo S, Bustos JR, Zambrano MM, Arzola JML. MycoFiser: a new machine learning-based classifier for fungal ITS sequences. BMC Res Notes. 2016;9(1):402.
19. Schloss PD, Westcott SL, Raybin T, Hall JR, Hartmann M, Hollister EB, Lesniewski RA, Oakley BB, Parks DH, Robinson CJ, et al. Introducing mothur: open-source, platform-independent, community-supported software for describing and comparing microbial communities. Appl Environ Microbiol. 2009;75(23):7537–41.
20. Altschul SF, Gish W, Miller W, Myers EW, Lipman DJ. Basic local alignment search tool. J. Mol. Biol. 1990;215:403–10.
21. Govindan G, Nair AS. New feature vector for apoptosis protein subcellular localization prediction. Advances in Computing and Communications. 2011;190:294–301.
22. Breiman L. Random forests. Machine Learning. 2005;64:5–32.
23. Sarkar IN, Trizna M. The Barcode of Life Data Portal: bridging the biodiversity informatics divide for DNA barcoding. PLoS One. 2011;6(7):e14689.
24. Kamath U, De Jong K, Shelu A. Effective automated feature construction and selection for classification of biological sequences. PLoS ONE. 2014;9(7):e99982.
25. Zhan X, Lee J, Chasin LA. The effect of nonsense codons on splicing: a genomic analysis. RNA. 2006;12(6):637–9.
26. Meher PK, Sahu TK, Rao AR. Identification of species based on DNA barcode using k-mer feature vector and Random forest classifier. Gene. 2016;592(2): 316–24.
27. Bhinda K, Sukulski M, Kuchero V. Spaced seeds improve k-mer-based metagenomic classification. Bioinformatics. 2015;31(22):3584–92.
28. Hong L. BioSeqClass: Classification for biological sequences. In: R package version 1.32.0, 2016.
29. Platt JC. In: Scholkopf B, Burges C, Platt J, Smola AJ, editors. Fast Training of support vector machines using sequential minimal optimization, Advances in Kernel Methods - Support Vector Learning. Cambridge MA: MIT Press; 1998. p. 185–208.
30. Quinlan R. C4.5: Programs for machine learning. In: Morgan Kaufmann Publishers. San Mateo CA: Morgan Kaufmann; 1993.
31. Cohen WW. Fast effective rule induction. Twelfth International Conference on Machine Learning (ICML). 1995;95115–23.
32. John GH, Langley P. Estimating continuous distributions in Bayesian classifiers. Eleventh Conference on Uncertainty in Artificial Intelligence, San Mateo, CA: Morgan Kaufmann. 1995:338–45.
33. Chaudhary A, Kolhe S, Kamal R. An improved random forest classifier for multi-class classification. Information Processing in Agriculture. 2016;3(4):215–22.
34. Breiman L, Friedman JH, Olshen RA, Stone CJ. Classification and Regression Trees. CRC Press. 1984.
35. Meher PK, Sahu TK, Rao AR. Prediction of donor splice sites using random forest with a new sequence encoding approach. BioData Mining. 2016:94.
36. Liaw A, Wiener M. Classification and regression by randomForest. R News. 2002;2(3):18–22.
37. Henderson J, Salzberg S, Fasman KH. Finding genes in DNA with a Hidden Markov Model. Journal of Computational Biology. 1997;4(2):127–41.
38. Van Velzen R, Wetschek E, Felici G, Bakker FT. DNA barcoding of recently diverged species: relative performance of matching methods. PloS ONE. 2012;7(1):e30490.
39. Farris JS. Estimating phylogenetic trees from distance matrices. The American Naturalist. 1972;106(951):645–68.
40. Saitou N, Nei M. The neighbor-joining method: a new method for reconstructing phylogenetic trees. Molecular Biological Evolution. 1987;4(4):406–25.
41. Austerlitz F, David O, Schaeffer B, Bleakley K, Olteanu M, Leflois R, Veullie M, Laredo C. DNA barcode analysis: a comparison of phylogenetic and statistical classification methods. BMC Bioinformatics. 2009;14:510.
42. Altschul SF, Madden TL, Schäffer AA, Zhang J, Zhang Z, Miller W, Lipman DJ. Gapped BLAST and PSI-BLAST: a new generation of protein database search programs. Nucleic Acids Research. 1997;25(17):3389–402.
43. DasGupta B, Konwar KM, Mandou Ii, Shvartsman A. DNA-BAR: distinguisher selection for DNA barcoding. Bioinformatics. 2005;21(16):3424–6.
44. Wetschek E, Van Velzen R, Felici G, Bertolazzi P. BLOG 2.0: a software system for character-based species classification with DNA Barcode sequences. What it does, how to use it. Molecular Ecology Resources. 2013;13(6):1043–6.
45. Dinca V, Zakharov EV, Hebert PD, Vila R. Complete DNA barcode reference library for a country’s butterfly fauna reveals high performance for temperate Europe. Proceedings of the Royal Society B. Biological Sciences. 2011;278(1704):347–55.
46. Tanabe AS, Toju H. Two new computational methods for universal DNA barcoding: a benchmark using barcode sequences of bacteria, archaea, animals, fungi, and land plants. PloS One. 2013;8(10):e76910.
47. Hibbett DS, Ohman A, Glotzer D, Nuhn M, Kirk P, Nilsson RH. Progress in molecular and morphological taxon discovery in Fungi and options for formal classification of environmental sequences. Fungal Biology Reviews. 2011;25(1):38–47.
48. Bachy C, Dolan JR, López-García P, Deschamps P, Moreira D. Accuracy of protist diversity assessments: morphology compared with cloning and direct pyrosequencing of 18S rRNA genes and ITS regions using the conspicuous tintinnid ciliates as a case study. ISME Journal. 2013;7(2):244–55.
49. Toju H, Yamamoto S, Sato H, Tanabe AS, Gilbert GS, Kadokawa K. Community composition of root-associated fungi in a Quercus-dominated temperate forest: co-dominance of mycorrhizal and root-endophytic fungi. Ecology and Evolution. 2013;3(5):1281–93.
50. Zhang AB, Savolainen P. BPSI2.0: A C/C++ Interface program for species identification via DNA barcoding with a BP-Neural Network by calling the Matlab engine. Molecular Ecology Resources. 2008;9(1):104–6.
51. Zhang AB, Feng J, Ward RD, Wan P, Gao Q, Wu J, Zhao WZ. A new method for species identification via protein-coding and non-coding DNA barcodes by combining machine learning with bioinformatic methods. PloS One. 2012;7(2):e30986.
52. Huang Y, Niu B, Gao Y, Fu L, Li W. CD-HIT Suite: a web server for clustering and comparing biological sequences. Bioinformatics. 2010;26:680–2.
53. Baten A, Halgamuge SK, Chang B, Li J. Splice site identification using probabilistic parameters and SVM classification. BMC Bioinformatics. 2006;7:1–15.