Efficient classification of sugarcane genomes

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Abstract
A Phylogenetic tree construction to know to the relationship of the ancestral association of species. The genome sequences, outlining the transmission of functional and genetic classification. Analysing the quantitative conduct of phylogenetics in the conservation of biodiversity and the successful heuristics of obtaining an accurate distribution of trees plays a predominant role. The study to know higher accuracy from efficient algorithm to deducing phylogenetic relationship among Sugarcane (Saccharum) species. A sample of 431 Saccharum genome sequences was drawn from NCBI dataset. Efficient algorithms like Maximum Likelihood Estimation (MLE) method and Unweighted Pair Group Method with Arithmetic Mean (UPGMA) method were considered to construct the phylogenetic tree. The maximum likelihood with Tamura Nei model, Kimura 2-parameter model and achieves the highest precision, while MLE with Maximum likelihood with Jukes-Cantor model achieves the least. The computational biology of statistically results is justifiable and compares the functional relationship between various models in which error percentage has been reduced. The same algorithms perform on individual species under different models such as maximum likelihood with Kimura 2-parameter model and Tamura Nei model more efficient than others to differentiate the species genomic sequences and group them to correct taxon.

Keywords: MLE (Maximum Likelihood Estimation), Phylogenetic tree, UPGMA (Unweighted Pair Group Method with Arithmetic Mean)

1. Introduction
“Amongst the sugarcane we are safe” say the Chinese with sugarcane (Saccharum spp.) symbolizing bravery, independence and protection (DeBernardi, 2009) [3]. The name sugarcane is used to refer to a group of tall perennial tropical grass species which were domesticated for sugar production, and have been classified inconsistently (Paterson et al., 2013) [17]. Sugarcane (Saccharum spp. hybrids) is the most important sugar-producing crop and raw biofuel material in the world (Zhou et al., 2018) [22], which belongs to the grass family Gramineae. The major commercial sugarcane cultivars are all complex interspecies hybrids with allopolyploidy and aneuploidy with a huge genome (Zhou et al., 2014; Garsmeur et al., 2018) [21, 8]. Sugarcane has been known for more than 2,200 years and it was one of the first plants to inspire humans to develop technology (Goldstein and Mintz, 2015) [9]. Sugarcane (Saccharum officinarum) family Gramineae (Poaceae) a widely grown crop in India. It employs over a million people, directly and indirectly contributing to the national exchequer. Sugar cane originated in New Guinea but Sugarcane plants spread along with human migration routes to the Indian subcontinent and other countries. Now cross with some wild sugarcane species produced commercial sugarcane with high sugar contents. Cultivation of sugarcane in India from Vedic period around 1400 to 1000 B.C. It is now widely accepted that India is the original home of Saccharum species. It belongs to family Gramineae, class monocotyledons and order grumaceae subfamily panicoidae, tribe andropogoneae, and sub tribe saccharinin.

1.1. Genome sequencing is the process of determining the complete DNA sequence of an organism's genome at a single time. This involves sequencing all of a chromosomal DNA as well as DNA confined in the mitochondria and chloroplast. Need for the study of the phylogenetic tree is that in recent days almost all evolutionary relationships are inferred from molecular sequence data. Introduced a space of phylogenetic bushes and described the space among two timber to be the length of the shortest path between them in that area (Billera et al., 2001) [3]. Reported phylogenetic inference in building a tree-based at the measured pairwise distance of all species (Fisher and Lindberg, 2006) [7]. As we know that the DNA is the inherited material can analyse easily, quickly, and inexpensive with less time and researchers can also investigate the phylogeny to determine their rates of evolution and can also study the direction of evolution by determining were a disease organism, such as bacterium or virus.
Where in the case, where it is not possible to obtain genetic material, the morphological measurement can be used to infer evolutionary relationships. However, this approach is less reliable than using molecular data because as we know that sometimes the same morphological trait can arise from multiple independent evolutionary lineages and also in bioinformatics phylogenetic tree of proteins and DNA are useful and the multiple sequence alignment, structure conservation, homology detection, and paralogous attributes estimation could be as close to the true value. The evidence presented ancestrs and the phylogenetic is used to assess DNA evidence presented in court cases to inform situations.

2. Material and methods

To acquire the nucleotide sequence of Saccharum species were sought over the database NCBI. The data sequences collected in FASTA format and aligned in a mega format. Total 431 sequences were taken from 11 Saccharum species for the tree construction and analysis. The 11 Saccharum species utilized for the investigation are represented in table1.

### Table 1: The 11 Saccharum species utilized for the investigation

| Sl. No. | Species                     | Sequence Total |
|--------|-----------------------------|----------------|
| 1      | S. arundinaceum             | 40             |
| 2      | S. asper                    | 31             |
| 3      | S. barberi                  | 40             |
| 4      | S. edule                    | 40             |
| 5      | S. officinarum              | 40             |
| 6      | S. procernum                | 40             |
| 7      | S. ravennae                 | 40             |
| 8      | S. robustum                 | 40             |
| 9      | S. sinese                   | 40             |
| 10     | S. spontaneum               | 40             |
| 11     | S. villosum                 | 40             |
| Total  |                            | 431            |

2.1 Unweighted Pair Group Method with Arithmetic Mean

The UPGMA (Unweighted Pair Group Method with Arithmetic Mean) algorithm, which is mainly used in bioinformatics for constructing the evolutionary tree and where in this UPGMA method is the simple agglomerative or hierarchical clustering method for the development of constructing taxonomic phenograms, which replicate the phenotypic similarities between the operational taxonomical units (OUTs). Phenogram is created by considering at each step, then the nearest two clusters are pooled into a higher-level cluster. Let the clusters be A and B, then the distance between the clusters is taken to be the average of all distances that is “x” in A and “y” in B, hence the mean distance between the cluster:

\[
\frac{1}{|A||B|} \sum_{x \in A} \sum_{y \in B} d(x,y)
\]

The maximum likelihood and parsimony algorithms, the phylogenetic tree was Built under UPGMA.

2.2 Maximum likelihood algorithm

In a statistical way that the maximum likelihood estimation (MLE) is a method of estimating the parameters of a given observation. Let us assume that there is a sample of \( x_1, x_2, x_3, \ldots, x_n \) of N independent and identically distributed observations, which are obtained from the distribution of unknown probability density function \( f_\theta(.) \). Summarized that the function \( f_\theta \) belongs to a certain family of distributions \( \{ f(\cdot, \theta) : \theta \in \Theta \} \) (where \( \theta \) is a vector of parameters) called the parametric model, so that \( f_\theta = f(\cdot, \theta_0) \). The value \( \theta_0 \) is the unknown parameter which is referred to as the true value of the parameter vector. It is necessary to estimate the estimator which would be as close to the true value \( \theta_0 \) as possible. Either or both the observed. Variables \( x_i \) and the parameter \( \theta \) can be the vectors. Before calculating the maximum likelihood estimation, one should specify the joint density function for all the observations of independent and identically distributed sample hence, the joint density function is

\[
f(x_1, x_2, x_3, \ldots, x_n | \theta) = f(x_1 | \theta) \times f(x_2 | \theta) \times \ldots \times f(x_n | \theta)
\]

Then the likelihood function that is the log-likelihood is given by,

\[
\ln L(\theta; x_1, x_2, x_3, \ldots, x_n) = \sum_{i=1}^{n} \ln f(x_i | \theta)
\]

And the average log-likelihood is:

\[
\bar{l} = \frac{1}{n} \ln L
\]

Where, \( \bar{l} \) indicates that an estimator, certainly, the \( \bar{l} \) estimates the expected log likelihood for the single observation in the model. Hence, by finding the value of \( \theta \) the maximum likelihood estimates \( \theta_0 \) can be estimated which maximizes \( \bar{l} = (\theta; x) \). Hence, this method of estimation defines a maximum likelihood estimator (MLE) of \( \theta_0 \),

\[
\hat{\theta} = \left\{ \arg \max_{\theta \in \Theta} \bar{l}(\theta; x_1, x_2, x_3, \ldots, x_n) \right\}
\]

If maximum exists, then the MLE estimate is the same irrespective of maximizing the likelihood or the log-likelihood function, because the log is a monotonically increasing function.

2.3 Distance-based method

Mainly the distance models are used in constructing a phylogeny as non-parametric distance method. Using the matrix of pair-wise distances, non-parametric distances were initially applied to phonetic data. There are various pair-wise distances, such as Euclidean distance which is applied to discrete morphological characters. For the Constructed phylogenetic tree, the following distance-based models were used: Jukes-Cantor model, Kimura 2-parameter model, Tajima-Nei model and P distance model.
2.3.2 Kimura 2-parameter model
This model basically differentiates between transitions and transversion, that is a conversion of A to G, i.e., from purine to pyrimidine, or C to T, i.e., from pyrimidine to pyrimidine which is called as transition and whereas when the conversion from a purine to pyrimidine or vice versa then it is called as transversion. In Kimura's original description of the model, they \( \alpha \) and \( \beta \) were used to denote the rates of these types of substitutions, but it is more usual that the rate of transversions is set to 1 and the \( K \) is used to indicate the transition transversion rate ratio (as indicated below). Even the Kimura 2-parameter model assumes that all of the bases are equally frequent.

\[
\pi_A = \pi_G = \pi_U = \pi_T = 0.25
\]

Rat matrix \( Q = \begin{pmatrix}
* & k & 1 & 1 \\
3 & k & 1 & 1 \\
1 & 1 & * & k \\
1 & 1 & k & *
\end{pmatrix}
\]

The Kimura two-parameter distance is given by:

\[
K = -\frac{1}{2} \ln \left(1 - 2p - q \right) \sqrt{1 - 2p}
\]

Where, \( p \) is the proportion of sites which indicates the transitional differences and \( q \) is the proportion of sites that show transversional differences.

2.3.3 Tamura-Nei model
In this model by considering the differences in substitution rate between nucleotides and the inequality of nucleotide frequencies, (Tamura-Nei., 1993). Mainly it differentiates between transitional substitutions rates between purines and trans versional substitution rates between pyrimidines. It also assumes equality of substitution rates among nucleotide sites. The Tamura-Nei model is:

\[
\begin{align*}
A & = - \beta_{gt} & & \beta_{gc} & & \beta_{gc} \\
T & = \beta_{ga} & & - \alpha_{2gc} & & \beta_{gc} \\
C & = \beta_{ga} & & \alpha_{2gt} & & - \beta_{gc} \\
G & = \alpha_{1ga} & & \beta_{gt} & & \beta_{gc} & & -
\end{align*}
\]

Where,
\[
\beta = \text{Transitional frequencies between two sequences}
\]
\[
\alpha = \text{Transversion frequencies between any two sequences}
\]
\[
d = -k_1 \ln(w_1) - k_2 \ln(w_2) - k_1 \ln(w_3)
\]
\[
s = -k_1 \ln(w_1) - k_2 \ln(w_2) - (k_3 - 2g_R g_y) \ln(w_3)
\]
\[
v = -2g_R g_y \ln(w_3)
\]
\[
R = s/v
\]

\( P_1 \) and \( P_2 \) are the proportions of transitional differences between nucleotides between purines and pyrimidines i.e., is between A to G and T to C respectively, \( Q \) is the Proportion of transversional differences, \( g_A', g_G', g_C', g_T' \), are the respective frequencies of A, C, G, and T.

\[
W_1 = 1 - P_1/K_1 - Q/2g_R, \quad W_2 = 1 - P_2/K_2 - Q/2g_Y, \quad W_3 = 1 - Q/2g_R
\]
Var (d) = \[c_1^2 P_1 + c_1^2 P_2 + c_1^2 Q - (c_1 P_1 + c_2 P_2 + c_4 Q)^2\] / L
Var (s) = \[c_1^2 P_1 + c_1^2 P_2 + c_1^2 Q - (c_1 P_1 + c_2 P_2 + c_5 Q)^2\] / L
Var (v) = \[c_1^2 Q (1 - Q)\] / L
Var (R) = \[c_1^2 P_1 + c_1^2 P_2 + c_1^2 Q - (c_1 P_1 + c_2 P_2 + c_6 Q)^2\] / L
\[c_1 = 1/\omega_1,\quad c_2 = 1/\omega_2,\quad c_3 = 1/\omega_3,\quad c_4 = k_1 c_1 / 2 g_p + k_2 c_2 / 2 g_v + k_3 c_3 / 2 g_v - c_3,\quad c_5 = c_1 / v,\quad c_6 = c_1 / v,\quad c_7 = c_2 / v,\quad c_8 = (c_5 c_3) / v\]

\[Var (R) = \frac{e}{v (L - v)} / L,\quad R = s / v\]

Where,
- d: Transitions + Transversions: Number of different nucleotide sites.
- s: Transitions only: Number of nucleotide sites with transitional differences.
- v: Transversions only: Number of nucleotide sites with transversional differences.
- R = s / v: Transition/transversions ratio.
- L: No of valid common sites: Number of compared sites.
- P and Q are the proportion of sites showing transversional and transversional differences, respectively.

Calculating F – measure through information retrieval approach: The information retrieval methods works on the two-way contingency table prepared form in the following form.

| Actual class | True Positive (TP) | False Negative (FN) | False Positive (FP) | True Negative (TN) |
|--------------|-------------------|---------------------|---------------------|-------------------|
|              |                   |                     |                     |                   |

For classification works, the term true positives, true negatives, false positives and false negative compare results of the classifier under test with trusted external judgements. The term positives and negatives refer to the observation and true and false refer to the expectation. These can be explained by the given formula:

\[Precision = \frac{TP}{TP + FN} \times 100,\quad Recall = \frac{TP}{TP + FN} \times 100\]

\[Accuracy = \frac{TP + TN}{TP + FN} \times 100\]

F-Measure (Probabilistic interpretation)

F- Measure is the harmonic mean between the precision and the recall, hence it can be interpreted by considering the two probabilities namely:

1. Precision is considered as the probability of a retrieved document is significant.
2. The recall is considered as the probability of relevant document which is retrieved.

\[F = 2 \times \frac{Precision \times Recall}{Precision + Recall} \times 100\]

Using the given methods, a total of 4 phylogenetic trees were constructed and results are shows in results section. Also, the information retrieval methods are applied to find the best classification among the procedure employed.

3. Results and discussion

Constructing phylogenetic tree: The construction of the phylogenetic tree for the examination includes the acquisition of the sugarcane genome sequences from the NCBI dataset with the FASTA and MEGA format, and the phylogenetic tree constructed for the given algorithms was clarified under the following headings.

Table 2: Classification measures for various distance models.

| S. No. | Models | True Positive | False Positive | True Negative | False Negative |
|--------|--------|---------------|----------------|---------------|----------------|
| 1      | MLE (Jukes Cantor) | 20427         | 2769           | 64202         | 5698           |
| 2      | MLE (Tamura Nei model) | 79936         | 7145           | 1322          | 4693           |
| 3      | MLE (Kimura 2 Parameter) | 65161         | 5597           | 19468         | 2870           |
| 4      | UPGMA (P distance model) | 82092         | 969            | 2485          | 7550           |

Table 3: Classification measures for various distance models.

| S. No. | Models | Accuracy (%) | Precision (%) | Recall (%) | F-Measure (%) |
|--------|--------|--------------|---------------|------------|---------------|
| 1      | MLE (Jukes Cantor) | 90.90         | 88.06         | 78.18      | 82.83         |
| 2      | MLE (Tamura Nei model) | 88.54         | 91.79         | 94.45      | 93.10         |
| 3      | MLE (Kimura 2 Parameter) | 90.90         | 92.08         | 95.78      | 93.89         |
| 4      | UPGMA (P distance model) | 90.84         | 98.83         | 91.57      | 95.06         |

The tree obtained for the aggregate 431 sugarcane genome sequencing of 11 distinctive Saccharum species for MLE with Tamura Nei distance measure has appeared in Fig 1. From Table 2, it was observed that they had been classified adversely with the 79936 true positives, 7145 false positives, 1322 true negative and 4693 false negatives. In view of this classification, the evolutionary measures are computed and from the Table 3, it was observed that the accuracy is 88.54 per cent, precision is 91.79 per cent, recall with 94.45 per cent and the harmonic mean between the precision and the recall that is F-measure is 93.10 per cent.
The present investigation it was inferred that the maximum likelihood with Kimura 2-parameter model and Tamura-Nei model give the minimum measure value with least accuracy. In the study of Confusion matrix that is 11×11 contingency table from Table 4 for the individual species under a different model in this model Saccharum villosum from Table 6 give the high F-measure value.

4. Conclusion
The examination demonstrated that maximum likelihood with Kimura 2-parameter model and Tamura-Nei model gives the highest results while Maximum likelihood with Jukes-Cantor model give the minimum. F measure will take the greatest functional relationship percentage between different models demonstrates the best outcomes where we can compare computational biology of statistical outcomes with the high F measure of 11 species of sugarcane with different method.

5. Future line of work
Phylogenetic trees have already witnessed applications in numerous practical domains, such as in conservation biology.
(illegal whale hunting), epidemiology (predictive evolution), forensics (dental practice HIV transmission), gene function prediction and drug development. Other applications of phylogenies include multiple sequence alignment, protein structure prediction, gene and protein function prediction, and drug design.

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