Evaluating DNA barcoding using five loci (matK, ITS, trnH- psbA, rpoB, and rbcL) for species identification and phylogenetic analysis of Capsicum frutescens

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ABSTRACT

The Capsicum genus comprises several species that are closely related to each other in terms of morphology. This usually leads to species misidentification when relying solely on their physical traits. Due to this, it becomes necessary to develop a method that will authenticate and identify the species, for which the present study was conducted to develop barcodes for Capsicum frutescens utilizing five different genetic loci (matK, ITS, trnH-psbA, rpoB, and rbcL) and comparing it with other Capsicum species. The present study evaluates the delimiting effect of these barcodes in effectively identifying C. frutescens and distinguishing between the three Capsicum species (annuum, frutescens, and chinense). It will be a unique study that will develop barcoding data using five genetic markers for C. frutescens. After analysis, it was found that the combination of rbcL + ITS proved to be a good barcode to differentiate between the three morphologically similar sisters Capsicum species, while matK, rbcL, and rpoB genetic loci were suitable for identifying C. frutescens. The present work will be a strong base, on which molecular identification of C. frutescens can be performed, and it will also aid in differentiating the morphologically close resembling “annuum-frutescens-chinense” group.

1. INTRODUCTION

DNA barcoding is an indicator used to build databases of DNA sequences that can be further exercised as DNA markers intended for “authentication” of the plant, along with studying the plant characters. DNA barcoding was originally based on differences in the mitochondrial Cytochrome c oxidase (cox-I) gene sequences, which successfully enabled the recognition of various flora and fauna [1]. However, it was later realized a need to establish other genes as barcodes as the cox-I gene was found to be less effective in identifying flowering plants [2]. Plant working group (PWG) has approved matK + rbcL as primary barcodes with ITS2 as one of the additional loci for distinguishing plants at the Third International Barcoding Conference, the Consortium for the Barcode of Life [3].

The family Solanaceae consists of a wide range of flowering plants with approximately 2,500 different plant species with significant benefits for humanbeings in terms of cost and productivity. This family includes many commercially valuable plants, Nicotiana tobacco, Capsicum annuum, Solanum lycopersicum, and Solanum melongena which are a few of them [4]. The genus Capsicum comprises the five most domestically cultivated species out of the overall 37 known species. These includes C. annuum L., C. frutescens L., C. chinense Jacq., C. pubescens., and C. baccatum L [5]. The identification of all these species is usually made on the basis of the differences in their morphological characteristics, such as the shape of the fruit, color of the corolla, length of pedicel, and the number of growth at each pedicel [5,6]. Unfortunately, the Capsicum species annuum, frutescens, and chinense appear similar in their physical characters, making it difficult to distinguish between them morphologically. As most of the plants, physical characteristics depend on various environmental factors. The three species are commonly known as the “annuum-chinense-frutescens” group due to their close association [1,5]. Various attempts have been made to characterize different Capsicum species utilizing their morphological traits, enzyme loci, restriction fragment length polymorphism, random amplified polymorphic DNA (RAPD), amplified fragment length polymorphism, direct or directed amplification of minisatellite region DNA amplified using the polymerase chain reaction (DAMD-PCR), cleaved amplified polymorphic sequence, simple sequence repeat length polymorphism, and inter-simple sequence repeats to determine genetic variations between and within different Capsicum species [7]. An effort was also made to distinguish C. frutescens from C. chinense using RAPD markers, where it was suggested C. frutescens and C. chinense to be

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different species [8]. However, a strong base describing the molecular diversity between these three closely resembling Capsicum species employing different markers used in DNA barcoding is yet to be established. Along with this, a strong genetic base that can be used to identify C. frutescens is still missing and needs to be addressed. With this as an objective, the present research is carried out to characterize C. frutescens using five different DNA markers.

The current research aimed to amplify matK, rbcL, ITS, rpoB, and trnH-psbA loci from C. frutescens and identify the best suitable barcode for identification of C. frutescens. To construct, a phylogenetic tree of the sequences obtained and comparing with the sequences of other plants belonging to Capsicum species and analyzing the delimitation strength of these barcodes. The study will assist in authenticating C. frutescens, along with its identification. It will also give an idea about the usefulness of these five genetic loci in differentiating between the three Capsicum species: annuum, frutescens, and chinense.

2. METHODS AND MATERIALS

2.1. Sample Collection

The seeds of C. frutescens were collected from Thiruvananthapuram, Kerala (8° 31’ 26.9004” N 76° 56’ 11.8968” E) India. The seeds were then grown in the Institute of Science Garden Mumbai, and fresh leaves were utilized for DNA isolation.

2.2. DNA Extraction

Genomic DNA extraction was carried out from young leaves using the cetyltrimethylammonium bromide (CTAB) method with few modifications [9]. The leaves were pulverized in a mortar using liquid nitrogen, and the ground leaves were then mixed with a CTAB isolation buffer. The quality of DNA extracted was, then, checked in 0.8% agarose gel. The extracted DNA was then quantified using NanoDrop Spectrophotometer (Thermo Fischer).

2.3. PCR Amplification and Sequencing

PCR was performed to amplify five genetic markers from C. frutescens which included two intergenic spacer regions (trnH-psbA, rpoB-trnCGAR), two plastid gene regions, namely, Ribulose 1, 5 bisphosphate (RbcL), and matK gene and a nuclear region, namely, inter transcribed spacer (ITS). The PCR was carried out using 25 µL reaction volume with each reaction mixture containing 90–100 ng of DNA template, 2.5 U Taq Polymerase enzyme (ThermoFischer), 2.5 mM MgCl2, 1X Taq DNA polymerase buffer, 0.2 mM dNTPs, and 0.5 µM primers ordered from Eurofins Genomics India Pvt. Ltd. Primer sequence used for amplifying different DNA regions of the gene is mentioned in Table 1, and their annealing temperatures are given in Table 2. The amplified genes were then separated on 1.5% agarose gel stained with 0.1 µg/ml ethidium bromide using 1X TAE buffer. The separated amplicons were later purified further to remove any possible contamination using Qiagen PCR clean-up kit. The samples were then, sent for sequencing at Eurofins Pvt. Ltd. (India).

2.4. DNA Sequencing and Sequence Analysis

Post amplification, the samples were analyzed on agarose gel electrophoresis and were sequenced using the Sanger sequencing method. The nucleotide sequences obtained for C. frutescens were then compared to sequences available in NCBI using basic local alignment search tool (BLAST). The sequences obtained were later aligned using multiple sequence alignment with hierarchical clustering [10] and edited using Finch TV.

### Table 1: List of primers used for PCR amplification.

<table>
<thead>
<tr>
<th>Gene</th>
<th>Primer sequence</th>
<th>Primer reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>ITS</td>
<td>Forward primer</td>
<td>TCCGTAGGTAACCTGC</td>
</tr>
<tr>
<td></td>
<td>Reverse primer</td>
<td>TCTCCGTTATGATATG</td>
</tr>
<tr>
<td>matK</td>
<td>Forward primer</td>
<td>CGATCTATTCATTCAAT</td>
</tr>
<tr>
<td></td>
<td>Reverse primer</td>
<td>AGTTCACACGAAGAGTCG</td>
</tr>
<tr>
<td>rbcL</td>
<td>Forward primer</td>
<td>GTAAATCAAGTCCACCRG</td>
</tr>
<tr>
<td></td>
<td>Reverse primer</td>
<td>CACCRCATTYGAACTGG</td>
</tr>
<tr>
<td>trnH-psbA</td>
<td>Forward primer</td>
<td>GTATAGCATGAACCTGCTG</td>
</tr>
<tr>
<td></td>
<td>Reverse primer</td>
<td>CGCGCATGGGATTCGGC</td>
</tr>
<tr>
<td>rpoB</td>
<td>Forward primer</td>
<td>CKACAAAAAYCCTCRAATTG</td>
</tr>
<tr>
<td></td>
<td>Reverse primer</td>
<td>CACCRCATTYGAACTGG</td>
</tr>
</tbody>
</table>

Nucleotide sequence divergences were calculated pair-wise using the Kimura-2-parameter model, and the distance matrices were generated using a maximum likelihood (ML) tree with 1000 bootstrap replications and also parameters such as conserved sites, variables sites, parsimony informative sites, and GC content of the sequences were estimated using molecular evolutionary genetic analysis (MEGA) X software [11].

3. RESULTS

3.1. PCR Result and Quality Analysis of DNA Sequences

All the five regions of C. frutescens were successfully amplified by PCR and found to be good quality. The result indicates that the primer set utilized for the study was able to amplify respective regions and gave a sharp band on Agarose gel electrophoresis, which is required for successful DNA sequencing. All the five nucleotide sequences obtained for C. frutescens were submitted to the NCBI Gene Bank database. Their accession numbers are given in Table 3.

### Table 2: PCR reaction conditions with respective annealing temperatures utilized for DNA amplification.

<table>
<thead>
<tr>
<th>Locus</th>
<th>Length of amplicons (base pairs)</th>
<th>Tm (degree)</th>
<th>Cycle conditions</th>
</tr>
</thead>
<tbody>
<tr>
<td>ITS</td>
<td>548</td>
<td>58°C</td>
<td>95°C/1</td>
</tr>
<tr>
<td>matK</td>
<td>862</td>
<td>54°C</td>
<td>32.8</td>
</tr>
<tr>
<td>rbcL</td>
<td>993</td>
<td>58°C</td>
<td>72°C/7</td>
</tr>
<tr>
<td>trnH-psbA</td>
<td>553</td>
<td>58°C</td>
<td>4°C/O</td>
</tr>
<tr>
<td>rpoB</td>
<td>183</td>
<td>58°C</td>
<td></td>
</tr>
</tbody>
</table>

3.2. Sequence Analysis

All the five genes nucleotide sequences (matK, rbcL, ITS, rpoB, and trnH-psbA) obtained for C. frutescens were aligned with the retrieved sequences of Capsicum annuum and Capsicum chinense (see Supplementary material) obtained from NCBI database. It was found that ITS was able to differentiate between all the three species of Capsicum most efficiently. The retrieved sequences varied in length from 548 bp for C. frutescens to 653 bp for C. chinense and...
636 bp for *C. annuum*. After alignment, it was observed that both *C. frutescens* and *C. annuum* showed gaps from position 100–107 and from position 111–115 along with a gap was also observed at position 254 when compared to *C. chinense*. However, interestingly, at position 251, all the three species of *Capsicum* showed different nucleotides, where *frutescens* showed the presence of guanine, while adenine was found for *chinense*, whereas *annuum* showed presence of cytosine nucleotide. It was seen that ITS sequence of *C. frutescens* showed 64% of G + C content. Conserved sites for ITS were found to be 458; variables were 105, parsimony informative sites were 17, and singleton sites were 17.

Sequence alignment of matK showed certain dissimilarities in many regions, but there were two specific regions at positions 7 and 913, where *C. frutescens* showed different nucleotides compared to the other two species. At position 7, *C. frutescens* showed substitution of thymine nucleotide, while other two species showed presence of cytosine nucleotide, while at position 913 *frutescens* showed the presence of cytosine, while the other two species showed a gap at the same position. Alignment of the matK sequences also showed that species *frutescens* and *chinense* are closely related as they both showed gaps at similar locations, while *annuum* did not show any gaps at these locations. matK sequence of *C. frutescens* showed 845 conserved sites and 16 variable regions, while no parsimony informative sites or singleton sites were observed with 32.8% of G + C content.

The aligned sequences of rbcL showed *frutescens* to have a substitution at position 573 with guanine nucleotide, while the other two species showed the same nucleotide adenine at that position. While the alignment also showed three other positions, where *C. chinense* showed substitutions. rbcL sequence of *C. frutescens* showed no parsimony-informative and singleton sites, while it had 989 conserved sites and four variable sites with 43.3% G + C content.

Next barcode analyzed was trnH-psbA; the aligned trnH-psbA nucleotide sequence of all the three *Capsicum* species displayed a notable change at certain positions. It was observed that *C. frutescens* and *chinense* showed a gap at position 360 while, interestingly, the *annuum* did not show any gap at this position. *C. frutescens* trnH-psbA sequence showed 27.4% G + C content and had no variable sites, parsimony informative sites or singleton sites with 553 conserved sites. While nucleotide alignment of rpoB showed a gap at position 5 in *C. frutescens*, while the other two species showed that the presence of adenine nucleotide along with this *C. frutescens* also showed substitution of nucleotides at four more positions, while *chinense* and *annuum* showed similar nucleotides at the same position. rpoB sequence of *C. frutescens* showed the presence of 175 conserved sites, nine variable sites, and singleton sites with no parsimony informative sites with 33.3% G + C content.

### 3.3. Phylogenetic Analysis

A BLAST was performed on all the five genetic marker sequences obtained for *C. frutescens* and were aligned using Multiple Sequence Comparison by Log-Expectation in the MEGA X. The nucleotide sequences of *Capsicum* genus were used to evaluate the evolutionary relationship using ML method with a 1000 replications bootstrap resampling. Outgroup belonging to the Solanaceae family were selected for all analyses. The first genetic marker studied was matK [Figure 1]. All the *Capsicum* species and the outgroup selected Withania somnifera were clustered together in one group, while *C. frutescens* was present in a separate monophyletic group.

#### Table 3: List of characteristics features to all five genetic markers from C.

<table>
<thead>
<tr>
<th>Barcode</th>
<th>Accession number</th>
<th>Number of conserved sites</th>
<th>Number of variable sites</th>
<th>Number of parsimony informative sites</th>
<th>Number of singleton sites</th>
<th>G+C %</th>
</tr>
</thead>
<tbody>
<tr>
<td>ITS</td>
<td>MT643918.1</td>
<td>458</td>
<td>105</td>
<td>17</td>
<td>86</td>
<td>64</td>
</tr>
<tr>
<td>matK</td>
<td>MN528466.1</td>
<td>845</td>
<td>16</td>
<td>none</td>
<td>none</td>
<td>32.8</td>
</tr>
<tr>
<td>rbcL</td>
<td>OK085708</td>
<td>989</td>
<td>4</td>
<td>none</td>
<td>none</td>
<td>43.3</td>
</tr>
<tr>
<td>psbA</td>
<td>OK663603</td>
<td>553</td>
<td>none</td>
<td>none</td>
<td>none</td>
<td>27.4</td>
</tr>
<tr>
<td>rpoB</td>
<td>ON368037</td>
<td>175</td>
<td>9</td>
<td>none</td>
<td>9</td>
<td>33.3</td>
</tr>
</tbody>
</table>

**Figure 1:** Phylogenetic relationship among the *Capsicum* species and outgroup taxa: The ML phylogenetic tree was reconstructed using the matK sequence data. The tree with the highest log likelihood (–2299.78) is shown.
analysis performed was of ITS [Figure 2], which showed different clusters; the first cluster showed species *frutescens, annuum, chinense,* and *eximium* to be closely related and the other cluster had *pubescens* and *baccatum* species, while the outgroup *Lycium torreyei* was seen to be present in a separate monophyletic clade. The result obtained was in correlation with previously observed results for species differentiation in *Capsicum* using ITS genetic marker, where it was noted ITS to be insufficient in differentiating among different *Capsicum* species, mainly between *C. frutescens* and *C. chinense* [18].

Phylogenetic analysis of rbcL [Figure 3] nuleotide sequences showed a well- resolved tree, where *C. frutescens* belonged to a monophyletic group, while *Capsicum annuum* belonged to a separate cluster, along with *Capsicum toverii* and the third cluster was made up of *Capsicum chinense* and *eximium*. It also showed a fourth cluster with *Capsicum pubescens* and *baccatum*, while the outgroup *Solanum Americanum* showed a separate cluster. Phylogeny analysis of *trnH- psbA* sequences [Figure 4] showed a tree with only two clusters, the first cluster contained all the *Capsicum* species, including *annuum, chinense,* and *frutescens*, while the outgroup formed a monophyletic cluster. Dendrogram representation of *rpoB* [Figure 5] showed a phylogenetic tree with three monophyletic clusters, where the first cluster had all the *Capsicum* species selected for analysis except *C. frutescens* which belonged to the second cluster, whereas the third cluster showed the outgroup that was selected.

The phylogenetic tree of *matK* shows it to be effective in differentiating *C. frutescens* from other *Capsicum* species as *C. frutescens* formed its separate branch with a length of 0.320, while all the other *Capsicum* species had a branch length of 0.303. It also showed 845 conserved sites and 16 variable regions with a GC content of 32.8%, while sequence analysis of ITS showed presence of highest number of variable sites with 64% GC content, while dendrogram analysis reveals that ITS was able to classify all the three *Capsicum* species in a separate clade and while they share the same ancestor and formed sister clades, they also showed branch length that were very close to one another suggesting them to be closely related.

![Figure 2](image1.png) - Phylogenetic relationship among *Capsicum* species and outgroup taxa: The ML phylogenetic tree was reconstructed using the ITS sequence data. The tree with the highest log likelihood (–1398.41) is shown.

![Figure 3](image2.png) - Phylogenetic relationship among *Capsicum* species and one outgroup taxa: The ML phylogenetic tree was reconstructed using the rbcL sequence data. The tree with the highest log likelihood (–1502.33) is shown.
The phylogenetic analysis of rbcL sequences gave very good results in terms of the conserved site with only four variable sites and 43.3% GC content. rbcL also proved to be a very good barcode in terms of its ability to differentiate between the three closely related sisters Capsicum species. The resulting phylogenetic tree placed frutescens, annuum, and chinense species all in three different clades making it effectively suitable to be used as a potential barcode. This again coincides with the results obtained in previous study, where phylogenetic analysis of rbcL marker placed C. frutescens in a monophyletic clade [19]. Whereas trnH-psbA did not resolve any Capsicum species as all the species were placed in the same clade with the same branch length, while the outgroup was placed in a different branch. While the phylogenetic tree of rpoB showed, C. frutescens has a wide difference when compared to other Capsicum species and placed frutescens under separate clade while keeping other Capsicum species and the outgroup species in the same cluster. It also showed that all the species involved in phylogenetic analysis shared a common ancestor. From the results obtained, it can be said that barcodes ITS and rbcL could be used as markers for differentiating the three Capsicum species (annuum, frutescens, and chinense), while matK and rpoB markers can be used as potential barcodes for species identification of C. frutescens.

4. DISCUSSION

Identification and discrimination of species are the first step in plant taxonomy, and DNA barcoding provides a rapid and efficient method to discriminate species [20]. In the present study, we have investigated five potential DNA barcoding loci, namely, matK, rbcL, ITS, rpoB, and trnH-psbA, to identify C. frutescens, and to examine the discriminating ability of these genetic loci in distinguishing the three morphological similar Capsicum species. In the present study, five genetic loci from C. frutescens have been sequenced and compared with the available
sequences of different *Capsicum* species in the GeneBank database of NCBI. The potential of the loci as a barcode was determined using multiple sequence alignment and their phylogenetic analysis. Sequence analysis of different loci from *C. frutescens* showed *psbA* to possess the highest number of conserved regions with zero variable sites; this was followed by *rbcL* and *rpoB*, which showed four and nine variable sites, respectively. High conserved sites suggest a low level of evolution in these sequences; this can also be observed in the phylogenetic tree that was created, where *psbA* and *rpoB* did not separate different *Capsicum* species into clusters, but *rbcL* was able to separate *C. frutescens* from other *Capsicum* species. This result observed was contradictory to the result observed by Jarret (2008), where *rpoB* placed *frutescens* species together with all other *Capsicum* species in the dendrogram [1]. Whereas *rbcL* dendrogram showed differentiation of *Capsicum* species into separate clusters, the branch length was close, suggesting the species to be closely related. The Consortium for Barcode of life PWG has recommended a combination of two loci *matK* + *rbcL*, to be used as a barcode for plants as a single barcode could not be identified as a universal barcode [21]. The result obtained in the present study proved to be in alignment with this conclusion as the *matK* locus was also able to place *C. frutescens* into a separate cluster interestingly, it was not able to differentiate among the remaining *Capsicum* species and was all placed in the same clusters.

Sequence analysis of *matK* correspondingly showed 16 variable sites for *C. frutescens*. Multiple sequence alignment of *matK* and *rbcL* also gave substantial results, where *C. frutescens* showed substitutions and gaps at specific positions. The highest number of variable sites was observed for ITS which gave a well-resolved phylogenetic tree. ITS successfully placed the three morphologically similar *Capsicum* species into three different clusters, proving it to be an effective barcode for analysis of *Capsicum* species and also identification of *C. frutescens* along with *matK* + *rbcL* markers. After evaluating the barcodes, the combination of three barcode markers *matK* + *rbcL* + *rpoB* can be proposed as a suitable barcode for identifying *C. frutescens*. In a previous China Plant BOL Group study, it was observed that the combination of *rbcL* + *matK* + ITS barcode markers gave 77.4% species discrimination [22]. The present study combination of *rbcL* + ITS loci was observed to be successful in distinguishing the three closely related *Capsicum* species of *annuum-frutescens-chinense*.

5. CONCLUSION

The present research attempts to analyze five different DNA markers as a way to authenticate *C. frutescens*. The present study employs DNA barcoding approaches and provides significant findings for determining the phylogeny and connection among different *Capsicum* species. The current study attempts to show the possibility of differentiating the three *Capsicum* species, namely *annuum*, *frutescens*, and *chinense* using five different gene markers as they are morphologically very similar. The results indicate that specific differences exist between the three *Capsicum* species. It was observed that *matK*, *rbcL*, and *rpoB* were useful in identifying *C. frutescens*, whereas *rbcL* and ITS successfully differentiated the three morphologically similar *Capsicum* species. This knowledge will not only help taxonomists to identify *C. frutescens* but also will provide essential insights into the differences existing in the *Capsicum* genus on the molecular level.

6. AUTHORS’ CONTRIBUTIONS

All authors made substantial contributions to conception and design, acquisition of data, or analysis and interpretation of data; took part in drafting the article or revising it critically for important intellectual content; agreed to submit to the current journal; gave final approval of the version to be published; and agreed to be accountable for all aspects of the work. All the authors are eligible to be an author as per the International Committee of Medical Journal Editors (ICMJE) requirements/guidelines.

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8. CONFLICTS OF INTEREST

The authors report no financial or any other conflicts of interest in this work.

9. ETHICAL APPROVALS

This study does not involve experiments on animals or human subjects.

10. DATA AVAILABILITY

All data generated and analyzed are included within this research article and the sequences are available online in the NCBI database.

11. PUBLISHER’S NOTE

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