Analysis of Phylogeny and Evolutionary Divergence of RbcL Sequence of Barleria longiflora L.f

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Abstract: Barleria longiflora L.f belongs to the family Acanthaceae which is endemic distribution. Presence of this plant in Erattamalai hill, located in Tiruchirappalli District of Tamil Nadu, India. The rbcl gene is widely used in systematic studies to resolve divergences at many taxonomic levels. The collected B. longiflora were subjected to sequence analysis of rbcL gene and the same was deposited in Gene Bank and an accession number has been assigned for the same (Accession No. KR861703.1). The data was further analyzed for the construction of neighbor joining tree and to infer the evolutionary divergence among the maximum identical sequences deposited in Gene Bank and an accession number has been assigned for the same (Accession No. KR861703.1). The results of the search revealed that the rbcL sequence of B. longiflora has not been sequenced earlier from any part of this world. It is obvious that the rbcL sequence of B. longiflora may be used for the identification of this species reported from any part of the world through BLAST analysis if the identical sequences are submitted to Gene Bank in future.

Keywords: ribulose-1,5-bisphosphate carboxylase/oxygenase large subunit gene, Barleria longiflora, Erattamalai hill,

1. Introduction

The large and primarily tropical plant family Acanthaceae contains at least 4000 species, placing it among the top 12 or so most diverse families of flowering plants. The group is part of the order Lamiales, which includes familiar plants such as snapdragons (Antirrhinum, Plantaginaceae), mints (Lamiaceae), African violets (Gesneriaceae), and monkey flowers (Mimulus, Phrymaceae). Notably, as a result of the largely tropical distribution of Acanthaceae, species-level diversity (as well as some generic diversity) remains poorly understood, and there is little doubt that many new species remain to be discovered. A wide variety of essential and medicinally active ingredients have been identified and extracted from some members of the family. Many scientists have worked on some members of the family and have discovered some very essential active ingredients present in them [1]. Traditional methods of plant identification based on morphological criteria are difficult to apply accurately due to subjective biases. Particularly, in the case of medicinal plants, the use of chromatographic profiles of marker compounds to standardize botanical preparations is also of limited value because the medicines have varied sources and chemical complexity, which is affected by growth, storage conditions, and harvest times[2].

DNA-based identification (barcoding) is simple, does not require taxonomic expertise, and is free from subjective errors, which is not the case in morphological identification. Valid identification of unknown samples is the main goal of barcoding (Hebert and Gregory, 2005)[3], despite ongoing criticism of the feasibility or even necessity of DNA barcoding for general taxonomic purposes (Will et al., 2005; Spooner, 2009)[4,5]. The most common gene used to provide sequence data for plant phylogenetic analyses is the plastid-encoded rbcL gene[6,7]. This single copy gene is approximately 1430 base pairs in length, is free from length mutations except at the far 3’ end, and has a fairly conservative rate of evolution. The function of the rbcL gene is to code for the large subunit of ribulose 1, 5 bisphosphate carboxylase/oxygenase (RUBISCO or RuBPCase). The sequence data of the rbcL gene are widely used in the reconstruction of phylogenies throughout the seed plants. However, it is apparent that the ability of rbcL to resolve phylogenetic relationships below the family level is often poor[8]. Thus, interest exists in finding other useful DNA regions that evolve faster than does rbcL to facilitate lower-level phylogenetic reconstruction. The Consortium for the Barcode of Life (CBOL) plant working group has recommended a standard barcode comprising ribulose-1,5-bisphosphate carboxylase/oxygenase large subunit (rbcL) for the barcoding of all land plants [9].

2. Materials and Methods

Young leaves of the respective species were collected in sterile Ziploc bags and stored at -20 °C until further use.

DNA isolation using NucleoSpin® Plant II Kit (Macherey-Nagel) About 100 mg of the tissue is homogenized using liquid nitrogen and the powdered tissue is transferred to a microcentrifuge tube. Four hundred microlitres of buffer PL1 is added and vortexed for 1 minute. Ten microlitres of RNase A solution is added and inverted to mix. The homogenate is incubated at 65°C for 10 minutes. The lysate is transferred to a Nucleospin filter and centrifuged at 11000 x g for 2 minutes. The flow through liquid is collected and the filter is discarded. Four hundred and fifty microlitres of buffer PC is added and mixed well. The solution is transferred to a Nucleospin filter and centrifuged at 11000 x g for 1 minute and the flow through liquid is discarded. Four hundred microlitre buffer PW1 is added to the column, centrifuged at 11000 x g for 1 minute and flow though liquid is discarded. Then 700 µl PW2 is added, centrifuged at 11000 x g and flow through liquid is discarded. Finally 200 µl of PW2 is added and centrifuged at 11000 x g for 2 minutes to dry the silica membrane. The column is transferred to a new 1.7 ml tube and 50 µl of buffer PE is
added and incubated at 65°C for 5 minutes. The column is then centrifuged at 11,000 x g for 1 minute to elute the DNA. The eluted DNA was stored at 4°C.

Agarose Gel Electrophoresis for DNA Quality check. The quality of the DNA isolated was checked using agarose gel electrophoresis. 1 µl of 6X gel-loading buffer (0.25% bromophenol blue, 30% sucrose in TE buffer pH-8.0) was added to 5 µl of DNA. The samples were loaded to 0.8% agarose gel prepared in 0.5X TBE (Tris-Borate-EDTA) buffer containing 0.5 µg/ml ethidium bromide. Electrophoresis was performed with 0.5X TBE as electrophoresis buffer at 75 V until bromophenol dye front has migrated to the bottom of the gel. The gels were visualized in a UV transilluminator (Genei) and the image was captured under UV light using Gel documentation system (Bio-Rad) (Figure 1). PCR amplification reactions were carried out in a 20 µl reaction volume which contained 1X Phire PCR buffer (contains 1.5 mM MgCl2), 0.2mM each dNTPs (dATP, dGTP, dCTP and dTTP), 1 µl DNA, 0.2 µl Phire Hotstart II DNA polymerase enzyme, 0.1 mg/ml BSA and 3% DMSO, 0.5M Betaine, 5pM of forward and reverse primers. The PCR amplification was carried out in a PCR thermal cycler (GeneAmp PCR System 9700, Applied Biosystems). Forward primer ATGTCACCACAAACAGAGACTAAAGC and Reverse primer GTAAATCAAGTCCACCRCG. PCR amplification profile for rbcL 98°C -30 sec, 98°C -5 sec, 60°C -10 sec and 72°C -15 sec for 30 cycles, 72°C - 60 sec and stored at 4°C.

Agarose Gel electrophoresis of PCR products were checked in 1.2% agarose gels prepared in 0.5X TBE buffer containing 0.5 µg/ml ethidium bromide. 1 µl of 6X loading dye was mixed with 5 µl of PCR products and was loaded and electrophoresis was performed at 75V power supply with 0.5X TBE as electrophoresis buffer for about 1-2 hours, until the bromophenol blue front had migrated to almost the bottom of the gel. The molecular standard used was 2-log DNA ladder (NEB). The gels were visualized in a UV transilluminator (Genei) and the image was captured under UV light using Gel documentation system (Bio-Rad). Consists of two hydrolytic enzymes, Exonuclease I and Shrimp Alkaline Phosphatase (SAP), in a specially formulated buffer for the removal of unwanted primers and dNTPs from a PCR product mixture with no interference in downstream applications. Five micro litres of PCR product is mixed with 2 µl of ExoSAP-IT and incubated at 37°C for 15 minutes followed by enzyme inactivation at 80°C for 15 minutes [10].

Sequencing using BigDye Terminator v3.1 sequencing reaction was done in a PCR thermal cycler (GeneAmp PCR System 9700, Applied Biosystems) [11]. The PCR mix consisted of the following components: PCR Product (ExoSAP treated) - 10-20 ng. Primer - 3.2 pM (either Forward or Reverse) Sequencing Mix - 0.28 µl , 5x Reaction buffer -1.86 µl, Sterile distilled water- make up to 10µl. The sequencing PCR temperature profile consisted of a 1st cycle at 96°C for 2 minutes followed by 30 cycles at 96°C for 30 sec, 50°C for 40 sec and 60°C for 4 minutes for all the primers.

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maximum Log likelihood for this computation was -1466.297. The analysis involved 20 nucleotide sequences. Clustered sequences showed 1794 conserved sites, 335 variable sites, 74 parsim-info sites, 203 singleton sites of 529bp. Further the results revealed that there were 1365 zero-fold sites, 412 two-fold sites and 175 four-fold sites. Of the 529bp sites, 663 sites showed 100 % coverage.

The results of NJ analysis of 697bp fragment of the rbcL gene belonged to *B. longiflora* with the twenty sequences obtained through BLAST showed different branch lengths in the Phenogram. Maximum identical sequences were not available for *B. longiflora* in this NJ analysis (Fig. 1). The evolutionary history was inferred using the NJ method. The optimal tree with the sum of branch length was 20.33. The tree was drawn to scale, with branch lengths (next to the branches) in the same units as those of the evolutionary distances used to infer the phylogenetic tree. The evolutionary distances were computed using the Maximum Composite Likelihood method [17] and were in the units of the number of base substitutions per site.

A detailed estimate of evolutionary divergence of rbcL sequence of *B. longiflora* was 0.026. The maximum evolutionary distance observed between *Barleria prionitis* and *B. acuminata* was 3.865 (Table 1). Therefore, it is concluded that rbcL sequence of *B. longiflora* may be used for the identification of this species reported from any part of the world through BLAST analysis if the identical sequences are submitted to Gene Bank in future.

**Table 1:** Estimate of Evolutionary Divergence of rbcL sequence of *B. longiflora* with their similar sequences through BLAST search

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References


