Phylogenetic study of mangrove associate grass *Myriostachya wightiana* (Nees ex Steud.) Hook. f. using *rbcL* gene sequence

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**ABSTRACT**

*Myriostachya* is a monotypic genus in the family Poaceae, with the only known species *Myriostachya wightiana* (Nees ex Steud.) Hook.f. It is a mangrove associate grass primarily distributed along the muddy streams and channels in intertidal mangrove swamps of India, Bangladesh, Sri Lanka, Myanmar, Thailand and Sumatra. Molecular identification and evolutionary studies of *M. wightiana* is unreported till now. Therefore, in this study, the phylogenetic analysis of *M. wightiana* was established with related family members by using chloroplast *rbcL* gene-based systematics. The molecular phylogeny was accomplished by DNA extraction, PCR amplification and sequencing of the *rbcL* gene and phylogenetic analysis. The genomic DNA was extract using the CTAB method and the *rbcL* gene amplification is by using the F-5’ATGTACACAAAAAAGAAGAAACGATCTC3’ and R-5’CTTCGGCACAAAATAAGAAACGATCTC3’ primers. Phylogenetic analysis of *M. wightiana* was performed by multiple sequence alignment with UPGMA, and the Maximum-parsimony phylogenetic tree was constructed using MEGAx. *Myriostachya wightiana* *rbcL* gene sequence shows the highest similarity to *Paspalum* species, and in the phylogenetic tree *M. wightiana* has a close branch with *Paspalum vaginatum*. The evolutionary divergence from *M. wightiana* is maximum (0.49) to *Sorghum propinquum* and minimum (0.01) to *Oryza officinalis* and *Oryza punctata*. This study concluded that *M. wightiana* has a strong morphological and phylogenetic relationship with salt-tolerant *Paspalum* sp.

**Introduction**

*Myriostachya* is monotypic genus in the Poaceae family, with *Myriostachya wightiana* (Nees ex Steud.) Hook. f. being the only species (1). The species is tropical with, its native range from the Indian Subcontinent to West Malesia. It is widely located in the intertidal mangrove swamps of India, Bangladesh, Sri Lanka, Myanmar, Thailand and Sumatra (2). In India, it is mainly distributed on the southeast coast of the Bay of Bengal. *M. wightiana* is a large, densely clumped perennial grass growing up to 3 meters. It frequently occurs along with *Acanthus ilicifolius* L., *Nypa fruticans* Wurmb., and *Porteresia coarctata* (Roxb.) Tateoka. The species grows well in saline water habitat rather than fresh water due to structural adaptations like the thick epidermis, sclerenchymatous vascular system, salt secretion glands, conspicuous metaxylem and broad phloem region in stem and leaf, dense cortex and lignified root exodermis (3).

The taxonomic position of *M. wightiana* is recorded based on the morphological features. However, the morphology-based systematics and evolutionary studies of Poaceae are not reliable due to the higher number of taxa, simplicity of floral architecture and vegetative structure, dynamic and mosaic evolution (4). Because of these challenges, DNA-based molecular systematics are being used to generate promising results. According to the available literature, the evolutionary studies of *M. wightiana* have not been carried out yet. Thus, this study aims to build phylogenetic relationships of *M. wightiana* by DNA dependent molecular systematics. DNA barcoding is a relatively quick and accurate method for identification of any plant or animal species (5). DNA barcoding can be used in molecular systematics to identify the new species in conjugation with conventional taxonomic approaches (6). The fundamental principle of DNA sequence utilisation in phylogenetic analysis is the occurrence of nucleotide changes over time. Therefore, the estimation and reconstruction of evolutionary relationships between different organisms are possible (7).

In plants, the chloroplast genome is quite helpful for evolutionary and phylogenetic experiments, especially above the species level because of its relatively abundant DNA material, single-copy genes,
Materials and Methods

Sample collection

Healthy and young *M. wightiana* (Fig. 1) plant material were collected from the mangroves (Fig. 2) of Bhavanapadu (Long: 18° 33’ 52” to 18° 32’ 11” N; Lat: 84° 21’ 26” E to 84° 18’ 22” E) which is located in the northeast of Andhra Pradesh, adjoining the Bay of Bengal, India. The collected plant samples were aseptically transferred to the zip bags and transported to the laboratory. The plant species was authenticated by Dr. S. Hara Sreeramulu, Taxonomist, Department of Botany, Dr. V. S. Krishna College. Visakhapatnam and the voucher specimen (No. 00564/AP) was deposited at the Herbarium, Department of Botany, Andhra University.

DNA extraction

Total DNA content was extracted from the leaves of *M. wightiana* was carried out by the CTAB method (11). Surface sterilised healthy and young leaves were grounded using liquid nitrogen and 500 µg of ground leaf powder was mixed with 750 µl of CTAB, 20 µl of mercaptoethanol in a 2 ml Eppendorf tube. The mixture was vortexed intermittently and then incubated in a water bath at 65 °C for 45 min. After incubation, the mixture was allowed to cool until it reaches room temperature. Subsequently, 750 µl of chloroform/isoamyl alcohol (24:1) was added to the mixture, and the tube was centrifuged at 12000 rpm (Eppendorf, 5810R) for 10 min and the upper aqueous phase was separated into a new tube. The chloroform/isoamyl alcohol extraction was repeated twice to the aqueous phase and centrifuged for 5 min. To the resultant supernatant, 0.1 ml of 3M sodium acetate (pH 4.6), 2 vol. of 95% ethanol was added and incubated at -20 °C for 1 hr to precipitate DNA. Then the precipitate was centrifuged at 12000 rpm for 10 min. The DNA pellet was washed twice with 750 µl of 70 % ethanol, then centrifuged for 10 minutes at 10000 rpm. The resulting DNA pellet was rewashed twice with 96 % ethanol and dried in a desiccator for 15 min. The DNA pellet was stored in a -20 °C freezer.

Sequence alignment and Phylogenetic analysis

The evolutionary analysis of *M. wightiana* was conducted by constructing a phylogenetic tree with some major species of Poaceae. *Myriostachya wightiana* rbcL gene sequence was subjected to the BLASTn in the NCBI server to identify the homologous sequence or species. From the NCBI database, 33 species were selected based on their degree of homology with the target rbcL gene sequence. The selected 33
species’ rbcL gene sequences were derived from the nucleotide NCBI database. In the selected thirty-three species for phylogenetic analysis, thirty species belong to the Poaceae family, and the remaining three species were used as an outgroup that consists of marine algae Saccharina latissimi (L.) C.E.Lane, C.Mayes, Druehl and G.W.Saunders., Undaria pinnatifida (Harv.) Suringar and Porphyra haitanensis T.J.Chang and B.F.Zheng, 1960. which belongs to Laminariaceae, Alariaceae and Bangiaceae respectively. The marine algae were used as an out group to known the evolutionary progress from the marine algae to marine grasses and to terrestrial grasses. Multiple sequence alignment was conducted with an advanced cluster method UPGMA using Muscle embedded in MEGAX to search homology of rbcL gene sequences between the M. wightiana and the selected 33 species. To analyse the variations among the sequences, a distance matrix was determined, and based on the differences expressed in the distance matrix; a maximum-parsimony tree was constructed using MEGAX. The evaluation of phylogenetic tree topologies was done by the bootstrap method with 1000 replicates for all nodes (15).

**Results and Discussion**

**DNA extraction and quantification**

The result was given as Mean ± Standard Deviation obtained from three independent experiments. As part of DNA barcoding, the complete genome of M. wightiana was successfully extracted. The isolated DNA appeared as a prominent band on 1% agarose gel (Fig. 3A) and the isolated total DNA content was measured as 663 ± 54 µg/gm. DNA barcoding is one of best way to classify new species and also to collect a database of the reference sequences (16).

**PCR amplification and sequence analysis**

The PCR amplified M. wightiana chloroplast rbcL gene was successfully run by the 1.5% agarose gel electrophoresis. In the agarose gel (Fig. 3B), the first well shows marker DNA and the remaining three wells indicate amplified rbcL gene product. The thick and single bands in the agarose gels indicate that the rbcL gene amplification was done successfully, and the size of the amplified M. wightiana rbcL gene was approximately ±650bp. The yield and consistency of DNA bands in the agarose gels determines the universality of selected primers and their discriminating strength. According to one report, universal primers identify the regions in the rbcL gene of Angiosperms and display a high degree of universality in terrestrial plants (14). The direct nucleotide sequencing of PCR amplified products is now emerging as an important field of evolutionary studies and systematics (17). An ideal DNA barcode can be recovered and gives maximum discrimination among species with a single pair of primers suitable for bidirectional sequencing with a minor sequence modification. The chloroplast rbcL gene contains the least number of variable regions and it is referred to as a best-characterised gene that is easily retrievable with common PCR primers. It was proposed that, out of the nuclear and plastid genomes, plastid genomes are used for phylogenetic research in plants since they are thought to have similar ancestors (18). Our research used the chloroplast rbcL gene sequence, considering that the rbcL gene is quickly amplified and sequenced in many terrestrial plants and impacts in phylogenetic studies by placing the species in a correct genus and plant family.
M. wightiana chloroplast rbcL gene nucleotide composition was computed by Seqstate V.1.21 server (19). The amplified conserved region of the rbcL gene (GenBank Accession No. KY293284) has 604 nucleotides and was estimated to have a molecular weight of 373 KDa. The nucleotide composition of the rbcL gene consists of 164 bp Adenine (A), 173 bp Thymine (T), 140 bp Guanine (G), 127 bp Cytosine (C), and the percentage of GC was measured as 44.2. The Extinction coefficient of the M. wightiana rbcL gene is estimated to be 9805831 Mol$^{-1}$ cm$^{-1}$. Kusumi and Tachida (20) reported that the GC content in the plants differs from 28-42%. It has been reported that the GC content of the rbcL gene in wild Solanum sp. is 43.9% (21). The current findings showed that the rbcL genes in M. wightiana have comparatively low GC content.

Multiple sequence alignment and phylogenetic tree analysis

The BLAST search results showed similar rbcL gene sequences of other species with their percentage of identity against the M. wightiana rbcL gene sequence. The BLASTn results revealed that the M. wightiana rbcL gene sequence showed the highest similarity to Paspalum sp., with an identity of 99% and an E-value of 0.0. From the hits, a total of 33 rbcL gene sequences from different species were selected to construct evolutionary relationships through multiple sequence alignment using the UPGMA program in MEGAX. The multiple sequence alignments indicated a variable number of deletions and insertions in the chloroplast rbcL. The phylogenetic parameters such as variance, parsimony sites, and overall mean distance were calculated as 0.52, 0.32 and 0.1 respectively in the rbcL alignments of family Poaceae. The evolutionary divergence of M. wightiana with its other species of Poaceae was determined. A satisfactory result was established by using the rbcL gene as a marker to evaluate the phylogenetic relationship among the grass species. The evolutionary divergence among sequences at the generic level varies from 0.01-0.49. M. wightiana showed the highest evolutionary divergence with the species of Sorghum propinquum (Kunth) Hitchc. (0.49), Undaria pinnatifida (Harv.) Suringar (0.33) and Porphyra haitanensis T.J.Chang and B.F.Zheng. Whereas M. wightiana shows the least evolutionary divergence with the species of Oryza officinalis Wall. (0.01), O. punctata Rotschy ex Steud. (0.01), O. glaberrima Steud. (0.02) and O. nivara S.D.Sharma and Shastry (0.02). Multiple sequence alignments of the rbcL gene sequences revealed that the rbcL gene is highly conserved throughout the Poaceae family.

In the phylogenetic tree (Fig. 4), there were five main clades. The first main clade consists of two subclades. The first subclade of main clade-I composed Paspalum sp. and the second subclade composed of Zea and Sorghum species. The second and third main clades consist of Triticum and Oryza species. The fourth main clade composed only Sorghum propinquum. The fifth main clade consists of marine macroalgae Porphyra haitanensis, Saccharina latissimi and Undaria pinnata. The phylogenetic tree indicated that the M. wightiana found a close branch with Paspalum vaginatum Sw. (GenBank Accession No. LN907995). In the phylogenetic tree, the clades are organised mainly with several species in the same genus according to their similarities. M. wightiana is known to be the nearest species to seashore grasses in phylogenetic analysis.
Conclusion

An ideal taxonomic recognition of species is essential for the proper management of any organism. Species-level evolutionary information of a plant can be provided by the chloroplast rbCL gene-based phylogenetic analysis. The present study of rbCL gene sequence and phylogenetic analysis explored the evolutionary divergence and relatedness of the *M. wightiana*. The rbCL gene sequence and multiple sequence alignment revealed that the *M. wightiana* showed 99% homology to *Paspalum* sp. The evolutionary divergence from *S. propinquum* to *M. wightiana* was estimated as 0.49, which was found maximum and the minimum was for *O. officinalis* and *O. punctata* that are 0.01. From the molecular phylogeny by the rbCL gene, it was concluded that the

![Fig. 4. Maximum Parsimony tree of *Myriostachya wightiana* and other species based on the rbCL gene. Bootstrap values are indicated on the branches.](image-url)
M. wightiana has a strong relationship to salt-tolerant grasses like *Paspalum* sp.

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**Authors’ contributions**

MKK carried out isolation of rbcL gene, sequencing, multiple sequence alignment, construction of the phylogenetic tree and drafting of the manuscript. BVS participated in the design and planning of research work, helped to write the article and corrected the manuscript. The final manuscript was read and approved by all the authors.

**Conflict of interests**

The authors declare that there are no conflicts of interest.

**Supplementary file**

**Table 1.** Estimates of evolutionary divergence among the selected species.

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