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Application Note

DNA Barcode ITS Effectively Distinguishes the Medicinal Plant Boerhavia diffusa from Its Adulterants

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Abstract

Boerhavia diffusa (B. diffusa), also known as Punarnava, is an indigenous plant in India and an important component in traditional Indian medicine. The accurate identification and collection of this medicinal herb is vital to enhance the drug's efficacy and biosafety. In this study, a DNA barcoding technique has been applied to identify and distinguish B. diffusa from its closely-related species. The phylogenetic analysis was carried out for the four species of Boerhavia using barcode candidates including nuclear ribosomal DNA regions ITS, ITS1, ITS2 and the chloroplast plastid gene psbA-trnH. Sequence alignment revealed 26% polymorphic sites in ITS, 30% in ITS1, 16% in ITS2 and 6% in psbA-trnH, respectively. Additionally, a phylogenetic tree was constructed for 15 species using ITS sequences which clearly distinguished B. diffusa from the other species. The ITSI demonstrates a higher transition/transversion ratio, percentage of variation and pairwise distance which differentiate B. diffusa from other species of Boerhavia. Our study revealed that ITS and ITS1 could be used as potential candidate regions for identifying B. diffusa and for authenticating its herbal products.

Keywords: Adulterant; Boerhavia diffusa; ITS; DNA barcoding; Punarnava

Introduction

Boerhavia is one of the highly polymorphic genus in Nyctaginaceae family [1]. About 40 species are distributed in tropical, subtropical and temperate regions. Among these, 6 species are reported in India and Boerhavia diffusa (B. diffusa) is indigenous [2]. B. diffusa is described as Punarnava by an Indian system of medicine, Ayurveda [3]. Roots and whole plants of B. diffusa are used in the Ayurvedic and Unani systems of medicine in Arabian countries [4] and many tribal communities in India still use it for the treatment of jaundice and various other liver disorders. It has anti-inflammatory, diuretic, fibrinolytic, anti-convulsant properties [5–8] and also used as carminatives [9–10]. The two pharmaceutically important alkaloids, Punarnavine-1 and Punarnavine-2, belonging to the group of quinolizidine were separated from *B. diffusa* [11–12].

B. diffusa is known to be extensively adulterated with other species like Boerhavia erecta, Boerhavia repanda, Boerhavia coccinea and Boerhavia verticillata. B. diffusa have taxonomical conflicts with B. coccinea, Boerhavia repens, Boerhavia tetranda and Boerhavia albiflora, making it difficult to distinguish from one another [13–14]. The species B. verticillata display similar morphological features and phytochemical properties with B. diffusa, but they differ by their habits [15]. Determination of plant specimens by DNA barcodes will be an effective, reliable and simple pharmacognostic tool to resolve the confusion in morphological identification. Due to different rates of evolution, nuclear ribosomal internal transcribed spacer (ITS) regions have become the routine marker in evolutionary studies at different taxonomic levels [16,17]. There is a report using the chloroplast intergenic spacer psbA-trnH for identifying

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the *Dendrobium* species of Chinese pharmacopoeia and *psbA-trnH* is recommended as an ideal DNA barcode candidate [18]. Recently the sequence variations are used to develop specific markers for the identification and authentication of drugs and herbal formulations [19].

The objective of the present study is to evaluate an ideal barcode candidate for distinguishing and authenticating the species *B. diffusa* from its common adulterants.

Results

Genomic DNA was isolated from the species of *B. diffusa*, *B. repanda*, *B. erecta* and *B. verticillata* and used for PCR amplification of the *ITS* and *psbA-trnH*. The obtained sequences were submitted to GenBank. The size and accession number for the gene *ITS* and *psbA-trnH* is shown in Table S1. Additionally, *ITS* sequences from 11 species of *Boerhavia* were taken from the GenBank (Table S2) and used for sequence alignments.

Multiple sequence alignment and pairwise alignment analysis were performed for nuclear *ITS* and chloroplast *psbA-trnH* (Figure S1). The *ITS* region consists of *ITS1*, 5.8S rDNA and *ITS2*. The ribosomal sites of 5.8S rRNA and 28S rRNA are highly conserved. The regions *ITS1* and *ITS2* were compared by multiple sequence alignment, where *ITS1* showed more variation than *ITS2*. Phylogenetic analysis using *ITS1* and *ITS2* indicated *B. diffusa* and *B. erecta* in the same clade while *B. verticillata* and *B. diffusa* was shown in the same clade when using *psbA-trnH* region for phylogenetic analysis (**Figure 1**). The tree also constructed using *ITS* region clearly distin-

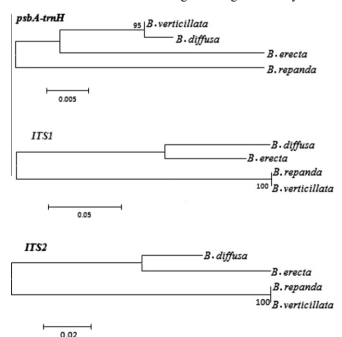


Figure 1 Phylogenetic trees of the four *Boerhavia* species constructed using *ITS1*, *ITS2* and *psbA-trnH*

Phylogenetic trees were constructed by Minimum Evolution method using ITS1, ITS2 and psbA-trnH, respectively, for the four species of Boerhavia, including B. diffusa, B. erecta, B. repanda and B. verticillata.

guished the morphologically similar species *B. diffusa* from the 14 other species of *Boerhavia* as shown in the **Figure 2**.

We further analyzed the nucleotide variations of *ITS* and *psbA-trnH* between different species. Percentage of variation shown in **Figure 3** indicated that *ITS* demonstrated higher inter-specific divergence. The Wilcoxson rank test indicated significant variation between the species for *ITS1* when compared to *ITS2* and *psbA-trnH*. BLAST 1 and distance method also indicated that *ITS1* showed higher identification percentage at species level (**Table 1**).

Discussion

Recent molecular methods like DNA barcoding have been extensively used for species identification, diversity, forensic medicine and ecological studies [20–21]. It also plays an important role in the identification of traditional medicinal herbs. ITS2 has been effectively used in differentiating morphologically similar species like Swartzia grandifolia and Swartzia longicarpa and also in solving the controversial species Caranga rosea and Caranga sinica of the family Fabaceae [22]. Medicinal plant species like the family Polygonaceae [23] and the genus Dendrobium [24] have been identified using ITS2 region. In addition, ITS1 was used to demonstrate that species of Amonum villosum belongs to the family Zingiberaceae [25].

In our study, multiple sequence alignment of *ITS1* and *ITS2* from four *Boerhavia* species showed that *B. diffusa* had a unique basepair variation, which can distinguish it from the other three species, despite the fact that they share many morphological similarities. On the other hand, although *psbA-trnH* distinguishes some species of Polygonaceae [18], less sequence variation in *psbA-trnH* was revealed among the four species of *Boerhavia*. This result is consistent with a previous report that *psbA-trnH* does not show any variation for closely-related Cycad species [26]. Hence, *ITS1* may be a better barcode region for distin-

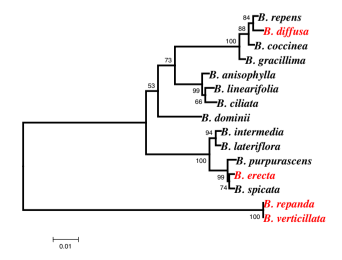


Figure 2 Phylogenetic tree of the 15 Boerhavia species constructed using ITS

Phylogenetic tree was constructed by Minimum Evolution method for the 15 species of *Boerhavia* using *ITS* region.

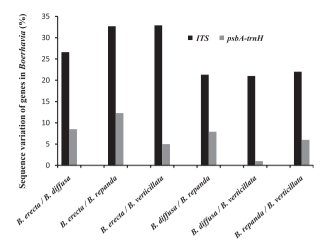


Figure 3 Sequence variation of ITS and psbA-trnH between four Boerhavia species

Percentage of nucleotide variations between different *Boerhavia* species, including *B. diffusa*, *B. erecta*, *B. repanda* and *B. verticillata*, was indicated for *ITS* (dark gray) and *psbA-trnH* (light gray), respectively.

Table 1 Validation of ITS1, ITS2 and psbA-trnH from four Boerhavia species

Gene name	Correct identification (%)		
	BLAST 1		Distance
	At genus level	At species level	At species level
ITS1	98	94	93.2
ITS2	99	92	91.1
psbA-trnH	94	91	76.86

guishing the species of *Boerhavia*, although *ITS2* has been widely used to distinguish the plant species [27]. This study clearly indicates that DNA barcoding using candidate like *ITS1* is a reliable method for differentiating *B. diffusa* from the other three species, which can also be applied to rapid identification of medicinal plants and their adulterants or substitutes.

Materials and methods

Sample collection

Four species of *Boerhavia* (*B. diffusa*, *B. erecta*, *B. repanda*, *B. verticillata*) were collected from the regions of Western Ghats (one of the hotspots), Coimbatore, India. The species were collected and identified by the taxonomist.

DNA extraction, PCR amplification and DNA sequencing

Fresh leaves from each plant species were used for isolating total genomic DNA by CTAB method [28]. The *ITS* and *psbA-trnH* gene amplifications were performed using Taq DNA polymerase with the primers indicated below. The forward and reverse primers for *ITS* and *psbA-trnH* used were *ITS* F 5'-GGAAGGAGAAGTCGTAACAAGG-3', *ITS* R 5'- TCCTCCGCTTATTGATATGC-3' [28] and

psbA-trnH F 5'-GTTATGCATGAACGTAAGCTC-3', psbA-trnH R 5'-CGCGCATGGTGGATTCAAATCC-3', respectively. The forward primer of ITS region from [29] was modified at the position of 6 $(T \rightarrow G)$ and 8 $(A \rightarrow G)$. The PCR program was as follows, an initial denaturation at 94 °C 5 min, followed by 35 cycles of 94 °C 1 min, 57 °C 30 s, 72 °C 1 min and final extension at 72 °C 10 min. PCR products were resolved by gel electrophoresis, purified and subjected to sequencing. The obtained ITS and psbA-trnH sequences were deposited in the GenBank of NCBI database (Gen-Bank Accessions: HQ386701, HQ386689, HQ386691, HO386695, HO386696, HO407399, HO386690 and JF423303) (Table S1). Sequences of ITS genes from 11 additional species of Boerhavia were obtained from GenBank (Table S2).

Sequence alignment and phylogenetic analysis

The DNA sequences were compared and aligned using the programs ClustalW [30] and MULTALIGN (http://www.multalin.toulouse.inra.fr/multalin/). Further, the DNA sequences were subjected to BLAST (http://www.ncbi.nlm.nih.gov/blast/blast.cgi) for better identification of sequence at species level. Phylogenetic trees were constructed with the Minimum Evolution method using MEGA 4.0. The intra-specific variation between the species was calculated using MEGA 4.0 [31] and StatsDirect was used to calculate the Wilcoxon signed rank [32].

Authors' contributions

SR supervised the research. RVS collected and identified the specimens. DS, DS, RKS and JCJ carried out the experimental study. Dhivya Selvaraj prepared the manuscript and SR revised it. All authors read and approved the final manuscript.

Competing interests

The authors have declared that no competing interests exist.

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Supplementary material

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.gpb.2012.03.002.

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