DNA Barcoding of Endangered Sacred Plant (Stereospermum suavelons) in Cuddalore District, Tamil Nadu, India

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Abstract
In the present investigation, DNA barcodings were made for the sacred trees of Main (Big) Temples in Cuddalore district, namely, *Stereospermum suavelons* to document their reliable identification, conservation, discrimination, similarities and evolutionary trend among them and with their related taxa for future use. DNA isolation from leaf samples of present study species was carried out by using a modified CTAB method and good isolation was got for the species studied. Gradient PCR amplification was performed for the isolated DNA using matK gene and the primers matK472F & matK1248R. The amplification success was 90-95%. PCR amplification was tested with 1 % agarose gel electrophoresis using ethidium bromide and the products were confirmed. The PCR products were sent to Xcelris Labs Ltd. Sydney House, Premchand Nagar Road, Bodakdev, Ahmedabad 380054, India, for DNA sequencing and sequences were got for the species with the success rate of 95 %. Pairwise sequence alignments were made with BLAST and multiple sequence alignments are made with ClustalW, and based on the sequence alignments, dendrograms were constructed using software Mega 5 and Neighbor joining method to study the phylogenetic aspects of the species studied and with their related taxa.


Introduction
DNA Barcoding is a new tool for the science of taxonomy at genotypic level. DNA barcoding as a scientific idea, initiated by Paul Herbert in 2003 has been attracting international attention for its significance in advancing the taxonomy of life forms. In other words, DNA barcoding is a taxonomic method that uses a short genetic marker in an organism’s DNA to identify it as belonging to a particular species. It differs from molecular phylogeny in that the main goal is not to determine classification but to identify an unknown sample in terms of a known classification. Although barcodes are sometimes used in an effort to identify unknown species or assess whether species should be combined or separated, such usage, if possible at all, pushes the limits of what barcodes are capable of plants.

Although a few million species have been discovered and identified, many millions more, including microorganisms, remain to be discovered and documented. Whether a species is endemic to a geographical region or is widely distributed, barcoding of life is truly international in scope.

In addition, DNA sequences have potential for routine plant identification in conjunction with a high throughput and bioinformatics system and optimal for the taxonomically wide plant identification with high relatedness within taxon variability and lack of confident assignment of orthology to overcome these hurdles (Blaxter, 2004). Some DNA regions are sufficiently conserved to permit primer sets for PCR amplification and contain sufficient informative sequence variation to discriminate up to the species level. For plants, many genes have been proposed as DNA barcodes till now. Applications include, for example, identifying plant leaves even when flowers or fruit are not available, identifying the diet of an animal based on stomach contents or feces, and identifying products in commerce (e.g. Herbal supplements or wood).

As barcode has multiple applications, has been used for ecological surveys (Dick & Kress, 2009), cryptic taxon identification (Lahaye et al., 2008), and confirmation of medicinal plant samples (Xue & Li, 2011). The DNA Barcoding has inspired a global initiative dedicated to create a library of new knowledge about species diversity, making that knowledge accessible and applying that knowledge to create tangible benefits. The uses and global benefits of DNA barcoding include i) Controlling agricultural pest, ii) identifying disease vectors, iii) Sustaining natural resources, iv) protecting endangered species and v) Monitoring water quality.

Universal *matK* primers for DNA barcoding
The chloroplast maturase K gene (*matK*) formerly known as *orf*K is one of the most variable coding genes of angiosperms and has been suggested to be a “barcode” for land plants. However, *matK* exhibits low amplification and sequencing rates due to low universality of currently available primers and mononucleotide repeats. To resolve these technical problems, the entire *matK* region has been evaluated to find a region of 600–800 bp that is highly variable, represents the best of all these technical problems, the entire *matK* region has been evaluated to find a region of 600–800 bp that is highly variable, and avoids the mononucleotide repeats. The following are the important universal *matK* primers used widely. 1. *matK*-2.1aF ATCCATCTGGAAATCTTAGTTC Royal Botanic Gardens, Kew (2007) 2. *matK*-5R GTTCTAGCACAAGAAAGTCG Royal Botanic Gardens, Kew (2007) 3. *matK*-1018R GTACYACYGAARKATYBAGYSCACZhang et al. (2006) 4. *matK*-390F CGATCTATTCATTCAATATTT
Sacred trees

Sacred Trees (Sthalavrikshas) have a unique place in the fabric of Indian heritage. Trees symbolize knowledge and spirituality. In ancient times, the trees were worshipped along with Gods and established in the temples as Sthalavrikshas, i.e., trees associated with the deity in the temple and became an inseparable part of the faith. Such trees were guarded and their saplings from mother plants were given to other temples.

Sacred trees are important not only for their botanical, economical, medicinal, environmental, religious and mythical values but also form an important biological heritage of our nation that plays role in the conservation of environment and biodiversity.

The Sthalavrikshas constitute a part of genetic resources for the conservation of species diversity. Propagation of Sthalavrikshas in temples contributes to the conservation of floral diversity. Some trees are important for their economic role in ship building or in the timber industry, some for providing homes for various animals, birds and others for their medicinal and air purifying qualities.

*Stereospermum suavelons* (Bignoniaceae) is a medicinal tree species native to India (Troup R.S., 1986). The Bignoniaceae having about 100 genera with 800 species, are known for their antimicrobial, antiprotozoal, and anti-inflammaratory properties (Binutu O.A., et al., 1996; Onegi B, et al., 2002). Both the timber (Sandermanns W. 1957) and the root heartwood (Joshi k.c. et al., 1977) of *S. suaveolens* were found to contain lapachol, elicitor of contact dermatitis (Schulz K.H. et al., 1977)

Moreover, barks, flowers, roots and leaves of *S. suaveolens* are used by traditional healers, rural communities and pharmaceutical companies for remedies of diseases like heating, vomiting, eructation, piles, acidity, diarrhoea, gonorrhoea, loss of taste, malaria and other fevers (Troup R. S., 1986).

Materials and Methods

Sample collection

The materials used and the methods followed in the present investigation are described below. In Cuddalore district, young leaf samples of sacred trees, namely, *Stereospermum suavelons* DC. Pathiri (Bignoniaceae) were collected from Padaleshwarar Temple, Cuddalore for the isolation of genomic DNA.

DNA isolation, PCR amplification, DNA sequencing

DNA isolation was carried out by using modified CTAB (Cetyl Trimethyl Ammonium Bromide) method (Sambrook, J., and Russell, D. W. 2001), which was designed for plants producing large amounts of secondary metabolites, exudates, milky or latex and oil substances. As the chloroplast maturase K gene (matK gene) is one of the most variable coding genes of angiosperms and suggested to be a “barcode” for land plants, in the present study, the same was used.

Gradient PCR was performed using isolated genomic DNA of this study to determine the optimum annealing temperatures of the Primers used, namely *matK*472F (5′-CCC RTY CAT CTG GAA ATC TTG GTT C-3′) and *matK*1248R (5′-GCT RTR ATA ATG AGA AAG ATT TCT GC-3′). This primer was obtained from Genei, Bangalore. The PCR reaction mixture was consisted of 10x Taq buffer, dNTP mix 100 mM, Taq polymerase (3U / µl) and 5 – 50 ng of template DNA. Thermal cycling conditions for gradient PCR were as follows: Initial DNA denaturation at 94°C for 3 minutes, followed by 40 cycles of final DNA denaturation at 94°C for 30 second, primer annealing temperature at 48-52°C for 40 second, DNA strand extension at 72°C for 1 minute, and final extension at 72°C for 10 minutes. The PCR products were verified by electrophoresis in 1 % agarose gel stained with ethidium bromide.

The PCR products were sent to the Xcelris Labs Ltd. Sydney House, Premchand Nagar Road, Bodakdev, Ahmedabad 380054, and India for DNA sequencing and the sequences were obtained. All the obtained sequences were submitted to gene bank. (Note: In 10 µl sequencing reactions, using 3 µl sample and 0.25 µl BigDye Terminator V 3.1, methods ABI – 3730 XL sequencer gave good amplification and sequences).

Sequence Analysis

Each obtained sequence was entered into Gene Bank’s BLAST search function (http://blast.ncbi.nlm.nih.gov/Blast.cgi), using Megablast parameter to assess the similarities and differences between the coded sequences of the sacred plants of the present study and with other related taxa. The similarity was also examined for the closest matches among them.

Phylogenetic analysis

All the sequences were aligned using ClustalW and the genetic distances were computed using MEGA 5 software, and the phylogenetic tree was built using NJ method. The bootstrap test with 1000 replicates was applied to assess the reliability of phylogenetic trees.
Results

DNA isolation

For the isolation of genomic DNA of the species studied, the modified protocol of Cetyl Trimethyl Ammonium Bromide method (CTAB method) was used and good isolation was obtained for the sample.

PCR amplification

The Gradiant PCR amplification was performed for the isolated genomic DNA of five species of sacred trees mentioned earlier by using matK gene and its primers, namely matK472F and matK 1248R, and the amplifications were obtained for the species studied.

DNA Sequence

The amplification of Gradient PCR product was strong enough for isolation of bands or direct sequencing and in the present study, the DNA sequences were done at Xcelris Labs Ltd. Sydney House, Premchand Nagar Road, Bodakdev, Ahmadabad 380054, India. The methods of ABI – 3730 XI sequencer gave a success rate of 90-95 % and read length of 700 bases or more.

*Stereospermum suavelons* DC.

AGCAATTTTTTATTTAACAGGATTCTTTCTCAACGGAATGATTTGAAATTGGAATAGTCTTAATACTCCA
AAGAAAGCAGTTCCTCTTTTCTAAAAAGAAATCAAGTTATTCTTATTTATATATTTCTATG
TATGGAAATGAAATCCATTCTTTTTCTTTGTCCTCTACTGTAACCAAGACTTGTCTTCTAAATCA
GAGGCTCTTCTCTGAAACAGGATCTTAATTTCTATTGAAAAATAGAAGCTATTGTGAACTCTAAG
TTAAGGAATTTTCAGGGCAACCTATAGTTTCGTCAGGAACCTTGCTGCTATTATTAAGGTATCAA
AGAAATCCCTCCTTCCTAAAAGGGAGCCTTCTCTCTCCTCCTGAAATCAATGGAATGTTACTCCTTAC
TTTTTCAGAATGCGCATTCTTTCTGTGTTCTCATCAGAAGGATTTATATATAGACTAATCCATTCAAAT
CATTCCCTTAATTTTTCGTCAGCTTCTTTTCTAAAATGGCAATAACCCCTACGTGATACGGGAGTCAAG
TTCTAGAAAATTCATTCTCCTAATCTAAATTGCTATTTAGAGGTTCGATACCTTCTACCTTTTTCTTTAAT
TTTTTCGTCATTCTCGGTCGAGAACAAGCAATTTTGTGAAATTGGAACCTTCTCTACCTTTTTCTTTCTCATG
ATTCATTGCGAGACAAGCA

Fig. 1 - Padaleshwar Temple (Cuddalore)  
Fig. 2 – Sthalavriksham – *Stereospermum suavelons* DC
Fig. 3a. DNA Sequences of *Stereospermum suavelons* DC

Fig. 3b. Electropherogram of *Stereospermum suavelons* DC
Fig 4: BLAST Formatting Results of *Stereospermum suavelons* DC
MULTIPLE SEQUENCE ALIGNMENT

Fig. 5: Multiple Sequence Alignment of *Stereospermum suavelons* DC using clustalW method.

Table 1: BLAST identification of the sample (*Stereospermum suavelons*)

<table>
<thead>
<tr>
<th>Accession No.</th>
<th>Accession Description</th>
<th>E value</th>
<th>Max ident</th>
</tr>
</thead>
<tbody>
<tr>
<td>JF270837</td>
<td><em>Kigelia Africana</em> voucher OM217 maturase K(matK) gene, partial cds</td>
<td>0.0</td>
<td>98 %</td>
</tr>
<tr>
<td>JF270862</td>
<td><em>Markhamia zanzibarica</em> voucher OM629 maturase K (matK) gene, partial cds</td>
<td>0.0</td>
<td>98 %</td>
</tr>
<tr>
<td>HQ384519</td>
<td><em>Catalpa aff. speciosa</em> Olmstead 88-003 trnK gene, partial sequence; and maturase (matK)gene, partial cds; chloroplast</td>
<td>0.0</td>
<td>97 %</td>
</tr>
<tr>
<td>HQ384523</td>
<td><em>Eccremocarpus scaber</em> trnK gene, partial sequence; and maturase (matK) gene, partial cds; chloroplast</td>
<td>0.0</td>
<td>98 %</td>
</tr>
</tbody>
</table>
Table 2: The sequences retrieved from NCBI – gene bank for the study of the sample has given in the table.

<table>
<thead>
<tr>
<th>S. NO.</th>
<th>ACCESSION NO.</th>
<th>SPECIES</th>
<th>AUTHORS</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>JF270837</td>
<td>Kigelia africana</td>
<td>Yessoufou et al., 2012</td>
</tr>
<tr>
<td>2</td>
<td>JF270862</td>
<td>Markhamia zanzibarica</td>
<td>Yessoufou et al., 2012</td>
</tr>
<tr>
<td>3</td>
<td>HQ384519</td>
<td>Catalpa aff. speciosa Olmstead</td>
<td>Refulio-Rodriguez, N.F. and Olmstead, R.G. 2012</td>
</tr>
<tr>
<td>4</td>
<td>HQ384523</td>
<td>Eccremocarpus scaber</td>
<td>Refulio-Rodriguez, N.F. and Olmstead, R.G. 2012</td>
</tr>
</tbody>
</table>

Morphologically it was identified that the sample 2 is Stereospermum suavelons. The matK gene sequence for Stereospermum suavelons was not available in gene bank and the present study sequence showed more similarity to the Kigelia africana species, belonged to Bignoniaceae family. The Stereospermum suavelons also coming under the same family. The other species used in the present study were Markhamia zanzibarica, Catalpa aff. speciosa Olmstead, Eccremocarpus scaber and Kigelia Africana. All these species were belonging to Bignoniaceae family (Fig. 7). The Bootstrap value 76 – 93 % was shown in the tree. (Fig. 6)

Fig. 6: Phylogenic tree represents the evolutionary relationship between the plant species of Stereospermum suavelons DC.

Discussion
DNA barcoding
DNA barcoding is a molecular-based identification system, recently introduced in the scientific community. The method is not completely new to science, but the real innovation is not in the discrimination system itself.

DNA barcoding is promising in providing a practical, standardized, species-level identification tool that can be used for different study including forensic analysis (Lahaye et al. 2008) and animal diet determination when the food is not identifiable by morphological criteria.

Given the pace of advancement in technology, it is not unrealistic that in the span of a few years we may be using a barcoding tool for routine identifications, discovering new species, solving ecological puzzles, controlling the pathways of invasive species, and for quality control in the food and herbal industries.

The characterization of nucleotide and amino acid substitution along the gene may also provide information on site-dependent probabilities of nucleotide substitutions. Such information could provide a guide to the regions to be used in phylogenetic analysis since methods of phylogenetic inference assume the probabilities of replacements are independent of site (Clegg et al., 1994).

Sacred trees
In India, communities have been involved as custodians or stewards in conserving germplasm of crop plants, horticulture, medicinal plants and some species of animals in situ over the years. For example, some species are protected by the communities in the form of sacred groves, which may contain endangered species. Communities do not use plant species or parts of the species in the sacred groves. Some species of plants have religious significance among Hindus. For example, Aegle marmelos, popularly called Bilva tree, is conserved by planting it in the premises of Hindu temples. Another example: Ocimum tenuiflorum, popularly called Tulsi, has some medicinal properties. It is protected by planting it in the backyards of households to have ready access to the plant for medicinal purposes. In fact, Hindu accords a sacred status to the plant and worships it so that it is not neglected.

Scientists recognize the need to describe and document the wide-ranging biodiversity that remains to be explored in India. They stated that DNA barcoding is a valuable technology for this purpose. In India, as in many other countries, there is a perception that taxonomy is a less attractive specialty (Prathapan et al., 2006). Perhaps one of the reasons is that scientists seem to attach more ‘glamour’ to a research career in molecular biology (Haribabu 2000). Research relating to DNA barcoding may change the situation and fill the shortfall in the number
of taxonomists. This calls for focused training programs that attract young scientists.

Scientists mentioned that a democratic decision has to be taken regarding which species in the country have to be barcoded, given the fact that there are endemic, endangered and commercially significant species. Since the legislation treats all bio-resources as national resources, accessing the species for barcoding has to be based on formal approvals at different levels. The scientists are of the view that the barcoded information has to be kept in the public domain except in the case of some endemic species which have commercial applications. In this connection, they strongly argue that in any international collaboration Indian scientists should not part with samples of species to be barcoded to collaborating scientists in foreign countries. The samples should be kept in a national repository, or bio – bank and the sequence information may be shared with scientists in other countries. Based on the vital reasons discussed above, it is necessary to take DNA barcode studies on sacred trees to document their reliable discrimination and identification for future use as they are socially, economically, culturally and religiously important one, and many of them are in the state of endemic and endangered. The Sacred Trees preserved through millennia by our ancestors as potential bio resources should be respected and conserved for the future generations. Hence, the present study was undertaken.

DNA Extraction

Generally, DNA extraction is performed using mainly fresh young leaves. The lower amplification and sequencing success in a study could be explained by the fact that the quality of DNA extracted from young leaves might be influenced by several factors, such as drying and storage conditions and preservation procedures used in herbaria (Erkens et al. 2008).

One of the early studies showed that leaflets were clipped from live plants, dried in silica gel, and then stored at - 80°C. Whole genomic DNA was extracted using DNeasy Plant Mini Kits (Qiagen, Valencia, CA) or a modified CTAB method (Doyle JJ, Doyle JL. 1987) from fresh or frozen tissue.

Today, as literatures revealed, a modified Cetyl Trimethyl Ammonium Bromide method (CTAB method) is followed in majority of the DNA barcode studies to isolate DNA from plant samples and in this study also a modified CTAB method suggested by Sambrook and Russal (2001) was followed to isolate DNA from fresh leaf samples of six sacred trees studied.

**PCR amplification**

The Gradient PCR amplification was performed for the isolated genomic DNA of the present study using the matK primers, namely matK472F and matK 1248R and the Gradient PCR was performed to determine the optimum annealing temperatures of the primer matK472F (5‘-CCC RTY CAT CTG GAA ATC TTG GTT C-3’) and matK1248R (5‘-GCT RTR ATA ATG AGA AAG ATT TCT GC-3’). In the early studies, the matK primers 472F and 1248R gave 94% and 96% amplification success rates respectively. In this study also it was confirmed as they gave 95% of amplification rate (Yu et al., 2011). In the present study, the PCR reaction mixture consisted of 10x Taq buffer, dNTP mix 100 mM, Taq polymerase (3U / µl) and 5 – 50 ng of template DNA and the thermal cycling conditions for gradient PCR were as follows: Initial DNA denaturation at 94°C for 3 minutes, followed by 40 cycles of final DNA denaturation at 94°C for 30 second, primer annealing temperature at 48-52°C for 40 second, DNA strand extension at 72°C for 1 minute, and final extension at 72°C for 10 minutes. The PCR products were verified by electrophoresis in 1 % agarose gel stained with ethidium bromide. In the early studies, depending upon the plant materials, there were slight changes in the temperature as well as time duration for denaturation, annealing and extension and they were as follows. Initial DNA denaturation ranged from 93 to 95°C for 1 to 4 minutes and 35 to 40 cycles. Final denaturation ranged from 93 to 95°C for 30 seconds. For primer annealing, it was between 48 to 55°C for 30 to 40 seconds. For DNA extension, it was seen as 72°C for 45 seconds to 1 minute and for final extension it was noted as 72°C for 5 to 10 minutes (Kevin et al., 2011; Xiaorong Guo et al., 2011; Jing YU et al., 2011).

Important criteria for evaluating the suitability of DNA barcode are amplification and sequencing success (CBOL Plant Working Group 2009). In this regard, the coding matK gene region had the best performance in this study with 90 - 95% success in sequencing rate and this result is in consistent with the previous studies on land plants that have reported rates from 78.5 % to 99 % (Kress & Erickson 2007; Fazekas et al. 2008; Gonzalez et al. 2009; Hollingsworth et al. 2009; Kress et al. 2009).

**Barcode gene(s) and primer(s)**

DNA barcoding, a concept that has recently become popular, is characterized by using one or a few DNA fragments to identify different species (Kress et al., 2005). The mitochondrial cytochrome c oxidase subunit 1 (COI) gene was selected to be a DNA barcode for animal species (Hebert et al., 2003). However, consensus is still to be reached regarding gene fragments for a plant DNA barcode, although the Consortium for the Barcode of Life (CBOL) Plant Working Group (2009) has suggested matK + rbcL. The difficulty in selecting specific gene(s) to be a plant barcode is due to the imperfection of any gene from either the chloroplast, mitochondrial, or nuclear genomes. The plant mitochondrial genes evolve slowly, and therefore, are ineffective for distinguishing between different plant species. Plant nuclear genes often occur in multiple copies and are highly variable, making the design of universal primers difficult. The search for a plant DNA barcode has focused on genes of the chloroplast
The matK gene with its underlying features represents a molecule that has strong potential in providing insight into evolutionary and systematic problems at various levels. The 1500–bp size and the high rate of substitutions make the gene a valuable source of information for addressing systematic and evolutionary questions at various taxonomic levels. The presence of a relatively conserved 3’ region and a less conserved 5’ region provide two sets of characters from the matK that can be used at a different taxonomic hierarchy from the tribal to the kingdom level.

As matK is one of the most rapidly evolving plastid coding regions and it consistently shows high levels of discrimination among angiosperm species and suggested as the best plastid option for a DNA barcode sequence that has good priming sites length and interspecific variation, in the present study it was used to barcode the species studied.

**matK primer**

A desirable DNA barcode must simultaneously possess enough sequence variations for species identification and adequate conserved flanking sites for the designation of the universal primers (Stoechle 2003). In systematics, the full-length matK sequence is amplified and sequenced with primers designed within the trnK region (Wang et al., 2006; Li & Zhou, 2007); however, for DNA barcoding, a fragment of 600–800 bp is usually sufficient.

The success rate of amplifying the matK regions with the universal primer pairs in higher plants were the highest among all the candidate DNA barcodes (Kress et al., 2005; CBOL, 2009). In the present study, the matK primer pairs, namely matK472F, matK1248R were used along with primer 390F/1326R as a control to get good amplification and sequencing of matK region of the samples studied and good amplification and sequence were got for the species.

Although matK472F + matK1248R worked reasonably well for the samples tested in this study, its usefulness for all angiosperms needs to be evaluated. To avoid using too many degenerate bases, one has to ignore rare base substitutions. More specific primers can be designed by modifying matK472F and matK1248R according to the desired sequence.

**DNA sequence**

DNA sequencing has emerged as one of the most utilized molecular approaches for inferring phylogenies because of the direct comparison of the nucleotide sequences and the relative ease of interpreting the sequence information. The cycle sequencing method used in this study directly sequenced the DNA by the PCR method.

**Sequence analysis**

Currently, there is no standardize method for comparing unknown sequences to reference sequences. The generally used BLAST algorithm (Altschul et al. 1997) is not specifically designed for barcoding, but it has repeatedly been used for this purpose in recent years (e.g. Ford et al. 2009), and comparisons based on test datasets show that it does not perform worse than other methods (Little & Stevenson, 2007). A disadvantage of BLAST is that there are no statistical methods that can give a measure of the accuracy of identifications (Munch et al. 2008). However, the E-value and maximum identity are two statistics that can be used as an informal measure of the likelihood of an identification being correct. In general, one can assume that the closer a hit approaches 100% in sequence identity (and an E-value of 0), the more likely it is to have been correctly identified to species as well. However, there is a possibility that hits scoring 100% in sequence identity may be incorrect, if there are closely related species in the target geographical area that were not included in the reference database. Based on the reasons, in the present study too, the BLAST algorithm was used to identify, compare and discriminate the species studied and it showed fare results.

**Sequence retrieval**

Sequence retrieval leads to the question of what quality of match is required to use barcodes for identification. A match of 100% between a query sequence and a reference sequence is unambiguous at one level – each base pair is exactly matched. However, if the query sequence is 150 base pairs long, and the reference sequence is 2000 base pairs long, the 100% match might not be a meaningful one. The match might be along a part of the gene region that is highly conserved, with little to no variation among many species.

**Phylogenetic tree**

In the present investigation, the DNA sequences were analyzed using BLAST for pairwise sequence alignment and using clustalW for the multiple sequence alignment. The BLAST also used for identification of species using matK gene. The matK had greater resolving power of single region barcode, dominated under NJ method. In this, the basic sequence statistics, including nucleotide frequencies, transition/transversion (ns/nv) ratio and variability
in different regions of the sequences were computed by MEGA, which is an integrated tool for automatic and manual sequence alignment to construct phylogenetic tree, to estimate the rates of molecular evolution and to test evolutionary hypothesis.

In the present study, the phylogenetic trees (Dendrograms) were constructed for the species studied and their related taxa by using Neighbor joining (NJ) method to study the identification, discrimination, closeness and the evolutionary trend among them and the constructed trees fulfil the needs of this study.

**Conclusion**

DNA Barcoding is an effective taxonomic tool for the identification of sacred trees of *Stereospermum suavelons*. In the present investigation, DNA barcodings were made for the sacred trees of Main (Big) Temples in Cuddalore district, namely *Stereospermum suavelons* to document their reliable identification, discrimination, similarities and evolutionary trend among them and with their related taxa for future use.

**Fig. 7:** Overall Phylogenetic tree represents the evolutionary relationship between the plant species of *Stereospermum suavelons* DC.

**Reference**


complements the non-coding trnH-psbA spacer region. PLOS one 2: e508.


