Phylogenetic assay and pollen structure of few species of the genus amaranthus L.

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Abstract

In Amaranthaceae, morphology based identification methods are usually time consuming and may sometimes lead to misidentification and always may not provide good resolution at the species levels. The phenotypic variability of the taxa may lead to misidentifications and creation of new false identity. DNA sequencing has been used to explain evolutionary relationships for more than 20 years in molecular systematics. The aims of DNA barcoding include identification of known specimens/species and new discovery of unknown plant species for enhancing taxonomy for the good of the science and betterment of society. The study basically emphasised on palynological studies and molecular profiling of Amaranthus species using universal markers rbcL and matK. The pollen morphology of the species of Amaranthus shows significant differences in polar length and equatorial diameter whereas the aperture was pantoporate with evenly distributed microspines. The phylogenetic assay showed bootstrap value of 96 and 98 for matK while 79 and 98 for rbcL dendrograms. All barcodes yield quality sequences.

Keywords: DNA Barcoding, *rbc*L, *mat*K, Palynology.

Introduction

The flowering plant family Amaranthaceae which corresponds to the classical family Amaranthaceae Juss. 69 genera and 772 species as well as the twice larger related family Chenopodiaceae have been subject to repeated taxonomical revisions from the time they were first described (1789 and 1799) respectively to the present¹⁻³. Recently it has been proposed to combine them into one large family Amaranthaceae as a result of molecular analysis⁴. The morphology of pollen grains, is an important source of information for plant systematics. In certain cases, palynological data have been crucial for taxonomic conclusions⁵. But for closely related species of the genus Amaranthus L., such data can lead to confusions regarding the classification though pollen grains morphological studies shows very slight variation. Traditionally, most plant identifications are based on morphological characters, but such identification is not always reliable and efficient⁶.

DNA barcoding basically relies on short and standardized gene regions for the identification of plant species. The agricultural and horticultural applications of barcoding such as for marketplace regulation and copyright protection remain poorly explored. This study examines the use of effectiveness/quality of the standard plant barcode markers (*matK* and *rbcL*) for the identification of plant species that are medicinal and are of interest for pharmaceutical industry both in private and public nurseries and to authenticate the medicinal plants. A wide variety and different range of molecular techniques including

the random amplified polymorphic DNA (RAPD), amplified fragment length polymorphism in plant (AFLP), restriction fragment length polymorphism of the selected taxa (RFLP), microsatellite and single nucleotide polymorphism (SNP) have been proposed to identify plant species/specimen and cultivars⁷ 11. DNA barcoding method has emerged as a relatively new/novel and perhaps more universal tools with which to analyze diversity of both plants and animals and to fix specimens to their respective species even in the absence/ unavailability of key morphological diagnostic features^{12,13}. Although there are still some reserves against the performance of DNA barcoding as compared, for example, to morphology, an early study, through a thorough comparison of DNA barcoding and morphology-based species identification recorded a number of limitations to the morphology particularly when it comes to cryptic species². As a taxonomic tool this technique has widely acceptable and also has been successfully used in large scale biodiversity projects where regional flora and fauna are documented including regulated and threatened taxa14-¹⁷. Although a number of plant loci including, trnHpsbA, rpoc1, rpoB, trnL, rbcL and matK were initially proposed as potential plant barcodes based on assessments of similarity with genebank and recoverability, sequence derived quality and levels of plant species discrimination, the Consortium for the Barcode of Life recommended the 2-loci combination of rbcL + matK as the most standard plant barcode for analysis 18-23.

The DNA barcode data generated in the present study will serve in the future in commercial agricultural and medicinal plant Vol. **6(6)**, 20-25, June (**2017**)

industries for the purpose of control of counterfeited product, and could also serve in ecological studies of local flora as demonstrated elsewhere²⁴⁻²⁶. In addition, the evolutionary processes such as hybridization especially interspecific hybridisation and different polyploidy are common in plants, so such species boundaries are difficult to define^{27,28}. Thus, screening for single or multiple regions using appropriate primers are necessary for DNA barcoding studies in the nuclear and plastid genomes in plants that has been an important point of research. Since matK and rbcL sequences from Amaranthaceae were the most abundant in GeneBank, they were chosen for the study. Similarly authentication is a critical aspect of research in selection of plants for study, so an attempt was done to authenticate the Amaranthus L. using the palynological and DNA barcodes data inorder to make a unique identity among the plants.

Materials and methods

The fresh leaves of of Amaranthus spinosus L., Amaranthus caudatus L., Amaranthus tricolor L., Amaranthus dubius Mart., and Amaranthus viridis L. were used for isolating genomic DNA. Pollen grains were collected from the anther of 15 flowers of each species and fixed in glacial acetic acid for acetolysis as per the technique²⁹. The sculpturing pattern, values of P(pollen axis ratio) and E(equatorial diameter) and aperture number were viewed and data were measured using the Scanning Electron Microscope (SEM) JSM-6390 LA coated with gold, examined and photographed using JSM-6390 LA. All the experiments were evaluated statistically with SPSS version 20.0, the results were represented in mean±SEM (standard error of mean). One way analysis of varience (ANOVA) followed by DMRT to find out any significant difference in pollen characters among five Amaranthus sps. resulting from analytical experiments carried out. P value less than 0.05 were adopted as statistically different.

The DNA was isolation by using branded Sigma kit GenElute Plant Genomic DNA Mini-preparation Kit. GeneAmp PCR System 9700, Applied Biosystems, a PCR thermal cycler for PCR amplification, using the primers of *rbc*L and *mat*K. The primer details were given in table 1 and PCR amplification data for conditions provided are given in Table-2.

Table-1: The universal primers rbcL and matK and their sequences.

Target gene	Primer	Direction	Sequence strand $(5' \rightarrow 3')$
matK	390 f	forward	CGATCTATTCATTCAA TATTTC
maik	1326r	reverse	TCTAGCACACGAAAGT CGAAGT
nh aI	rbcLa_f	Forward	ATGTCACCACAAACA GAGACTAAAGC
rbcL	rbcL724 _rev	Reverse	GTAAAATCAAGTCCAC CRCG

Table-2: PCR amplification profile.

matK		rbcL	
98 °C	-30 sec	98 °C	-30 sec
98 °C 50 °C	$\begin{array}{c} -5 \text{ sec} \\ -10 \text{ sec} \end{array}$	98 °C 60 °C	- 5 sec -10 sec \\ \delta 0 cycles
72 °C	-15 sec	72 °C	-15 sec
72 °C 4 °C	- 60 sec - ∞	72 °C 4 °C	- 60 sec -∞

Sequencing reaction was done in a PCR thermal cycler named GeneAmp PCR System 9700, Applied Biosystems) using the BigDye Terminator v3.1 Cycle sequencing Kit of Applied Biosystems, USA model. The sequence quality was checked Applied Biosystems Sequence Scanner Software v1. Using Geneious Pro v5.6 Sequence alignment and required editing of the obtained sequences were carried out ³⁰. The DNA sequences of *Amaranthus* spp. under study were subjected to BLAST analysis for better identification at the species level. Sequences obtained were aligned and compared using Multiple Sequence Alignment software program of BioEdit Sequence Alignment Editor, CLUSTAL W Multiple Alignment^{31,32}. DNA barcodes namely *rbc*L and *mat*K for constructing neighbouring tree model using MEGA 7.0, and a tree was constructed using a combination of rbcL and matK³³.

Results and discussion

The SEM photomicrographs of radial, longitudinal axis, pore aperture, aperture number and distance are shown in Figure-1. The pore structure is similar to the Type II of that is the pores possess microspines, granulate surface and are evenly spread, with numerous pores³⁴. The present palynological result support existing data which have indicated that Amaranthus is pantoporate in terms of pore number. Most of the species examined in Amaranthaceae have pantoporate with Amaranthus type of pollen³⁵. The aperture of the pollen examined showed great variation in Amaranthus tricolor L., were their size and number is reduced as reported that there is a *Amaranthus* type of porate aperture in the family Amaranthaceae³⁶. The pollen grains of Amaranthus species are spheriodal to oblate spheroidal, pantoporate and pantotreme, with this a key is prepared inorder to make a comparison and identification among the species.

Shape- Spheroidal
Diameter 22.12-22.21μm
Aperture distance 3.23 -3.66 μm Amaranthus viridis L.
Shape- Oblate-spheroidal
Diameter 21.80-21.84μm
Aperture distance 1.00 -1.22 μm Amarantus spinosus L.
Diameter 18.00-19.86 μm
Aperture distance 2.05-2.57 μm Amarantus tricolor L.

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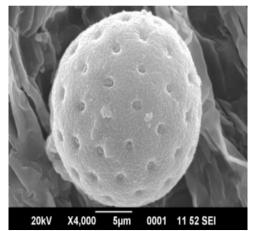


Figure-1: SEM photograph of Amaranthus L. pollen.

This study demonstrated differences in pollen characteristics among the *Amaranthus* species. However, pollen morphology/palynological data will have limited use in species identification because of similarities across the species. To

increase the usefulness of pollen morphology in species identification, additional analysis of naturally occuring population as well as sampling from different geographical regions would be needed to account for the study³⁷. The statistical analysis of pollen characters using SPSS version 20.0 by Duncan's Multiple Range test ($\alpha/p = 0.05$) showed that there is a significant difference between the species of *Amaranthus* L. as because the p <0.05. Thus it infers that the palynological characters are significant among five selected species. Similar attempt was made to pollen morphological differences in *Amaranthus* L. species and the hybrids formed by interspecific hybridisation where the mean value with same letter are not significant as reported³⁷.

From the DNA barcoding studies using the universal primers *rbcL* and *matK*, the study revealed the correct differentiation among *Amaranthus* species. It has been proved experimentally that DNA markers can act as a powerful/authentic tool for identification of cultivars and species for phylogenetic evaluation^{38,39}.

The accession numbers of the DNA sequences submitted and size of the sequences are given in the Table-5. The tree constructed by neighbour joining tree feature using MEGA 7 is given in the Figure-2.

Table-4: Pollen characters among 5 *Amaranthus* species

Species	Amarantus viridis L.	Amarantus spinosus L.	Amarantus dubius Mart.	Amarantus caudatus L.	Amaranthus tricolor L.
Pore distance	1.254 ^d ±0.006	1.19 ^d ±0.022	1.172°±0.19	1.226 ^d ±0.01	1.146 ^e ±0.02
Aperture distance	3.448°±0.081	2.924°±0.146	2.672 ^d ±0.07	2.73°±0.06	2.534 ^d ±0.07
Diameter(1)	20.872 ^b ±0.85	20.634 ^b ±0.766	19.91°±0.48	20.006 ^b ±0.49	19.652°±0.45
Diameter(e)	21.086 ^b ±0.321	21.376 ^b ±0.257	21.308 ^b ±0.32	21.386 ^b ±0.36	21.364 ^b ±0.37
Aperture number	24.2°±0.74	25.8°±0.374	28 ^a ±0.44	30°±0.89	16.8°±0.58

Each value that is expressed in as mean \pm Std. Error done in triplicates. Data analysed by SPSS version 20.0 by Duncan's Multiple Range test (α /p=0.05). Mean values followed by the different Superscript in the columns are significantly different among the pollen characters.

Table-5: Accession number of sequence submitted in the GenBank

Tuble et l'iceession number of	sequence submitted in the Genbar	1	
Plant samples	Place of Collection	rbcL Accession No.	matK Accession No.
A.viridis	Edakochi	KJ773261	KJ772535
A.spinosus	Thevara	EF590496	EF590394
A.dubius	Thrissur	KX090210	KX090202
A.caudatus	Boat jetty	KX090209	KC747133
A. tricolor	Kodugallur	JF940812	JF953165

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Table 0. Scuuchce data Holli Celibalik 101.) iliculciilai Diai	Sequence data from GenBank for 5 medicinal p	Jants
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Species	Gene	%of match with Gen Bank
A.viridis	matK	99% with 4 species of Amaranthus
	rbcL	99-97% to other Amaranthus spp.
A.spinosus	matK	99% match with 3 species
	rbcL	100% similar
A.dubius	matK	100-99% similar
	rbcL	100-99% to multiple Amaranthus spp.
A.caudatus	matK	98% to 4 Amaranthus spp.
	rbcL	100-99% to multiple Amaranthus spp.
A. tricolor	matK	100% to 3 Amaranthus spp.
	rbcL	100% with multiple Amaranthus spp.

[%] match shows how closely the barcode sequences matched with the other accessions in GenBank for Amaranthus L.

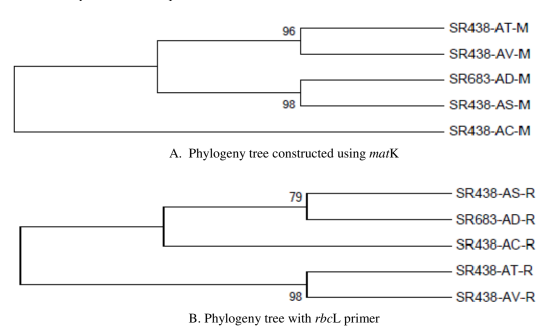


Figure-2: Phylogenetic tree constructed using two universal barcodes.

The currently available DNA sequences in the GenBank demonstrate the different challenge of discrimination power among the species. Table-5 lists out the barcoding sequences of five pharmaceutically important medicinal plants and indicates how well each sequence matches or identifies the selected plant. The results can vary by species and by gene region or target site. For example, using the *rbc*L shows good percentage of similarity with some *Amaranthus* species than *mat*K. Both *mat*K and *rbc*L nucleotide sequences had been previously successful for determination of phylogenetic relationships among taxa of other angiosperm plants ⁴⁰⁻⁴².

Using the Neighbor-Joining method the evolutionary history was inferred⁴³. As with the case of tree constructed using matK, the sum of branch length = 0.01871936 is shown in the optimal tree. While tree constructed using rbcL primer showed branch length of 0.00720922. The percentage of replicate trees in which the associated taxa clustered together in the bootstrap test (with

100 replicates) are shown next to the selected branches⁴⁴. The tree is drawn to scale, with branch lengths in the same units as those of the evolutionary distances that are used to infer the phylogenetic tree constructed. The analysis basically involved 5 nucleotide sequences and all positions containing gaps and missing data were avoided. Evolutionary analyses were conducted in MEGA7³³.

The robustness of the trees and branch support was estimated by bootstrap analysis. In the present study the phylogenetic assay showed bootstrap value of 96 and 98 for *mat*K while 79 and 98 for rbcL dendrograms. The boot strap value was found to be more for *mat*K than *rbc*L primer. The *mat*K sequence provide good resolution within many angiosperm orders from earlier reports. Combination or combined analysis of *mat*K and other rapidly evolving DNA regions with available multi-gene data sets will have the strong potential to enhance the resolution and internal support in deep level angiosperm phylogenetics and

provide additional insights into angiosperm evolution⁴². Similarly from the dendrogram of *rbc*L and *mat*K, *A. spinosus* and *A. dubius*, *A. tricolor* and *A. viridis* showed same clade and *A. caudatus* showed distinct clade as shown in Figure-2. Thus use of these molecular markers in addition to the classical methods (palynology/morphology) provides more positive identification technique to locate new varieties and both can be used to authenticate the plants among the species level in more scientific way for providing better outcome.

Conclusion

The palynological data as well as molecuar data using *rbc*L and *mat*K sequences shows that certain taxa *Amaranthus* L. show similarly and dissimilarly though they are morphologically similar. Constructing phylogeny tree using bioinformatics tool serve as a useful source of new information to both complement and evaluate morphological methods and as an aid to identifying those traditional morphological features that are taxonomically significant. Also from the findings the medicinal plants of pharmacologically important can be uniquely identified using DNA barcoding and provide quality control and standardization of the plant material supplied to the pharmaceutical industry for preparation of new drug formulation. Such identification is useful for avoiding the entry of different adultrants and substitutes for the preparation of Ayurvedic medicinal formulations.

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