Development of DNA barcodes for selected Acacia species by using rbcL and matK DNA markers

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A R T I C L E   I N F O

Article history:
Received 19 May 2020
Revised 28 July 2020
Accepted 11 August 2020
Available online 19 August 2020

Keywords:
Desertification
Acacia
Genetic diversity
Afforestation
DNA barcoding

A B S T R A C T

Acacia species are very important tree species in tropical and subtropical countries of the World for their economic and medicinal benefits. Precise identification of Acacia is very important to distinguish the invasive species from rare species however, it is difficult to differentiate Acacia species based on morphological characters. In addition, precise identification is also important for wood characterization in the forest industry as these species are declining due to illegal logging and deforestation. To overcome these limitations of morphological identification, DNA barcoding is being used as an efficient and quick approach for precise identification of tree species. In this study, we selected two chloroplast and plastid base DNA markers (rbcL and matK) for the identification of five selected tree species of Acacia (A. albida, A. ampiceps, A. catechu, A. coriacea and A. tortilis). The genomic DNA of the selected Acacia species was extracted, amplified through PCR using specific primers and subsequently sequenced through Sanger sequencing. In matK DNA marker the average AT nucleotide contents were higher (59.46%) and GC contents were lower (40.44%) as compared to the AT (55.40%) and GC content (44.54%) in rbcL marker. The means genetic distance K2P between the Acacia species was higher in matK (0.704%) as compared to rbcL (0.230%). All Acacia species could be identified based on unique SNPs profile. Based on SNP data profiles, DNA sequence based scannable QR codes were developed for accurate identification of Acacia species. The phylogenetic analysis based on both markers (rbcL and matK) showed that both A. coriacea and A. tortilis were closely related with each other and clustered in the same group while other two species A. albida and A. catechu were grouped together. The species A. ampiceps remained ungrouped distantly, compared with other four species. These finding highlights the potential of DNA barcoding for efficient and reproducible identification of Acacia species.

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1. Introduction

DNA barcoding is a widely accepted technique that focuses on nucleotide sequence based identification of plant species in an efficient and precise manner (Group et al., 2009). DNA barcoding have been developed as a system for species recognition and identification by using specific regions of DNA sequences (Hebert et al., 2003; Asif and Cannon, 2005; Newmaster et al., 2006; Kress and Erickson, 2007; Ratnasingham and Hebert, 2007; Fazekas et al., 2008; Lahaye et al., 2008; Pettengill and Neel, 2010). One of the major challenges faced by barcoding, is the ability to resolve sister species within a large geographical ranges. It has been suggested that a system based on any one or small number of chloroplast genes will fail to differentiate taxonomic groups with extremely low amount of plastid variations while it will be effective in other groups. Steven et al. (2009) successfully utilized barcoding in discriminating multiple populations among a sister species complex in pantropical Acacia subgenus across three subcontinents. The use of three chloroplast regions i.e. rbcL, matK and trnH-psbA successfully discriminated sister species within both genera and differentiated biogeographical patterns among populations from India, Africa and Australia. These findings clearly established the power of DNA barcoding for taxonomic and biogeographical studies, for identifying cryptic species as well as biogeographic patterns for resolving classification at the rank of genera and species level. The matK gene has ideal size, high rate of substitution, a large proportion of variation at nucleic acid level at the first and second codon positions, low transition/transversion ratio and the presence of mutational conserved regions. These features of matK gene are exploited to resolve family and species level relationships. Using matK, about 90% amplification was achieved in angiosperm by using single pair of primer while with gymnosperm the amplification percentage was 83% and cryptogram was 10% (Lahaye et al., 2008). Similarly, rbcL has been greatly utilized for phylogenetic analysis offering better results at family and class levels however, it showed moderate discrimination power at the species level (Fazekas et al., 2008; Lahaye et al., 2008; Hollingsworth et al., 2009; Chen et al., 2010).

Acacia is an economically important genus, belongs to family Fabaceae. There are 1380 species of Acacia and with about two third are native to Australia while others are found in tropical and subtropical regions of the World (Maslin et al., 2003; Orchard and Maslin, 2003; Muhammad et al., 2017, Muhammad et al., 2018). Acacia species are economically very important and well adopted to arid conditions and in agroforestry. These species are used for fuelwood, pulp, fodder, fiber, timber, medicine, enhancing the fertility of degraded soils due to its ability of fixing nitrogen, environmental amelioration, gums and tannins (Midgley and Turnbull, 2003). The genus Acacia consists of three subgenera; found in all tropics i.e. Acacia, Aculeiferum and Phyllodineae. These subgenera are differentiated based on morphological and genetic variation (Maslin et al., 2003). The morphological identification however is very difficult due to the large diversity of invasive, natural hybrids and rare cryptic species (Steven et al., 2009).

Consortium for the Barcode of life (CBOL) has reported that the matK and rbcL are the main barcode markers for accurate identification of plants and trees (Li et al., 2011). Therefore, the main objective of this study was to identify various Acacia species by using sequence specific markers (matK and rbcL) and to develop DNA barcodes.

2. Material and methods

2.1. Sample collection and DNA extraction

The research was conducted in the Center for Advance studies in Agriculture and Food security (CAS-AFS), University of Agriculture, Faisalabad, Punjab Pakistan. The open pollinated seeds from five Acacia species (A. albida, A. amplexica, A. coriacea, A. catechu and A. tortilis) were collected from premises of Pakistan Forest Institute (PFI), Peshawar, Khyber PakhtunKhwa (KPK), Pakistan. Acacia species with their local names, scientific name and their origin has been given in Table 1. The genomic DNA was extracted from the seedlings by using cetyl-trimethyl ammonium bromide (CTAB) method. The quantification and purity of extracted DNA were measured using NanoDrop (ND-8000) (Thermo-Scientific, Waltham, MA) and 0.8% agarose gel electrophoresis.

2.2. PCR amplification with matK and rbcL primers

High quality template DNA (20ng) was used for PCR amplification, using selected primers (Table 2). Amplifications were performed in PCR tubes using a C–1000 Touch Thermocycler (Bio-Rad). PCR reaction was carried out in a total volume of 20μl containing 20 ng/μl genomic DNA template, 0.2 mM of each primer, 2 μl Taq buffer (100 mM Tris-HCl, 500 mM MgCl2, pH = 8.3), 0.5μl Taq DNA polymerase (5U/μl, Thermo Scientific Americia), 2.5 mM MgCl2, (25 mM), 0.2 mM dNTPs (10 mM) and 7.5 μl double distilled deionized water. The optimized PCR profile for both rbcL and matK comprised of an initial denaturation at 95 °C for 1 min, subsequently 30 cycles starting with 95 °C denaturation for 30 sec, annealing for rbcL and matK at 50 °C and 45 °C respectively for 30 sec, followed with a final extension at 72 °C for 1 min. The amplified PCR products were resolved on 1.3% agarose gel and stained with ethidium bromide for 20 minutes. Amplicon sizes were confirmed by comparing with 1Kb DNA ladder (Thermo Fisher Scientific, USA). The PCR products was purified for Sanger sequencing by using FavorPrep™ PCR Cleanup Mini Kit (FAVORGEN, Cat# FAPCK001-1).

2.3. Sequencing and data analysis

PCR amplicons were purified and submitted for sequencing through Sanger method (Eurofins Genomics, Germany Gmbh).

| Common/ local name | Scientific name | Origin | Ecology |
|-------------------|----------------|--------|---------|
| Sufaid kiker      | Acacia albida   | Southwest Africa | Tropical thorn to subtropical regions and precipitation of 250 to 400 mm/yr |
| Australian kiker  | Acacia amplexes | Northwest Australia | Subtropical and tropical zones and precipitation of 400 to 600 mm/yr |
| Catha             | Acacia catechu  | Pakistan, India and Nepal | Sub humd to subtropical climate and precipitation of 500 to 2700 mm/yr |
| Desert oak        | Acacia coriacea | Northern Australia | Tropical and subtropical of Australia and precipitation of 300 to 500 mm/yr |
| Samaor            | Acacia tortilis | Africa | Arid to semi-arid and precipitation of 100 to 1000 mm/yr |

| Loci  | Primer | Sequences     |
|-------|--------|---------------|
| matK  | F      | 5’-CTTACATCTGGAAATCTTGGTT-3’ |
| rbcL  | F      | 5’-ATGCTACCATCGAAGGTTTC-3’ |
| rbcL  | R      | 5’-CCATGACTCCATCGCA-3’ |
After sequencing, the resultant sequences were edited manually and multiple sequence alignment was performed using bioinformatics software MEGAX 10.1 developed by Penn State University, USA. Most of the mismatched sites and gaps were excluded using SeqMan software (DNAStar). The sequencing data was further utilized to identify AT (Adenine + Thymine) and GC (Guanine + Cytosine) contents and SNPs for each species. In addition, data was also used to develop DNA barcodes for each species by using online DNA Barcode Generator (QR barcode) software (https://www.the-qrcode-generator.com/). Several DNA regions were used to generate effective DNA barcodes for Acacia Species as previously reported by different researches (Fazekas et al., 2008; Lahaye et al., 2008). Phylogenetic relationship among species was developed through cluster analysis in R Core Team (R Core Team 2013).

3. Results

The specific genomic fragments of all selected species of *Acacia* were successfully amplified by using both *rbcL* and *matK* primers with an average read length of 750bp and 950 bp (Figs. 1 and 2). Similarly, DNA sequencing for both *rbcL* and *matK* amplicons generated high quality sequences. Nucleotide composition of amplicons from selected species of *Acacia* revealed variability in AT and GC contents. In amplicons of *matK*, nucleotide composition of AT and GC for *Acacia* species *Acacia albida*, *A. ampliceps*, *A. catechu*, *A. coriacea* and *A. tortilis* were (62.7%, 57.3%, 66%, 44.1% and 67.2%) and (37.2%, 42.6%, 33.9%, 55.8% and 32.7%) respectively (Table 3 and Fig. 3). In *rbcL*, AT and GC contents were (56.9%, 54.5%, 52.2%, 56.8% and 56.6%) and (43%, 45.5%, 47.7%, 43.2% and 43.3%) respectively as compared to *matK*. Overall, in both markers (*rbcL* and *matK*), the average AT contents was higher (57.43%) than average GC (42.49%) contents (Fig. 3 and Table 3). In *rbcL*, the minimum, maximum, mean and standard error of K2P genetic distance between *Acacia* species were calculated to be 0.078%, 0.368%, 0.230% and 0.0140% respectively. Similarly, in *matK*, the minimum, maximum, mean and standard error of K2P genetic distance between species were 0.048%, 1.01%, 0.704% and 0.088% respectively (Table 4). Moreover, for sequence data analysis, the obtained sequences were subjected to BLAST (Basic local alignment search tool) in the NCBI nucleotide database, which showed maximum similarity range from 92 — 98% for *rbcL* and 88–97% for *matK*. In addition, NCBI database sequences with maximum similarity to the queries were downloaded for comparison to the experimental species. For comparison of *matK* sequences, the reference sequences, accession numbers and gene bank name for the *Acacia* species were *Faidherbia albida* (JF265429.1), *Senegalia catechu* (MH560438.1) and *Vachellia nilotica* (KX119249.1). The reference

![Fig. 1. PCR amplification products with matK primers resolved on 1.3% agarose gel electrophoresis. M representing DNA ladder lines, N is a negative control, lane 1–5 *Acacia albida*, *Acacia ampliceps*, *Acacia catechu*, *Acacia coriacea* and *Acacia tortilis*. Lane 6-8 are duplicate loaded samples of *Acacia*. An amplicon of 930 bp was amplified.](image1)

![Fig. 2. PCR amplified products using rbcL specific primers. M representing DNA ladder, lane 1–5 representing PCR amplification of *Acacia albida*, *Acacia ampliceps*, *Acacia catechu*, *Acacia coriacea* and *Acacia tortilis* respectively. Lane 6-10 are duplicate loaded products. An amplicon of 750 bp was amplified in all tested species.](image2)

![Fig. 3. Nucleotide composition in amplicons of *Acacia* species using matK and rbcL.](image3)
sequence for \textit{rbcL} marker matched with \textit{Faidherbia albida} (HM020737.1), \textit{Acacia grandicornuta} (EU812049.1), \textit{Vachellia tortilis} (MK290437.1) and \textit{Acacia cyclops} (JQ412187.1). No similarity was found in \textit{A. tortilis} and \textit{A. coriacea} for \textit{rbcL} and similarly, for \textit{matK} no sequence homology was observed for \textit{A. ampliceps} (Table 5). DNA sequences were subsequently used to develop phylogenetic relationship between \textit{Acacia} species.

After sequencing, all the resultant sequences of \textit{rbcL} and \textit{matK} were aligned in order to generate dendrogram which depicts genetic relationship among the selected \textit{Acacia} species. Phylogenetic tree based approach was used to classify the selected tree species of \textit{Acacia} using \textit{rbcL} and \textit{matK} markers. All the sequences were edited manually as well as using SeqMan software i.e. similar, mismatch sites and all the gaps were removed for proper screening of SNPs. Moreover, only those sequences were selected for detection of SNPs, which have shown differentiation in only a single base pair in all experimented tree species of \textit{Acacia}. A total of 42 unique SNP variant sites were found in five selected \textit{Acacia} species for differentiation through \textit{rbcL} marker. Based on \textit{rbcL} data analysis, the number of unique sites observed in the selected five \textit{Acacia} species were as following, \textit{A. albida} (0), \textit{A. ampliceps} (1), \textit{A. coriacea} (17), \textit{A. catechu} (2) and \textit{A. tortilis} (22) (Table 6). As described earlier that in \textit{matK} based characterization, the minimum, maximum, means and standard error of K2P genetic distance between the \textit{Acacia} species were, 0.048%, 1.010%, 0.704% and 0.088% respectively (Table 4). Further, phylogenetic characterization with \textit{matK} marker, revealed that \textit{A. coriacea} and \textit{A. tortilis} were linked closely, while the other two species \textit{A. albida} and \textit{A. catechu} were clustered together and the specie \textit{A. ampliceps} remained ungrouped and consequently found dissimilar to all other four \textit{Acacia} species (Fig. 5). Phylogenetic relationship among selected species of \textit{Acacia} using both markers showed that \textit{A. coriacea} and \textit{A. tortilis} were clustered together compared \textit{A. ampliceps}. Similarly, \textit{A. albida} and \textit{A. catechu} were also closely linked and grouped together combined analysis (Fig. 6). Through these unique SNPs, we developed barcode for selected \textit{Acacia} species by using \textit{rbcL} and \textit{matK} sequences separately. Both \textit{rbcL} and \textit{matK} markers identified \textit{Acacia} species differently supporting use of different DNA markers for species differentiation. When used, these barcode independently differentiated \textit{Acacia} species thus can be applied to identify species through barcode scanning apps on smart phones (see Fig. 7).

4. Discussion

\textit{Acacia} and their hybrids are an emerging forest tree species for fuelwood and pulpwood production due to high growth rate and

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### Table 3

| Markers | Acacia Species | AT (%) | GC (%) |
|---------|----------------|--------|--------|
| \textit{matK} | \textit{Acacia albida} | 62.7 | 37.2 |
|          | \textit{Acacia ampliceps} | 57.3 | 42.6 |
|          | \textit{Acacia coriacea} | 66 | 33.9 |
|          | \textit{Acacia catechu} | 44.1 | 55.8 |
|          | \textit{Acacia tortilis} | 67.2 | 32.7 |
|          | Average | 59.46% | 40.44% |

### Table 4

| Markers | Minimum | Maximum | Means | Standard error |
|---------|---------|---------|-------|----------------|
| \textit{rbcL} | 0.078 | 0.368 | 0.230 | 0.040 |
| \textit{matK} | 0.084 | 1.010 | 0.704 | 0.088 |

### Table 5

| Markers | Species | Gene Bank/BLAST | Max. ID(%) | Accession no. |
|---------|---------|----------------|------------|---------------|
| \textit{rbcL} | \textit{Acacia albida} | \textit{Faidherbia albida} | 96.78 | JF265429.1 |
|          | \textit{Acacia catechu} | \textit{Senegalia catechu} | 98.40 | MH560438.1 |
|          | \textit{Acacia ampliceps} | \textit{Vachellia nilotica} | 92.55 | KX119249.1 |
|          | \textit{Acacia tortilis} | No sequence similarity | – | – |
|          | \textit{Acacia coriacea} | No sequence similarity | – | – |
| \textit{matK} | \textit{Acacia albida} | \textit{Faidherbia albida} | 95.22 | HM020737.1 |
|          | \textit{Acacia catechu} | \textit{Acacia grandicornuta} | 88.52 | EU812049.1 |
|          | \textit{Acacia ampliceps} | No sequence similarity | – | – |
|          | \textit{Acacia tortilis} | \textit{Vachellia tortilis} | 96.14 | MK290437.1 |
|          | \textit{Acacia coriacea} | \textit{Acacia cyclops} | 97.39 | JQ412187.1 |
Markers Italic nucleotides show unique SNP variations observed in this study.

The barcode of selected species of Acacia based on variable regions of \textit{rbcL} and \textit{matK} markers. Italic nucleotides show unique SNP variations observed in this study.


table

| Marker | Acacia albida | Acacia ampluscepts | Acacia coriacea | Acacia catechu | Acacia tortilis |
|--------|--------------|-------------------|----------------|---------------|----------------|
| \textit{rbcL} | T | T | G | T | T |
| C | C | C | T | C | G |
| A | A | T | A | A | C |
| A | A | A | A | C | T |
| A | A | A | C | A | C |
| A | T | A | A | T | T |
| C | C | C | C | T | C |
| T | T | T | T | C | G |
| G | G | G | G | T | A |
| A | A | A | A | C | C |
| G | G | G | G | T | A |
| C | C | C | C | T | T |
| T | T | T | T | A | G |
| C | C | A | C | C | T |
| T | T | T | T | C | T |
| A | A | G | A | A | T |
| C | C | G | C | C | A |
| A | A | G | A | A | A |
| A | A | C | A | T | T |
| A | T | A | A | T | T |
| T | T | G | T | T | T |
| T | T | T | T | T | A |
| T | T | T | T | A | A |
| \textit{matK} | A | A | A | T | A |
| A | A | A | A | A | A |
| T | G | T | T | T | T |
| T | T | A | T | T | G |
| A | G | A | A | A | A |
| A | A | G | A | A | A |
| T | C | T | C | T | C |
| T | C | T | C | C | C |
| A | T | A | A | A | A |
| C | A | A | A | A | A |
| T | C | T | T | T | T |
| T | G | T | T | T | G |
| G | A | G | G | G | A |
| C | T | C | C | C | C |
| T | G | G | G | G | G |
| T | G | A | G | G | G |
| A | C | A | A | T | T |
| A | A | T | A | A | A |
| A | G | G | G | G | G |
| C | A | C | C | C | C |

High adaptability to various environments. Acacias are more desirable than Eucalypts due to low water requirement thus suitable for arid and semiarid regions of the World. However, for future sustainable biomass, fuelwood and pulp supply breeding of improved materials with desirable traits e.g. low lignin, low tannin, longer fiber, high wood density and straight bole require precise selection of superior trees. Precise identification of \textit{Acacias} based on DNA markers is highly desirable, to select superior species with beneficial characters. In post-genomic era with rapidly advancing sequencing technologies, molecular markers offer a reliable and reproducible method for identification of tree species based on DNA sequences. To overcome the limitations of morphological markers to differentiate closely related species, we used \textit{matK} and \textit{rbcL} markers for efficient and reliable identification of \textit{Acacia} species.

High quality DNA extracted from the fresh leaves of selected \textit{Acacia} species was used as template and expected read lengths were amplified using gene specific primers. Successful DNA isolation discussed in this study, has been correlated with earlier studies Kumar et al. (2018) and Hu et al. (2009). In addition, our results also support earlier studies that observed no variation in sequence length for \textit{rbcL}, with successful PCR amplification and sequencing results (Kress et al., 2005; Roy et al., 2010) sometimes even with 100% success rate (Maia et al., 2012).

In contrast to \textit{rbcL}, the success rate of amplification and sequencing of \textit{matK} fragment was 57.24 ± 4.42% and 50.82 ± 4.36%, respectively. A similar study by Kress et al. (2009) reported that \textit{matK} had the lowest overall rate of recovery (69%) in amplification and sequencing. Similarly, the success rate of amplification in Kress and Erickson (2007) was only 39.3% for 96 species and 46 genera. Sass et al. (2007) used \textit{matK} to amplify \textit{Cycus}, however, with a lower success rate of only 24%. The possible reason behind this low amplification with \textit{matK} was that different features of the \textit{matK} gene are difficult to amplify and sequencing with universal and shows poor results (Chase et al., 2007; Hollingsworth, 2008). Previously several researches have suggested that success rate of amplification with \textit{matK} was highly variable, ranging from 40% to 97% (Zhang et al., 2012; Kress and Erickson, 2007). Therefore, universality of primers is recognized as an important criterion for evaluating the effectiveness of DNA barcodes (CBOL 2009).

The average AT nucleotide composition for \textit{rbcL} of all selected species of \textit{Acacia} was found higher to be 55.4% than GC content, 44.54%. Similarly, in \textit{matK}, average AT contents (59.46%) were higher as compared to the GC contents (40.44%). Similar findings have been reported by Dhakad et al. (2017) who found higher AT content (55.7%) than GC content (44.2%) in \textit{Acacia} sp. using \textit{rbcL}. In addition, Wijayasirawarden et al. (2017) also reported that nucleotide composition of AT was higher (68.98%) than GC (31.01%) contents for \textit{matK}. The possible explanation of higher AT contents than GC contents in both markers could be due to high

Table 6 (continued)

| Marker | Acacia albida | Acacia ampluscepts | Acacia coriacea | Acacia catechu | Acacia tortilis |
|--------|--------------|-------------------|----------------|---------------|----------------|
| A | G | A | A | A |
| G | A | G | G | A |
| T | C | T | T | T |
| T | A | T | T | T |
| A | T | A | A | A |
| G | T | G | G | G |
| A | T | A | A | A |
variability in nucleotide composition and higher nucleotide substitution rate in these genes.

Evaluation of intraspecific and interspecific divergence among species is a useful approach for assessing potential of DNA barcodes (Newmaster et al., 2008; Yu et al., 2014; Puillandre et al., 2012). The interspecific distances were calculated with K2P model utilizing pairwise comparison to trace evolutionary relationship between species. The K2P genetic distance between species summarized in (Table 4) showed that means genetic distance was higher (0.704%) in *Acacia* species using *matK* marker as compared to (0.04%) with *rbcl*. In a similar study, Nithaniyal et al. (2014). Nithaniyal et al. (2014) reported interspecific divergence range from 0.0 to 1.8% and 0.0 to 2.6% for *rbcl* and *matK* respectively. The poor interspecific divergence could be the result of frequent natural hybridization reported in *Acacia* species. Natural hybridization in *Acacia’s* is well documented and various hybrids with different ploidy levels have been identified in *Acacia nilotica* and *Acacia senegal* (Blakesley et al., 2002; Khatoon and Ali, 2006; Odee et al., 2015).

DNA barcode is a short DNA sequence based system specialized for one or more loci which can be used as a complementary unit for precise identification (Kress and Erickson, 2007). In our study, we propose to develop barcodes for proper identification of *Acacia* species; thereby a phylogenetic tree based approach was also used for validation of DNA barcodes obtained through SNPs analysis. Our results revealed that *rbcl* and *matK* generated strong barcodes for the proper identification of tested *Acacia* species. In phylogenetic analysis based on *rbcl* sequences *A. albida*, *A. coriacea* and *A. tortilis* were distantly related and clearly differentiated from each other as compared to *A. catechu* (Fig. 4). However, differentiation between *A. catechu* and *A. amплiceps* was poorly resolved thus required more markers for their identification. In case of *matK*, *A. coriacea* and *A. tortilis* were closely related with each other whereas *A. catechu*, *A. amплiceps* and *A. tortilis* were closely related with each other whereas *A. amплiceps* and *A. tortilis* were distantly related and clearly differentiated from each other as compared to *A. catechu* (Fig. 6).

**Fig. 4.** Phylogenetic relationship between different *Acacia* species differentiated based on *rbcl* genetic marker.

**Fig. 5.** Phylogenetic relationship between different *Acacia* species differentiated based on *matK* genetic marker.

**Fig. 6.** Phylogenetic relationship between different *Acacia* species differentiated based on both *matK* and *rbcl* genetic marker.
albida and A. ampliceps were clearly differentiated in a barcode tree (Fig. 5). Phylogenetic relationship between different Acacia species was also generated using both matK and rbcL genetic markers. In combined analysis using both markers, A. coriacea and A. tortilis were closely related as compared to the A. ampliceps, and A. albida and A. catechu were also grouped together thus closely related to each other. Finally, DNA sequences based QR codes were developed, which were scannable with smartphone applications, similar to barcode scanner in supermarkets. Researchers have developed similar barcodes for different species. For example, Bhagwat et al. (2015) reported the suitability of matK and rbcL markers for generating a barcode and differentiation of Dalbergia species. Similarly, Li et al. (2011) also generated the barcode for Aquilaria species.

Most of the genomes have been sequenced, so molecular approaches based on DNA sequences may help in efficient and reproducible identification of tree species. In the past, morphological characters were utilized for identification however morphological markers are not always reproducible and identifiable. Therefore, it is very difficult to identify the species on the basis of visible morphological characters that are almost indistinguishable from each other. Rapid developments in technology, bioinformatic tools, next generation sequencing and availability of genome sequences for most of species, have imparted more power for species identification using DNA sequence based markers. The rbcL has low resolution and high universality while, in contrast, matK has high resolution and low universality among the different species.
Therefore, using both matK and rbcL genetic regions, may help to discriminate Acacia species. 5.

Conclusions

The molecular assay performed through both selected DNA markers rbcL and matK provided reproducible results. Both DNA markers (rbcL and matK) were utilized effectively, to develop DNA barcodes for Acacia species. The molecular characterization using rbcL and matK markers, can be utilized to tag more SNPs based variations in different Acacia species.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Acknowledgement

This research was partially supported by Higher Education Commission, Islamabad, Pakistan under project NRPDU-6350 to Dr Aftab Ahmad. Authors also thankful the support of the Research Center for Advanced Materials Science (RCAMS) at King Khalid University, Abha Saudi Arabia through a grant RCAMS/KKU/08-20. We thank our colleagues from cotton Biotechnology Laboratory for their assistance in research.

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