Emergence of Plastidial Intergenic Spacers as Suitable DNA Barcodes for Arid Medicinal Plant Rhazya stricta

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Abstract

The desert plant Rhazya stricta has anticancer and antimicrobial properties, and is widely used in indigenous medicines of Saudi Arabia. However, the therapeutic benefits rely on an accurate identification of this species. The authenticity of R. stricta and other medicinal plants and herbs procured from local markets can be questionable due to a lack of clear phenotypic traits. DNA barcoding is an emerging technology for rapid and accurate species identification. In this study, six candidate chloroplastid barcodes were investigated for the authentication of R. stricta. We compared the DNA sequences from fifty locally collected and five market samples of R. stricta with database sequences of R. stricta and seven closely related species. We found that the coding regions matK, rbcL, rpoB, and rpoC1 were highly similar among the taxa. By contrast, the intergenic spacers psbK-psbI and atpF-atpH were variable loci distinct for the medicinal plant R. stricta. psbK-psbI clearly discriminated R. stricta samples as an efficient single locus marker, whereas a two-locus marker combination comprising psbK-psbI + atpF-atpH was also promising according to results from the Basic Local Alignment Search Tool and a maximum likelihood gene tree generated using PHyML. Two-dimensional DNA barcodes (i.e., QR codes) for the psbK-psbI and psbK-psbI + atpF-atpH regions were created for the validation of fresh or dried R. stricta samples.

Keywords

Rhazya Stricta, Medicinal Plant, DNA Barcoding, matK, rbcL, rpoB, rpoC1, atpF-atpH, psbK-psbI, Two-Dimensional DNA Barcode, QR Code
1. Introduction

*Rhazya stricta* of the Apocynaceae family is an important folkloric medicinal plant of Arabia [1]. The genus *Rhazya* includes only two species, *Rhazya stricta* and *Rhazya orientalis* (syn. *Amsonia orientalis*) [2] and both of these are unambiguously distributed, as the natural habitats of *R. stricta* are the coastlines and arid regions of Arabian Peninsula and the Indian subcontinent [3] while those of *R. orientalis* are northwest Turkey and northeast Greece [4]. But these two sister species exhibit similarity in possessing therapeutically significant secondary metabolites. About 100s of alkaloids [2] and flavonoids from *R. stricta* contain innumerable pharmacological properties [5] [6] [7]. *R. stricta* is considered as a potential chemopreventive [8] [9], antifungal [10], and antidiabetic [11] agent and also produces analgesic and sedative effects [7]. Likewise the metabolites originating from *R. orientalis* have various anti-cancer and anti-tumor properties [12]. This study targeted the locally available samples of *R. stricta* species hence its sole sister species *R. orientalis* was not included in this study.

Like other medicinal plants, *R. stricta* is procured either from the vast deserts or from herbal markets, where it goes by the vernacular name “Harmal”, and is found in a dried or powdered form. Although authenticity is among the most significant aspects related to pharmacological products [13] the absence of distinct phenotypic traits for *R. stricta* impedes the identification of this plant. Unfortunately, contamination and adulteration of medicinal plant products are quite common in the herbal markets, which have even caused severe diseases and death in some cases [14]. It is essential to carry out molecular level identification of medicinal herbs before they are used as a therapeutic agent [15]. Thus, there is a dire need for accurately identifying *R. stricta* at the molecular level in addition to the traditional taxonomical examination to protect consumers.

Within the past two decades, we have witnessed advances in molecular level techniques for accurately discerning specimens, even in the absence of diagnostic morphological characteristics. These techniques have proved unambiguous for the authentication of medicinal plants [16]. One such molecular tool is DNA barcoding, which can offer quick authentication compared with traditional taxonomy [17]. This technology identifies unknown species by using standardized DNA segments as universal product codes. It targets highly conserved DNA sequences in which minor nucleotide polymorphisms have evolved [18]. These nucleotide variations aid in the creation of unique DNA barcode markers, which can be used to validate a sample from a given species [19].

Whereas the 5' end of the mitochondrial gene for cytochrome c oxidase 1 is used for the standard barcode for animals [20], this region is unsuitable for use in identifying plants owing to a low substitution rate [21]. A variety of complex evolutionary processes in plants (e.g., hybridization and polyploidy) complicate the distinctions for defining species boundaries [22]. Consequently, after years of tremendous efforts, not one single region has worked as an identifier for all of the plant species tested [23] [24], and it is unlikely that a single universal plant DNA barcoding marker exists [25].
Various regions of plant DNA have been studied for barcoding [26] [27] [28] [29], including the coding and noncoding regions of plastid genomes (rbcL, psbA-trnH, trnL-trnF, and matK) and nuclear regions (5S, 16S, 18S, and ITS) [30]. These efforts have led to six plastid DNA regions as leading candidates for a suitable plant barcode, namely, matK, rbcL, rpoB, rpoC1, and psbK-psbI and atpF-atpH intergenic spacers [29]. Indeed, some of the loci from chloroplastid DNA, including matK, rbcL, atpF-atpH, psbK-psbI, rpoB, and rpoC1, have shown promising results for identifying medicinal plants. For example, the matK region, which was proposed as a standard barcode for flowering plants after studying 1,084 plant species [28], produced positive results for the medicinal plant Rauvolfioideae (a subfamily of Apocynaceae) [31]. rbcL was recommended for species discrimination [29], owing to the successes for its amplification and sequencing [27]. Moreover, taxonomic confusions among Cupressaceae, Cornaceae, Ericaceae, and Geraniaceae specimens were resolved via rbcL sequences [32]. Sequences for rpoB and rpoC1 were informative in identifying Ochradenus arabicus [19], and Chase et al. [26] suggested they be used in combination with matK for DNA barcoding.

Promising results have also been obtained with the noncoding intergenic spacers, which have evolved comparatively rapidly and exhibit sequence divergence and high rates of insertion/deletion [27]. The intergenic spacer atpF-atpH could be used for validating medicinal plant material [33] and for distinguishing all three species in the genera Landoltia and Spirodela [34]. The intergenic spacer psbK-psbI shows potential for barcoding of the flora of Kruger National Park, South Africa [28] and its use has been endorsed over that of matK and other loci by the CBOL Plant Working Group, 2009 [29] due to its capability for discriminating species.

Nevertheless, a single locus DNA barcode has been proposed to lack sufficient variation for discriminating closely related taxa [35]. Hence, to enhance the identification capabilities of barcode markers, we investigated a multi-locus DNA barcode [27] [28]. We also determined the abilities of the six markers proposed by the CBOL Plant Working Group for discriminating R. stricta at the molecular level.

2. Materials and Methods

2.1. Ethics Testimony

The R. stricta plant is indigenous to the region and was found growing in the surrounding deserts and on roadsides; hence, no permission was required for sample collection.

2.2. Plant Material

Ten fresh leaf samples of R. stricta were collected from each of five different locations in the Western province of Saudi Arabia (at 21°21.042', 39°33.054'; 21°24.849', 39°44.044'; 21°29.622', 39°35.823'; 21°26.817', 39°31.109'; and 21°26.423', 39°31.853') to exclude the presence of any subspecies. We sampled
plants during the winter season, when they are actively growing. Plants were identified at King Abdul Aziz University on the basis of morphological markers. Samples were immersed in liquid nitrogen and stored at -80˚C until DNA extraction. Five raw dried and powdered samples of *R. stricta*, were obtained from the local herbal market to check the efficiency of the proposed DNA barcode.

### 2.3. DNA Extraction, Amplification, and Sequencing

*R. stricta* is a poisonous medicinal plant with high levels of alkaloids and flavonoids that can make DNA extraction challenging. To overcome this, genomic DNA was extracted from young leaf tissues [36] using a DNeasy plant mini kit (Qiagen, Germany) according to the manufacturer’s instructions with minor modifications (elution buffer AE was heated to 65˚C and elution times were increased by 30 min). Following the same modification DNA was extracted from the *R. stricta* market samples. All of the extracts were stored at −20˚C until their use as templates for PCR. The concentrations and purities of the extracted DNA were quantified using a Nano Drop 2000 spectrophotometer (Thermo Scientific, Wilmington, DE, USA). The extracted genomic DNA was also examined by electrophoresis on a 1% agarose gel in 1× TBE buffer and stained with ethidium bromide (0.5 mg⋅mL⁻¹).

Target loci sequences were amplified in a total volume of 30 μL, consisting of 15 μL of GoTaq green master mix (for final concentrations of 200 μM for each deoxynucleoside triphosphate and 1.5 mM MgCl₂), 1 μM each of forward and reverse primers, 25 - 250 ng of sample genomic DNA, and deionized distilled water. The primers (Bioneer Corp., South Korea) and parameters used for amplifying chloroplast loci are shown in Table 1. As initial amplifications failed for some loci (e.g., *matK*), which has been reported for samples in the order Gentianales [37], the thermocycling conditions were optimized using touchdown PCR (Table 1). This resulted in successful amplification of target regions.

The amplicons were analyzed on a 1.5% agarose gel as described above for genomic DNA analysis using a 100-bp DNA ladder (Promega Corp., USA) as a molecular marker. Gel images were analyzed on a gel documentation system (Biospectrum 410; UVP). PCR products were sent to Macrogen, Inc. (Seoul, South Korea) for bidirectional sequencing using the same primers to resolve ambiguities [38].

### 2.4. Data Analysis

Sequence chromatograms were viewed via Flinch TV 1.4.0 (Geospiza, Inc., Seattle, WA, USA) to analyze base calls and quality values. Further analysis and alignments were carried out using Geneious 9.1 software [39]. Sequence similarities were identified using the BLAST algorithm (http://blast.ncbi.nlm.nih.gov/blast.cgi) to judge the identification capability of the six-barcode regions. High Bit scores, Grades (a weighted score for the hit comprising the E-value, pairwise identity, and coverage), and a cutoff E-value of 1e20 were taken into account. Sequences of the barcode regions for *R. stricta* and
for seven highly similar species (related genera from the Apocynaceae family) that performed the best were downloaded from the GenBank database to assess their discriminatory efficiency (Table 2). We studied species from different genera, as no data from the chloroplast regions of other species in the genus *Rhazya* (other than *R. stricta*) appeared in our GenBank database search. These sequences were edited and aligned in Geneious Pro 9.1 [39], using the multiple alignments tool MUSCLE plug-in [40]. For alignment, sequence orders from the Blast search were preserved. GC contents and the percentages of identical nucleotide sites were analyzed with the same software.

We used the tree-base method of identification with phylogenetic distance (via gene tree) to study the divergences of different barcode markers. Here, a query was allocated to the species with whom it clustered. In this study, phylogenetic distances were analyzed using PHyML 3.0 software [41]. On the basis of the HKY85 model of nucleotide substitution, this software generated a maximum likelihood gene tree with a 1000-replicate bootstrap test to authenticate *R. stricta*. After these analyses, all the sequences were deposited into the GenBank nucleotide database (Table 3).

In addition to the single locus barcodes, a double locus combination was investigated by concatenating the candidate barcode regions using Geneious Pro 9.1 software. The mean intra-and interspecific genetic distances were calculated using MEGA 6.0 [42] with the Kimura 2-parameter model. The proposed DNA barcodes were validated on five dried powered market samples of *R. stricta*.

### 2.5. Two-Dimensional DNA Barcoding

The nucleotide sequences of the candidate barcode markers, which validated *R. stricta* species efficiently, were converted to two-dimensional DNA barcode images using an open-source PHP QR coding method [43].

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**Table 1.** Primers and amplification conditions for chloroplastid regions.

<table>
<thead>
<tr>
<th>Primer</th>
<th>Nucleotide sequence (5′ → 3′)</th>
<th>Thermocycling conditions</th>
</tr>
</thead>
<tbody>
<tr>
<td>matK_XF1</td>
<td>TAATTTCGATCAATTCTATTCA</td>
<td>94°C for 4 min, 40 cycles of 94°C for 50 s, 52°C for 1 min, and 72°C for 1 min, 72°C for 10 min</td>
</tr>
<tr>
<td>matK_MALP_R1</td>
<td>ACAAGAAAGTGAAGTAT</td>
<td></td>
</tr>
<tr>
<td>rbcLa F</td>
<td>ATGTCCACACAAACAGACTAAAGC</td>
<td>94°C for 1 min, 30 cycles of 94°C for 45 s, 51°C for 45 s, and 72°C for 5 s, 72°C for 5 min</td>
</tr>
<tr>
<td>rbcLa R</td>
<td>CTTCCTGCTACCATGACTAATCTGCC</td>
<td></td>
</tr>
<tr>
<td>rpoB F</td>
<td>ATGCAACGTCAAGCAGTTC</td>
<td>94°C for 4 min, 40 cycles of 94°C for 30 s, 53°C for 40 s, and 72°C for 40 s, 77°C for 7 min</td>
</tr>
<tr>
<td>rpoB R</td>
<td>GATCCCCAGCATCAAATCTCC</td>
<td></td>
</tr>
<tr>
<td>rpoC1-1F</td>
<td>GTGGATACACTTCTCTGATAATGG</td>
<td>94°C for 4 min, 40 cycles of 94°C for 30 s, 53°C for 40 s, and 72°C for 40 s, 72°C for 7 min</td>
</tr>
<tr>
<td>rpoC1-3R</td>
<td>TGAGAAACATAAGTAAAAGGCC</td>
<td></td>
</tr>
<tr>
<td>atpF-atpH F</td>
<td>AACTCGCAGACACTCCCTTT</td>
<td>94°C for 3 min, 35 cycles of 94°C for 30 s, 51°C for 40 s, and 72°C for 40 s, 72°C for 10 min</td>
</tr>
<tr>
<td>atpF-atpH R</td>
<td>GGRGTTGGTCAAGGTACTGC</td>
<td></td>
</tr>
<tr>
<td>psbK-psbI F</td>
<td>TTAGGCATTTGGTGACCAAG</td>
<td>95°C for 2 min 30 s, 35 cycles of</td>
</tr>
<tr>
<td>psbK-psbI R</td>
<td>AAAGTTTGAGATAGAAGCAT</td>
<td>95°C for 30 s, 51°C for 1 min, and 72°C for 1 min, 72°C for 10 min</td>
</tr>
</tbody>
</table>

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Table 2. Sequence sources for \textit{atpF-atpH} and \textit{psbK-psbI} markers from GenBank.

<table>
<thead>
<tr>
<th>No.</th>
<th>Species</th>
<th>Accession no.</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>\textit{Rhazya stricta}</td>
<td>KJ485849</td>
</tr>
<tr>
<td>2.</td>
<td>\textit{Catharanthus roseus}</td>
<td>KC561139</td>
</tr>
<tr>
<td>3.</td>
<td>\textit{Pentalinon luteum}</td>
<td>KJ953909</td>
</tr>
<tr>
<td>4.</td>
<td>\textit{Nerium oleander}</td>
<td>KJ953907</td>
</tr>
<tr>
<td>5.</td>
<td>\textit{Secamone afzelii}</td>
<td>KF539845</td>
</tr>
<tr>
<td>6.</td>
<td>\textit{Echites umbellatus}</td>
<td>KJ953904</td>
</tr>
<tr>
<td>7.</td>
<td>\textit{Wrightia natalensis}</td>
<td>KJ953913</td>
</tr>
<tr>
<td>8.</td>
<td>\textit{Marsdenia astephanoides}</td>
<td>KF539849</td>
</tr>
</tbody>
</table>

Table 3. Accession numbers for \textit{Rhazya stricta} sequences submitted to GenBank.

<table>
<thead>
<tr>
<th>Target region</th>
<th>Accession no.</th>
</tr>
</thead>
<tbody>
<tr>
<td>matK</td>
<td>KX602160</td>
</tr>
<tr>
<td>rbcL</td>
<td>KX602163</td>
</tr>
<tr>
<td>rpoB</td>
<td>KX602164</td>
</tr>
<tr>
<td>rpoC1</td>
<td>KX602165</td>
</tr>
<tr>
<td>atpF-atpH</td>
<td>KX602155</td>
</tr>
<tr>
<td>psbK-psbI</td>
<td>KX602162</td>
</tr>
</tbody>
</table>

3. Results

3.1. Success of DNA Isolation and Amplification

Although the extraction of DNA was challenging from some samples, DNA was successfully extracted using the modifications described in the Methods. All six of the chloroplast barcode loci were successfully amplified after optimizing the thermocycling conditions. The properties of these six loci are given in “Table 4”.

3.2. BLAST Analysis and Multiple Alignments

The BLAST analysis of the six barcode chloroplast loci (\textit{matK}, \textit{rbcL}, \textit{rpoB}, \textit{rpoC1}, \textit{atpF-atpH}, and \textit{psbK-psbI}) showed 100% identities and query covers with the \textit{R. stricta} sequence (GenBank no. KJ485849). In addition, the sequences from the loci of \textit{R. stricta}, viz., \textit{matK}, \textit{rbcL}, \textit{rpoB}, and \textit{rpoC1}, showed maximum (95%, 98%, 97%, and 98%, respectively) similarities with the other taxa examined (of the Apocynaceae family). However, \textit{atpF-atpH} and \textit{psbK-psbI} sequences showed ≤90.6% and ≤87.5% similarities, respectively, to the other taxa sequences (given in Table 5). They were found to be distinct for \textit{R. stricta}.

There were no variations in the intraspecific alignments of sequencing products for the six loci (from fifty \textit{R. stricta} samples). The mean intraspecific distances (calculated by MEGA 6) were zero, verifying that all of the fifty \textit{R. stricta} samples belong to the same species with no genotypic variation.

The interspecific divergences (investigated after carrying out a multiple
alignment between *R. stricta* and the seven species from Apocynaceae) for *matK*, *rbcL*, *rpoB*, and *rpoC1* regions were low due to the high percentages of identical sites (*Table 4*). Nevertheless, interspecific divergences were high for *atpF-atpH* and *psbK-psbI* loci due to variations in *R. stricta* sample sequences (*Table 4*). The identical sites percentages (calculated using Geneious Pro 9.1) for these two loci were much lower than for the coding loci (*Table 4*). “Figure 1” highlights the variable informative sites in the *psbK-psbI* locus from *R. stricta* sample sequences. The mean interspecific distance was highest for *psbK-psbI* (7.80%) compared with those of the other loci.

### 3.3. Maximum Likelihood Tree Identification

The maximum likelihood tree [41] for the *psbK-psbI* region clustered sample sequences of *R. stricta* (query sequences) with the GenBank *R. stricta* sequences into a single independent clade that was highly supported (**Figure 2(a)**). The case was similar for the double locus (*psbK-psbI + atpF-atpH*) sample sequences...
Figure 1. Multiple-sequence alignments highlighting the variable informative sites in the *psbK-psbI* locus from *R. stricta* sample sequences.

Figure 2. Maximum likelihood trees of *psbK-psbI* (a) and *psbK-psbI + atpF-atpH* (b) sequences for *R. stricta*. The branch labels display maximum likelihood bootstrap values.
Hence, the similarity-based method and tree-based identification indicate that *psbK-psbI* is an appropriate DNA barcode region for identifying *R. stricta*. The *psbK-psbI* intergenic region aided by *atpF-atpH* also displayed promising results, supporting its use as a double locus barcode for *R. stricta*. These two proposed DNA barcodes (*psbK-psbI* and *psbK-psbI + atpF-atpH*) clearly identified dried powdered *R. stricta* samples acquired from the market.

### 3.4. Generation of Two-Dimensional Barcodes

The purpose of DNA barcoding is to accurately and quickly identify medicinal plants and their products at the molecular level to benefit the consumer. With this aim, the nucleotide sequences of the DNA barcode markers (*psbK-psbI* and *psbK-psbI + atpF-atpH*) were considered analogous to the products’ barcodes in a supermarket and were converted to two-dimensional barcodes (i.e., QR codes) ([Figure 3](#)). The colors in the bars represent nucleotides of the marker, and the numbers at the ends denote the lengths of the markers. After scanning the two-dimensional barcode on a mobile terminal, a message can be sent to a DNA barcode database, where the authenticity of a plant or specimen, such as *R. stricta*, can be checked. This will aid the ability of consumers with no prior knowledge of DNA barcoding to accurately identify medicinal plants and their products.

### 4. Discussion

Approximately 80% of the world’s population are inclined to use herbal medicines for their primary care [44]. Similarly, the Saudi population exhibits a keen interest in the medicinal species of the local flora [45]. For example, the numerous pharmacological properties of *R. stricta* make this a significant medicinal plant in this part of the world [46]. Although it is easily available from its local natural habitat, identifying this plant in herbal markets is difficult due to the lack
of distinct morphological traits and the variability in storage conditions and product ages. In an effort to safeguard consumers’ health, we investigated a method of DNA barcoding for validating *R. stricta* at the molecular level, irrespective of its physical state.

DNA barcoding is considered the renaissance of taxonomy [47]. This molecular tool emerged rapidly over the past two decades for species discrimination [25], and it has outperformed other diagnostic tools for identifying and authenticating species [48]. Numerous studies have not only affirmed its competency for species identification, but also highlighted its strengths in defining species boundaries and flagging new species [20] [49]. Owing to its ease and swiftness, this tool has been used extensively for distinguishing medicinal plants [50] [51] [52]. It is predicted to play a significant role in identifying medicinal plant products in the future [53].

It was proposed that an ideal DNA barcode (i) can be easily amplified and sequenced, (ii) does not exceed 1 kb in size, and (iii) has higher interspecific than intraspecific variation [54]. We selected six barcoding loci to examine with these criteria in mind. The *rbcL*, *rpoB*, *rpoC1*, *psbK-psbI*, and *atpF-atpH* loci were easily amplified and sequenced. The amplification of *matK* was challenging initially, which has been found in medicinal *Uncaria* species [55] and temperate flora comprising 436 species in 269 genera of land plants [56]. By contrast, the *psbK-psbI* region amplified easily, similar to that for *Panax* species (Ginsengs) [57]. The successful amplification of intergenic spacers (*psbK-psbI* and *atpF-atpH*) results from the ability to use universal primers for the highly conserved coding sequences on either side of the locus [58].

In this study, the sequences of *R. stricta* for *matK*, *rbcL*, *rpoC1*, and *rpoB* were similar to those for the other taxa. Accordingly, *matK* did not aid in species identification in other studies, suggesting that it may be nonfunctional in some taxa [29] [17]. The *rpoC1*, *rpoB* [59] [60], and *rbcL* [61] loci similarly have shown very low species discrimination in other investigations. The noncoding intergenic spacer regions, *psbK-psbI* and *atpF-atpH*, are promising barcode markers for *R. stricta* due to their faster divergence than genes in otherwise slow-evolving plant species [62] [27]. Indeed, the *psbK-psbI* region exhibited efficient barcode recovery and species discrimination in a number of studies [29] [63]. Zuo *et al.* [58] endorsed *psbK-psbI* over coding regions, as it is a more informative and powerful marker for species identification. This locus discriminates the Kruger National Park flora [28] and taxa of *Orchidaceae* in Korea [64]. Similarly, the *atpF-atpH* intergenic spacer has been endorsed as a supplementary locus [65] as it was found to successfully authenticate medicinal plant materials in a study involving 17 barcode regions [33]. Another option that was proposed was to combine *psbK-psbI* with other loci, such as *matK* and *atpF-atpH* [66] [28] [65].

DNA barcoding takes advantage of polymorphisms that lead to gaps in sequences. Polymorphisms in *psbK-psbI* and *atpF-atpH* regions were observed as gaps upon amplicon alignment that were found to be distinct to *R. stricta* and
might serve for barcoding this species. In addition, the psbK-psbI and concatenated psbK-psbI + atpF-atpH regions displayed 64.1% and 72.4% similarities (on the basis of nucleotide sites), respectively, to the other species sequences on alignment. Hence, these loci are considerably unique with regard to R. stricta. Indeed, a 100% monophyletic differentiation of the R. stricta species was found for the single locus (psbK-psbI) and double locus combination (psbK-psbI + atpF-atpH) as shown by the maximum likelihood trees. Moreover, the psbK-psbI and psbK-psbI + atpF-atpH loci successfully identified market samples of R. stricta.

5. Conclusion

In this study, we found that matK, rbcL, rpoC1, and rpoB coding regions are not effective markers for R. stricta, as they were similar to the sequences of other species. By contrast, intergenic spacer regions psbK-psbI and atpF-atpH contained sequence polymorphisms that are unique to R. stricta. We propose the use of psbK-psbI as an optimal single locus barcode and psbK-psbI + atpF-atpH as a complementary double locus combination for creating DNA barcodes to identify the medicinal plant R. stricta. We generated two-dimensional barcodes (QR codes) for these loci to facilitate molecular authentication of R. stricta via an easy and inexpensive method for consumers. This method would also greatly benefit market supervision of other medicinal plants of Saudi Arabia and its neighboring countries. It can be a preliminary step towards generating a database of barcode markers for economically significant flora of this arid region.

Acknowledgements

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