RAPD Based Assessment of Genetic Diversity of *Adhatoda vasica* Leaves from Different Sub-Continents of India

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**ABSTRACT**

The present work assessed the genetic divergence amongst the accessions of vasaka collected from different sub-climatic zones of India by RAPD (Randomly Amplified Polymorphic DNA) using twenty random decamer primers (OPA 1-OPA 20) as the plant was found to enjoy its therapeutic efficacy in Ayurvedic and traditional system of medicines. The dendrogram constructed for cluster analysis using an un-weighted pair group method with arithmetic means (UPGMA) grouped the accessions into 2 major clusters based on win boot. Out of the twenty random primers used for studying genetic divergence sixteen primers were found to be polymorphic. Out of 20 primers 3 were found to be 100% polymorphic generating a total of 313 amplification products with an average of 19.5 products per polymorphic primer. Genetic relationships among accessions were evaluated by generating a similarity matrix based on Jaccard’s co-efficient ranging from 0.60 to 0.91. Results showed that both environmental and genetic factors were effective in observing variations. The degree of genetic variations detected among the accessions of vasaka suggested that RAPD approach seemed to be best suited for assessing with high accuracy the genetic relationships among distinct *A. vasica* accessions.

**Key words:** RAPD, Genetic diversity, *Adhatoda vasica*, variation, Polymorphism, Primer.

**INTRODUCTION**

*Adhatoda vasica* (Acanthaceae) commonly known as vasaka, Malabar nut tree in English and arusa or adula in local Hindi language is a primary herb of Ayurvedic system of medicines and has been used in indigenous in India for over last 2000 years. *A. vasica* is a medium sized shrub found up to an altitude of 1300 m distributed throughout tropical and temperate regions.⁵

Leaves of *A. vasica* are generally administered in yogic practices to clear the respiratory passages as well as in the preparations including cough syrups.⁶ They were reported to contain the quinazoline alkaloids vasicine, vasicinone and deoxyvasicine.⁴ Some of the chemical compounds found in the leaves and roots of this plant includes essential oils, fats, resins, sugars, gums, amino acids, proteins and vitamin C.⁵ Since a vast majority of medicinal plants have been recklessly exploited to a greater extent so it makes sense to rationalize the use of some important medicinal plant through screening and validation of germplasm. In the present research work it will be prudent to evaluate the genetic component in *A. vasica* for conservation and management of genetic diversity as the species has been included in the list of endangered and threatened species of India over a decade.⁶ The Indian sub-climatic zones plant specialist group has also identified *A. vasica* for the detailed study and protection along with other species like *Abrus precatorius* Linn, *Costus speciosus*, *Centella asiatica* (Linn.), *Gloriosa superb* Linn, *Rauwolfia serpentina* Benth. ex Kurz., *Saraca indica* de Wilde, *Streblus asper* Lour., *Tribulus terrestris* L. and *Withania*
Moreover, frequent adulteration of vasaka leaves with *Alianthus beddomei* (Clarke), *A. altissimia* (Mill) Swingle, *A. glandulosa* (Desf) and *A. excelsa* Roxb. Leaves has also made the condition even worsened as did the Traditional Chinese Medicines (TCM) contaminated with *Aristolochia* causing an epidemic of sub-acute intestinal nephropathy and necessitating kidney transplantation of the users in Belgium therefore further necessitating the need for the proper identification and conservation of the particular plant species *A. vasica*.

To combat the mentioned difficulties the present research was aimed to establish the level of genetic relatedness between the different accessions of *A. vasica* by RAPD so as to identify and maintain the efficacy of the plant. The technique chosen is a PCR based simple, reliable and cost effective technique and acting as a powerful tool for the analysis of plant genome in comparison to other molecular techniques i.e. AFLP (Amplified Fragment Length Polymorphism) RFLP (Restricted Fragment Length Polymorphism), SSR (Simple Sequence Repeat) SCAR (Sequence Characterized Amplification Regions) VNTRs (Variable Number of Tandem Repeats) and non-PCR based Restriction hybridization Techniques employing the use of restriction end nucleases and Hybridization methods as it is simple to operate and does not involve radioactive labelling. The RAPD technique requires a very low amount of genetic material and provides more of the detailed genetic information due to either increased variability of loci or the greater no of available loci. In addition it has been used to identify polymorphism for elicit information on divergence, variation, diversity analysis, phylogeny, quantitative traits, marker assisted selection etc.

### MATERIALS AND METHODS

#### Plant material

In the present study genetic diversity was analysed using RAPD primers among the different accessions of vasaka collected from different geographical locations of India. A definite criterion was adopted for the collection of the samples as the collection of young leaves was carried out in the month of March under flowering conditions from various sub-climatic zones at different altitudes from various geographical locations and transferred in a laboratory in an ice-box and stored at -20°C. The collection sites included viz. New Delhi (1) Hisar (2) Kurukshetra (3) Nahan (4) Chandigarh (5) Banaras (6) Solan (7) Dehradun (8) Patiala (9) Dalhousie (10). The study was conducted at Bioactive Natural Product Laboratory, Jamia Hamdard, New Delhi, located at 77° (longitude) and 28°8° (latitude). The plants collected were identified by Dr. Altaf ahmad, Taxonomist in Department of Botany and deposited the voucher specimens in Department of Pharmacognosy and Phytochemistry, Jamia Hamdard, New Delhi, India.

Ten cultivars of *A. vasica* were utilized to carry out the present study. The accessions numbers and codes given to these cultivars were mentioned in Table 1.

#### Reagents and chemicals

The chemicals and reagents used in the isolation of DNA were

- CTAB extraction buffer [2% (w/v) CTAB; 20 mM EDTA, pH 8.0; 100 mM Tris-HCl, pH 8.0; 1.4 M NaCl];
- CTAB/NaCl solution [10% (w/v) CTAB; 0.7 mM NaCl];
- CTAB precipitation solution [1% (w/v) CTAB; 50 mM Tris-HCl, pH 8.0; 10 mM EDTA];
- High salt TE buffer [10 mM Tris-HCl, pH-8.0; 0.1 M EDTA, pH 8.0; 1.0 M NaCl];
- TE buffer [10 mM tris-HCl, pH-8.0; 0.1 M EDTA, pH 8.0; 1.0 M NaCl];
- Tris-EDTA buffer [10 mM Tris-HCl, pH-8.0; 0.1 M EDTA, pH 8.0; 1.0 M NaCl];
- RNase and DNase free water.

### Table 1: *A. vasica* accessions from different sub-climatic zones sampled for RAPD

<table>
<thead>
<tr>
<th>Acc. Code</th>
<th>Place of collection</th>
<th>Latitude (N°)</th>
<th>Longitude (E°)</th>
<th>Altitude (m)</th>
</tr>
</thead>
<tbody>
<tr>
<td>AVND</td>
<td>New Delhi</td>
<td>28°63°</td>
<td>77°22°</td>
<td>305</td>
</tr>
<tr>
<td>AVHR</td>
<td>Hisar</td>
<td>29°9°</td>
<td>75°43°</td>
<td>221</td>
</tr>
<tr>
<td>AVKU</td>
<td>Kurukshetra</td>
<td>29°58°</td>
<td>76°53°</td>
<td>260</td>
</tr>
<tr>
<td>AVNH</td>
<td>Nahan</td>
<td>30°33°</td>
<td>77°21°</td>
<td>932</td>
</tr>
<tr>
<td>AVCH</td>
<td>Chandigarh</td>
<td>30°43°</td>
<td>76°47°</td>
<td>321</td>
</tr>
<tr>
<td>AVBS</td>
<td>Banaras</td>
<td>25°22°</td>
<td>83°00°</td>
<td>81</td>
</tr>
<tr>
<td>AVSN</td>
<td>Solan</td>
<td>30°90°</td>
<td>77°09°</td>
<td>1445</td>
</tr>
<tr>
<td>AVDN</td>
<td>Dehradun</td>
<td>30°19°</td>
<td>78°04°</td>
<td>1880</td>
</tr>
<tr>
<td>AVPL</td>
<td>Patiala</td>
<td>30°20°</td>
<td>76°24°</td>
<td>252</td>
</tr>
<tr>
<td>AVDL</td>
<td>Dalhousie</td>
<td>32°38°</td>
<td>75°58°</td>
<td>2040</td>
</tr>
</tbody>
</table>
DNA extraction (Isolation of DNA by CTAB method)

Five hundred micro litres of CTAB extraction buffer was heated to 65°C followed by the addition of 2-mercapto-ethanol (10 µl) to this mixture just before the extraction process. 0.5 g of fresh plant material and 100 mg of dried sample were frozen in liquid nitrogen (-196°C) in a sterile mortar and ground to a fine powder. The ground frozen tissue was incubated at 65°C for 1 h in a water bath and mixed uniformly by gentle inversion. Chloroform: Isoamyl alcohol (24:1,v/v) was added in equal volume to the homogenate and mixed gently by inversion, followed by centrifugation at 10,000 rpm (7500 × g) for 15 minutes. The upper phase was pipette into a sterile eppendorff tube. This step was repeated twice with addition of 10% CTAB in second repeated step. One volume of CTAB precipitation solution heated at 65°C was added to the recovered supernatant. A precipitate was formed at this stage and mixture got centrifuged at 2700 rpm (500 × g) for 15 min. DNA sample and DNA loading dye were mixed in 5:1 ratio for each well and loaded with the help of micropipette. Electrophoresis was conducted at 90 volts for 1 h in 0.5 X TAE buffer. The µgel was then stained with ethidium bromide solution (0.5 µg/ml). After de-staining in distilled water, the gels were viewed and stored in gel documentation system (alpha imager EC).

DNA quantification and agarose gel electrophoresis

The quantification of genomic DNA was achieved using a spectrophotometer (UV-Visible spectrophotometer, Shimadzu, Japan). The yield was determined by measuring the absorbance at 260, 280 and 320 nm. The level of DNA purity was determined by the ratio of absorbance obtained at 260/280 nm.

Table 2: Optimization of amplification reaction mixtures (15 µl) containing PCR ingredients

<table>
<thead>
<tr>
<th>Parameters/Reagents</th>
<th>Optimized concentrations</th>
<th>Tested range (U)</th>
<th>Volume used (µL)</th>
<th>Remarks</th>
</tr>
</thead>
<tbody>
<tr>
<td>PCR buffer with MgCl₂ (mM)</td>
<td>15 mM</td>
<td>1-5</td>
<td>2.5</td>
<td>Deviation increased non-specificity and affected yields of products</td>
</tr>
<tr>
<td>Deoxy-nucleotide triphosphate (mM)</td>
<td>2.5 mM</td>
<td>0.2-1.4</td>
<td>1.5</td>
<td>Increased concentration reduces free Mg²⁺ ion which interferes with enzyme</td>
</tr>
<tr>
<td>Primer concentration (10 pmol/µL)</td>
<td>10 pmol</td>
<td>0.1-1.5</td>
<td>1.0</td>
<td>Lower concentration leads to absence of amplification whereas higher concentration leads to dimer formation</td>
</tr>
<tr>
<td>Milli Q water</td>
<td>-</td>
<td>-</td>
<td>8.0</td>
<td>Absence of amplification at lower concentration and presence of smear at higher concentration</td>
</tr>
<tr>
<td>Extracted DNA (ng)</td>
<td>50 ng</td>
<td>5-200</td>
<td>1.0</td>
<td>Lower concentration led to improper amplification whereas increased concentration showed decreased specificity</td>
</tr>
<tr>
<td>Taq DNA polymerase (units/µL)</td>
<td>one unit/µL</td>
<td>0.1-1.0</td>
<td>1.0</td>
<td></td>
</tr>
</tbody>
</table>

RAPD-PCR analysis

A total of 20 random decamer primers (custom synthesized by Bangalore Genei Pvt. Ltd., GCs content > 50%) were used for RAPD analysis. DNA amplification reactions were performed in 15 µl reaction volumes (2.5 µl PCR buffer with MgCl₂, 1.5 µl dNTPs, 1.0 µl primer, 1.0 µl Taq DNA polymerase, 1.0 µl DNA template and 8.0 µl of milli Q water) (Table 2). Amplification reaction was carried out in a Bio-Rad Thermal cycler with a following thermal profile comprising of one cycle of 2 min at 94°C (initial denaturation) followed by 38 cycles of 30 sec at 92°C (denaturation), 60 sec at 32°C (primer annealing) and 90 sec at 72°C (final extension) which
were previously optimized. PCR products were kept at 4°C. Amplified PCR products were separated on 1.5% (w/v) agarose gel in 1X TAE buffer (pH 8.3) stained with Ethidium bromide in a final concentration of 10 µg/ml. Electrophoresis was performed at 90 volt for 2 h and then visualized the gel, photographed and analyzed. Gene ruler™ 1 kb DNA ladder (Bangalore Genei Pvt. Ltd.) was used as molecular size marker. The reproducibility of the amplification products was checked twice for each polymorphic primer. A control PCR tube containing all the components but no genomic DNA was run with each primer to check the contamination. The bands that did not show any fidelity were eliminated. Only reproducible fragments were scored while non-reproducible with the faintly stained fragments were discarded.

Data analysis

Evaluation of fragment patterns in RAPD analyses were analyzed using Nei genetic similarity index that determined the genetic distances between the genotypes of *A. vasica*. The co-efficient on x axis represented the similarity indices (DICE) of the different genotypes chosen in the current study. Reproducible bands were scored manually as 1 or 0 for the presence or absence of the bands across all the Adhatoda accessions for each primer. Only reproducible fragments were scored while non-reproducible with the faintly stained fragments were discarded.

### Results

#### Reproducibility of amplifications patterns

DNA amplifications with RAPD primer were repeated at least thrice to ensure reproducibility. The bands were considered reproducible and scorable only after observing and comparing them in three separate amplifications for each primer. Clear and intense bands were scored whereas faint bands against background smear got rejected and not considered for further analysis.

#### Scoring and data analysis

For each accession, each fragment/band that was amplified using RAPD primers was treated as a unit character. Unequivocally scorable and consistently reproducible amplified DNA fragments were transformed into binary characters matrices (1 for presence, 0 for absence). The commercial software package NTSYS-PC was used to develop similarity matrices. These data were then used to construct dendrogram for cluster analysis based on un-weighted pair group method with arithmetic mean (UPGMA) using computer programme WINDIST.

#### Genetic diversity between accessions

The similarity matrix of genetic distance was used to show the relationship amongst the ten accessions of *A. vasica*. Genetic variability studies in *A. vasica* collected from different locations of India had been carried out using RAPD markers. DNA was isolated by CTAB method. Measurement of absorbance at 260 nm and at 280 nm provided validation of the purity of nucleic acid in all the samples: A260/A280 ratios above 1.8 for DNA or 2.0 for RNA indicated pure samples; lower ratio values indicated the presence of protein or other contaminants. The ratios of optical density of 260/280 of all samples found in the range of 1.7-1.9.

### Table 3: Quantification of DNA in different accessions of *A. vasica*

<table>
<thead>
<tr>
<th>A. vasica samples</th>
<th>Optical density (λ)</th>
<th>Ratio 260/280 nm</th>
<th>DNA conc. (µg/ml)</th>
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<tbody>
<tr>
<td></td>
<td>260 nm</td>
<td>280 nm</td>
<td></td>
</tr>
<tr>
<td>AVND</td>
<td>0.04</td>
<td>0.021</td>
<td>1.90</td>
</tr>
<tr>
<td>AVHR</td>
<td>0.05</td>
<td>0.027</td>
<td>1.85</td>
</tr>
<tr>
<td>AVKU</td>
<td>0.07</td>
<td>0.037</td>
<td>1.89</td>
</tr>
<tr>
<td>AVNH</td>
<td>0.02</td>
<td>0.011</td>
<td>1.81</td>
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<tr>
<td>AVCH</td>
<td>0.09</td>
<td>0.05</td>
<td>1.80</td>
</tr>
<tr>
<td>AVBS</td>
<td>0.03</td>
<td>0.016</td>
<td>1.87</td>
</tr>
<tr>
<td>AVSN</td>
<td>0.09</td>
<td>0.048</td>
<td>1.87</td>
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<tr>
<td>AVDN</td>
<td>0.03</td>
<td>0.038</td>
<td>1.84</td>
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<td>AVPL</td>
<td>0.09</td>
<td>0.048</td>
<td>1.87</td>
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<tr>
<td>AVDL</td>
<td>0.07</td>
<td>0.036</td>
<td>1.94</td>
</tr>
</tbody>
</table>
Sayeed et al., RAPD analysis of *Adhatoda vasica* Linn in Indian sub-continents

Figure 1: Gel picture showing the RAPD amplification patterns generated by OPA-6
(M-Marker, Names of Samples: 1-New Delhi; 2-Hisar; 3-Kurukshetra; 4-Nahan; 5-Chandigarh; 6-Banaras; 7-Solan; 8-Dehradun; 9-Patiala; 10-Dalhousie)

Figure 2: Gel picture showing the RAPD amplification patterns generated by OPA-7
(M-Marker, Names of Samples: 1-New Delhi; 2-Hisar; 3-Kurukshetra; 4-Nahan; 5-Chandigarh; 6-Banaras; 7-Solan; 8-Dehradun; 9-Patiala; 10-Dalhousie)

Figure 3: Gel picture showing the RAPD amplification patterns generated by OPA-11
(M-Marker, Names of Samples: 1-New Delhi; 2-Hisar; 3-Kurukshetra; 4-Nahan; 5-Chandigarh; 6-Banaras; 7-Solan; 8-Dehradun; 9-Patiala; 10-Dalhousie)

Figure 4: Gel picture showing the RAPD amplification patterns generated by OPA-12
(M-Marker, Names of Samples: 1-New Delhi; 2-Hisar; 3-Kurukshetra; 4-Nahan; 5-Chandigarh; 6-Banaras; 7-Solan; 8-Dehradun; 9-Patiala; 10-Dalhousie)

Figure 5: Gel picture showing the RAPD amplification patterns generated by OPA-18
(M-Marker, Names of Samples: 1-New Delhi; 2-Hisar; 3-Kurukshetra; 4-Nahan; 5-Chandigarh; 6-Banaras; 7-Solan; 8-Dehradun; 9-Patiala; 10-Dalhousie)

Figure 6: UPGMA dendrogram based on Dice (Nei and Li, 1979) method from RAPD data
indicated high purity of DNA (Table 3). The coefficient on x axis represented the similarity indices (DICE) of the different accessions chosen in the current study. The genetic distance between the studied accessions was illustrated by the UPGMA dendrogram (Figure 6). Of all the samples analyzed accessions AVBS was marked by the highest distance from the remaining accessions. The most similar accessions were found to be AVDL and AVDN followed by AVHR and AVPL. The values of Nei’s genetic similarity validated the above findings. The genetic similarity derived from the data of RAPD marker for analysis all the accessions varied from 0.60 between AVSN and AVBS to 0.91 between AVDN and AVDL.

**Level of polymorphism**

To assess the genetic diversity of A. vasica, samples were collected from different regions in India and twenty random declares primers were used to amplify the genomic DNA isolated from the leaves and out of them five primers OPA-06, OPA-07, OPA-11, OPA-12, OPA-18 (Figures 1-5) showed the maximum of polymorphism. These primers generated total of 313 fragments of which 295 (94%) were polymorphic exhibiting a high degree of diversity amongst the accessions. The high level of polymorphism observed in the present material was similar to that observed in studies of Mangifera indica21, Olea europea22 and Malpighia glabra Linn.23 Primers (OPA 1-OPA 20) employed exhibited a wide variation in polymorphism ranging from 0-100% (Table 4). This was understandable as product amplification depends upon the sequence of random primers and their compatibility with genomic DNA. The number of markers detected by each primer found to be dependent on primer sequence and the extent of genetic variation, which was genotype specific.24 Out of the 20 primers used 8 of them gave satisfactory and reproducible amplification patterns exhibiting high degree of polymorphism as shown below in Figures 1-5 where OPA were operons and employed here as RAPD primers for carrying out the amplification of DNA in powdered leaf samples of A. vasica; M denoted the Molecular marker as 1 kb ladder and 1-10 denoted the different accessions of vasaka collected from different geographical locations.

**Dendrogram analysis**

The similarity matrix representing Jaccard’s coefficient was used to cluster the data following the UPGMA algorithm. To estimate the potential of individual primers for the characterization of variability in A. vasica, data obtained from the individual primers were...
Table 5: Dice similarity coefficient for RAPD data: Dice (Nei and Li, 1979)

<table>
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<th></th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
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<tbody>
<tr>
<td>1</td>
<td>1.000</td>
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<td>5</td>
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<td>6</td>
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<td>7</td>
<td>0.789</td>
<td>0.871</td>
<td>0.735</td>
<td>0.786</td>
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<td>1.000</td>
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<tr>
<td>8</td>
<td>0.848</td>
<td>0.853</td>
<td>0.716</td>
<td>0.882</td>
<td>0.813</td>
<td>0.644</td>
<td>0.863</td>
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<td>9</td>
<td>0.890</td>
<td>0.903</td>
<td>0.809</td>
<td>0.834</td>
<td>0.828</td>
<td>0.711</td>
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<td>10</td>
<td>0.829</td>
<td>0.821</td>
<td>0.709</td>
<td>0.885</td>
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<td>0.635</td>
<td>0.848</td>
<td>0.911</td>
<td>0.876</td>
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</tr>
</tbody>
</table>

1-New Delhi 2-Hisar 3-Kurukshetra 4-Nahan 5-Chandigarh 6-Banaras 7-Solan 8-Dehradun 9-Patiala 10-Dalhousie

processed separately (not shown). Different primers grouped the accessions in two major clusters. The dendrogram constructed for pooled data showed two major clusters. The dendrogram (Figure 6) based on SI (Table 5) showed distinct separation of the collected accessions though morphologically they were similar and inseparable. All the accessions were found in one cluster except AVBS while it was found in other cluster. The accessions which were found in another cluster were further sub divided into sub-clusters comprising of all the accessions except AVKU. The sub-cluster formed comprised of AVCH and was further subdivided into two trunks that in turn sub divided into upper sub cluster 1(USC1) and lower sub cluster 1 (LSC1). USC I comprised of collections from AVDL, AVDL, AVNH, AVND and the LSC I comprised of accessions from AVHR, AVPL, and AVSN. AVBS was marked by highest genetic distance from other accessions and hence clearly separated from them.

**DISCUSSION**

Due to easiness, rapidity and simplicity the RAPD technique has been used widely for differentiating a large number of medicinal species from their close relatives or adulterants including Panax species, Coptis species, Astragalus species, *Lycium barbarum* L., *Panax ginseng* and Echinacea species. These characters were especially advantageous for the identification of any herbal drug because of little DNA existing in the dried material. The significance of present work was mainly focused in differentiating genuine samples from the adulterated ones and to establish the level of genetic relatedness between them as our RAPD marker proved to be easily reproducible under wide variation of amplification conditions. Results were not affected at all with the changes in the origin of primer, the taq polymerase and the thermal cycler used in the experiment.

Various accessions of *A. vasica* collected from various locations as described earlier were subjected to RAPD studies as the same had already been found to be useful in differentiating the accessions of *Taxus wallichiana*, *Juniperus communis* L., *Allium schoenoprasum* L., *Andrographis paniculata* collected from different geographical regions. High degree of polymorphism was exhibited by the *vasaka* accessions and this might be attributed due to deletion, addition, substitution of base within the priming site sequence and this was in accordance with previously reported findings by Deshwal et al., 2005 as he reported 14 (58%) polymorphic primers with 73 amplification products in Neem accessions. The number of polymorphic primers and fragments generated may vary as product amplification depends upon the sequence of random primers and extent of genetic variations which in turn is genotype specific.

We reported the genetic diversity values in the range of 0.60 - 0.91 in *A. vasica*. The range of genetic diversity values broadly indicated the degree of heterogeneity and homogeneity in different accessions of plant species. The samples from AVBS and AVSN showed significant genetic diversity with similarity co-efficient value of 0.60 followed by that of collections from AVBS and AVDL. This was understandable as these samples were located far apart (> 2500 km), at different altitudes and belonging to two different geographical regions clearly showed that the climatic conditions and physical parameters might affect the plant genome and as the plant was adapted and these changes were inherited through genome to next generation. However the accessions AVDL and AVDN collected from Dalhousie and Dehradun displayed the maximum genetic similarity with a similarity co-efficient value of 0.91 followed by collection from AVHR and AVPL locations clearly stated the fact that collections from the almost similar altitudinal heights and geographical locations possess almost similar characteristics. The dendrogram obtained also estab-
lished the genetic relatedness among different accessions and separated all the accessions with the exceptions of AVDL and AVDN, AVHR and AVPL and thus considered as closely related genetically.

The gene diversity in *A. vasica* was comparatively of narrow range (0.60 - 0.91) with higher mean gene diversity value. The higher mean gene diversity could be explained as the samples were collected from different altitudes located at different geographical locations. Despite the collections from varying altitudes *A. vasica* showed somewhat lower range of genetic diversity which implied conservation of germplasm and lower level of heterogeneity. One of the possible reasons may be that the species has been endangered by human interventions.32 The collections from AVSN and AVBS showed a significant genetic diversity between them with a co-efficient value of 0.60. This is understandable as these samples were located far apart (>2500 km) at different altitudes and belong to different geographical locations clearly showed that the climatic conditions and physical parameters might affect the plant genome as the plant was adapted and these changes were inherited through one genome to next generation.

Based on the above findings there were the collections comprising of both high as well as low similarity indexes. High similarity indices suggested that the individuals in the population had close genetic relationship among them that in turn was reflection of adaptation to environment and beneficial to propagation, resources conservation and domestication of wild species. This situation could arise in natural populations where there was a possibility of free/random pollen flow and fertilization. The genetic similarity was closely related with their geographical locations.33 However, Cluster analysis had clearly indicated that there was eco-geographical isolation between the samples collected from different locations as sometimes even the geographically isolated individuals tend to accumulate genetic variations during the course of environmental adaptations.34

The present study suggested that RAPD is appropriate for the analysis of genetic variability in closely related genotypes. Moreover it could differentiate the plants collected from distant places belonging to same agro-climatic sub zones.

**CONCLUSION**

The present study confirmed the suitability of RAPD as a reliable, simple, easy to handle and as an elegant tool in molecular diagnosis of different accessions of *A. vasica*. RAPD hereby proved to be useful in molecular profiling of different accessions of *A. vasica*. The variations observed in the genetic diversity could be due to the use of random primers as well as the variations that naturally occur in the genotypes. The high amount of polymorphism (%) observed in the study could also be due to inclusion of vasaka genotypes with popping and non-popping characters.

The low level of genetic diversity observed in different accessions suggested the regional approach for the conservation of *A. vasica*. The species or at least a large part of its genetic diversity may be lost in the near future due to its medicinal and other uses and its consequent exploitation if appropriate conservation measures are not adopted. Since single or even a few plants will not represent the whole genetic variability in Adhatoda, there appears to be a need to maintain sufficiently large populations in natural habitats to conserve genetic diversity and to avoid genetic erosion.

**CONFLICT OF INTEREST**

The authors confirms that this article has no conflict of interest.

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