Detection of Molecular and Phytochemical Variations in Tamilnadu Landraces of Gymnema sylvestre Genotype: An Important Anti-Diabetic Plant

U. Jinu¹, M. Thiyagarajan¹, S. Javed², A. A. Alatar², and P. Venkatachalam¹,³

ABSTRACT

Gymnema sylvestre, a potential anti-diabetic medicinal plant, is used in many herbal drug formulations. The medicinal properties have been credited to the presence of gymnemic acid, which is effective in controlling blood glucose and lipid levels in the body. Molecular and biochemical variations of seven populations of G. sylvestre were investigated using DNA markers and phytochemical fingerprints of gymnemic acid content. Results obtained through these techniques were used in a comparative analysis and revealed a correlation between genetic variations and biochemical content analysis. Thirteen primers were used for RAPD-PCR analysis revealing 112 bands, of which 62 were found to be polymorphic. The percentage of polymorphic bands detected ranged from 11.1 to 84.6% with an average of 54.2%. The average number of amplified bands per primer was 8.62 while the mean number of polymorphic bands per primer was 5.42. A dendrogram of the genetic similarities among the populations was constructed using the genetic distance coefficients and the seven accessions were clustered into four major groups. The total gymnemic acid content level varied significantly among these groups from 3.84 to 7.12 mg g⁻¹ DW. A principle coordinate analysis confirmed the results of clustering. Maximum yield of gymnemic acid content (7.12 mg g⁻¹ DW) was recorded with L7 accession. Results suggest that there might be a positive correlation between genetic makeup and gymnemic acid content; however, this has to be further investigated by using other molecular approaches.

Keywords: Cluster analysis, Genetic diversity, Gymnemic acid, Random amplified polymorphic DNA.

INTRODUCTION

India is endowed with enormous natural wealth in terms of ecosystems, species richness, and genetic diversity, enlisting it in the 12 mega biodiversity hotspots of the world. Large numbers of plant species are being used for medicinal purposes in various traditional as well as modern medical systems. Herbal drug and pharmaceutical industry is one of the fastest growing industries for herbal-based medicine production. Currently, there is an increasing realization of health hazards associated with indiscriminate use of modern medicines. However, lack of scientific inputs in this area has also limited its potential uses and benefits. Moreover, unscientific wild harvesting of important medicinal plants have resulted in over exploitation of natural population and a threat to important gene pools.

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Gymnema sylvestre R. Br. (Apocynaceae), a woody shrub, found in the tropical and subtropical regions of Indian subcontinent, possesses great medicinal potential. It is also known as “Gur-Mar” in Hindi, literally meaning the “sugar killer” because of its effectiveness in the treatment of diabetes in the traditional medical systems of the Indian subcontinent. Recent scientific studies have confirmed its effectiveness in diabetes mellitus because of its hypoglycemic, anti-hyperlipidemic and anti-oxidant properties (Tiwari et al., 2014; Jain and Devi, 2016). These activities have been credited to gymnemic acid, a group of complex triterpenic glycosides. Gymnemagenin, a common aglycone of gymnemic acids produced after acidic and basic hydrolysis, inhibits glucose absorption and is commonly used by several workers as an analytical marker to determine the quality of Gymnema plant materials (Puratchimani and Jha, 2004).

Molecular and biochemical markers have opened a new avenue to investigate the genetic diversity. The identified markers could serve as basis for selection of elite genotypes and future improvement (Karaca and Ince, 2008). Also, Ince and Karaca (2015) have been successfully applied used the touchdown direct amplification of minisatellite-region DNA polymerase chain reaction (Td-DAMD-PCR) technique for identification of genetic diversity in Dianthus. Correlations between the metabolic contents and DNA fingerprints among geographically distinct plant populations have been reported in several species, Fructus xanthii (Han et al., 2008), Ocimum gratissimum (Vieira et al., 2001), Tanacetum vulgare (Keskitalo et al., 2001). However, no report is available on combined diversity analysis using molecular and biochemical methods in Gymnema spp. G. sylvestre shows high level of variability in bioactive compounds levels among the accessions collected from different locations (Pandey and Yadav, 2010). Variation in gymnemic acid content may affect the quality of pharmaceutical products and herbal medicine preparation. Although independent studies of either genetic or phytochemical profile analysis of various medicinal plants including Gymnema have been reported (Tripathi et al., 2012; Shilpha et al., 2013; Mahar et al., 2013), to the best of our knowledge, correlation studies for genetic and biochemical markers have not been done. Therefore, it is one of the prerequisites to conduct genetic diversity study for the analysis of gymnemic acid contents level among the different Gymnema accessions growing from various agroclimatic conditions. The genetic diversity study is highly essential for the selection of the high gymnemic acid yielding varieties for industrial applications in the future. Earlier reports suggest that a positive correlation exists between genetic diversity and the level of bioactive molecules, as recorded in various medicinal plants (Hu et al., 2007; Han et al., 2008; Azizi et al., 2012).

It is hypothesized that: (i) Identification of potential genotype for higher yield of gymnemic acid content may be beneficial for pharmaceutical industry, and (ii) This will be highly useful for selective cultivation of best genotypes for sustainable production. The present study aimed at establishing the correlation between genetic diversity with the variability in gymnemic acid yield among different accessions of G. sylvestre, so as to identify the potential accession with increased accumulation of bioactive molecules by estimation of genetic variability using molecular as well as phytochemical fingerprinting analysis.

**MATERIALS AND METHODS**

**Plant Materials**

Seven accessions of G. sylvestre were collected from various locations in Tamilnadu, India (Table 1). Young leaves were collected and used for DNA extraction and HPLC analysis. The accessions were selected to represent a wide range of geographical as well as agro-climatic conditions, based on distribution of G. sylvestre within Tamilnadu, India.
Table 1. Accession and collection details of G. sylvestre plants with longitude and latitude of the study area.

<table>
<thead>
<tr>
<th>Accessions ID</th>
<th>Area of the study</th>
<th>Latitude</th>
<th>Longitude</th>
</tr>
</thead>
<tbody>
<tr>
<td>L1</td>
<td>Horticulture Research Station, TNAU, Yercaud, Salem Dt, TN</td>
<td>11° 77’ N</td>
<td>78° 20’ E</td>
</tr>
<tr>
<td>L2</td>
<td>ABS Botanical garden, Kaaripatti, Salem (Dt), TN</td>
<td>11° 65’ N</td>
<td>78° 28’ E</td>
</tr>
<tr>
<td>L3</td>
<td>Centre for Siddha medicinal garden, Mettur, Salem Dt. TN</td>
<td>11° 47’ N</td>
<td>77° 48’ E</td>
</tr>
<tr>
<td>L4</td>
<td>TAMCOL Medicinal Garden, Kolli Hills, Namakkal Dt. TN</td>
<td>11° 14’ N</td>
<td>78° 20’ E</td>
</tr>
<tr>
<td>L5</td>
<td>Forest Research Centre, Kanjimalai, Salem, Dt. TN</td>
<td>11° 37’ N</td>
<td>78° 30’ E</td>
</tr>
<tr>
<td>L6</td>
<td>Yelgri Hills, Thirupattur, Vellore Dt. TN,</td>
<td>12° 34’ N</td>
<td>78° 38’ E</td>
</tr>
<tr>
<td>L7</td>
<td>Morappur, Salem Dt. TN</td>
<td>11° 11’ N</td>
<td>77° 19’ E</td>
</tr>
</tbody>
</table>

**Preparation of Hydro-Methanolic Extracts**

The coarse powder (25 g) was extracted with 500 mL of hydro-methanol (20:80) using a cold extraction method. The extract was filtrated using Whatman filter paper (No. 1) and then concentrated at 40°C using a rotary evaporator. The residue was preserved in ultra-deep freezer (-80°C) until further biochemical analysis.

**Acid Base Hydrolysis Method**

The leaf extract (50 mg) was weighed accurately and dissolved in 50% (v/v) methanol to make 50 mL. To 10 mL of this solution, 2 mL of 12% (w/v) potassium hydroxide (KOH) was added and heated on a boiling water bath for 1 h. After cooling, 5.5 mL of 4N HydroChloric acid (HCl) was added and heated on a boiling water bath for 1 h. Thereafter, on cooling, pH was adjusted to 8.0 using 12% (w/v) KOH, volume was maintained to 100 mL with 50% (w/v) methanol, filtered and subjected to HPLC quantification of biochemical compound.

**DNA Isolation**

Total DNA was extracted from leaves using a modified CTAB method based on the protocol of Doyle and Doyle (1990). DNA quality as well as integrity was analyzed on 0.8% (w/v) agarose gel electrophoresis. The DNA band was visualized under UV transilluminator and the image was captured by using gel documentation system (Alpha Inotech Gel Imaging System, USA). For each DNA sample, a series of assays were carried out to estimate the optimum DNA concentrations (15-25 ng µL⁻¹) for PCR amplifications. This was accomplished by comparing the dilutions to a known DNA standard after separation on 0.8% (w/v) agarose gel and stained with ethidium bromide. The DNA was diluted with sterile TE buffer to 10-15 ng µL⁻¹ and used for PCR amplification.

**High Performance Liquid Chromatography (HPLC) Analysis**

Quantification analysis was performed using HPLC (Shimadzu, Tokyo, Japan, model LC10A) instrument equipped with Shimadzu ApD-M 10 Avp Photodiode Array Detector (PDA) in order to determine peak purity and similarity test of gymnemagenin. HPLC grade solvents (Sigma Alrich, USA) were pre-filtered using a Millipore (Billerica, MA, USA) system and analysis was performed on a water column (Milford, MA, USA) C18 Spherisorb S10 ODS2 (250×4.6 mm, id 10 µm). The mobile phase was methanol:water (0.1% TFA) (50:50) at a flow rate of 0.6 mL min⁻¹. The detection wavelength was 210 nm, which was close to the absorption maxima for compound. The injection volume for sample was 20 µL and column temperature was 26°C.
DNA Amplification by Polymerase Chain Reaction (PCR)

Random oligonucleotide primers (Operon Inc, USA) were used for RAPD analysis (Williams et al., 1990). The reaction was carried out in a volume of 20 μL consisted of 2 μL, 10X PCR buffer (10 mM Tris HCl (pH 8.3), 50 mM KCl, 1.5 mM MgCl$_2$), 2 μL, 1.5 mM dNTPs (dATP, dGTP, dCTP and dTTP), 250 nM random decamer primers (1.0 μL), 0.5 units of Taq DNA polymerase, 2 μL of genomic DNA (15 ng) and finally added 13 μL of sterile water. DNA amplifications were performed in a PCR thermal cycler (Cyber lab, USA) under the amplification profile of initial denaturation at 94°C for 4 minutes, followed by 40 cycles consisting of denaturation at 94°C for 1 minute, annealing at 37°C for 1.30 minutes, extension at 72°C for 2 minutes with a final extension at 72°C for 7 minutes. After PCR cycles, loading buffer was added to the amplified products and they were analyzed on a PCR thermal cycler (Cyber lab, USA) under the amplification profile of initial denaturation at 94°C for 4 minutes, followed by 40 cycles consisting of denaturation at 94°C for 1 minute, annealing at 37°C for 1.30 minutes, extension at 72°C for 2 minutes with a final extension at 72°C for 7 minutes. After PCR cycles, loading buffer was added to the amplified products and they were analyzed on a 1.5% (w/v) agarose gel electrophoresis containing 0.5 μg mL$^{-1}$ ethidium bromide using 1X TAE buffer. Electrophoresis was performed at 60V for 2 hours until the bromophenol blue dye front migrated to the bottom of the gel. The molecular standard used was the lambda DNA double digested by EcoRI/Hind III. The gels were visualized under UV transilluminator and photographed with Alpha Inotech gel documentation system (USA). In order to optimize the DNA amplification rate, different concentrations of Taq DNA polymerase (0.1–0.5 units) and MgCl$_2$ (1.5 to 2.5 mM) were used for PCR amplification reactions. Among the random oligo nucleotide primers, those that exhibited clear banding pattern after PCR amplification were selected for further RAPD analysis (Venkatachalam et al., 2008).

DNA Fingerprinting Data Scoring and Analysis

RAPD banding patterns were analyzed by biostatic type scoring. The amplified DNA band was scored ‘+’ as presence and ‘-’ as absence for all prominent bands within a fingerprint profile (Collard and Mackill, 2009). Therefore, a sequence of ‘1’ and ‘0’ was generated for each primer/species to form a data matrix. DNA fragment sizes were estimated from the agarose gel by comparison with PCR molecular weight marker. RAPD-PCR amplification with each primer was performed with 5 replicates and repeated three times to obtain consistent banding pattern. A dendrogram was generated by cluster analysis using Unweighted Paired Group Method with Arithmetic average (UPGMA) based on Jaccard’s similarity co-efficient analysis. The fit of dendrogram obtained was checked by bootstrapping.

Phylogenetic Tree Construction

For evaluation purposes, the genetic relationship between different locations of Gymnema plant species and DNA bands from seven accessions produced by different random primers were scored and used for construction of phylogenetic tree. To identify the amplified DNA fragment size, double standard DNA marker (lambda DNA double digested by EcoRI/Hind III) was used to determine the size of each amplified DNA fragment. RAPD fragments were assigned a DNA length and recorded as a binary matrix for each individual as presence (1) or absence (0) of a given band. Phylogenetic relationship of G. sylvestre plant species was generated using Hierarchical clustering of DARwin 5.0 software based on UPGMA method.

RESULTS AND DISCUSSION

Gymnema sylvestre R.Br, is considered as a potential herb for diabetes treatment, because of the presence of important blood glucose and lipid regulating substances, specially gymnemic acid. However, G. sylvestre is a highly cross-pollinated species,
showing high levels of genetic variation among populations. The wild harvesting is thus less effective. It is also essential to conserve its gene pool from rapid disappearance caused by excess collection from the natural habitat. Therefore, identification of elite and potential genotypes is essential for sustainable industrial application. Genetic diversity analysis and its correlation to phytochemical yield would be one of the prerequisites for selection of potential and high yield germplasm accession and implementation of effective commercial crop cultivation for extraction of large-scale secondary metabolites. Although various molecular techniques are being used for genetic diversity analysis, the Randomly Amplified Polymorphic DNA (RAPD) analysis is one of the most effective PCR based methods as it is rapid, simple, cost effective tool, and reveals high levels of polymorphism among populations.

Assessment of Genetic Diversity by RAPD-PCR Analysis

Genomic DNA was isolated from healthy leaf tissue samples using modified CTAB method and used for RAPD-PCR analysis. PCR amplification yielded prominent bands and clear lanes with 10-15 ng DNA in the reaction mixture. PCR assay was carried out to obtain the optimum Taq DNA polymerase concentration, among the various concentrations tested, the intensity of DNA amplicons at 0.5 unit was found to be the best concentration for amplification of most stable/reproducible bands. The yield of RAPD products declined if the MgCl₂ concentration was increased from 1.5 mM to 2.5 mM in the reaction mixture and smeared lanes appeared in the electrophoresis profile pattern. One of the reasons may be due to the inhibition of Taq DNA polymerase activity in the presence of excess Mg²⁺ in the reaction mixture. In the present study, MgCl₂ at 1.5 mM concentration was found to be the best for PCR amplification using RAPD-PCR markers in DNA samples of *G sylvestre* (Figure 1).

The presence and absence of the DNA bands in the RAPD profile for six random primers is presented in Figure 1. RAPD profile generated using selected random primers was found to be clear and reproducible banding patterns and consistent for all the 7 accessions. For preliminary screening, about one hundred oligo-nucleotide primers were tested for amplification of the genomic DNA from the seven accessions of *G. sylvestre* plant species (L1 to L7). The RAPD primer sequence, total DNA bands, and polymorphic bands frequency details are represented in Table 2. Among the hundred primers screened, 13 random oligonucleotide primers (OPA04, OPA10, OPA13, OPB15, OPB17, OPB18, OPC06, OPC09, OPC16, OPC20, OPD03, OPD06 and OPD08) produced clear and reproducible DNA fingerprints for all the seven accessions of *G. sylvestre*. All the primers revealed DNA polymorphism among 7 populations and 62 amplicons were found to be polymorphic (Table 2). It is interesting to note that the average of DNA polymorphism noticed was 54.2% among the 7 accessions collected from different regions of Tamilnadu (20 to 280 km distribution, Table 2). The highest percent of polymorphic bands (84.6%) was noticed with OPA04 primer followed by OPC16 (81.8%) (Figure 1), while the lowest percentage polymorphism obtained was 11.1% with OPA13 (Figure 1). The number of polymorphic bands ranged from 1 to 11 with a mean average of 4.78 per primer. RAPD has been applied for determination of genetic diversity in *Withania somnifera* accessions collected from various geographical locations (Dharmar and Britto, 2011). Similarly, an attempt was made to determine the genetic variability among 17 accessions of *G. sylvestre* using 20 primers by RAPD-PCR technique and about 50% polymorphism...
Table 2. Primers selected for RAPD analysis, the total number of bands amplified by each primer and polymorphism among 7 *G. sylvestre* accessions collected from different geographical locations of Tamilnadu.

<table>
<thead>
<tr>
<th>Primers code</th>
<th>Nucleotide Sequence 5’ to 3’</th>
<th>Total No of bands</th>
<th>No of polymorphic bands</th>
<th>Polymorphism (%)</th>
<th>DNA bands size (in bp)</th>
</tr>
</thead>
<tbody>
<tr>
<td>OPA04</td>
<td>AATCGGGCTG</td>
<td>13</td>
<td>11</td>
<td>84.6</td>
<td>1509-200</td>
</tr>
<tr>
<td>OPA10</td>
<td>GTGATGCAG</td>
<td>13</td>
<td>7</td>
<td>53.8</td>
<td>895-200</td>
</tr>
<tr>
<td>OPA13</td>
<td>CAGCACCAC</td>
<td>9</td>
<td>1</td>
<td>11.1</td>
<td>1520-150</td>
</tr>
<tr>
<td>OPB15</td>
<td>GGAGGGTTT</td>
<td>6</td>
<td>3</td>
<td>50.0</td>
<td>698-350</td>
</tr>
<tr>
<td>OPB17</td>
<td>AGGGACCAG</td>
<td>11</td>
<td>7</td>
<td>63.6</td>
<td>1980-400</td>
</tr>
<tr>
<td>OPB18</td>
<td>CCAACGACT</td>
<td>9</td>
<td>4</td>
<td>44.4</td>
<td>1894-450</td>
</tr>
<tr>
<td>OPC06</td>
<td>GAACGACTC</td>
<td>6</td>
<td>2</td>
<td>33.3</td>
<td>1680-250</td>
</tr>
<tr>
<td>OPC09</td>
<td>CTCACCGTCC</td>
<td>10</td>
<td>6</td>
<td>60.0</td>
<td>1606-250</td>
</tr>
<tr>
<td>OPC16</td>
<td>CACACTCCAG</td>
<td>11</td>
<td>9</td>
<td>81.8</td>
<td>1872-200</td>
</tr>
<tr>
<td>OPC20</td>
<td>ACTTGCACCAC</td>
<td>3</td>
<td>2</td>
<td>66.7</td>
<td>876-400</td>
</tr>
<tr>
<td>OPD03</td>
<td>GTGGGCTCCA</td>
<td>7</td>
<td>3</td>
<td>42.8</td>
<td>1350-350</td>
</tr>
<tr>
<td>OPD06</td>
<td>ACCCTGACGG</td>
<td>3</td>
<td>2</td>
<td>66.7</td>
<td>980-450</td>
</tr>
<tr>
<td>OPD08</td>
<td>GTGTGCCCCA</td>
<td>11</td>
<td>5</td>
<td>45.4</td>
<td>1980-400</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td></td>
<td><strong>112</strong></td>
<td><strong>62</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Mean</strong></td>
<td></td>
<td><strong>8.62</strong></td>
<td><strong>4.78</strong></td>
<td><strong>54.2%</strong></td>
<td></td>
</tr>
</tbody>
</table>

Figure 1. RAPD pattern generated using various selected oligo-nucleotide primers. (Accession ID: L1- Yercaud Hills; L2- Kaaripatti; L3- Mettur; L4- Kollihills; L5- Kanjamalai Hills; L6- Yelgri Hills; L7- Morappur). Arrow (  ) indicates the presence of DNA band.
was reported by Balamuralikrishna et al. (2012). Earlier, Nair and Keshavachandran (2006) reported that the percent of DNA polymorphism was 73.2% recorded by using 15 RAPD primers. Lal et al. (2013) reported that RAPD fingerprinting profile generated by using 5 random primers revealed 100% polymorphism among 12 G. Sylvestre accessions collected from various locations of Gujarat. Recently, Mouna et al. (2014) reported considerable genetic diversity within the G. sylvestre populations collected from Karnataka region and about 85% of polymorphism was noticed with 12 accessions examined by ISSR marker analysis.

Genomic DNA from the seven different populations was amplified using random primers to detect the polymorphic DNA as molecular markers (Table 2). PCR amplification using OPA10, OPB15, OPB17, OPB18, OPC06, OPC09, OPC20, OPD03, OPD06 and OPD08 random primers produced different major DNA bands within 7 accessions. Figure 1 shows that populations L2, L3, L6 and L7 have a specific 1.6 kb DNA band amplified by OPA04 primer, while this band was absent in L1, L4, and L5 populations. In the case of OPC18 primer, a specific DNA fragment with the size of 1.2 kb was amplified in L4, L6, and L7 populations and it was not amplified in L1, L2, L3 and L5 populations (Figure 1).

For Principal Coordinates Analysis (PCoA), dissimilarity matrix values were calculated using a binary data matrix with Jaccard simple co-efficient analysis to ordinate 7 accessions from different locations on a scattered plot. The genetic distance values were used for hierarchical cluster analysis using UPGMA agglomeration method using Darwin 5.0 software. According to PCoA results, the codes of the accessions from each location were plotted on the four components corresponding to different regions, where the 7 populations were collected from Tamilnadu. The PCoA displayed the separation of populations (L2, L1), (L4, L3), (L6, L5) and (L7) according to genetic distances and based on dissimilarity matrix values on the scattered plot graph (Figure 2). It is noteworthy to mention that PCoA

![Figure 2. Principal coordinates analysis of RAPD markers from seven accessions of G. sylvestre (Accession ID: As defined in Figure 1).](image-url)
values migrated from negative to positive side on the scattered plot graph, suggesting the increase of means in high degree of similarity. Similar result was also reported recently in *Rauvolfia serpentina* (Nair et al. 2014).

The ratios of the common DNA markers and genetic distance coefficients among the populations were calculated using Jaccard co-efficient. The genetic distance coefficients among the populations were distributed between 0.22 and 0.51 (Table 3). A dendrogram generated from Jaccard similarity matrix and the UPGMA method revealed clear genetic relationships among the *Gymnema* accessions. A dendrogram of genetic similarities among the populations was constructed using the genetic distance coefficients shown in Table 3. Based on the clustering analyses, the populations were classified into four major distinct clusters: group I (populations L4 and L3), group II (populations L2 and L1), group III (populations L6 and L5) and group IV (population L7) (Figure 3). It is important to mention that accessions from different geographical origins were formed in the same clusters, for instance the accession L5 (Kanjamalai Hills) with L6 -Yelgri Hills and L3-Mettur with L4-Kolli Hills in the UPGMA dendrogram together notwithstanding the distance of 285 and 135 km between them. Interestingly, two geographically very near to accessions viz., L1-Yercaud Hills and L2-Kaaripatti were clustered into the same group and, on the contrary, the L7-Morappur population was very close to L1-Yercaud Hills but formed

Table 3. Similarity index matrix based on Jaccard’s simple coefficient analysis revealed by RAPD markers.

<table>
<thead>
<tr>
<th>Accession ID&lt;sup&gt;a&lt;/sup&gt;</th>
<th>L1</th>
<th>L2</th>
<th>L3</th>
<th>L4</th>
<th>L5</th>
<th>L6</th>
<th>L7</th>
</tr>
</thead>
<tbody>
<tr>
<td>L1</td>
<td>0.000</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>L2</td>
<td>0.222</td>
<td>0.000</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>L3</td>
<td>0.267</td>
<td>0.280</td>
<td>0.000</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>L4</td>
<td>0.400</td>
<td>0.384</td>
<td>0.319</td>
<td>0.000</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>L5</td>
<td>0.360</td>
<td>0.387</td>
<td>0.346</td>
<td>0.428</td>
<td>0.000</td>
<td></td>
<td></td>
</tr>
<tr>
<td>L6</td>
<td>0.350</td>
<td>0.358</td>
<td>0.337</td>
<td>0.475</td>
<td>0.246</td>
<td>0.000</td>
<td></td>
</tr>
<tr>
<td>L7</td>
<td>0.306</td>
<td>0.272</td>
<td>0.337</td>
<td>0.397</td>
<td>0.510</td>
<td>0.507</td>
<td>0.000</td>
</tr>
</tbody>
</table>

<sup>a</sup>L1- Yercaud Hills; L2- Kaaripatti; L3- Mettur; L4- Kolli Hills; L5– Kanjamalai Hills; L6- Yelgri Hills, L7- Morappur.

Figure 3. Un-weighted Pair Group Method with Arithmetic average (UPGMA) tree showing the genetic relationships between seven accessions of *G. sylvestre* as determined by RAPD markers.
separate cluster in the dendrogram and showed more genetic distance. Similar results were also reported earlier in *Rauvolfia serpentina* (Nair et al., 2014).

Based on the phytochemical analysis, the L7 accession remained having higher level of variation in the content rate than other accessions. The present results showed the existence of genetic variations among the 7 *G. sylvestre* accessions and the maximum distance recorded was 0.51 for L7 accession. Therefore, this population could be utilized for the production of high quality herbal formulations and also may be selected for large scale cultivation and supply of quality materials to pharmaceutical industry. Balamuralikrishna et al. (2012) reported that there was considerable association between molecular and biochemical diversity with respect to gymnemic acid content in *G. sylvestre*. Shahnawaz et al. (2012) studied the genetic diversity of *G. sylvestre* using RAPD and ISSR markers and reported the genetic diversity of 22 *Gymnema* populations spread across Western Ghats of Maharashtra. Similar genetic variability was also noticed in the present study. A relationship between genetic variability and geographic distribution has been reported earlier in other medicinal plants (Grass et al., 2006; Zong et al., 2008; Rahimmalek et al., 2009; Djabou et al., 2012; Sundaresan et al., 2012). In the present study, RAPD results provided useful insights to understand the existence of genetic variation through evolutionary dynamics among 7 populations of *G. sylvestre* across different geographical locations within Tamilnadu.

**Determination of Phytochemical Diversity by HPLC Analysis**

In identification of genetic variability using DNA fingerprints, there will not be any change irrespective of the tissue used for analysis, but the phytochemical content will largely vary depending on the tissue type used, due to the influence of physiological nature and environmental factors (Nair et al., 2014). In this context, it would be more appropriate to justify applying both DNA and phytochemical-based methods for determination of genetic diversity. In this context, it would be more appropriate to justify applying both DNA and phytochemical-based methods for determination of genetic diversity effectively. In order to find out the differences in phytochemical constituents level from *G. sylvestre* and also to provide comprehensive information on biochemical quality, the main bioactive molecule i.e., gymnemiacid was selected and used for analysis from 7 populations distributed in various locations within Tamilnadu (from 20 to 285 km distance). In the present study, the acid–base hydrolysis extraction method was used for extraction of higher yield of gymnemagenin content from the leaves of *G. sylvestre*. Gymnemagenin is a glycone of gymnemic acid and is separated after acidic and basic hydrolysis. Also, addition of water in methanol increased the yield of gymnemagenin to a certain extent by improving its solubility. The phytochemical fingerprint and gymnemiacid content level of 7 accessions was determined by using HPLC at an Ultra violet wavelength of 210 nm (Figure 4). Among the populations tested, the higher level of gymnemiacid content recorded was 7.18 mg g⁻¹ DW with accession L7, followed by L1 (6.97 mg g⁻¹ DW) and L2 (6.89 mg g⁻¹ DW), whereas the low level of gymnemiacid content observed was 3.84 mg g⁻¹ DW with L5 population (Figure 5). It is very important to mention that the DNA fingerprinting profile data showed maximum genetic variation in the UPGMA dendrogram for L7 accession that had the high gymnemic acid content followed by L1 and L2 populations. Results showed a strong positive correlation between molecular and phytochemical content variability in *G. sylvestre*. Phytochemical analysis clearly showed the presence of rich biochemical diversity among the 7 populations. The present results provide the evidences for existence of phytochemical diversity among the accessions tested and the best accession could be selected based on the level of bioactive compound for bio-resource conservation and pharmaceutical utilization.
Figure 4. Chromatograms of methanolic extracts of *G. sylvestre*. HPLC chromatograms showing presence of gymnemic acid at 2.8 minutes. (Accession ID: As defined in Figure 1).

Figure 5. Distribution of gymnemic acid in *G. sylvestre* (Accession ID: As defined in Figure 1).

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Molecular and Phytochemical Variations in Gymnema of this important medicinal herb. Earlier, Han et al. (2008) observed a correlation between genetic diversity and variations in phenolic contents of Fructus xanthii. Balamuralikrishna et al. (2012) also investigated the genetic diversity and compared with gymnemic acid content in G. sylvestre and observed a correlation between genetic and biochemical diversity. Recently, Nair et al. (2014) demonstrated that there was a considerable variation in the reserpine content levels among the populations tested and a maximum reserpine content noticed was 1.312 g 100 g⁻¹ with R. serpentina accession 7. It is also reported that the level of reserpine content from different accessions of this plant varied with geological conditions and the occurrence of variability might be largely due to changes in environmental conditions of the selected location (Nair et al. 2014).

The results of the present study reveal some relationships among genetic makeup, gymnemic acid content, and place of origin of each G. sylvestre population. The genetic relationships between populations from the same region were closer than those between populations from different regions. There were significant differences in the bioactive compound levels in accessions collected from the various locations of Tamilnadu (Table 1), suggests that genetic factors may also play a critical role in determining the level of bioactive compounds in the plants. Similar results were also reported earlier (Tao et al., 2009; Hu et al., 2007; Han et al., 2008; Azizi et al., 2012). Based on the phytochemical chromatograms and molecular markers, seven G. sylvestre populations were perfectly matched in PCoA groupings. Results strongly suggest that the level of gymnemic acid in each population was positively correlated with the data generated by genetic diversity analysis. Similarly, Ince and Karaca (2015) have successfully applied the e-microsatellite DNA markers to identify medicinal value species and quantitative detection of dry herbal products in Salvia species. In this context, the present findings clearly suggest the necessity of comparative analysis of both bioactive component level and genetic variability among the populations originating from different geographical regions to obtain ideal medicinal plant accession for pharmaceutical industry applications. Taken together, the present results reveal that the L7 (Morappur) accession, which showed maximum genetic diversity in the dendrogram, is positively correlated with the highest level of bioactive compound (Gymnemic acid) compared to other accessions. Moreover, one of the bands (1.6 kb) amplified with OPA 4 primer (Figure 1) corresponded closely with high yielding accessions. This can be further confirmed by using other molecular tools for screening high yielding genotypes.

CONCLUSIONS

In conclusion, the combined molecular and biochemical fingerprint analysis showed the existence of genetic and phytochemical diversity among 7 Gymnema populations originating from different geographical regions. Fingerprinting analysis showed a relationship between genetic distance and chemotype variations among the accessions tested. Dendrogram result strongly indicated that the maximum level of genetic diversity was with L7 accession, which had the highest level of gymnemic acid content (7.18 mg g⁻¹ DW) compared to other accessions. The influence of the geographic origin on genetic diversity and biochemical contents in G. sylvestre plants could provide an important insight to select the potential genotype for large-scale cultivation, and L7 population deserves special consideration for pharmaceutical applications due to the presence of higher gymnemic acid content. We also found a band that showed high correlation with the gymnemic acid production; this can be used as a marker. Further, this band can be sequenced to find out the genetic basis of this correlation. To the best of our knowledge, this is the first report to provide valuable information on existence of correlations between the genetic diversity and chemotype pattern in this important medicinal plant species.
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تشخیص تغییرات ملکولی و فیتوشیمیایی در نژادهای بومی Tamilnadu از ژنوتیپ Gymnema sylvestre

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چکیده

گیاهی است با پاتئسیل دارویی ضد دیابتی ی. جینو، م. تیاگارجان، س. جاود، ا. العطار، و پ. ونکاتاچالام

استفاده می شود. خواص دارویی آن به حضور اسید زیمیمیک (gymnemic acid) نسبت داده می شود که در G. sylvestre کنترل گلوکز خون و سطح چربی بدن می تواند در این نوع و تغییرات ملکولی و فیتوشیمیایی در جمعیت DNA با استفاده از نشانگر sylvestre و اثرات فیتوشیمیایی محتوای اسید زیمیمیک بررسی شد. نتایج به دست آمده از این روش در یک ثبت رنگ تغییرات استفاده شد و آشکارساخت که بین تغییرات زننکی و تجزیه محتوای مواد بوشیمیایی همبستگی وجود دارد. سبب آغازگر برای تجزیه RAPD-PCR می باشد. این نتایج در با داشت استفاده از ضرایب فواصل زیمیمیک و 7 درصد میزان افزایش توانایی میانگین معمول در زنبور عسل، به دست آمده، یک دندروگرام (dendrogram) از نشانگر زننکی با استفاده از ضرایب فواصل زننکی ساخته شد و 7 نمونه ثبت شده در 4 گروه اصلی ساخته شد. در میان یک گروه ها، كل محتوای اسید زیمیمیک بین 3/84mg/g DW تا 12mg/g DW گروه ها، كل محتوای اسید زیمیمیک بین 3/84mg/g DW تا 12mg/g DW نتین تغییر زیادی ومعناداری میکرد. نتایج و تحلیل مختصر تا نتایج دسته بندی مزبور را تایید کرد. حالی بسیاری همراه اسید زیمیمیک در L7 (4/8mg/g DW) نتین ثبت شده شد. نتایج حاکی از آن است که بین ساختار زننکی و محتوای اسید زیمیمیک احتمالا یک همبستگی مشت و موجود دارد، با این همه، این مطلب یاد با استفاده از دیگر روشهای ملکولی بیشتر پرسید.