

1
2 **Utilising inter-Random Amplified Polymorphic DNA, assess genetic**
3 **variability in indigenous medicinal plant Heart-leaved Moonseed**
4 **(*Tinospora cordifolia*)**
5
6

7 **Abstract**

8 The Ayurvedic medical system, which uses herbal preparations for its traditional
9 remedies, has been used in India since ancient times. One of the powerful immunomodulators
10 is *Tinospora cordifolia*, also known as AMRITA, which is credited to this miracle medication
11 for giving its user youth, longevity, and vigour.

12 RAPD has been widely and successfully used to identify several species of
13 medicinally significant plant. The need for research into and protection of therapeutic plant
14 species is growing. The perseverance of the evaluation was to assess the genetic diversity
15 among 10 accessions of *T. cordifolia* obtained from various districts in Tamil Nadu. 18
16 decamer primers were used in the analysis. Nine of them had repeatable, different
17 polymorphism patterns in their primers. RAPD investigation, it was discovered that the DNA
18 recovered from several *T. cordifolia* accessions exhibited random amplification and various
19 levels of genetic polymorphism. In the sample from Salem district the most genetic
20 polymorphism was found.

21 **Key words:** *Tinospora cordifolia*, Tamil Nadu, RAPD, Genetic diversity, Medicinal plants

22 **Introduction**

23 Since ancient times, people have employed plants as medicine. The Ayurvedic
24 medical system, which uses herbal preparations for its traditional remedies, has been used in
25 India since ancient times. The therapeutic benefits of these medicinal plants have been
26 thoroughly investigated in a number of scientific fields, including pharmacology, cosmetics,
27 fragrance, cuisine, etc. (Dubey et al., 2004). The term "Return to Nature" can be used to
28 describe the recent shift in Indian medicine from allopathic to herbal therapy. Since
29 millennia, medicinal plants have been highly valued for their curative qualities in treating
30 ailments and preventing disease. Plants have the capacity for both natural healing and acting
31 as potential antibacterial agents. According to estimations from the World Health
32 Organization (WHO), 80% of people worldwide depend on traditional ayurvedic medicines
33 to maintain a healthy way of living. (Foster et al., 2005).

34 T. cordifolia is also referred as "Guduchi" (Singh, 2008), although it is most often
35 referred to as "Giloya" a mythological Hindu term that refers to the holy concoction that
36 prevents the godly spirits from ageing and keeps them youthful eternally. *T. cordifolia*, a
37 herbaceous vine that grows 300 metres above sea level and found throughout the tropical
38 Indian subcontinent and China, is a member of the Menispermaceae family. Although *T.*
39 *cordifolia* is a resilient plant that can be cultivated in virtually any temperature, it loves a
40 warm environment for the best development. The greatest time to grow this plant is during
41 the rainy season, which is from July to August. This medicinal plant has the amazing capacity
42 to generate aerial roots that may extend 10 metres down and 10 metres up from the earth.
43 These aerial roots are crucial to the plant's ability to grow again. This plant doesn't actually
44 die; instead, it either drops to the ground or expands to a new "life line," reinstating itself. *T.*
45 *cordifolia* is a climber that needs help from the outside to flourish. Neem is known as NEEM
46 GILOY when it helps *T. cordifolia* flourish. Giloy and neem share a similar chemical makeup
47 and exhibit superior medicinal qualities when combined (Sharma et al., 2008).

48 *T. cordifolia* has a number of documented medical applications in Ayurveda.
49 According to "Ayurveda," the nutrient-rich starch extracted from *T. cordifolia*'s roots and
50 stems can treat diarrhoea and dysentery. Ayurvedic "Rasayanas" derived from *T. cordifolia*
51 strengthen the immune system and increase the body's resistance to numerous illnesses
52 (Singh et al., 2003). The fresh plant components are thought to be more effective at treating
53 illnesses. The plant's aqueous extract, known as Indian quinine, is effective in lowering high
54 body temperatures brought on by a cold or dyspepsia. Along with these other uses, it is
55 widely used for spermatorrhea, impotence, sclerotic disorders, urinary tract infections,
56 acidosis, simple debility, and sexually transmitted illnesses. In addition to being used to cure
57 leprosy and malaria, *T. cordifolia* roots also serve as natural sedatives (Kapoor, 2001).

58 A contemporary polymerase chain reaction-based approach for identifying genetic
59 changes across and within the same species is called random amplified polymorphic DNA
60 (RAPD). The DNA sequences of RAPD markers are amplified using a single, short, and
61 randomly chosen oligonucleotide primer (usually 10 bases long). RAPD has been widely and
62 successfully used to identify several species of medicinally significant plant. The need for
63 research into and protection of therapeutic plant species is growing. The following goal have
64 been set for the current study by taking into account the significance of *T. cordifolia* as a
65 medicinal plant and RAPD as a potent technique to evaluate the genetic polymorphism of
66 medicinal plants.

67 **Materials and Methods**

68 **Plant Collection**

69 *Tinospora cordifolia* leaves were collected in Tamil Nadu, India, from a variety of
 70 biological zones and altitudes ranging from 8 to 378 m MSL. Entirely of the plant samples
 71 were carefully categorised and brought back from the fields in ice boxes. The current
 72 research includes ten accessions of *T. cordifolia* from Trichy, Dindigul, Ariyalur, Ranipet,
 73 Vellore, Cuddalore, Coimbatore, Nagapattinam, Dindigul, Thirupathur (Table 1). Altitude
 74 was used as a reference point among other factors while examining genetic variation.

75 **Table 1. Samples from various Tamil Nadu locations collected for RAPD analysis**

| S. No. | Place | District | Latitude | Longitude | Altitude (MSL) | Code |
|--------|---------------|--------------|-------------------------|-------------------------|----------------|------|
| 1. | Thittacheri | Nagapattinam | 10 ⁰ 52'09"N | 79 ⁰ 47'08"E | 8 m | TCNT |
| 2. | Virudhachalam | Cuddalore | 11 ⁰ 30'52"N | 79 ⁰ 19'31"E | 33 m | TCCV |
| 3. | Jayankondam | Ariyalur | 11 ⁰ 12'45"N | 79 ⁰ 21'45"E | 61 m | TCAJ |
| 4. | Kalavai | Ranipet | 12 ⁰ 46'01"N | 79 ⁰ 25'01"E | 132 m | TCRK |
| 5. | Thalaivasal | Salem | 11035'10"N | 78045'30"E | 162 m | TCST |
| 6. | Vedasandur | Dindigul | 10 ⁰ 31'50"N | 77 ⁰ 56'54"E | 212 m | TCDV |
| 7. | Gudiyatham | Vellore | 12 ⁰ 56'12"N | 78 ⁰ 52'46"E | 268 m | TCVG |
| 8. | Kinethukadavu | Coimbatore | 10 ⁰ 48'39"N | 77 ⁰ 01'20"E | 308 m | TCKK |
| 9. | Vaniyampadi | Thirupathur | 12 ⁰ 41'23"N | 78 ⁰ 37'09"E | 341 m | TCTV |
| 10. | Palani | Dindigul | 10 ⁰ 26'57"N | 77 ⁰ 30'58"E | 378 m | TCDP |

76 **Extraction of genomic DNA using the modified CTAB technique**

77 *Tinospora cordifolia* leaves yielded DNA (5 gms). Using a pre-cooled mortar and
 78 pestle, leaves were pounded into a fine powder and then dissolved in liquid nitrogen. The
 79 powder was added to a centrifuge tube with 7 ml of preheated (65⁰ C) DNA extraction buffer,
 80 and the mixture was forcefully agitated to create a homogeneous suspension. The suspension
 81 was incubated at 65⁰ C for an hour, with inversion mixing performed every ten minutes. The
 82 mixture was extracted for around 15 minutes using an equal amount of phenol and
 83 chloroform (24:1). After incubation, centrifuged the samples at 10,000 rpm for 10 minutes.
 84 Transferring the top aqueous layer with care to a different centrifuge tube 0.6 litre of cold
 85 ethanol was added to this and stirred by inverting. Overnight, the samples were incubated at -
 86 20°C. The next morning, the samples were centrifuged for 10 minutes at 10,000 rpm after
 87 being mixed by inverting them. The particle was washed three times with 70% ethanol after

88 the supernatant was discarded. Overnight, The pellet was dried by air. After drying, the
 89 dissolved the pellet in 50 l of TE.

90 **PCR-RAPD- amplification**

91 The Gene Amp PCR System 9700 R (Applied Biosystems, Carlsbad, USA) was used
 92 for the PCR amplification, with the following amplification parameters: initial denaturation at
 93 94°C for 5 min, followed by 45 cycles of denaturation at 94°C for 1 min, primer annealing at
 94 37°C for 1 min, amplification at 72°C for 2 min, and a final extension at 72°C for 5 min.
 95 Amplified DNA was electrophoresed in a 1.4% agarose gel using a 1 TAE buffer
 96 electrophoresis at 50 V for three hours, and ethidium bromide was used to observe the
 97 electrophoresis. Gene Ruler™, a 1-kb DNA ladder, was applied as usual.

98 **Table 2 Primers list**

| S. No. | Codes | SEQUENCE | Tm ⁰ c |
|--------|----------|------------------------|-------------------|
| 1. | OPA - 01 | 5' GA GT CT CA GG 3' | 32.00 |
| 2. | OPC - 02 | 5' GG TC TA GA GG 3' | 32.00 |
| 3. | OPC - 14 | 5' GG AG TA CT GG 3' | 32.00 |
| 4. | OPA - 03 | 5' AA GC CT CG TC 3' | 32.00 |
| 5. | OPA - 09 | 5' GT CC CG AC GA 3' | 32.00 |
| 6. | OPC - 04 | 5' AG CC AG CG AA 3' | 32.00 |
| 7. | OPC - 15 | 5' TC TG TG CT GG 3' | 32.00 |
| 8. | OPA - 13 | 5' GA CC GC CT TG T 3' | 32.00 |
| 9. | OPA - 17 | 5' GG GT AA CG CC 3' | 32.00 |
| 10. | OPA - 20 | 5' GT TT GC GA TC C 3' | 32.00 |
| 11. | OPG - 11 | 5' GT GA GA CG GA 3' | 32.00 |
| 12. | OPC - 11 | 5' AC CG AG CG AT 3' | 32.00 |
| 13. | OPG - 02 | 5' TG TG TG CT AA 3' | 32.00 |
| 14. | OPA - 04 | 5' GA TT GC CA AG T 3' | 32.00 |
| 15. | TC - 01 | 5' GG CT AA CG GC 3' | 32.00 |
| 16. | TC - 02 | 5' GG AC TG CA GA 3' | 32.00 |
| 17. | TC - 03 | 5' AG TC AG CC AC 3' | 32.00 |
| 18. | OPG - 24 | 5' CA GG CC CT TC 3' | 32.00 |

100 The presence or lack of bands was used to assess genetic similarities. DNA bands
101 may be seen and rated as present (1), absent (0), or not present, and a binary matrix was
102 created. MVSP software was used for data processing, and genetic diversity was investigated.

104 **RESULTS AND DISCUSSION**

105 Researchers have identified 152 species in India that might be used to create
106 ayurvedic medications, and *T. cordifolia* is one of these potentially useful plants (Srivastava
107 et al., 2011). RAPD markers have the capacity to detect and define the genetic diversity of
108 plants (Rout, 2002). Proper plant management of its genetic resources is aided by the many
109 genetic components found in the crop and detected by markers reported Saha and co-workers
110 in 2010. Despite the availability of several molecular markers, RAPD markers are explored in
111 the current research because of their implications.

113 **Sample collection**

114 *T. cordifolia* stems were gathered in ten different places in Tamil Nadu, ranging from
115 8.00 to 378.00 meters above (Table 1) over the years of 2022. Plant samples were collected
116 from Thittacheri village of District Nagapattinam at the peak of 8m above MSL (Mean Sea
117 Level), one town from municipality Cuddalore i.e Virudhachalam at the peak of 33m above
118 MSL one town from municipality Ariyalur i.e Jayankondam at the peak of 61m above MSL
119 one town from municipality Ranipet i.e Kalavai at the peak of 132m above MSL, one town
120 from municipality Salem i.e Thalaivasal at the peak of 162m above MSL, one town from
121 municipality Dindigul i.e Vedasandur at the peak of 212 m above MSL, one village from
122 District Vellore i.e Gudiyatham at the peak of 268m above MSL, one town from
123 municipality Coimbatore i.e Kinethukadavu at the peak of 308m above MSL, one village
124 from District Thirupathur i.e Vaniyampadi at the peak of 341m above MSL, and one village
125 from District Dindigul i.e Palani at the peak of 378m above MSL. The RAPD analysis and
126 antibacterial properties of *T. cordifolia* have both been the subject of several investigations.
127 For instance, Mohapatra and Rout, 2005 reported fresh plant samples were gathered from
128 Kanyakumari District Tamil Nadu, while leaves of *T. cordifolia* were taken from
129 Ambasamudram , Tirunelvi, Sankarankoil, Tenkasi and Alangulam, in District Tamil Nadu
130 (Britto et al., 2010).

131 ***Tinospora cordifolia* molecular profiling with RAPD markers**

132 The genetic link and degree of relatedness among the several *T. cordifolia* accessions
133 were discovered using RAPD primers. 18 decamer primers were first screened for

134 polymorphism analysis, including TC - 01, TC - 02, TC - 04, OPA - 01, OPA - 03, OPA - 04,
 135 OPA - 09, OPA - 13, OPA - 17, OPA - 20, OPC - 02, OPC - 04, OPC - 11, OPC - 14, OPC -
 136 15, OPC - 24, and OPG - 11 (Table 2). Nine primers were chosen from among them based on
 137 the highest level of polymorphism found in all DNA samples extracted from *Tinospora*
 138 leaves collected from various places in Himachal. This was also utilised to identify and
 139 examine the genetic connections among the 10 plant accessions. The resultant amplified
 140 fragments were examined for genetic variants and relationships based on the bands are
 141 present or not. DNA bands were either present means indicted as one (1) or absent means
 142 indicted as zero (0), depending on data. After RAPD, the bands were seen under a UV
 143 transilluminator, and gel documentation system photographs were made, as shown in Figure
 144 3. (A-H). All of the analysed samples revealed polymorphism. Less polymorphic primers
 145 included OPC – 15, OPC - 04, OPA -13 and OPA - 09 whereas markers OPA – 03 and OPC -
 146 14 had fair to outstanding polymorphism, as did primers OPA-01 and OPC-2. Further cluster
 147 analysis was performed using the RAPD primer OPA-01. Table 3 provides an overview of
 148 primer codes, sequences, number of amplified alleles, number of polymorphic alleles, number
 149 of monomorphic alleles, polymorphic per cent and amplification status.

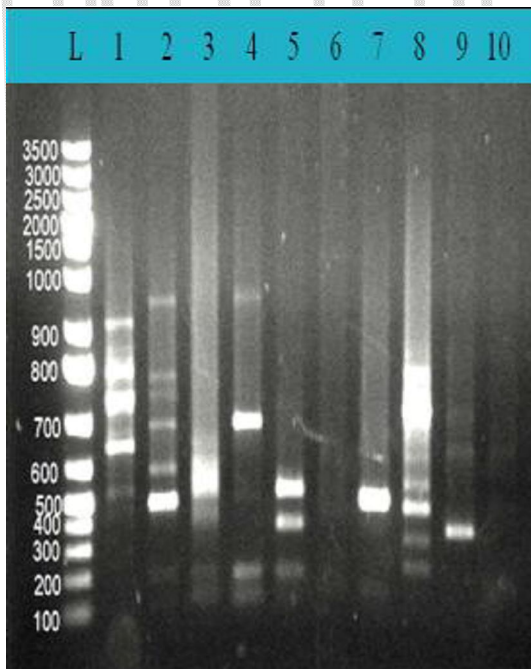
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152 **Table 3. *T. cordifolia* sample RAPD analysis using primers**

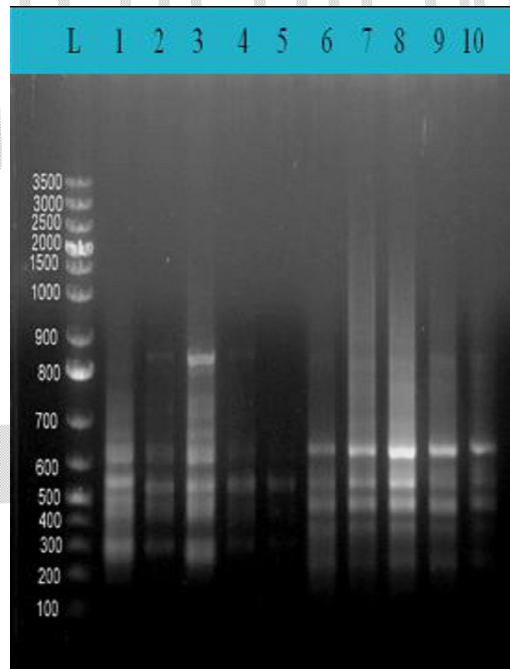
| S. No. | CODES | No. of amplified alleles | No. of polymorphic alleles | No. of monomorphic alleles | polymorphic % | Amplification Status |
|--------|----------|--------------------------|----------------------------|----------------------------|---------------|----------------------|
| 1 | OPA - 01 | 6 | 6 | 0 | 100 | Amplified |
| 2 | OPC - 02 | 5 | 4 | 1 | 80 | Amplified |
| 3 | OPC - 14 | 6 | 6 | 0 | 100 | Amplified |
| 4 | OPA - 03 | 4 | 4 | 0 | 100 | Amplified |
| 5 | OPA - 09 | 5 | 5 | 0 | 100 | Amplified |
| 6 | OPC - 04 | 4 | 4 | 0 | 100 | Amplified |
| 7 | OPC - 15 | 4 | 4 | 0 | 100 | Amplified |
| 8 | OPA - 13 | 4 | 4 | 0 | 100 | Amplified |
| 9 | OPA - 17 | - | - | - | - | No Amplification |

| | | | | | | |
|----|----------|---|---|---|---|------------------|
| 10 | OPA - 20 | - | - | - | - | No Amplification |
| 11 | OPG - 11 | - | - | - | - | No Amplification |
| 12 | OPC - 11 | - | - | - | - | No Amplification |
| 13 | OPG - 02 | - | - | - | - | No Amplification |
| 14 | OPA - 04 | - | - | - | - | No Amplification |
| 15 | TC - 01 | - | - | - | - | No Amplification |
| 16 | TC - 02 | - | - | - | - | No Amplification |
| 17 | TC - 03 | - | - | - | - | No Amplification |
| 18 | OPG - 24 | - | - | - | - | No Amplification |

153

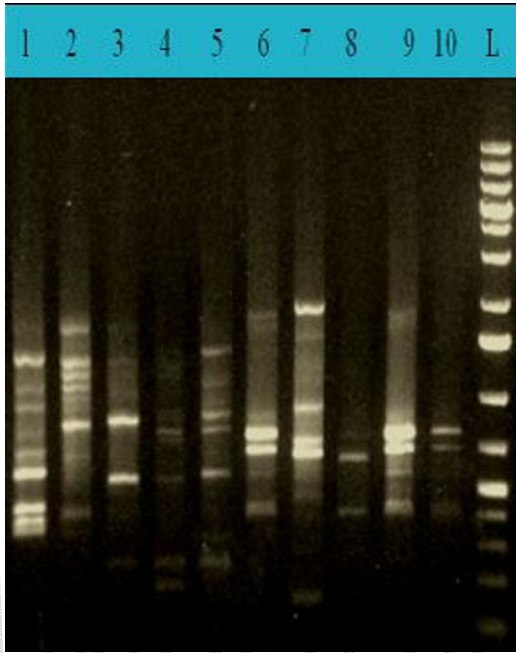


I* OPA 01

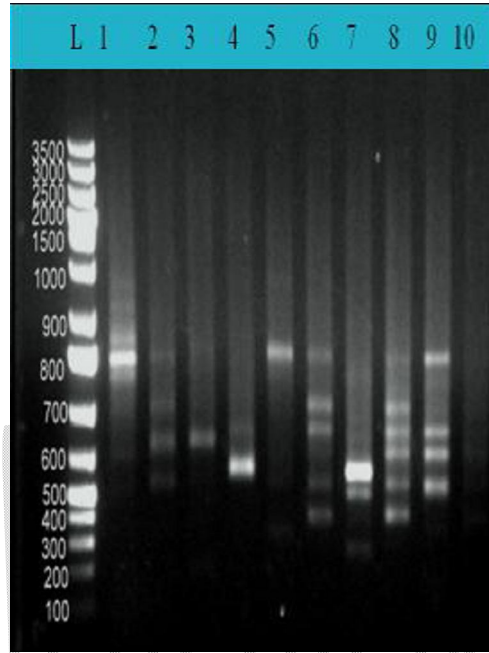


II* OPC 02

154

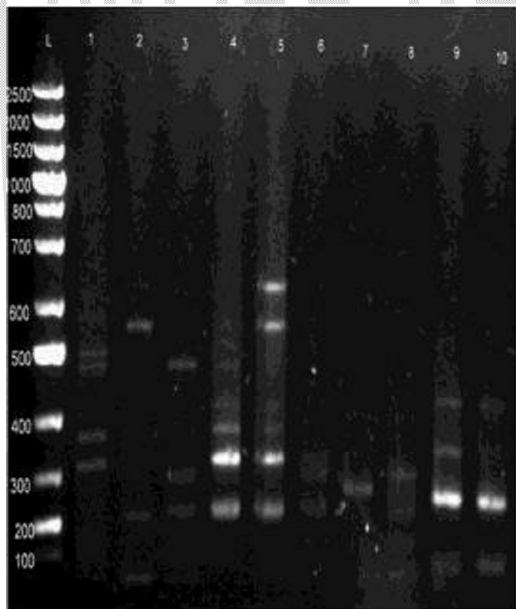


III* OPC14

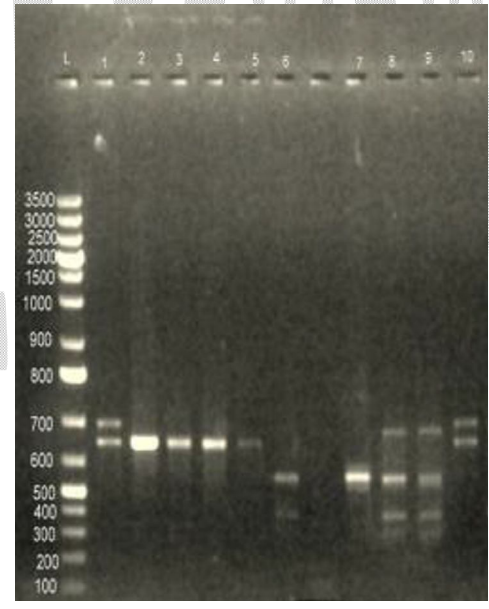


IV* OPA03

155

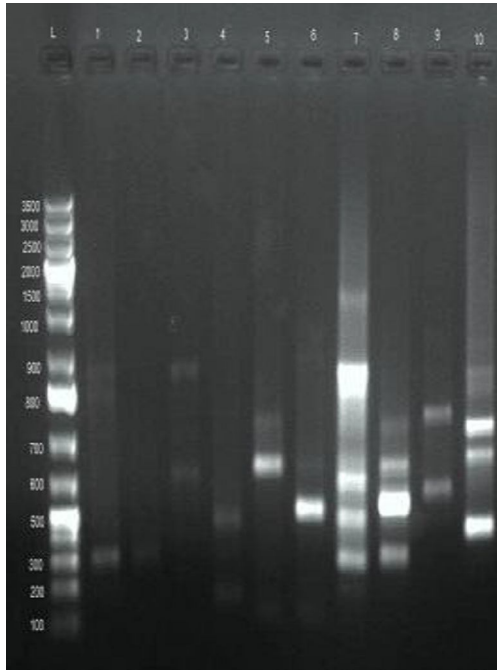


V* OPA - 13

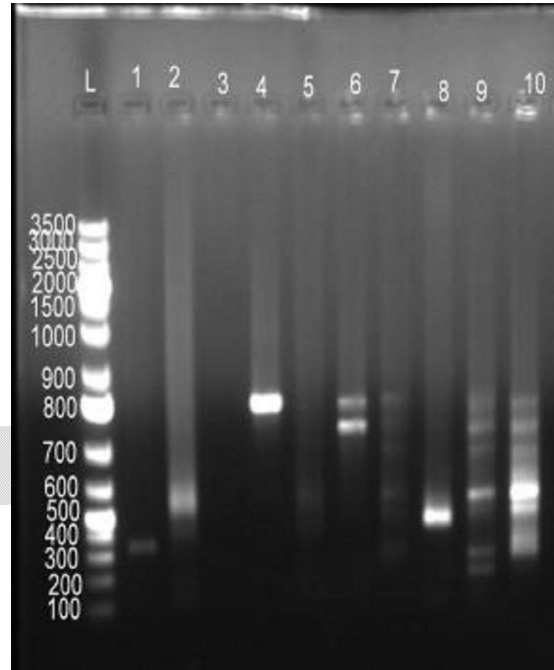


VI* OPA - 04

156



VII* OPC - 04



VIII* OPC - 01

157

158 **Figure 1:** I and II demonstrate tha samples 1, 2, 3, 4, 5, 6, 7, 8, 9 and 10 underwent random
 159 amplification using the RAPD primers OPA01 and OPC2, which demonstrates the greatest
 160 polymorphism; C and D demonstrate that samples 1, 2, 3, 4, 5, 6, 7, 8, 9 and 10 underwent
 161 random amplification using the RAPD primers OPC14 and OPA03, which demonstrates the
 162 finest polymorphism; OPA-13, OPA-4, OPC-4, and OPC-1 RAPD primers, random
 163 amplification was performed in samples 1, 2, 3, 4, 5, 6, 7, 8, 9, and 10, and the results
 164 indicated minimal polymorphism (E, F, G, and H).

165 *L (**Ladder**) Sample from Lane 1 comes from Thittacheri, Nagapattinam; Sample from lane 2
 166 comes from Virudhachalam, Cuddalore; Sample from Lane 3 comes from Jayankondam,
 167 Ariyalur; Sample from Lane 4 comes from Kalavai, Ranipet; Sample from Lane 5 comes
 168 from Thalaivasal, Salem ; Sample from Lane 6 comes from Vedasandur, Dindigul; Sample
 169 from Lane 7 comes from Gudiyatham, Vellore; Sample from Lane 8 comes from
 170 Kinethukadavu, Coimbatore; Sample from Lane 9 comes from Vaniyampadi, Thirupathur;
 171 Sample from Lane 10 comes from Palani, Dindigul

172

173 Initial RAPD results showed that 9 primers RAPD, namely OPC 2, OPA 01, OPC 14,
 174 OPA 13, OPA 03, OPA 09, OPC 1 and OPC 4 with 10 distinct samples of *T. cordifolia*
 175 accessions obtained from several districts in Tamil Nadu, exhibited amplification and varying

176 degree of genetic polymorphism. In the sample taken from Thalaivasal in the District of
177 Salem, primer OPA01 demonstrated good polymorphism, while samples taken from other
178 locations revealed significant genetic variety. Figure 1A displayed all of the distinct bands
179 produced by amplifying using OPA 01 primer. Using Alfaview software, the amplified
180 fragment sizes were estimated and discovered to be between 200 and 900 bp. Thus, the
181 genetic relationship and degree of polymorphism of the various *T. cordifolia* samples were
182 assessed using the illuminating OPA-01 primer as well as the sample obtained from Salem
183 had the highest level of polymorphism.

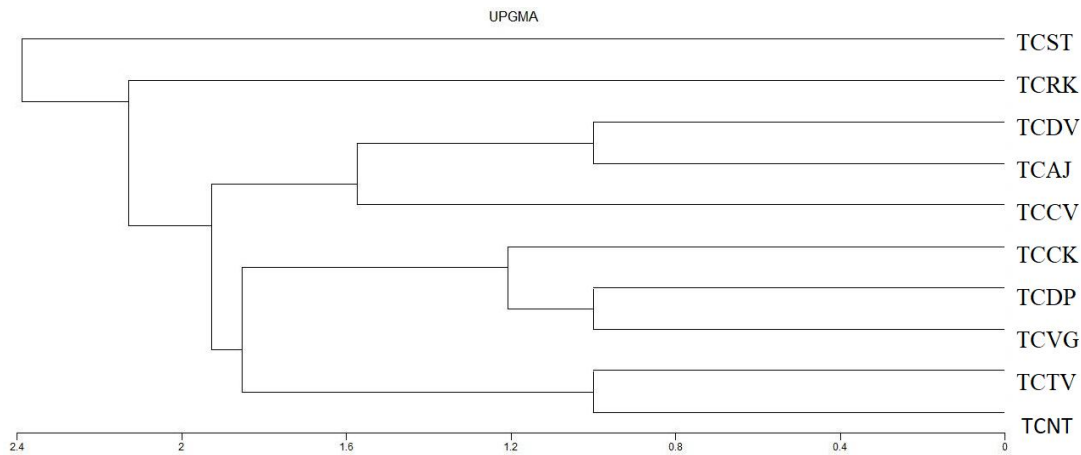
184 Using RAPD markers, the genetic diversity of 15 Bhubaneswar-collected *T. cordifolia*
185 clones was evaluated. There were 40 decamer primers used. To determine the genetic links
186 among 15 clones, 15 primers were utilised, and these primers produced unique fragments of
187 DNA (Rout, 2006). Using RAPD markers, Shinde et al. (2010) discovered limited amount of
188 *T. cordifolia*'s genetic diversity. Twenty Western Himalayan *T. cordifolia* samples were
189 chosen for genetic analysis using RAPD markers. Out of the 120 decamer oligonucleotide
190 primers employed, four RAPD primers only OPC-13, OPA-16, OPC-07 and OPC-05
191 exhibited minimal genetic variation of *T. cordifolia* (Sharma et al., 2006). 40 germplasm
192 were gathered from different parts of India, including Delhi, Kerala, Himachal Pradesh,
193 Assam, and Jammu & Kashmir. Three species of *Tinospora* were polymorphized by five out
194 of nine primers (Ahmad et al., 2009). The unweighted paired group method (UPGM)
195 dendrogram produced by Ishnava et al. (2009) demonstrated the extent of genetic diversity
196 among several *T. cordifolia* germplasm.

197 **Cluster analysis and Similarity matrix**

198 Ten samples' dendrogram demonstrating their genetic relatedness shows on fig. 2. The
199 samples TCDP, TCST, TCRM1, and TCCK were discovered to be genetically related and to
200 have reduced polymorphism. While distantly linked to samples TCDP and TCST, sample
201 TCDV was genetically distinct from samples TCRM1 and TCCK. Despite being distantly
202 related to sample TCVG, TCTV and TCRK shared genetic similarities. The samples TCDP
203 and TCST were closely linked to TCRK, TCTV, and TCVG. Genetically, TCVG and TCDV
204 were not related. The greatest polymorphism samples, TCNT and TCAJ, were distantly
205 linked to each other and to all other samples. The genetic makeup of TCAJ was distinct from
206 all other samples, demonstrating the most genetic variety among the *T. cordifolia* accessions
207 that had been gathered.

208

209



210

211 **Figure 2.** Ten samples' dendrogram demonstrating their genetic relatedness

212

213 **Conclusion**

214 Ten accessions of *T. cordifolia* were gathered for the current study from diverse
 215 locations throughout several districts of Tamil Nadu. The leaf samples of *T. cordifolia* that
 216 were procured from several areas in Tamil Nadu contained PCR amplifiable genomic DNA.
 217 Although leaves reacted to the improved CTAB technique of DNA separation, the leaves
 218 consistently produced large amounts of DNA. Using RAPD (random amplified polymorphic
 219 DNA) markers, the genetic polymorphism of multiple *T. cordifolia* samples was examined.
 220 Nine primers out of the 18 primers were capable of spotting differences between the 10 *T.*
 221 *cordifolia* samples that were obtained. Following analysis of RAPD, it was discovered that
 222 the DNA recovered from several *T. cordifolia* plants exhibited random amplification and
 223 various levels of genetic polymorphism. In the sample from Salem district the most genetic
 224 polymorphism was found.

225 **Reference**

- 226
- 227 • Ahmad SM, Hoot SB, Qazi PH, Verma V. 2009. Phylogenetic patterns and genetic
 228 diversity of Indian *Tinospora* species based on chloroplast sequence data and cytochrome
 P450 polymorphisms. *Plant Systematics and Evolution*. 56:751-767.
 - 229 • Britto AJD, Stephan TLR, Petchimuthu KP, Kumar BJR, Mary RS, Dharmar K. 2010.
 230 Inter Population Genetic Variability in *Tinospora cordifolia* (wild.) Miers ex Hook. F. &
 231 Thoms. (Menispermaceae), through RAPD marker. *Scientia Acta Xaveriana*. 0976-
 232 1152:57-64.
 - 233 • Dubey NK, Kumar R, Tripathi P. 2004. Global promotion of herbal medicines: India's
 234 opportunity. *Current Science*. 86: 37-41.

- 235 • Foster B.C, Arnason J.T, Briggs C.J. 2005. Natural health products and drug disposition.
236 Annual review of pharmacology and toxicology. 45: 203-226.
- 237 • Ishnava K and Mohan JSS. 2009. Assessment of Genetic Diversity in Medicinal Climber
238 of *Tinospora cordifolia* (Willd.) Miers (Menispermaceae) from Gujarat, India. Asian
239 Journal of Biotechnology. 1(3):93-103.
- 240 • Kapoor LD. 2001. Hand book of Ayurvedic Medicinal plants. 18–19.
- 241 • Mohapatra A and Rout GR. 2005. Identification and analysis of genetic variation among
242 rose cultivars using random amplified polymorphic DNA. Z Naturforschung. 60: 611-
243 617.
- 244 • Rout GR. 2002. Direct plant regeneration from leaf explants of *P'lumbago* species and its
245 genetic fidelity through RAPD markers. Annals of *Applied Biology*. 140(3): 305-313.
- 246 • Rout GR. 2006. Identification of *Tinospora cordifolia* (Willd.) Miers ex Hook F &
247 Thomas using RAPD markers. Z Naturforsch C. 61:118–22.
- 248 • Saha S, Dey ST, Adhikari S, Ghosh PD. 2010. *In vitro* multiple shoot regeneration and
249 analysis of genetic fidelity of *Mentha piperita* L. Bionature. 30(2): 71-81.
- 250 • Sharma AC, Shanker LK, Tyagi, Singh M, Rao CV. 2008. Herbal medicine for market
251 potential in India: An overview. African Journal of Plant Science. 1: 26-36.
- 252 • Sharma R, Kumar V, Mohapatra T, Khandelwal V. 2006. A simple and non-destructive
253 method of direct-PCR for plant systems. Journal of Plant Biology. 55: 114-122.
- 254 • Shinde VM and Dhalwal K. 2010. DNA Fingerprinting of *Tinospora cordifolia* using
255 RAPD analysis. Journal of Global Pharma Technology. 56:751-767.
- 256 • Singh A. 2008. A note on variation of active principles in Indian medicinal plants and
257 TIM formulations. Journal of Ethnobotanical Leaflets. 12: 603–606.
- 258 • Singh SS, Pandey SC, Srivastava S, Gupta VS, Patro B, Ghosh AC. 2003. Chemistry and
259 medicinal properties of *Tinospora cordifolia* (Guduchi). Indian Journal of Pharmacology.
260 35:83—91.
- 261 • Srivastava P. 2011. *Tinospora cardifolia* (*Amrita*)- A miracle herb and lifeline to many
262 diseases. International Journal of Medicinal and Aromatic plants. 1 (2):57-61.