



Molecular Characterization of Mulberry Germplasm through ISSR Markers

**Aruna N. ^{a*}, Mavilashaw V P. ^{b++}, Saranya, M. ^{c++},
Krishnamoorthy S V. ^{a#} and Parthiban K T ^{d†}**

^a Department of Sericulture, Forest College and Research Institute, Mettupalayam, Tamil Nadu, India.

^b Department of Agriculture Entomology, The Indian Agriculture College, Radhapuram, Tirunelveli, Tamil Nadu, India.

^c Department of Entomology, Adhiparasakthi Agricultural College, Kalavai, India.

^d Forest College and Research Institute, Mettupalayam, Tamil Nadu, India.

Authors' contributions

This work was carried out in collaboration among all authors. All authors read and approved the final manuscript.

Article Information

DOI: <https://doi.org/10.9734/ijpss/2025/v37i95702>

Open Peer Review History:

This journal follows the Advanced Open Peer Review policy. Identity of the Reviewers, Editor(s) and additional Reviewers, peer review comments, different versions of the manuscript, comments of the editors, etc are available here: <https://pr.sdiarticle5.com/review-history/142867>

Original Research Article

Received: 28/06/2025

Accepted: 23/08/2025

Published: 27/08/2025

ABSTRACT

Inter simple sequence repeat (ISSR)-PCR is a technique, which involves the use of microsatellite sequence as primers in a polymerase chain reaction to generate multilocus markers. Twenty-four mulberry accessions were obtained from the Central Sericulture Germplasm Resource Center, Hosur, Tamil Nadu were screened using Inter simple sequence repeat technique, with eight primers. Out of eight primers, only three primers viz., UBC 807, UBC 809 and UBC 811 produced 100 per cent polymorphism in mulberry accessions. The highest Polymorphism information content value was observed with the primer UBC 807 and the lowest PIC value was observed with the primer UBC 811. The mulberry accessions A18 (MI-0685) and A8 (MI-0535) were found to be the

⁺⁺ Assistant Professor;

[#] Professor (Agricultural Entomology);

[†] Professor (Agroforestry);

*Corresponding author: E-mail: aruna6371@gmail.com;

Cite as: Aruna N., Mavilashaw V P., Saranya, M., Krishnamoorthy S V., and Parthiban K T. 2025. "Molecular Characterization of Mulberry Germplasm through ISSR Markers". *International Journal of Plant & Soil Science* 37 (9):249–257. <https://doi.org/10.9734/ijpss/2025/v37i95702>.

closest at the similarity coefficient of 0.93. Mean genetic diversity and PIC were 0.35 to 0.29, respectively. The genetic similarity coefficient ranged from 0.913 A18 (MI-0685) to 0.535 A8 (MI-0535). The phylogenetic tree was obtained using the UPGMA method using the total number of amplified Inter simple sequence repeat fragments. The result from the cluster analysis indicated that the mulberry accessions MI-0017, MI-0763, MI-0663, MI-0652, MI-0349, V-1 (check variety), MI-0034 and MI-0768 were distinct from other accessions.

Keywords: Mulberry; molecular marker; genetic diversity; inter simple sequence repeat.

1. INTRODUCTION

"Mulberry (*Morus indica*. L) a deep rooted perennial deciduous herb belonging to the family *Moraceae* is a high biomass producing foliage plant. It is exploited on a commercial scale for silk production. It has different ploidy levels and most of the genotypes available are diploids ($2n=28$), but the plant exhibits higher ploidy levels *i.e.*, $3n=42$, $4n=56$, $6n=84$ and docasaploid $22n=308$ " (Basaviah et al., 1989). "Leaf quality is an important parameter used for evaluation of mulberry plants aimed at selecting superior varieties for rearing performance. Around 60% of the total cost of cocoon production goes towards mulberry production alone in Sericulture" (Veerapuram et al., 2013). "Germplasm resources are very much required for the continuous improvement of crop plants, where genetic characterization is essential for scientific germplasm conservation and characterization permits the estimation of genetic relatedness and diversity" (Keshavamurthy et al., 2013). "In order to assess the characterization among species, DNA markers have gained popularity among conservationists, geneticists and breeders basically because of its stability and ease to generate maximum information in comparatively less time. PCR based markers have the potential to screen large number of samples quickly to elucidate accurate genetic information" (Girish et al., 2015). "Recently, DNA based molecular markers have increasingly been applied in tree improvement programmes for quantification of genetic diversity. These markers have little been affected by environment and developmental stages" (Haines and Martin, 1995). "Markers also have important immediate application in supportive research for tropical hardwoods and non-industrial species mainly for quantification of genetic diversity. There are several methods based on DNA markers *viz.*, RFLP (Restriction Fragment Length Polymorphism), RAPD (Randomly Amplified Polymorphic DNA), SSR (Simple Sequence Repeats), ISSR (Inter Simple Sequence Repeats), AFLP (Amplified Fragment Length

Polymorphism) etc. Among these techniques, ISSR marker is very significant because this marker in its simplest form, does not require genomic knowledge to design arbitrary sequence oligonucleotide primers" (Gonzalez et al., 2005). But the use of molecular markers for genetic diversity studies in mulberry underscores the research needs.

2. MATERIALS AND METHODS

2.1 Plant Materials

Table 1 lists the accessions of mulberry used in the current study and their origins. Twenty-four mulberry materials, were obtained from the Central Sericulture Germplasm Resource Center, Hosur, Tamil Nadu.

2.2 Extraction and Quantification of Genomic DNA

It is necessary to check the quality and concentration of DNA for carrying out PCR reaction. Genomic DNA was quantified using the spectrophotometer and the concentration of nucleic acids were determined by measuring the sample OD (optical density) at 260 nm against blank. The ratio A260/A280 was used to estimate the purity of nucleic acid. Pure DNA had a ratio approximately ~1.8-2.0. The quality of DNA was also checked using 0.8 per cent agarose gel electrophoresis. The DNA was then diluted to appropriate concentration for use in molecular analysis.

2.3 Polymerase Chain Reaction Using ISSR Primers

"Extracted and quantified genomic DNA was subjected to polymerase chain reaction (PCR) using a thermal cycler (Eppendorf master cycler personal, Germany) for performing ISSR profiling" (Williams et al., 1990) to identify polymorphism from the collected germplasm. This included random decamers as primers and

the list of primers (Table 2) used for the polymorphic markers towards identification and characterization of the germplasm and the varieties within the species.

DNA from the collected twenty four test accessions along with V-1 as check variety was amplified using a set of 15 arbitrary oligonucleotide decamer primers. Amplification reaction were in volume of 25.0 µL of sterilized double distilled water, 2.5 µL of 10 X buffer A, 1.0 µL of Dntp (2.5 Mm each), 1.0 µL of primer (1 µM) and 0.3 µL (~ 1.0 U) of Taq polymerase in 4.0 µL of the template DNA. Amplification were performed using 0.2 ml PCR tubes (Axygen Inc., USA) in an Eppendorf master cycler personal programmed for 40 cycles consisting of 30 seconds in 94 °C; one minute in 36 °C and 2.5 minutes in 72 °C denaturing, annealing and extension respectively. These cycles were proceeded by a step for an initial denaturation of five minutes at 94 °C and the cycles followed by a final extension of ten minutes at 72°C. It was then programmed to maintain at 4°C until usage for electrophoresis.

2.4 Electrophoresis of PCR Products and Its Documentation

Amplified products along with the 6X tracking dye were resolved electrophoretically based on their molecular weight by running the products on a 1.2 per cent agarose gel matrix using a submarine electrophoresis with 1X TAE buffer (Sambrook and Russell, 2001) at 5 V cm⁻¹. After the run, the gel was stained using ethidium bromide (Sharp et al., 1973) solution for five to ten minutes and was destained using distilled water for two to three minutes. Ethidium bromide fluoresced orange upon UV illumination (310-320 nm). The gel image was documented using a Herolab documentation unit (Herolab 442 K, E.A.S.Y., Germany).

2.5 Data Analysis

Documented gel was used to score polymorphism by the presence or absence of the bands. Also, monomorphic bands were scored to calculate the percentage of polymorphism.

Table 1. List of mulberry test accessions under *Morus indica* used in this study

Sl. No	Accession Number	Name	Origin
1	MI-0017	Sujanpur – 5	Punjab
2	MI-0395	Tingari	Uttar Pradesh
3	MI-0349	Garobadha – 2	Meghalaya
4	MI-0034	Sujanpur – 1	Punjab
5	MI-0425	Deharadun Local -11	Tamil Nadu
6	MI-0490	Kakathodepaalam	Kerala
7	MI-0652	Pouri – 2	Punjab
8	MI-0535	Araku Local - 2	Himachal Pradesh
9	MI-0543	Palampur Local	Tamil Nadu
10	MI-0536	Ananthagiri Local	Tamil Nadu
11	MI-0558	Kollihills – 2	Uttaranchal
12	MI-0613	Chandrapuri	Jammu and Kashmir
13	MI-0615	Ukhimath	Jammu and Kashmir
14	MI-0651	Rajouri	Jammu and Kashmir
15	MI-0495	Lingamala – 1	Uttar Pradesh
16	MI-0658	Roop Nagar	Himachal Pradesh
17	MI-0677	Durgapur – 3	Rajasthan
18	MI-0685	Khakad – 2	Rajasthan
19	MI-0686	Mount Abu – 1	Kerala
20	MI-0763	Saranath – 2	Tripura
21	MI-0718	Chozhiakadu	Uttar Pradesh
22	MI-0768	Mangari	Uttar Pradesh
23	MI-0663	Chauntra - 2	Rajasthan
24	MI-0845	Rajapur – 2	Tripura
25	V-1 (check variety)	Victory - 1	

Table 2. List of primers used for ISSR amplication

S.No	Primers	Primer sequence 5'-3'
1	UBC 807	AGAGAGAGAGAGAGAGT
2	UBC 809	AGAGAGAGAGAGAGAGG
3	UBC 811	GAGAGAGAGAGAGAGAC
4	UBC 820	GTGTGTGTGTGTGTGTC
5	UBC 821	GTGTGTGTGTGTGTGTT
6	UBC 834	AGAGAGAGAGAGAGAGYT
7	UBC 841	GAGAGAGAGAGAGAGAYC
8	UBC 856	GTGTGTGTGTGTGTGTCTA

2.6 Scoring of Bands

“Clear and unambiguous bands of ISSR markers were scored based on the presence or absence of the corresponding band among the genotypes. The scores ‘1’ and ‘0’ indicated the presence and absence of bands respectively. In case of binary coding, a data matrix computing of ‘1’ and ‘0’ was formed and subjected to cluster analysis. Sequential Agglomerative Hierarchical Non-overlapping (SAHN) clustering was performed on similarity matrix using dice coefficient for the binary data (obtained through scoring) utilizing the Unweighted Pair Group Method with Arithmetic Average (UPGMA) method. This was done by using NTSYSpc version 2.02” (Rohlf, 2002).

“The confidence limits for the groupings by dendrogram were computed by using WINBOOT, a program for performing bootstrap analysis of binary data to determine the confidence limits of UPGMA based dendrograms” (Yap and Nelson, 1996). Boot straping involved repeated sampling with replacement of the characters in matrix of operational taxonomic unit (OTU) characters to create numerous bootstrap matrices of the same size as the original matrix.

2.7 Computation of Similarity Indices

From the binary code matrices obtained from the ISSR markers, the Jaccard's similarity indices were computed for the twenty four test accessions of *Morus indica* using the SIMQUAL programme of NTSYSpc version 2.02 as follows, Jaccard (1908) similarity index.

$$SI_{ij} = a/(n-d)$$

Where,

SI_{ij} = Similarity between *i*th and *j*th samples

a = Number of electromorphs/traits commonly shared

d = Number of electromorphs/traits absent in *i*th and *j*th samples but present in other samples

n = (b+c) i.e. the number of unmatcheds

b = Number of electromorphs/traits present in *i*th samples but absent in *j*th samples

c = Number of electromorphs/traits present in *j*th samples but absent in *i*th samples.

2.8 Polymorphosim Information Content

Polymorphism information content (PIC) or expected heterozygosity scores for each ISSR marker was calculated based on the formula given by Anderson et al. (1993).

$$PIC = 1 - \sum_{j=1}^n P_{ij}^2$$

Where,

P_{ij} is the frequency of *j*th allele for the marker *i* and the summation extends over *n* alleles.

3. RESULTS

The details of the nucleotide sequence of selected primers used to generate PCR products, summary of total number of polymorphic DNA fragments generated and percentage of polymorphism. Out of 8 primers used, only three Primers (UBC 807, UBC 809 and UBC 811) showed polymorphism in twenty four accessions.

3.1 Polymorphism Revealed by ISSR Markers

For ISSR markers, the polymorphic information content values ranged from 0.29 to 0.35. The mean PIC value for polymorphic bands was 0.32 with the highest PIC value of 0.35 with primer UBC 807 and the lowest PIC value of 0.29 with primer UBC 811 (Table 3, Plate 1a, b, c).

3.2 Genetic Variation and Cluster Analysis Based on ISSR Markers

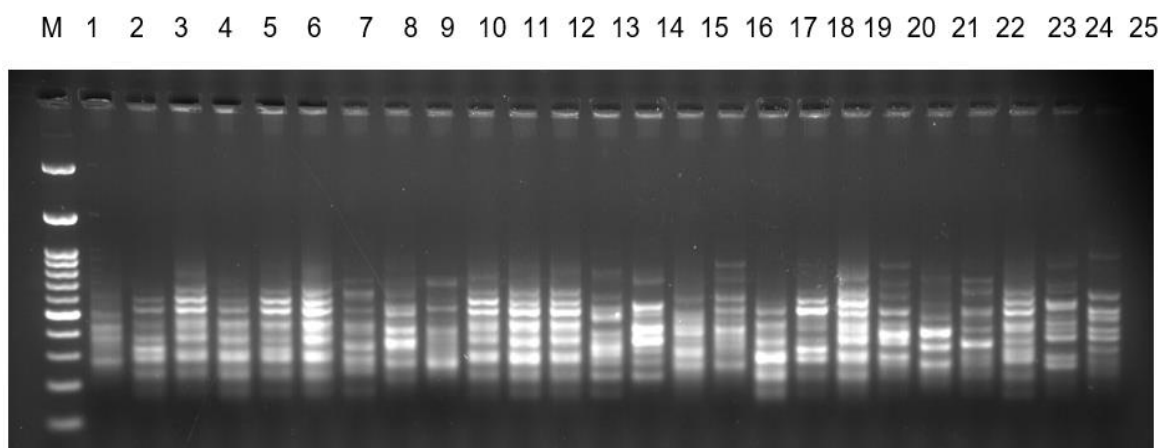
The genetic distance between the analyzed individuals was calculated by simple matching method to obtain a matrix of genetic distance. The Jaccard's co-efficient of similarity values ranged from 0.913 to 0.535 which indicated high level of genetic variation among the genotypes studied. The highest value of similarity coefficient of 0.913 was found between the A18 (MI-0685)

and A8 (MI-0535) and it was followed by similarity coefficient of 0.535 between the genotypes A20 (MI-0763) and A4 (MI-0034) (Table 4, Fig. 1). The average linkage between the genotypes was used for constructing a phylogenetic tree. The UPGMA cluster analysis indicated the presence of distinct major clusters. Among the clusters, cluster II and IV constituted only one accession each. The result provided valid guidelines for collection, conservation and identification of mulberry genetic resources.

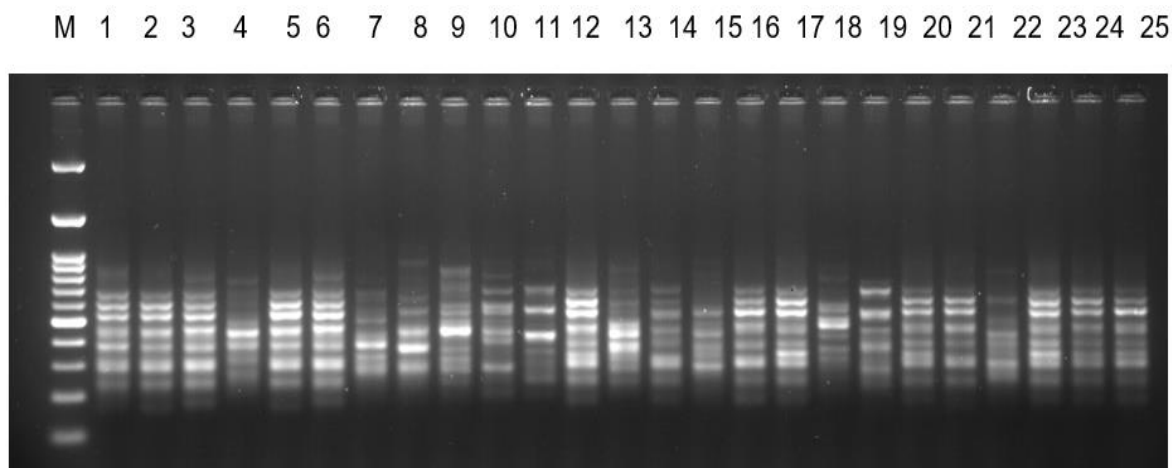
Table 3. Random ISSR primers used for DNA amplification of test mulberry accessions

Sl. No	Primers	Sequence 5' – 3'	Total number of bands	PIC values
1	UBC 807	AGAGAGAGAGAGAGAGT	10	0.35
2	UBC 809	AGAGAGAGAGAGAGAGG	9	0.31
3	UBC 811	GAGAGAGAGAGAGAGAC	7	0.29
Mean				0.32

a. UBC 807 (AGAGAGAGAGAGAGAGT)



b. UBC 809 (AGAGAGAGAGAGAGAGG)



c. UBC 811 (GAGAGAGAGAGAGAC)

M 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23 24 25

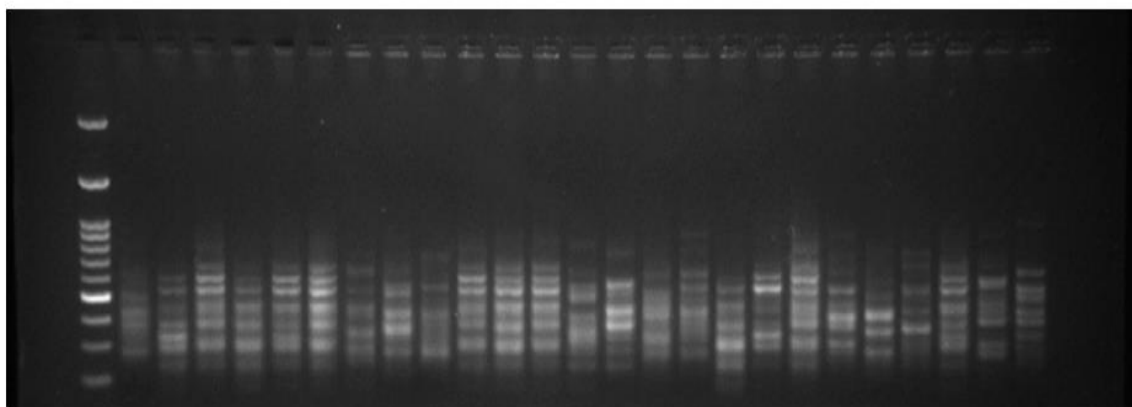


Plate 1. ISSR profile of test mulberry accessions

Table 4. ISSR similarity matrix of test mulberry accessions

Accs	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19	20	21	22	23	24	25
1	1.000																								
2	0.596	1.000																							
3	0.625	0.686	1.000																						
4	0.591	0.568	0.660	1.000																					
5	0.637	0.698	0.921	0.672	1.000																				
6	0.711	0.655	0.824	0.719	0.862	1.000																			
7	0.618	0.615	0.614	0.653	0.627	0.758	1.000																		
8	0.728	0.644	0.750	0.670	0.816	0.885	0.716	1.000																	
9	0.649	0.618	0.796	0.654	0.836	0.783	0.583	0.830	1.000																
10	0.706	0.649	0.758	0.714	0.827	0.898	0.724	0.883	0.779	1.000															
11	0.633	0.576	0.711	0.637	0.779	0.850	0.650	0.866	0.762	0.847	1.000														
12	0.736	0.593	0.728	0.680	0.796	0.898	0.694	0.883	0.810	0.864	0.847	1.000													
13	0.586	0.673	0.792	0.648	0.867	0.810	0.631	0.766	0.781	0.807	0.821	0.775	1.000												
14	0.727	0.576	0.750	0.703	0.789	0.862	0.714	0.847	0.771	0.827	0.810	0.859	0.767	1.000											
15	0.701	0.614	0.694	0.709	0.762	0.864	0.689	0.881	0.716	0.862	0.844	0.862	0.771	0.824	1.000										
16	0.719	0.576	0.741	0.666	0.779	0.850	0.677	0.898	0.824	0.847	0.830	0.847	0.728	0.810	0.844	1.000									
17	0.694	0.637	0.716	0.672	0.763	0.852	0.663	0.900	0.796	0.913	0.833	0.850	0.762	0.783	0.816	0.833	1.000								
18	0.689	0.576	0.741	0.666	0.776	0.650	0.650	0.931	0.824	0.816	0.894	0.847	0.758	0.842	0.877	0.894	0.833	1.000							
19	0.677	0.649	0.789	0.627	0.659	0.636	0.666	0.915	0.842	0.864	0.879	0.833	0.839	0.827	0.830	0.847	0.881	0.879	1.000						
20	0.820	0.584	0.649	0.535	0.644	0.758	0.607	0.793	0.672	0.724	0.736	0.785	0.631	0.732	0.750	0.771	0.771	0.771	1.000						
21	0.696	0.607	0.719	0.566	0.789	0.600	0.627	0.647	0.771	0.796	0.642	0.796	0.767	0.769	0.762	0.779	0.877	0.842	0.892	0.763	1.000				
22	0.578	0.574	0.660	0.641	0.701	0.688	0.596	0.793	0.777	0.683	0.694	0.663	0.709	0.701	0.736	0.724	0.728	0.785	0.771	0.631	0.701	1.000			
23	0.800	0.660	0.706	0.631	0.775	0.879	0.701	0.929	0.789	0.877	0.859	0.910	0.754	0.807	0.842	0.859	0.894	0.859	0.877	0.886	0.824	0.719	1.000		
24	0.678	0.561	0.732	0.625	0.771	0.813	0.666	0.630	0.785	0.842	0.762	0.810	0.750	0.836	0.807	0.824	0.796	0.824	0.842	0.701	0.771	0.684	0.789	1.00	
25	0.636	0.603	0.722	0.641	0.701	0.775	0.716	0.762	0.664	0.741	0.694	0.741	0.620	0.732	0.677	0.754	0.728	0.754	0.711	0.660	0.672	0.586	0.719	0.77	1.00

1	MI-0017	11	MI-0558	21	MI-0718
2	MI-0395	12	MI-0613	22	MI-0768
3	MI-0349	13	MI-0615	23	MI-0663
4	MI-0034	14	MI-0651	24	MI-845
5	MI-0425	15	MI-0495	25	V-1(Check variety)
6	MI-0490	16	MI-0658		
7	MI-0652	17	MI-0677		
8	MI-0535	18	MI-0685		
9	MI-0543	19	MI-0686		
10	MI-0536	20	MI-0763		

4. DISSCUSSION

“Assessment of genetic diversity is a prerequisite for efficient conservation and utilization of genetic resources. During the past two decades, several high-throughput PCR-based technologies such as inter-simple sequence repeats (ISSR) and amplified fragment length polymorphisms (AFLP) have been developed to assay genetic

polymorphism at the DNA level” (Shyamsundar et al., 2014). “An advantage of ISSRs is that in their simplest form, no prior genomic knowledge is required to design arbitrary sequence oligonucleotide primers” (Gonzalez et al., 2005). “With the large array of molecular analytical techniques available, it has become possible to provide an accurate and unambiguous tool for evaluation of genetic diversity and identification

of germplasm" (Simmons et al., 2007) and (Li et al., 2008). "Assessment of genetic variation within the species based on morphological and polypeptide descriptions, provide more speed, accurate and detailed information" (Williams et al., 1990). "DNA marker studies had been successfully employed in plants to understand provenance variation" (Chalmers et al., 1992), geotypic identity (Keil and Griffin 1994), molecular characterization of germplasm (Maechado et al., 1996), molecular systematics (Harris, 1995) genetic diversity (Varghese et al., 1997) and DNA finger printing studies of plants (Caetano-Anolles et al., 1991).

"ISSR has been used for diversity assessment by many researchers in tropical and temperate plants. Extensive studies have already been reported in mulberry" (Karku et al., 1996). ISSR technique was used to study the genetic structure and relationship between species and populations. Similarity, DNA based ISSR techniques have been successfully utilized in many genera and species of mulberry Vijayan et al. (2004a) and (Ahmed sumbul, 2025). The variation in genetic distance among the different mulberry accessions revealed by ISSR tech-niques reflected a high level of polymorphism at the DNA level. Earlier studies by Sharma et al. (2000) using the AFLP tech-nique also showed a

large genetic variation among different *Morus* genotypes.

In the present study, mulberry accessions showed 25 reproducible bands from 3 primers. The primers produced 100 per cent polymorphism (UBC 807, UBC 809 and UBC 811). The highest PIC values were observed with the primer UBC 807 and the lowest PIC values was observed with primer UBC 811. Polymorphic information content were documented by (Emine orhan et al. 2024). "The present investigation revealed that mulberry accessions used for diversity through ISSR, showed moderate level of genetic variations. Polymorphism in mulberry accessions were documented" (Girish et al., 2015) and (Miran A et al., 2023). Similar findings were noticed by (Keshavamurthy et al., 2013) among 26 diverse germplasm of mulberry collected from germplasm bank at Bangalore University, India. The overall percentage of polymorphism was found to be 55.83 per cent by ISSR marker. Among the germplasm, S 36, V1, S54 and M5 showed close similarity. Zhao et al. (2007) further reported a higher value of 0.4912 within cultivated species of mulberry for the same. Aggarwal et al. (2004) also reported "a higher Ho value of 0.59 across 45 mulberry genotypes from species of diverse origin using SSR markers".

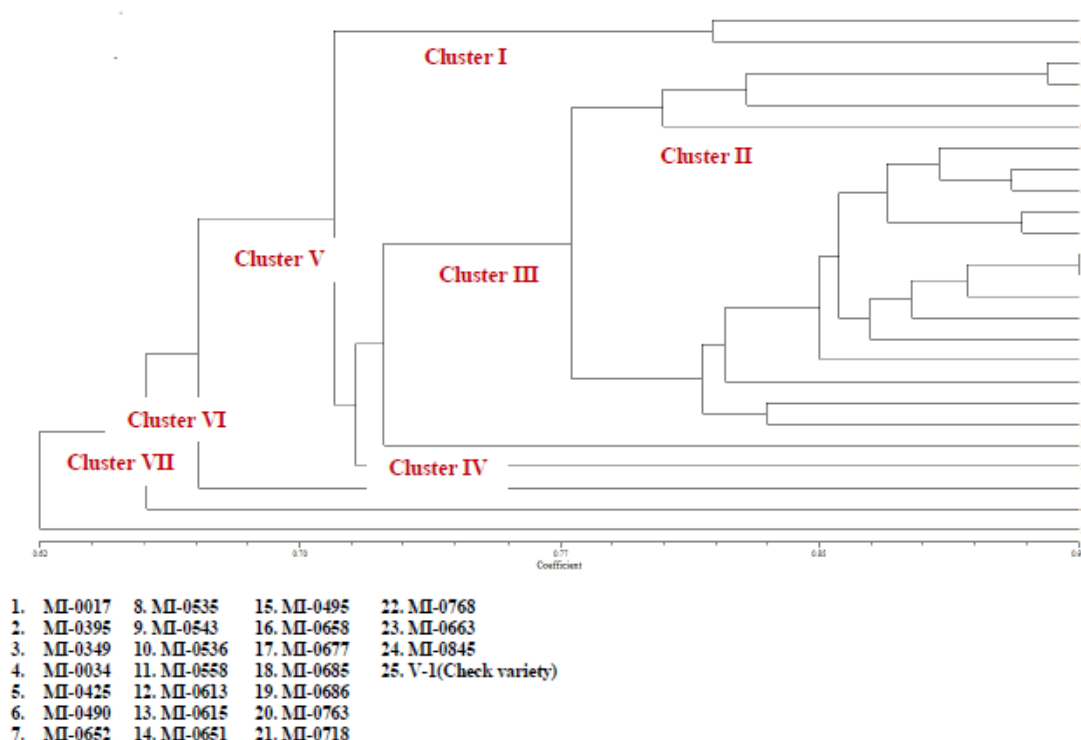


Fig. 1. Dendrogram of test mulberry accessions based on jaccard's similarity coefficient

In mulberry accessions, the average linkage between the accessions was used for constructing a phylogenetic tree. The association amongst different mulberry accessions was presented in the form of dendrogram. The accessions A18 (MI-0685) and accession A8 (MI-0535), were closest at the similarity coefficient 1.000. The accessions A20 (MI-0763) and A4 (MI-0034) were the farthest or most diverse and dissimilar to all others. Similar grouping pattern within species and related genotypes are evident from the earlier report in mulberry (Keshavamurthy et al., 2013) and (Girish et al., 2015). Koidzumi (1973) considered that *M. lhou*, *M. multicaulis* and *M. latifolia* are similar and belong to a single species. The percent polymorphic loci revealed an average of 74.47% in all the mulberry accessions studied using SSR markers. This value was lower than 85 and 91% shown in RAPD and DAMD markers by Bhattacharyya and Ranade (2001). Earlier, Sharma et al. (2000) had reported a higher average value of 81.2%. Additionally, a higher value of 86% was reported by Vijayan et al. (2004a) and Zhao et al. (2007). "These values suggest a high level of genetic variability among the accessions studied. Such high level of genetic diversity can be attributed to the mode of reproduction of these accessions. Vegetatively propagated species such as mulberry often have minute unobserved changes in passage of time and accumulation of these changes can lead to significant changes in plant population. Such variability could come from mutation" (Vijayan, 2009) and/or long cultivation periods (Bhattacharyya and Ranade 2001). "Mulberry is a highly heterozygous and outcrossing species and therefore it is expectable that its accessions exhibit a high level of polymorphism" (Awasthi et al., 2004).

5. CONCLUSION

The results obtained by ISSR analysis of mulberry accessions were in general agreement with our morphological classification, suggesting that ISSR is a simple and effective molecular marker technique and could be successfully applied to the study of genetic relationships, to the conservation and identification of mulberry collections, and to plant breeding. Our results also suggest that molecular tools should be used for the classification of the genus *Morus*.

DISCLAIMER (ARTIFICIAL INTELLIGENCE)

Author(s) hereby declare that NO generative AI technologies such as Large Language Models

(ChatGPT, COPILOT, etc.) and text-to-image generators have been used during the writing or editing of this manuscript.

COMPETING INTERESTS

Authors have declared that no competing interests exist.

REFERENCES

- Aggarwal, Udayakumar, D., Hendre, P., Sarkar, A., Singh and Li. (2004). Isolation and Characterization of six novel microsatellite markers for mulberry (*Morus indica*). *Molecular Ecology*, 4, 477-479.
- Ahmed Sumbul (2025). Characterization of Genetic Diversity of Mulberry (*Morus alba*) Genotypes Growing Naturally in Northeastern Türkiye (Kelkit Valley) Using Morphological, Biochemical, and Molecular Markers. Nature based solutions to improve the sustainability of Horticulture Ecosystem. 11(3), 298.
- Anderson, Churchill, G., Autrique, J., Tanksely, S and Sorrells, M. (1993). Optimizing parental selection for genetic linkage maps. *Genome*, 36, 181-186.
- Awasthi, Nagaraja, G., Naik, G., Kanginakudru, S., Thangavelu, K and Nagaraja, J. (2004). Genetic diversity in mulberry (Genus *Morus*) as revealed by RAPD and ISSR marker assays,. *BMC Genetics*, 5, 1.
- Basaviah, Dandin, S and Rajan, M. (1989). Microsporogenesis in hexaploid *Morus serrata* Rxb. *Cytologia*, 54, 747-751.
- Bhattacharyya and Ranade, S. (2001). Molecular distinction amongst varieties of mulberry using RAPD and DAMD profiles *BMC Plant Biol.* , 1(3), 1471-2279.
- Emine Orhan, Meleksen Akin, Sadiye P. Eydurhan and Sezai Ercisli. (2024). Molecular characterization of mulberry genotypes and species in Turkey. *Notulae Botanicae Horti Agrobotanici Cluj-Napoca.*, 48(2), 579-557.
- Girish, Dandin, S., Tikader , A and Pinto, M. V. (2015). Molecular diversity of wild mulberry (*Morus* spp) of Indian subcontinent. *Indian journal of Biotechnology*, 14, 334-343.
- Gonzalez, Wong, A., Delgado-Salinas, A., Papa, R and Grepts, P. (2005). Assessment of inter simple sequence repeat markers to differentiate sympatric wild and domesticated populations of common bean. *Crop Science*, 45(2), 606-615.

- Haines and Martin, B. (1995). Biotechnologie et production soutenue des bois tropicaux. Etude sur documents effectuée pour l'OIBT. *Council International des Bois Tropicaux*, Janvier., 20-23.
- Jaccard. (1908). Nouvelles recherches sur la distribution florale. *Bull. Soc. Vaud. Nat*, 44, 223-270.
- Karku, Hume, M., Karjalainen, P., Karvonen, K., Karkkainen, K., Nedeand, D and Savolainen, O. (1996). *Do molecular markers reflect pattern of differentiation in adaptive traits of conifers. Theor. Appl. Genet.*
- Keshavamurthy, Prakash, B. M., Hittalmani, S and Puttaraju, H. P. (2013). DNA marker-assisted evaluation of cultivated and local mulberry genotypes of southern India. *Crop Breeding and Applied Biotechnology*, 9, 239-245.
- Koidzumi. (1973). Synopsis specierum generis Mori. *Bull Imp SericultExp Stat*, 11, 1-50.
- Miran A. El- Haggag, Yasmin A.Mahagoub, Hesham. M Aly, Nabila. M. Ghazy, Fathy K. El-Fiky, Amr M. El- Hawiet (2023). DNA barcodes, ISSR, RAPD and SCAR markers as potential quality control tools for molecular authentication of black and white mulberry. *Food control*, vol 152.
- Rohlf. (2002). *NTSYS-pc: Numerical taxonomy system ver.2.1. Exerter Publishing Ltd., Setauket, New york.*
- Sambrook and Russell, D. (2001). Molecular cloning. *A laboratory manual 3/e (Vol.I to III). Cold Spring Harbor Laboratory Press, Cold Spring Harbor, New York.*
- Sharma, Sharma, R and Machii, H. (2000). Assessment of genetic diversity in a *Morus* spp germplasm collection using fluorescence-based AFLP markers. *Theor. Appl. Genet*, 101, 1049-1055.
- Sharp, Sugden, B and Sambrook, J. (1973). Detection of two restriction endonuclease activities in *Haemophilus influenzae* using analytical agarose-ethidium bromide electrophoresis. *Biochemistry*, 12, 3055-3063.
- Veerapuramarayanappa, Murthy, Y., Ramesh, H. L., Lokesh, G., Munirajappa and Yadav, R. D. (2013). Assessment of six mulberry (*Morus*) germplasm varieties through moulting and bioassay parameters by using cross breed silkworms *Bombyx mori* L. for commercial exploitation in Kolar district, Karnataka. *Int. Res. J. Biological Sci.*, 2(9), 69-75.
- Vijayan, Kar, P., Tikader, A., Srivastava, P., Awasthi, A., Thangavelu, K and Saratchandra, B. a. P. B.-. (2004a). Molecular evaluation of genetic variability in wild populations of mulberry (*Morus serrata* Roxb.), *Plant Breeding*, 123, 568-572.
- Vijayan. (2009). Approaches for enhancing salt tolerance in mulberry (*Morus* L). *Plant Omics J*, 2(1), 41-59.
- Williams, Kubelik, A., Livak, K., Rafalski, J and Tingey, S. (1990). *DNA polymorphisms amplified by arbitrary primers are useful as genetic markers*. *Nucleic Acids Res.*
- Yap and Nelson, R. (1996). *WINBOOT: a program for performing bootstrap analysis for binary data to determine the confidence limits of UPGMA based dendrograms*. . IRRI Disc. Ser. No. 14. Int. Rice Res. Inst., Manila, Phillippines.
- Zhao, Zhou, Z., Miao, X., Zhang, Y., Wang, S., Huang, J and Huang, Y. (2007). A comparison of genetic variation among wild and cultivated *Morus* species (*Moraceae: Morus*) as revealed by ISSR and SSR markers. *Biodivers. Conserv*, 16, 275-290.

Disclaimer/Publisher's Note: The statements, opinions and data contained in all publications are solely those of the individual author(s) and contributor(s) and not of the publisher and/or the editor(s). This publisher and/or the editor(s) disclaim responsibility for any injury to people or property resulting from any ideas, methods, instructions or products referred to in the content.

© Copyright (2025): Author(s). The licensee is the journal publisher. This is an Open Access article distributed under the terms of the Creative Commons Attribution License (<http://creativecommons.org/licenses/by/4.0>), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

Peer-review history:
The peer review history for this paper can be accessed here:
<https://pr.sdiarticle5.com/review-history/142867>