

Genetic Divergence Studies in *Ailanthus excelsa* Using D^2 Analysis

ABSTRACT

Aims: To estimate the genetic diversity studies among the biometric attributes of 30 progenies in *Ailanthus excelsa* Roxb.

Place and Duration of Study: The study has conducted at Forest College and Research Institute, TNAU, Mettupalayam during 2015-2018.

Methodology: The D^2 statistics was adopted for the estimation of genetic divergence. Using D^2 statistical results, the clustering of progenies was done. The progenies were grouped into different clusters using 'GENERES' statistical package on the basis of D^2 values according to Tocher's method as suggested by Rao

Results: The 30 progeny of *Ailanthus excelsa* has grouped into nine clusters and among the nine clusters, the cluster IV has ten progenies. The maximum intra cluster distance was exhibited by the cluster VIII followed by cluster IV. The maximum inter cluster distance was in cluster III which indicated the presence of wider genetic distance between *Ailanthus excelsa* progenies. Among the growth attributes, volume index contributed maximum percentage towards genetic divergence.

Conclusion: The results of 30 progeny of *Ailanthus excelsa* showed the presence of wider genetic distance between *Ailanthus excelsa* progenies.

11
12
13
14
15
16
17
18
19
20
21
22
23
24
25
26
27
28
29
30
31

Keywords: *Ailanthus excelsa*, Biometric attributes, Genetic resources, Diversity, Genetic distance, D^2 clustering

1. INTRODUCTION

Ailanthus excelsa Roxb. is a tree belonging to family Simaroubaceae, indigenous to Central and Southern India and commonly it is known as Tree of Heaven. It is a large deciduous tree and will be growing 18-25 m tall with straight trunk and 60 to 80 cm in diameter. It is mainly used for making plywood as well as match splint production [1]. Due to the demand of both plywood and match wood this study has conceived. Rapid socio-economic changes are having profound impacts on all sectors including forestry. Societal transformations are changing people's perceptions of forests, while growing and often conflicting demands for forest-derived goods and services have increased the complexity of forest management. Concerns over climate change, escalating energy prices and deepening water deficits have moved forestry into the spotlight of global and national development. Currently, the forest area in the country is around 23.81 per cent and in the state of Tamil Nadu it is around 17.59 per cent which is much low against the demanded requirement of 33.0 per cent. The productivity in terms of MAI is also one of the lowest comparing to the global average [2]. The annual estimated production of wood from forest is estimated to be 3.173 million m^3 and the annual potential production of wood from outside the forests is

32 estimated to be 42.77 million m³ [2]. The country's timber imports value is growing at 12 per
33 cent per annum and is likely to increase in years ahead. The liberalization of imports has
34 benefited the domestic timber market, otherwise faced paucity of the desired wood in the
35 required quantity and quality. However, there is a potential to increase the domestic
36 production of industrial wood through tree planting, afforestation and reforestation
37 programmes [3].

38 Hence shrinking forest area associated with low productivity established a total
39 mismatch between the demand and supply of both domestic and industrial wood
40 requirement besides creating environmental disequilibrium [4]. The current supply of raw
41 materials for industries like match wood, pulpwood, plywood, furniture and biomass energy
42 in India particularly in Tamil Nadu is far behind the demand. Hence, to meet the growing raw
43 material demand and also to meet the National Forest Policy (1988). Guidelines, the
44 industries must expand sharply its plantation programme. There are over 400 small-scale
45 sector Splints and Veneer Industry involved in the manufacturing of veneers and splints in
46 southern India of which 75% are located in Kerala [5]. Per capita consumption of matches in
47 India increased steadily from 2.45 sticks per capita in 1970 to 8.35 in 2013. There are wide
48 fluctuations in the annual growth rate in the consumption of matches varying from as low as
49 3 per cent (before 1970) to as high as 28 per cent. The rising levels of income, growing
50 urbanization, swelling numbers of smokers, and changes in fuel consumption patterns
51 indicates that the future rate of growth could be higher than the 6 per cent as supported by
52 past trends [6].

53 The major raw materials used in the production of safety matches are soft woods.
54 Safety matches manufactured in India are of the standard type with wooden veneer or
55 cardboard boxes and wooden splints. Historically the Indian match industry depended on
56 imported wood including Aspen (*Populus tremula*) from Sweden, Canada, America, and
57 Russia; Cotton Wood (*Populus deltoides*) from Canada; Balsam Poplar (*Populus
58 balsamifera*) from Manchuria; and Linden (*Tilia japonica*) from Japan. But the government
59 quickly moved to encourage the use of indigenous woods by restricting the import. Even
60 though there are number of alternative match wood species are available to replace the
61 imported wood, *Ailanthus excelsa* occupies predominant position because of its suitability for
62 the production quality match splints. However there is no systematic evaluation or
63 improvement programme in order to utilize the existing genetic variation among broader
64 genetic base population which warrants a systematic tree improvement programme in
65 *Ailanthus excelsa* which will also address the shortage of suitable raw material to the match
66 industries.

67

68 **2. MATERIALS AND METHODS**

69

70 **A. MATERIALS**

71 The species *Ailanthus excelsa* was chosen as the experimental material for the
72 present study which consists of 30 progenies established as a progeny evaluation trial.

73

73 **B. METHODS**

74 ***Estimation of Morphometric attributes***

75

75 ***Source of progenies***

76

76 The predominant eleven *Ailanthus excelsa* distributed districts of Tamil Nadu viz.,
77 Coimbatore, Tirupur, Erode, Salem, Theni, Dindugal, Viruthunagar, Darmapuri, Krishnagiri,

78 Villupuram, and Karur were surveyed and a total number of 30 candidate plus trees were
 79 selected. These selected Candidate Plus Trees (CPTs) were given with the accession number
 80 as FCRI AE. The details on the actual locations of the 30 selected candidate plus trees were
 81 presented in table 1.

82 **Table 1. Details of *Ailanthus excelsa* genetic resources and their location**

Sl. No.	District	Sources	Name of sources	Latitude	Longitude
1	Coimbatore	Akkarai sengapalli	FCRI AE 1	11 ⁰ 19'28"N	77 ⁰ 04'53"E
2	Coimbatore	S. Pungampalayam	FCRI AE 2	11 ⁰ 03'24"N	77 ⁰ 19'51"E
3	Coimbatore	Cherannagar – 1	FCRI AE 3	11 ⁰ 03'05"N	76 ⁰ 56'32"E
4	Coimbatore	Cherannagar – 2	FCRI AE 4	11 ⁰ 03'05"N	76 ⁰ 56'32"E
5	Coimbatore	Teachers colony	FCRI AE 5	11 ⁰ 09'37"N	76 ⁰ 56'33"E
6	Coimbatore	Annur – 1	FCRI AE 6	11 ⁰ 14'03"N	77 ⁰ 06'19"E
7	Coimbatore	Annur – 2	FCRI AE 7	11 ⁰ 14'03"N	77 ⁰ 06'19"E
8	Coimbatore	Alamelu mangapuram	FCRI AE 8	11 ⁰ 02'45"N	76 ⁰ 58'40"E
9	Coimbatore	Vaikalpalam	FCRI AE 9	10 ⁰ 58'53"N	76 ⁰ 55'17"E
10	Tirupur	Pogalur	FCRI AE 10	11 ⁰ 15'25"N	77 ⁰ 02'26"E
11	Tirupur	Samundipuram	FCRI AE 11	11 ⁰ 07'28"N	77 ⁰ 18'60"E
12	Tirupur	Kulathu thottam	FCRI AE 12	11 ⁰ 03'33"N	77 ⁰ 15'56"E
13	Tirupur	Salakkudi	FCRI AE 13	10 ⁰ 41'04"N	77 ⁰ 36'22"E
14	Tirupur	Chettipalayam	FCRI AE 14	11 ⁰ 08'38"N	77 ⁰ 20'28"E
15	Erode	Appachimar madam	FCRI AE 15	11 ⁰ 19'51"N	77 ⁰ 28'47"E
16	Erode	Perundurai	FCRI AE 16	11 ⁰ 16'26"N	77 ⁰ 35'18"E
17	Salem	Pethanayakkanpalayam	FCRI AE 17	11 ⁰ 38'51"N	78 ⁰ 30'20"E
18	Salem	Idapadi	FCRI AE 18	11 ⁰ 35'05"N	77 ⁰ 50'20"E
19	Theni	Uthamapalayam	FCRI AE 19	9 ⁰ 48'20"N	77 ⁰ 19'40"E
20	Theni	Thevaram	FCRI AE 20	9 ⁰ 53'44"N	77 ⁰ 16'31"E
21	Theni	Bodi	FCRI AE 21	10 ⁰ 01'00"N	77 ⁰ 21'00"E
22	Dindugal	Kallimandayam	FCRI AE 22	10 ⁰ 35'28"N	77 ⁰ 44'11"E
23	Viruthunagar	Srivilliputhur	FCRI AE 23	9 ⁰ 30'44"N	77 ⁰ 38'03"E
24	Darmapuri	Harur	FCRI AE 24	12 ⁰ 03'05"N	78 ⁰ 28'49"E
25	Darmapuri	Papparettipatti	FCRI AE 25	11 ⁰ 54'49"N	78 ⁰ 21'57"E
26	Krishnagiri	Oothangarai	FCRI AE 26	12 ⁰ 15'57"N	78 ⁰ 32'07"E
27	Villupuram	Thiruvakkarai	FCRI AE 27	12 ⁰ 01'34"N	79 ⁰ 39'06"E
28	Villupuram	Mathangadipattu	FCRI AE 28	11 ⁰ 57'59"N	78 ⁰ 45'28"E
29	Villupuram	Pudupattu	FCRI AE 29	11 ⁰ 58'21"N	78 ⁰ 53'52"E
30	Karur	Salikaraiipatti	FCRI AE 30	10 ⁰ 45'04"N	78 ⁰ 10'70"E

83
 84
 85
 86
 87
 88
 89
 90
 91
 92
 93
 94

95
96
97
98
99
100
101
102
103
104
105
106
107
108
109
110
111
112

Determination of genetic diversity

The data recorded at 6 MAP in *Ailantus excelsa* progeny evaluation trial were used for diversity analysis.

Determination of genetic divergence

The D^2 statistics was adopted for the estimation of genetic divergence [7]. Using D^2 statistical results, the clustering of progenies was done.

D^2 statistics

The D^2 statistics was carried out using the traits viz., plant height, diameter at breast height and volume. The mean squares and the mean products were estimated between groups and within components by one-way analysis of variance, covariance and the significance were tested at progeny level. A variance – covariance was formed from the above and subjected to pivotal condensation to obtain the linear function for transformation of character mean values (x) to a set of independent variables (uncorrected mean) value (y).

The difference between any two mean values for each pair of progeny was squared and added to give the D^2 values. For each character Cumulative D^2 values in all the possible combination of progeny were estimated.

$$y_1 = x_1$$

$$y_2 = x_2 - a_2 x_1$$

$$y_3 = x_3 - a_{32} y_2 - a_{31} y_1$$

$$y_p = x_p - a_{pp-1} y_{p-1} \dots a_{p1} y_1$$

130 where,

131 x_1 = normalized variables

$$a_{ij} = b_{ij}/v(y_j) S < -1$$

$$v(y_j) = \lambda \sum a_{(ij)} b_{ij} - b_{ij} = \lambda_{ij} - 1/atbt$$

$$\lambda_{ij} = \text{Covariance of } i \text{ and } j^t = j^i$$

132 All possible $\frac{n(n-1)}{2} D^2$ values were calculated by taking sum of difference between pair
 133 of corresponding 'y' values taking two progenies at a time.

134 **Determination of clusters or grouping**

135 The progenies were grouped into different clusters using 'GENERES' statistical
 136 package on the basis of D^2 values according to Tocher's method as suggested by Rao [8].

137

138 **Tocher's method**

139 All the $\frac{n(n-1)}{2} D^2$ values were clustered by using Tocher's method [8].

140

141 **Average intra and inter cluster distances**

142 On completion of clustering, the intra and inter cluster relationships were studied
 143 and the mutual relationship between clusters and their distances were represented. The
 144 average intra cluster distances was measured using the formula

$$D^2 = D^2/n$$

145 Where D^2 was the sum of distances between all possible combinations of the
 146 progeny included in a cluster whereas the average inter cluster divergences were arrived at
 147 by taking into consideration of all the component D^2 values possible among the numbers of
 148 the two clusters. Then the genetic distance 'D' between the clusters were obtained from
 149 square root of the average D^2 values.

150

151 **3. RESULTS AND DISCUSSION**

152

153 Observations on morphometric traits viz., survival percentage, plant height, basal
 154 diameter, number of branches and volume index and biochemical attributes viz., chlorophyll
 155 'a', chlorophyll 'b', total chlorophyll and chlorophyll a / b ratio were recorded in 30 progenies
 156 of *Ailanthus excelsa*. The morphometric traits were measured at four growth periods
 157 viz., initial, 2 MAP, 4 MAP and 6 MAP whereas biochemical attributes were recorded only 6
 158 MAP. The data were subjected to genetic diversity analysis and the results were presented
 159 below.

160 **Genetic divergence**

161 The genetic divergence was analyzed using multivariate analysis among the 30
 162 progenies with computer based "GENRES" statistical package. The D^2 were computed for all
 163 positive pairs. The morphometric traits viz., plant height, basal diameter, number of branches
 164 and volume index were used for divergence and clustering analysis. The 30 progenies of
 165 *Ailanthus excelsa* were resolved into nine genetically distinct clusters.

166 **Intra and inter cluster average distance**

167 The average intra and inter cluster values among the nine clusters are presented in
 168 Table 2. The progenies resolved within the intra cluster VIII has high genetic distance of
 169 13.78, while the least genetic distance of 0.21 was observed in the cluster III. The highest
 170 inter cluster genetic distance was recorded between the cluster III and IX (80.88). The
 171 minimum inter cluster genetic distance was recorded between the cluster I and V (4.56).

172

173 **Table 2.** Inter (diagonal) and intra cluster estimates of *Ailanthus excelsa* progenies based on
 174 morphometric attributes.

Cluster	1	2	3	4	5	6	7	8	9
---------	---	---	---	---	---	---	---	---	---

I	1.12 (1.06)	15.94 (3.99)	63.18 (7.94)	5.90 (2.43)	4.56 (2.13)	8.89 (2.98)	6.17 (2.48)	9.02 (3.00)	11.72 (3.42)
II		7.67 (2.77)	32.96 (5.74)	15.70 (3.96)	6.48 (2.54)	33.99 (5.83)	11.30 (3.36)	13.45 (3.66)	42.83 (6.54)
III			0.21 (0.46)	59.09 (7.68)	44.69 (6.68)	77.24 (8.78)	50.85 (7.13)	51.88 (7.20)	80.88 (8.99)
IV				10.87 (3.29)	6.55 (2.55)	16.60 (4.07)	9.46 (3.07)	11.08 (3.33)	20.74 (4.55)
V					0.60 (0.77)	18.29 (4.27)	4.38 (2.09)	4.91 (2.21)	25.72 (5.07)
VI						8.83 (2.97)	13.56 (3.68)	22.61 (4.75)	8.72 (2.95)
VII							2.56 (1.60)	11.58 (3.40)	25.63 (5.06)
VIII								13.78 (3.71)	28.19 (5.31)
IX									0.00 (0.00)

175

176 Plant diversity is a variety and variability of a plant in an ecosystem
 177 [9]. Most forest trees are long lived, out breeding and generally highly heterozygous, which
 178 have developed a number of natural mechanisms to maintain heterozygosity and *intra*
 179 specific variations. These genetic mechanisms combined with the often variable
 180 environment, in which forest trees occur, have contributed to the fact that, with a few
 181 exceptions, forest trees seem to be among the most genetically variable of all organisms
 182 studied to date [10].The extent and pattern of genetic diversity in forest trees are influenced
 183 by their native system and the movement of genes between dispersed populations of the
 184 same species. Measuring genetic diversity in trees has typically been done by either
 185 provenance testing [9] or electrophoresis analysis of the enzymes [11] and [12] and also by
 186 DNA based molecular techniques [13][14].In the current study, genetic diversity existed
 187 among the 30 selected genotypes of *Ailanthus excelsa* had been assessed through D²
 188 analysis which resolved the 30 progenies into nine clusters.

189

190 **Cluster components**

191 The multivariate analysis grouped 30 progenies into nine clusters. The cluster
 192 members and number of progenies constituting each cluster are furnished in Table 3.
 193 Among the nine clusters, the cluster IV resolved with ten progenies *viz.*, FCRI AE 7, FCRI
 194 AE 8, FCRI AE 11, FCRI AE 12, FCRI AE 13, FCRI AE 15, FCRI AE 17, FCRI AE 18, FCRI
 195 AE 19, and FCRIAE 20. Whereas, Cluster II had five progenies *viz.*, FCRI AE 2, FCRI AE 3,
 196 FCRI AE 4, FCRI AE 10and FCRI AE 14 and Cluster I and VI constituted only three
 197 progenies each (FCRI AE 1, FCRI AE 4, FCRI AE 9and FCRI AE 21, FCRI AE 22, FCRI AE
 198 24).The cluster III, cluster VII and cluster VIII had two progenies *viz.*, FCRI AE 6, and FCRI
 199 AE 16; FCRI AE 23 and FCRIAE 29 and FCRIAE 26 and FCRI AE 28 respectively. The
 200 cluster IX consisted only one progeny (FCRI AE 25).

201

202 **Table 3.** Clustering pattern of *Ailanthus excelsa* progenies for morphometric attributes.

Cluster No	Number of progenies	Members
I	3	FCRI AE 1, FCRI AE 4, FCRI AE 9
II	5	FCRI AE 2, FCRI AE 3, FCRI AE 5, FCRI AE 10, FCRI AE 14
III	2	FCRI AE 6, FCRI AE 16
IV	10	FCRI AE 7, FCRI AE 8, FCRI AE 11, FCRI AE 12, FCRI AE 13, FCRI AE 15, FCRI AE 17, FCRI AE 18, FCRI AE 19, FCRI AE 20
V	2	FCRI AE 27, FCRI AE 30
VI	3	FCRI AE 21, FCRI AE 22, FCRI AE 24
VII	2	FCRI AE 23, FCRI AE 29
VIII	2	FCRI AE 26, FCRI AE 28
IX	1	FCRI AE 25

203

204 The cluster mean for different biometric traits was estimated and furnished in the
 205 Table 4. The maximum cluster mean for plant height (69.73 cm) was observed in cluster III,
 206 whereas, the least cluster mean for plant height (38.67 cm) was exhibited by the cluster IX.
 207 The highest performance in basal diameter was exhibited by the cluster III which accounts
 208 3.88 cm followed by cluster VIII (3.17 cm) whereas, the minimum was observed for the
 209 cluster IX (1.65 cm) and in no. of branches the maximum was observed in cluster VI (0.85)
 210 and minimum found in cluster VIII (0.00). In case of volume index, the cluster mean was
 211 highest for cluster III (1056.78) and the lowest was exhibited by the cluster IX (105.68).

212

213

214

Table 4. Cluster mean values of *Ailanthus excelsa* progenies for morphometric attributes.

Cluster	Plant height (cm)	Basal diameter (cm)	No. of Branches	Volume index (cm³)
I	45.30	2.27	0.22	238.06
II	58.45	3.05	0.02	561.59
III	69.73	3.88	0.33	1056.78
IV	47.52	2.51	0.14	320.80
V	50.31	2.92	0.11	454.30
VI	40.42	2.10	0.85	183.56
VII	50.79	2.69	0.66	377.56
VIII	46.22	3.17	0.00	503.58
IX	38.67	1.65	0.44	105.68

215

216 Clustering methods have the goal of separating a pool of observations in many
217 subgroups to obtain homogeneity within and between the formed subgroups. D² statistics is
218 an important tool in plant breeding for estimating genetic divergence [15]. The exploitation of
219 heterosis and success in getting desirable segregates in a breeding programme largely
220 depends on the degree of divergence in a chosen population [16]. Genetic diversity is
221 essential to meet the diversified goals of tree breeding such as breeding for cultivation,
222 increasing yield, wider adaptation, desirable quality, pest and disease resistance. The
223 genetic divergence analysis estimates the extent of diversity existed among selected
224 genotypes [17].

225 The application of D² clustering technique in *Ailanthus excelsa* resolved the thirty
226 genotypes into nine clusters. Among the nine clusters, the clusters IV were the biggest with
227 ten progenies. Similarly, earlier studies in *Ailanthus excelsa*, 30 progenies were grouped into
228 eight clusters, of which group A formed the largest cluster containing ten progenies followed
229 by group B with five progenies [18]. In *Acacia nilotica* also by D² clustering technique, 27
230 seed sources were grouped into five clusters (A, B, C, D and E) which showed that group A
231 was the largest in size and possessed 21 seed sources. Group B and C included two seed
232 sources each and Group D and E included only one seed source each [19]. Similarly, 80
233 batches of teak had been grouped into eight clusters, of which group A formed the largest
234 cluster containing 46 batches [20].

235 In the present investigation, it could be seen that the progenies from different
236 locations got clubbed together to form a single major cluster as evident in cluster IV and
237 therefore the pattern of divergence was not depend on the geographic locations. The above
238 findings also confirmed the earlier report of Bagchi [20] in Teak; *Eucalyptus*[21]; *Leucaena*
239 *leucocephala*[22] and *Melia dubia*[18] and [23]. The inclusion of geographically divergent
240 provenances in the same cluster may be attributed to the fact that the factors other than
241 geographic distribution might be responsible for their genetic similarity [24]. Hence the
242 divergent progenies used in the current project and grouped under one cluster might be due
243 to the factor other than the geographical distribution as evidenced in *Santalum album*[25]
244 and *Prunus armeniaca*[26] which supported the results of this work.

245 The intra and intercluster analysis indicated that the cluster IX showed that there is
246 no intra cluster generalized distance since it contained only one progeny. The maximum
247 intra cluster distance was shown by the cluster VIII. The maximum inter cluster distance was
248 recorded between cluster III and II which indicated the presence of wider genetic distance
249 between *A. excelsa* progenies. Such inter and intra cluster distance among *Pinus gerardiana*
250 genotypes was also evidenced which support the current conclusion [27].

251 **Contribution of characters towards genetic divergence**

252 The number of times each character ranking first was counted and percentage
253 contribution towards divergence was calculated and presented in Table 5. Volume index
254 contributed maximum percentage towards divergence (50.34 %) followed by plant height
255 (30.11 %) and number of branches (10.34 %). The minimum percentage contribution
256 towards divergence was recorded by basal diameter (9.19 %).

257
258 **Table 5.** Percentage contributions of morphometric traits of *Ailanthus excelsa* progenies to
259 genetic divergence.

S.No	Character	No. of first rank	% Contribution
------	-----------	-------------------	----------------

1.	Plant height	131	30.11
2.	Basal diameter	40	9.19
3.	No. of branches	45	10.34
4.	Volume index	219	50.34
Total		435	100.00

260
261
262
263
264
265
266
267
268
269
270
271

Volume index contributed maximum towards genetic divergence followed by plant height and the minimum by basal diameter. Paramathma [28] reported similar results in six Eucalyptus species and twelve interspecific hybrids; Bagchi [20] in *Tectona grandis*; Manga and Sen [29] in *Prosopis cineraria*; Tewari *et al.*[30] in *Dalbergia sissoo*; Chauhan and Sehgal [31] in *P. roxburghii* and Vennila [21] in Eucalyptus also reported contribution of volume index along with other morphometric traits towards genetic divergence among the genotypes tested which might be due to the existence of broader genetic base. Kumar [18] also reported similar results in *Ailanthus excelsa* genetic resources. Based on the past work and present finding, the contributions of volume for genetic divergence indicated that this factor could be used as an index for *Ailanthus excelsa* tree improvement programme.

272 4. CONCLUSION

273
274
275
276
277
278
279
280
281
282

The multivariate analysis grouped 30 progeny of *Ailanthus excelsa* genetic resources into nine clusters. Among the nine clusters, the cluster IV has ten progenies. The maximum intra cluster distance was exhibited by the cluster VIII followed by cluster IV. The maximum inter cluster distance was in cluster III which indicated the presence of wider genetic distance between *Ailanthus excelsa* progenies. Among the growth attributes, volume index contributed maximum percentage towards genetic divergence followed by plant height and number of branches while the basal diameter recorded minimal contribution to the divergence.

283 REFERENCES

- 284
285
286[1] Orwa C, Mutua A, Kindt R , Jammadass R, and Anthony S. Agroforestry Database: a
287 tree reference and selection guide version 4.0. 2009.
288[2] Forest Survey of India. India State of Forest Report. 2011: 10-11.
289[3] Manoharan TR. Natural resource accounting: Economic valuation of intangible
290 benefits of forests. RIS Discussion Paper, Research and Information System for the
291 Non-aligned and other Developing Countries, New Delhi. 2000.
292[4] Parthiban KT, Seenivasan R, Vennila S, Anbu PV, Kumar P, Saravanan V, Umesh
293 Kanna S, Rajendran P, Subbulakshmi V and Durairasu P. Designing and augmenting
294 pulpwood supply chain through contract tree farming. **Indian J. Ecol.** 2011: **38**
295 **(Special Issue): 41-47.**

- 296[5] Bansal AK, Rangaraju TS and Shankar KS. Matchsticks from bamboo. **Journal of**
297 **Bamboo and Rattan**, 2002: **1(4)**:333-340.
- 298[6] FAO. Global Forest Resources Assessment 2015, Food and Agriculture
299 Organization, Rome. 2015.
- 300[7] Mahalanobis PC. On the generalized distance in statistics. Proceedings, National
301 Institute of Science, India. 1928: 49-55.
- 302[8] Rao CR. Advanced statistical methods in biometrical research. John Wiley and Sons,
303 New York. 1952: 357-363.
- 304[9] Tewari SK and Singhania DL. Character association and path analysis in grain
305 sorghum. **Sorghum Newsletter**, 1994: **27**: 16-17.
- 306[10] Libby WJ. Genetic resources and variation in forest trees. **In: Improving**
307 vegetatively propagated crops. Academic Press, USA. 1987: 199-209.
- 308[11] Kertadikara AWS and Prat D. Genetic structure and mating system in teak
309 (*Tectona grandis*L.) provenances. **Silvae Genetica**, 1995: **44(2-3)**:104-110.
- 310[12] Mandal AK and Gupta BN. Isozyme differentiation in two subspecies of
311 *Acacia nilotica*. Proc. of Indian National Science Academy, Part B. **Biological**
312 **Sciences**, 1996: **61(1)**:39-42.
- 313[13] Williams JGK, Kubelik AR, Livak KJ, Rafalski JA and Tingey SV. DNA
314 polymorphisms amplified by arbitrary primers are useful as genetic markers. **Nucleic**
315 **Acids Res.**, 1990: **18**:6531-6535.
- 316[14] Mohapatra SK and Singhal RP. Role of molecular markers in biodiversity
317 conservations. **Ann. For.**, 2000: **8(1)**: 1-7.
- 318[15] Mohd AA, Reshi Z and Siddiqi TO. Genetic divergence in half-sib progenies
319 of *Pinus wallichiana* A.B. Jackson plus trees in the Kashmir Himalaya, India.
320 **Tropical Ecology**, 2011:**52(2)**: 201-208.
- 321[16] Paramathma M and Surendran C. Exploitation of heterosis for afforestation in
322 *Eucalyptus*. **In: Proceeding of the International Symposium on Hybrid Breeding and**
323 **Genetics**, 9-14 April, Noose Lake, FRI, Australia. 2000.
- 324[17] Mondal MAA. Improvement of potato (*Solanum tuberosum*
325 L.) through hybridization and invitro culture technique. **Ph.D. Thesis**. Rajshahi
326 University, Rajshahi, Bangladesh. 2003.
- 327[18] Kumar P. Genetic evaluation, Growth characterization and clonal
328 propagation studies in *Melia dubia* Cav. **Ph.D Thesis**, Tamil Nadu Agricultural
329 University, Coimbatore. 2011.
- 330[19] Bagchi SK. Seed source variation in *Acacia nilotica*. Genetic divergence in 8
331 months old seedlings. **Ann. For.**, 1999: **7(1)**:45-55.
- 332[20] Bagchi SK. Genetic divergence in *Tectona grandis*. **Ann. For.**, 2000: **8(1)**:
333 25-37.

- 334[21] Vennila S. Pulpwood traits, genetic and molecular characterization of
335 *Eucalyptus* genetic resources. **Ph.D Thesis**, Tamil Nadu Agricultural University,
336 Coimbatore. 2009.
- 337[22] Chavan Sangram and Keerthika A. Genetic variability and association studies
338 among morphological traits of *Leucaena leucocephala* (Lam.) de Wit. Genetic
339 resources. **Research Journal of Agriculture and Forestry Sciences**, 2013:1(8):23-
340 29.
- 341[23] Saravanan V. Genetic evaluation and wood characterization of *Melia dubia*
342 for pulp, anatomical, mechanical, and energy properties, **Ph.D. Thesis**, Tamil Nadu
343 Agricultural University, Coimbatore.2012.
- 344[24] Subramanian KN, Nicodemus A and Radhamani A. Teak improvement in
345 India. **Forest Genetic Resources**, 1994: **22**: 33-36.
- 346[25] Manoj Kumar K.Genetic divergence, isozyme pattern and micropropagation
347 studies in sandal (*Santalum album*).**M.Sc. Thesis**, Tamil Nadu Agricultural
348 University, Coimbatore. 1994.
- 349[26] Singh NB and Chaudhari VK. Multivariate analysis of genetic divergence in
350 wild apricot (*Prunus armeniaca* Linn.). **Indian J. For.**, 1992: **15(3)**: 211-216.
- 351[27] Kant A, Dutt V and Sharma DR. Genetic variability in phenotypic characters
352 of *Pinus gerardiana*.**Indian Forester**: 2006: 681-690.
- 353[28] Paramathma M.Studies on genetic inheritance and interspecific crosses of
354 *Eucalyptus*. **Ph.D. Thesis**, Tamil Nadu Agricultural University, Coimbatore.1992.
- 355[29] Manga VK and Sen DN. Genetic diversity among different genotypes of
356 *Prosopis cineraria* Druce. **Indian J. For.**, 2000: **23(3)**: 291-295.
- 357[30] Tewari S, Subhanjana K, Shukla AK and Pandey SBS. Genetic Divergence in
358 Shisham (*Dalbergia sissoo* Roxb). **Indian J. For.**, 2002:**25(1)**: 21-24.
- 359[31] Chauhan SK and Sehgal RN. Genetic divergence among progenies of
360 Himalayan long leaf pine.**Indian Journal of Forestry**, 2001: **24**: 65-71.
- 361