### MUTIVARIATE ANALYSIS IN STEM AMARANTH (Amaranthus tricolor)

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#### ABSTRACT

The genetic divergence among 22 genotypes of stem amaranth was estimated using D² and Principal Component Analysis. The genotypes were grouped into four clusters. Cluster I, II, III and IV composed of two, four, seven and nine genotypes in succession. No relationship was found between divergence and geographic distribution of the genotypes. Maximum inter cluster distance (12.326) was observed between cluster I and III and it was minimum (3.526) between cluster I and II. The crosses between the genotypes of cluster I with that of cluster III and cluster II with cluster III would exhibit high heterosis and also likely to produce new recombinants with desired characters in stem amaranth. The yield contributing characters were leaves per plant, petiole length, stem diameter, leaf weight per plant and stem weight per plant. Leaf width, petiole length and 1000 seed weight showed maximum contribution to the total divergence. The results obtained by D² analysis were confirmed by Principal Component Analysis.

**Key words:** Stem amaranth (*Amaranthus tricolor*); D<sup>2</sup>; PCA; cluster analysis; stem yield.

# INTRODUCTION

Amaranth (Amaranthus tricolor L.) is considered to be the cheapest dark-green vegetables in tropical markets and is often described as poor man's vegetable (Shanmugavelu, 1989). Vegetable amaranth occupies a predominant position as the cheapest source of vitamins and minerals. A wide range of variability in respect of morphological trait is available in Bangladesh. But, till now, very few systematic assessment of genetic divergence has been done.

Genetic resources are, in the sense, the building blocks, fundamental not only to crop improvement program, but also for the very survival of the species in time and space (Swaminathan, 1983). Variability is essential for any plant breeding program. A new variety as per requirement of the farmers can be developed from an assembled diverse genetic stocks of any crop. So success of any breeding program depends much on the genetic variability available to the breeder and judicious selection of parents. Genetic diversity is one of the important tools to quantify genetic variability in both cross and self-pollinated crops (Griffing and Lindstrom, 1954; Murty and Aurunachalam, 1966 and Gaur *et al.*, 1978). The quantification of genetic diversity through biometrical procedures has made it possible to choose genetically diverse parents for a successful hybridization program (Rao, 1952; Jain *et al.*, 1975 and Rahman *et al.*, 2005). Tomoka (1991) reported

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that evaluation of genetic diversity is important to know the source of gene for a particular trait within the available germplasm. So the breeder must have clear understanding about the nature and magnitude of variability among the collected genetic stocks. But studies pertaining to multivariate analysis of amaranth have scarcely been studied. Hence, the present study was undertaken to choose of genetically diverged parents to obtain desirable recombinant in segregating generations.

#### MATERIALS AND METHODS

The experiment was conducted with 22 diverse stem amaranth genotypes collected from PGRC, BARI, East West Seed Co. (Lal Teer Seed Ltd.), BRAC and different parts of the country during April to July 2007. All the genotypes were grown in a Randomized Complete Block Design (RCBD) with three replications at the experimental field, Department of Horticulture, Bangabandhu Sheikh Mujibur Rahman Agricultural University (BSMRAU), Gazipur, Bangladesh. The plot size was 1.5 m x 1.5 m. The seed drilling was done between rows at 30 cm apart and seeds were sown continuously in the rows. Seedlings were thinned maintaining a spacing of 30 cm x 10 cm after 15 days of sowing. All the recommended package and practices were followed to raise good crop. The crop was harvested when started flowering. Data were recorded during harvesting time on plant height (cm), leaves per plant, leaf length (cm), leaf width (cm), petiole length (cm), stem diameter (mm), primary branches per plant, leaf weight per plant (g), stem weight per plant (g), leaf - stem ratio, fibre (%), stem yield (t/ha) and 1000 seed weight (g) during seed harvesting. Plot means over three replications were used to statistical analysis. Genetic diversity was studied following Mahalanobis's (1936) generalized distance (D2) extended by Rao (1952). Clustering of genotypes was done according to Tocher's method (Rao, 1952) and Principal Component Analysis (Rao, 1964) for graphical representation of the genotypes. All the statistical analyses were carried out using GENSTAT 5.3 program in IBM computer.

### RESULTS AND DISCUSSION

Analysis of variance revealed that the differences among 22 genotypes were significant for all the thirteen characters under study indicating the presence of notable genetic variability among the genotypes. The  $D^2$  values ranged from 3.526 to 12.326 and principal component scores also indicated a high degree of genetic diversity among the genotypes.

### Principal component Analysis (PCA)

Thirteen (13) characters were considered for genetic diversity analysis. Eigen values of thirteen principal axes and percentage of total variation accounting for them obtained from the principal component analysis are presented in Table 1. The results revealed that the first principal axis largely accounted for the variation among the genotypes which alone contributed 39.96% of the variations followed by the 2<sup>nd</sup> axis (18.51%) while the first four Eigen values for the principal component axes of genotype accounted for 82.96% of the total variation among 13 characters describing 22 genotypes while the former two accounted for 58.47%.

### **Cluster analysis**

On the basis of D<sup>2</sup> analysis, 22 genotypes of stem amaranth were grouped into four clusters (Table 2). Cluster IV contained the highest number of genotypes (nine) followed by cluster III having seven genotypes while cluster I and II having two and four

genotypes, respectively. The clustering pattern of the genotypes under this study revealed that the genotypes collected from the same district were grouped into different clusters. The genotypes of Gazipur district were distributed in three different clusters (Table 2).

Table 1. Eigen values and percentage of variation for corresponding 13 component characters in stem amaranth genotypes

Principal	Eigen velves	Percentage	
component axis	Eigen values	Total variation accounted for	Cumulative
A	5.1954	39.96	39.96
В	2.4059	18.51	58.47
C	1.7939	13.80	72.27
D	1.3893	10.69	82.96
E	0.7394	5.69	88.65
F	0.5072	3.90	92.55
G	0.4549	3.50	96.05
Н	0.2927	2.25	98.30
I	0.1354	1.04	99.34
J	0.0403	0.31	99.65
K	0.0331	0.25	99.90
L	0.0125	0.10	100.00
M	0.00	0.00	100.00

Table 2. Distribution and place of collection of 22 stem amaranth genotypes in four clusters

Cluster	Number of genotypes	Genotypes	Sources/ Place of collection
I	2	SA003	East West Seed Co. Ltd. Gazipur
		SA013	Alamgir Seed, Siddik bazaar, Dhaka
II	4	SA001	East West Seed comp. Ltd.Gazipur
		SA007	Hamid Seed Vander, Jessore
		SA014	Samrat Seed Vander, Siddik bazaar
		SA028	Gazipur
III	7	SA018	BARI, Gazipur
		SA019	BARI, Gazipur
		SA023	Comilla
		SA033	Plant Genetic Resource Center, BARI, Gazipur
		SA037	Plant Genetic Resource Center, BARI, Gazipur
		SA039	Plant Genetic Resource Center, BARI, Gazipur
		SA040	Plant Genetic Resource Center, BARI, Gazipur
IV	9	SA005	Agro Seed, Rangpur
		SA006	Bismillah Seeds, Naogaon
		SA011	Hamid Seed Store, Jessore
		SA015	Sufala Seed, BRAC
		SA020	Faridpur
		SA026	Sylhet
		SA027	Zenaidah
		SA029	Gazipur
		SA035	Plant Genetic Resource Center, BARI, Gazipur

Similar results also reported by Waghmode *et al.* (1997) in *A. hypochondriacus*, Pan *et al.* (1992) in *Amaranthus tricolor*, Joshi and Rana (1995) in grain amaranthus, Quamruzzaman *et al.* (2008) in French bean and Islam (2008) in Hyacynth bean.

Shanmugam and Rangaswamy (1982) reported that falling of materials of same origin into different clusters was an indication of broad genetic base of the genotypes belonging to that origin.

The maximum inter cluster distance was observed between cluster I and III (12.326) followed by cluster II and III (9.082) (Table 3). Intermediate or moderate inter cluster distance was observed between I and IV (7.661) followed by cluster III and IV (5.852) and II and IV (5.485). The minimum inter cluster distance was observed between cluster I and II (3.526) indicating that the genotypes of these clusters were genetically closed or more homogenous. The intra-cluster distance varied from 0.639 to 1.115 comprising two and four genotypes, respectively.

Kanthaswamy (2006), reported that 74 amaranth genotypes of were grouped into 12 clusters, Oboh (2007) observed four clusters for 16 amaranth (*A. hybridus*), Shukla *et al.* (2010) obtained six clusters for 39 amaranth strains and Akhter *et al.* (2013) discriminated 17 amaranth genotypes into four clusters.

Table 3. Intra (bold) and inter cluster distances (D<sup>2</sup>) of 22 stem amaranth genotypes of stem amaranth

Cluster	I	II	III	IV
I	0.639			
II	3.526	1.115		
III	12.326	9.082	1.069	
IV	7.661	5.485	5.852	1.032

The mean values of cluster I ranked first for fibre (%), leaf-stem ratio and 1000 seed weight and second for primary branches per plant and leaf length (Table 4). Where as minimum mean values for leaf width, petiole length, stem diameter, leaf weight, stem weight and stem yield was observed. Cluster II comprising four genotypes had the highest cluster mean for primary branches per plant and second for leaf width, petiole length, leaf-stem ratio and fibre (%), while the minimum mean values for plant height, leaves per plant, leaf length and 1000 seed weight was observed.

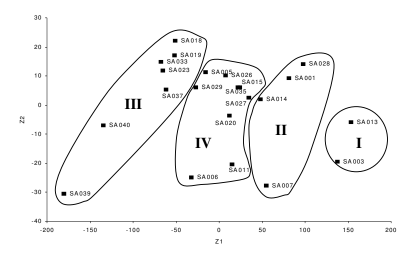
Table 4. Cluster mean values for different characters of stem amaranth genotypes

Characters	Cluster			
Characters	I	II	III	IV
Plant height (cm)	120.00	119.56	128.62	131.75
Leaves per plant	16.10	14.75	21.29	19.48
Leaf length (cm)	20.28	19.44	20.10	20.46
Leaf width (cm)	9.67	10.99	11.19	10.78
Petiole length (cm)	10.05	10.97	11.54	10.66
Stem diameter (mm)	18.16	19.15	22.62	20.94
Primary branches per plant	2.39	2.50	1.65	2.26
Leaf weight per plant (g)	66.33	73.81	109.79	92.10
Stem weight per plant (g)	172.26	243.98	389.91	303.70
Leaf :Stem ratio	0.39	0.30	0.28	0.30
Fibre (%)	0.99	0.95	0.88	0.87
1000 seed weight (g)	0.78	0.73	0.76	0.75
Stem yield (t/ha)	57.36	81.24	129.84	101.13

Cluster III comprising seven genotypes had the highest cluster mean for leaves per plant, leaf width, petiole length, stem diameter, leaf weight per plant, stem weight per plant and stem yield (t/ha) and second for plant height and 1000 seed weight, where as minimum mean values for primary branches per plant and leaf-stem ratio was observed. These results indicated this cluster possessed such characters which were responsible for higher yield. So that for higher yield selection of superior genotypes can safely be done in this cluster. Cluster IV comprising nine genotypes had the highest cluster mean for plant height and leaf length and second for leaves per plant, stem diameter, leaf weight per plant, stem weight per plant, leaf-stem ratio and stem yield (t/ha). Where as the minimum mean value for fibre (%) was observed in this cluster.

Considering yield performance, cluster distance and cluster mean the genotypes SA003 and SA013 from cluster I, the genotypes SA014 and SA028 from cluster II, genotypes SA023, SA033, SA037, SA039 and SA040 from cluster III and genotypes SA005, SA006 and SA029 from cluster IV may be considered better parents for future hybridization program.

Based on principal component axes I and II, a two dimensional scattered plotting diagram ( $Z_1$  and  $Z_2$ ) of the genotypes are presented in Figure 1, reflecting the position of the genotypes. The scattered diagram revealed apparently showed four clusters. Distantly located genotypes of different clusters were SA003 and SA013for cluster I and SA018, SA019, SA023, SA033, SA037, SA039 and SA040 for cluster III. Pattern of distribution of genotypes in the scattered diagram revealed that considerable variability exist among the genotypes.



**Fig 1.** Scatter distribution of 22 genotypes based on their principal component scores super imposed with clustering

### Contribution of the characters towards divergence of the genotypes

The results of PCA revealed that in vector I  $(Z_1)$  the important characters responsible to genetic divergence in the major axis of differentiation were leaves/plant (0.110), primary branches per plant (0.307), and leaf weight per plant (0.117) having positive vector values (Table 5). While in vector II  $(Z_2)$  which was second axis of differentiation, stem weight per plant (0.022), leaf-stem ratio (15.428), fiber (8.247%) and stem yield (t/ha) (0.066) having positive vector values played a major role while rest of the characters played a minor role in the second axis of differentiation. The role of

leaf width, petiole length, and 1000 seed weight in both the vectors indicated the important component of genetic divergence among the 22 genotypes of stem amaranth.

# D<sup>2</sup> and PCA

The clustering pattern of  $D^2$  analysis in Table 2, has followed the same trend of distribution of the genotypes in PCA (Fig 1).  $D^2$  and PCA were found to be an alternative method in giving the information regarding the clustering pattern. Moreover, the PCA provides information on contribution of the characters towards divergence.

Table 5. Latent vectors for 13 principal component characters of stem amaranth genotypes

Characters	Vector-I	Vector-II
Plant height (cm)	-0.051	-0.172
Leaves per plant	0.110	-0.255
Leaf length (cm)	-0.138	-0.619
Leaf width (cm)	0.145	0.458
Petiole length (cm)	0.478	0.612
Stem diameter (mm)	-0.243	-1.088
Primary branches per plant	0.307	-0.201
Leaf weight per plant (g)	0.117	-0.008
Stem weight per plant (g)	-0.046	0.022
Leaf: stem ratio	-25.840	15.428
Fibre (%)	-3.093	8.247
1000 seed weight (g)	2.311	20.495
Stem yield (t/ha)	-0.139	0.066

The crosses involving parents belonging to the maximum divergent clusters were expected to manifest maximum heterosis and also wide variability in genetic architecture. Ramnujam *et al.* (1974) in mungbean and Mian and Bhal (1984) in chickpea reported that parental clusters separated by medium D² values exhibited significant and positive heterosis for seed yield and some of its components in mungbean. Thus crosses between the genotypes of cluster I with that of cluster III and cluster II with cluster III would exhibit high heterosis and also likely to produce new recombinants with desired characters in stem amaranth.

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