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# **Stability and Molecular Characterization of Heat Tolerant Genotypes of Chickpea (***Cicer arietinum* **L.)**

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#### *Authors' contributions*

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

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*<u>Original Research Article</u>* 

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# **ABSTRACT**

Twenty eight diverse genotypes sown in three different dates were screened using thirty three SSR primers. Twelve morphological characters recorded. The current study was conducted at all India Coordinated Research Project on Chickpea at R.A.K., College of Agriculture, Sehore (M.P.) during *Rabi* 2020-21, and 2021-22. The molecular work was carried out at Plant Molecular Biology Laboratory, Department of Plant Molecular Biology and Biotechnology, College of Agriculture, Rajmata Vijayaraje Scindia Krishi Vishwa Vidyalaya, Gwalior (M.P.). The component G×E interaction were found significant for flower initiation, days to 50% flowering, days to pod initiation, plant height, days to maturity, number of pods per plant, number of empty pods per plant, number of seeds per plant, biological yield per plant, harvest index, 100 seed weight and seed yield per plant. The highest gene diversity was found in TA-135 (0.7474) followed by GAA-44 (0.7219), GAA-40 (0.7015), STMS-2 (0.6939), TA-71 (0.6709), NCPGR-1 (0.6403) and TA-18 (0.3648). Based on a dendrogram all the 28 genotypes were grouped into three major clusters, in which cluster I contained 2 genotypes, cluster II contained 5 genotypes and cluster III encompassed remaining 21 genotypes. Genotypes RVG 204, JG-14, and RVSSG-61 were found stable for favourable and unfavourable sowing conditions, while ICC-4958, JG-11, JG-12, RVG-203, RVG-204, RVSSG-52, JG-74, RVSSG-71 showed consistent performance during unfavourable sowing conditions for seed yield per plant. The important traits and marker based diversity and stability has been discussed in this research paper.

*Keywords: Chickpea; heat stress; stability analysis; molecular diversity.*

#### **1. INTRODUCTION**

Chickpea (*Cicer arietinum* L.) is a cool season food legume grown in more than 50 countries across all continents. Among pulses, chickpea is one of the most important protein-rich food legumes majorly grown under rainfed condition. Pure lines, hybrids, synthetics, or any other material utilised for breeding typically have genotypic and environmental interaction present under all conditions, which makes breeding difficult and prevents the advancement of the crop improvement programme [1]. The selection of superior genotypes for both new crop production and improved cultivar development can be seriously affected by a significant G×E interaction for a quantitative trait like seed yield [2]. A temperature of  $35^{\circ}$ C was found to be critical in differentiating heat tolerant and sensitive genotypes in chickpea under field conditions [3]. High temperature during the reproductive stage is a major cause of yield loss due to partial or complete pollen sterility. In chickpea, temperatures at or exceeding 35°C also affected male reproductive tissue (anther and pollen), function (pollen germination and tube growth) and pod set [4]. Heat killing temperature in chickpea was found 44.3°C for 41 minutes [5]. Stigma receptivity was also affected at high temperature (≥40/30˚C) through oxidative stress in the leaves which causes failure of fertilization [6]. Above 45°C no germination is

observed due to lack of embryo growth [7]. Temperature is an important factor controlling crop growth and development [8] by affecting wide range of physiological processes and altering plant-water relationship. Therefore, it is essential to examine a crop's performance in various conditions in to find genotypes that provide high yield across a variety of environments. These genotypes will be very helpful for maximising their potential for the development of stable and high-yielding cultivars. Due to the growth of irrigation facilities in MP, farmers are planting more chickpea, and they favour the early genotypes with high yields that are heat tolerant It has been demonstrated that molecular markers are essential to crop development programmes. These markers act as effective and potent tools for the marker-assisted selection of traits that are significant from an agronomic aspect. Understanding the genetic foundation of chickpea variations would help breeders plan future crossing programmes and focus their efforts in a way that would increase the genetic diversity of such types. The goal of the current study was to discover how the G×E interaction affected the morphological and yield-attributing features of plants growing in both normal and heat-stress settings. Additionally, to examine the molecular diversity of each chickpea genotype in order to determine the best to use it looking forward in breeding programmes.

# **2. MATERIALS AND METHODS**

The current study was conducted at all India Coordinated Research Project on Chickpea at R.A.K., College of Agriculture, Sehore (M.P.) during *Rabi* 2020-21, and 2021-22. The molecular work was carried out at Plant Molecular Biology Laboratory, Department of Plant Molecular Biology and Biotechnology, College of Agriculture, Rajmata Vijayaraje Scindia Krishi Vishwa Vidyalaya, Gwalior (M.P.) during 2021-2022. Sehore, is situated on 27°12 north latitude and 77°0 east longitude at an altitude of 498.77 meters from mean sea level in Vindhyan Plateau of Madhya Pradesh. The average annual rainfall varies from 1000 to 1200 mm, concentrated mostly from June to September. The temperatures vary from 4.0ºC minimum in January to 42ºC maximum in May. The experimental material comprised of 28 genotypes, which were grown in a RCBD with two replications on three different dates (Tables 1 and 2). Consisting of 2 rows of 2m length, the row to row distance was 30 cm and plant to plant spacing was 10 cm. The fertilizer dose 20:60:20 NPK Kg/ha was applied at the time of sowing and recommended package of practices were adopted for optimum crop growth further plant protection under irrigated condition was applied when required. Field observations were recorded on single plant basis on five selected plants from each plot of each replication for 12 morphological characters, which were flower initiation, days to 50% flowering, days to pod initiation, plant height, days to maturity, number of pods per plant, number of empty pods per plant, number of seeds per plant, biological yield per plant, harvest index, 100 seed weight and seed yield per plant. The data were statistically analyzed in accordance with method described by Eberhart and Russell [1].

For molecular analysis, DNA from 2g of fresh young leaf tissue was collected in the winter season of 2021-2022 and was immediately frozen in liquid nitrogen and stored at -80ºC. Isolation of DNA was carried out using modified CTAB method. Thirty three SSR primers were screened, out of which only seven were polymorphic (Table 4). PCR analysis was taken up by having preparation of 3 min at 95ºC followed by 35 cycles of denaturation at 95ºC for 20s, annealing for 20 s at 50-55ºC and 1.5 min. initial elongation at 72 ºC and 10 min. elongation at 72ºC and finally hold at 15 min were performed [9].

Band patterns for each of the microsatellites markers were recorded for each genotype by assigning a letter to each band. Alleles were numbered as A/A, B/B, C/C, D/D, E/E, F/F, G/G sequentially from the smallest to the largest sized band. Only clear and detectable bands were scored for data analysis. The PCR products from SSR analyses were scored quantitatively are present or absent of amplicons. DNA bands were scored '1' for its presence and '0' for its absence. For Clustering, UPGMA was used based on the similarity matrix generated on combined data. Polymorphic information content for each SSR primer pair was calculated.

# **3. RESULTS AND DISCUSSION**

Analysis of variance revealed significant variance due to genotype against pooled deviation for all the characters days to flower initiation, days to 50% flowering, days to pod initiation, plant height, days to maturity, number of pods per plant, number of empty pods per plant, number of seeds per plant, biological yield per plant, harvest index, 100 seed weight and seed yield per plant indicating the presence of genetic variability for the traits under investigation.

The component GxE linear were found significant for all the characters indicated that the genotypes interacted considerably to environmental condition and major portion of G×E interaction was attributed to linear component in respect of these traits. Non-liner component (pooled deviation) was also found to be significant for most of the characters (Table 3). Yadav et al. [10], Babbar and Tiwari [11] have also recorded significantly G×E interaction for most of the yield and its contributing traits in chickpea.

When the overall mean, regression coefficient and mean square deviation from regression are taken into consideration, genotype JG-14 were found to be stable for days to flower initiation, with mean values greater than population mean and regression coefficient lesser than one with deviation from regression. It means days to flower initiation had less susceptibility for these genotypes against change of environmental condition in the expression of this character. Looking to the above parameters the genotypes RVG-203, RVKG-121, RVSJKG-102 for days to 50% flowering; JG-12,JG-14,RVG-202, for days to pod initiation; JG-12,RVG-202,RVG-203,RVG-204 for plant height; ICC-4959, JG-14, RVSSG-71 for days to maturity, ICC-4958, JG12,



# **Table 1. Description of chickpea genotypes used in the experiment**

# **Table 2. Sowing season and timing of experimental material**



# **Table 3. Stability analysis of variance of pooled data for different morphogical traits in chickpea**



*Note: \* and \*\* significant at 5% and 1% level of probability, respectively.*



# **Table 4a. Stability parameters for various morphological traits in chickpea**

Genotypes	Number of empty			Number of seeds per			<b>Biological Yield per</b>			<b>Harvest index</b>			100 seed weight			Seed vield per		
	pods per plant			plant			plant									plant		
	X	Bi	$s^2$ di	$\mathbf{x}$	bi	$s^2$ di	X	Bi	$s^2$ di	X	Bi	$s^2$ di	X	Bi	$s^2$ di	X	Bi	$s^2$ di
<b>ICC-4958</b>	2.69	1.79	$-0.59$	34.50	$-0.08$	$-5.15$	16.50	$-0.10$	$-0.30$	35.50	0.90	$-3.01$	25.30	$-0.73$	4.88	5.77	0.15	$-0.145$
<b>RVG-202</b>	3.14	0.09	0.75	45.10	1.91	33.90	25.10	1.48	19.00	32.10	3.06	2.21	26.00	$-1.16$	2.02	8.60	2.29	2.800
RVSSG-51	4.69	6.22	5.72	20.20	1.34	30.30	11.90	1.46	2.50	45.30	0.30	21.10	18.30	1.90	9.13	5.59	1.41	0.323
$JG-74$	7.19	0.11	5.83	19.30	1.08	$-3.90$	8.56	0.63	0.31	33.10	1.11	$-3.62$	15.40	2.43	$-0.90$	2.97	0.55	$-0.190$
<b>RVG-204</b>	2.48	0.61	0.09	43.50	1.12	$-6.24$	20.40	$-0.17$	$-0.76$	43.00	0.41	0.66	27.20	$-0.26$	4.41	8.80	0.01	0.148
$JG-14$	3.47	1.41	$-0.84$	35.40	1.19	$-4.43$	21.90	0.75	10.20	33.60	3.11	229.0	25.60	$-0.31$	$-0.91$	7.27	1.61	0.769
RVSSG-75	4.33	1.22	$-0.78$	23.60	1.11	47.50	16.80	1.44	43.10	36.70	1.51	7.01	25.10	1.09	0.38	6.38	1.53	5.020
$JG-11$	3.81	$-2.18$	2.11	40.90	$-0.02$	$-5.10$	18.70	0.37	3.46	42.90	0.32	3.07	25.50	0.64	7.05	7.88	0.46	0.273
<b>JG-315</b>	6.25	1.09	2.02	21.00	1.17	8.03	12.80	0.98	4.68	35.90	1.56	0.08	14.00	0.29	$-0.91$	4.98	0.98	0.346
<b>RVG-203</b>	2.86	3.46	$-0.85$	45.10	0.09	$-5.12$	20.50	$-0.06$	$-0.34$	40.80	0.44	10.90	25.50	$-0.80$	6.48	8.15	0.11	0.603
<b>JAKI-9218</b>	4.91	1.62	$-0.39$	28.20	1.48	67.20	17.70	1.63	$-0.85$	43.50	0.77	2.46	21.60	0.55	$-0.05$	8.03	1.55	0.604
$JG-12$	4.64	$-2.08$	1.22	45.00	0.10	$-4.79$	18.70	0.11	$-0.86$	37.50	$-0.04$	$-2.95$	24.50	0.20	$-0.14$	6.79	0.05	$-0.289$
$JG-130$	9.36	2.46	8.98	36.80	2.62	115.0	18.70	2.65	50.50	42.20	0.17	21.00	19.70	2.13	$-0.43$	8.07	2.38	10.00
$JG-6$	8.33	2.17	1.66	22.10	0.90	23.50	13.00	0.79	6.48	38.80	2.09	20.20	18.30	1.44	$-0.68$	5.54	0.95	3.010
<b>RVSSG-52</b>	3.97	0.72	$-0.62$	19.40	0.57	$-1.60$	14.30	0.77	$-0.64$	32.70	0.20	$-4.64$	21.70	0.90	$-0.37$	4.64	0.54	$-0.287$
<b>ICC-4812</b>	7.06	0.40	0.74	30.10	1.66	13.90	11.90	1.02	$-0.33$	29.90	1.87	0.57	12.60	1.26	$-0.90$	3.87	0.95	$-0.264$
RVSSG-71	9.08	$-1.21$	18.80	22.30	0.91	9.90	12.20	0.51	42.10	27.40	0.51	116.0	16.20	1.99	6.25	3.09	0.38	$-0.266$
<b>BGD-112</b>	2.78	$-0.03$	0.11	16.50	0.87	17.10	11.60	1.03	13.90	29.90	0.96	8.68	14.10	1.29	3.57	3.57	0.78	0.379
RVSSG-68	9.06	$-4.73$	148.00	43.20	1.42	0.58	11.80	0.54	6.94	33.40	2.13	0.37	10.50	0.66	$-0.53$	4.45	0.70	1.770
RVSSG-61	4.06	0.85	$-0.68$	45.40	$-0.01$	$-3.54$	23.60	1.24	$-0.82$	38.20	1.45	$-1.27$	40.60	1.04	2.57	9.25	1.45	$-0.157$
JGK-5	4.03	3.16	$-0.68$	11.80	0.41	0.68	13.80	0.98	$-0.18$	39.20	0.86	5.18	42.90	1.06	0.10	5.60	0.95	0.188
<b>PKV-4</b>	5.72	7.83	6.38	15.70	0.66	$-6.12$	19.40	2.73	2.38	44.70	$-0.06$	$-4.74$	42.00	2.19	$-0.86$	8.44	2.37	0.102
<b>KRIPA</b>	4.14	$-1.70$	1.46	26.10	1.57	$-6.25$	15.20	1.08	17.50	30.10	$-0.25$	99.70	23.50	1.58	$-0.49$	4.14	0.61	$-0.284$
<b>RVKG-111</b>	4.33	$-0.14$	$-0.37$	24.90	1.59	20.40	16.50	1.60	4.75	33.80	2.05	$-3.33$	23.80	1.91	$-0.80$	6.61	1.55	1.190
<b>RVKG-121</b>	4.66	$-0.55$	1.02	31.90	1.56	32.60	18.90	0.98	54.70	35.80	1.72	7.65	22.00	0.94	$-0.33$	7.27	0.92	6.700
RVSJKG-102	3.58	1.36	5.14	18.60	0.70	7.31	15.00	0.93	$-0.37$	35.40	1.33	$-0.43$	24.30	4.10	70.30	5.66	0.88	$-0.237$
RVSSG-36	3.02	1.70	$-0.81$	29.70	0.94	$-5.73$	17.30	1.56	0.72	37.40	0.32	$-4.01$	20.70	0.47	5.38	6.81	1.21	1.160
RVSSG-63	5.42	2.35	$-0.56$	24.00	1.13	8.35	15.30	1.05	$-0.91$	40.30	$-0.80$	1.61	33.30	1.19	0.12	5.88	0.69	0.0002

**Table 4b. Stability parameters for various morphological traits in chickpea**





#### *Tare et al.; Int. J. Environ. Clim. Change, vol. 13, no. 2, pp. 55-65, 2023; Article no.IJECC.96696*



Where, DFI: Days to flower initiation, D50%F: Days to 50% flowering, DPI: Days to pod initiation, DM: Days to maturity, PH: Plant height, NPPP: Total number of pods per plant, NEPP: Number of *effective pods per plant, NSPP: Number of seeds per pod, 100 SW: 100 seed weight, BY: Biological yield per plant, HI: Harvest index, and SYPP: Seed yield per plant*

RVSSG52, RVG-202, RVG-203, RVSSG-61, RVSSG-68, RVKG-102 for number of pods per plant; JG-11, JG-12 for empty pods per plant, RVSSG-52, ICC-4958, JG-11, JG-12, RVG-203 for number of seeds per plant, ICC-4958, RVSSG-52, JG-11, JG-12, RVG-203, RVG-204, for seed yield per plant, ICC-4958, JG-12, RVG-203, RVG-204, for biological yield per plant, JG-12, RVSSG-52 for harvest index and JG-14, JG-202, JG-12 for hundred seed weight were found to be stable in poor environmental conditions respectively. It indicated that these genotypes should be given due consideration at the time of formulation of breeding programme specially for mid late sown and very late sown conditions (Table 4a, 4b).

Twenty eight genotypes with higher/lower mean values than grand mean were divided into three groups based on stability parameters viz., mean, regression coefficient and squared deviation, (Table 5) according to the methodology followed by Ramanujam [12]. Genotypes falling in group I have desirable mean, regression coefficient value around unity with non-significant squared deviation. Under group II, genotypes with significantly less than unity regression value and non-significant squared deviation were taken, indicating suitability towards unfavourable environments. Again, the genotypes with significantly more than unity regression was also classified under group II indicating its suitability towards favourable environments. Finally, genotypes falling in group III and cannot be predicted as they exhibited significant squared deviation, irrespective of the regression coefficient values.

According to the grouping (Table 5), the genotypes RVG 204, JG-14, and RVSSG-61 were found stable in unfavourable condition for most of the traits under study. Under group II (bi<1) the genotype ICC-4958 was found to be stable for days to maturity, number of pods per plant, number of seeds per plant, and biological<br>yield per plant, perform better under perform better under unfavourable conditions.

Genotype ICC-4958, JG-11, JG-12, RVG-203, RVG-204, RVSSG-52, JG-74, RVSSG-71 were observed to exhibit constant performance during unfavourable conditions for seed yield per plant. The genotype RVSSG-68 placed under group II (bi>1) and was stable in favourable conditions for days to flower initiation, days to 50% flowering, days to pod initiation, number of seeds per plants, and for harvest index; while the genotype PKV-4 was stable in favourable conditions for the traits seed yield per plant, biological yield per plant, days to maturity, and for hundred seed weight.

At molecular level out of 33 primers only 7 SSR primers were highly polymorphic and rest other primers were monomorphic, these 7 polymorphic SSR primers were used for screening of all the genotypes in the present study. The polymorphic information content among the markers ranged from 0.3426 (TA-18) to 0.7035 (TA-135) with the mean value of 0.5990. TA-135 (0.7035) showed highest polymorphic information content as well as highest gene diversity (0.7474). The study revealed that all 28 diverse genotypes were grouped into three major clusters (Fig. 1). Bhardwaj et al. [13] also grouped different chickpea lines into two clusters in their study using molecular markers. In which, cluster I contained 2 genotypes, cluster II contained 5 genotypes and cluster III encompassed remaining 21 genotypes. Cluster I included two genotypes namely JG-74 and JG-11. Cluster II was divided into 2 subgroups – II A with 4 genotypes and II B with one genotype (JG-14). Cluster II A was further divided into small subgroups (subgroup C and subgroup D). The subgroup D contains only one genotype – RVG-203.

The cluster C was again divided into two subgroups - E and F. E with two genotypes – RVKG-121 and RVG-204, F with only one genotype- RVSSG-75. Cluster III included 21 genotypes, which was divided into small subgroups – subgroup G with one genotype RVG-202 and subgroups H. Subgroup H was again divided into small subgroups – I with one genotype – RVSSG-51 and subgroup J. Subgroup J was further divided into subgroup K and subgroup L. Subgroup K with only one genotype ICC-4958. Subgroup L was further divided into subgroup M with two genotypes (JG-315 and BGD-112) and subgroup N with subgroup O and subgroup P. Subgroup O with 7 genotypes (RVSSG-36, RVKG-102, RVSSG-52, JG-12, RVSSG-63, 1CC-4812 and RVSSG-71) and Subgroup P with 9 genotypes (RVSSG-68, JG-130, JG-6, PKV-4, KRIPA, RVKG-111, JGK-5, RVSSG-61 and JAKI-9218). Many genotypes, which were derived even from diverse parents, were clustered together because of selections during the advancement of generations. In this study, Kabuli and Desi lines did not grouped into two broad categories. This indicates that the Kabuli and Desi lines have not evolved in wide



**Fig. 1. Cluster dendrogram showing the genetic relationships between 28 genotypes of chickpea based on the alleles detected by 32 microsatellite markers**

isolation and only few genes are involved in their differentiation; similar to the observations made earlier [14-16]. In this study RVSSG-75 makes a different sub cluster, indicating it is quite different to rest of the lines.

The highest gene diversity was found in TA-135 (0.7474) followed by GAA-44 (0.7219), GAA-40 (0.7015), STMS-2 (0.6939), TA-71 (0.6709), NCPGR-1 (0.6403) and TA-18 (0.3648). The power and potential of SSR markers for a wide range of applications in genetic and breeding of chickpea has been well demonstrated by

Flandez-Galvez et al. [17], but still substantial numbers of chickpea microsatellites are not available in public domain. Microsatellite genotypic data from a number of loci have potential to provide unique allelic profiles or DNA fingerprints for establishing genotypes identity as well as in development of molecular linkage map of chickpea.

#### **4. CONCLUSION**

In the present study, the population structure and dendrogram analysis gave out 3 major clusters showing the varietal distribution, Which can be

used efficiently for crossing program and variety development. The Genotypes ICC-4958, JG-11, JG-12, RVG-203, RVG-204 were overall best performing genotypes.

# **COMPETING INTERESTS**

Authors have declared that no competing interests exist.

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