STUDY OF GENETIC ADVANCE AND BROAD SENSE HERITABILITY FOR GRAIN YIELD AND YIELD COMPONENTS OF CHICKPEA (CICER ARIETINUM L.) GENOTYPES

Gudiyada Parida1, Siamak Saghfi2,5, Alireza Eivazi2,6, Abolfazl Akbarzadeh2,4
Taras Kavetskyy2,8,9, Immi Aliyeva2,10, Rovshan Khalilov2,3,7

1Ethiopian Institute of Agricultural Research, Addis Ababa, Ethiopia
2Joint Ukraine-Azerbaijan International Research and Education Center of Nanobiotechnology and Functional Nanosystems, Drohobych, Ukraine & Baku, Azerbaijan
3Department of Biophysics and Molecular Biology, Faculty of Biology, Baku State University, Baku, Azerbaijan
4Biotechnology Research Center, Tabriz University of Medical Sciences, Tabriz, Iran
5Department of Plant Physiology, Faculty of Biology, Baku State University, Baku, Azerbaijan
6Seed and Plant Improvement Research Department, West Azerbaijan Agricultural and Natural Resources Research and Education Center, AREEO, Urmia, Iran
7Institute of Radiation Problems, National Academy of Sciences of Azerbaijan, Baku, Azerbaijan
8The John Paul II Catholic University of Lublin, Lublin, Poland
9Drohobych Ivan Franko State Pedagogical University, Drohobych, Ukraine
10Department of Environmental Engineering, Azerbaijan Technological University, Ganja, Azerbaijan

Abstract. Genetic variability is a prerequisite to crop improvement. The objective of the present study was to assess and quantify the genetic variability, estimate heritability and genetic advance for yield and yield contributing characters of released and pipeline chickpea varieties based on agro- morphological traits. A total of 16 chickpea materials (8 improved varieties, 7 advanced lines, and one local check) were planted in Randomized Complete Block Design (RCBD) with three replications at Shambu, Hawa Galan, Mata, Alaku Belle, and Badesso, in western Ethiopia. Important agronomic data were collected and subjected to analysis using appropriate software. The combined analysis of variance (ANOVA) indicated highly significant differences (P≤0.01) among chickpea genotypes for grain yield and other agronomic traits except for seed per pod and branch per plant, indicating the existence of ample genetic variability among present chickpea genotypes. GCV and PCV for the character such as grain filling duration, days to flowering and hundred seed weight, showed very small difference demonstrating that the observed variations for these traits were mostly due to genetic factors with little impact of environment. The higher heritability coupled with higher genetic advance was noted for hundred seed weight, pods per plant, seed per pod and grain yield signifying that the ease of phenotype-based selection for the improvement of those traits.

Keywords: Chickpea (Cicerareitinum L.), Genetic variability, PCV, GCV, Selection.

Corresponding Author: Siamak Saghfi, Ph.D., Department of Plant Physiology, Faculty of Biology, Baku State University, Z. Khalilov 23, Baku, Azerbaijan, e-mail: s.saghfi@yahoo.com

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1. Introduction

Among pulses, chickpea (*Cicer arietinum* L.) is the fourth leading grain legume primarily grown for food and feed as well as soil fertility amendment in Ethiopia (CSA, 2015). Besides, Ethiopia is known as secondary center of origins for different crop in general and for chickpea in particular. Regardless of its importance and presence of immense genetic variability, its yield is by far below its potential. This yield gap can be altered through proper manipulation of variable genotypes and thereby using in a breeding program. The main objective of any breeding programs in the world is to produce high-yielding and better-quality lines for release as cultivars to farmers and commercial growers. The prerequisite to achieving this goal is to find sufficient amount of variability.

Since long-time plant breeders were aware that predominant portion of phenotypic variation were accounted from environmental variation than from genotypes and genotypes by environmental interactions. Thus, in-depth knowledge of genetic variation has paramount importance for yield and its component improvement in any crop as the observed variability is a combined effect of genetic, environments and various interactions between genes and environments, of which only the former one is heritable. Genotypic variability is, therefore, the component of variation which is due to the genotypic differences among individuals within a population and is the main concern of plant breeders all over the world (Tadesse et al., 2016).

Selection would generate promising genotypes only if the source germplasms used in crossing are genetically diverse (Quban et al., 2010). When genetically diverse parents were chosen for crossing, there is a likelihood of producing heterotic progenies and advance required genetic recombination.

However, the amount of genetic variability present in any genotypes will not merely dictate the selection of superior genotypes. Therefore, in addition to the existence of genetic variability, the extent to which the characters in questions are transmitted from parents to offspring is very important to draw realistic conclusion vis-à-vis superior selections (Falconer & Mackay, 1996). Transmission of a quantitative trait from parents to offspring by itself is highly influenced by genotype by environment interaction and traits with low heritability might not be improved through selection. Additionally, the estimates of heritability alone do not lonely deliver an idea about the expected gain in the next generation but should be considered in conjunction with estimates of genetic advance, the change in the mean value among successive generations (Shukla et al., 2006; Johnson et al., 1955).

Assessment of genetic variability using appropriate tools such as genetic coefficient of variation, heritability estimates, and genetic advance is, therefore, undeniably indispensable to boost yield as well as the quality of any crop and/or forage in any breeding program (Atta et al., 2008).

The objective of the present study was, therefore, to assess and quantify the genetic variability of chickpea germplasm grounded on agro-morphological traits and evaluating heritability and genetic advance among desi type chickpea genotypes for yield and yield contributing characters.
2. Materials and methods

Experimental sites, designs and experimental materials

A field experiment was conducted at five locations viz., Shambu, Hawa Galan, Mata, Alaku Belle and Badesso, western Ethiopia, during the 2016/17 main cropping season. A total of 16 desi type chickpea varieties viz., 8 cultivars released over three decades, 1 local variety and 7 advanced lines collected from Debre Zeit Agricultural Research Center (DZARC) were used (Table 1).

Table 1. Passport description of the test genotypes

<table>
<thead>
<tr>
<th>Genotype Codes</th>
<th>Genotype names</th>
<th>Status</th>
<th>Year of release</th>
</tr>
</thead>
<tbody>
<tr>
<td>G1</td>
<td>Akaki</td>
<td>Released</td>
<td>1995</td>
</tr>
<tr>
<td>G2</td>
<td>Dalota</td>
<td>Released</td>
<td>2013</td>
</tr>
<tr>
<td>G3</td>
<td>Dimtu</td>
<td>Released</td>
<td>2012</td>
</tr>
<tr>
<td>G4</td>
<td>Dubie</td>
<td>Released</td>
<td>1978</td>
</tr>
<tr>
<td>G5</td>
<td>Local</td>
<td>Local variety</td>
<td>-</td>
</tr>
<tr>
<td>G6</td>
<td>Mariye</td>
<td>Released</td>
<td>1985</td>
</tr>
<tr>
<td>G7</td>
<td>Minjar</td>
<td>Released</td>
<td>2010</td>
</tr>
<tr>
<td>G8</td>
<td>Natoli</td>
<td>Released</td>
<td>2007</td>
</tr>
<tr>
<td>G9</td>
<td>Teketay</td>
<td>Released</td>
<td>2013</td>
</tr>
<tr>
<td>G10</td>
<td>DZ-2012-CK-0032</td>
<td>Advanced line</td>
<td>-</td>
</tr>
<tr>
<td>G11</td>
<td>DZ-2012-CK-0034</td>
<td>Advanced line</td>
<td>-</td>
</tr>
<tr>
<td>G12</td>
<td>DZ-2012-CK-0233</td>
<td>Advanced line</td>
<td>-</td>
</tr>
<tr>
<td>G13</td>
<td>DZ-2012-CK-0237</td>
<td>Advanced line</td>
<td>-</td>
</tr>
<tr>
<td>G14</td>
<td>DZ-2012-CK-0312</td>
<td>Advanced line</td>
<td>-</td>
</tr>
<tr>
<td>G15</td>
<td>DZ-2012-CK-0313</td>
<td>Advanced line</td>
<td>-</td>
</tr>
<tr>
<td>G16</td>
<td>DZ-2012-CK-20113-2-0042</td>
<td>Advanced line</td>
<td>-</td>
</tr>
</tbody>
</table>

The experiment was laid out in a randomized complete block design with three replicates and with plot size of 3m length and 1.8 m width. All other crop management practices and recommendations were applied uniformly to all varieties as recommended for the crop.

Statistical Analysis

Days to 50% flowering, days to 90% maturity, grain filling duration, pods per plant, seed per pod, plant height, branch per plant, hundred seed weight, and grain yield data were collected as suggested in IBPGR, ICRISAT, ICARDA (1993) and were subjected to analysis using appropriate software. Analysis of variance (ANOVA) using randomized complete block design was computed for all measured characters using SAS (SAS Institute, 2002). Genotypic ($\sigma^2_g$), genotype by location interaction ($\sigma^2_gl$) and phenotypic variances ($\sigma^2_p$) were obtained from the analysis of variance (ANOVA) table according to Comstock and Robinson (1952) as follows,

Genotypic variance

$$ (\sigma^2_g) = \frac{MS_g - MSE}{lr} $$

where, $MS_g$ is mean square of genotype, $MSE$ is mean square of error, $l$ is number of locations and $r$ is replication.

Genotype by location interaction variance was estimated as:

Genotype by location interaction variance

$$ (\sigma^2_gl) = \frac{MS_{gl} - MSE}{r} $$
where, $Msgl$ is mean square of genotype by location interaction, $Mse$ is mean square of error, and $r$ is replication.

Phenotypic variance ($\sigma^2_p$) = $\delta^2_g + \delta^2_e + \delta^2_{gl}$

where, $\delta^2_g$ is genotypic variance, $\delta^2_e$=environmental variance, and $\delta^2_{gl}$ is genotype by location interaction.

Heritability in the broad sense ($H^2$) was estimated according to Falconer (1989) as follows,

$$H^2 = \frac{\delta^2_g}{\delta^2_p} \times 100$$

where, $\delta^2_g$ is genotypic variance and $\delta^2_p$ is phenotypic variance.

The mean values for genetic analyses were used to determine phenotypic coefficient of variation (PCV), Environmental coefficient of variations (ECV), and genotypic coefficient of variation (GCV), as described by Singh and Choudhury (1985) as follows,

$$GCV = \frac{\sqrt{\delta^2_g}}{\bar{x}} \times 100$$

where, $\delta^2_g$ is genotypic variance and $\bar{x}$ is grand mean of a character.

$$PCV = \frac{\sqrt{\delta^2_p}}{\bar{x}} \times 100$$

where, $\delta^2_p$ is phenotypic variance and $\bar{x}$ is grand mean of a character.

$$ECV = \frac{\sqrt{\delta^2_e}}{\bar{x}} \times 100$$

where, $\delta^2_e$ is error variance and $\bar{x}$ is grand mean of a character.

Genetic advance (GA) was calculated with the method suggested by Allard (1960); Singh and Choudhary (1985) as follows,

$$GA = K \cdot \sqrt{\delta^2_p} \times H^2$$

where $K$ is 2.06 at 5% selection intensity, $\sqrt{\delta^2_p}$ is square root of phenotypic variance and $H^2$ is Heritability.

Genetic advance as a percentage of the mean (GAM) was computed as,

$$GAM (%) = \frac{GA}{\bar{x}} \times 100$$

where, $GA$ is genetic advance and $\bar{x}$ is grand mean of a character.

3. Results and discussion

Genetic variability

The combined analysis of variance (ANOVA) for grain yield and other agronomic traits of sixteen chickpea genotypes grown at five locations in 2016/17 was shown in Table 2.

The analysis of variance (ANOVA) indicated highly significant differences ($P \leq 0.01$) among chickpea genotypes for grain yield and other agronomic traits except for seed per pod and branch per plant demonstrating the presence of ample genetic variability among chickpea genotypes. Once the presence of genetic variability is secured it is the task of the breeder to separate the heritable portion from the non-heritable part to plan for the proper breeding program. This might give an opportunity to plant breeders to improvement of characters with high heritability via selection.
Table 2. Combined analysis of variance for grain yield and other agronomic traits of sixteen chickpea genotypes grown at Shambu, Hawa Galan, Badesso, Mata, and Alaku Belle in 2016.

<table>
<thead>
<tr>
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</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>D2F</td>
<td>D2M</td>
<td>GF</td>
<td>PPP</td>
<td>SPP</td>
<td>PH</td>
<td>BR</td>
<td>HSW</td>
<td>GYL</td>
</tr>
<tr>
<td></td>
<td>Mean squares</td>
<td>10.44</td>
<td>43.83**</td>
<td>96.53**</td>
<td>84.84**</td>
<td>0.22**</td>
<td>4.74**</td>
<td>3.32**</td>
<td>22.8*</td>
</tr>
<tr>
<td></td>
<td>p</td>
<td></td>
<td></td>
<td></td>
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<td></td>
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</tbody>
</table>
| Key: - *** = Very highly significant at P ≤ 0.001, ** = highly significant at P ≤ 0.01, * = significant at P ≤ 0.05, ns = not significant at P = 0.05, a Numbers in parentheses are degrees of freedom associated with the corresponding source of variation, Rep(Loc) = Replications within locations, D2F= Days to flowering, D2M= Days to maturity, GF= grain filling duration, PPP= pods per plant, SPP=seed per pod, PH=plant height BR=branch per plant, HSW=hundred seed weight, GYL= grain yield

Estimates of variance and genetic parameters

Splitting the total phenotypic variance into the genotypic, genotype by environment interaction, and error variances were done found on their expected mean squares (EMS) as per the fixed effects model. The genotypic variance (σ^2g) account slow proportion of the total phenotypic variance (σ^2p) as compared to the variances due to error (σ^2e) in most of the traits studied (Table 3). This shows that the observed phenotypic differences were not solely due to inherent genetic differences among the genotypes but also the substantial portion was due to environmental factor and the interplay between environment and genotype. Genotypic variance, phenotypic variance, genotypic coefficient of variability (GCV), environmental coefficient of variation (ECV), phenotypic coefficient of variability (PCV), broad sense heritability, and genetic advance for nine (9) agronomic characters are indicated in Table 3.

Estimates of phenotypic coefficients of variation (PCV) and genotypic coefficients of variation (GCV) ranged from 2.24% (for days to maturity) to 33.60% (for pod per plant) and 6.71% (for days to maturity) to 20.67% (for hundred seed weight), respectively. Likewise, estimates of environmental coefficient of variation ranged from 1.83 (for days to maturity) to 28.89 (for pods per plant) (Table 3). Quban et al. (2010) also reported lowest phenotypic and environmental coefficient of variation for days to maturity in chickpea.

Deshmukh et al. (1986) classified PCV and GCV values as low (0-10%), moderate (10-20%) and high (>20%). Reliant on this suggestion, high GCV was observed for hundred seed weight, and moderate GCV was observed for seed per pod, pods per plant, and grain yield. The low value of a genotypic coefficient of variation (GCV) and phenotypic coefficient of variation (PCV) was noted for characters such as days to 50% flowering and days to maturity. Similar result was reported in numerous findings (Amin et al., 1992; Panwar & Singh, 2000; Bergale et al., 2001; Sharma et al., 2006; Dwivedi et al., 2008; Kumar et al., 2014).

The result of GCV expresses the true genetic potential of the genotypes. In the present study, the PCV values were higher than the corresponding GCV suggesting the existence of substantial environmental variation. This result was completely in agreement with the result of combined analysis of variance (ANOVA) table. However, traits such as grain filling duration, days to flowering and hundred seed weight, which
showed very small difference indicating the observed variations for these traits were mostly due to genetic factors with little environmental factors.

Table 3. Variance components, genotypic, environment, and phenotypic coefficients of variation, and heritability estimates of grain yield and other agronomic traits of sixteen chickpea genotypes grown at Shambu, Hawa Galan, Badesso, Mata and Alaku Belle in 2016/17

<table>
<thead>
<tr>
<th>Trait</th>
<th>σ²g</th>
<th>σ²gl</th>
<th>σ²e</th>
<th>σ²p</th>
<th>GCV (%)</th>
<th>ECV (%)</th>
<th>PCV (%)</th>
<th>H² (%)</th>
<th>GA</th>
<th>GAM</th>
</tr>
</thead>
<tbody>
<tr>
<td>Days to flowering</td>
<td>7.28</td>
<td>1.53</td>
<td>6.67</td>
<td>15.48</td>
<td>4.27</td>
<td>4.08</td>
<td>6.23</td>
<td>47.04</td>
<td>3.81</td>
<td>6.03</td>
</tr>
<tr>
<td>Days to maturity</td>
<td>0.98</td>
<td>2.29</td>
<td>6.39</td>
<td>9.65</td>
<td>0.71</td>
<td>1.83</td>
<td>2.24</td>
<td>10.11</td>
<td>0.65</td>
<td>0.47</td>
</tr>
<tr>
<td>Grain filling duration</td>
<td>12.01</td>
<td>4.96</td>
<td>13.02</td>
<td>29.99</td>
<td>4.6</td>
<td>4.79</td>
<td>7.27</td>
<td>40.05</td>
<td>4.52</td>
<td>5.99</td>
</tr>
<tr>
<td>Pods per plant</td>
<td>24.11</td>
<td>4.71</td>
<td>81.76</td>
<td>110.58</td>
<td>15.69</td>
<td>28.89</td>
<td>33.60</td>
<td>21.8</td>
<td>4.72</td>
<td>15.08</td>
</tr>
<tr>
<td>Seed per pod</td>
<td>0.06</td>
<td>0.00</td>
<td>0.06</td>
<td>0.12</td>
<td>19.91</td>
<td>19.91</td>
<td>27.77</td>
<td>51.43</td>
<td>0.37</td>
<td>29.84</td>
</tr>
<tr>
<td>Plant height</td>
<td>13.02</td>
<td>16.14</td>
<td>30.94</td>
<td>60.11</td>
<td>6.97</td>
<td>10.74</td>
<td>14.96</td>
<td>21.67</td>
<td>3.46</td>
<td>6.68</td>
</tr>
<tr>
<td>Branch per plant</td>
<td>0.04</td>
<td>0.03</td>
<td>0.69</td>
<td>0.71</td>
<td>5.07</td>
<td>19.92</td>
<td>20.18</td>
<td>6.31</td>
<td>0.11</td>
<td>2.64</td>
</tr>
<tr>
<td>Hundred seed weight</td>
<td>37.49</td>
<td>2.27</td>
<td>5.97</td>
<td>45.73</td>
<td>20.67</td>
<td>8.25</td>
<td>22.82</td>
<td>81.98</td>
<td>11.42</td>
<td>38.54</td>
</tr>
<tr>
<td>Grain yield</td>
<td>0.66</td>
<td>0.54</td>
<td>0.13</td>
<td>0.25</td>
<td>14.93</td>
<td>21.62</td>
<td>29.74</td>
<td>25.2</td>
<td>0.26</td>
<td>15.59</td>
</tr>
</tbody>
</table>

Key: \( \sigma^g \) = genotypic variance, \( \sigma^g l \) = variance due to g×l interaction, \( \sigma^e \) = error variance, \( \sigma^p \) = phenotypic variance, \( GCV \) = genotypic coefficient of variation, \( ECV \) = environmental coefficient of variation, \( PCV \) = phenotypic coefficient of variation, \( H^2 \) = heritability in broad sense, \( GA \) = genetic advance, \( GAM \) = genetic advance as the percentage of the mean.

Estimation of broad sense heritability, genetic advance (at 5% selection intensity) and the corresponding genetic advance as a percent of the mean were estimated for each measured character. Heritability estimates are classified as low (5-10%), medium (10-30%) and high (30-60%) (Dabholkar, 1992). The broad sense heritability (\( H^2 \)) values more than 80% was observed for hundred seed weight. This implies that the expected gain from selection would be high if this trait is used as selection criteria in chickpea breeding.

Traits having high heritability combined with genetic advance could result in a better genetic gain through selection since the variation that prevails in such a trait is due to additive gene action (Johnson et al., 1955; Vimal & Vishwakarma, 1998). Hundred seed weight, pods per plant, seed per pod and grain yield showed high heritability combined with high genetic advance as a percentage of the mean which could be used as a powerful tool in phenotypic selection as such characters are controlled by the additive genes and less influenced by the environment.

Genetic gain (GAM) that could be expected from selecting the top 5% of the genotypes as a percent of the mean, varied from 0.47% for days to maturity to 38.54% for hundred seed weight. According to Johnson et al. (1955), genetic advance as a percent of the mean was categorized as high (≥20%), moderate (10-20%) and lower (0-10%). Depending on this delineation, seed per pod and hundred seed weight showed high genetic advance as a percent of the mean, whereas pods per plant and grain yield showed moderate genetic advance as a percent of the mean. Therefore, selection based on traits with a high-level genetic advance as a percent of the mean will result in the improvement of the performance of the genotypes.

4. Conclusion

It can be concluded that direct selection can be done for most of the yield attributing traits since it exhibited high genetic variability and high range of variation.
Traits with high heritability and genetic advance will grant superior genotypes through phenotypic based selection.

Acknowledgments

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