Original Research Article

Genetic Diversity and Characterization of Pigeon Pea Germplasm

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A B S T R A C T

The study is planned to study variability, genetic divergence and characterization of pigeonpea germplasm at NARP, Agricultural Research Station, Navsari Agriculture University, Bharuch. Two genotypes i.e. NP 09-41 and NP-09-46 recorded white colour and the same genotypes are identified as high yielder too. While other genotypes are light, reddish or dark brown in colour These traits can be used for identifying individual germplasm. Ten genotypes were grouped into two major clusters. Five genotypes grouped under 1st cluster and the another five genotypes grouped under IInd cluster. NP-09-46, 43, 49, 47 and 50 falls under first cluster while NP-09-41, 48, 44, 45 and 42 falls under another cluster. Based on per se performance genotype viz NP-09-43, Np-09-41, NP-09-46 and NP-09-49 recorded highest yield per hector and medium duration genotype recorded as NP-09-45 and 47. Hence, it can be considered that high yielding genotypes are grouped under Its cluster while in second cluster medium yielding genotypes are grouped representing high genetic diversity among the germplasm.

Keywords
Pigeonpea, Germplasm, Genetic diversity, Yield

Introduction

Pigeonpea (Cajanus cajan (L.) Millspaugh) (2n= 22) is the second most important pulse crop of India after chickpea, commonly known as arhar, redgram and tur. Pigeonpea/redgram is a grain legume belonging to the Cajaninae sub-tribe of the economically important leguminous tribe Phaseoleae. Based on the natural genetic variability in local germplasm and the presence of numerous wild relatives, Van der Maesen (1990) concluded that India is probably primary centre of origin. Pigeonpea is an important pulse crop that performs well in semi-arid tropics where moisture availability is unreliable or inadequate (Reddy et al., 1993). India is the largest producer of pigeonpea in the world sharing approximately 70% of the production and covering 74% of the area. Pigeonpea has a unique place in Indian farming and India accounts for about 90% of the global production. It is the second most important pulse crop next to chickpea, covering an area of around 4.42 m ha (occupying about 14.5% of area under pulses) and production of 2.86 mt (contributing to 16% of total pulse production) and productivity of about 707 kg/ha (FAOSTAT, 2011). Although India leads the world both in
area and production of pigeonpea, its productivity is lower than the world average. One of the factors responsible for the poor productivity of pigeonpea is the lack of improved cultivars. Research for genetic improvement of this crop is required to raise yield levels effectively through widening genetic base. Information on genetic divergence among the plant materials is vital to a plant breeder for an efficient choice of parents for hybridization. It is an established fact that genetically diverse parents are likely to contribute desirable segregants and or to produce high heterotic crosses. Hence, this study is planned to study variability, genetic divergence and characterization of redgram germplasm at NARP, Agricultural Research Station, Navsari Agriculture University, Bharuch

### Materials and Methods

The experiments for the present study were conducted in the NARP, Agricultural Research Station, Navsari Agriculture University, Bharuch Kharif 2019. The experimental material comprised of 10 pigeonpea germplasm. The experiment was laid out in Randomised Block Design with two replications. Observations were recorded for eight quantitative traits. They are days to maturity (DM), plant height (PH) measured in cm, number of branches per plant (NBP), number of pods per plant (NBP), and seed yield kg per hectar (YPH). The morphological character i.e. colour of the seed coat was noted to distinguish the genotypes from each other. R version 3.1.3 package ‘gplots’ was used to draw heat map for diversity study. Hierarchical clustering and heatmap analysis were performed for mean-centered and standardized data in R. The ‘hcclus’ function was used for hierarchical clustering. Heatmap.2 function used to draw heatmap’ for given genotypes and traits analysed.

The genotype which matures within the range of 150-160 days is considered as medium duration genotype. The different variability parameters like mean, minimum, maximum is presented in table 1.

### Results and Discussion

In India, medium duration genotypes are mostly preferred among the pigeonpea growing farmers.

Days to maturity (DM), plant height (PH) measured in cm, number of branches per plant (NBP), number of pods per plant (NBP), and seed yield kg per hectar (YPH) For plant height the range was recorded from 109 cm to 161 cm, for number of branches it is varying from 14 to 22. In case of days to maturity it is 125 days to 185 days which means great variation among genotype like short duration NP-09-42 and NP-09-48 long duration is present. It is interesting to note that the average mean of genotype falls under medium duration. Two genotypes NP_09-47 and NP-09-45 falls under medium duration category in present study. The range for yield for hectar is 915 to 1523 kg/ha. Wide range for seed yield per hectar is recorded among the genotypes. The highest yield was recorded for genotype NP_09-43 followed by NP-09-41, NP-09-46 and NP-09-49 respectively. These genotypes proved good yielder than the others and can be utilized in future breeding programme. Only two genotypes i.e. NP-09-41 and NP-09-46 recorded white colour and the same genotypes are identified as high yielder too. While other genotypes are light, reddish or dark brown in colour. Similar results were being reported by Upadhyaya et al., (2007) and Manyasa et al., (2008) for growth habit, base flower color, pattern of streaks, pod colour, pod form and seed color pattern. These traits can be used for identifying individual germplasm. Other objective of the present study was to identify
the high yielding and medium duration genotypes through cluster analysis. Hierarchical clustering of genotypes based on the Euclidian distances for the eight traits is represented. Heat map approach gives an opportunity to visualize the expression of the each trait in particular the environments. Heat map and dendrogram displaying relatedness of genotypes and traits under study based on their Euclidean distances. Red and yellow correspond to low and high diversity for expressed traits. White colour represents median levels of expression. The color key indicates the correlation between diversity and colors. The histogram represents a distribution of each value under observation for particular trait. Based on the Ward’s clusters analysis (Squared Euclidean distance). The dendrogram of 10 genotypes of Pigeonpea is presented in Fig.1. Ten genotypes were grouped into two major clusters. Five genotypes grouped under Ist cluster and the another five genotypes grouped under IInd cluster.

Table.1 The mean performance of the genotype along with the Colour of the seed coat trait

<table>
<thead>
<tr>
<th>Genotypes</th>
<th>PH (cm)</th>
<th>NBP</th>
<th>DM</th>
<th>NPP</th>
<th>YPH(kg/ha)</th>
<th>Seed coat colour</th>
</tr>
</thead>
<tbody>
<tr>
<td>NP-09-41</td>
<td>149</td>
<td>22</td>
<td>169</td>
<td>238</td>
<td>1498</td>
<td>white</td>
</tr>
<tr>
<td>NP-09-42</td>
<td>119</td>
<td>14</td>
<td>125</td>
<td>251</td>
<td>915</td>
<td>Light brown</td>
</tr>
<tr>
<td>NP-09-43</td>
<td>161</td>
<td>19</td>
<td>165</td>
<td>357</td>
<td>1523</td>
<td>Dark brown</td>
</tr>
<tr>
<td>NP-09-44</td>
<td>154</td>
<td>18</td>
<td>168</td>
<td>272</td>
<td>1081</td>
<td>Reddish brown</td>
</tr>
<tr>
<td>NP-09-45</td>
<td>142</td>
<td>17</td>
<td>158</td>
<td>281</td>
<td>1185</td>
<td>Light brown</td>
</tr>
<tr>
<td>NP-09-46</td>
<td>159</td>
<td>19</td>
<td>172</td>
<td>384</td>
<td>1448</td>
<td>White Reddish</td>
</tr>
<tr>
<td>NP-09-47</td>
<td>112</td>
<td>16</td>
<td>158</td>
<td>387</td>
<td>1432</td>
<td>Brown</td>
</tr>
<tr>
<td>NP-09-48</td>
<td>158</td>
<td>20</td>
<td>185</td>
<td>285</td>
<td>913</td>
<td>Light brown</td>
</tr>
<tr>
<td>NP-09-49</td>
<td>149</td>
<td>17</td>
<td>178</td>
<td>363</td>
<td>1388</td>
<td>Brown Reddish</td>
</tr>
<tr>
<td>NP-09-50</td>
<td>109</td>
<td>16</td>
<td>165</td>
<td>342</td>
<td>1232</td>
<td>Brown</td>
</tr>
<tr>
<td>Mean</td>
<td>141</td>
<td>17</td>
<td>164</td>
<td>316</td>
<td>1264.5</td>
<td></td>
</tr>
<tr>
<td>Max</td>
<td>161</td>
<td>22</td>
<td>185</td>
<td>387</td>
<td>1523</td>
<td></td>
</tr>
<tr>
<td>Min.</td>
<td>109</td>
<td>14</td>
<td>125</td>
<td>238</td>
<td>915</td>
<td></td>
</tr>
</tbody>
</table>

Fig.1 The dendrogram of ten pigeonpea germplasm showing diversity among the pigeonpea genotypes
NP-0946, 43, 49, 47 and 50 falls under first cluster while NP-09-41, 48, 44, 45 and 42 falls under another cluster. Based on per se performance genotype viz., NP-09-43, NP-09-41, NP-09-46 and NP-09-49 recorded highest yield per hectar and medium duration genotype recorded as RG-187 and 190. Hence, it can be considered that high yielding genotypes are grouped under 1st cluster while in second cluster medium yielding genotypes are grouped representing high genetic diversity among the germplasm. Earlier in pigeonpea Thombre et al., (2000), Rekha et al., (2011), Praveen Pandey et al., (2013) and Rupika and Bapu (2014) studied genetic divergence either in among working germplasm and identified most divergent genotypes for the yield and yield attributing traits in redgram.

Use of diverse parents in hybridization programme can serve the purpose of combining desirable genes or to obtain recombination.

Crosses between divergent parents usually produce greater heterosis than those between closely related ones hence, identified genotypes can be utilised for the development of high yielding and medium duration varieties in pigeonpea.

References


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