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# Identification of High-yielding and Late Leaf Spot Disease-tolerant Families in Groundnut (*Arachis hypogaea* 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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#### ABSTRACT

The foliar fungal disease late leaf spot (LLS) caused by *Cercospora personata* is a major and widely distributed disease. Cause substantial yield loss and in combination with the diseases increased up to 70 *per cent* in India. Therefore, to reduce the effect of disease and yield penalty effective chemical control mainly depends on multiple fungicide applications which are costly for resource-poor farmers and also raise environmental and health concerns. Therefore, to reduce the cost of production the development of resistant cultivars is an eco-friendly concept, with this research gap study was conducted to identify groundnut families with high yield and resistance to late leaf spot disease. From 60 F<sub>7</sub> families of three crosses (GKVK-16 × KCG-2, GKVK-13 × KCG-2 and GKVK-6 × KCG-2 along with checks KCG-6, KCG-2 and TMV-2) which were evaluated in

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augmented design during *Kharif* 2017 in disease plot and control plot, disease screening was done using a modified 9-point scale. Disease scoring was done at 60, 90 and 120 days after sowing (pod filling stage). The disease scores were mainly based on the extent of leaf area damage. Results depicted that among the 60 families, 19 families showed consistence performance both at normal and disease plots. However, three families showed differences in the pod yield both at normal and disease plots. The families that exhibited resistance to LLS disease showed *per cent* yield reduction that ranged from 0-35%, and moderately resistant families exhibited a yield reduction from 35-50%, however, 50-90% yield reduction was observed in the families which showed moderate susceptibility, further more than 100% yield reduction was noticed in the families which are susceptible to LLS disease in three crosses of groundnut. High values of GCV and high PCV, high heritability coupled with high GAM were observed for PDI at 60<sup>th</sup>, PDI at 90<sup>th</sup> and PDI at 120<sup>th</sup> days after sowing in all three crosses. Among 14 superior families six, among 13 superior families eight, and from three superior families one family showed resistance to LLS disease from the cross GKVK-16×KCG-2, GKVK-13×KCG-2 and GKVK-6×KCG-2 respectively. Therefore, these identified families will be forwarded for muti-location disease and further yield stabilization.

Keywords: Groundnut; late leaf spot disease; per cent disease index; disease sore.

#### 1. INTRODUCTION

Groundnut (Arachis hypogaea L.) is a selfpollinated, cleistogamous annual herb belonging to the family Fabaceae or Leguminaceae with a chromosome number of 2n=40. It is the most important oil seed crop of India and the world in terms of area and production. It is vernacularly known as peanut, monkey nut, earth nut and pigmy nut. Cultivated groundnut is classified into two subspecies, subsp. fastigiata and subsp. hypogaea. The subsp. fastigiata contains four botanical varieties, var. vulgaris, var. fastigiata, var. peruviana, and var. aequatoriana. The subsp. hypogaea contain two varieties, var. hypogaea and var. hirsuta. Each of these botanical types has different plant, pod and seed characteristics [1].

There are many factors that affect the yield of the crop among them are the biotic factors mainly foliar fungal diseases like late leaf spot (LLS) caused by Cercospora personata is a major and widely distributed disease. In groundnutpredominant areas, foliar diseases like late leaf spot and rust cause substantial yield loss and in combination with the diseases increased up to 70 per cent in India [2]. The disease generally appears 55 to 60 days after sowing and can cause more than 50 % loss in yields in groundnut-producing areas of Karnataka (Oteng et al. 2023). Seventeen potential candidate genes were predicted at ± 300 kbp of the stable/prominent SNP positions (Oteng et al. 2023). To reduce the disease effective chemical control is mainly relied upon multiple fungicide applications Khangura et al. [3], which are costly for resource-poor farmers and also raise

environmental and health concerns. Therefore, the development of resistant cultivars to the diseases could be effective in decreasing production costs, improving production quality and reducing detrimental effects of chemicals and this is the only cost-effective and environmentally eco-friendly method to overcome the diseases. A total of 30 functional nucleotide polymorphisms or genic SNP markers were detected, among which eight genes were found to encode leucine-rich repeat (LRR) receptor-like protein kinases and putative disease resistance proteins [4]. There are many reasons which are attributed to low yield levels viz., lack of improved high yielding cultivars, cultivation under shallow soils of low fertility, uneven rainfall distribution, continuous cropping without rotation of crop, low plant population, non-availability of improved varieties and incidence of foliar diseases and pests are cited as the major limiting factors in most of the groundnut growing regions. Keeping all the above points in view, an attempt has been made to identify superior families performing better with less yield reduction under disease condition [5].

#### 2. MATERIALS AND METHODS

#### 2.1 Experimental Site and Plant Material

The present study was conducted at the experimental field, K-Block, Department of Genetics and Plant Breeding, GKVK, University of Agricultural Sciences, Bangalore located between  $13^{\circ}N$  latitude and  $77^{\circ}35$ ' E longitude and an altitude of 899 m above mean sea level (MSL), during *kharif* 2017. The details of the material used and the techniques adopted in the

present investigation for recording of observations and analysis of data are briefly presented in this chapter. The experimental material for the present study comprised of  $F_6$  and  $F_7$  families of three crosses *viz.*, GKVK-16 × KCG-2, GKVK-13 × KCG-2 and GKVK-6 × KCG-2 and checks KCG-6, KCG-2 and TMV-2.

#### 2.2 Late Leaf Spot Disease Screening

The  $F_6$  plants of three crosses, their parents and checks were raised in plant to progeny rows and evaluated in augmented design for late leaf spot disease during *kharif* 2017 (Fig. 1). The TMV-2 was used as spreader row for natural disease incidence. Screening of Late leaf spot disease was carried out by visual screening method Fig. 2 using modified 9-point scale for late leaf spot

(Table 1) given by Subrahmanyam et al. [2]. The scores were converted into Percentage Disease Index (PDI).

#### 2.3 Disease Scoring for Late Leaf Spot

For late leaf spot disease screening, visual screening (Fig. 2) and modified 9-point scale as given by Subrahmanyam et al. [2] (Table 1) was used. The visual scores (1-9) and the extent of leaf area destroyed (0-100 %) are linearly related to each other. Disease scoring was done at 60, 90 and 120 days after sowing (pod filling stage). The disease scores were mainly based on the extent of leaf area damage. The scores were converted into *Per cent* disease Index (PDI) by using the following formula.

PDI(%) =  $\frac{\text{Sum of the individual ratings}}{\text{No of plants assessed}} \times \frac{100}{\text{maximum disease rating}}$ 



Field view of control plot of F7 generation



Field view of disease plot of F7 generation



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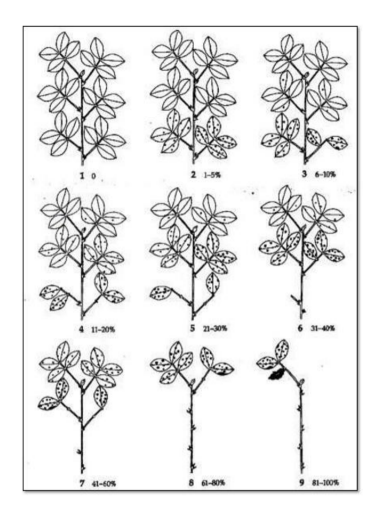


Fig. 2. Standard reference for scoring late leaf spot disease resistance [2]

Table 1. Modified 9-Point scale used for field-screening of groundnut genotypes for Late Leaf
Spot resistance [2]

Disease	Description	Disease severity %
score		-
1	No disease	0
2	Lesions present largely on lower leaves, no defoliation	1-5
3	Lesions present largely on lower leaves, very few on middle leaves; defoliation of some leaflets evident on lower leaves.	6-10
4	Lesions present on lower and middle leaves but severe on lower leaves, defoliation of some leaflets evident on lower leaves	11-20
5	Lesions present on lower and middle leaves, over 50 % of Defoliation of lower leaves	21-30
6	Severe lesions on lower and middle leaves; lesions present but less severe on top leaves; extensive defoliation of lower leaves; some defoliation on middle leaves	31-40
7	Lesions on all leaves but less severe on top leaves; defoliation of all lower and middle leaves	41-60
8	Defoliation of all lower and middle leaves; severe lesions on top leaves evident	61-80
9	Almost all leaves defoliated, leaving bare stem; some leaflets may remain, but show severe leaf spot	81-100

#### 3. RESULTS AND DISCUSSION

#### Pod 3.1 Effect of LLS Disease on Yield in $F_7$ Generation in the Cross the three Crosses of Groundnut

The 60 families which were evaluated both at normal and disease stress condition were compared each other to dissect out influence of the LLS disease at 90 DAS (days after sowing) on the pod vield, in three crosses and data presented in the Tables 2, 3 and 4. Results depicted in the tables indicated that among the 60 families 19 families showed consistence performance both at normal and disease plot, which indicated that LLS disease in the sick plot has less effect on the pod yield, therefore these families could be resistant for LLS disease However three families showed difference in the pod yield both at normal and disease plot, which confirms that the pod yield of these families at disease plot is less than the normal plot and per cent yield reduction at disease plot is very high. Further, PDI value of these families were also high. Therefore, these families could be susceptible for LLS disease [6].

The families exhibited resistant to LLS disease showed per cent yield reduction that from 0-35%, moderately resistant ranged exhibited vield families reduction from 35-50%, however 50-90% yield reduction was observed in the families which showed susceptibility, moderate further more than 100% yield reduction was noticed in the families which are susceptible to LLS disease in three crosses of groundnut (Survawanshi et al. 2023).

The PDI at  $60^{\text{th}}$ ,  $90^{\text{th}}$  and  $120^{\text{th}}$  days was estimated for the LLS disease, in 60 families of three crosses and presented in the Figs. 3, 4 and 5. The Fig. 3 indicated that the PDI values at  $120^{\text{th}}$  day were higher than the PDI at  $90^{\text{th}}$  and  $60^{\text{th}}$  day. Further some families like 6, 16, 18 and 22 exhibited lower PDI value at  $120^{\text{th}}$  day in cross GKVK - 16 × KCG – 2, suggest that these families were less effected by LLS disease spores even at 120 days after sowing therefore these families could be considered as resistance

families for LLS disease. The Fig. 4 indicated that the PDI values at  $120^{th}$  day were higher than the PDI at  $90^{th}$  and  $60^{th}$  day. Further some families like 5 and 15 showed lower PDI value at  $120^{th}$  day in cross GKVK -13 × KCG -2, indicates these families were less influenced by LLS disease spores during 120 days of cropping period therefore these families could be considered as resistance families for LLS disease. The Fig. 5 indicated that the PDI values at  $120^{th}$  day were higher than the PDI at  $90^{th}$  and  $60^{th}$  day in cross GKVK -6 × KCG -2, which indicates the families of this cross is more affected by LLS disease. Hence, families presented in this cross were susceptible for LLS disease [6].

#### 3.2 Estimates of Genetic Variability Parameters for LLS Disease in F<sub>7</sub> Generation in three Crosses of Groundnut

The amount of genetic variability for LLS disease in the F<sub>7</sub> generation was estimated by various genetic parameters viz., mean, range, genotypic coefficient of variation (GCV), phenotypic coefficient of variation (PCV), heritability  $(h^2)$  and genetic advance as per cent of mean (GAM) and presented in Table 5. There was a fluctuation in the mean values of PDI at 60<sup>th</sup>, PDI at 90<sup>th</sup> and PDI at 120<sup>th</sup> days after sowing in all three crosses of groundnut. High values of GCV and high PCV, high heritability coupled with high GAM were observed for PDI at 60<sup>th</sup>, PDI at 90<sup>th</sup> and PDI at 120<sup>th</sup> days after sowing in all three crosses. indicated the presence of a higher magnitude of variation for these traits. Narasimhulu et al. [7], Padmaja et al. [8]. This individual plant scoring can be practiced. Similar results were reported by Sun et al. [9] and Puppala et al. [10]. High heritability coupled with high GAM suggested characters additive that are under genetic control and selection will be effective [11]. According to Varshney et al. [12], disease score is the best selection criterion in the field for use in the breeding program due to high heritability and its measures, which is in accordance with the present results. Wankhade et al. [4] identified eight genes that were found to encode leucine-rich repeat (LRR) receptor-like protein kinases and putative disease resistance proteins [13].

SI. No.	Families	Pod yield per plant(g)under disease control condition	Pod yield per plant (g)under disease stress condition	<i>Per cent</i> yield reduction due to LLS disease	PDI of LLS disease@90DAS	LLS Disease response
1	C1-40-1-5-1-8	63.60	48.53	31.05	29.67	R
2	C1-213-6-2-4-1	28.48	12.45	128.76	65.00	S
3	C1-213-6-2-2-4	49.37	35.46	39.23	40.67	MR
4	C1-74-4-3-3-1	57.81	24.69	134.14	75.23	S
5	C1-47-9-10-1-3	36.45	32.45	12.33	27.33	R
6	C1-213-6-2-7-4	28.58	22.45	27.31	19.89	R
7	C1-40-1-4-2-12	28.33	15.67	80.79	47.52	MS
8	C1-74-4-3-3-4	32.78	11.03	197.19	75.89	S
9	C1-213-2-1-1-2	48.36	38.11	26.90	24.78	R
10	C1-46-1-10-1-4	20.45	13.11	55.99	50.18	MS
11	C1-213-6-2-4-9	36.26	24.22	49.71	44.00	MR
12	C1-213-6-2-7-1	47.48	35.25	34.70	45.21	MR
13	C1-40-1-4-5-1	34.43	20.00	72.15	50.48	MS
14	C1-47-9-10-1-1	56.24	38.11	47.57	36.58	MR
15	C1-46-1-10-1-9	58.91	33.64	75.12	46.28	MS
16	C1-74-4-3-3-2	63.60	48.33	31.60	04.39	R
17	C1-40-1-5-1-7	49.56	28.12	76.24	49.33	MS
18	C1-137-4-7-5-1	57.66	44.78	28.76	15.78	R
19	C1-43-3-8-3-1	37.93	26.89	41.06	36.00	MR
20	C1-213-6-2-7-2	62.28	35.44	75.73	45.89	MS
21	C1-213-2-8-4-16	61.79	45.98	34.38	27.60	R
22	C1-137-4-7-5-9	57.39	53.78	6.71	07.38	R
23	C1-213-2-1-3-1	34.41	20.33	69.26	38.12	MR
24	C1-137-4-7-5-3	45.52	25.56	78.09	49.77	MS

### Table 2. Effect of LLS disease on pod yield in F7 generation of the cross GKVK-16×KCG-2 in Groundnut

Note: PDI- Per cent disease index,

LLS- Late leaf spot

SI. No	Families	Pod yield per plant(g) under disease control condition	Pod yield per plant (g) under disease stress condition	<i>Per cent</i> Yield reduction due to LLS disease	PDI of LLS disease@90DAS	LLS Disease response	
1	C2-84-6-6-5-6	35.48	20.22	75.47	49.33	MS	
2	C2-84-6-6-5-5	39.11	24.23	61.41	40.36	MR	
3	C2-42-2-5-1-6	46.38	35.56	30.43	33.93	R	
4	C2-49-4-9-1-1	48.33	38.15	26.68	31.58	R	
5	C2-42-2-5-1-7	44.27	42.00	5.40	17.33	R	
6	C2-57-6-7-7-1	25.46	13.93	82.77	51.33	MS	
7	C2-49-2-5-2-2	50.78	45.22	12.30	13.22	R	
8	C2-63-3-1-1-2	30.50	19.79	54.12	42.22	MR	
9	C2-63-1-4-1-1	29.40	15.26	92.66	50.39	MS	
10	C2-49-4-9-1-4	32.90	20.39	61.35	42.61	MR	
11	C2-47-9-6-2-4	26.90	14.39	86.94	45.24	MS	
12	C2-84-6-1-2-1	30.00	22.16	35.38	25.44	R	
13	C2-70-5-2-3-1	24.00	13.65	75.82	51.78	MS	
14	C2-59-1-3-4-1	41.14	28.36	45.06	38.93	MR	
15	C2-75-5-6-1-3	33.56	26.78	25.32	8.11	R	
16	C2-59-1-3-4-5	39.20	30.11	30.19	25.39	R	
17	C2-47-9-2-1-1	29.00	19.33	50.03	37.47	MR	
18	C2-48-6-6-1-2	36.30	25.14	44.39	35.93	MR	
19	C2-59-1-3-4-2	38.54	23.14	66.55	45.33	MS	
20	C2-75-5-6-3-3	45.56	28.33	60.82	49.33	MS	
21	C2-70-5-2-6-3	40.78	36.23	12.56	17.28	R	
22	C2-49-2-7-3-8	26.24	13.50	94.37	50.22	MS	
23	C2-63-3-1-3-6	29.04	16.89	71.94	49.39	MS	
24	C2-63-3-5-6-7	18.33	14.39	27.38	21.89	R	

### Table 3. Effect of LLS disease on pod yield in F7 generation in the cross GKVK-13×KCG-2 of Groundnut

Note: PDI- Per cent disease index,

LLS- Late leaf spot

SI. No	Families	Pod yield per plant(g) @disease control condition	Pod yield per plant (g) @disease stress condition	Per cent Yield reduction due to LLS disease	PDI of LLS disease@90DAS	LLS Disease response	
1	C3-72-5-4-2-2	28.99	11.33	155.87	49.33	MS	
2	C3-88-4-5-1-5	35.21	34.00	03.56	38.93	MR	
3	C3-32-1-7-4-1	55.33	42.33	30.71	25.36	R	
4	C3-2-3-6-1-4	37.00	22.53	64.23	44.93	MR	
5	C3-62-2-10-2-4	36.32	18.41	97.28	20.38	R	
6	C3-88-4-5-1-3	48.33	32.97	46.59	41.80	MR	
7	C3-88-4-5-1-6	28.00	19.66	42.42	40.39	MR	
8	C3-32-1-7-4-6	30.00	22.50	33.33	37.07	MR	
9	C3-72-5-4-2-6	32.00	22.45	42.54	43.47	MR	
10	C3-130-5-6-2-4	49.00	25.33	93.45	46.23	MS	
11	C3-62-2-10-2-1	30.00	21.33	40.65	43.00	MR	
12	C3-40-1-4-1-7	25.00	11.00	127.27	49.33	MS	

#### Table 4. Effect of LLS disease on pod yield in F7 generation in the cross GKVK-6×KCG-2 of Groundnut

Table 5. Estimates of genetic variability parameters for LLS disease in F<sub>7</sub> generation in three cross of groundnut

Crosses	Days	Mean	Range		Standardized Range	GCV (%)	PCV (%)	h² <sub>(bs)</sub> %	GAM %
	-		Min	Max					
GKVK-16 × KCG-2	60DAS	13.55	6.32	20.45	1.04	12.36	17.08	72.36	12.36
	90DAS	39.71	10.52	45.52	0.88	23.52	28.93	80.36	25.52
	120DAS	67.25	15.63	88.56	1.08	36.45	51.73	70.45	31.78
GKVK-13 × KCG-2	60DAS	17.68	7.28	22.89	0.88	16.89	21.48	78.63	20.12
	90DAS	36.41	12.75	48.18	0.97	25.63	29.95	85.56	34.52
	120DAS	69.07	14.64	83.12	0.99	32.52	41.98	77.45	22.56
GKVK-6 × KCG-2	60DAS	16.44	4.25	18.22	0.84	25.36	33.61	75.45	13.45
	90DAS	40.01	11.74	42.75	0.77	33.52	50.36	66.56	23.45
	120DAS	71.45	16.28	87.12	0.99	42.52	53.74	79.12	27.45

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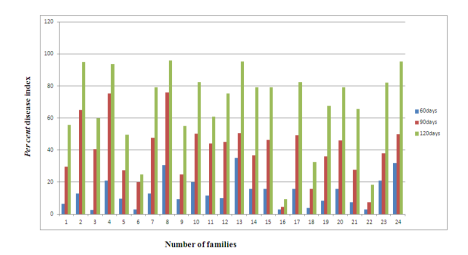
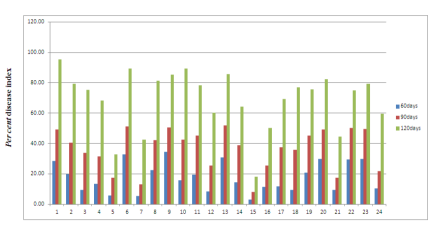


Fig. 3. Rate of infection of LLS disease at different DAS during cropping period in F<sub>7</sub> generation of the Cross GKVK-16×KCG-2



Number of families

Fig. 4. Rate of infection of LLS disease at different DAS during cropping period in F<sub>7</sub> generation of the Cross GKVK-13×KCG-2

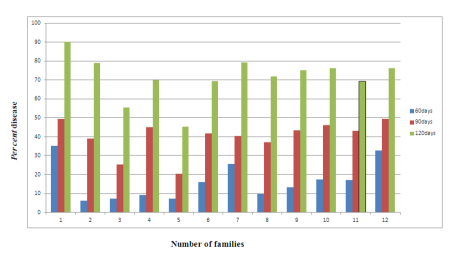


Fig. 5. Rate of infection of LLS disease at different DAS during cropping period in F<sub>7</sub> generation of the Cross GKVK-6×KCG-2

#### 3.3 Superior Selected Families for LLS Disease in the F<sub>7</sub> Generation in the Three Crosses of Groundnut

From among the 14 superior families six families (C1-213-2-1-1-2, C1-74-4-3-3-2, C1-213-2-8-4-16, C1-40-1-5-1-8, C1-137-4-7-5-9, C1-137-4-7-5-1) showed resistance to LLS disease, three (C1-213-6-2-2-4, C1-213-6-2-7-1, C1-47-9-10-1-1) were moderately resistant to LLS disease, four (C1-213-6-2-7-2, C1-40-1-5-1-7, C1-137-4-7-5-3, C1-46-1-10-1-9) were moderately susceptible and C1-74-4-3-3-1 showed susceptible to LLS disease in the cross GKVK-16×KCG-2. From among the 13 superior families eight families (C2-42-2-5-1-6, C2-49-4-9-1-1, C2-42-2-5-1-7, C2-59-1-3-4-5, C2-70-5-2-6-3, C2-49-2-5-2-2, C2-75-5-6-1-3. C2-42-2-5-1-6) showed resistance to LLS disease. three (C2-84-6-6-5-5. C2-48-6-6-1-2, C2-59-1-3-4-1) were moderately resistant to LLS disease, three (C2-59-1-3-4-2, C2-84-6-6-5-6, C2-75-5-6-3-3) were moderately susceptible to LLS disease in the cross GKVK-16×KCG-2. From among the three superior families. family C3-32-1-7-4-1 showed family C3-88-4-5-1-3 showed resistance. moderate resistance, and family C3-130-5-6-2-4 showed moderately susceptible to LLS disease in the cross GKVK-6×KCG-2.

#### 4. CONCLUSION

In the present investigation, 60 families were evaluated in the augmented design along with three checks to identify superior families for LLS disease. Results depicted that among the 60 families, 19 families showed consistence performance both at normal and disease plots. However, three families showed differences in the pod yield both at normal and disease plots. High values of GCV and high PCV, high heritability coupled with high GAM were observed for PDI at 60<sup>th</sup>, PDI at 90<sup>th</sup> and PDI at 120<sup>th</sup> days after sowing in all three crosses. Among 14 superior families six, among 13 superior families eight, and from three superior families one family showed resistance to LLS disease from the cross GKVK-16×KCG-2, GKVK-13×KCG-2 GKVK-6×KCG-2 and respectively. Therefore, these identified families will be forwarded for multi-location disease and further yield stabilization.

#### **COMPETING INTERESTS**

Authors have declared that no competing interests exist.

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