Effect of Different Sowing Dates on Pest Incidence in Chickpea

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A B S T R A C T

Global warming and climate change will have a major bearing on pest incidence and pest associated losses in field crops. Therefore, we studied pest incidence in chickpea across sowing dates to understand the effect of climatic factors on pest incidence on five genotypes of chickpea. The egg laying by the pod borer, Helicoverpa armigera decreased across sowing dates from October to December, with a slight increase in oviposition was observed in the January sown crops. ICC 3137 was most preferred for egg laying (9.5 eggs/5 plants), followed by KAK 2 (6.8 eggs/5 plants). The incidence of H. armigera decreased with a delay in time of sowing (60.0 larvae/5plants in the October sown crop to 21.9 larvae/5plants in the December sown crop). However, a slight increase was observed in the January sown crop (34.8 larvae/5plants). The highest incidence of H. armigera larvae was recorded on ICC 3137 (55.1 larvae/5plants), and the lowest on ICCV 10 (29.9 larvae/5plants).

Keywords
Chickpea, Climate change, Pest incidence, Helicoverpa armigera, Spodoptera exigua, Campoletis chlorideae

Introduction

Chickpea (Cicer arietinum L.) also known as Bengal gram or gram, is the second most important food legume in Asia, North Africa, and Mexico. Recently, it has also become an important grain legume crop in North USA, Canada, and Australia. It is grown on 13.5 million hectares worldwide, with an average production of 8.8 million tonnes. India is the largest producer of chickpea in the world sharing 71.0 and 67.2% of the total area (9.6 m ha) and production (8.8 mt), respectively (FAOSTAT, 2013). Several biotic and abiotic constraints limit the production and productivity of chickpea, of insect pests are a major constraint to increase the production and productivity of chickpea (Sharma 2005
and Yadav et al., 2006; Sharma et al., 2011). Losses due to insect pest damage are likely to increase as a result of changes in cropping patterns, and global warming.

The pod borer, *Helicoverpa armigera* (Hubner), is one of the most important constraints in chickpea production (Sharma, 2005). Its population peaks generally correspond to the full bloom and pod formation stage of the crop in the post rainy season. Temperature, relative humidity (Yadava and Lal 1988, Yadava et al., 1991), rainfall (Tripathi and Sharma 1985), predators (Thakur et al., 1995, Gunathilagaraj 1996) and parasitoids (Bhatnagar 1980, Srinivas and Jayaraj 1989, Thakur et al., 1995) affect the incidence and population densities of *H. armigera* on chickpea. Information on pest incidence under field conditions across sowing dates can be used to assess the effect of different climatic variables on pest incidence and grain yield. Therefore, we studied the effect of climatic factors on pest incidence and grain yield on five genotypes of chickpea.

**Materials and Methods**

Five chickpea genotypes (2 resistant - ICCL 86111 and ICCV 10, 2 commercial cultivars - JG 11 and KAK 2, and 1 susceptible genotype - ICC 3137) were sown across four planting dates between October - January at monthly intervals during 2012 - 14 post rainy seasons under field conditions. The experiment was laid out in randomized complete block design (RCBD) with three replications for each genotype, in a plot of four rows 2 m long (with a spacing of 60 cm between the rows and 10 cm between plants with in a row). Data were recorded on numbers of insects/plant. at fortnightly intervals in each planting. Data were also recorded on leaf feeding (leaf damage rating on a 1 to 9 scale (1 = <10% leaf area damaged, and 9 = > 80% leaf area damaged) (Sharma et al., 2005). The incidence/abundance of different insect pests was correlated with the climatic factors (average temperature, open pan evaporation, rainfall, sunshine hours, solar radiation, wind velocity, and relative humidity during the observation period). The crop was raised under normal agronomic practices, and there was no insecticide application in the experimental plots.

Weather data during the experimental period was obtained from the agro meteorology station at the ICRISAT farm. Data on rainfall, temperature, relative humidity, open pan evaporation, sunshine hours, solar radiation and wind velocity during the experimental period was correlated with lead damage, and egg and larval density (Incidence) during the experimental period.

**Results and Discussion**

**Oviposition by *H. armigera* females on different genotypes of chickpea**

There were significant differences in the numbers of *H. armigera* eggs across different dates of sowing in both the seasons, as well as across the seasons. The egg laying by the *H. armigera* females decreased as the sowing dates advanced from October to December (19.9 – 5.2 eggs/5 plants in 2012/13; 9.2 – 3.7 eggs/5 plants in 2013/14 and 13.9 – 4.3 eggs/5 plants across the seasons), but a slight increase in oviposition was recorded in the January sown crop (5.9 eggs/5 plants in 2012 –13, 4.3 eggs/5 plants in 2013 – 2014, and 5.1 eggs/5 plants across the seasons). More number of eggs were recorded in 2012 –13 than in 2013 –14. Highest numbers of eggs were observed in the crop sown in October in both the seasons.

There were significant differences in oviposition on different genotypes across sowing dates, and the interaction effects were nonsignificant. Among the genotypes tested,
ICC 3137 had the highest number of eggs across the seasons (11.3 eggs/5 plants, in 2012 - 13; 7.7 eggs/ 5 plants in 2013 - 14 and 9.5 eggs/5 plants across the seasons), while the oviposition was recorded on JG 11 (6.3 eggs/ 5 plants) in 2012 – 13, and on ICCV 10 and ICCL 86111 (3.5 eggs/ 5 plants) in 2013 - 14. Across seasons, ICC 3137 was most preferred for egg laying (9.5 eggs/5 plants), followed by KAK 2 (6.8 eggs/5 plants). ICCV 10 and JG 11 (5.9 eggs/5 plants) were relatively non-preferred for egg laying (Fig. 1).

**Variation in density of *H. armigera* larvae on different genotypes of chickpea across sowings**

The incidence of *H. armigera* larvae was highest in the crop sown in October (80.7 larvae/5plants), and lowest in the December sown crop (20.1 larvae/5plants) in 2012 – 13. In the 2013 – 14 cropping season, the incidence of *H. armigera* was quite high in the crop sown in November (40.7 larvae/5plants), October (39.3 larvae/5plants) and January (38.3 larvae/5plants), but low in the December sown crop (23.8 larvae/5 plants). Across seasons, the incidence of *H. armigera* declined as the sowing date was advanced from October (60.0 larvae/5plants) to December (21.9 larvae/5plants), but increased in the January sown crop (34.8 larvae/5 plants).

There were significant differences in numbers of *H. armigera* larvae across genotypes in both the seasons, but the interaction effects were nonsignificant. Highest number of *H. armigera* larvae were recorded on ICC 3137 (51.9 larvae/5plants), followed by KAK 2 (46.6 larvae/5plants) and ICCL 86111 (41.8 larvae/5plants). The lowest incidence of *H. armigera* larvae was recorded in ICCV 10 (28.2 larvae/5plants), followed by JG 11 (38.3 larvae/5plants). In 2013 – 14 post rainy seasons, the *H. armigera* larval density was significantly higher on ICC 3137 (58.3 larvae/5plants) and KAK 2 (37.9 larvae/5plants) than on ICCV 10 (31.7 larvae/5plants), JG 11 (30.1 larvae/5plants and ICCL 86111 (24.7 larvae/5plants). Across seasons, highest incidence was recorded on ICC 3137 (55.1 larvae/5plants), and the lowest on ICCV 10 (29.9 larvae/5plants). The larval density decreased from October to December, but a slight increase was observed in the crop sown in January. Across seasons, lowest larval density was recorded on ICCV 10 (15.5 larvae/5plants) in the December sown crop, and highest on ICC 3137 (84.6 larvae/5plants) in the October sown crop (Fig. 2).

**Oviposition by beet armyworm, *S. exigua* on different genotypes of chickpea**

There were no significant differences in the numbers of *S. exigua* egg masses across the sowings in the 2012 - 13 cropping season. No egg masses were observed in the October sown crop in 2012 - 13. Highest egg laying was recorded in the January sown crop (0.4 egg masses/5 plants). The number of egg masses differed significantly across sowing dates in the 2013 - 14 cropping season. In 2013 - 14, significantly highest numbers of egg masses were recorded in the December sown crop (1.3 egg masses/5 plants), but the differences in egg laying were nonsignificant in the crops sown in October, November and January. Similar trend was observed across seasons. The highest numbers of egg masses were recorded in the December sown crop (0.7 egg masses/5 plants), and greater egg laying was recorded in 2013-14 than in 2012 - 13 cropping season.

No egg laying was observed on ICCL 86111, while a fewer egg masses were recorded on ICCV 10 (0.3 egg masses/ 5plants) in the January sown crop, and in JG 11 in the November and January sown crops. The number of egg masses deposited on different genotypes differed significantly during the
2013 - 14 cropping season, and highest number of egg masses (1.7 egg masses/5 plants) were recorded on KAK 2, while no eggs were recorded in ICCV 10. Across seasons, highest number of S. exigua egg masses (1.0 egg masses/5 plants) were recorded on KAK 2, followed by ICC 3137 (0.4 egg masses/5 plants) and ICCL 86111 (0.4 egg masses/5 plants). The interaction effects were non – significant across the seasons. No egg masses were recorded in the October sown crop in both the seasons, except on KAK 2 in the 2013 – 14 cropping season (Fig. 3).

Population of beet armyworm, S. exigua larvae on different chickpea genotypes

In the 2012 – 13 cropping season, the numbers of S. exigua larvae were highest in the crop sown in January (16.1 larvae/5 plants), followed by the December (11.6 larvae/5 plants), November (10.1 larvae/5 plants) and October (4.7 larvae/5 plants) sown crops. During the 2013 – 14 cropping season, the numbers of S. exigua larvae were significantly higher in the crop sown in January (15.5 larvae/5 plants), followed by the December sown crop (11.6 larvae/5 plants). Significantly lower larval population was recorded in the November (1.3 larvae/5 plants) and October (2.0 larvae/5 plants) sown crops. Across the seasons, the S. exigua incidence was significantly greater in the January sown crop (15.8 larvae/5 plants) than in the crops sown in October, November and December. The January sown crop was most affected by S. exigua larvae in both the cropping seasons, as the crop grew and matured during the warm months of February to May. The larval incidence was comparatively greater in the 2013 - 14 than in 2012 – 13 cropping season.

There were no significant differences in the numbers of S. exigua larvae on different genotypes in the 2012 – 13 cropping season. KAK 2 had the maximum numbers of S. exigua larvae (15.6 larvae/5 plants), followed by ICCL 86111 (11.6 larvae/5 plants), JG 11 (9.3 larvae/5 plants) and ICC 3137 (8.8 larvae/5 plants). Less S. exigua larval numbers were recorded on ICCV 10 (7.8 larvae/5 plants). During the 2013 – 14 cropping season, there were no significant differences among the genotypes tested. However, the highest numbers of S. exigua larvae were observed on JG 11 (12.1 larvae/5 plants), followed by ICC 3137 and ICCL 86111 (5.1 larvae/5 plants). Across seasons, the highest numbers of S. exigua larvae were recorded on KAK 2 (12.9 larvae/5 plants) and lowest on ICC 3137 (7.0 larvae/5 plants).

The interaction effects between the genotypes and sowing dates were not significant. The lowest (2.5 larvae/5 plants) incidence was recorded in ICCV 10 in the November sown crop, and highest in KAK 2 in the January sown crop (27.2 larvae/5 plants). Highest numbers of egg masses were also recorded on KAK 2 – Kabuli type genotype, suggesting that it is highly susceptible to S. exigua. KAK 2 was found to be highly susceptible to S. exigua, while ICC 3137 was highly susceptible to H. armigera. ICCV 10 was relatively resistant to both H. armigera and S. exigua. The S. exigua incidence was observed mostly in the early stages of the crop, irrespective of the planting dates (Fig. 4).

Variation in parasitization of H. armigera by the larval parasitoid Campoletis chlorideae

During the 2012 – 13 cropping season, greater numbers of cocoons of C. chlorideae were observed in the December sown crop (3.4 cocoons/5 plants), followed by the October sown crop (2.4 cocoons/5 plants). Lowest parasitization (0.1 cocoons/5 plants) were
recorded in the January sown crop. In the 2013 – 14 cropping season, maximum parasitization (5.7 cocoons/5 plants) was recorded in the October sown crop, and the lowest (0.4 cocoons/5 plants) in the January sown crop. Across seasons, highest (4.0 cocoons/5 plants) activity of the parasitoid was recorded in the October sown crop, and the lowest (0.2 cocoons/5 plants) in the January sown crop, suggesting that the parasitoid is mostly active during the cooler part of the winter season.

Fig.1 Oviposition by *H. armigera* females on different genotypes of chickpea in relation to temperature and RH under natural infestation in the field

![Graph of H. armigera eggs](image)

**H. armigera eggs**

- ICC 3137
- ICCL 86111
- ICCV 10
- JG 11
- KAK 2
- Temp (°C), Max
- Min Temp (°C)
- RH 1 (%)
- RH 2 (%)

![Graph of H. armigera larvae](image)

**H. armigera larvae**

- ICC 3137
- ICCL 86111
- ICCV 10
- JG 11
- KAK 2
- Temp (°C), Max
- Min Temp (°C)
- RH 1 (%)
- RH 2 (%)

Fig.2 Abundance of *H. armigera* larvae on different genotypes of chickpea in relation to temperature and RH under natural infestation in the field
**Fig. 3** Oviposition by *S. exigua* females on different genotypes of chickpea in relation to temperature and RH under natural infestation in the field

![Graph showing oviposition by *S. exigua* females](image1)

**Fig. 4** Abundance of *S. exigua* larvae on different genotypes of chickpea in relation to temperature and RH under natural infestation in the field

![Graph showing abundance of *S. exigua* larvae](image2)
There were no significant differences in the numbers of *C. chloridea* cocoons on different genotypes in both the seasons. However, highest numbers of cocoons were recorded on ICC 3137 (2.6 cocoons/5 plants), and the lowest on KAK 2 and JG 11 (2.0 cocoons/5 plants). The interaction effects were not significant (Fig. 5).

In the early sown crop, which developed and matured during the cooler part of the post rainy season, there were significant differences in genotypic susceptibility to pod borer damage, but the differences between the genotypes were less apparent in *H. armigera* larvae in the late sown crops. Though the numbers of *H. armigera* larvae decreased with the planting dates, the extent of damage by *H. armigera* increased across the planting dates in both cropping seasons, which could be ascribed to warmer conditions during crop development and maturity. Parasitization of *H. armigera* larvae by *C. chloridea* also decreased with the planting dates, resulting in a decreased in biological control of *H. armigera* larvae, and increased crop damage. Damage by *H. armigera* increased with an increase in temperature as a result of reduction in the dry matter and grain yield.

Shankar et al., (2014) reported that numbers of *S. exigua* and *H. armigera* larvae were maximum on ICC 3137 at the vegetative, flowering and podding stages in both the seasons, while ICCL 86111 harboured the lowest numbers of *H. armigera* and *S. exigua* larvae. More *H. armigera* moths were trapped during March to April (Mahapatra et al., 2007), and November sown crops suffered less pod damage than that sown in December (Prasad et al., 1989; Begum et al., 1992). Delayed sowing of chickpea is risky under rainfed conditions due to inadequate stored soil moisture, and increased risk of damage by *H. armigera*. (Prasad and Singh 1997). Oviposition by *H. armigera* was low in the crop sown between December to Mid-February due to cold conditions in Pakistan (Shah and Shahzad, 2005), whereas Ali et al., (2009) observed that the numbers of eggs laid
by \textit{H. armigera} differed significantly across sowings on different genotypes of cotton, but there were no significant differences in larval density and damage across genotypes and sowing dates.

The \textit{H. armigera} larval population was high in early sown crops (October 15\textsuperscript{th} to November 1\textsuperscript{st}) than in and delayed sowings (November 1\textsuperscript{st} to 30\textsuperscript{th}) (Anwar et al., 1994). The genotypic response to damage by \textit{H. armigera} varies across seasons and locations (Sharma et al., 2003). The genotypes (ICC 506EB, ICC 12476, ICC 12477, ICC 12478 and ICC 12479) that are not preferred for oviposition also suffer low leaf damage by \textit{H. armigera} (Narayanamma et al., 2007).

The abundance of \textit{H. armigera} decreased with an increase in temperature, but plant damage increased with a rise in temperature. This may be due to better plant growth in early sowings than in the late sown crops due to inadequate soil moisture and dry weather conditions, which retarded the plant growth, with less pod setting, and consequently resulting in poor grain yield. The vegetative growth and the dry matter production decreased with an increase in temperature due to water stress.

The numbers of \textit{C. chlorideae} cocoons decreased with an increase in temperature. Higher temperatures resulted in reduced efficacy of control agents of \textit{H. armigera}, which may also have contributed to increase in plant damage. Patnaik and Senapati (1996) observed a negative correlation between mean temperature range and larval incidence of \textit{H. armigera}. A positive association was observed between \textit{H. armigera} and \textit{S. exigua} larvae, and similar results were earlier reported by Sharma (2012b). Positive correlation has earlier been observed between \textit{H. armigera} larval incidence and the maximum and the minimum temperatures (Sharma et al., 2005.; Upadhyay et al., 1989; Pandey, 2012). Ugale et al., (2011) reported that moth emergence was negatively correlated with the maximum (r = -0.62) and minimum temperature (r = -0.75), but there was no association with relative humidity. Minimum temperature and rainfall exerted a negative influence on pheromone trap catches of \textit{H. armigera} (Prasad et al., 1989). The population of \textit{H. armigera} and \textit{S. exigua} larvae was negatively correlated with relative humidity across genotypes. However, a significant and negative correlation has earlier been reported between \textit{H. armigera} larval density and maximum relative humidity (Sharma et al., 2005; Upadhyay et al., 1989; Pandey, 2012 and Shah and Shahzad, 2005).

Densities of eggs and of different larval instars of \textit{H. armigera} were significantly and negatively correlated with the maximum relative humidity, but not with the minimum relative humidity. Extremes of temperature, humidity and other weather factors (e.g., wind and hailstorm) might result in mortality of eggs, larvae and pupae of most of insect species (Pearson, 1958 and Qayyum and Zalucki, 1987). Pest outbreaks are more likely to occur with stressed plants as a result of weakening of plants' defensive system, and thus, increasing the level of susceptibility to insect pests. Global warming will lead to earlier infestation by \textit{H. armigera} in North India (Sharma, 2010a), resulting in increased crop loss. Climate change may also alter the interactions between the insect pests and their host plants (Sharma, 2014). Relationships between insect pests and their natural enemies will change as a result of global warming, resulting in both increases and decreases in the status of individual pest species. Changes in temperature will also alter the timing of diurnal activity patterns of different groups of insects and changes in inter specific interactions could also alter the effectiveness of natural enemies for pest management (Hill and Dymock, 1989).
Global warming and climate change will influence survival, development and population dynamics of *H. armigera*, and this will have a major bearing on extent of crop losses, and timing of different components of pest management to minimize the losses due to this pest. Future studies should focus on simultaneously testing the effects of multiple environmental factors on insect-plant interactions, to gain a realistic perspective of how global climatic changes may impact the production of secondary chemicals and its potential implications for co evolutionary associations between the interacting plant and insect species.

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**References**


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