Assessment of phenological and biochemical basis of resistance induced by organic, conventional and conservation practices in groundnut (*Arachis hypogaea* L.) to *Spodoptera litura*

B Thirupam Reddy, RS Giraddi and DN Kambrekar

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**Abstract**

The studies on biophysical factors at 60 DAS of groundnut revealed lower leaf succulency (72.37\%) and higher leaf thickness (6.52 mg/cm\(^2\)) imparted induced resistance against pest infestation in plants receiving organic nourishment compared to conventional and conservation practices. The biochemical constituent, phenol recorded higher amount (0.53 mg/g fr.wt at 60 DAS) in plants nourished with organics resulting into lower pest incidence owing to induced resistance. The lower amounts of sugars, free amino acids, soluble proteins and prolines also imparted resistance to the plants receiving organics whereas higher quantities recorded in conventional practices resulted in susceptibility to pests. Morphological characters like plant height, leaf area index, etc. were enhanced in organically nourished plants.

**Keywords:** Groundnut, *Spodoptera litura*, leaf thickness and phenol

**Introduction**

Nutrient management plays a vital role in the growth and yield of crops. Chemical fertilizers are the major source of nutrients, but its escalating cost coupled with increasing demand and adverse effect on soil health have made them a problematic resource so far. Indiscriminate use of pesticides for management of various pests and diseases also has lead to pesticide resistance, pest resurgence, secondary pest outbreaks, destruction of beneficial insects and other natural fauna, accumulation of undesirable residues resulting in environmental pollution and severe health hazards. This necessitated for adoption of eco-friendly farming with efficient use of organics and biologicals in crop production.

Groundnut is an important oilseed crop in the World and is an important source of digestible proteins, cooking oil and also vitamins (Savage and Keenan, 1994) [26]. Insect pests are important constraints in achieving higher groundnut yields. In India, the annual yield losses by insect pests in groundnut are about 15 per cent which accounts for about 1.6 million tonnes and 25.27 billion rupees (Dhaliwal *et al.*, 2010) [26]. Among the insects, polyphagous pest *S. litura* (Lepidoptera: Noctuidae) is an important pest of groundnut besides affecting tobacco, cotton, pulses and several vegetables crops (Singh and Jalali, 1997) [29]. It has been reported that an infestation level of one larva per plant during the seedling or flowering stage resulted in 20\% yield loss (Dhir *et al.*, 1992) [9]. Severe outbreak of the pest results in 30–40\% loss in pod formation (Joshi 2005). In India, transitional tract of Karnataka (Dharwad) has been identified as hot spot for *S. litura* during kharif season, where yield loss to an extent of 66.6 per cent was reported in groundnut (Kulkarni, 1989) [16]. Larvae feed gregariously on leaves and fresh growth causing extensive damage (Patil, 2000) [20]. Though many effective insecticide molecules are suggested to combat *Spodoptera*, they are not eco-friendly and add to the cost of cultivation especially in semi-arid tropics where farmers Grow groundnut as subsistence crop. This paper is mainly focus on impact of organic nutrients induced resistance in groundnut to *S. litura*. Organic practices enhance and maintain soil organic carbon status for obtaining sustainable crop yields. The increasing complexities of insect pest management and current discernment of problems associated with chemical control of insect pests, have led to an
Plant succulence was expressed as relative water content (RWC). A composite sample of leaf discs was taken and the fresh weight was determined, followed by flotation on water for up to 4 hrs. The turgid weight was then recorded and the leaf tissue is subsequently oven-dried to a constant weight at about 85°C and dry weight was taken as described by Barrs and Weatherley (1962). RWC was calculated by using the formula:

\[ RWC = \frac{\text{Fresh weight} - \text{dry weight}}{\text{Turgid weight} - \text{dry weight}} \times 100 \]

Leaf area

Leaf area was measured by disc method as suggested by Vivekanandan et al. (1972). Leaflets were separated from petiole and fifty discs of known size were prepared through a cork borer from randomly selected fifty leaves from five plants. The discs and the remaining parts of the 50 leaves as well as the remaining leaves were oven dried at 75 to 80°C to a constant weight and leaf area was calculated by using the formula given below.

\[ \text{LA} = \frac{\text{Wa} \times A}{\text{Wd}} \]

LA- Leaf area of five plants in cm²
Wa- Oven dry weight of all the leaves (inclusive of 50 discs) in g
Wd- Oven dry weight of 50 discs in g
A- Area of 50 discs in cm²

Leaf thickness

Leaf thickness was expressed in specific leaf weight (SLW).

\[ \text{SLW (mg/cm}^2) = \frac{\text{Leaf weight per plant (mg)}}{\text{Leaf area per plant (cm}^2)} \]

Material and Methods

Collection of Experimental Data on Biophysical Parameters

Five plants were selected randomly from each plot for recording observations on various biophysical parameters at 60 DAS in groundnut crop in different farming systems.

Extraction of Plant Tissues in Alcohol

The estimation of metabolites requires their complete extraction from tissues. The activities of the enzymes, which synthesize and utilize them, need to be stopped at once to get reliable values. The plant constituents possess different solvents. Though water is the universal solvent, it does not penetrate tissues quickly enough to stop enzymatic activity. In this context alcohol especially hot alcohol, is the choicest solvent for extraction.

Clarification of Alcoholic Extracts

Dark coloured alcohol extracts of the tissues create a great problem in analytical procedure. Heavy metal salts were used for clarification of alcoholic extracts.

Reagents and Procedure

Saturated solution of neutral lead acetate and saturated solution of disodium hydrogen phosphate were used. Two ml of saturated lead acetate solution was added drop wise to 10 ml of the coloured alcoholic extract and saturated solution of di-sodium hydrogen phosphate was added drop wise till the precipitation is completed. The above solutions were mixed thoroughly and kept overnight and filtered through Whatmann No. 1 filter paper and made up to 25 ml with alcohol. The extract was stored in a refrigerator at 4°C. This alcoholic extract was used further for analysis of reducing sugar, non-reducing sugar, total sugar and phenols.

Estimation of Total Phenols

Estimation of total phenols present in plant samples was determined following Folin-Ciocalteau Reagent (FCR) method (Bray and Thorpe, 1954) [5].

Reagents and Procedure

Folin-Ciocalteau reagent (1%) and Sodium carbonate (2%) reagents were used. One ml of each alcohol extract was taken in a test tube to which one ml of folin-ciocalteau reagent followed by two ml of sodium carbonate solution (2%) were
added. The tubes were shaken well and heated in a hot water bath for exactly one minute and then cooled under running tap water. The blue coloured complex developed was diluted to 15 ml with distilled water and its absorbance was read at 650 nm in a spectrophotometer. The amount of phenols present in sample was calculated from 3.63 Estimation of sugars. The estimation of total sugars and reducing sugars was done by following Nelson-Somogyi’s method. The reducing sugar was estimated following Nelson’s modification of Somogyi’s method (Nelson, 1944) [17].

Reagents Used
1. Solution A
Twenty five grams of anhydrous sodium carbonate, 25 g of sodium potassium tartrate, 20g of sodium bicarbonate and 200 g of sodium sulphate were dissolved separately in distilled water and volume was made upto one litre.

2. Solution B
Fifteen gram of copper sulphate was dissolved in distilled water to which one or two drops of concentrated sulphuric acid were added and made up to 100 ml volume with distilled water. Solutions A and B were mixed in 24:1 (v/v) proportion just before use.

3. Arsenomolybdate reagent
Twenty five gram of ammonium molybdate was dissolved in 450 ml of distilled water. Twenty one ml of concentrated sulphuric acid was added and mixed with above solution. Three gram of sodium orthoarsenate was dissolved in 25 ml distilled water. Both the solutions were mixed with stirring and placed in an incubator at 37 °C for 24-48 hrs. The reagent was stored in an amber coloured bottle.

Procedure
One ml of each sample (alcohol extract) was pipetted out into test tubes. To each one ml of extract, one ml of mixture of solution A and B was added and test tubes were heated on a hot water bath for 20 min. After cooling under running tap water, one ml of arsenomolybdate reagent was added with immediate mixing. The volume of the above solution was made up to 15 ml, to dilute the density of the blue coloured complex. The absorbance of the solution was read at 510 nm in a spectrophotometer. The amount of reducing sugars was determined by using standard curve prepared with different concentrations of standard glucose.

Acid-hydrolysis of non-reducing sugar and its estimation as reducing sugar: Non-reducing sugar was first hydrolyzed with the help of diluted mineral acid like hydrochloric acid. Then the hydrolyzate was neutralized and the reducing sugar was estimated by Nelson-Somogyi’s method.

Reagents Used
1. 0.1 and 1 N hydrochloric acid and 1 N sodium hydroxide.
2. Phenolphthalein indicator solution in alcohol.

Procedure
One ml of each alcohol extract was taken in test tubes and to it 1.0 ml of 1 N hydrochloric acid was added and the test tubes were then kept in a hot water bath at 50 °C for 20 min. After cooling one drop of indicator was added and mixed well. To the solution 1 N sodium hydroxide was added drop wise till the colour turned to pink due to excess alkali. The excess alkali was neutralized with 0.1 N hydrochloric acid till the solution became colourless. Then, the volume was made up to 5.0 ml. From the above 5.0 ml solution one ml was taken and reducing sugar present in the hydrolyzate was estimated by Nelson-Somogyi’s method. The reducing sugar in the hydrolyzate was a measure of total sugar, the quantity of reducing sugar was subtracted from this value and it was multiplied by a conversion factor of 0.95.

Estimation of Free Amino Acid
The estimation of total free amino acids was done following Ninhydrin method (Moore and Stein, 1984).

Reagents Used
1. Ninhydrin reagent: Stannous chloride (0.8 g) was dissolved in 500 ml of 0.2 M citrate buffer (pH 5.0) and to this 20 g of ninhydrin in 500 ml of methyl cellosolve (2-methoxyethanol) was added.
2. Diluent Solvent: Equal volumes of water and n-propanol is mixed.

Procedure
Alcohol free extract of plant samples was pipetted out into separate test tubes together with the reagent blank and 1ml of ninhydrin reagent was added and the volume was made up to 2 ml by using distilled water. Test tubes were heated in a boiling water bath for 20 minutes. Five ml of diluent solution was added and mixed well and kept for 15 min. Absorbance of purple colour was measured against reagent blank at 570 nm by using spectrophotometer. The amount of total free amino acids in samples were estimated using the standard curve prepared using different concentrations of standard leucine.

Estimation of Proline
Proline concentration was determined using the method of Bates et al. (1973) [4].

Reagents Used
1. Acid ninhydrin: 1.25g ninhydrin was warmed in 30 ml glacial acetic acid and 20 ml 6 M phosphoric acid with agitation until dissolved and stored at 4°C and used within 24 hr.
2. 3% Aqueous Sulpho Salicylic Acid
3. Glacial Acetic Acid
4. Toluene
5. Proline

Procedure
Fresh leaves (0.5g) were homogenized in 10 ml of 3% aqueous sulpho salicylic acid. The homogenate was filtered through Whatman No. 2 filter paper. Two ml of the filtrate was mixed with an equal volume of acetic acid and acid ninhydrin and incubated for 1 h at 100 °C. The reaction was terminated by placing the tube in an ice bath and extracted with 4 ml of toluene. The extract was vortexed for 20 seconds and the toluene layer was separated and warmed to room temperature. The intensity of red colour developed was measured by using a spectrophotometer at 520 nm. The amount of proline in the test sample was measured from the standard curve prepared using pure proline and proline content was expressed on fresh weight-basis using the formulae:

$$\text{mg proline per ml x ml toluene} = \frac{115.5 \times \text{g sample}}{\mu \text{moles per g tissue}}$$
Where, 115.5 is the molecular weight of proline.

Estimation of Soluble Proteins
Soluble proteins were estimated following Lowry’s method (Lowry et al., 1951).

Reagents Used
1. Alkaline copper reagent:
Solution A (2% sodium carbonate in 0.1 N NaOH), Solution B (1% sodium potassium tartrate), Solution C (0.5% copper sulfate). Solutions A, B, C were mixed in 100: 1: 1 proportion just before use.
2. Folin–Ciocalteau Reagent (FCR) 1 N

Procedure
One ml of each alcohol extract was taken in test tube and 5.0 ml of alkaline copper reagent was added to it and then it was mixed thoroughly and allowed to react for 10 minutes followed by addition of half a ml of 1 N FCR and kept in dark for 30 minutes. The blue coloured complex developed was diluted to 15 ml with distilled water and its absorbance was read at 660 nm in a spectrophotometer. The amount of soluble proteins present in sample was calculated from a standard curve prepared using different concentrations of standard protein solution prepared using Bovine serum albumin (BSA).

Results and Discussion
Impact of biophysical factors on the activity of S. litura in groundnut as influenced by organic and inorganic sources of nutrients.

A) Leaf succulency
The nourishment in organic farming system significantly influenced the leaf succulency in different treatments of groundnut crop. The treatments viz., organic sources of nutrition registered significantly less leaf succulency (72.37%) as against the maximum leaf succulency (76.89%) recorded in conventional treatment. Conservation treatment practices recorded moderate levels (71.28 to 74.59%) of leaf succulency. Rao (2002) [24] opined that the nitrogen content of the leaves in organically manured treatments of soybean was comparatively lower than the straight fertilized treatments, resulting in lesser leaf succulency thus reducing the incidence of S. litura. The correlation coefficients obtained between leaf succulency and larval populations of S. litura (0.81) were significant and positive indicating its role in imparting resistance to the host. Increase in the plant nitrogen through application of straight fertilizers increased the level of leaf succulency and this might have facilitated the increased feeding of defoliators due to thinner cell walls or higher turgidity of the tissues. Elanchezhyan et al. (2009) [10] obtained significant positive relationship between total chlorophyll and moisture content with shoot damage caused by Leucinodes orbonalis in brinjal. Similar association was reported by the earlier workers (Jinsa, et al., 2012; Abhilash et al., 2018) [12, 1] who reported increased palatability of the food material with more moisture content in case of susceptible varieties.

B) Leaf Thickness
The treatments showed significant difference in leaf thickness, with maximum thickness in organic treatments that recorded 6.52 mg/cm² in groundnut as against minimum of 5.86 mg/cm² in conventional treatments, where as in conservation treatments it ranged from 6.13 to 6.41 mg/cm². Jinsa et al. (2012) [12] reported maximum leaf thickness in soybean grown under organic sources of nutrition compared to conventional, which can be attributed to higher accumulation of epicuticular wax content in plants nourished with organics. The relationship between leaf thickness and larval population (r= -0.69) was negative and significant, expressing its role in imparting resistance against defoliator pests. Jinsa et al. (2012) [12] also observed the lowest incidence of major pests in soybean plants, having maximum leaf sheath thickness, treated with organic sources of nutrients.

C) Leaf area
The results with regard to leaf area in groundnut under different treatments revealed that maximum leaf area (767.95 cm²) was recorded in conventional treatment as against treatment with organic sources of nutrients which recorded comparatively minimum leaf area (696.33 cm²) and in Conservations treatments leaf area was observed in between 726.46 to 751.75 cm². The correlation studies exhibited positive and significant relationship between leaf area and larval population (r=0.77). The present results are in agreement with Amjad et al. (2003) [2] who gave an account on positive correlation with leaf area and soybean looper infestation in soybean.

D) Plant Height
Significantly higher plant height was observed in organic system treatment (18.47 cm.) as against conventional (16.21 cm.), where as plant height in conservation treatments was in between organic and conventional treatments. The relationship between plant height and larval population of S. litura were positive but, non-significant (r= 0.38). Similar results were also reported by Ramachandra reddy et al. (1998) [22], Sharma et al. (2002), Ramgopal et al. (2003) [23] and Shwetha (2007) [28].

Impact of biochemical factors on the activity of S. litura in groundnut as influenced by organic and inorganic sources of nutrients.

1) Total Phenols
In the present investigation, the crop nourished with organics showed higher leaf phenol content of 0.53 mg/g, than the conventional (0.41 mg/g) (Table-1). This might be a contributing factor for low incidence of pests in crops that received organics. Conservation treatments also showed higher phenol content compared to inorganically treated plots. Total foliar nitrogen increased as much as four fold with fertilizers (Rauzi, 1978) and increase in plant nitrogen content through fertilizers was reported to reduce the level of phenols (Jones, 1976). The correlation studies showed a significant negative relationship (r = -0.737 in groundnut) (Table -2) between phenol content and defoliator population. Phenols are known to play an important role in plant defence against insects. Jinsa et al. (2014) [13] observed a negative correlation between total phenolic content and defoliators in soybean.
both in inorganic and organic forms, recorded moderate levels, 4.21 to 4.83 mg per g, which are next to organic source of treatments. The free amino acids had a significant positive relationship with that of larval population (r = 0.77) indicating its contribution in inducing susceptibility against \textit{S. litura}. The organic sources of nutrients showed considerable impact in decreasing the free amino acid content of plants and in turn reduced the incidence of defoliator pests of groundnut. This is in line with earlier findings that lower amount of free amino acids in cotton imparted resistance to leaf hopper attack (Singh and Agarwal, 1988) \cite{30}.

4) Proline

The leaves samples of soybean and groundnut plants in conventional agriculture treatments which received maximum pest damage recorded maximum proline content of 3.125 \textmu M per g. This might be due to the rapid accumulation of proline in relation to stress condition caused by heavy insect infestation. Whereas, in plants nourished with organics, proline content was less 0.97 \textmu M per g. The proline content in conservation treatment ranged from 0.98 to 1.09 \textmu M per g. The correlation between proline content and pest infestation revealed a significant positive relationship (r = 0.743 in groundnut). Similar relationship between proline content and pest incidence was revealed by Jinsa \textit{et al.} (2014) \cite{13}, who observed lower proline content in soybean resistant to caterpillar damage. Similar results were also obtained by Roy \textit{et al.} (1988) who reported higher proline accumulation due to rice brown plant hopper attack in rice.

5) Soluble Proteins

The crop receiving organics recorded significantly less protein content of 20.66 mg per g 4.56 mg per g in groundnut as against 4.12 mg per g in groundnut in conventionally grown plants and 3.16 to 3.89 mg per g in conservation agriculture treatments. The plants receiving organic amendments showed low amounts of proteins thus making the plant unpalatable to insects and rendered them unsuitable for insect nutrition. Further, the correlation studies revealed a significant positive relationship (r = 0.689) between soluble proteins and pest infestation and damage. This is in agreement with the observation of Jinsa \textit{et al.} (2014) \cite{13} who stated that proteins were important constituents in insect nutrition and

2) Sugars

Among different biochemical constituents sugars (Reducing sugars and non-reducing sugars) contribute greatly to susceptibility of the host. The results revealed that the treatments with organic sources of nutrition possessed reduced quantities of total sugars at the critical stages of crop growth (4.72 mg/g) as against conventional treatment (5.39 mg/g). However, in case of conservation treatments, it varied from 5.12 to 5.24 mg per g. The correlation studies revealed positive and significant relationship between reducing sugars and larval population (r= 0.80). The results are in conformation with Chhabra \textit{et al.} (1990) \cite{6}, Halder \textit{et al.} (2006) \cite{11} and Jinsa \textit{et al.} (2014) \cite{13} who reported a positive significant correlation between total sugars, reducing sugars and non-reducing sugars and pod damage in gram and defoliators on soybean respectively.

3) Free Amino Acids

The results with regard to free amino acids in groundnut grown under different systems revealed that organic sources of nutrients showed considerable impact in decreasing the free amino acid content of plants to 4.47 mg per g. This in turn reduced pest infestation in groundnut whereas, in plants which received NPK as inorganic fertilizers (conventional) recorded high amounts of free amino acids (5.09 mg/g). However, conservation treatments which received nutrients

\begin{table}
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\begin{tabular}{|c|c|c|c|c|c|c|c|c|}
\hline
\textbf{Farming systems} & \textbf{Biochemical parameters} & \textbf{Biophysical parameters} \\
\hline
 & Total sugar (mg g fresh wt\(^{-1}\)) & Reducing sugar (mg g fresh wt\(^{-1}\)) & Phenols (mg g fresh wt\(^{-1}\)) & Proline (\textmu M/g fresh wt\(^{-1}\)) & Soluble protein (mg g fresh wt\(^{-1}\)) & Amino acid (mg g fresh wt\(^{-1}\)) & Leaf succulence (\%) & Leaf thickness (mg/cm\(^2\)) & Leaf area (cm\(^2\)) & Plant height (cm) \\
\hline
Organic & 4.72\(^{a}\) & 1.06\(^{a}\) & 0.53\(^{bc}\) & 0.97\(^{b}\) & 4.56\(^{a}\) & 4.47\(^{b}\) & 72.37\(^{a}\) & 6.52\(^{b}\) & 696.33\(^{b}\) & 18.47\(^{b}\) \\
Broad bed furrow with residue retention & 5.21\(^{b}\) & 1.11\(^{a}\) & 0.48\(^{bc}\) & 0.98\(^{a}\) & 3.16\(^{c}\) & 4.83\(^{c}\) & 71.28a & 6.13\(^{a}\) & 751.75\(^{bc}\) & 17.13\(^{a}\) \\
Broad bed furrow with residue incorporation & 5.12\(^{b}\) & 1.16\(^{bc}\) & 0.42\(^{b}\) & 0.98\(^{a}\) & 3.89\(^{d}\) & 4.21\(^{a}\) & 74.59\(^{a}\) & 6.27\(^{bc}\) & 763.82\(^{bc}\) & 17.56\(^{a}\) \\
Flat bed with residue retention & 5.24\(^{b}\) & 1.17\(^{bc}\) & 0.47\(^{b}\) & 1.02\(^{d}\) & 3.46\(^{b}\) & 4.67\(^{c}\) & 72.16\(^{a}\) & 6.18\(^{bc}\) & 726.46\(^{a}\) & 17.26\(^{a}\) \\
Flat bed with residue incorporation & 5.16\(^{b}\) & 1.06\(^{b}\) & 0.51\(^{c}\) & 1.09\(^{d}\) & 3.87\(^{c}\) & 4.22\(^{d}\) & 71.84\(^{a}\) & 6.41\(^{b}\) & 740.71\(^{b}\) & 17.18\(^{b}\) \\
Conventional & 5.39\(^{b}\) & 1.32\(^{c}\) & 0.41\(^{b}\) & 1.25\(^{c}\) & 4.12\(^{b}\) & 5.09\(^{c}\) & 76.89\(^{b}\) & 5.86\(^{c}\) & 767.95\(^{b}\) & 16.21\(^{b}\) \\
S.Em (\pm) & 0.06 & 0.05 & 0.01 & 0.02 & 0.07 & 0.05 & 1.00 & 0.11 & 11.09 & 1.08 \\
CD @1% & 0.24 & 0.19 & 0.04 & 0.08 & 0.27 & 0.20 & 4.07 & 0.44 & 45.13 & 4.40 \\
\hline
\end{tabular}
\caption{Biochemical and biophysical traits in groundnut as influenced by organic, conservation and conventional practices}
\end{table}

\begin{table}
\centering
\begin{tabular}{|c|c|c|}
\hline
\textbf{Biochemical and biophysical factors} & \textbf{Leaf larval population (r value) (60 DAS)} \\
\hline
Total sugar & 0.808\(^{a}\) \\
Reducing sugar & 0.749 \\
Non reducing & - \\
Phenol & -0.737 \\
Proline & 0.743 \\
Soluble protein & 0.689 \\
Amino acid & 0.775 \\
Leaf area & 0.775 \\
Thickness & -0.692 \\
Leaf succulence & 0.819 \\
Plant height & 0.386\(^{bc}\) \\
\hline
\end{tabular}
\caption{Correlation of biochemical and biophysical traits with \textit{Spodoptera litura} population in groundnut as influenced by organic, conservation and conventional practices}
\end{table}
the strains with higher soluble protein would naturally have high defoliator infestation.

Conclusion
Biophysical characters in groundnut like leaf succulence and leaf area revealed significant positive relationship with incidence of defoliators but leaf thickness was observed to have significant and negative relationship. Higher phenols, lower free amino acids, lower total sugars and reducing sugars, lower protein content were present in plants nourished with organics, followed by conservation farming and conventional farming system. These metabolites are on the opposite in conventional farming. These accumulated biochemical constituents and their quantities in plant tissues impart resistance v/s pests in crops.

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