Tritrophic Interactions of Cowpea [Vigna unguiculata subsp. unguiculata (L.)], Aphids [Aphis craccivora (Koch)] and Coccinellids [Menochilus sexmaculatus (Fab.)] under eCO2 and eTemp


A B S T R A C T

Experiments were conducted to understand the direct and indirect effects of temperature and elevated CO2 (eCO2), on tritrophic interactions of cowpea (Vigna unguiculata subsp. unguiculata L.), legume aphid Aphis craccivora Koch and coccinellid predator Menochilus sexmaculatus Fab. Reduction of the leaf nitrogen (6%), amino acid (6%) and protein (7%) of cowpea foliage with increased carbon (13%) and C:N ratio (21%) at eCO2 over aCO2 indicated the dilution of biochemical constituents at first trophic level. Shortened development time, DT and increment of reproductive rate, RR at eCO2 over ambient CO2(aCO2) was significant with increase in temperature from 20 to 35 °C. Reduction of the mean degree day, DD requirement of both nymphal (75.79 ± 15.163) and adult stages (157.15 ± 67.04) at eCO2 over aCO2 and same was reflected in the sumation DD for both the stages at eCO2 (232.96 ± 80.32) and aCO2 (247.07 ± 64.77) across six temperatures. The ‘r_m’ and ‘R_r’ increased gradually with increase in temperature followed the non-linear trend and reached maximum values at 27 °C with shortened ‘Tm’ across 20 to 35 °C temperatures at eCO2 indicating the significant variation of growth and development at the second trophic level. Decreased grub duration (23%) with increased predation capacity (19%) of M. sexmaculatus on A. craccivora at eCO2 over ambient was noted, indicating the incidence of A. craccivora is likely to be higher with increased predation in the future climate change scenario.

Introduction

Atmospheric CO2 concentration has been steadily increasing and reached 380–400 ppm in 2013. It is anticipated to double by the end of the 21st century and an increase is predicted by 1.7–4.9 °C in mean global temperature by 2100 (IPCC, 2013). The average increase in temperature was found to be 0.78 (0.72 to 0.85) °C between the periods of 1850–1990 and 2003–2012. Ambient carbon dioxide (aCO2) concentration exceeded 400 ppm and future estimations predict an increase up to 550 ppm within a few decades (IPCC, 2014). The agriculture sector is more vulnerable to climate variability and climate change and the elevated CO2 and temperature would threaten the food security of the people across the globe. It is well documented that crop plants get stimulated with elevated CO2 concentrations and thus increased biomass and higher photosynthesis (Yu et al., 2012) that in turn influences the growth and development of insect herbivores (Hunter, 2001). Elevated atmospheric CO2 (eCO2) influences crop growth by altering C: N ratio, increasing C (Gutierrez et al., 2008) and decreasing N (Dixon et al., 1993). The quality and quantity of the food available to the insect herbivores get changed due to eCO2 (Chen et al., 2007) and temperature; and thus the insect pests prefer to go for increased feeding rate to meet compensatory nutritional requirements (Hunter, 2001).

Cowpea (Vigna unguiculata subsp. unguiculata L.) is an important legume crop with higher protein content and is grown both in alfisols and vertisols. The crop is preferred as vegetable and even after processing cowpea, peanut, pigeonpea etc. The incidence of aphids is common in the early vegetative stage of the crop and its infestation will continue till crop maturity. Both nymphs and adults suck the sap from plant parts (leaves, inflorescence and tender pods) and cause enormous yield losses. In the majority of the cases A. craccivora also acts as a vector in transmitting the important viral diseases (Wightman and Wightman, 1994).

The response of insect species to eCO2 varies individually across different species and it is understood that the multiplication of aphids is...
higher at eCO2 conditions (Newman, 2005) and can cause significant economic losses (Cannon, 1998). It is known (Hughes and Bazzaz, 2001) that the response of aphids to eCO2 is ‘species-specific’ in nature. The growth and development of aphids is influenced by the temperature and complete understanding will be of immense help to know the population dynamics. It is well known that insect performance is affected both directly and indirectly by temperature and eCO2 (Stiling and Cornelissen, 2007). Harrington et al. (2001) reported that the direct effect of temperature will be greater than rising CO2. The interactive and concurrent effects of both parameters are more evident and complex; therefore clear understanding will be of immense help to predict insect pest dynamics in future climate change.

The occurrence of the important generalist predator, Menochilus sexmaculatus (Fabricius) is very common on A. craccivora on cowpea. Level of coccinellid predation on aphids is in the range of 110.60 to 132.53 nos. /day and is a host dependent factor (Patel, 2015). The significant effects of temperature on tritrophic interactions is well known but scarce information is available on impacts of eCO2 (Gao et al., 2010). The significant effect of eCO2 and eTemp on growth and development of A. craccivora and its predator (M. sexmaculatus) on cowpea across three trophic levels was hypothesized here by conducting an array of experiments.

Materials and methods

Elevated atmospheric CO2 (eCO2) of 550 ± 25 ppm and ambient CO2 (aCO2) concentration of 380 ± 25 ppm were maintained in open top chambers (OTC) of 4X4X4 m dimensions, constructed at ICAR-Central Research Institute for Dryland Agriculture, Hyderabad (17.38 N, 78.47 E). Four OTCs were used for experimentation purpose (Srinivasa Rao et al., 2012b) with two replications of eCO2 and aCO2 in each chamber. During the month of June, 2016 cowpea (cv C 152) seeds were sown and the crop was maintained by following the recommended package of practices except plant protection. Methodology adopted for study on three trophic levels, i.e., estimation of changes in cowpea crop (trophic-I), in the growth and development of A. craccivora (trophic-II) and on predator (trophic-III) are given here under.

Biochemical analyses of cowpea foliage (Trophic-I)

The carbon, N contents, C:N and polyphenols were estimated in the cowpea foliage by adopting the standard protocols. Carbon and N were measured using a CHN analyzer (Model NA 1500 N, Carlo Erba Strumentazione, Italy) after drying of leaf samples (Jackson, 1973). Condensed tannins were determined by the Folin-Denis method by estimating the tannic acid equivalents (TAE) (Anderson and Ingram, 1993). For this method, leaf samples were dried at 40 °C for 48 h. Dried leaf samples were ground to powder, and phenolics were extracted with CH3OH. The concentration of polyphenols in the extract was determined by spectrophotometry using tannic acid as the standard, and the results were expressed as percentage tannic acid equivalents. The soluble protein was estimated by Lowry et al. (1951). Soluble proteins were extracted from 0.5 g samples (leaf and seed) using distilled water. Aliquot 0.1 and 0.2 ml were taken and made up to 1 ml with distilled water and a tube with 1 ml of water alone serving as a blank. Five ml of alkaline copper reagent was added and kept for 10 min. Then 0.5 ml of 1 N Folin Ciocalteau Reagent (FCR) was added and kept in a dark place for 30 min. Later blue color develops and read the intensity at 660 nm. Soluble proteins were calculated using Fraction V of Bovine Serum Albumin (BSA) standards (20–100 μg). Estimation of free amino acids gives an indication about physiological status and the health condition of plants. Ninhydrin method (Moore and Stein, 1949) was used for the estimation of free amino acids and is colorimetrically measured at 570 nm to explore physiological status and the health condition of plants.

Cowpea-aphid interactions (Trophic-II)

Field collected adults of A. craccivora from infested plants of cowpea were maintained in insect laboratory of CRIDA at optimum conditions of temperature (25 ± 2 °C), humidity (75 ± 5%) and 14 L/10D cycle conditions. Stock culture of A. craccivora was maintained continuously on cowpea plants in separate growth chambers for use in experimental studies.

Feeding trials of A. craccivora on cowpea

Feeding trials were conducted in CO2 growth chambers (PERCIVAL I-36L). Aphids were reared on leaves of potted plants of cowpea obtained from respective CO2 conditions. Aphids were reared individually in petridishes (110 mm diameter and 10 mm height) adopting the ‘cut leaf’ method (Srinivasa Rao et al., 2013a). A group of A. craccivora adults were placed in a closed petridishes and were maintained in CO2 growth chambers at six constant temperatures (20, 25, 27, 30, 33 & 35 ± 0.5 °C) at 75 ± 5% relative humidity (RH) and 14L/10D hour photoperiod.

Biology of A. craccivora on cowpea

Adults of A. craccivora used for study on biology were reared in treatment environments for two generations to rule out the maternal effects. One day old nymphs were placed singly on the leaf in single petriplate in each CO2 condition (380 & 550 ppm) and allowed to grow until the start of reproduction to record the development time (DT). Ten replicates per set and three sets were used under each treatment with each aphid serving as one replicate. It means that 30 replications were adopted. Time taken by each nymph from its birth to the onset of reproduction was accounted for development time. The freshly moulted adults of A. craccivora were kept individually in petridish each consisting of a cowpea leaf for observation on reproduction. Number of nymphs produced per day were also noted at 12 h interval and removed after counting to prevent overcrowding. Parameters of aphid biology viz., development time (DT), total life cycle (TLC) and fecundity were recorded as per the standard procedure (Srinivasa Rao et al., 2013a) for each of the eCO2 and eTemp conditions.

Estimation of degree days requirement on cowpea

Data on biological parameters were analyzed to estimate thermal constant for each stage of aphid. Degree days requirement was calculated considering the average development time at six different temperatures used in the experiment. Linear regression co-efficient (b) and constant (a) were estimated after plotting the reciprocal of development time (developmental rate) over temperature. The standard formula of Υ = a + bx was used to estimate the threshold development (Tb) and degree day calculation was based on developmental rate (Zahiri et al., 2010) was adopted.

Estimation of life table parameters of A. craccivora on cowpea

Various life table parameters were calculated using TWosex - MS Chart software (Chi, 2005). To facilitate raw data analysis, life table analysis, a user-friendly computer program, was used (http://140.120197.173/Ecology/prod02.htm). Intrinsic rate of increase (rni), generation time (T), net reproductive rate (Ro) and finite rate of increase (λ) were estimated using cowpea foliage from CO2 and aCO2 separately at above mentioned six temperatures.

Cowpea-aphid- coccinellid interactions (Trophic-III)

Culture of Predator (M. sexmaculatus) was maintained by following standard procedure (Solangi et al., 2007) on A. craccivora feeding on cowpea. Adults of M. sexmaculatus were collected from field and selected mating pairs were reared on A. craccivora nymphs in laboratory in wide mouth plastic jars. Feeding trials of M. sexmaculatus on cowpea reared A. craccivora were conducted at eCO2 and aCO2 conditions at
optimum temperature of \(27 \pm 1^\circ\text{C}\) to understand the impact of CO2 on predator. Each mating pair of coccinellids was selected from the stock culture and was kept in petridishes of 9 cm diameter per replication (10 replications/each CO2 condition).

Grubs and adults of \textit{M. sexmaculatus} were released independently on ten \textit{A. craccivora} adults (3 day old) in a petridish on cowpea leaves by maintaining the corresponding CO2 conditions. The predators were allowed to lay eggs. Freshly hatched \textit{M. sexmaculatus} grubs were released on \textit{A. craccivora} and observations viz., grub duration (days) and predation capacity (%) of \textit{M. sexmaculatus} were recorded at 24 hour period.

**Statistical analysis**

One way ANOVA was adopted for analyzing the data on biochemical constituents. The data on DT, RT, TLC and fecundity of aphid at six constant temperatures at two CO2 conditions were analyzed by using two factorial analysis considering the CO2 and temperatures levels as main and sub-factors. Results presented are mean value of each determination (treatment) ± standard deviation. The mean values of life main and sub-factors. Results presented are mean value of each determination (treatment) ± standard deviation. The mean values of life main and sub-factors. Results presented are mean value of each determination (treatment) ± standard deviation. The mean values of life main and sub-factors. Results presented are mean value of each determination (treatment) ± standard deviation. The mean values of life table parameters of \textit{A. craccivora} at six constant temperatures were plotted and a non linear relationship (standard quadratic form: \(Y = ax^2 + bx + c\)) was worked out taking the plotted and a non linear relationship (standard quadratic form: \(Y = ax^2 + bx + c\)) was worked out taking the plotted and a non linear relationship (standard quadratic form: \(Y = ax^2 + bx + c\)) was worked out taking the plotted and a non linear relationship (standard quadratic form: \(Y = ax^2 + bx + c\)) was worked out taking the plotted and a non linear relationship (standard quadratic form: \(Y = ax^2 + bx + c\)) was worked out taking the plotted and a non linear relationship (standard quadratic form: \(Y = ax^2 + bx + c\)) was worked out taking the plotted and a non linear relationship (standard quadratic form: \(Y = ax^2 + bx + c\)) was worked out taking the plotted and a non linear relationship (standard quadratic form: \(Y = ax^2 + bx + c\)) was worked out taking the plotted and a non linear relationship (standard quadratic form: \(Y = ax^2 + bx + c\)) was worked out taking the plotted and a non linear relationship (standard quadratic form: \(Y = ax^2 + bx + c\)) was worked out taking the plotted and a non linear relationship (standard quadratic form: \(Y = ax^2 + bx + c\)) was worked out taking the plotted and a non linear relationship (standard quadratic form: \(Y = ax^2 + bx + c\)) was worked out taking the plotted and a non linear relationship (standard quadratic form: \(Y = ax^2 + bx + c\)) was worked out taking the plotted and a non linear relationship (standard quadratic form: \(Y = ax^2 + bx + c\)) was worked out taking the plotted and a non linear relationship (standard quadratic form: \(Y = ax^2 + bx + c\)) was worked out taking the plotted and a non linear relationship (standard quadratic form: \(Y = ax^2 + bx + c\)) was worked out taking the plotted and a non linear relationship (standard quadratic form: \(Y = ax^2 + bx + c\)) was worked out taking the plotted and a non linear relationship (standard quadratic form: \(Y = ax^2 + bx + c\)) was worked out taking the plotted and a non linear relationship (standard quadratic form: \(Y = ax^2 + bx + c\)) was worked out taking the plotted and a non linear relationship (standard quadratic form: \(Y = ax^2 + bx + c\)) was worked out taking the plotted and a non linear relationship (standard quadratic form: \(Y = ax^2 + bx + c\)) was worked out taking the plotted and a non linear relationship (standard quadratic form: \(Y = ax^2 + bx + c\)) was worked out taking the plotted and a non linear relationship (standard quadratic form: \(Y = ax^2 + bx + c\)) was worked out taking the plotted and a non linear relationship (standard quadratic form: \(Y = ax^2 + bx + c\)) was worked out taking the plotted and a non linear relationship (standard quadratic form: \(Y = ax^2 + bx + c\)) was worked out taking the plotted and a non linear relationship (standard quadratic form: \(Y = ax^2 + bx + c\)) was worked out taking the plotted and a non linear relationship (standard quadratic form: \(Y = ax^2 + bx + c\)) was worked out taking the plotted and a non linear relationship (standard quadratic form: \(Y = ax^2 + bx + c\)) was worked out taking the plotted and a non linear relationship (standard quadratic form: \(Y = ax^2 + bx + c\)) was worked out taking the plotted and a non linear relationship (standard quadratic form: \(Y = ax^2 + bx + c\)) was worked out taking the plotted and a non linear relationship (standard quadratic form: \(Y = ax^2 + bx + c\)) was worked out taking the plotted and a non linear relationship (standard quadratic form: \(Y = ax^2 + bx + c\)) was worked out taking the plotted and a non linear relationship (standard quadratic form: \(Y = ax^2 + bx + c\)) was worked out taking the plotted and a non linear relationship (standard quadratic form: \(Y = ax^2 + bx + c\)) was worked out taking the plotted and a non linear relationship (standard quadratic form: \(Y = ax^2 + bx + c\)) was worked out taking the plotted and a non linear relationship (standard quadratic form: \(Y = ax^2 + bx + c\)) was worked out taking the plotted and a non linear relationship (standard quadratic form: \(Y = ax^2 + bx + c\)).

**Results**

**Impact of CO2 on biochemical constituents of cowpea (Trophic-I)**

In this study, the leaf N concentration was lower in cowpea foliage obtained from eCO2 conditions over aCO2. Nitrogen content was significantly (\(F_{1,4} = 26.27; \ p < 0.01\)) lower under eCO2 (5.08 ± 0.08) than ambient. In contrast, carbon content was significantly higher (38.20 ± 1.43) (\(F_{1,4} = 23.23; \ p > 0.05\)) and influenced by CO2 (Fig. 1). However, the relative C: N ratio (\(F_{1,4} = 27.78; \ p < 0.05\)) was considerably higher (7.52 ± 0.256) in eCO2 cowpea foliage than in ambient (6.23 ± 0.70). No significant difference was observed in polyphenols content measured in terms of tannic acid (\(F_{1,2} = 0.07; \ p < 0.01\)) equivalents at both CO2 conditions (Fig. 1). Amino acids and proteins (\(F_{1,2} = 5.83; \ p < 0.01\)) are lower in eCO2 (0.0016 & 27.223) when compared to ambient (0.0017 & 29.011).

**Impact of eCO2 and temperature on \textit{A. craccivora} (Trophic-II)**

Variations in biological parameters of \textit{A. craccivora} on cowpea at two CO2 conditions and six constant temperatures are presented in Table 1. Development time (DT) was significantly reduced by the first factor, CO2 (\(F_{1,29} = 4954.26; \ P < 0.001\)) and second factor, temperature (\(F_{5,290} = 8944.18; \ P < 0.001\)) indicating the reduction of DT with elevated CO2 and an increase in temperature. Increased reproductive time (RT) (19.50 ± 0.628) with CO2 (\(F_{1,29} = 1392.63; \ P < 0.001\)) and TLC (\(F_{5,290} = 9106.12; \ P < 0.001\)) decreasing with the increase in temperature. Increased fecundity (\(F_{5,290} = 1611.72; \ P < 0.001\)) was also observed with the increase in temperature.

**Table 1**

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**Fig. 1.** Variation in biochemical constituents of cowpea foliage at eCO2 conditions.
female was also impacted by CO$_2$ ($F_{1,92} = 342.14; P < 0.001$) and temperature ($F_{5,290} = 1611.72; P < 0.001$). A substantial increase in production of offspring was noticed at eCO$_2$ (74.90 ± 6.90) over aCO$_2$ (60.67 ± 8.06). The highest offspring production was noticed between 20 and 27°C and started declining with increase in temperature (30–35°C). In contrast, adults required higher degree days of development rate of nymphs and adults which on temperature (Fig. 2).

**Thermal constants and degree days for A. craccivora stages on cowpea**

Linear regression was used to estimate the temperature thresholds and thermal requirements of nymph and adult stages of *A. craccivora* independently. The best fit linear regression equation for nymph ($Y = 0.0278x - 0.4587, R^2 = 0.89$) and adult ($Y = 0.0101x - 0.1687, R^2 = 0.93$) was noted. The mean lower development thresholds for both stages were 7.7 and 20.5°C on eCO$_2$ foliage. In contrast, on aCO$_2$ foliage, similar and significant trends with nymphal and adult stages of *A. craccivora* were noted. The mean lower development thresholds for nymph were 16.50 at eCO$_2$ than aCO$_2$ (8.34°C) by regressing development rate of nymphs and adults which on temperature (Fig. 2).

Accumulated heat units for nymph stage varied from 58.86–95.71 DD on eCO$_2$ as against 85.83–117.86 DD on aCO$_2$ with temperature range of 20–35°C. In contrast, adults required higher degree days of 88.33–264.50 at eCO$_2$ as against 86.66–189.33 on aCO$_2$ cowpea foliage at temperatures of 20–27°C. The mean degree-day requirement of nymph (75.79 ± 15.163) and adult (157.15 ± 67.04) at eCO$_2$ varied over aCO$_2$ with similar trends reflected after summation of degree days of both nymphal and adult stages at eCO$_2$ (232.96 ± 80.32) and aCO$_2$ (247.07 ± 64.77).

**Construction of life tables**

Impact of eCO$_2$ across different temperatures on life table parameters of *A. craccivora* are given in Table 2. It is evident that $r_{m,n}$ increased with increase in temperature from 20°C showing proportional trend that started declining from 30°C. The $R_{m,n}$ of *A. craccivora* was higher at 27°C with 62 offspring at eCO$_2$. The $r_{m,n}$ and $R_{m,n}$ increased gradually with increase in temperature and reached maximum values at 27°C. The reduction of $T$ was evident from 11 days at 20°C with minimum of 3.29 days at 35°C at eCO$_2$ and followed the non-linear trend. The $\lambda'$ was found to be highest 1.45 at 27°C and followed the decreasing trend with an increase in temperature. The association between four life table parameters viz., $r_{m,n}$, $R_{m,n}$, $T$ and $\lambda'$ and temperature at both eCO$_2$ and aCO$_2$ conditions was found to be non-linear (Fig. 3).

Quadratic form of equation with higher $R^2$ (0.85) at eCO$_2$ was the best noticed between $r_{m,n}$ and temperature. Other parameters viz., $R_{m,n}$, $T$ and $\lambda'$ followed similar trend. When the first order derivative of equation is taken as ‘zero’, the temperature at which higher $R_{m,n}$ was estimated to be 24.20°C on eCO$_2$ foliage as against 23.29 (aCO$_2$). The $r_m (27.80)$ and $\lambda$ (26.48) showed declining trend at 20–35°C at eCO$_2$ and similar nonlinear trends were observed at aCO$_2$ which are as follows ($r_m = -0.002x^2 + 0.1138x - 1.2354, R^2 = 0.85$, $(R_m = 0.3309x^2 + 15.416x - 135.39, R^2 = 0.83)$, $(T = 0.0157x^2 - 0.3799x + 12.064, R^2 = 0.93)$ and $(\lambda = -0.0029x^2 + 0.1597x - 0.805, R^2 = 0.86)$. When the first order derivative is equated to ‘zero’ and to solve the $x' (variable)$ the $r_m$ and $\lambda$ started declining at 28.45°C and 27.22°C on eCO$_2$ foliage and found to be higher than eCO$_2$ (Fig. 3).

**Impact of eCO$_2$ on *M. sexmaculatus* (Trophic-III)**

The data on impact of CO$_2$ on two vital parameters of coccinellids which were allowed to feed on *A. craccivora* of cowpea foliage at both elevated and ambient conditions depicted in Fig. 4. Duration of grubs was found to be significantly ($F_{1,9} = 20.43; P < 0.001$) reduced at eCO$_2$ (16) over aCO$_2$ (18.70) and in contrast, the predation capacity of coccinellids at both nymph and adult stages increased at eCO$_2$ (82) over aCO$_2$ (78) condition ($F_{1,9} = 5.88; P < 0.001$).

**Discussion**

Experiments conducted to assess and quantify the impact of eCO$_2$ and temperature across three trophic levels of cowpea, *A. craccivora*, and *M. sexmaculatus* revealed the significance of two major dimensions of climate change. Earlier studies (Cornelissen, 2011; Srinivasa Rao et al., 2014) also reported the significant effect of both eCO$_2$ and temperature on other insect pests.

At first trophic level, reduction of leaf N content (6%) amino acid (6%) and protein (7%) in cowpea foliage with increased C (13%) and CN ratio (21%) at eCO$_2$ over aCO$_2$. A similar reduction of leaf N and protein concentrations by more than 12% (Ainsworth and Long, 2005) and ($\lambda = -0.0029x^2 + 0.1597x - 0.805, R^2 = 0.86$). When the first order derivative is equated to ‘zero’ and to solve the ‘$x'$ (variable) the $r_m$ and $\lambda$ started declining at 28.45°C and 27.22°C on eCO$_2$ foliage and found to be higher than eCO$_2$ (Fig. 3).

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**Discussion**

Experiments conducted to assess and quantify the impact of eCO$_2$ and temperature across three trophic levels of cowpea, *A. craccivora*, and *M. sexmaculatus* revealed the significance of two major dimensions of climate change. Earlier studies (Cornelissen, 2011; Srinivasa Rao et al., 2014) also reported the significant effect of both eCO$_2$ and temperature on other insect pests.

At first trophic level, reduction of leaf N content (6%) amino acid (6%) and protein (7%) in cowpea foliage with increased C (13%) and CN ratio (21%) at eCO$_2$ over aCO$_2$. A similar reduction of leaf N and protein concentrations by more than 12% (Ainsworth and Long, 2005) and ($\lambda = -0.0029x^2 + 0.1597x - 0.805, R^2 = 0.86$). When the first order derivative is equated to ‘zero’ and to solve the ‘$x'$ (variable) the $r_m$ and $\lambda$ started declining at 28.45°C and 27.22°C on eCO$_2$ foliage and found to be higher than eCO$_2$ (Fig. 3).
Srinivasa Rao et al., 2012a).

Impact of eCO₂ and temperature on second trophic level indicated that DT of A. craccivora decreased by 1–5 days as with increase in temperature from 20 to 35 °C (67%) and was more evident at eCO₂. A similar trend was reported with other species of aphids i.e., A. gossypii in cotton and cucumber (Mohammad, 2010) and Rhopalosiphum maidis (Fitch) in barley (Xie et al., 2014) with CO₂ and temperature. The response of aphids to eCO₂ is mainly with species specificity and is able to overcome the disadvantage of eCO₂ by shortening DT. In addition to CO₂, the temperature also influences the developmental rate of insect pests because of their physiological sensitivity (Aleosfsoor and Fekrat, 2014).

Increased reproductive time of aphids and production of offspring for longer periods was more evident in the present study with elevated CO₂. Slight increment of RT in aphids with elevated CO₂ was observed. Insects change their physiology from time to time to get adaptation to changing climate. Earlier studies conducted by the authors (Mathukumalli et al., 2017) on A. craccivora in groundnut crop at both CO₂ levels showed the similar reduction of DT and increment of RT of aphids at eCO₂ over ambient across different temperatures.

It is reported that aphids proliferate in a higher number to overcome both indirect and direct effect of CO₂ and temperature. Increased fecundity was reported by Oehme et al. (2011) in case of Myzus persicae on Triticum aestivum. Similar finding was reported by the authors with A. craccivora on cowpea (Srinivasa Rao et al., 2013a) and groundnut (Mathukumalli et al., 2017) crops. More production of offspring with temperature in the range of 20–27 °C was noticed. Further increase in temperature caused reduction of progeny production which is in agreement with Lange et al. (2006). The present study quantifies the impact of host mediated effect of eCO₂ and direct effect of temperature on A. craccivora on cowpea and the corroborative evidences for change in growth and development of aphids were identified.

Decreased N (6%), amino acids (6%) and protein content (7%) with increased C (13%) and C:N ratio (21%) at eCO₂ over aCO₂ was noticed in cowpea foliage and these changes have significantly influenced the growth and development of A. craccivora on cowpea. Decreased N (3%) and increased C: N ratio (13%) and polyphenols (2%) in foliage of groundnut having significant influence on growth and development of insect herbivores have well documented (Srinivasa Rao et al., 2015) at eCO₂ and castor (Srinivasa Rao et al., 2013b). The variation of biological of aphids is attributed to changing nutritional quality of food due to change in biochemical composition of cowpea foliage at eCO₂ conditions.

Lower degree days requirement for completion of nymph and adult stages at eCO₂ (75.79 ± 15.16 DD) differed from those of aCO₂ (156.15 ± 67.04 DD) possibly due to to shortening of DT. Similar
lower thermal constants were reported for *Aphis glycines* Matsumura (McCormack et al., 2004), *R. maidis* (Kuo et al., 2006) and *A. craccivora* (Berberet et al., 2009). Summation of degree days requirement of both nymph and adult stages together also revealed a similar trend at eCO₂ (232.96 ± 80.32) and aCO₂ (247.07 ± 64.77). Findings also showed lower thermal requirement for nymphs and higher for adults feeding on foliage at eCO₂ over aCO₂ at temperatures of 20–35 °C. The overall degree day requirement of nymphs and adults of *A. craccivora* on cowpea at eCO₂ were similar to that of *A. craccivora* on groundnut at eCO₂ (Mathukumalli et al., 2017) except lower threshold levels of nymphs which were found to be higher on cowpea than groundnut. This might be due to higher CN variation in cowpea (17%) than that of groundnut (12%) reported earlier.

Increased rm (54%) and R0 (32%) with reduced T (28%) was noticed and followed an increasing trend with temperature in the range of 20 °C–27 °C and later followed descending trend and was more evident in case of eCO₂ than aCO₂. Several workers reported the influence of CO₂ (Kuo et al., 2006) and temperature (Aleosfoor and Fekrat, 2014) on life table parameters of various aphids and it was found that non-linear trend was more evident. In our study eCO₂ influenced the ‘rm’ of *A. craccivora* which was contradictory to Flynn et al. (2006) who reported that eCO₂ did not cause the increase of ‘rm’ of *R. padi* and *M. persicae*.

Tritrophic interactions studied were at one temperature (27 °C) considered to be optimal. *M. sexmaculatus* were allowed to feed on *A. craccivora* reared on cowpea plants at eCO₂ and aCO₂ and the two vital parameters viz., grub duration and predation capacity calculated to vary significantly between eCO₂ and aCO₂. The findings reflected the influence of CO₂ and temperature on coccinellids tentatively and further investigations are required to quantify the tritrophic interactions across six temperatures and two CO₂ levels.

Decreased grub duration (23%) and increased predation capacity (19%) of *M. sexmaculatus* on *A. craccivora* at elevated over eCO₂ in the present study was similar to increased aphid consumption rate of adult *Harmonia axyridis* (Pallas) in elevated CO₂ treatment reported by Chen et al. (2005). Prey preference of natural enemy varies with host quality mediated at eCO₂. Chen et al. (2007) reported increased aphid abundance (*Sitobion avenae* Fab.) at elevated CO₂ over ambient CO₂ with 26–42.6% increase and with higher predation (23%). Reports of significantly increased consumption of aphids by first-instar *Chrysoperla sinica* (Tjeder) under high CO₂ than under low CO₂ levels and variation in consumption of aphids with instars of coccinellids exist (Gao et al., 2010).

Conclusions

Direct effects of temperature at all three trophic levels and host-mediated effects of eCO₂ at second and third trophic levels imply an increasing incidence of aphids and its predators under anticipated changing climate with positive predator-prey density relations upholding the generalized ecological processes. However, rates of adaptation of organisms at each trophic level of crop-pest-predator and interaction thereof to changing climate accounting other abiotic variables such as rainfall would determine the status of aphids as pests or potential of coccinellids as predator. Parallels noticed across crops relating to influence of eCO₂ and eTemp signify an universal trend of impact on other trophic levels and the established results of tritrophic (dicot plant-assimilate reducing insects & predator) interactions can serve as model system for similar organisms in respect of each trophic levels.

Author’s contribution

Conceived and designed the experiments-MSR, SV. Performed the experiments OS. Analyzed the data OS, PS, AVM. Contributed reagents/materials/analysis tools- MV, MM, KSR. Wrote the paper- MSR, PS, SV.

Conflict of interest

The authors declare that they have no conflict of interest with any scientist or any firm/agency.

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