Lethal and sub-lethal effects of *Beauveria bassiana* (Cordycipitaceae) strain NI8 on *Chrysoperla rufilabris* (Neuroptera: Chrysopidae)

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**Abstract**

A Mississippi Delta native strain (NI8 ARSEF88889) of *Beauveria bassiana* (Bals.-Criv.) Vuill. (Cordycipitaceae), isolated from *Lygus lineolaris* (Palisot de Beauvois) (Hemiptera: Miridae), was tested on green lacewings, *Chrysoperla rufilabris* (Burmeister) (Neuroptera: Chrysopidae) at 4 spray concentrations (7.02 × 10⁴, 10⁵, 10⁶, and 10⁷ spores per mL) to evaluate effects on reproductive rates and adult life expectancy of this insect predator. The application method simulated atomized spray, and concentrations tested were similar to those used to measure impacts of the fungus on *L. lineolaris*. Significant effects of *B. bassiana* on *C. rufilabris* adults were found, and the severity of impact depended on the concentrations tested. *Beauveria bassiana* impacted all demographic measurements of *C. rufilabris* reproduction and survival. Intrinsic and finite rates of increase and gross and net reproductive rates of adults treated with the highest concentrations tested were significantly decreased, whereas doubling time increased for adults treated with the lowest test concentrations. Based on these observations, *C. rufilabris* will be affected by sprays of *B. bassiana* targeted at *L. lineolaris* if adults are present at the time and location of treatment. The measured lethal concentration, LC₅₀, of 2.11 viable spores per mm² compares to an LC₅₀ of 2.75 spores per mm² determined previously for *L. lineolaris*. Higher concentrations of spores per mm² were required for sporulation (SRₐ) of the entomopathogenic fungus on *C. rufilabris* (13.60 viable spores per mm²) than concentrations required for mortality (LC₅₀).

**Key Words**: lacewing; entomopathogenic fungus; demographic parameter; life expectancy; solid diet

**Resumen**

Se hizo un biosenso utilizando una cepa nativa del Delta de Mississippi (NI8 ARSEF88889) de *Beauveria bassiana* (Bals.-Criv.) Vuill. (Cordycipitaceae), aisladada de *Lygus lineolaris* (Palisot de Beauvois) (Hemiptera: Miridae), en una crisopa, *Chrysoperla rufilabris* (Burmeister) (Neuroptera: Chrysopidae) a 4 concentraciones de pulverización (7.02 × 10⁴, 10⁵, 10⁶, and 10⁷ esporas por mL) para evaluar su efecto sobre la tasa reproductiva y duración de la vida de los adultos de este predador de insectos. El método de aplicación simuló pulverización atomizada y las concentraciones probadas fueron similares a las utilizadas para medir el impacto del hongo en *L. lineolaris*. Se encontraron efectos significativos de *B. bassiana* en adultos de *C. rufilabris* y la gravedad del impacto dependería de la concentración probada. *Beauveria bassiana* impactó todas las medidas demográficas de la reproducción y sobrevivencia de *C. rufilabris*. La tasa de aumento intrínseco y finito y las tasas de reproducción bruta y neta de adultos tratados con las concentraciones de prueba más altas disminuyeron significativamente, mientras que el tiempo de duplicación aumentó para adultos tratados con las concentraciones de pruebas más bajas. Basándose en estas observaciones, *C. rufilabris* si será afectado por aerosoles de *B. bassiana* dirigidos a *L. lineolaris* si los adultos están presentes en el momento y lugar del tratamiento. La concentración letal medida, LC₅₀, de 2.11 esporas viables por mm² se compara con una LC₅₀ de 2.75 esporas por mm² determinada previamente para *L. lineolaris*. Se requirieron mayores concentraciones las esporas por mm² para la respuesta de esporulación (SRₐ) del hongo entomopatógeno sobre *C. rufilabris* (13,60 esporas viables por mm²) de las concentraciones que se necesitaron para la respuesta de mortalidad (LC₅₀).

**Palabras Clave**: crisopa; hongos entomopatogenos; parámetros demográficos; esperanza de vida; dieta solida

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Host-predator-entomopathogen interactions in agricultural systems can be synergistically or antagonistically harmful to beneficial arthropods, other non-target insects, and ecological communities (Fuentes-Contreras & Niemeyer 2000; Roy & Cottrell 2008; Meyling et al. 2011). Therefore, the successful use of *B. bassiana* for targeted pest control depends not only on high efficacy against insect pests, but also potential selectivity and low virulence against non-target insects. There are several studies that have demonstrated that *B. bassiana* has been employed with success against a variety of insects in a number of different agro-ecosystems with no significant ecological implications (Lipa 1985; Kimtova & Bajan 1982; Hajek et al. 1987; Weiser 1987; Groden & Lockwood 1991). More recently, Rossini et al. (2014) observed high compatibility of *B. bassiana* and the parasitoid *Cotesia flavipes* Cameron (Hymenoptera: Braconidae) when applied against *Reticulitermes* (Isoptera: Rhinotermitidae), *Metasamia hemipterus* (Coleoptera: Curculionidae), and *Sphenophorus levius* Vaurie (Coleoptera: Curculionidae). Similarly, several studies have shown under laboratory conditions that application of commercial concentrations of *B. bassiana* is compatible with beneficial insects. Thungrabae and Tongma (2007) indicated that *B. bassiana* was found to be non-pathogenic to several natural enemies including *Coccinella septempunctata* L. (Coleoptera: Coccinellidae), *Chrysoperla carnea* Stephens (Neuroptera: Chrysopidae), and *Dicyphus tamaninii* Wagner (Hemiptera: Miridae), and the beneficial soil-dwelling insect *Heteromorus nitidus* (Templeton) (Collemboleta: Entomobryidae). Al mazra’awi (2007) exposed honey bee, *Apis mellifera* L. (Hymenoptera: Apidae), to high inoculum densities of *B. bassiana*, which resulted in very low mortality that was not different from the untreated control regardless of the isolate tested. Two of the strains tested were isolated from *Lygus lineolaris* (Palisot de Beauvois) (Hemiptera: Miridae) collected in Arkansas and New York. Todovora et al. (1996) fed *Coleomegilla maculata* lengi Timberlake (Coleoptera: Coccinellidae) with *B. bassiana* infected Colorado potato beetle *Leptinotarsa decemlineata* Say (Coleoptera: Chrysomelidae) and *B. bassiana* contaminated pollen and found no mortality of *C. maculata*. Levya et al. (2011) found that larvae and pupae of *Chrysoperla exotera* (Navás) (Neuroptera: Chrysopidae) submerged in *B. bassiana* contaminations of spores per g) was suspended in 50 mL of 0.04% Tween-80 (Sigma-Aldrich P8074, St. Louis, Missouri) and diluted to obtain final concentrations of 7 × 10^7 spores per mL. The inoculum viability was measured according to the methodology of Portilla et al. (2014a, 2016). The inoculum concentration (1.20 × 10^5 spores per g) was suspended in 50 mL of 0.04% Tween-80 (Sigma-Aldrich P8074, St. Louis, Missouri) and diluted to obtain final concentrations of 7 × 10^5 spores per mL. The inoculum viability was measured according to the methodology of Portilla et al. (2014a, 2016). Lower test concentrations (7 × 10^4, 10^5, and 10^4) for this study were extrapolated based on dilution of the highest concentration (7 × 10^4). Resulting data were analyzed by analysis of variance (SAS 2013). Aliquots (6 mL) of the highest concentration suspension (7 × 10^5) provided 395 viable spores per mm³ on the targeted sprayed area when applied using a Potter Precision Laboratory Spray Tower (Burkard Scientific, Uxbridge, UK) following the procedures of Portilla et al. (2014a).

**Materials and Methods**

**Colonies of Chrysoperla Rufilabris**

*Chrysoperla rufilabris* adults used in this study were obtained from a commercial supplier (Biocentrol Net Work, Brentwood, Tennessee). About 400 adults (2–3 d old) were received overnight. To ensure copulation, insects were maintained collectively in the original container obtained from the commercial supplier (3 L cylindrical cardboard cotton covered with organydy cloth). A sponge with sugar-water solution (10%) was placed in an 11 cm diameter Petri dish inside the cage. Insects were held in a growth chamber at 25 °C, 55% relative humidity (RH), and a photoperiod of 12:12 h L:D until first oviposition was observed.

**Culture of Beauveria Bassiana Strain N18**

The N18 strain of *B. bassiana* was obtained from stored sources of spore powder maintained at the United States Department of Agriculture (USDA), Agricultural Research Service (ARS), Southern Insect Research Unit (SIMRU). N18 is produced at SIMRU regularly for the *L. lineolaris* research program (Portilla et al. 2016). The inoculum concentration (1.20 × 10^5 spores per g) was suspended in 50 mL of 0.04% Tween-80 (Sigma-Aldrich P8074, St. Louis, Missouri) and diluted to obtain final concentrations of 7 × 10^5 spores per mL. The inoculum viability was measured according to the methodology of Portilla et al. (2014a, 2016). Lower test concentrations (7 × 10^2, 10^3, and 10^2) for this study were extrapolated based on dilution of the highest concentration (7 × 10^5). Resulting data were analyzed by analysis of variance (SAS 2013). Aliquots (6 mL) of the highest concentration suspension (7 × 10^5) provided 395 viable spores per mm³ on the targeted sprayed area when applied using a Potter Precision Laboratory Spray Tower (Burkard Scientific, Uxbridge, UK) following the procedures of Portilla et al. (2014a).

**Bioassay Procedure**

Serial dilutions of 4 test concentrations of N18 strain (1.2 × 10^5, 10^6, and 10^7 spores per mL) were prepared to treat *C. rufilabris* females and evaluate the effect of the N18 strain on their reproductive rates. To avoid cross infection, only 4–5 d-old adult females received from the commercial supplier were used. Selected females were sprayed with N18 using the direct inoculation method (atomized spray delivery) described by Portilla et al. (2014a, 2016). Treated insects were held in a growth chamber at 25 °C, 55% RH, and a photoperiod of 12:12 h L:D. Each assay treatment (individual concentration) was replicated 4 times with 10 adult females per replicate (200 females total). Control insects were sprayed similarly (12.5 kPa per inch²) with 6 mL of water (water control). Treatments of N18 concentrations (1.20 × 10^5, 10^6, and 10^7 spores per mL) were similar
delivered in a 6 mL spray volume of B. bassiana solution. After application, C. rufilabris females were placed individually into a 29.7 mL cups with a solid diet developed for L. lineolaris bioassays (Portilla et al. 2014a). No additional food source was provided. Females were examined daily for mortality and oviposition. The numbers of eggs oviposited every day by each female were counted. Females with eggs were removed and placed in a new diet cup until the last female died. Dead insects were retained in individual diet cups for 10 d and were observed daily for sporulation.

REPRODUCTIVE RATES OF C. RUFILABRIS

Fertility life tables were calculated according to Portilla et al. (2014b). Life fertility tables were determined by selecting age class (x) and the number of females surviving to age x (N). Using these parameters the following model was determined: I = N/N0 (where I is the proportion of females surviving to age x, and where N0 is the number of initial females) (Carey 1993). The daily calculation of age-specific survival rate (lx) and age specific fecundity (mx) was used to estimate net reproductive rate (R = \sum x\, l\, m\, x / \sum x \, l\, x), doubling time (DT = 1n(2)/r), mean generation time (T = \sum x\, l\, x / \sum x \, l\, x), intrinsic rate of increase (r = \sum x\, e^{-l\, x}\, m\, x = 1) and finite rate of increase (λ = e^{\, r\, x}) (Carey 1993; Krebs 2001). Calculations were done by assuming a 1:1 sex ratio, an immature survival of 0.7 (due to their cannibalistic nature) and a developmental time (egg to adults) of 25 d based on quality assessment data obtained from C. rufilabris producers in California (Silvers et al. 2002) and published work by Nordlund & Morrison (1990, 1992), Legaspi et al. (1994), and Giles et al. (2000). By using the fertility tables, reproductive values were calculated, which is defined as the contribution in population numbers that 1 newly hatched individual will make over the remaining life of the female where y and x are age and w is the age of the last successful reproduction (Krebs 2001).

STATISTICAL ANALYSES

One-way ANOVA followed by the Tukey Honest Significant Difference test was used to compare fertility table parameters on C. rufilabris sprayed with different B. bassiana concentrations. Non-parametric estimates of the survival function of C. rufilabris females were compared between treatments by using PROC LIFETEST procedure in SAS (SAS 2013). Statistical differences in the survival of C. rufilabris females were declared based on the log-rank statistic and by using the PROC GLM procedure to detect differences between concentrations at 3, 5, and 10 d after application. Mortality and sporulation data for each group of C. rufilabris females and each concentration were analyzed by PROBIT (SAS 2013) using common logarithm (log to the base 10) of the concentration value.

Results

TIME-MORTALITY RESPONSE OF C. RUFILABRIS TO B. BASSIANA STRAIN NI8

The survivorship of C. rufilabris females treated at different concentrations of B. bassiana strain NI8 is shown in Fig. 1A. Survival was measured through daily post-treatment observations until all females died. Survival rates of treated females varied among the 4 B. bassiana concentrations, where those at higher concentrations died faster than those at lower concentrations. The earliest mortality recorded was observed at the highest concentration (7 × 10^7) followed by 7 × 10^6 at 2 and 3 d after treatment, respectively. Mortality at lower concentrations (7 × 10^4 and 7 × 10^5) was recorded 4 and 5 d after treatment, yet the first mortality in the water control was not recorded until 9 d after application. Mortality analyzed by the test of equality with the strata statement in −log (survival probability) PROC LIFETEST indicated significant differences among concentrations (Log-Rank X^2 = 23.99, df = 4, p < 0.0001) (Fig. 1A).

MORTALITY-RESPONSE AND SPORULATION RESPONSE OF CHYSOPERLA RUFILABRIS TO B. BASSIANA STRAIN NI8

The B. bassiana strain NI8 was pathogenic to C. rufilabris females. However, the levels of mortality and resulting sporulation in cadavers were highly variable between concentrations (Fig. 2). Mortality 3 d af-
Fig. 2. Cumulative mortality of Chrysoperla rufilabris females at 3, 5, and 10 d exposed to Beauveria bassiana strain NI8 at different concentrations (spores per mm²) under laboratory conditions. Insects were fed with a Lygus species solid diet after being sprayed with fungus. Columns within the group labeled with a different letter were significant different at P = 0.05 (Tukey Honest Significant Difference test).

ter spray at the lowest concentration (7 × 10⁴) was 2.5-fold lower than that observed at the highest concentration (7 × 10⁴), but no significant differences were found among those insects exposed to water alone (F = 1.47, df = 4, 199; p = 0.2135). Mortality at 5 d (F = 4.73, df = 4, 199; p = 0.0012), and 10 d (F = 1.47, df = 4, 199; p < 0.0001) after spray was significantly different among treatments. The percentage of individuals resulting in sporulating cadavers at 10 d was significantly affected by spor concentration (F = 43.34, df = 4, 199; p < 0.0001). Sporulation increased with concentration tested (Table 1). Time of sporulation after death was significantly different among treatments (F = 57.87, df = 4, 115; p < 0.0001). Sporulation took longer at lower concentrations. Analyses of concentration–mortality and sporulation responses are shown in Table 2. Chrysoperla rufilabris females were highly affected at low concentrations of B. bassiana (LC₅₀ = 2.11 viable spores per mm²); higher concentration were needed for sporulation (SR₀ = 3.60 viable spores per mm²).

**EFFECTS OF B. BASSIANA NI8 STRAIN ON THE FERTILITY TABLE PARAMETERS OF C. RUFILABRIS**

All demographic measurements for C. rufilabris females obtained from the water controls were significantly higher than those for insects treated with different concentrations of B. bassiana except for doubling time (DT) (F = 2.13, df = 4, 19; p = 0.1402) which, did not differ among treatments. However, high variation among treatment values of DT were observed (Table 3). Water control females doubled their populations in 6.28 ± 0.39 (SE) d and females sprayed with the highest concentration of B. bassiana doubled their population in 20.11 ± 17.21 (SE) d. Total egg production varied among the 4 test concentrations (F = 18.13, df = 4, 19; p < 0.0001). Egg production from females exposed to water control (822.25 ± 141.70 [SE] eggs per 10 females) was 1.68-fold (488.5 ± 156.74 [SE]) and 7.18-fold (114.5 ± 70.35 [SE]) greater than those obtained from females exposed to the lowest and highest concentrations of B. bassiana, respectively. The highest intrinsic rate of increase (rₘ) was found in females sprayed with water alone (0.111 ± 0.003 [SE]) and rₘ values varied significantly among treatments (F = 10.41, df = 4, 19; p = 0.0007). Daily rate of increase of 1.12 females per female per d, a doubling time of 6.28 ± 0.39 (SE), a gross fecundity (Rₒ) of 127.15 ± 44.39 (SE) for female and male eggs per female, and a mean generation time (T) of 31.89 ± 1.81 (SE) d were observed for females exposed to water alone. The mean generation time of females from the water control was significantly higher than all other treatments (F = 3.41, df = 4, 19; p = 0.0289) with a prolonged mean age of reproduction of about 1 and 4 d longer than that of females sprayed with lowest and highest concentrations of B. bassiana, respectively. Females sprayed with the lowest concentration had a gross fecundity (Rₒ) of 66.47 ± 31.19 (SE) eggs per female; those sprayed with the highest concentration had a gross fecundity of 175.1 ± 9.39 (SE) eggs per female. Significantly shorter longevity also was found (F = 21.90, df = 4, 199; p < 0.0001) in treated insects. Females sprayed with the highest concentration lived 6 d shorter than those females sprayed with the lowest concentration and 13 d shorter than females sprayed with water alone (Table 3). Figure 1A, B, and C showed that trends of survival (S(t)), fecundity function (L(m)), and reproductive values (V) were inversely related to spore concentrations. Higher concentrations resulted in lower survival and reproduction.

**Discussion**

The significant differences in −log survival probability among concentrations indicated that C. rufilabris females obtained lethal concentrations of conidia directly from the B. bassiana spray (Fig. 1A). Low mortality and survival noted for insects in the water controls suggests that the Lygus species diet (Portilla et al. 2014a) may be an acceptable diet for rearing C. rufilabris females. Preliminary assays (data not shown) indicated that this predator survived better on the Lygus diet than when females were fed individually with a nutrient-rich slurry consisting of brewer’s yeast, sugar, and water (1:1:1) (Cohen & Smith 1998). Cohen (1993, 1995) explained the extra-oral digestive nature of feeding by Neuropteran predators; predators such as C. rufilabris thrive on solid lipid- and protein-rich diets. Portilla et al. (2016) similarly demonstrated that the Lygus diet facilitated a comparison of pathogenesis and sporogenesis phases of 3 B. bassiana strains tested against Megacopta cribraria F. (Heteroptera: Plataspidae).

Mortality and sporulation are the main evaluation factors used to determine levels of B. bassiana pathogenicity (Portilla et al. 2016). Results presented in this investigation indicated that under laboratory conditions C. rufilabris adult females are highly susceptible to

**Table 1.** Mean (± SD) percentage sporulation in Chrysoperla rufilabris sprayed with 4 concentrations of Beauveria bassiana strain NI8 and fed with a solid Lygus species diet.

| Variable                  | Water control | 7 × 10⁴ | 7 × 10⁵ | 7 × 10⁶ | 7 × 10⁷ |
|---------------------------|---------------|---------|---------|---------|---------|
| Sporulation (%)           | 0 ± 0d        | 17.50 ± 3.84dc | 22.50 ± 4.22c | 67.96 ± 4.74b | 90.00 ± 3.03a |
| Sporulation after dead (d)| 0 ± 0a        | 3.71 ± 1.38b   | 3.44 ± 1.13c | 2.25 ± 1.28b | 2.17 ± 0.87c  |

Means ± standard deviation (SD) followed by the same letter in each row are not significantly different (P < 0.05 Tukey Honest Significant Difference test).
B. bassiana infection by direct exposure. Infectivity and sporulation of entomopathogenic fungi has been shown to increase under high humidity in field, laboratory, and green house conditions (Barson 1976). However, the number of conidia acquired by the host is probably the key factor that increases propagation of conidia by sporogenesis. Mortality and sporulation levels gradually increased when concentrations of conidia increased, even when the humidity condition that occurred in the closed diet cup (>80% RH) was consistent for all treatments (Table 1; Fig. 2). The LC50 reported in this study (Table 2) showed that C. rufilabris mortality could be affected at very low concentrations of B. bassiana strain N18 (2.11 viable spores per mm²), which is comparable to that found for C. lineolaris using the same strain (2.75 viable spores per mm²) (Portilla 2014). Both C. rufilabris and L. lineolaris need higher concentrations of conidia for sporulation (SR), but those needed for sporulation in C. rufilabris (13.60 spores per mm²) were 5.4-fold greater than those needed for L. lineolaris (5.81 spores per mm²) (Portilla 2014). Other chrysopids (Neuroptera: Chrysopidae) including Chrysoperla externa Hagen (Pessoa et al. 2005), C. carnea (Thungrabeb & Tongma, 2007), and Chrysoperla exterior (Navás) (Leyva et al. 2011) also have shown to have concentration-response dependencies to entomopathogenic fungi. Pessoa et al. (2005) observed that C. externa third instar larvae were affected by suspensions of 1.0 × 10⁹ conidia mL of B. bassiana; but, there was no fungal effects on egg viability or developmental time of first and second instar larvae. Leyva et al. (2011) obtained similar results when C. exterior was exposed to different concentrations of B. bassiana. No significant sporulation was measured with 1 × 10⁴ and 1 × 10⁵ on any immature stages, but adults showed 10% mortality 4 d after application with concentrations of B. bassiana.

Table 2. Mortality-response (LC₅₀) and sporulation-response (SR₅₀) of adult female of Chrysoperla rufilabris treated with Beauveria bassiana strain N18 applied at 4 concentrations (± 95% CI [confidence interval]).

| Variable          | Concentration response (spores per mm²) | Slope ± SE | LC₅₀/SP₀ (95% CI) | Slope* | GoFb | Ratio response |
|-------------------|----------------------------------------|-----------|-------------------|---------|------|----------------|
| Mortality         | 0.271 ± 0.047                          | X² = 33.560, P < 0.0001 | 2.110 (0.252–10.832) | X² = 1.1013, P = 0.3551 | 1.000 | 6.443 |
| Sporulation       | 0.396 ± 0.056                          | X² = 48.960, P < 0.0001 | 13.595 (3.397–49.025) | X² = 0.9695, P = 0.4718 | 1.000 |

*Test for slope, significance indicates concentration affects mortality or sporulation (SE = standard error)

**Test for goodness of fit (GoF), significance indicates error from Probit trend is greater than expected for simple binomial response

Measurements of fundamental reproductive components are essential for understanding the population dynamics of C. rufilabris when exposed to B. bassiana. Based on the present results, applications of B. bassiana to C. rufilabris adult females will decrease survival and reproduction. Exposure to higher concentrations will exhibit greater effects (Fig. 1A, B, C; Table 3). According to Donegan (1989) temperature, starvation, and nutrition stresses significantly affect susceptibility of C. carnea to B. bassiana, but nutrition is the most important. With the present study, it should be noted that the use of Lygus diet in this research could impact some aspects of C. rufilabris female biology and behavior such as longevity and estimates of production. However, results in this study were comparable to those on the quality assessment of C. rufilabris producers in California (Silvers et al. 2002), where a female fed with artificial diet deposited more than 200 eggs in her 4 to 6 wk lifespan under laboratory conditions, which is similar to the gross fecundity of 127.15 eggs per female obtained in an approximate 4 wk period (25.17 ± 8.95 SE d) for the water control. It should also be noted that the egg production in the present study was obtained from females that were exposed to males only from emergence to the mating period (2 d after received from commercial supplier). This could explain the shorter longevity obtained in infected females.

In general, the reproductive estimates shown here assumed a hypothetical cohort subjected throughout its lifetime from egg to adult females mortality that could be measured for an actual population of C. rufilabris exposed to the entomopathogenic fungus B. bassiana. The speed at which a population increased (rₚ), is the most important parameter (Carey 1993; Krebs 2001) and C. rufilabris individuals in the water control obtained the highest intrinsic rate value (0.111). The rₚ calculation of C. rufilabris agrees closely with Jokar & Zarabi (2012).

Table 3. Life table statistic for Chrysoperla rufilabris sprayed with Beauveria bassiana strain N18 at different concentrations and fed with a Lygus species solid diet.

| Statistic                        | Water control | 7 × 10⁴ | 7 × 10⁵ | 7 × 10⁶ | 7 × 10⁷ |
|---------------------------------|---------------|---------|---------|---------|---------|
| Longevity (after treated)       | 25.17 ± 8.84a | 18.65 ± 6.67b | 18.85 ± 6.85b | 13.02 ± 5.78c | 12.30 ± 5.95c |
| Total egg¹                      | 822.25 ± 141.70a | 488.50 ± 156.74b | 297.75 ± 44.46bc | 199.00 ± 96.35bc | 114.50 ± 70.35c |
| Gross fecundity²                | 127.15 ± 44.39a | 66.47 ± 31.19ab | 39.59 ± 17.54b | 28.43 ± 15.38ab | 17.51 ± 9.39b |
| Net fecundity³                  | 63.57 ± 22.19a | 33.23 ± 19.60ab | 19.79 ± 8.02b | 14.21 ± 6.69ab | 8.76 ± 5.19b |
| Net reproductive rate⁴          | 28.82 ± 4.49a | 17.09 ± 6.41b | 10.42 ± 1.78bc | 6.96 ± 3.32bc | 4.08 ± 2.32c |
| Mean generation time⁵           | 31.89 ± 1.81a | 30.02 ± 2.15ab | 29.74 ± 2.09ab | 28.41 ± 0.55ab | 27.22 ± 0.48b |
| Doubling time⁶                  | 6.28 ± 0.39a | 7.87 ± 1.76a | 8.83 ± 0.12a | 12.26 ± 3.65a | 20.57 ± 17.21a |
| Intrinsic rate of increase⁷     | 0.111 ± 0.003a | 0.091 ± 0.018ab | 0.078 ± 0.001abc | 0.062 ± 0.020bc | 0.048 ± 0.020c |
| Finite rate of increase⁸       | 1.120 ± 0.007a | 1.090 ± 0.010ab | 1.080 ± 0.001abc | 1.060 ± 0.022bc | 1.050 ± 0.025c |

Means ± SE (standard error) followed by the same letter in each row are not significantly different (P < 0.05 Tukey Honest Significant Difference test)

¹10 females per replicate
²Total offspring per female
³Females per female at age x
⁴Daughters per newly hatched female
⁵Mean age of reproduction (d)
⁶Time required for (λ) to doubling number
⁷Rate of natural increase (daughters per female per d)
⁸Individuals per female per d
when \( C. \ carnea \) was reared under laboratory conditions (\( r_m \) values of 0.074, 0.162, and 0.185) and fed with different media. Amarasekare & Shearer (2013) reported similar \( r_m \) values calculated for \( C. \ carnea \) and \( Chrysoperla \) johnsoni Henry (Neoptera: Chrysopidae) of 0.161 and 0.132, respectively.

This laboratory experiment provides information needed to understand the effect of \( B. \ bassiana \) on \( C. \ rufilabris \). Strain NI8 affects this predator by direct mortality effects and indirect reproductive impacts. The \( r_m \) values reported in this study may vary under field conditions, where chrysopids directly interact with pests and the environment. Further studies are under way that will examine the pathogenicity of \( B. \ bassiana \) strain NI8 to predator and other non-targets arthropods under field conditions. Decisions to deploy NI8 as a biological control for \( L. \ lineolaris \) in different host environments should be based on an overall assessment of ecological and economic benefits and costs.

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