Moringa water extract promising additive to prolong the activity of baculovirus under field-sunlight conditions in Egypt

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(With 1 figure)

Abstract

Baculoviruses are considered as effective biopesticides except of being not active under sunlight conditions. The aim of this study is to evaluate the capability of moringa extract to prolong virus activity under Egyptian field conditions especially that Moringa proved to be strong protective material under previous investigation under laboratory conditions the addition of moringa filters were tested on tomato plant foliage. Results are based on leaf bioassay using Spodoptera littoralis test insect and its nucleopolyhedrovirus (SplINPV) as standard materials. The Original Activity Remaining (OAR) and Lethal Infectivity Time to 50% (LIT50) were estimated after exposure to natural sunlight. cacao and green tea were tested as comparative materials, which proved to be effective as virus protective agent in earlier investigations. The results showed that moringa additive at 10% sustained 50% of virus activity for 193.53 hours and 62.05 and 23.023 hours post application for cacao and green tea; respectively. While virus alone treatment lasts for only 17.551 hours. Moringa generally available, relatively cheap; it also has been tested and proved to be non-toxic, safe, and friendly to the environment. The obtained results showed the activity of moringa water extract in prolonging the virus activity under field application.

Keywords: antioxidants, baculovirus activity, field application, virus protection, moringa.

Aditivo promissor do extrato aquoso de moringa para prolongar a atividade de baculovírus sob condições de campo-luz solar no Egito

Resumo

Os baculovírus são considerados como biopesticidas eficazes, exceto por não estarem ativos sob condições de luz solar. O objetivo deste estudo é avaliar a capacidade do extrato de moringa para prolongar a atividade do vírus sob condições de campo egípcias, tendo em vista que Moringa provou ser um material protetor forte sob investigação anterior em condições de laboratório a adição de filtros moringa foram testados na folhagem de plantas de tomate. Os resultados são com base em bioensaios foliares utilizando o inseto-teste Spodoptera littoralis e seu vírus de poliedrose nuclear (VPNSI) como materiais padrões. A Atividade Original Restante (AOR) e o Tempo de Infectividade Letal até 50% (LIT50) foram estimados após a exposição à luz solar natural. Cacau e chá verde foram testados como materiais comparativos, que se mostraram eficazes como agentes protetores do vírus em investigações anteriores. Os resultados mostraram que a moringa aditiva a 10% sustentou 50% da atividade viral por 193,53 horas e 62,05 e 23,023 horas após a aplicação de cacau e chá verde, respectivamente. Enquanto o tratamento sozinho do vírus dura apenas 17,551 horas, a moringa geralmente está disponível, e é relativamente barata; e a mesma também foi testada e provou ser não tóxica, segura e propícia ao meio ambiente. Os resultados obtidos mostraram a atividade do extrato aquoso da moringa no prolongamento da atividade do vírus sob aplicações em campo.

Palavras-chave: antioxidantes, atividade de baculovírus, aplicação em campo, proteção contra vírus, moringa.

1. Introduction

Several lepidopterous insects are important pests attacking agricultural crops in Egypt. Insect viruses, as biocontrol agents, may play an important role in pest control, thus reducing chemical insecticides. Baculoviruses especially NPV’s are considered promising microbial agents for the biological control of lepidopterous pests. Baculoviruses are safe and selective bio-insecticides, and used worldwide against many insect pests, especially lepidopterous (Singh et al., 2005). During the last 40 years several virus preparations have been developed and distributed. The effectiveness of viral products depends on many biological factors such as infectivity of the pathogen, the host insect behavior.
and the host plant species of the insect (Raymond et al., 2005). These viruses are proved safe due to their specificity. However, their use is hampered by their susceptibility to inactivation by ultraviolet (UV) in sun-light which is a major limitation for field application.

The long-term persistence of baculoviruses relies on their survival in the external environment in the form of occlusion bodies which provide a degree of protection from UV in sunlight. Many pathogens persist in reservoirs, i.e., microhabitats where survival is enhanced, due to protection from the degrading effects of UV irradiation (Krieg et al., 1981; Lasa et al., 2007).

The effect of sunlight and UV light particularly in the wavelength range of 280-320 nm (Chaudhari and Shapiro, 1983) Spli nuclear polyhedrosis virus can be increased by the addition of a caffeinated to apply in filed that lead scientists all over the world to use the economical point of view are lacking.

Studies proved certain protective effects of some material extract, brewers’ yeast (Saccharomyces cerevisiae), light, and ultraviolet radiation (Ignoffo and Garcia, 1990). The persistence of a nucleopolyhedrosis virus can be increased by the addition of certain dyes i.e., cerulean blue (Reddy and Divakar, 2001) Congo red (Baskaran et al., 1998; Shapiro, 1989) Indian ink (Krieg et al., 1980). Fluorescent brightener (Martignoni and Iwai, 1985; Rabindra et al., 1989; Killick, 1990; Koa and Huang, 1992; Shapiro, 1992; Arivudainambi et al., 2000). Congo red (Baskaran et al., 1998). Different groups of chemicals such as shade formulation (Ignoffo and Garcia, 1996). Mixing petroleum spray oils (Mensah et al., 2005). Time of application (Alves et al., 2001). The role of the cationic nature of the berberine molecule and the importance of its absorbance spectrum in (Cohen et al., 2001). Also starch-encapsulated (Ignoffo et al., 1991). The addition of extracts of moringa (Shapiro, 1984). Dar-mol molasses, Shade and Coax (Shapiro et al., 1983). charcoal, yeast extract, brewers’ yeast (Jaques, 1971). Although previous studies proved certain protective effects of some material information about the efficiency in field application and the economical point of view are lacking.

All previous materials found to be expensive or difficult to apply in field that lead scientists all over the world to use natural derived plant antioxidants, the addition of a caffeine derived green tea (Shapiro et al., 2008). An extract of mango leaves (Muralibaskaran et al., 2000) the potentials of vitamins folic acid, pyridoxine, and riboflavin (Ramakrishnan and Chaudhary, 1991) diludin and ionol (Zarin and Eglite, 1985) pantothenic acid, pyridoxine, folic acid and riboflavin protected the virus. Pantothenic acid and pyridoxine (Shapiro, 1985).

Folic acid, riboflavin, pyridoxine, eucalyptus (Deotale et al., 2003). Nano zinc oxide and nano aluminum oxides (El-Helaly and Sayed, 2015) mixing with synergistic effect like spinosa (El-Helaly and El-bendary, 2013) different natural derived plant antioxidants (El-Helaly et al., 2009, 2013) mixing different natural antioxidants (El-Helaly, 2016) this investigation was to complete my promising laboratory results investigations moringa and rice bran in laboratory conditions (El-Helaly, 2013) which proved that moringa promising additive worth to be evaluated under field conditions.

2. Material and Methods

2.1. Test insect

The test insect used in the present investigation is the Egyptian cotton leaf worm, Spodoptera littoralis. It was established on the semi-synthetic diet described by Shorey and Hale (1965) with the exclusion of formaldehyde from diet ingredients and replacing the methyl p-hydroxy benzene with p-methyl benzene.

2.2. Virus inocula

Spodoptera littoralis multiple embedded nucleopolyhedrovirus, Egyptian isolate (SpliMNPV) was the test virus used in the investigation.

2.3. Additives

Moringa, cacao and green tea plant-derived materials were evaluated as UV-protective additives to SpliMNPV suspension. Five or ten gram of each dry plant material was soaked in 100 mL distilled water for 24 hours then blended and filtrated through three layers of muslin and. The filtrate was the stock additive to the tested virus concentration (1x 10⁶ PIB/s/mL) to prepare a final concentration of 5 or 10% of the material additive according to the method described by Shapiro et al. (2008).

2.4. Field experiment

An area under tomato plants about 1/4 feddan (1 feddan = 4200 m²) was used and one small scale field test was set up to confirm laboratory results (El-Helaly, 2013). Two different concentrations of tested additives were prepared (5 and 10% w/v) 12 hr. before spraying and kept in the fridge ready to use. The virus inocula SpliNPV was diluted in distilled de-ionized water and the suspension was adjusted to 10⁶ OBs/mL (=LC 90-95). At the time of field application, the virus and tested additives were thoroughly mixed together and the measured volume was transferred into a hand sprayer. The sticker arabic gum (0.25%) was added to each NPV dilute as a sticking agent. Virus suspension treatments were applied separately to tomato foliage using one litre hand sprayer. Untreated leaves and virus treated leaves were randomly collected at 0, 1, 2, 4, 7, and 10 days post application and kept individually in plastic bags at room temperature until tested. Each leaf was placed into a glass bottle, on which 10 neonate larvae were allowed
to feed for 48 hr. before transferred daily to fresh leaves from the same treatment. Larval mortality was recorded at day 0, 1, 2, 4, 7, 10 days thereafter until day 14. Virus persistence was calculated as % OAR (percentage of Original Activity Remaining) based upon 100% mortality at ‘0’ day post treatment. Potency was calculated for each treatment. Bioassay tests were repeated in five replicates with 10 larvae per treatment (Shapiro et al., 2008). The hydrogen ion concentration (pH) values of all tested natural additive products were measured at 1% concentration using Beckman zeromatic SS-3 pH meter (Beckman instrument, Fullerton, and CA). The absorption values were calculated according to the method described by Finney (1971). Original activity remaining percentages (OARs %) were determined for each treatment according to Muro and Paul (1985) in which NPV- caused larval mortality post UV exposure were divided by NPV- caused larval mortality pre UV exposure and multiplied by 100.

3. Results and Discussion

This experiment was designed to compare and confirm the protective potency of moringa when used with virus sprayed on tomato foliage in the field under the natural conditions of UV in sunlight. In the case of using Sp/LiNPV alone treatment, the recorded rates of mortality among S. littoralis neonate larvae after exposure periods of 10, 24, 48, 96 and 168 hours to natural conditions were 76.00, 42.00, 2.08, 0.00 and 0.00%, respectively, (the calculated LIT50 was only 17.55 hours) (Table 1). In the case of moringa additive at 5%, the rates of mortality were 96.00, 94.00, 86.00, 44.00 and 34.00 respectively, (the calculated LIT50 was 120.915 hours). With cacao additive at 5%, the rates of mortality were 100.00, 84.00, 60.41, 28.00 and 6.00% (the calculated LIT50 was 48.266 hours). With green tea 5% additive, rates of mortality were 76.00, 68.00, 42.00, 14.00 and 4.00%, respectively, (the LIT50 was only 23.47 hours). In the case of moringa additive at 10%, the rates of mortality were 100.00, 96.00, 88.00, 64.00 and 54.00%, respectively, (the calculated LIT50 was 193.53 hours). For cacao at 10% additive; the rates of mortality were 97.77, 89.79, 68.08, 38.00 and 12.00% (LIT50 reached 62.05 hours). With green tea 10% additive the rates of mortality were 78.26, 67.34, 46.44, 22.44 and 2.08%, respectively, (the LIT50 was 23.02 hours). As the data shows that moringa 10% additive increased the persistence of 50% of its activity for almost eight days (193.53 hours 8,063 days), compared to (62.05 hours = 2.58 days) and (23.47 hours = 0.977 days) for cacao and green tea, respectively. (Table 1 the lower concentration 5% also moringa gave the best protection of persistence of 50% of its activity for almost five days (120.915 hours = 5.033 days) compared with (48.266 hours = 2.011 days) and (23.47 hours = 0.959) days) for cacao and green tea, respectively (Figure 1), while virus

<table>
<thead>
<tr>
<th>Irradiation periods (hrs.)</th>
<th>Mortality % due to Sp/LiNPV alone</th>
<th>Mortality % due to Sp/LiNPV alone+</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>moringa</td>
<td>cacao</td>
</tr>
<tr>
<td>Zero time</td>
<td>100.00</td>
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<td>(38/50)</td>
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<td>42.00</td>
<td>84.00</td>
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<tr>
<td>(21/50)</td>
<td>(84/84)</td>
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<td>48</td>
<td>2.08</td>
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<tr>
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<tr>
<td>(0/50)**</td>
<td>(0/50)</td>
<td>(0/50)</td>
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<tr>
<td>LIT50 hrs.</td>
<td>17.551</td>
<td>48.266</td>
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<tr>
<td>Slope +/−</td>
<td>0.4822</td>
<td>0.1240</td>
</tr>
<tr>
<td>Potency(Fold)</td>
<td>6.889</td>
<td>2.747</td>
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</table>

*Refers to either distilled water or additives alone at 5% or 10%; **Between brackets are the no. of virus dead larvae/total no. tested.
treatment alone gave only (17.551 hours = 0.7312 day). Folds of protection (Figure 1 and Table 1) in comparison to virus alone (17.551 hours), it was found that both moringa concentrations were superior where it gave 6.889 folds and 11.026 folds for 5% and 10%; Respectively followed by 2.747, 3.535, 1.337 and 1.311 folds for cacao %%, cacao 10%, green tea 5% and green tea 10%; Respectively. It is also worth to be mentioned that from these data could concluded that both moringa and cacao were more effective when their concentrations increased vice versa green tea gave stable low protection even when its concentration was increased. OAR was also calculated (Table 2) and it reveal the same previous trend of protection where moringa 10% was the best candidate with 54.00% OAR `168 days post application followed by 34.00, 12.24, 6.81, 4.08 and 2.08 OAR % for moringa 5%, cacao 10%, cacao 5%, green tea 10% and green tea 5%; Respectively. When these results compared with my laboratory experiments at previous investigation (El-Helaly, 2013) it was evident that moringa was best protective additive compared to other antioxidants in the enhancement of SpliMNPV activities under the suppressive effect of UV. Testing the polarity of hydrogen ion at the tested concentration (5% or 10%) revealed no apparent role to the pH ion in relation to viral activity as well as counting the polyhedra in mixture of virus + additive. The most effective additive moringa had a pH degree 6.90 followed by 6.85 and 7.1 for cacao and green tea respectively (almost neutral). Also, Argauer and Shapiro (1997) in this respect found no correlation between pH and activity enhancement.

The mode of action of UV-protective additives is measured by its efficiency in absorbance of the ultraviolet light; UV-B region, 280-320 nm, UV-A region, 320-400 nm or both of them (Shapiro, 1989). The success of additive substances was thought to be due to its good absorption in the ultraviolet UV-B as well as UV-A (Shapiro, 1985).

In the present study, the obtained results revealed that all tested additives without significant differences showed a high rate of absorption (from 210-400 nm) especially in UV region of the inactivation 300-320 nm at the 1% concentration (it was undoable to test higher concentrations). In conclusion, there was no correlation between absorption at the UV destructive range and the additive material responsible for protection of virus; this could be referred to using crude materials containing huge complexes of chemical compounds.

Table 2. Effect of Moringa, cacao and green tea additives at both 5 and 10% concentrations on Original Activity Remaining (OAR) of Spodoptera littoralis NPV sprayed on tomato foliage and bioassay against cotton leaf worm neonate larvae.

<table>
<thead>
<tr>
<th>Irradiation periods (hrs.)</th>
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<tbody>
<tr>
<td></td>
<td>moringa</td>
<td>cacao</td>
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<tr>
<td></td>
<td>5%</td>
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<tr>
<td>Zero time</td>
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<tr>
<td>Control*</td>
<td>0.00</td>
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</tbody>
</table>

*Refers to either distilled water or additives alone at 5% or 10%.

Figure 1. Effect of moringa, cacao and green tea additives at both 5 and 10% concentrations on Potency (Fold) of Spodoptera littoralis NPV sprayed on tomato foliage and bioassay against cotton leaf worm neonate larvae.
Finally, moringa found to be the best additives with a marginal difference from the second best candidate (cacao). I recommend to add moringa at 10% concentration for all virus treatment in field and for industrial it should be added for commercial formulation as far. Future studies should be done in order to evaluate the stability of adding antioxidants in formulation under different degrees of temperature and humidity toward getting new stable viral formulation under Egyptian sunny conditions.

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References


KRIEG, A., GRONER, A., HUBER, J. and MATTER, M., 1980. The effect of medium- and long-wave ultraviolet rays (UV-B and UV-A) on insect-pathogenic bacteria and viruses and their...


