Histopathology of cotton bollworm midgut infected with *Helicoverpa armigera* cytoplasmic polyhedrosis virus

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Submitted: July 3, 2012; Approved: April 4, 2013.

Abstract

This research was carried out to examine cytopathological effects of *Helicoverpa armigera* Cytoplasmic polyhedrosis virus (HaCPV) on infected midgut cotton bollworm (*Helicoverpa armigera*) using transmission and scanning electron microscope. The symptoms on infected host larvae of the host, compared with healthy ones, were getting swollen with milky-white and fragile Histopathological examinations showed infection with HaCPV small polyhedral inclusion bodies (PIB) after 1 or 2 days which were observed in columnar cells of midgut. Virions were partially or completely occupied in a polyhedral matrix to form polyhedral inclusion bodies (PIB) at periphery of virogenic stroma. PIBs were measured 0.5 to 3.5 μm and virions about 46 nm in diameter. Microvilli of infected columnar cells were affected and degenerated immediately prior to rupture of the cell. Some infected columnar cells ruptured to release PIB into the gut lumen 3 days after infection. In addition, PIB were found in goblet cells, 5 or 6 days after infection. Infected goblet cells degenerate to such an extent that only a few of the original microvillus-like cytoplasmic projections and cell organelles were left. These cytopathic effects caused in the midgut by HaCPV on cotton bollworm larvae are essentially similar to those have been reported for lepidoperan and dipteran infection by CPV.

**Key words:** cytoplasmic polyhedrosis virus, *Helicoverpa armigera*, histopathology, HaCPV.

Introduction

Most insect reoviruses described to date belong to the family Reoviridae within the genus *Cypovirus* (Cytoplasmic polyhedrosis virus- CPV). These viruses have been reported as pathogens of the midgut epithelium in numerous lepidopteran and dipteran species, because these insects are, respectively, of economic or medical importance (Mertens et al., 2004). CPVs are so-named because they produce large polyhedral occlusion bodies that occlude virions in the cytoplasm of infected cells. They have been recognized as an important entomopathogen, especially among lepidopteran insects because of their potential for biological control (Martignoni and Iwai, 1981). CPVs infect midgut cells of a wide range of insects (Payne and Rivers, 1976). Inclusion bodies after ingestion by insects, dissolve and break, and release infectious virions which enter and replicate in the cytoplasm of midgut epithelial cells (Mertens et al., 1999, 2004). Infection is usually limited to the insect gut wall. Infection frequently results in death or loss of fitness of the host which is reported to be important in regulating host populations (Dwyer et al., 2004). Most CPV infections produce chronic disease with low mortality, although some are pathogenic (Mertens et al., 1999, 2004). *Helicoverpa armigera* Cytoplasmic polyhedrosis virus (HaCPV) (Chinese strain) is a mixture of CPV, so that Belloncik et al. (1996) using cell culture, separated type 14., Li et al. (2006) separated type 5 from a mixture of HaCPVs. They reported that HaCPV is virulent to *H. armigera* and has the potential to be used as a bioinsecticide. HaCPV had negative impact on growth and development of *H. armigera*, and resulted in reduced pupation and pupal weight and an extended developmental period, especially in early instars larvae (Marzban et al., 2009). HaCPV
Sub lethal effects may be due to the diversion of host energy to support or combat the pathogen (Marzban, 2012; Sikorowski and Thompason, 1979; Wiygul, and Sikorowski, 1978, 1991). *H. armigera* is one of the most serious insect pests of cotton, corn, vegetables, and other crops in the Old World including Iran. It has a history of developing resistance to almost all of the insecticides used for its control. Transgenic cotton incorporating Cry1Ac gene derived from *Bacillus thuringiensis* Berliner is one of the most exciting advances made in cotton pest management in recent years. Resistance monitoring of *H. armigera* field populations suggested that there has been some decline in the susceptibility to Cry1Ac in the field (Gunning et al., 2001; Lietal., 2004; Shen et al., 1998). Therefore, alternative control measures, such as biological control agents, are required. HaCPV is a highly infectious insect pathogen (Lietal., 2006; Martignoni and Iwai, 1981) and a candidate for biocontrol of *H. armigera*, especially in combination with *B. thuringiensis* toxins (Iwashita, 1971; Kawai and Miyajima, 1971; Katagiri et al., 1978; Ying, 1986). Presently there is no description of the histopathology of HaCPV on cotton bollworm, *H. armigera*. This study was undertaken to examine the pathological effects of HaCPV on the midgut of cotton bollworm, which forms part of the groundwork for evaluation of HaCPV potential in the integrated control of this pest, for dealing with the decline in the susceptibility of cotton bollworm to Cry1Ac toxin.

**Materials and Methods**

**Insect**

*H. armigera* used in this study obtained from the Wang Mo laboratory, Hua Zhong Agricultural University, Wuhan. For eliminating surface contaminations, the eggs, of each generation, were disinfected by immersing in 2% formaldehyde for 15 min at room temperature, then washed several times with tap water and finally rinsed with sterile distilled water. The eggs were allowed to air-dry on tissue paper and left to hatch in 10 x 6 cm plastic bags at 26 °C .The larvae were individually fed artificial diet at 26 °C and 65% RH, with a 14:10 h photoperiod (Bot, 1966). Adults were fed with a diet sweetened with 10% honey solution.

**Preparation and purification**

HaCPV was provided by Dr Jiang Zhong (Fudan University, Shanghai) that it is a mixture type CPV. *H. armigera* larvae were reared on an artificial diet. The eggs were disinfected, air-dried and incubated as described above. The larvae and adults, also, were fed same as explained above.

The first-instar larvae were infected with HaCPV by spraying a suspension of 3 x 10⁷ polyhedra on the artificial diet and allowed to feed normal artificial diet until newly moulted second-instar (3 day old) larvae. On day 7 after infection, midguts were homogenized in deionized water and strained through 35-μm mesh nylon cloth to remove large debris. The filtrated product was layered on top of HS-40 Ludox continues gradient and centrifuged at 16000 g for 45 min. The resulting band containing purified virus was recovered, washed in sterile distilled water three times and maintained in 0.1 mM NaOH at 5 °C.

**Treatment**

*H. armigera* larvae were placed individually in glass diet tubes (1 x 6 cm) containing a 1-cm diameter cotton leaf discs on moist sterile tissue paper. The discs were disinfected with 0.5% sodium hypochlorite for 10 min and immediately washed three times in sterile water. The discs were treated with 5 μL of viral suspension (3 x 10⁷ polyhedra) with a micropipette prior to adding larvae. Control larvae were fed leaf discs treated with sterile distilled water. Those larvae that consumed whole leaf disc were used for electron microscopy studies.

**Electron microscopy**

Foregut, midgut, hindgut, abdominal lipids, and malpighian tubules of the larvae were dissected at 1, 2, 3, 4, 5 or 6 days after treatment. For transmission electron microscopy (TEM), tissues were fixed in 2.5% glutaraldehyde, washed with PBS and postfixed in cold, buffered, 1% osmium tetroxide, dehydrated in an ethanol series, infiltrated and embedded in Spurr’s resin. Ultrathin sections were obtained with a Leica UC6 ultramicrotome equipped with glass knives, and stained with 0.5% ethanolic uranyl acetate and lead citrate, and examined using a JEM-1230 TEM at 80 kV. For scanning electron microscopy (SEM), tissues were fixed overnight in 2.5% glutaraldehyde, washed as before, dehydrated in an ethanol series, isoamyl acetate for substitution and dried by critical point drying method. Dried tissues were mounted, coated with gold-palladium, and examined using a Hitachi HH-3400 SEM at 30 kV.

**Results**

**Symptomology**

Virus infection noticeably affects the development, feeding, or behavior of the infected larvae. After about 48 h post infection, the larvae become sluggish, move little, and cease to feed. After 3-4 days, the larvae midgut becomes hypertrophied and overgrown with milky-white aspect and fragile (Figures 1 and 2). After about 7 days the larvae die, the body contents liquefies and the skin remained intact without marked change in color as in nuclear polyhedrosis infection.

![Figure 1 - Gut of healthy larvae of *Helicoverpa armigera*](image-url)
Histopathology

Examination of larval gut tissues by electron microscopy revealed viral infection in the cytoplasm of foregut, midgut, and hindgut. There was no evidence of virus particles or polyhedra in the nuclei. Viral particles were localized in regions of the cytoplasm that were tightly packed with inclusion bodies and lacked typical cytoplasmic organelles. In some of the epithelial cells of midgut, these cypovirus areas occupied more than two-thirds of the cytoplasm, although other regions of even heavily infected cells looked unchanged, containing all of the normal cellular organelles. No ultrastructural changes and no polyhedral inclusion body (PIB) were observed in the Malpighian tubules. In contrast, fat bodies of infected larvae contained large number of HaCPV particles. For this publication, the results of midgut infection are presented.

The midgut epithelium of cotton bollworm larvae consists of three types of cells: columnar and goblet cells, which make up most of the midgut epithelium, and the basal regenerative cells (Figure 3). Small PIB (< 1 μm) were observed in some of the columnar cells 1 or 2 days after treatment with HaCPV. The PIB sizes were 0.5 to 3.5 μm. The diameter of virions measures from 41 to 51 nm, with a mean of 46 nm.

On day 3 after infection, infected columnar cells were packed with PIB that virtually fill the whole cell (Figure 4). At this stage of infection the microvilli, which cover the apical ends of columnar cells, were not affected. However, at more advanced stages of infection and just prior to cell rupture, the apical microvilli deteriorated, revealing the bare and distended apices of infected cells (Figure 5). The distended cells eventually ruptured through their apices, releasing PIB into the gut lumen (Figure 6).

On day 4 after infection, deterioration of mitochondria and rough endoplasmic reticulum were observed (not shown). The mitochondria became swollen and eventually disintegrated. There is little or no rough endoplasmic reticulum within infected cells (not shown). At this stage, the nucleus in many of the infected columnar cells was still present (not shown). Examinations of sectioned tissues with the electron microscopes showed that infections occurred in the cells of the fat body. Because adjacent cells usually differed in the degree and stage of infection, the virus in cells in earlier stages of infection varied in size; those in terminally infected cells were uniformly large, an indication that individual virus grow during the infection. On day 5 or 6 after treatment, PIB were observed in some goblet cells (Figure 7). The process of infection of goblet cells is similar to that described for columnar cells. Infected goblet cells degenerate to such an extent that only a few of the original microvillus-like cytoplasmic projections and cell organelles were presented, as compared with control (Figures 7, 8).

Discussion

The midgut epithelium is mainly involved in the absorption of nutrients and other useful substances and their transport. In addition, it has an important role in the removal of unwanted and harmful substances from the body.
The histopathology of HaCPV in the midgut of the cotton bollworm was similar to that of reported for the corn earworm (Bong and Sikorowski, 1991) and the silkworm (Iwashita, 1971; Kawase and Miyajima, 1971) infected with cypoviruses, especially regarding to changes in the columnar cells. Small PIB were found in some midgut columnar cells of the cotton bollworm as early as one day after treatment. Penetration of the Cytoplasmic polyhedrosis virus particles into the cotton bollworm midgut incidented much earlier. In the silkworm, penetration of the virus particles into midgut cells occurs within 10 min of inoculation (Kobayashi, 1971) and small PIB are observed in the midgut cells 48 h later (Kawase and Miyajima, 1971). Apparently, formation of PIB is more rapid in the cotton bollworm than that in the silkworm.

The PIB in cotton bollworm are in general smaller (0.5 to 3.5 μm) than those reported for silkworm and other lepidopterous insects (Aizawa, 1971; Cunningham and Longworth, 1968), the PIB as large as 5 μm occur in larvae of the monarch butterfly, Danaus plexippus (L.) (Arnott et al., 1968). The virion (46 nm) in the cotton bollworm is smaller than the general size range (60-80 nm) reported for CPV (Fenner, 1976). The virion measures 69 nm in the silkworm (Aizawa, 1971) and 67 nm in monarch butterfly larvae (Arnott et al., 1968).

Kobayashi (Kobayashi, 1971) showed that the virogenic stroma is the developmental focus of viral synthesis where formation of PIB takes place. Formation of HaCPV PIB in the cotton bollworm is similar to that in the silkworm that infected with CPV. Empty capsids described by Arnott et al. (1968) in larvae of the monarch butterfly which infected with CPV are not observed in the cotton bollworm. In infected columnar cells, the microvilli, which play a vital role as the absorptive lining of the gut lumen, are severely affected. Whereas, microvilli of infected columnar cells are partially or completely absent and they contain large number of PIB. This differs from that reported for the corn earworm, Helicoverpa zea (Boddie) that infected with CPV (Bong and Sikorowski, 1991).

Most digestive and adsorptive functions of the insect occur in the midgut. So, its damage by HaCPV would adversely affect the normal growth and development of cotton bollworm. Marzban et al. (2009) found that exposure of H. armigera larvae to HaCPV have negative impact on its
growth and development which resulted in reducing pupation and pupal weight and an extended developmental period. Also, viral synthesis requires energy. The large number of PIB within infected cells indicates a great expenditure of the insect’s metabolic energy. Marzban (2012) observed that HaCPV-infected cotton bollworm reduced significantly not only body weight of larvae, but also glycogen, soluble protein, and total lipid content. This is indicative of extensive diversion of normal metabolism for synthesis of viral materials adversely affecting normal functions of the insect.

As infection advances, columnar cells becomes filled with PIB, and so distended that they rupture through their apices releasing PIB into the gut lumen.

Infected columnar cells of the cotton bollworm exhibit additional changes. The mitochondria and rough endoplasmic reticulum degenerate 3 or 4 days after the onset of infection. Kobayashi (Kobayashi, 1971) reported similar observations for rough endoplasmic reticulum in CPV-infected silkworm midgut, but found that mitochondria remained normal except for several enlarged ones near the virogenic stroma. In advanced stages of infection, the nuclei of infected cells are obscured by the large number of PIB.

Goblet cells are infected by HaCPV 5 or 6 days after treatment and undergo serious changes leading them to deterioration. Goblet cells are responsible for potassium ion transport from haemolymph to the gut lumen (Aizawa, 1971). These histopathological data is part of researches evaluating the potential of HaCPV as a biocontrol agent in the integrated pest management of the cotton bollworm.

Acknowledgments

The authors are grateful to Dr. M. Motazeri of Iranian Research Institute of Plant Protection and two anonymous reviewers for their critical review and comments on the manuscript. We are indebted to Dr. Jiang Zhong (Fudan University) who most generously provided cytoplasmic polyhedrosis virus of Helicoverpa armigera for this study; also extend our appreciation to Dr. Wang Mo (Hua Zhong agricultural University) for providing H. armigera pupae. This research was funded by “973” Projects (2006CB100204) and the State Key Research Programs of Science and Technology Ministry (2006BAD08A07-05) of PR China.

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