Toxicological and ultrastructural effects of chitin synthesis inhibitors (lufenuron and chlorfluazuron) on third larval instars integument of Chrysomya albiceps (Diptera: Calliphoridae)

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ABSTRACT

Background: Chrysomya albiceps is of medical and veterinary importance as larvae cause cutaneous myiasis in both man and animals. Chitin synthesis inhibitors (CSI) are commonly used in control of dipterous flies causing myiasis. Several compounds are utilized to interfere with chitin deposition and molting processes during development.

Objectives: In comparison to chlorfluazuron (CF), the current study aims to investigate the toxicological and ultrastructural effects of the CSI, lufenuron (LF) on the biological parameters of the third larval instar (L3) of C. albiceps.

Material and Methods: Early L3 of C. albiceps were fed on diets mixed with four concentrations of LF and CF. Average larval mortality rates were subjected to probit model analysis for calculating LC$_{25}$, LC$_{50}$ and LC$_{90}$. Larval deformation and mortality, percent pupation, adult emergence and chitinase enzyme activity were recorded. Using transmission electron microscope (TEM), ultrastructural study was carried out on non-treated and treated L3 to evaluate LF effects on the integument and muscle layer.

Results: By recording LC$_{50}$ values for LF and CF (146 and 194 ppm, respectively), LF showed more toxic effects on L3 than CF, and at a lower concentration. Reduction in pupation percentage, complete cessation of adult emergence from pupae and decrease in chitinase activity were observed after treatment with all concentrations of both compounds. Ultrastructural changes after treatment with LC$_{90}$ of LF indicated tegumental, nuclear and mitochondrial toxicological effects, and muscle fibers disorganization.

Conclusion: LF proved to be a successful CSI in controlling myiasis causing C. albiceps L3.

Keywords: chitinase, chlorfluazuron, Chrysomya albiceps, integument, lufenuron, myiasis, ultrastructure.

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INTRODUCTION

Chrysomya albiceps is a species belonging to the blowfly family, Calliphoridae$^{[1,2]}$. It is of great medical and sanitary importance as the adults feed on decaying organic matter, human and animal excreta. Its medical importance is associated with cutaneous myiasis produced in both man and animals$^{[3,4]}$. It results in economic damage to cattle breeding by causing primary myiasis$^{[5]}$. This blowfly species is incriminated in spreading of pathogenic microorganisms$^{[6]}$. In South Africa, it is involved in spreading carbuncles, caused by Bacillus anthracis$^{[7]}$.

The wide use of conservative insecticides has led to many problems such as development of resistant strains, environmental drawbacks, and harmful effects on non-target insects as natural enemies, and pollinators$^{[8]}$. Accordingly, it is urgent to search for effective and safe alternative insecticides. Insect growth regulators (IGRs) are insecticides that disturb the activity of the endocrine system regulating molting and metamorphosis processes. Some IGRs inhibit chitin synthesis and deposition in insects, interfering with formation of exoskeletons. These CSI compounds are used as alternatives to traditional insecticides. They are safely biodegradable compounds$^{[9]}$ with minimum effect on natural enemies and their effect is specific to target pests$^{[10]}$. This may explain their classification as biorational insecticides$^{[11]}$. Usage of CSIs deteriorates reproduction and development of adult insects$^{[12,13]}$. Benzoylphenylureas (BPUs), a group of insecticides that belong to the chitin synthesis inhibitors, act mostly on larvae through inhibition of chitin biosynthesis causing abortion of molting$^{[14-17]}$. The possible target of BPUs is the sulfonyleurea receptor in integument epidermal cells that facilitates chitin vesicles transportation$^{[18]}$.

Lufenuron (Match 10%), besides being a CSI insecticide, additionally acts as a juvenile hormone as well as ecysteroid agonist. Moreover, it causes the formation of abnormal new cuticles and death of the insects$^{[12,20]}$. Chlorfluazuron (5%), also an insect growth regulator of the BPUs group, is used to control chewing pests and affects various insect pests, especially lepidopteran larvae$^{[11]}$. The present study evaluates the toxic effect of LF and CF on C. albiceps L3 mortality, larval deformations, pupation percent, and
The two CSIs were exposed to four different concentrations; 120, 140, 160, and 180 ppm for both LF and CF. The procedures were replicated five times for each concentration. Twenty larvae were used for each replicate. Two ml of each concentration were added to 20 gm of meat in the glass jar (6 cm x 9 cm). Water was added to the meat as control experiment. Larvae were transferred to jars containing treated meat. Experiments were carried out under laboratory conditions at 27±2°C, 65±5% relative humidity and 16:8 light:dark cycle. Larval mortality was recorded 24 h after treatments; dead L3 were discarded, and live ones were transferred to other jars containing meat for feeding till pupation. The pupae were counted and moved to new clean plastic jars containing dry wheat bran. The jars were covered by muslin secured with a rubber band. The pupae were sieved from the wheat bran and transferred to rearing cages (30 x 30 x 30 cm) for adult emergence. Emerging adults were supplied with sucrose granules, 10% sucrose solution and offered fresh beef meat (as source of protein and egg deposition medium).

The chitin synthesis inhibitors: The two CSIs were kindly obtained from the Laboratory of Physiology, Plant Protection Research Institute, Dokki, Giza. The LF (Match 10% EC-CAS No. CG A-184699) formula is composed of: N-[2,5-dichlor-o-4-(1,1,2,3,3 hexa-fluoro-propoxyl) phenyl] amino 2,6 difluobenzamid. The CF or Atabraon (5% EC) tested in the present study has the IUPAC name: N-[3,5-dichloro-4-[3-chloro-5-(trifluoromethyl) pyridin-2-yl] oxophenyl carbamoyl]-2,6 difluoro benzamid.

Bioassay and morphological studies: Early L3 of C. albiceps were exposed to four different concentrations; 120, 140, 160, and 180 ppm for both LF and CF. The procedures were replicated five times for each concentration. Twenty larvae were used for each replicate. Two ml of each concentration were added to 20 gm of meat in the glass jar (6 cm x 9 cm). Water was added to the meat as control experiment. Larvae were transferred to jars containing treated meat. Experiments were carried out under laboratory conditions at 27±2°C, 65±5% relative humidity and 16:8 light:dark cycle. Larval mortality was recorded 24 h after treatments; dead L3 were discarded, and live ones were transferred to other jars containing meat for feeding till pupation. The pupae were counted and moved to jars containing autoclaved wheat bran, then placed in cages till adult emergence. Number of emerged adults were counted. Morphological aberrations of treated L3 were determined. Photos of normal and deformed L3 were taken by Canon-Power Shot–G12 camera fixed to a Stereo- microscope (Optika, Italy) after 24 h of feeding on treated diet.

Determination of chitinase activity: The activity of chitinase enzyme was determined according to method described by Ishaaya and Casida. Briefly, 3,5-dinitrosalicylic acid was used for analysis of glucose from chitin digestion. The remaining undigest chitin was centrifuged (Model: PLC-012E, Taiwan) for 15 min at 6,000 rpm. The supernatant was removed for determination of N-acetyl-β-D-glucosamine (NAGA) produced by chitin digestion. The enzyme activity was expressed as µg NAGA x1000/min/gm.

Ultrastructural techniques: Being the most effective CSI, the histopathological effects of LC50 of LF on ultrastructure of the integument of normal and treated larvae was examined using TEM in comparison with untreated L3. Five L3 specimens were dissected, cuticle with underlying muscles were cleaned from any fat bodies and divided into small portions using a sharp blade. The samples were immersed in 3% gluteraldehyde for 24 h, phosphate buffer (pH 7.3) for 1 h, and 1% osmium tetraoxide for 1-2 h. They were dehydrated in series of ethanol to propylene oxide. Semi-thin sections were cut and stained with 0.25% tolidine blue, then examined by light microscopy. Subsequently, thin sections (80 nm) were cut, stained with uranyl acetate and lead citrate, and photographed in a JEOL JEM-2100 TEM (Japan) at 80 Kv in the Electron Microscope unit, Faculty of Agriculture, Al Mansoura University.

Statistical analysis: Data were recorded in Costat statistical software (Cohort software, Berkeley). Lethal concentration was calculated. Data were analyzed using Chi-square test, least significance differences test (LSD), and one-way analysis of variance (ANOVA) at 95% of the upper confidence limit (UCL) and lower confidence limit (LCL). Statistical significance was considered when P<0.05.

Ethical consideration: This study was performed using commercially available chitin synthesis inhibitors (CSIs). No human specimens were examined. No in vivo experiments were conducted. All experimental issues were approved by the Zoology Department, Faculty of Sciences, Zagazig University.

RESULTS

Toxicological and morphological studies: The sensitivity of C. albiceps L3 to the two CSIs under investigation was demonstrated by LC50 values of 146 and 194 ppm for LF and CF, respectively. The insecticidal efficacy increased with increased concentrations. It was observed that LF is 1.3 times more toxic against C. albiceps L3 than CF (Table 1).
Chitin synthesis inhibitors in myiasis

Chitinase activity: Effect of treatment of L3 of *C. albiceps* with LF and CF on chitinase activity was elucidated (Table 3). After 24 h of LF treatment with the median lethal concentration, LC_{50}, a significant reduction (*P*<0.01) in chitinase activity was detected. This activity was recorded as 227.33±3.66 µg NAGA x1000/min/gm fresh weight and reduced by 38.73%. In CF treatment, a significant reduction (*P*<0.01) in activity was estimated as 212.33±1.2 µg NAGA x1000/min/gm fresh weight with reduction percentage of 42.76% in enzyme activity compared with that in control (371±4 µg NAGA x1000/min/gm fresh weight).

Table 1. Toxicity of LF and CF on L3 of *C. albiceps* after 24 h of treatment.

<table>
<thead>
<tr>
<th>Concentrations</th>
<th>LF (ppm)</th>
<th>CF (ppm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>LC_{90} (LCL-UCL)</td>
<td>130 (105-138)</td>
<td>162 (153-169)</td>
</tr>
<tr>
<td>LC_{50} (LCL-UCL)</td>
<td>146 (129-167)</td>
<td>194 (184-210)</td>
</tr>
<tr>
<td>LC_{10} (LCL-UCL)</td>
<td>184 (182-250)</td>
<td>302 (263-387)</td>
</tr>
</tbody>
</table>

Statistical analysis

| Slope ± SE | 12.9 ± 1.0 |
| Chi square (df=4) | 19.03 |
| Relative potency | 1.328 |

LF: Lufenuron; CF: Chlorfluazuron; ppm: Part per million; LCL: Lower confidence level; UCL: Upper confidence limit; SE: Standard error; df: Degree of freedom.

From Table (2), it is apparent that increased concentrations led to increase in larval mortality and deformities, and reduced pupation percentage. Following treatment with 120, 140, 160, and 180 ppm LF, a significant increase (*P*=0.000) in the recorded mean number of larval mortalities were 2.00±0.70, 8.40±0.54, 11.20±1.64, and 19.20±0.83, respectively. While significant elevation (*P*=0.000) in the mean number of larval mortalities were 1.00±0.70, 1.80±1.64, 4.20±0.44, and 8.60±2.19 after treatment with the same concentrations of CF (Table 2). Significant larval deformations were recorded following application of LF, and CF compared with control. The concentration of 160 ppm induced 3.40±0.81 and 1.4±0.89 larval deformities after treatment by LF and CF, respectively. Whereas 120 ppm scored 0.40±0.54 and 1.0±0.00 after treatment with both CSIs, respectively. Significant reduction in pupation percent was recorded after application of both compounds. It was observed that 180 ppm LF has the most pronounced effect by scoring reduction percentage of 96.9% followed by 72.7% after application with 160 ppm of the same compound when compared with control. Significant reduction in pupation percentages (50.50 and 27.30%) were induced by 180 and 160 ppm of CF, respectively (Table 2). Following treatment with all concentrations of the two CSI, complete block of adult emergence was observed.

Treatment of L3 with the two CSIs showed similar larval morphological abnormalities when compared with normal larvae (Fig. 1a). Darkening of the larval cuticle was clearly observed (Fig. 1b, c). Treatment led to formation of twisted or C-shaped larvae (Fig. 1d). Furthermore, larvae with dark brown color were observed (Fig. 1e), while others showed tanned or hardened cuticle with black coloration (Fig. 1f). Presence of larval pupal intermediates was dominant. These intermediates retained some parts of their last larval skin at their anterior end (Fig. 1g); while others failed to complete their pupal period and died maintaining the reduced size (Fig. 1h) and showed shrunken cuticle (Fig. 1i).

Table 2. Effects of different concentrations of the LF and CF on the biological aspects of *C. albiceps* L3.

<table>
<thead>
<tr>
<th>Concentrations</th>
<th>Larval mortality (Mean ± SD)</th>
<th>Larval deformities (Mean ± SD)</th>
<th>Pupation reduction (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>LF CF</td>
<td>LF CF</td>
<td>LF CF</td>
</tr>
<tr>
<td>120 ppm</td>
<td>2.00 ± 0.70pq</td>
<td>1.00 ± 0.70i</td>
<td>0.40 ± 0.54hi</td>
</tr>
<tr>
<td>140 ppm</td>
<td>8.40 ± 0.54i</td>
<td>1.80 ± 1.64i</td>
<td>0.80 ± 0.44i</td>
</tr>
<tr>
<td>160 ppm</td>
<td>11.20 ± 1.64i</td>
<td>4.20 ± 0.44i</td>
<td>3.40 ± 1.81i</td>
</tr>
<tr>
<td>180 ppm</td>
<td>19.20 ± 0.83*</td>
<td>8.60 ± 2.19s</td>
<td>0.40 ± 0.54b</td>
</tr>
<tr>
<td>LSD</td>
<td>1.11</td>
<td>1.65</td>
<td>1.14</td>
</tr>
<tr>
<td><em>P</em> value</td>
<td>0.0000*</td>
<td>0.0000*</td>
<td>0.0000*</td>
</tr>
<tr>
<td>Control</td>
<td>0.00 ± 0.00</td>
<td>0.20 ± 0.20</td>
<td>19.80 ± 0.20</td>
</tr>
</tbody>
</table>

LF: Lufenuron; CF: Chlorfluazuron; LSD: Least significance differences test; *: Significant. Each datum represents the mean and standard deviation (SD) of five replicates. All concentrations showed significant results (*P*<0.05) compared with control. Means within the same column followed by the same superscript are not significantly different (LSD test, *P*>0.05); while means of other concentrations with different symbols within the same column are significant (*P*<0.010).
Table 3. Effect of treatments with LC$_{50}$ of LF and CF on chitinase activity in *C. albiceps* L3.

<table>
<thead>
<tr>
<th>Treatments</th>
<th>Chitinase activity</th>
<th>Statistical analysis</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean ± SD</td>
<td>Inhibition%</td>
</tr>
<tr>
<td>Water (control)</td>
<td>371.00 ± 6.92</td>
<td>0</td>
</tr>
<tr>
<td>LF (LC$_{50}$)</td>
<td>227.33 ± 6.35</td>
<td>38.73</td>
</tr>
<tr>
<td>CF (LC$_{50}$)</td>
<td>212.33 ± 2.08</td>
<td>42.76</td>
</tr>
</tbody>
</table>

LF: Lufenuron; CF: Chlorfluazuron; *: Significant.

**Ultrastructural studies:** The structure of the normal integument of *C. albiceps* larva consists of inner basement membrane, epidermis composed of a single epithelial cell layer, outer cuticle differentiated into an outer epicuticle, exocuticle and an inner endocuticle. Beneath the integument is a clear muscle layer (Fig. 2a). The endocuticle consists of characteristic successive sheets or lamellae (Fig. 2b-d); and cuticle lamellae that consist of microfibrils of chitin arranged horizontally and oriented at a definitive angle to each other (Fig. 3a, b). Such a structure arrangement is referred to as helicoidal architectures. The endocuticle lamellae are interrupted by numerous vertical columns of oriented microfibrils in characteristic structures called pore canal (PC) fibers (Fig. 1c). The PC fibers are chitinous in structure and extend from a structure called the apical membrane protrusion (AMP) (Figs. 2c and 3a). A special pattern of pigmentation was observed in larval endocuticle (Fig. 3c).

The normal epidermis consists of a single layer of columnar epithelial cells with large nuclei (Figs. 2a and 4a) that have obvious free nerve endings (Fig. 3d). The epidermal cells have a distinct supply of numerous tracheal structures (Fig. 4b). The nucleus of each cell is extremely large with well-defined condensed chromatin (Fig. 4a-d). Numerous round shaped mitochondria are located adjacent to the nucleus (Fig. 4c). The cytoplasm contains a distinct rough endoplasmic reticulum (RER), and the lysosomal structures are scattered throughout the cytoplasm (Figs. 4d). Lucent vesicles or areas of rarefied cytoplasm is diagnostic for this species (Figs. 4d and 5a). Microtubules (Fig. 5a) which provide a device for intracellular transport in the cell were obviously observed. These participate in many cellular functions such as the motion of secretory granules and macromolecules inside epidermal cells. Presence of Golgi cristae indicate the secretory function of the epidermal cells for chitin deposition (Figs. 4d, and 5a).

The ultrastructural alterations of cuticle following treatment with median lethal concentration of LF presented as vacuolization (Fig. 5b-d) especially at the exocuticle and between cuticular lamellae at the
endocuticle (Fig. 5c, d). The PC fibres were moderately distorted after larval treatment (Fig. 5d). Borders between cuticular lamellae disappeared from some regions (Fig. 6a). Vacuolation in the cytoplasm of epidermal cells was clearly obvious (Fig. 6b). Autophagic vacuoles and cytoplasm condensation were noticed as signs of apoptosis (Fig. 6b, d). Nuclear changes presented as coagulation in chromatin (Fig. 6c). Strong blebbing of nuclear membrane (Fig. 6d) was a diagnostic cytopathological sign of apoptosis; the chromatin became peripherally located, marginal to the nuclear membrane (Fig. 7 c, d), and most cell organelles disappeared (Fig. 7 b, c). Treatments also induced other cytological deformations of the nucleus which appeared as irregular shape and strong clumping of chromatin material (Fig. 7 b). Vacuolation in the cytoplasm, swallowing endoplasmic reticulum (Fig. 7 a-d), and fragmentation in RER were observed (Figs. 6 b and 7 b). Presence of numerous lysosomes portrayed signs of detoxification due to lysis by foreign substances (Fig. 7 b, d). This was observed as elongation of mitochondria (Fig. 7 d). Damaged organelles and un-degraded debris in autolysosomes were observed (Fig. 6d). Additionally, another sign of apoptosis detected after treatment with LF, was the appearance of phagolysosome (Fig. 6 b) and autophagic vacuoles (Fig. 7 d).

A clear muscle layer is located beneath the epidermal cell layer in normal larvae. This muscle layer has a typical cytological striated structure and clearly distinguished Z lines (Fig. 8a). It helps in contraction and relaxation during larval molting. After treatment with LF there was vacuolation, disorganization and destruction of muscle fibres. Furthermore, abnormal morphology of mitochondria of muscular tissue seen as destruction in inner mitochondrial membrane and mitochondrial cristae were clear (Fig. 8b).

Fig. 4. a) TEM photograph of epidermal cell of normal L3 of C. albiceps showed large irregular nucleus with nuclear membrane, obvious chromatin, rough endoplasmic reticulum and mitochondria. b) TEM photograph of cuticle and epidermal cell of normal L3 of C. albiceps showing numerous tracheae; c) numerous mitochondria near the nuclear membrane and rough endoplasmic reticulum; d) rough endoplasmic reticulum, numerous lysosomes, mitochondria, secretory granules or dense vesicles, Golgi cristae and lucent vesicle.

CH: Chromatin; CL: Chitin lamellae; GC: Golgi cristae; GA: Golgi apparatus; MT: Mitochondria; N: Nucleus; LV: Lucent vesicle; LY: lysosome; RER: Rough endoplasmic reticulum; SG: Secretory granule.

Fig. 5. a) TEM photograph of epidermal cell of normal L3 of C. albiceps showed microtubules, secretory granules or dense vesicles, Golgi cristae and lucent vesicle. b-d) TEM photograph of cuticle in L3 of C. albiceps treated with LC50 LF showed b) vacuolation at endocuticle and between cuticular lamellae; c) exocuticle with vacuoles; d) vacuolation between lamellae of the endocuticle and in pore canal region.


Fig. 6. a) TEM photograph of cuticle of C. albiceps L3 after treatment with LC50 of LF showed disappearance of borders between lamellae of endocuticle; b-d) TEM photograph of epidermal cell of C. albiceps L3 after treatment with LC50 of LF showed b) fragmentation of rough endoplasmic reticulum, numerous lysosomes, secretory granules or dense vesicles, phagolysosome and vacuole; c) disappearance of chromatin, splitting of nucleolus and disappearance of most cell organelles; d) disappearance of chromatin and strong blebbing of nuclear membrane and autolysosomes.

AL: Autolysosomes; D: Desmosome; N: Nucleus; Nu: Nucleolus; LY: Lysosome; PL: Phagolysosome; PG: Pigment; RER: Rough endoplasmic reticulum; V: Vacuole.
DISCUSSION

Nowadays, IGRs are considered effective in the field of insect control by inhibiting the life cycle. They do not persist for long due to their quick biodegradation with low mammalian toxicity. Among them, CSIs inhibit molting process and lead to production of faulty cuticle[26]. The present investigation reported that the larval mortality basically resulted from molting failure with incomplete life cycle and death within the cuticle. Similar results were recorded with Tribolium castaneum[27], Spodoptera littoralis[28,29] and Musca domestica[22,30]. In addition, LF proved to be more toxic at all concentration levels than CF inducing larval mortality, larval deformation, and reduction of pupation percent. Treatment of L3 of C. albiceps with the two CSIs showed high significant mortality after treatment with 180, 160 and 140 ppm of LF and with 180 ppm CF. Furthermore, extremely high significant reduction in pupation percentage was recorded after treatment with 180 and 160 ppm of LF. The results clearly proved that after treatment with the two CSIs, all pupae were dead, and the adult emergence completely ceased.

Previous studies reported the toxic effect of LF and CF against different insects. Percentage of pupal mortality in T. castaneum increased at higher concentrations of CF[31]. Pupal and adult mortalities were reported after treatment of M. domestica with LF while the larval stages were not affected[32]. Lufenuron was highly active against Lobesia botrana eggs[33]. High significant mortality in larvae, pupae, and adults of T. castaneum was recorded at different concentrations of CF. This result could be due to the anti-feeding effect of CF[27]. Lufenuron inhibited adult emergence of S. littoralis after larval treatment[34], and was more potent than CF and chromafenzoide when applied on one day old eggs of Pectinophora gossypiella[35]. Lufenuron proved to be effective against yellow sugarcane borer Diatraea flavipennella[28], and was more toxic against second instar larvae of S. littoralis than flufenoxuron[37]. Chlorfluazuron affected the larval development of Bradysia odoriphaga[36]. In contrast, LF was the least toxic compared with flufenoxuron and hexaflumuron when used against M. domestica[22]. In fact, the majority of insecticides induced variable effects against different insects, because evaluation of the insecticide toxicity is species specific[39].

The two CSIs under investigation in the present study induced L3 abnormalities. The highest percentage of larval deformations was caused by LF. The survived larvae showed two forms of abnormalities, larvae with darkened cuticle and twisted or c-shaped larvae with inability to complete molting and development. Furthermore, small sized and shrunken larvae were clearly observed. Similar deformations were recorded on larvae of M. domestica[22,31,33], T. castaneum[27], L. botrana[24], S. littoralis[24,37], and on B. odoriphaga[38]. Furthermore, the two tested CSIs treatments led to formation of larval-pupal intermediates. These intermediates retained some parts of the last larval skin at their anterior end. They were often completely sclerotized, while others failed to complete their pupal period and died. Similar results were induced after treatment of Ephestia figulilella last instar larvae with hexaflumuron and LF causing deformity in both pupal and adult stages and in some cases produced larval-pupal intermediates. Distorted pupae failed to form the pupal skin, and larval pupal intermediates were
clearly observed\cite{48}. Some treated larvae acquired dark pigmentation at the posterior end of the abdomen\cite{50}. Larval deformation after treatment of \textit{M. domestica} L3 by LF presented as irregular and elongated shapes\cite{50}. Otherwise, LF and flufenoxuron treatments induced full darkened color and curved shaped larvae\cite{59}. Both LF and CF impaired moulting process through significant reduction of chitinase enzyme that is essential in apolysis. This enzyme helps to digest the main constituents of the old endocuticle with the aid of protease enzyme\cite{49}, so that the normal development and transformation into adult stage of \textit{C. albiceps} is blocked. This clarifies why the treated larvae fail to complete their life cycle and acquire the adult stage. This result is in accordance with Bayoumi \textit{et al.},\cite{42} who observed 77.7% and 14.2% reduction in the cuticle chitinase activity after treatment of \textit{S. littoralis} larvae with flufenoxuron and chlorfluanuron, respectively\cite{42}. Reduction in activity of chitinase in \textit{Pectinophora gossypiella} larvae was recorded one-day post treatment of eggs with LF, CF and chromafenzoide\cite{51}. The researchers concluded that the reduction may be due to blocking and inhibition of the enzyme active site.

The insect cuticle is a mechanical barrier that maintains homeostasis and protects the insect against infection and desiccation. The ultrastructural changes of the cuticle revealed structural alterations of the lamellae of endocuticle that caused misplacement of its archetypal structure. Blebbing of nuclear membrane and disappearance of most cell organelles was observed. Another sign of apoptosis is the appearance of phagolysosomes. The use of chitin synthesis inhibitors against insects clearly induces deformities of the cuticle\cite{41,42}. Insects become more vulnerable to contagions by viruses\cite{42,43} and intestinal bacterial\cite{43}.

The larval cuticle of \textit{T. castaneum} unveiled main structural alterations and loss of multilayered array of the procuticle by benzoylureas\cite{40}. Ultrastructural investigations elicited disruption in deposition of procuticle after diflubenzuron application on \textit{Oxya japonica}\cite{40}. Both polyoxin D and diflubenzuron elicited a similar effect on \textit{Lacella cuprina} larvae as the procuticle lacked the normal lamellar arrangement\cite{40}. Analogous results were recorded with the house fly after treatment with chlorfluanuron\cite{41}. The ultrastructural deteriorations of cuticle and epidermis of the sixth instar pharate of \textit{Choristoneura fumiferana} induced by edysone agonist, RH-5992 were revealed. After 24 h the cuticle lacked the usual lamellar arrangement, and the epidermal cells showed alterations in their organelles. Golgi apparatus showed hypertrophied circular vesicle revealing the secretary activity due to secretion of detoxification substances\cite{51}. \textit{Aedes aegypti} larvae treated with novaluron showed abortive discontinuous cuticle that detached from the epidermis with the latter’s degeneration in some cases\cite{39}. Pyriproxyfen treatment causes vacuolization in cuticle and destruction of cuticular lamellae in addition to destruction of nuclear envelop and increase of lysosomes in \textit{Culex pipiens} larvae\cite{64}. Diflubenzuron and LF applications lead to defects in chitin synthesis and organization of \textit{Drosophila melanogaster}\cite{51}. Presence of vacuoles and numerous mitochondria were recorded in 7 days old nymph of \textit{Schistocerca gregaria} following treatment with teflubenzuron. Vacuoles were observed in the epidermal cells and an increase in the number of mitochondria. Mitochondria supply energy important for almost all chemical reactions and hydrogen ion transport mechanisms in the cell\cite{51}. The appearance of vacuoles and autophagic vacuoles following application by CSIs in the current investigation were probably induced as a result of toxification process\cite{52}. Also, vacuoles and phagic vacuoles that appeared in epidermal cells were thought to be an immune response in larvae to exposure to pyriproxyfen\cite{51}. A spectacular increase in RER was detected in \textit{Chrysodeixis chalcites} larvae after treatment with tebufenozide. Furthermore, clear increase of nucleus volume was observed, and presence of numerous mitochondria were demonstrated\cite{53}. Treatment of \textit{Aubeonynus mariae franciscacae} adults with hexaflumuron resulted in obvious disorganization in ultrastructural features of the integument of embryos. The lamellar arrangement of endocuticle was absent. This may be due to defective deposition of chitin leading to death\cite{40}. Treatment of L3 of house fly with LF led to reduction in endocuticle and the epithelial cell layer became disorganized. Moreover, it became difficult to differentiate between exo and endocuticle\cite{51}. After treatment of \textit{S. gregaria} with LF, vacuolation, disorganization and destruction of muscle fibres were observed\cite{61}.

In conclusion the ultrastructural changes induced by the chitin synthesis inhibitor LF on the cuticle, epidermis, and the underlying muscles of L3 of \textit{C. albiceps}, clarify why insects after application become unable to complete their life cycle. Thus, these compounds seem to be valuable in control of this myiasis producing insect and can be used in integrated pest management programs.

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