Impact of the photosensitizers copper and magnesium chlorophyllin on biological and biochemical parameters of *Bulinus truncatus* snail

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**ABSTRACT**

*Bulinus truncatus* snail is the intermediate host of *Schistosoma haematobium*, which causes serious damage to the urinary system of infected patients. To control it in its rapid interruption and/or elimination of the disease transmission would be efficient. Copper chlorophyllin (Cu-chl) and magnesium chlorophyllin (Mg-chl); two photosensitizers used in several biological applications, were bio-assayed against *B. truncatus* snails. Mg-chl proved to be more toxic to the snails than Cu-chl; their LC₉₀ values were 516.7 and 668.9 ppm, respectively. The sub lethal concentrations of each photosensitizer significantly suppressed the fecundity (Mₓ) and reproductive rate (R₀) of treated snails, which could be partially attributed to the recorded disturbances in their biochemical parameters. The reduction rates of R₀ for snails treated with LC₂⁵ Mg-chl and Cu-chl were 83.5% and 50.8%, respectively. Moreover, these photosensitizers exerted marked histological changes in the hermaphrodite gland of treated snails where spermatogonia and oogonia in the gland acini were degenerated, while the connective tissue was disintegrated. Both Mg-chl and Cu-chl exhibited toxic effect on *B. truncatus* snail and interfered with the biological parameters of it that could negatively interrupt the transmission of *S. haematobium*. Therefore, both chemicals could be considered in the control program of this parasite being cheap and environmentally safe.

**INTRODUCTION**

Schistosomiasis is one of the major communicable diseases with socio-economic and health importance in the developing world (Steinman et al, 2006). There are multiple parameters affecting schistosomiasis transmission, among which is the snail intermediate host (Abdel-Hamid and Meckawy, 2014). Snails control by molluscicides could be one of the methods of choice for schistosomiasis control (Rizk et al, 2012). Molluscicides of natural origin have several advantages over the synthetic compounds (Perrett and Whitfield, 1996). In 2004, Ibrahim et al demonstrated a great histological damage in the ovotestis of *Biomphalaria alexandrina* snails post exposure to the plants *Panicum repens* and *Solanum nigrum*.
that resulted in ceasing snails’ oviposition after 4 weeks of exposure. Moreover, Bakry (2009) added that methanol extract of the plants *Euphorbia splendens* and *Agave stylosa* negatively deteriorated the biological and physiological activities of *B. alexandrina* snails. Similarly, Mossalem *et al* (2013) stated that the antimalarial drug artemether has molluscicidal effect against *B. alexandrina* snails. Recently, in semi-field and field trials in Egypt, the plant molluscicide Luowei/TDS 4% proved to be a potent molluscicide against *B. alexandrina* and *Bulinus truncatus* snails (El-Emam *et al*, 2020). Most of the Photosensitizers are used nowadays to control noxious insects and some parasites. Photosensitization is a novel treatment involving the administration of a photoactive compound that accumulates in certain cells of the exposed organism, and if followed by the exposure to visible light, these cells will be killed (Luksiene, 2005). Chlorophyllin, the chlorophyll derivative, is among the recently promising photosensitizers used in this field.

Sodium/ Copper derivative is the most common form of chlorophyllin used as a food additive. The photodynamic process using chlorophyll and its derivatives have been promising agents in the pest control (Erzinger *et al*, 2011). Thus, field investigations using chlorophyll derivatives were carried out to control malaria, filaria and dengue fever vectors in infested epidemic swamps in Uganda, Ethiopia and Sudan (Abdel-kader and EL-Tayeb, 2012). In these investigations chlorophyll derivatives were added to the swamps infested with mosquito larvae *Anopheles gambiae*, thereafter the accumulated photoactive compound (photosensitizer) inside the larva body induced upon sunlight exposure an oxidation stress that resulted in their death. Additionally, Awad *et al* (2008) observed that the photosensitizer hematoporphyrin IX was lethal to mosquito larvae *Culex pipiens* after 5 days of exposure to $1 \times 10^{-4}$ M, and attributed this to the damage of organelles in the cells of treated larvae as a result of the high oxidative stress caused by the photosensitize effect.

Thus, photosensitizers and light (photosensitization) could help in developing novel and environmentally safe effective method for controlling the medically important snails. Therefore, molluscicidal activities of sodium copper chlorophyllin and sodium magnesium chlorophyllin were evaluated against biological and biochemical parameters of *Bulinus truncatus* snails.

**MATERIALS AND METHODS**

**Copper chlorophyllin and magnesium chlorophyllin:**

The photosensitizers’ sodium/copper chlorophyllin (Cu-chl) and sodium/magnesium chlorophyllin (Mg-chl) were kindly provided by prof. Tarek EL-Tayeb, National Institute for Laser Enhanced Science (NILS), Cairo University, Egypt.

**Snails**

Laboratory-bred *B. truncatus* snails used in the present study were from a colony maintained at the Medical Malacology Department, Theodor Bilharz Research Institute (TBRI), Imbaba, Giza, Egypt. They were maintained in aquaria containing de-chlorinated water (24±1°C) and fed oven-dried lettuce leaves. Water was changed once a week and dead snails were removed.
Bioassay tests

A stock solution of 1000 ppm from each of Cu-chl and Mg-chl was prepared, and dilutions were performed for determining LC10, LC25, LC50 and LC90 against *Bulinus truncatus* snails post incubation with these compounds and exposed to light. Therefore, those snails were incubated in the dark with the concentrations of 450,500,550,600,650 and 700 mg/L of copper chlorophyllin for 6 hours and with 250,300,350,400, 500, and 550 mg/L of magnesium chlorophyllin for 6 h (Ragheb *et al.*, 2013). After that they were exposed to light (desk lamp, 100 w/15cm height) for a 3 hours period.

Afterwards, snails were thoroughly washed and transferred to a clean de-chlorinated water to recover for 24 hours. For each concentration, 3 replicates were used, each of which 10 snails (6-8 mm)/L were utilized. Mortality of snails was recorded (WHO, 1993) and analyzed to obtain the lethal concentration values by probit analysis software (WHO, 1965 & Finney, 1970). For control groups, light and dark controls were allowed to run along with the test samples. In light control group, the tested snails were maintained in clean de-chlorinated water under the same experimental conditions. While the dark control involved incubation of the snails with the highest tested concentration of Cu-chl or Mg-chl in the dark under the same experimental conditions without light exposure.

**Effect on snail’s fecundity**

Three replicates of *B. truncatus* (6–8 mm), each containing 10 snails/L, were incubated in the dark with LC10 and LC25 of Cu-chl and Mg-chl once merely at the beginning of the experiment. The incubation period was 6 hours for Cu-chl and Mg-chl (optimum periods for snails to survive incubation) followed by exposure to light for 3 hours (desk lamp, 100 w/15cm height) (Ragheb *et al.*, 2013). Then, the snails were transferred to clean de-chlorinated water for recovery and observation during the following eight consecutive weeks under laboratory conditions (ceiling light, 25±1°C). The dark and light control groups were run parallel to the test groups. The light control snails were kept in clean de-chlorinated water without any treatment, and exposed to light followed by eight weeks of recovery. The dark control snails, incubated with Cu-chl and Mg-chl without light exposure, were transferred to clean de-chlorinated water for recovery. The survivorship of snails (Lx) and the number of laid eggs/snails (Mx) were recorded weekly; the reproductive rate (R0) was calculated at the end of the experiment. Throughout the experimental period the snails were fed oven-dried lettuce leaves and the aquaria were provided with pieces of foam sheets for egg deposition, meanwhile, water was changed weekly.

**Effect on biochemical parameters of snails**

The soft tissues of the surviving snails in the treated and control (light and dark) groups were removed from the shells and homogenized (1 g/2 mL de-chlorinated water) using UP 200H ultrasonic processor, whereas the suspensions were centrifuged at 4,000 rpm for 45 min at 25±1°C. The pellets were discarded while the supernatant was subjected to estimate the activities of transaminase’s enzymes (ALT
and AST) using the Reitman and Frankel (1957) technique, on the other hand, alkaline phosphatase (ALP) was performed according to Bessey et al. (1946), while total protein was determined according to the method of Doumas (1975). In addition, Albumin was determined according to Gustafsson (1976). Calculation of globulin was determined by subtracting the amount of albumin from the total protein.

**Effect on histological features of the hermaphroditic gland**

The snails were treated with LC$_{50}$& LC$_{90}$ of Cu-chl and Mg-chl as mentioned before. Light and dark control groups were simultaneously carried out. Three replicates (10 snails/ L for each) were used for both control and the tested groups. Thereafter, the snail’s hermaphroditic gland was dissected out of their shells and were fixed using Bouin's fixative, then embedded in wax blocks, sectioned (5-8µm), and stained with delafied haematoxyline and eosin (EL-Nahas and EL-Deeb, 2007). Similarly, sections of control snails’ hermaphroditic glands were prepared.

**Statistical analyses**

Statistical analyses were run on IBM compatible PC using SPSS for windows statistical package (SPSS, 2006). Lethal concentrations were calculated using probit analysis software. The mortality rates of experimental groups were compared using Pearson's Chi-square test. Values of biochemical parameters were expressed as mean ± SD. Student’s t-test was applied to locate significant changes between control and treated groups (Sokal and Rohlf, 1995).

**RESULTS**

The molluscicidal activity of Cu-chl and Mg-chl against *Bulinus truncatus* snails was concentration dependent (Table 1). Notably, Mg-chl was more toxic to *B. truncatus* snails than Cu-chl. Their LC$_{90}$ values were 516.7 and 668.9ppm, respectively after 6 hours of dark incubation and 3 hours exposure to light (desk lamp, 100 w/15 cm heights). The slope values indicated that the lethal concentration probability lines (LCP) of these compounds were steep, and their heterogeneity factor was less than 1.0, demonstrating the log-concentration-probit lines to be within the 95% confidence limits, and thus the model fitted the observed data (Kavita et al., 2017).

<table>
<thead>
<tr>
<th>Compound</th>
<th>LC$_{10}$ (ppm)</th>
<th>LC$_{25}$ (ppm)</th>
<th>LC$_{50}$ (ppm)</th>
<th>Confidence limits of LC$_{50}$ (ppm)</th>
<th>LC$_{90}$ (ppm)</th>
<th>Slope</th>
</tr>
</thead>
<tbody>
<tr>
<td>Copper chlorophyllin</td>
<td>473.9</td>
<td>520.1</td>
<td>571.4</td>
<td>492.2-623.7</td>
<td>668.9</td>
<td>1.15</td>
</tr>
<tr>
<td>Magnesium chlorophyllin</td>
<td>273.5</td>
<td>331.1</td>
<td>395.1</td>
<td>336.3-451.1</td>
<td>516.7</td>
<td>1.29</td>
</tr>
</tbody>
</table>

Regarding the egg-laying capacity of *B. truncatus* snails treated with Cu-chl and Mg-chl, the data in Tables 2&3 revealed a gradual decrease in their survivorship (Lx) during the experimental period (8 weeks). The Lx at the 8th week was 0.4 (40%)
and 0.2 (20%) for snails treated with LC25 of Cu-chl and Mg-chl, respectively, compared to 0.7 (70%) for light control group.

**Table (2): Survivorship and fecundity of Bulinus truncatus snails incubated with copper chlorophyllin (Cu-chl) for 6 hours then exposed to light for 3 hours.**

<table>
<thead>
<tr>
<th>Observation Period (week)</th>
<th>Light control</th>
<th>Cu-chl LC10 (ppm) dark control</th>
<th>Cu-chl LC10 (ppm) dark control</th>
<th>Cu-chl LC25 (ppm) dark control</th>
<th>Cu-chl LC25 (ppm) dark control</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Lx</td>
<td>Mx</td>
<td>LxMx</td>
<td>Lx</td>
<td>Mx</td>
</tr>
<tr>
<td>0</td>
<td>1.0</td>
<td>1.4</td>
<td>1.4</td>
<td>1.0</td>
<td>1.4</td>
</tr>
<tr>
<td>1</td>
<td>1.0</td>
<td>3.84</td>
<td>3.84</td>
<td>0.9</td>
<td>1.1</td>
</tr>
<tr>
<td>2</td>
<td>0.9</td>
<td>1.20</td>
<td>0.96</td>
<td>0.9</td>
<td>0.0</td>
</tr>
<tr>
<td>3</td>
<td>0.9</td>
<td>0.44</td>
<td>0.39</td>
<td>0.8</td>
<td>0.0</td>
</tr>
<tr>
<td>4</td>
<td>0.9</td>
<td>2.11</td>
<td>1.90</td>
<td>0.8</td>
<td>0.0</td>
</tr>
<tr>
<td>5</td>
<td>0.9</td>
<td>2.11</td>
<td>1.90</td>
<td>0.8</td>
<td>0.4</td>
</tr>
<tr>
<td>6</td>
<td>0.8</td>
<td>0.0</td>
<td>0.0</td>
<td>0.7</td>
<td>0.5</td>
</tr>
<tr>
<td>7</td>
<td>0.8</td>
<td>2.0</td>
<td>1.60</td>
<td>0.7</td>
<td>1.4</td>
</tr>
<tr>
<td>8</td>
<td>0.7</td>
<td>2.0</td>
<td>1.40</td>
<td>0.7</td>
<td>1.4</td>
</tr>
</tbody>
</table>

R<sub>0</sub>± S.D 13.39±1.10 5.02±0.53 2.28±0.45 8.68±0.55 6.59 ±0.96
Reduction (%) of R<sub>0</sub> 62.5 83.8 35.2 50.8

The fecundity (Mx) of snails incubated with Cu-chl and Mg-chl followed by exposure to light was decreased compared to light control group. Thus, after 8 weeks of recovery, the Mx values considering snails at LC<sub>10</sub> of Cu-chl and Mg-chl were 0.4 and 0.5 eggs/snails/week, respectively, compared to light control group with values of 2.0 eggs/snail/week. Moreover, the snails survived at LC<sub>10</sub> and LC<sub>25</sub> in each of the tested compounds ceasing egg-laying for 3 or 4 weeks during the recovery period. Similarly, dark control group in LC<sub>10</sub> of each compound suffered from ceasing egg-laying during the recovery period and thus, laid few eggs during this period.

**Table (3): Survivorship and fecundity of B. truncatus snails incubated with magnesium chlorophyllin (Mg-chl) for 6 hours followed with exposure to light for 3 hours.**

<table>
<thead>
<tr>
<th>Observation Period (week)</th>
<th>Light control</th>
<th>Mg-chl LC10 (ppm) dark control</th>
<th>Mg-chl LC10 (ppm) dark control</th>
<th>Mg-chl LC25 (ppm) dark control</th>
<th>Mg-chl LC25 (ppm) dark control</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Lx</td>
<td>Mx</td>
<td>LxMx</td>
<td>Lx</td>
<td>Mx</td>
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<tr>
<td>0</td>
<td>1.0</td>
<td>1.4</td>
<td>1.4</td>
<td>1.0</td>
<td>1.4</td>
</tr>
<tr>
<td>1</td>
<td>1.0</td>
<td>3.84</td>
<td>3.84</td>
<td>0.9</td>
<td>0.0</td>
</tr>
<tr>
<td>2</td>
<td>0.9</td>
<td>1.20</td>
<td>0.96</td>
<td>0.9</td>
<td>0.0</td>
</tr>
<tr>
<td>3</td>
<td>0.9</td>
<td>0.44</td>
<td>0.39</td>
<td>0.8</td>
<td>0.0</td>
</tr>
<tr>
<td>4</td>
<td>0.9</td>
<td>2.11</td>
<td>1.90</td>
<td>0.7</td>
<td>0.0</td>
</tr>
<tr>
<td>5</td>
<td>0.9</td>
<td>2.11</td>
<td>1.90</td>
<td>0.7</td>
<td>0.0</td>
</tr>
<tr>
<td>6</td>
<td>0.8</td>
<td>0.0</td>
<td>0.0</td>
<td>0.7</td>
<td>0.53</td>
</tr>
<tr>
<td>7</td>
<td>0.8</td>
<td>2.0</td>
<td>1.60</td>
<td>0.7</td>
<td>1.42</td>
</tr>
<tr>
<td>8</td>
<td>0.7</td>
<td>2.0</td>
<td>1.40</td>
<td>0.6</td>
<td>1.30</td>
</tr>
</tbody>
</table>

R<sub>0</sub>± S.D13.39±1.10 3.54±0.54 2.17±0.45 2.49±0.46 2.21±0.46
Reduction (%) of R<sub>0</sub> 73.5 83.8 81.4 83.5
Fig (1) Reduction (mean % ± S.D) of reproductive rate ($R_0$) of *Bulinus truncatus* snails incubated in the dark with copper chlorophyllin (Cu-chl) and magnesium chlorophyllin (Mg-chl), then exposed to light followed by eight weeks of recovery.

The values of reproductive rate ($R_0$) of treated snails were less than that of light control group, reflecting the reduction in their Lx and Mx values compared to those of the control snails (Fig 1). The reduction rates in $R_0$ of snails at LC$_{25}$ Cu-chl and Mg-chl were 50.8% and 83.5%, respectively.

Table (4) revealed an increase in total protein levels in snails’ tissues incubated with LC$_{50}$ Cu-chl, recording 108.1mg/g tissue compared to 44.3 mg/g tissue of light control group (p<0.01). On the other hand, the snails incubated with LC$_{90}$ Cu-chl suffered from significant reduction in their protein level (21.3mg/g tissue) in comparison with that of light control snails (p<0.01). It was noticed, also, that the pattern of deteriorations in globulin and albumin levels of snails incubated with Cu-chl was approximately similar to that of protein concentrations of these snail groups. Moreover, the levels of total protein, globulin and albumin of the snail groups incubated with Mg-chl were significantly less than those of light control group (p<0.01). The levels of total protein of snails incubated at LC$_{50}$ and LC$_{90}$ Mg-chl were 20.2 & 27.3 mg/g tissue, respectively, compared to 44.3 mg/g tissue of light control snails.

Concerning the activities of the enzymes AST, ALT and ALKP (Table 4), it was noticed that incubation of snails with LC$_{50}$ Cu-chl significantly reduced AST activity, while it was increased by incubation with LC$_{90}$ in comparison with the values of light control group. Similar observation was recorded with snails incubated with LC$_{50}$ and LC$_{90}$ of Mg-chl. The activities of AST for snails incubated with LC$_{90}$ of Cu-chl and Mg-chl were 23.3 and 59.4 U/g tissue, respectively, compared to 19.3 U/g tissue for light control group (p<0.01). For the activity of AKP, incubation of snails with Mg-chl significantly raised it in comparison with light control group, but generally, it was not significantly deteriorated by Cu-chl treatment, except in snail group incubated in dark with LC$_{90}$. The AKP activity for snails at LC$_{90}$ Mg-chl was 12.5 U/g tissue compared to 4.7 U/g tissue for light control group (p<0.01).
Impact of the photosensitizers; copper and magnesium chlorophyllin on Bulinus truncatus

Table (4): Effect of copper chlorophyllin (Cu-chl) & magnesium chlorophyllin (Mg-chl) on biochemical parameters of Bulinus truncatus snails.

<table>
<thead>
<tr>
<th>Treatment</th>
<th>AST U/g tissue</th>
<th>ALT U/g tissue</th>
<th>AKP U/g tissue</th>
<th>Total Protein mg/g tissue</th>
<th>Albumin mg/g tissue</th>
<th>Globulin mg/g tissue</th>
</tr>
</thead>
<tbody>
<tr>
<td>Light/control</td>
<td>19.32±0.44</td>
<td>10.26±0.11</td>
<td>4.7±0.36</td>
<td>44.3±0.31</td>
<td>14.8±0.25</td>
<td>29.5±0.25</td>
</tr>
<tr>
<td>Cu-chl LC&lt;sub&gt;50&lt;/sub&gt; dark control</td>
<td>16.32±0.44**</td>
<td>10.42±0.08</td>
<td>4.1±0.25</td>
<td>98.5±0.44**</td>
<td>39.6±0.36**</td>
<td>58.9±0.63**</td>
</tr>
<tr>
<td>Cu-chl LC&lt;sub&gt;50&lt;/sub&gt;</td>
<td>16.25±0.36**</td>
<td>10.48±0.14</td>
<td>4.44±0.08</td>
<td>108.1±0.44**</td>
<td>45.2±0.1**</td>
<td>62.8±0.44**</td>
</tr>
<tr>
<td>Cu-chl LC&lt;sub&gt;90&lt;/sub&gt; dark control</td>
<td>16.44±0.08**</td>
<td>10.64±0.08</td>
<td>11.63±0.11**</td>
<td>44.6±0.25**</td>
<td>25.0±0.25**</td>
<td>19.6±0.44**</td>
</tr>
<tr>
<td>Cu-chl LC&lt;sub&gt;90&lt;/sub&gt;</td>
<td>23.3±0.25**</td>
<td>15.42±0.11**</td>
<td>4.36±0.08</td>
<td>21.3±0.36**</td>
<td>3.8±0.25**</td>
<td>17.5±0.25**</td>
</tr>
<tr>
<td>Mg-chl LC&lt;sub&gt;50&lt;/sub&gt; dark control</td>
<td>16.26±0.11**</td>
<td>10.42±0.11</td>
<td>10.5±0.25**</td>
<td>17.6±0.25**</td>
<td>3.6±0.25**</td>
<td>13.9±0.36**</td>
</tr>
<tr>
<td>Mg-chl LC&lt;sub&gt;90&lt;/sub&gt;</td>
<td>16.52±0.14**</td>
<td>12.3±0.25**</td>
<td>5.9±0.25**</td>
<td>20.2±0.36**</td>
<td>6.9±0.44**</td>
<td>13.3±0.38**</td>
</tr>
<tr>
<td>Mg-chl LC&lt;sub&gt;90&lt;/sub&gt; dark control</td>
<td>23.34±0.11**</td>
<td>12.6±0.25**</td>
<td>11.36±0.08**</td>
<td>23.8±0.25**</td>
<td>6.9±0.57**</td>
<td>16.9±0.36**</td>
</tr>
<tr>
<td>Mg-chl LC&lt;sub&gt;90&lt;/sub&gt;</td>
<td>59.46±0.18**</td>
<td>15.66±0.08**</td>
<td>12.55±0.08*</td>
<td>27.3±0.25*</td>
<td>8.5±0.36**</td>
<td>18.8±0.44**</td>
</tr>
</tbody>
</table>

**= extremely significant, (p< 0.001).

Examination of the hermaphrodite gland histological sections of control B.truncatus snails monitored numerous acini connected with a connective tissue (Fig. 2A). Each acinus lined with germinal epithelial cells and containing mature ova along its periphery in addition to developed sperms in its lumen.

The hermaphrodite gland transverse sections of snails treated with LC<sub>50</sub> and LC<sub>90</sub> of Cu-chl and Mg-chl (Fig. 2, B-G) revealed varying degrees of degenerations, atrophy and rupture of different cell types, ova and sperms of the glands’ acini. Furthermore, the most prominent damages were clear in several vacuolated ova, degeneration and scattered sperms, as well as degeneration of the acinus’ germinal epithelial layer that led to ceasing snails’ oviposition for some weeks during the experimental period.
Fig (2) A- Light photomicrograph of a transverse section in the hermaphrodite gland of *Bulinus truncatus* (light control) stained with E&H, exposed to light (desk lamp 100 W/15 cm height) for 3 hours (x400). Showing: Ov= mature ova, Sp= developed sperms, Ac= compact acini, Ct= connective tissue, Ep= epithelial cells.

Fig 2: B- Light photomicrograph of hermaphrodite gland transverse section of *B. truncatus* snails treated LC50 Cu-chl and exposed to light, stained with E&H (x400) showing: degenerated ova (ov), deformed sperms (s), rupture of acinus cells (Rc) with vacuoles (v), degeneration of connective tissue (ct).
Fig 2: C-Light photomicrograph of hermaphrodite gland transverse section of *B. truncatus* snails treated LC90 Cu-chl and exposed to light, stained with E&H (x400) showing rupture of cells and vacuoles in the acinus components, degeneration of connective tissue (ct) and ova (ov), damaged and irregular sperms (s).

Fig2: D-Light photomicrograph of hermaphrodite gland transverse section of *B. truncatus* snails treated with LC$_{90}$ Cu-chl in the dark (dark control), stained with E&H (x400), showing: vacuolated ova (ov), damage in developmental stages of spermatogenia (ds), rupture of acinus cells (rc).
Fig 2: **E**-Light photomicrograph of hermaphrodite gland transverse section of *B. truncatus* snails treated with LC$_{50}$ Mg-chl and exposed to light, stained with E&H (x400) showing: degenerated spermatogonia (sp) and ova (ov), damaged sperms (s), atrophy and vacuoles in the acinus content (at).

Fig 2: **F**- Light photomicrograph of hermaphrodite gland transverse section of *B. truncatus* snails treated with LC$_{90}$ Mg-chl and exposed to light, stained with E&H (x400) showing: severe atrophy (at) in the acinus structure with vascular (v), scattered and irregular sperms (s), degenerated ova (ov), degeneration of connective tissue (ct) and epithelial cells (ep).
Impact of the photosensitizers; copper and magnesium chlorophyllin on *Bulinus truncatus*

**DISCUSSION**

The control method would be more efficient and economically valuable in case of selecting to treat the available molluscicide for the particular habitat of snails. The current study determined that Mg-chl was more toxic to *B. truncatus* snails than Cu-chl, a result that agrees with the data of Ragheb *et al* (2013) on the toxicity of Cu-chl and Mg-chl to *B. alexandrina* snails. This could be due to the fact that Cu-chl has a lower photo-reactivity than Mg-chl (Erzinger *et al*, 2011).

The survivorship (Lx) of *B. truncatus* snails after incubation with Cu-chl and Mg-chl, exposure to light and 8 weeks of recovery, was time and concentration dependent. Similar observations were noticed for *B. alexandrina* after exposure to the plants *Collistimon citrinus* and *Zingiber officinale* (EL-Emam *et al*, 2017) and *Haplophyllum tuberculatum* (Rizk *et al*, 2012). Moreover, the survival rate of nymphs of the milkweed bug (*Spilostethus pandurus*) was decreased with increasing the concentration of the photosensitizer hematoporphyrin (Elelimy *et al*., 2016).

The fecundity (Mx) and reproductive rate (R$_0$) of *B. truncatus* snail groups incubated with Cu-chl and Mg-chl, exposed to light and recovered for 8 weeks were significantly less than those of light control group. This could be partially due to their high mortality rates and long periods of ceasing egg-laying during the recovery period. In addition to the interruption of their physiological activities, confirmed by the deteriorations caused by the effect of tested compounds on total protein levels and the activities of enzymes AST, ALT and AKP in their tissues, the decrease or cessation of snails’ egg-laying capacity (Mx) and the net reproductive rate (R$_0$) could be figured out.

**Fig 2:** G- Light photomicrograph of hermaphrodite gland transverse section of *B. truncatus* snails treated with LC$_{90}$ Mg-chl in the dark (dark control), stained with E&H (x400) showing: deleterious effects in acinus structure with vacuoles (v), vacuolated ova (ov), atrophy and degenerated sperms (s).
The upper-mentioned observations coincided with the reduction in Mx and R₀ of *B. alexandrina* snails post incubation with Cu-chl and Mg-chl (Ragheb *et al.*, 2013). In their study, they attributed this condition to the harmful oxidative stress of those compounds on both the regulation of snails' oviposition and the disturbances of their sex hormones (progesterone, testosterone and estradiol) affecting their tissues. El-Ansary *et al* (2001) stated that disturbance of some enzymes in *B. alexandrina* snails treated with sub-lethal concentrations of molluscicides correlated with the reduction of their egg-laying capacity. In addition, Elelimy *et al.*, (2016) noticed that the biological parameters of milkweed bug *S. pandurus* deteriorated through increasing nymphal mortality. They realized a decreasing number of deposited eggs and a reducing percentage of eggs hatchability in post incubation with photosensitizer hematoporphyrin and exposure to sunlight in the summer season.

The total protein concentrations in tissues of *B. truncatus* snails incubated with Mg-chl and exposed to light were significantly reduced compared to light control group. This was, also recorded with the high concentration (LC₉₀) of Cu-chl. These results were confirmed by previous studies relating these changes to disturbances in the internal organs’ function of treated snails to compensate and overcome the toxic stress of such compounds; this phenomenon requires high energy that may stimulate protein catabolism (Morad 2005). The negative effect on protein levels in tissues of *B. alexandrina* and *B. truncatus* snails, post exposed to heavy metals was previously stated by Tolba *et al*., (1997).

The activities of the enzymes AST, ALT and AKP, as biomarkers, may provide more information on the molluscicide induced stress on molluscs. The present deteriorations in the activities of these enzymes in tissues of *B. truncatus* snails treated with Cu-chl and Mg-chl may be due to the destructive stress of such compounds on hepatic tissues and/or the snails’ trials to restore the amino acid balance in their body organs. This conclusion was previously recorded by EL-Emam and Ebeid (1989) in their study on *B. alexandrina* snails treated with mollotox. The high activities recorded for AST and ALT in *B. truncatus* treated with LC₉₀ Mg-chl agreed with the data of Ragheb *et al* (2013) on *B. alexandrina* incubated with Mg-chl and exposed to light. Furthermore, the activities of AST and ALT were altered in hemolymph and tissues of *Helisoma duryi* and *Lynenaea natalensis* snails by low concentrations of copper, therefore they could be used as biomarker for water pollution (Masola *et al*., 2003).

The histological results of *B. truncatus* snails treated with Cu-chl and Mg-chl revealed marked destruction and degeneration of the hermaphrodite gland acini, spermatogonia and oogonia, in addition to a disintegration of the acini connective tissue. The genital organs of treated snails may be sensitive to these sensitizers which could lead to minimize or cease their oviposition. This support the present records on ceasing the snails’ oviposition for some weeks upon exposure to these sensitizers that was mirrored on reduction of the snails’ reproductive rate (R₀). These results were previously stated by Ragheb *et al* (2013) on destruction of the hermaphrodite and
digestive glands cells of *B. alexandrina* snails treated with Cu-chl and Mg-chl and added that this might be due to the harmful effects exerted by such agents during the photosensitization process. Later on, in 2019, Ibrahim and Bakry stated that chlorophyllin extracted from deep-frozen leaves of *Moringa oleifera* plant exerted deleterious effects in the digestive gland of *B. alexandrina* snails treated with LC$_{25}$ of water soluble chlorophyllin, represented by deformation of secretory cells, disintegration of the digestive cells and rupturing the connective tissue between the gland tubules. Moreover, Elelimy *et al* (2016) recorded that light and electron microscopic studies on mid-gut regions of the milkweed bug *S. pandurus* adults resulting from nymphs treated with the photosensitizer hematoporphyrin revealed severe disintegration of cells, many vacuoles, disappearance of most cell organelles and rupturing or detaching of nuclear membrane with clumping of its chromatin material.

The foregoing data revealed considerable molluscidal activity of the photosensitizers Cu-chl and Mg-chl against *B. truncatus* snails. Moreover, these agents are capable of inducing significant deleterious effects on biochemical parameters of the treated snails and on their hermaphrodite gland tissues that was negatively reflected on their fecundity and reproductive rate. These harmful effects will greatly suppress the population size of the snail intermediate host of *S. haematobium* which could disturb and minimize schistosomiasis transmission. In addition, these photosensitizers are inexpensive and environmentally friendly. Therefore, they should be considered in the integrated control program of schistosomiasis to overcome the dangerous drawback of chemical molluscicides.

REFERENCES


