**MOLLUSCICIDAL ACTIVITY OF VARIOUS SOLVENT EXTRACTS FROM Solanum nigrum var. villosum L. AERIAL PARTS AGAINST Galba truncatula**

HAMMAMI H.**, MEZGHANI-JARRAYA R.**, DAMAK M.** & AYADI A.*

### Summary:
Molluscicidal activity of Solanum nigrum var. villosum (morelle velue) extracts and their fractions were tested against the mollusca gastropoda Galba truncatula intermediate host of Fasciola hepatica. The results indicated that the hydro-methanol (MeOH-H₂O) immature fruit extract possess the highest molluscicidal activity (LC₅₀ = 3.96 mg/L) against Galba truncatula compared with other tested compounds. After acido-basic treatment, the methanolic extract fraction isolated from the immature fruits and the richest in alkaloids was the most toxic (LC₅₀ = 1.65 mg/L). The fractions richest in saponosides obtained from the hydromethanolic and methanolic extracts of immature fruits showed interesting molluscicidal activities (LC₅₀ = 6.15 mg/L and LC₅₀ = 7.91 mg/L, respectively). The observed molluscicide activity could be attributed to the presence of alkaloids or saponosides. So, the immature fruits of Solanum nigrum var. villosum could be substrates of choice for molluscicide activity. In addition, total alkaloids and saponosides present in this plant deserve further investigations in order to identify the active principles and demonstrate their activities on mollusks in their natural habitat. According to the World Health Organization’s guidelines on screening for plant molluscicides, use of these fractions may add to the arsenal of methods to control snail transmitting fasciolosis in tropical and Third World countries where fasciolosis is a common disease.

KEY WORDS: molluscicidal activity, Galba truncatula, Fasciola hepatica, Solanum nigrum var. villosum L., alkaloids, saponosides.

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**Fasciolosis** caused by *Fasciola hepatica* (*F. hepatica*), a parasite trematode is of considerable medical and veterinarian importance. This old disease has a great potential of expansion in view of the large colonization capacities of its fasciolid causal agents as well as the freshwater lymnaeid snail vector species (Mas-Coma et al., 2001). The control of Fasciolosis depends on the understanding of the epidemiology of the disease, especially the biology, ecology and distribution of the snails that act as intermediate hosts. It is generally considered that snail control by mollusciciding is of considerable medical and veterinarian importance and can be a rapid and effective means of reducing Fasciolosis transmission (Madsen, 1985). The search for local plants with molluscicidal activity was considered more sustainable than the use of synthetic molluscicides. Plants from Solanaceae family with several species of *Solanum* have been found to have a significant and potentially usefull molluscicidal activity (Medina & Ritchi, 1980; Kloos et al., 1987; Beccouche et al., 2000; Wei et al., 2002; Silva et al., 2006). *Galba truncatula* (*G. truncatula*) has been identified as the main intermediate host of fasciolosis in southern Tunisia (Ayadi et al., 1997, Hammami & Ayadi, 1999; Hammami et al., 2007). Previous studies in our laboratories have shown the high molluscicidal activity.
activity of alkaloids especially the N-methylisosalsoline isolated from *Hammada scoparia*, Chenopodiaceae (Jararraya *et al*., 2009). It was also reported in earlier studies that crude water extracts of *Solamum nigrum* var. *villosum* has molluscicidal and antiparasitic activity against the mollusca gastropoda *G. truncatula* (Hammami & Ayadi, 2008).

In our on going researches for the isolation and the purification of more plant molluscicidal active products, we aimed our study to explore molluscicidal activities of various solvent extracts and fractions from *Solamum nigrum* var. *villosum* aerial parts against *G. truncatula*.

**MATERIAL AND METHODS**

**PLANT MATERIAL**

*Solanum nigrum* var. *villosum* L. 1753, a common annual herbaceous weed (Solanaceae family) locally known as “Enab-El-Dheeb”, was described in many fields of the south-western of Tunisia by Pottier Alapetite (1981). Samples of this plant were collected in June 2006 from Tozeur’s traditional oases, Tunisia and botanically identified by Prof. Ridha Krichen, Botanist at “École d’Horticulture de Chott Mariem, Sousse, Tunisie”. Voucher specimens (LFPMB10) have been deposited at the “Fungal and Parasitic Molecular Biology Laboratory, Faculty of Medicine, Sfax, Tunisia”. Leaves, ripe red and unripe green fruits collected in Tozeur, Tunisia, were carefully detached from the fresh plant and air-dried.

**EXTRACTIONS**

Air-dried leaves, ripe red and unripe fruits of *Solamum nigrum* var. *villosum* were extracted (during 24 hours each time) successively in a Soxhlet apparatus by solvents of increasing polarity: dichloromethane, methanol and then lixiviated with methanol-water (8:2, v-v) at room temperature.

All extracts were filtered through paper Whatman No.1 and then concentrated under reduced pressure to afford the dichloromethane extract (DE) and the methanolic extract (ME). For the methanol-water (8:2, v-v) extract, methanol was removed under reduced pressure and the remaining aqueous phase was lyophilized to afford MeOH-H$_2$O extract. Each extract was weighed (Table I) and kept in dark at 4 °C until use.

**PHYTOCHEMICAL SCREENING**

The preliminary phytochemical screening was performed according to the Harborne (1964, 1973) methods. Plant extracts (dichloromethane, methanol and methanol-water) were subjected to chemical tests for the presence of sterols, triterpenoids, carotenoids, tropolone, quinones, alkaloids, flavonoids and saponosides. A sample of each extract or fraction obtained is dissolved in a minimum amount of solvent adequate for a solution E. This will be added to various reagents according to the protocols experimental T$_1$, T$_2$, T$_3$, T$_4$, T$_5$ and T$_6$.

T$_1$: reaction of Liebermann: 1 ml E + 0.2 ml of acetic anhydride + 4 drops of H$_2$SO$_4$ (cc). Violet changing to blue-green indicates the presence of sterols and (or) triterpenoids.

T$_2$: reaction of Carr and Price: 1 ml E + 4 drops of SbCl$_3$ 20 % in CHCl$_3$. Coloring fleeting blue to blue violet could be a sign of the presence of carotenoids and (or) triterpenoids.

T$_3$: reaction of Wiustater: 1 ml of E + 0.2 ml of MeOH + 1 drop of FeCl$_3$ (0.005 M) + 0.6 ml water + 0.4 ml of CHCl$_3$. Red coloration of the chloroform layer indicates the presence of a tropolone nucleus.

T$_4$: reaction of Borntraeger: 2 ml E + 2 ml of NaOH (0.1 M). If the aqueous phase is colored of red to violet, we can deduce the presence of free quinons.

T$_5$: test of flavonoids: 1 ml of E + 2 ml EtOH-H$_2$O (1-1) + some drops of HCl + Mg chips. Orange to red-violet coloration and blue indicates the presence of flavonoids.

T$_6$: Mayer reaction: 1 ml of E + 0.5 ml HCl (0.1 N) + 5 drops of Mayer reagent. Formation of a white precipitate indicates the presence of alkaloids.

The result with a single reagent being not absolute because there are false positive, we used a second reagent for alkaloids: the Dragendorff reagent revealing red-orange spots of alkaloids on thin layer chromatography plates.

. Mayer reagent: (solution A: 13.5 g HgCl$_2$ + 20 ml water and solution B: 49.8 g KI + 20 ml water). The two solutions are mixed and diluted with water to one liter.

. Dragendorff reagent: formula Munier and Macheboeuf for alkaloids: (solution I: dissolve 0.85 g of under bismuth nitrate in a mixture of 10 ml glacial acetic acid + 40 ml water; solution II: 8 g KI + 20 ml water). Immediately before use, mix 15 ml of solution I and 15 ml of solution II, and add 20 ml of glacial acetic acid and make up to 100 ml with water. On TLC, the total steroidal saponins appear red-purple after spraying a solution of chloroform saturated antimony trichloride, and few minutes after heating the plate to 100-110 °C. All steroidal nuclei are located without distinguishing saponosides and glycoalkaloids (GA). The glycoalkaloids (GA) were revealed as orange spots on TLC by Dragendorff reagent.
The concentration of the chemical groups investigated was scored: - (no reaction), + (weakly positive reaction), ++ (important positive reaction), +++ (very important positive reaction).

**Fractionation of Active Extracts**

An acido-basic treatment of each active extract (except hydromethanolic extract of leaves) was performed as follows: m(g) of each extract were solubilized in hydrochloric solution (0.05 M) then extracted thrice by ethyl acetate (EtOAc). Following evaporation of the ethyl acetate, we obtained the extract A ($m_A$). The aqueous layer was alkalized by addition of ammonia then extracted by ethyl acetate until negative Mayer test. After evaporation of the ethyl acetate, we obtained the extract B ($m_B$) containing total alkaloids revealed as orange spots on TLC by Dragendorff reagent. Then, the remaining aqueous layer was extracted thrice by butanol. After evaporation of the butanol, we obtained the extract C ($m_C$) containing polar products such as saponins, which were revealed as red-purple spots on TLC by SbCl$_3$/CHCl$_3$ (Aubert et al., 1989). Finally, the remaining aqueous layer was lyophilized to produce the extract D ($m_D$).

**Snails**

Uninfected adult *G. truncatula* (3-5 mm in length) were collected locally from a deep draining canal isolated from definitive host, located in the Tozeur's traditional oases in February 2007 and transferred in laboratory aquaria and acclimatized for a minimum period of four days, in holding tanks containing aerated, dechlorinated tap water and washed sand, before being exposed to the aqueous extracts. The toxicity of these preparations was also tested on *G. truncatula* control organisms.

**Molluscidal Tests**

Evaluation of the molluscidal activity of *Solanum nigrum* var. *villosum* extracts and CuCl$_2$ (used as positive control) against snails was investigated as recommended by the World Health Organization (1965). The CuCl$_2$ solution was prepared by adding dechlorinated water to 1 g of CuCl$_2$ up to 1 liter. The extract or fraction of each plant part and of CuCl$_2$ was used to produce a series of concentrations so that the molluscidal potency of each sample could be evaluated. The snails were exposed, in groups of ten (five replicates) for 24 to 48 hours, to 500 ml of each concentration of one of the tested materials: leaf, ripe red and unripe green fruit extracts, or CuCl$_2$ (positive control product) shown in Table III.

*Galba truncatula* negative control organisms were similarly immersed in dechlorinated water. After exposure, snails were rinsed thoroughly in dechlorinated water and left for 48 hours in dechlorinated water before mortality was evaluated.

Mortality was recorded after 24 to 48 hours and dead animals were removed immediately to avoid contamination of animals. Snail mortality was established by the contraction of the body within the shell; no response to a needle probe was taken as evidence of death.

**Statistics**

The concentrations that would kill 50 % (LC$_{50}$), 90 % (LC$_{90}$) of the exposed snails and the confidence interval (95 % CI), were determined by the R language analysis which is an integrated suite of software facilities for data manipulation, calculation and graphical display (Ihaka & Gentleman, 1996).

**Results**

The mass and yields of different plant part extracts of *Solanum nigrum* var. *villosum* after 24 hours extraction with increasing polarity solvents were given in Table I.

The phytochemical tests performed for alkaloids and saponosides were positive for the methanolic extracts and important for the MeOH-H$_2$O (8:2, v-v) extracts. We detected sterols and terpenoids in dichloromethane extracts. The flavonoids were present in the hydromethanolic extracts (Table II).

The molluscidal activities of *Solanum nigrum* var. *villosum* extracts and the cupric chloride against the mollusca *G. truncatula* were given in Table III. Toxic

| Initial mass of plant parts (g) | Leaves (114 g) | Mature fruits (400 g) | Immature fruits (200 g) |
|-------------------------------|---------------|-----------------------|-------------------------|
|                               | Mass (g)      | Yield (%)             | Mass (g)               | Yield (%)             | Mass (g)       | Yield (%) |
| Dichloromethane extract (CH$_2$Cl$_2$) | 4.21 | 3.69 | 0.40 | 0.1 | 0.18 | 0.09 |
| Methanol extract (MeOH)        | 18.06 | 15.84 | 21.72 | 5.43 | 8.31 | 4.15 |
| MeOH-H$_2$O extract (8:2, v-v) | 0.05 | 0.04 | 2.45 | 0.61 | 0.24 | 0.12 |

Table I. – Mass and yields of different plant part extracts of *Solanum nigrum* var. *villosum* after 24 hours extraction with increasing polarity solvents.
activity of this plant was determined for dichloromethane, methanolic and hydro-methanolic extracts. The hydro-methanolic extract of the immature fruits, after 48 hours, showed the highest activity with values for LC$_{50}$ = 3.96 mg/L and LC$_{90}$ = 7.49 mg/L. By comparison, we obtained values of LC$_{50}$ = 15.19 mg/L and LC$_{90}$ = 18.85 mg/L for the methanolic extract and LC$_{50}$ = 34.41 mg/L and LC$_{90}$ = 50.47 mg/L for the dichloromethane extract. For leaves, the hydro-methanolic extract (LC$_{50}$ = 15.92 mg/L) and the methanolic extract (LC$_{50}$ = 17.51 mg/L) showed interesting toxicity compared with the dichloromethane extract (LC$_{50}$ = 31.37 mg/L). Mature fruit extracts show the lowest toxicities with LC$_{50}$ = 50.84 mg/L for the dichloromethane extract and LC$_{50}$ = 84.83 mg/L for the hydro-methanolic extract. The methanolic extracts of mature fruits were inactive. Cupric chloride, used as a reference substance, killed 90% of mollusks at 11.34 mg/L after six days of treatment. Control assays with dechlorinated water showed no effect on the snails (Table III). Haemolysis and hyper secretion of mucus were the common toxic responses of snails to the active tested materials. The molluscicidal activity of methanolic and hydro-methanolic extracts of Solanum nigrum after acido-basic treatment were reported in Table IV. The highest toxicity, after 48 hours, was shown with immature fruit fractions: the fractions B (LC$_{50}$ = 1.65 mg/L) and C (LC$_{50}$ = 7.91 mg/L) of the methanolic extract and the fraction C (LC$_{50}$ = 6.15 mg/L) of the hydromethanolic extract. Weaker molluscicidal activities were obtained for the fraction C of leaf methanol extract (LC$_{50}$ = 13.6 mg/L) and mature fruit hydromethanolic extract (LC$_{50}$ = 14.29 mg/L). The molluscicidal activity disappeared for the fraction A of each fractionated extract.

| Phytochemical test     | T$_1$ (sterols; triterpenoids) | T$_2$ (carotenoids; triterpenoids) | T$_3$ (tropolone) | T$_4$ (quinone; flavonoids) | T$_5$ (flavonoids) | T$_6$ (alkaloids; saponosides) |
|------------------------|---------------------------------|-----------------------------------|-------------------|-----------------------------|-------------------|--------------------------------|
| Dichloromethane        | Leaves                          | +                                 | -                 | -                           | -                 | -                              |
|                        | Mature fruits                   | +                                 | -                 | -                           | -                 | -                              |
|                        | Immature fruits                 | +                                 | -                 | -                           | -                 | -                              |
| Methanol               | Leaves                          | -                                 | -                 | -                           | +                 | +                              |
|                        | Mature fruits                   | -                                 | -                 | +                           | -                 | -                              |
|                        | Immature fruits                 | -                                 | -                 | +                           | +                 | ++                             |
| Methanol-H$_2$O (8:2, v-v) | Leaves                          | -                                 | -                 | -                           | +                 | +                              |
|                        | Mature fruits                   | -                                 | -                 | +                           | +                 | ++                             |
|                        | Immature fruits                 | -                                 | -                 | +                           | +                 | ++                             |

- : no reaction; + : weakly positive reaction; ++ : important positive reaction.

Table II. – Chemical groups present in the extracts from aerial parts of Solanum nigrum var. villosum.
CuCl₂ showed molluscicidal activity against the mollusca gastropoda *Galba truncatula* after 48 hours with LC₅₀ = 26.12 (19.35; 31.69) mg/L and LC₉₀ = 62.71 (49.83; 96.50) mg/L; and after 144 hours with LC₅₀ = 6.20 (5.42; 6.94) mg/L and LC₉₀ = 11.34 mg/L.

Table IV. – Yields, phytochemical tests and molluscicidal activities of *Solanum nigrum* var. *villosum* fractions, after 24 and 48 hours exposure time, against *Galba truncatula*. 

| Initial mass (mi) of extract (g) | Solvent / pH | Activity after 24 H exposure time | Activity after 48 H exposure time | Phytochemical tests |
|---------------------------------|--------------|----------------------------------|----------------------------------|---------------------|
|                                 |              | LC₅₀ (mg/L) (95 % CI)            | LC₉₀ (mg/L) (95 % CI)            | Dragendorff SbCl₅₀ |
| MeOH-H₂O / Mature fruits        |              |                                  |                                  |                     |
| mi = 2.02                       | EtOAc / acid medium | mₐ = 96                          | inactive                          | inactive            |
|                                 | EtOAc / alkali medium | mₐ = 20                          | inactive                          | inactive            |
|                                 | Butanol / alkali medium | mₐ = 28(8.2)                    | 15.54                            | inactiv            |
|                                 |                |                                   | 28(30)                           | inactive            |
|                                 | Aqueous lyophilized layer | mₐ = 1,440                      | 71.28                            | inactiv            |
| MeOH / Leaves                   | EtOAc / acid medium | mₐ = 182.1                       | 26.38                            | inactive            |
| mi = 0.69                       | EtOAc / alkali medium | mₐ = 3.6                         | 0.52                             | inactive            |
|                                 | Butanol / alkali medium | mₐ = 7.3                        | 10.57                            | inactive            |
|                                 |                |                                   | 12.59                            | inactive            |
|                                 | Aqueous lyophilized layer | mₐ = 340                        | 49.27                            | inactiv            |
| MeOH-H₂O / Immatures fruits     | EtOAc / acid medium | mₐ = 26.5                        | 0.18                             | inactive            |
| mi = 0.15                       | EtOAc / alkali medium | mₐ = 6.7                         | 4.47                             | inactive            |
|                                 | Butanol / alkali medium | mₐ = 23.6                      | 15.73                            | inactive            |
|                                 |                |                                   | 10.96                            | inactive            |
|                                 | Aqueous lyophilized layer | mₐ = 90                         | 60                               | inactive            |
| MeOH / Immatures fruits         | EtOAc / acid medium | mₐ = 81.8                        | 12.58                            | inactive            |
| mi =0.65                        | EtOAc / alkali medium | mₐ = 11.1                        | 1.7                              | inactive            |
|                                 | Butanol / alkali medium | mₐ = 181.6                     | 27.94                            | 14.69               |
|                                 |                |                                   | 48.07                            | 7.91                |
|                                 | Aqueous lyophilized layer | mₐ = 340.2                     | 52.34                            | 24.59               |
|                                 |                |                                   | 89.40                            | 20.46               |
| Dechlorinated water             |              | 0                                | 0                                | 0                   |
DISCUSSION

One of the major preventive steps against fasciolosis is the control of the vector snail populations. In recent years, several plant extracts have been studied against snail transmitted parasitic diseases and several species of *Solanum* have been found to have significant and potentially useful molluscicidal activity. Such activity varies greatly from species to species and even between different parts of the same plant (Silva et al., 2005).

During our study, only immature fruit and leaf extracts showed significant molluscicidal activities with LC$_{50}$ values that fell well below the upper threshold of 40 mg/L set for a potential molluscicide by the World Health Organization (WHO, 1993). So that, the immature fruit extracts have the most toxicity against the snail *G. truncatula* correlating with previous studies (Hammami & Ayadi, 2008). By comparison with immature fruit crude water extract of the same plant (*Solanum nigrum* var. *villosum* collected from Tozeur’s oases) obtained values (LC$_{50}$ = 41.2 mg/L and LC$_{90}$ = 66.6 mg/L) in our previous studies, the immature fruit hydro-methanolic extract was more efficacious and demonstrated the importance of the plant components separation.

This result was even supported by Ndamukong et al. (2006) when studying molluscicidal activity of *Solanum scabrum* against the snails Bulinus came-runensis and *B. truncatus* showing that methanolic extracts were more toxic than water extracts.

Methanolic extract was found even possessing a strong molluscicidal activity and exhibited potent effect on mollusks (Lahlou et al., 2002).

In the present study, methanolic and hydro-methanolic extracts of the aerial green parts (leaves and immature fruits) were the most toxic from the whole extracts tested. These findings are in agreement with other data in the literature. For instance, aerial parts methanolic extracts of *Solanum jabrense* were described to have molluscicidal activity with LC$_{50}$ = 56.0 mg/L and LC$_{90}$ = 80.3 mg/L (Silva et al., 2006).

It was even reported that the methanolic extract of the berries of *Solanum aculeastrum* also showed significant activity against host snails of schistosomiasis, the LC$_{100}$ = 50mg/L (Wanyonyi et al., 2003).

According to other studies, the cercaricidal activity was related to the flavonoids, terpenoids, and saponosides (Chifundera et al., 1993; Perret & Whitefield, 1995; Lahlou et al., 2002). It was also reported that the toxicity in *Solanum genera* was accorded to the presence of flavonoids (Silva et al., 2002) and steroid alkaloids in the fruits (Bhattacharyya, 1984; Valverde et al., 1993). Alkaloids of stems of *Solanum stipula-

ceum* were toxic against *Biomphalaria glabrata* LC$_{50}$ = 45.2 mg/L and LC$_{90}$ = 56.0 mg/L (Silva et al., 2006).

In our study, the fractionation has concerned only active extracts showing significant LC$_{50}$ below the upper threshold of 40 mg/L (Table III) fixed by WHO (1993) and that were positive for alkaloid and saponosides phytochemical tests (Table II). Leaf hydromethanolic extract having a fewer yield (0.04 %) was not fractionated (Table I).

During our study, after acido-basic treatment, we noticed that fractions showing molluscicidal activity contained generally alkaloid and/or saponosides (Table IV). Among the eight different total alkaloid and saponosides fractions tested, that from immature fruit methanolic extract was the most potent in *G. truncatula* (LC$_{50}$ = 1.65 mg/L) but it was presented in a weaker yield of 1.7 %. Even in other studies, it was reported that *S. nigrum* immature fruits are not a promising source of the alkaloids because of their small size and declining alkaloid content (Eltayeb et al., 1997). Other workers have reported steroidal alkaloids in *Solanum incanum* fruits at levels ranging from traces (Lin et al., 1990) to 0.031 % (Fukuhara & Kubo, 1991) on a fresh weight basis in mature fruits.

The fractions B less rich in alkaloids from mature fruit hydromethanolic (1 %) and leaf methanolic (0.52 %) extracts were not active against *G. truncatula* and that may be attributed to the fact that total alkaloids were not enough to produce molluscicidal activity.

The fraction C rich in saponosides was active in all our extracts tested and it presented important yields (ranging from 10.57 % to 52.34 %) and interesting molluscicidal activity. The fraction C from hydromethanolic immature fruit extract was found to be the next toxic against *G. truncatula* (LC$_{50}$ = 6.15 mg/L) and after that was classified the fraction C from immature fruit methanolic extract (LC$_{90}$ = 7.91 mg/L). We deduce that molluscicidal activity was related to polar fractions rich in alkaloids and/or saponosides and that the fraction richest in alkaloids from the immature fruits was more toxic than those rich in saponosides.

Saponosides fractions from leaves methanolic extract and mature fruits hydromethanolic extracts have shown weaker molluscicidal activity (LC$_{50}$ = 13.64 mg/L and LC$_{90}$ = 14.29 mg/L, respectively).

So, we concluded that immature fruits were best candidates for molluscicidal activity. This is in agreement with earlier studies carried out in our laboratories confirming the highest toxicity of crude water extracts of immature fruits of the same plant *Solanum nigrum* var. *villosum* (Hammami & Ayadi, 2008).

Immature fruits more rich in alkaloids were more toxic than mature fruits, this may be due to the high concentration of steroidal alkaloids. So, it was demonstrated
that small unripe fruits of *Solanum nigrum* had a high concentration of solasodine that decreased with fruit maturation (El Tayeb et al., 1997).

Phytochemical tests indicated the steroidal nature of the total alkaloids of *Solanum nigrum*. Several steroidal alkaloids including solasodine, solasonine and solamargine were isolated and identified in this plant (Schreiber, 1968; Ripperger & Schreiber, 1981; Drewes & Staden, 1995; El Tayeb et al., 1997). Steroidal saponins were previously isolated and identified as nigrumin I and II (Tsuyoshi et al., 2000). Six new steroidal saponins were also isolated by Zhou et al. (2006).

Hence, the observed molluscidal activity may be due to the presence of steroidal alkaloids. It was reported that steroidal alkaloids from *Solanum aculeastrum* possess strong molluscidal properties (Wanyonyi et al., 2002). In other studies, saponins and steroidal alkaloids were isolated from *Solanum melongena* L. fruits (Aubert et al., 1989).

CONCLUSION

In the present study, molluscidal activity of *Solanum nigrum* var. *villosum* varies between different parts of this plant and may be attributed to the nature of components. The analyses by thin layer chromatography achieved on the different extracts of this plant showed that the methanolic and hydro-methanolic ones were the richest in alkaloids and/or saponosides and the most active against *G. truncatula*. So, molluscidal activity may be correlated with the content of total alkaloid and/or saponosides found in the methanolic and the hydro-methanolic immature fruit extracts. This finding is in agreement with other data in the literature confirming that alkaloids from methanolic extracts of *Solanum aculeastrum* showed significant activity against host snails of schistosomiasis (LC₅₀ = 50 ppm) (Wanyonyi et al., 2003). However, molluscidal activity of *Solanum nigrum* var. *villosum* collected in Tunisia was more active and especially the hydro-methanolic extract (LC₅₀ = 3.96 mg/L) and the methanolic extract fraction richest in alkaloids (LC₅₀ = 1.65 mg/L) isolated from the immature fruits. Even the fractions richest in saponosides from the immature fruit hydromethanolic and methanolic extracts have shown interesting molluscidal activity (LC₅₀ = 6.15 mg/L and LC₅₀ = 7.91 mg/L, respectively). In our knowledge, this is the first study proving that immature fruit methanolic or hydromethanolic fractions rich in alkaloids and/or saponosides of *Solanum nigrum* var. *villosum* have important molluscidal activity against the snail gastropoda *G. truncatula*.

Green berries of *Solanum nigrum* var. *villosum* offers promised as a potential molluscidal control agent and the total alkaloids and saponosides from these plant merit further investigations to identify the active principal responsible of molluscidal properties and to further assess their activity in *G. truncatula* and its associated fauna in their natural habitats.

According to the World Health Organization’s guidelines on screening for plant molluscicides (WHO, 1983), use of these fractions may add to the arsenal of methods to control snail transmitting fasciolosis in tropical and third world countries where fasciolosis is a common disease.

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