Scolicidal effects of the *Nicotiana tabacum* L. extract at various concentrations and exposure times

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**Keywords:** Herbal medicine, Tobacco, Hydatid cyst, Scolicidal treatment, Immunotoxicity  

**ABSTRACT**  
**Background:** Tobacco is a plant that has anti-nematode and anti-inflammatory properties, due to that it contains compounds such as nicotine sulfate. It is utilized as a medicinal plant in the therapy of infectious diseases; also, it has been used to kill insects’ larvae and plant pests. **Objective:** The goal of this work was the evaluation of tobacco extract effects on the scolices of hydatid cysts. **Methods:** Hydatid cysts of the infected sheep livers were gathered from the Qazvin abattoir and their protoscolices were isolated under sterile conditions. Fertility and viability of protoscolices were confirmed before the experiments. The protoscolices were exposed to the ethanolic extract of *Nicotiana tabacum*, at different concentrations (3, 5, 10, 25, 50, and 100 mg/ml) and various exposure times (10, 20, 30, 40, 50 and 60 min). Sodium chloride 0.9% was selected as negative control. The viability of protoscolices was evaluated by staining via 0.1% eosin. **Results:** At 100 mg/ml, the tobacco extract caused the death of 91.19% of protoscolices after 10 minutes of contact, and other exposure times (20 to 60 minutes) resulted in the death of 100% of protoscolices. **Conclusion:** This medicinal plant showed a good scolicidal effect and it may be used for improving the procedure of hydatid cyst surgery. But, further studies are required.

1. Introduction

Human echinococcosis (also called hydatidosis and hydatid disease) results from the larval phases of cestodes (tapeworms) of the genus *Echinococcus*. Hydatidosis is one of the most crucial zoonotic illnesses and is prevalent in various regions of the globe. At present, surgery is the standard treatment for hydatid disorder [1-3]. One of the most notable risks of hydatid cyst surgery is the leakage of cyst contents and relapsing [4].

**Abbreviations:** ANOVA, Analysis of Variance; mg, Milligram; ml, Milliliter  
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Discovery of efficient and safe scolicidal candidates and administration of them into cysts can attenuate the dissemination risk of viable protoscolices. Different scolicidal candidates have been utilized for deactivation of the cyst ingredients, however, most produce adverse side effects [1, 5, 6]. Formalin, hydrogen peroxide, setrimide, pure alcohol, hypertonic saline, and silver nitrate have been utilized as protoscolicidal substances, however, their adverse effects have limited the applicability of these substances [5, 6]. Currently, the use of natural medications including *Allium sativum* L., *Zataria multiflora* Boiss., *Trachyspermum ammi* (L.) Sprague, *Zingiber officinale* Roscoe, *Coriandrum sativum* L. *Mentha pulegium* L., *Satureja khuzestanica* Jamzad, *Ziziphora Tenuior* L. extracts and *Olea europaea* L. have been considered as alternatives to the mentioned synthetic substances. Nonetheless, to ensure that they are harmless to the human, more research is needed [6-13].

Some herbals can provide immunomodulatory and anti-inflammatory effects and are therefore useful for treatment of a wide array of diseases [14-17]. Inflammation is the basis for many diseases, and reducing the inflammation while killing pathogens may provide an optimal approach [18, 19]. Tobacco extract can exert anti-worm, anti-inflammatory, anti-insect, and anti-fungal effects, mainly because of its chemical ingredients, such as resin, gum, potassium salts, various organic acids, and nicotine sulfate. In the present study, it was tested as a protoscolicidal substance [17, 20-25].

2. Materials and Methods

2.1. Preparation of Nicotiana tabacum L. plant extract and hydatid cyst samples

*Nicotiana tabacum* L. plant was purchased from a medicinal plants shop and its authenticity was confirmed by a botanist. A small sample of this plant was kept in the Institute of Medicinal Plants Herbarium with the herbarium code: 001 (IMPHM). Different parts of the dried plant including leaves, stems, and branches were crushed by a milling machine. In short, to prepare the plant extract, after drying and grinding them (half a kilogram), it was first extracted by ethanol and percolator. This process was repeated three times and then the resulting extracts were concentrated by vacuum distillation. The amount of dried extract was calculated; then, it was stored in a sealed plate in the refrigerator at 4 °C until the experiment.

The liver containing hydatid cysts was prepared from Qazvin abattoir, transferred to the parasitology department, and the contents of the cysts were suctioned in sterile settings and poured into sterile tubes separately. Calcified and purulent cysts were eliminated from this study. After disinfecting the surface of the cysts with betadine, the contents of the cysts were drained through 5 or 10 ml syringes and transferred into sterile test tubes. Cysts were examined for fertility by light microscopy. In the case of cyst fertility, the viability of protoscolices was evaluated through staining by 0.1 % eosin and movement assessment of candle flame cells. Cysts with a viability of more than 90 % were used for testing. [26, 27].

2.2. Evaluation of the protoscolicidal influence of plant extract

In this study, 6 concentrations of tested plant extracts (3, 5, 10, 25, 50 and 100 mg/ml) were exposed to protoscolices for 10, 20, 30, 40, 50 and 60 minutes. To prepare the mentioned concentrations, 30, 50, 100, 250, 500, and 1000 mg of plant extract were mixed with about 9 ml of normal saline in separate tubes and to facilitate the dissolution of the plant extract in normal saline, 0.3 ml of DMSO was transferred...
to each tube and the tubes were stirred with a magnet, and finally the volume of the mixture with normal saline reached 10 ml. 2.5 ml of the mixture was poured into each test tube, and 100 μl of the samples was added that contained about 1000 protoscolices. The ingredients of the tubes were incubated for 10, 20, 30, 40, 50 and 60 minutes at 37 °C. After the incubation period, we removed the superior part of the tubes.

1 ml of 0.1 % eosin was transferred into the precipitate obtained in each tube and mixed well. After 15 minutes of incubation, the surface was removed and the remaining sediment was laid on a slide and evaluated under a light microscope. By counting about 700 protoscolices, stained (dead) and unstained (live) protoscolices, percentage of dead protoscolices was determined. The control group (without plant extract) was used to ensure the accuracy of the experiments. The experiments were repeated three times [6, 12, 28, 29]. Examples of dead and live protoscolices are provided in Fig. 1 and 2, respectively.

Data were entered into the computer using Statistical Packages for Social Sciences (SPSS), (SPSS Inc., Chicago, USA) and the results were analyzed using one-way analysis of variance (ANOVA).

3. Results
Tobacco extract had little effect on protoscolices at all administered concentrations except for 100 mg/ml at all exposure times. At a concentration of 100 mg/ml, tobacco extract led to the death of 91.19 % of protoscolices following 10 minutes of contact. Moreover, longer exposure times (20 to 60 minutes) resulted in the elimination of 100 % of protoscolices (Table 1).
Table 1. Protoscolicidal effects of various concentrations of \textit{Nicotiana tabacum} L. extract in different exposure times

<table>
<thead>
<tr>
<th>Concentrations (mg/ml)</th>
<th>Experiments</th>
<th>10 (min)</th>
<th>20 (min)</th>
<th>30 (min)</th>
<th>40 (min)</th>
<th>50 (min)</th>
<th>60 (min)</th>
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<tbody>
<tr>
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<td>1</td>
<td>8.02 (56/698)</td>
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<td>8.98 (69/768)</td>
<td>11 (90/818)</td>
<td>10.47 (90/860)</td>
<td>11.84 (114/963)</td>
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<td>2</td>
<td>8.61 (72/836)</td>
<td>8.63 (66/765)</td>
<td>9.40 (82/872)</td>
<td>11.76 (108/918)</td>
<td>12.40 (122/984)</td>
<td>12.00 (90/750)</td>
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<td>3</td>
<td>5.49 (40/728)</td>
<td>5.73 (45/786)</td>
<td>7.86 (72/916)</td>
<td>9.44 (85/900)</td>
<td>11.00 (93/836)</td>
<td>10.53 (96/912)</td>
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<td>Total</td>
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<td>7.58 (185/2441)</td>
<td>8.72 (223/2556)</td>
<td>10.74 (283/2636)</td>
<td>11.34 (304/2680)</td>
<td>11.43 (300/2625)</td>
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<td>11.36 (108/951)</td>
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<td>9.47 (90/950)</td>
<td>9.76 (80/820)</td>
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<td>10.98 (84/765)</td>
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<td>10.86 (312/2873)</td>
<td>11.03 (258/2340)</td>
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<td>8.47 (63/744)</td>
<td>9.94 (80/805)</td>
<td>11.32 (96/848)</td>
<td>12.59 (90/715)</td>
<td>12.00 (96/800)</td>
<td>11.64 (88/756)</td>
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<td>11.11 (296/2665)</td>
<td>11.66 (295/2529)</td>
<td>13.68 (332/2427)</td>
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<td>20.16 (200/992)</td>
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<td>17.59 (152/864)</td>
<td>17.45 (156/894)</td>
<td>17.81 (176/988)</td>
<td>17.81 (171/960)</td>
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<td>18.29 (492/2690)</td>
<td>19.17 (527/2749)</td>
<td>19.20 (536/2792)</td>
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<td>21.01 (166/790)</td>
<td>24.94 (216/866)</td>
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<td>18.68 (130/696)</td>
<td>23.01 (168/730)</td>
<td>26.60 (225/846)</td>
<td>28.97 (288/994)</td>
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<td>17.44 (136/750)</td>
<td>18.40 (159/864)</td>
<td>24.82 (210/846)</td>
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<td>20.68 (493/2384)</td>
<td>25.45 (651/2558)</td>
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<td>100 (962/962)</td>
<td>100 (828/828)</td>
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<td>100 (735/735)</td>
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<td>92.93 (644/693)</td>
<td>100 (984/984)</td>
<td>100 (992/992)</td>
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<td>100 (950/950)</td>
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<td>89.94 (715/795)</td>
<td>100 (861/861)</td>
<td>100 (930/930)</td>
<td>100 (714/714)</td>
<td>100 (952/952)</td>
<td>100 (800/800)</td>
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<td>Total</td>
<td>91.19 (2123/2328)</td>
<td>100 (2807/2807)</td>
<td>100 (2750/2750)</td>
<td>100 (2616/2616)</td>
<td>100 (2669/2669)</td>
<td>100 (2732/2732)</td>
</tr>
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</table>

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DOIs: 10.5254/jmp21.82.111
Table 1. Protoscolicidal effects of various concentrations of *Nicotiana tabacum* L. extract in different exposure times (Continued)

<table>
<thead>
<tr>
<th>Concentrations</th>
<th>Experiments</th>
<th>10 (min)</th>
<th>20 (min)</th>
<th>30 (min)</th>
<th>40 (min)</th>
<th>50 (min)</th>
<th>60 (min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>(mg/ml)</td>
<td></td>
<td>7.10 (72/1014)</td>
<td>8.02 (56/698)</td>
<td>8.62 (60/696)</td>
<td>8.96 (90/1004)</td>
<td>9.33 (84/900)</td>
<td>10.47 (84/802)</td>
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<tr>
<td>Control</td>
<td>2</td>
<td>7.96 (78/980)</td>
<td>8.33 (70/840)</td>
<td>8.90 (78/876)</td>
<td>9.55 (102/1068)</td>
<td>10.28 (96/934)</td>
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<td>4.90 (40/816)</td>
<td>4.55 (36/792)</td>
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<td>5.91 (44/744)</td>
<td>6.28 (60/955)</td>
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<tr>
<td>Total</td>
<td></td>
<td>6.76 (190/2810)</td>
<td>6.95 (162/2330)</td>
<td>7.71 (184/2386)</td>
<td>8.38 (230/2816)</td>
<td>8.61 (240/2789)</td>
<td>9.47 (212/2238)</td>
</tr>
</tbody>
</table>

4. Discussion

Hydatidosis is widespread worldwide and causes the death of many patients. Surgery is currently considered the best method, but unfortunately, leakage of cyst contents is the most common cause of recurrence during surgery [2, 3, 5].

So far, several protoscolicidal chemicals have been proposed to solve this problem, but many of them have shown side effects that limit their use [1, 6, 30]. According to previous research, exposure with hypertonic salts, formalin, hydrogen peroxide, setrimide, pure alcohol, hypertonic saline, as well as silver nitrate can induce the death of protoscolices [5, 6, 31-34]. The presence of a substance with a rapid and complete protoscoliotoxic effect without any side effects and good cost-efficiency is regarded as part of the features of the best scolicidal solution; but, no perfect protoscoliotoxic solution has yet been explained. [35, 36]. Tobacco has been used as an herbal medicine, as an anti- worm, anti-inflammatory, anti-insect and anti- fungal [20-24].

In the present work, the efficiency of ethanolic extract of tobacco was explored on protoscolices. According to our findings, at a concentration of 100 mg/ml, 10 minutes of exposure resulted in the death of 91.19 % of protoscolices and longer exposure times (20 to 60 minutes) resulted in the death of 100 % of protoscolices. The findings of the current study showed that the ethanolic extract of tobacco comprises an efficient protoscolicidal candidate. Although according to this study and previous research, it seems that tobacco, due to its substances such as various organic acids and nicotine sulfate has protoscolicidal effects, it also affects the human immune system, nevertheless, the effects of tobacco scoliosis need further research [35]. We also suggest that the scolidal effect of different fractions of tobacco as well as different plant extracts like aqueous extracts, hydroglycolic extracts, glycerin extract, hydroalcoholic extract, be investigated to compare and find the most effective extract on protoscolices [37].

5. Conclusion

We showed a significant scolicidal capability for *Nicotiana tabacum* L. extract. This may help advance hydatic cyst surgery protocols and resolve shortcomings in the treatment of hydatosis, such as recurrence. Furthermore, we do not have enough information about the effect of this extract on different genomic variants of *Echinococcus granulosus*, which is an area that may need future experiments. In addition, it may
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be useful to evaluate the protoscolicidal actions of the alcoholic extract in vivo.

**Author contributions**

R.L., A.A., S.M., K.S., and R.H. drafted the manuscript. A.A. and R.H. conceptualized and supervised the study. R.L. and S.M. performed the experiments and analyzed the data. A.A. and K.S. analyzed the data. A.A. and K.S. critically appraised the manuscript.

**Conflict of interest**

The authors declare that there is no conflict of interest.

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Journal of Medicinal Plants  2022; 21(82): 111-118
چکیده
مقدمه: تنباکو گیاهی است که به دلیل داشتن ترکیباتی از قبیل تیکوکسین سولفات دارای خواص ضد تهیه و ضد الکل است. از این گیاه به علت لیمویی در دستان بیمارهای عفونی و همچنین برای کشتن لارو حشرات و آفات کیا به سادگی استفاده می‌شود. هدف از این مطالعه ارزیابی اثر عصاره گیاه تنباکو بر روی اسکولکسهای کیست هیداتیک بود. روش بررسی: کیست‌های هیداتیک از مولفان آنلود در کشت‌گاه فرآیند جمع آوری شد و پروتواسکولکس‌ها در شرایط استریل از این کیست‌ها جدا شدند. زنده بودن پروتواسکولکس‌ها با روش‌های آزمایشگاهی ثابت شد. پروتواسکولکس‌های زنده با غلظت‌های مختلف عصاره انتانولی گیاه تنباکو (0.5، 1، 5، 10 و 100 میلی‌گرم در میلی‌لیتر) در مدت زمان‌های مختلف (0، 10، 20، 30 و 60 دقیقه) مواجه شدند. از کلرید سدیم 10 درصد به عنوان کنترل منفی استفاده شد. میزان زنده بودن پروتواسکولکس‌ها پس از مواجهه با رنگ‌آمیزی با رنگ‌آبی از اسکولکس کاهش یافت. نتایج: عصاره تنباکو در غلظت 100 میلی‌گرم در میلی‌لیتر در 10 دقیقه باعث کشت شدن 91/19 درصد از پروتواسکولکس‌ها شد و در مدت زمان‌های مختلف کاهش میزان زنده بودن باعث کشته شدن 100٪ پروتواسکولکس‌ها شد. نتیجه‌گیری: این گیاه دارویی عاطفه‌ای که توانایی گزارش کمک‌کننده باشد، بهبرای برای مستحکم شدن هر چه بیشتر تازه به انجام تحقیقات بیشتر است.

اطلاعات مقاله
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کیست هیداتیک: درمان کیست هیداتیک
ایمتوکسیست‌ها: ANOVA
مخفف‌ها: ANOVA: آنالیز واریانس; mg: میلی‌گرم; ml: میلی‌لیتر
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