Scolicidal activity of taurolidine for the treatment of hydatid disease

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Abstract: Objectives: In this experimental study, we have evaluated in vivo and in vitro activities of taurolidine (TRD) against protoscolices of Echinococcus granulosus. Background: Scolicidal agent application is mandatory for the interventional treatment of hydatid cysts. Serious adverse effects of current scolicidal agents forced the scientist to search for less toxic chemicals. Methods: One milliliter of protoscolices suspension was administered into twelve Petri’s dishes, six of them containing 5 ml of isotonic saline, and six of them containing 5 ml (5 mg/ml) TRD. Forty, male Balb/c mice were divided into five equal groups. Group 1: protoscolices inoculated group; Group 2: protoscolices inoculated and thereafter immediately administered with a single dose intravenous (IV) 0.5 ml TRD (400 mg/kg); Group 3: protoscolices inoculated and thereafter immediately administered with a single dose intraperitoneal (IP) 0.5 ml TRD (400 mg/kg); Group 4: three months after inoculating the protoscolices, one day before the sacrifice single dose IV 0.5 ml TRD (400 mg/kg) administered; Group 5: three months after inoculating the protoscolices, one day before the sacrifice single dose IP 0.5 ml TRD (400 mg/kg) administered. Results: All the protoscolices were dead in TRD-added Petri’s Dishes in 90 minutes. In group 2 and group 3, no mouse had an intraabdominal hydatid cyst. Conclusion: The results of this study are encouraging us to suggest TRD as an alternative scolicidal agent. Further clinical studies are needed to define the effectiveness and the mode of application of TRD for the treatment of hydatid disease (Tab. 1, Fig. 4, Ref. 30). Full Text in PDF www.elis.sk.

Key words: taurolidine, hydatid disease, scolicidal agent, Echinococcus granulosus.

Hydatid disease (HD) is a zoonosis caused by larvae of Echinococcus granulosus which form cysts in different organs of human and herbivores. Despite its benign nature, Echinococcus granulosus causes serious financial and manpower loss and even death in case of complications (1). Medical treatment, percutaneous drainage and surgical treatment are current treatment options of intra abdominal HD (2, 3). Use of scolicidal agents is a mandatory step in the interventional treatment of hydatid cysts. By killing scolices, scolicidal agents assure the disinfection of the cyst and prevent contamination of the surgical field. Hypertonic sodium chloride, silver nitrate, povidone iodine, cetrimide are frequently used scolicidal agents. However, these scolicidal agents could cause acute or chronic toxicity (4, 5). An ideal scolicidal agent should be highly effective in low concentrations and in short time, should not have local or systemic side-effects, should not harm liver or bile ducts, should have low viscosity, should be prepared and conserved easily and be inexpensive (4-6). But there is no agent that fulfills all of these specifications.

Taurolidine (TRD), bis-(1,1-dioxo-perhydro-1,2,4-thiadiazinyl-4)-methan, consists of two taurinamide rings derived from the naturally occurring amino sulfonic acid, taurine and three mole-
cules of formaldehyde combining to form a two-ringed structure bridged by a methylene group. Intravenous (IV) five-day TRD treatment showed beneficial effects in septic patients without causing any side effects (7). TRD has also been used intraperitoneally for the treatment of peritonitis (8). In this experimental study, we have evaluated the in vivo and in vitro activities of TRD against protoscolices of Echinococcus granulosus.

Materials

This study was performed in the Experimental Research Laboratory of Istanbul University Cerrahpasa Medical Faculty after approval of the Istanbul University Experimental Medicine Research Institute Ethical Committee (Protocol No: 47/2008).

Preparation of protoscolices

The live protoscolices were harvested intraoperatively from liver hydatid cysts (Echinococcus granulosus) of patients by suction of the virgin cyst fluid. Infected or bile-stained aspirates were discarded. Submerged protoscolices were collected, washed with isotonic saline and decanted (not centrifuged) for three times to get a purified scolices suspension. Thus, a suspension of 1,000–1,200 scolices/ml was obtained. Then, the viability of scolices was evaluated based on their morphology, motility and vital dye (eosin) uptake.

In vitro study

One milliliter of this suspension was put in twelve Petri’s
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Dishes, six of them containing 5 ml of isotonic saline, and six of them containing 5 ml (5 mg/ml) TRD. One drop of 0.1 % eosin was added to each Petri’s dish to confirm or disprove the death of protoscolices. Ten different fields of each Petri’s dish were examined under 50x and 100x magnifications with a light microscope. The viabilities were calculated as percentage by counting live protoscolices in the microscope area. Protoscolices that conserved their ovoid shape, did not lose their ameboid movements and did not stain with eosin were assumed alive. The ones that were immotile, stained with eosin and whose rostella were retracted, thus becoming spherical were considered dead (Fig. 1–4).

**In vivo study**

After showing scolicidal activity of TRD in vitro, we designed an in vivo experiment. Forty young (eight-week old), male, Balb/c mice of 24–30 g weight, were divided into five equal groups (n=8 for per group) (Tab. 1). The mice were looked after in accordance with the Guide for the Care and Use of Laboratory Animals (NIH Publication no. 86–23, revised 1985), maintained in colony cages under controlled light, temperature and humidity conditions and permitted *ad libitum* access to standard lab chow and tap water during the experimental procedures. Protoscolices were inoculated intraperitoneally under aseptic conditions. Group 1: protoscolices inoculated group; Group 2: protoscolices inoculated and immediately thereafter administered with a single dose IV 0.5 ml TRD (400 mg/kg); Group 3: protoscolices inoculated and immediately thereafter administered with a single dose IP 0.5 ml TRD (400 mg/kg); Group 4: three months after inoculating the protoscolices, one day before the sacrifice a single dose IV 0.5 ml TRD (400 mg/kg) administered; Group 5: three months after inoculating the protoscolices, one day before the sacrifice a single dose IP 0.5 ml TRD (400 mg/kg) administered. The mice were sacrificed 90 days after the protoscolices inoculation. Intraperitoneal hydatid cyst development and viability of cyst content were evaluated.

**Statistical analysis**

Kruskal–Wallis and post hoc Bonferoni adjusted Mann–Whitney tests were used for the statistical analysis, while p<0.01 was considered significant.

**Results**

In our preliminary evaluation, we had observed that a single dose IV, IP and intrabiliary TRD (400 mg/kg) administration had caused no toxic effects upon the peritoneum, hepatocytes and cholangiocytes of the Balb/c mice.

**In vitro**, the movement of protoscolices stopped and the protoscolices got circular shape seven minutes after TRD application. Forty-five minutes after TRD administration, 50 % of the protoscolices and ninety minutes after the administration all protoscolices were dead in TRD-added Petri’s dishes (Fig. 3). However, at the end of the same period, 95 % of protoscolicesesiceses were alive in saline group.

When we evaluate the in vivo study, no mouse died during the experiment. In group 1, seven mice (7/8) had intraabdominal hy-
datid cysts which were located on the liver and intestines. These cysts contained live protoscolices. The mice in group 2 (0/8) and group 3 (0/8) had no intraabdominal hydatid cysts. These values for group 2 and group 3 are significantly lower than for group 1 (p<0.001). Seven mice had intraabdominal hydatid cysts in each of groups 4 (7/8) and 5 (7/8) but none of the hydatid cysts contained alive protoscolices.

**Discussion**

HD is currently an important public health problem especially in the developing countries including Turkey. Many clinical and experimental studies have been published about the treatment of the disease; nevertheless none of them has defined an ideal scolicidal agent (9). Protoscolices cause recurrent hydatid cysts in the liver. Inoculation of protoscolices into the peritoneal cavity of a young mouse is an experimental model of hydatidosis (10, 11).

Serious adverse effects of current scolicidal agents forced the physicians to search for less toxic chemicals to be used for this purpose. Formalin is one of the former drugs but it has serious systemic toxic effects and causes epithelial proliferation in biliary tract so it is not currently used in clinical practice (12). Ethyl alcohol at 98 % concentration is used for percutaneous treatment of hepatic hydatid cyst. Its effectiveness depends on concentration of alcohol while sclerosing cholangitis could occur after ethyl alcohol administration (13). Additionally, ethyl alcohol is not preferred during the surgical treatment of the hydatid cysts since it is flammable and volatile. Hypertonic sodium chloride and silver nitrate are used as scolicidal drug but they could cause acute pancreatitis and sclerosing cholangitis (4, 14). Hypernatremia is a foreseeable complication of hypertonic sodium treatment (15). Cetrimide-chlorhexidine combination has an adhesive action on small bowel (16). Hypertonic sodium chloride and alcohol inactive and degenerate the germinative membrane. Alcohol also obliterates residual cavity beside its scolicidal activity (15). Scolicidal effect of povindone-iodine which is used mostly in our country depends on dosage, it can also protect secondary echinococcosis (4, 17). Allergic reactions could be seen after application of povidine-iodine. Considering these adverse effects of the current scolicidal agents, the discovery of less toxic and much effective chemicals are urgently needed.

The scolicidal effects of TRD are under investigation. Ekci et al (18) suggested that TRD has no scolicidal activity. However, the design of the methodology was not compatible with the clinical scenario in this pilot study. Our *in vitro* application time for TRD was longer and we administered TRD intraabdominally after inoculating the protoscolices into the abdomen. They had inoculated TRD-treated protoscolices into the abdominal cavity and did not evaluate the exact application time (18). In the current experimental study, *in vitro* and *in vivo* scolicidal activities of TRD have been shown. TRD has a therapeutic effect against many infections causing peritonitis when administered at the doses that we used in the current study. Additionally, no serious adverse effects were reported after administration of TRD infusion in human studies (19). TRD is presented as an antibacterial and antineoplastic agent in literature. Active metabolites of TRD, taurinamide, interact with the structures of bacterial wall (20–23). Antioxidant effects of TRD could attenuate the immune response against the parasitic infections (24). TRD has beneficial effects on wound healing process too (25, 26). TRD could support physiologic mechanisms by reducing the harmful effects of the pathology. Parasitic infections could be harmful via producing endotoxins. TRD also neutralizes bacterial endotoxins, exotoxins, and lipopolysaccharides (27–29). TRD has anti-adherent and anti-inflammatory activities (7–8). One or more of these characteristics could be responsible for the scolicidal activity of TRD.

Formaldehyde is one of the degradation products of TRD. Carcinogenetic and cytotoxic effects of formaldehyde are known. However, the drug (Taurolin®, Boehringer Ingelheim, Germany) that we used in the study includes a stabilizer, polyvinylpyrrolidone, for reducing formaldehyde concentration (30). No cancer development and no carcinogenetic pathways were reported to be related to TRD treatment.

The results of this study are encouraging us to suggest TRD as an alternative scolicidal agent. Reaching the maximum scolicidal activity could take a longer time for TRD but it could be less toxic than other scolicidal agents. Further clinical studies are needed to define the effectiveness and the mode of application of TRD for the treatment of hydatid disease.

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