Topical ivermectin in the treatment of pediculosis capitis

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ABSTRACT

Background: Head lice infestation is considered a prominent issue because of its worldwide distribution, especially among children. Millions of children are infested with pediculosis capitis every year. The increasing rate of lice infestation has been reported even in the developed countries due to the resistance to known pediculicides. Besides, the louse is a vector for serious diseases, such as epidemic typhus fever. Objective: The aim of this work was the search for a new cheap and effective drug for pediculosis capitis. 2% ivermectin solution, a promising novel drug used for endoparasite and ectoparasite infections, was used in this study. Materials and Methods: This was a clinical trial study. The in vitro study was carried out in a period of five months. Samples of nits and lice were collected from female patients of the Department of Dermatology and Venereology, Baghdad Teaching Hospital. The study was conducted during a period from October 2002 through November 2003. One hundred fifty children from four primary schools in Hayy Al-A'amel, Baghdad, participated in this part of the study. The preparation was applied to the children by the author. One hundred children were tested with ivermectin solution and fifty children with rectified spirit. The first part of the study involved a parasitological evaluation (in vitro study) of pediculicidal and ovicidal activity, in which 4% ivermectin solution was discontinued because of contact dermatitis. Therefore, 2% ivermectin solution was used instead. Pediculicidal and ovicidal activity was examined half an hour afterward. The second part of the study involved a clinical evaluation (in situ study) of 2% ivermectin solution for half an hour. Results: Topical 2% ivermectin solution in rectified spirit is a good pediculicidal (100%) and a good ovicidal (88%) drug with good therapeutic efficacy (82% after the first application, 90% after the second application) against pediculosis capitis. Conclusion: The study described a novel topical preparation for pediculosis capitis, which was proven effective and safe.

Key words: Ivermectin; Pediculosis capitis; Iraq; Topical solution

INTRODUCTION

Population-based studies in European countries show a highly divergent prevalence of pediculosis capitis, ranging from 1% to 20% [1]. With an increasing rate of treatment failure, it is worthwhile to consider the issues of misdiagnosis, lack of adherence, inadequate treatment, reinestation, lack of ovicidal or residual killing properties of the pediculicide, and/or resistance of lice to the pediculicide [2]. Resistance to topical pediculicides is an emerging concern in most parts of the world [3]. The problem of resistance is directly related to the frequency of its use [4,5] in genotyping, demonstrating that head lice exhibit resistance to compounds such as DDT, the pyrethrins, and the pyrethroids [6]. Numerous treatment options have been used to control the spread of pediculosis capitis with various drawbacks. For instance, 1% lindane exhibits central nervous system toxicity and may cause severe seizures in children [7]. 5% benzyl alcohol was the first non-neurotoxic pediculicide, and pruritus, erythema, pyoderma, and ocular irritation are its usual side effects. It is also non-ovicidal [8]. The percentage of patients who were louse-free after treatment with a product containing tea tree oil and lavender oil and a head lice suffocation product was higher than with a product containing pyrethrins and piperonyl butoxide [9]. Eucalyptus oils have been used against permethrin-resistant pediculosis [10].

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Grapefruit juice [11], coconut, and anise spray have been found to be more effective than permethrin in pediculosis capitis, thereby demonstrating clinical resistance [12].

In head lice infestations that are difficult to treat, oral ivermectin may prove more effective [13] or as effective as topical 0.5% malathion lotion [14]. Oral ivermectin demonstrates high efficacy and tolerability in the treatment of pediculosis capitis. A significant number of children required a second dose to ensure complete eradication [15]. A single oral dose of ivermectin 400 μg/kg repeated throughout seven days has been shown to be more effective than 0.5% malathion lotion [13]. It has also been used successfully in school children [16]. In head lice infestations that are difficult to treat, oral ivermectin given twice every seven days had superior efficacy when compared with topical 0.5% malathion lotion [17]. After a single dose, complete healing was achieved in 77.5% and 87.5% of ivermectin and malathion groups, respectively. No major adverse effects were observed in either group. Oral ivermectin is a promising and effective approach for the treatment of head lice and might be an ideal substitute for conventional pediculicides [14]. Community-based treatment with oral ivermectin in susceptible, and poor sections of the society has given us promising results [18].

MATERIALS AND METHODS

In vitro parasitological evaluation: The activity of the test preparation was evaluated on the Pediculus capitis in ovicidal and pediculicidal tests. Samples of nits and lice were collected from female patients of the Department of Dermatology and Venereology, Baghdad Teaching Hospital. The study was conducted during a period from September 2002 through November 2003. 2% ivermectin solution 1 gm/50 cc rectified spirit was used and tested by parasitological evaluation (in vitro study). A control test was included as part of the parasitological evaluation as well. Rectified spirit was the control preparation. Ovicidal tests of each type of test and control preparation were tested half an hour after washing in tap water. A hundred fully mobile lice were transferred to another clean Petri dish. A hundred fully mobile lice were exposed to test preparations by immersion at room temperature. The number of live and dead lice was recorded. After three minutes, the lice were washed in tap water, then microscopically examined. A louse was considered dead if it lacked signs of internal and external movement. Fifty fully mobile lice were exposed to rectified spirit half an hour after washing, then microscopically examined. The percentage of dead lice represented the pediculicidal activity of the test preparation.

Ovicidal Test

Hair shafts with attached viable ova (nits) were removed with forceps and scissors, transferred onto a sterile Petri dish, closed with self-adhesive tape, and identified. An ovum was judged viable if it was plump, had an intact operculum and ideal eyespot, and was yellow or creamy white. Dark tan to brown or black ova with shrunken or shrunken shells were judged nonviable. An ovum that had the operculum popped open and a translucent shell without a brown residue was considered hatched. Nits were examined with a binocular microscope. Empty (hatched) or possibly dead (nonviable) nits were excluded from the ovicidal test, whereas the plump (viable) nits were transferred to another clean Petri dish. The viable ova, firmly fixed to their support (hairs), were exposed to the test preparation by immersion for half an hour at room temperature. Afterward, the nits were washed three times in tap water. Nits were allowed to dry at room temperature and incubated again in similar conditions at room temperature for two weeks for hatching. Next, unhatched nits were counted under the microscope. The percentage of unhatched nits represented the ovicidal activity of the test preparation. Then, the same was done but with the control preparation after the completion of the parasitological evaluation.

In Situ Study

Ivermectin solution was tested for its therapeutic efficacy on primary school children infested with pediculosis capitis. One hundred fifty children from four primary schools in Hayy Al-A’amel, Baghdad, participated in this part of the study.

The children were in grades one to six. The preparation was applied to the children by the author. One hundred children were tested with ivermectin solution and fifty children with rectified spirit.
To be eligible for participation, each patient had to have a diagnostically active head lice infestation confirmed by direct visual identification of live adult lice or nymphs. The presence of nits on the hair shafts was insufficient to qualify a child for inclusion in the study. A child was excluded if there were any other dermatological conditions present or if a pediculicide was used within two weeks of the initial evaluation.

The parents agreed not to wash the hair of their children for half an hour of solution application and not to use pediculicides, medicated shampoos, or lotions, other than ordinary shampoo, during the study. During the first visit, a visual estimation of live adult louse and nymph populations in the children’s hair was conducted and the test preparation was applied to the dry scalp in amounts sufficient to thoroughly wet the hair and skin of the infested areas. An amount of 25–50 mL was sufficient to wet the hair and scalp, but extremely long and thick hair sometimes required larger amounts. During the application of the test preparation, the children were instructed to protect their eyes from the preparation by holding a folded towel against their forehead. The hair was covered with a cup for half an hour, then washed.

The therapeutic index (i.e., eradication of live lice and nymphs) was evaluated on days seven and fourteen following the treatment. The children were examined for a minimum of five minutes for the presence of one or more live adult lice or nymphs on either evaluation. The first application of the preparation was given at the first visit, and the second application was given at the second visit (after one week) with the same procedure as on the first application. The parents of the children were instructed to boil or steam sheets, pillowcases, and other formats for about 15–20 minutes.

A clinical evaluation was performed 30–60 minutes following day fourteen. The children were examined for three dermal signs (edema, erythema, and rash) and five dermal symptoms (pruritus, burning, stinging, numbness, and tingling).

**Ethics Statement**

All authors hereby declare that the study has been approved by the scientific committee of the Scientific Council of Dermatology and have, therefore, been performed in accordance with the ethical standards defined by the Iraqi Board of Dermatology and Venereology.

**RESULTS**

A total of 150 lice were included in the study. A hundred were tested with ivermectin and fifty with rectified spirit.

The following pediculicidal activity was observed:

1. **2% ivermectin solution**
   After exposure for three minutes, all of the 100 ova were dead. The pediculicidal activity of the preparation was 100%.

2. **Rectified spirit**
   After half an hour, 7 ova out of the 50 were dead. The pediculicidal activity was 14%.

**Ovicidal Test Results**

A total of 150 eggs were included in the study.

1. Out of the 100 eggs, 88 eggs were dead. The ovicidal activity of the ivermectin solution was 88% after half an hour.
2. Out of the 50 eggs, 6 eggs were dead. The ovicidal activity of the rectified spirit was 12% after half an hour.

The lice were found to be able to survive for 2–4 days at 30°C without food (away from the human host).

**In Situ Study Results**

Hundred fifty primary school children infested with active pediculosis capitis were included in the study. One hundred of them were treated with 2% ivermectin solution at the first and second visit and fifty with a control preparation (rectified spirit). All children were female, and their age ranged from 6 to 12 years with a mean of 9 years.

Eighty-two children had no live lice at the first examination (one week after the first application of the test preparation). The therapeutic efficacy of the test preparation after one application was 82%, and 90% after two applications.

While out of the 50 children infested with pediculosis capitis treated with rectified spirit applied for half an hour before washing, 47 children had live lice (adults or nymphs) at the first examination (one week after the first application), and 43 children had live lice (adults or nymphs) at the second examination (one week after the second application). Therefore, the therapeutic efficacy
of the control preparation (rectified spirit) was 6% after the first application and 14% after the second application.

A burning sensation was reported in twelve children (12%) 15–30 minutes after the application of the test solution (2% ivermectin solution). No other dermal symptoms or signs were observed after the application of either preparation.

DISCUSSION

Ivermectin was chosen because of its role in the treatment of ectoparasitic infestations and its relative lack of toxicity as compared with other modalities by recent studies and, because of the increasing number of treatment failures, possibly due to the development of resistance, a change in treatment is to be considered.

Ivermectin is a new drug for the treatment of ectoparasites. To determine its effect on pediculosis capitis, in this study, we applied it as a topical 2% solution in rectified spirit and found that the in vitro pediculicidal activity of the ivermectin solution after three minutes was exceptionally good (100%) and that its ovicidal activity was good as well (88%).

After the first application of 2% ivermectin solution for half an hour, the therapeutic efficacy (percentage of dead lice and ova) was very good (82%) and, after the second application, it was still very good (90%).

A burning sensation was reported in twelve children 30 minutes after the application. These results support that topical ivermectin solutions produce a low frequency of side effects, whereas, with the use of lindane, toxicity has been reported [7].

The result of the study indicates that a 2% ivermectin solution is a good pediculicidal and ovicidal drug that appears to be a suitable alternative, especially considering the reported worldwide spread of resistance to both pyrethroids and malathion.

Therefore, 2% ivermectin solution is a good and cheap pediculicide with a low frequency of side effects. Its main advantages are low adverse reactions and rapid insecticide action, which requires a short contact time and makes it suitable for home treatment.

The study suggests that topical ivermectin may be a promising treatment for head lice and that a second dose on day seven might be appropriate.

2% ivermectin solution was found to have good ovicidal activity (88%) and, if the time of exposure is increased, the ovicidal activity may increase; therefore, a second application is of extreme importance.

In the study, the lice were able to survive for 2–4 days without food (away from the human host).

Causes of treatment failure: Active lice, adults and nymphs, in parasitological evaluation, exposed to the test preparation under optimal conditions with complete exposure of all lice to the preparation in a Petri dish, while the application of the preparation on the hair of the children may not be in close contact with all the lice. Some nits may need more than one week to hatch. Therefore, the nits will hatch on the second application, and this may lead to failure in treatment and reinfection since all failed cases in our study had live lice. Most of them were small (nymphs) but several were of a mature size (adults), suggesting that these children had been reinfested.

CONCLUSION

This is the first Iraqi study on the use of topical ivermectin for pediculosis capitis.

Topical 2% ivermectin solution in rectified spirit is a good pediculicidal (100%) and a good ovicidal (88%) drug for the in vitro treatment of pediculosis capitis. However, a second application seems to be necessary because of its incomplete ovicidal activity.

It is deemed safe and has no or few side effects.

It is a good pediculicide drug, with 82% therapeutic efficacy on the first application and 90% therapeutic efficacy on the second application, and a good choice for a primary treatment of pediculosis capitis and resistant cases.

At the time of the study, there had been few similar studies, which shows that there is a need for more clarification on Iraqi patients because of the geographical diversity of resistant cases.

Recommendations

1. Further studies regarding ivermectin solutions for longer periods of exposure.
2. Further comparative studies of ivermectin solutions with other pediculicides.
3. There is an urgent need for monitoring the development of resistance through official control of sales and prescriptions.
4. New products such as topical ivermectin are required and, once introduced, careful control of their use would be of benefit.
5. Control of head lice can be attempted by head shaving and wet combing.
6. We recommend the use of a fine-toothed louse comb in addition to treatment for its role in:
   a. The prevention of lice infestation;
   b. Anti-louse treatment methods as an accessory tool;
   c. The removal of nits.
Mass family treatment is necessary.

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Statement of Human and Animal Rights

All the procedures followed were in accordance with the ethical standards of the responsible committee on human experimentation (institutional and national) and with the 2008 revision of the Declaration of Helsinki of 1975.

Statement of Informed Consent

As per the national standard, written consent has been collected from the patient or a parent of the patient and preserved by the author(s).

REFERENCES