**Research Paper: Investigating the Efficacy of Sumac Topical Solution Against Permethrin-resistant Human Head Lice**

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**ABSTRACT**

**Background:** The present study aimed at determining the efficacy of applying Rhus coriaria (Sumac) solution for the treatment of Permethrin-resistant head louse in patients, who used permethrin for at least 2 consecutive periods, but have not been cured. 

**Methods:** This study is a before-after clinical trial performed on 100 patients with pediculosis aged between 2 and 50 years old and both sexes. All patients had used Permethrin at least twice consecutively (with at least 14 days interval) according to correct instructions (on the first and 7th day), but they have not been cured. Each patient received 60ml of Rhus coriaria solution for 3 consecutive days, and the treatment was repeated again for another 3 days; then, the patients were followed-up on the 4th, 10th, and 14th days after the treatment.

**Results:** The results showed a significant difference in the severity of head lice infection and itching before the treatment and 14 days after the treatment (P<0.001).

**Conclusion:** Rhus coriaria solution was more effective in eliminating head-louse infestations on 4, 10, and 14 days after the treatment and itching disappeared in most of the patients, while negligible complications were observed.

**Keywords:** Pediculus, Permethrin, Insecticide-resistance, Rhus coriaria, Sumac

**Article info:**
Received: 13 Jan 2019
First Revision: 10 Feb 2019
Accepted: 13 May 2019
Published: 01 Jul 2019
practical effectiveness in 1999 [1]; even the prescription of more drug levels was also not effective [2].

Resistance is an advanced mechanism that causes the survival of the insect against a deadly attack. This successful mutation can be transmitted over generations via DNA [2]. These resistant organisms can withstand against stimulants and destroyers agents.

Several mechanisms are responsible for permethrin resistance. Permethrin and pyrethroids affect voltage-sensitive sodium channel [6]. If these channels get insensitive to these drugs, the treatment will not be effective. Knockdown resistance (Kdr) is one of the most common mechanisms of drug resistance [7].

From 1984 to 1995, the permethrin efficiency was about (96%) to (100%). After it was shifted to Over The Counter (OTC), based on the reports, its efficacy reduced to (80%) and, then, gradually reduced to (55%) and (28%), in conditions, where treatments have been completed by nit combing [7]. In the United States, from 1996 to 1999, the permethrin cure rate was only (79%) [8].

Therefore, resistance has been identified as “an upward challenge” that is very terrible, making patients use potentially-harmful treatments like using more powerful or very toxic pesticides, which can be very dangerous for these patients or even cause death because of their side effects. In other words, these hazardous substances and drugs, including Dichloro Diphenyl Trichloroethane (DDT), Oil, Gasoline, and Detergents that have very severe complications, sometimes cause the death of the patients.

In addition, this dilemma will increase costs, drug complications, and the disappointment of treatment, which will lead patients to depression. Studies have shown that because of their low toxicity and fewer complications, the application of plants or herbs can be considered very good alternative pediculicides to be replaced with anti-llice drugs for the elimination of drug resistance. Therefore, there is a growing interest in using herbs in the treatment of head lice.

The results of a study on the effectiveness of Eucalyptus species on the treatment of permethrin-resistant Pediculus humanus capitis showed that Eucalyptus sideroxylon was more effective than all species [9].

The anti-pediculosis effect of 25 essential oils extracted from different plant species from Argentina, was investigated in the treatment of permethrin-resistant head lice. Cinnamomum porphyrium had the most efficacy [10].

In traditional medicine, 120 simple drugs have been named for the topical treatment of head lice, for example, Tamarix gallica oil, as well as 150 compound drugs, for example, the combination of Acorus calamus aromaticus, Cinnamomum Verum, and Carthamus tinctorius oil [11, 12]. It has been referred to Sumac (Rhus coriaria) and Olive oil in a book written by Ibn-e-Sina (Al-qanun), Al-Havi, a book written by Razi, and health drugs guidebook written by Ahmed Ali Khosravi [11-13].

Rhus coriaria or Sumac belongs to the Anacardiaceae herbaceous family [14]. It is found in different places of the world, including North Africa and various parts of Iran and Afghanistan [15]. It has been used in traditional medicine and cookery as a spice. In folk medicinal, it has been used for treating diarrhea [14]. Rhus coriaria consists of compounds such as tannins, phenolic acids, flavonoids, terpenoids, and essential oil composed of monoterpenes that have numerous biological activities, which are used as hair cleaning solutions or anti-dandruff agents for the improvement of dermatological problems [16, 17]. It has antimicrobial, antiviral, antifungal, antioxidant, anti-inflammatory, and anticancer activities [18, 19]; it is also used orally for reducing blood glucose, uric acid, and cholesterol levels [20-22].

Olive oil contains very high amounts of fatty acids and terpenoid species [23]. Even it is used locally for the treatment of head lice; however, in this study, the prepared solution has only low amounts of olive oil, and its main ingredient is Rhus coriaria (Sumac).

The oral and therapeutic uses of these ingredients can be attributed to this fact that they cause no complication [24]. No clinical trial has examined the efficacy of Rhus coriaria solution on pediculosis. This study was conducted as a clinical trial and it evaluated the efficacy and safety of Sumac solution on patients with mild to severe head lice.

2. Materials and Methods

The study was conducted in the form of the before-after clinical trial that was performed on 100 patients with pediculosis aged between the range of 2-50 years old and on both sexes. All patients had used permethrin at least for twice consecutively (with at least 14 days interval), according to correct instructions (on the first and seventh day) and they were not treated. The 2-week study was conducted.
Inclusion and exclusion criteria

The inclusion criteria included avoiding depriving patients of routine treatment, patients who had used permethrin treatment twice and more consecutively (with at least 14 days interval) according to the correct instructions (on the first and 7th day) and did not respond to treatment, and yet they had at least grade 1 infection (5-9 nit or live nymphs/lice); they were selected as subjects of the study according to the severity of the contamination [25]. Although the second line of treatment is based on Lindane therapy, the resistance to Lindane is also very high [2, 8]. Additionally, it causes neurotoxicity and seizure, especially at the younger age; therefore, it has been obsoleted in other countries and its use is not advised [2, 8].

The exclusion criteria included: 1. Any sensitivity to the contents of the solution; 2. Itching and skin ulcers on the head; 3. Secondary skin infection including yellow impetigo and chronic skin diseases such as psoriasis; 4. History of diseases related to immune deficiency; 5. pregnancy; 6. Participation in any clinical trials within the past 4 weeks or previous participation in this clinical trial [25, 26]. Obviously, the patients who did not want to continue the cooperation or did not use the medicine in accordance with its correct instructions (the treatment for less than twice every 3 sequential days), or the patients, who used another anti-lice drug simultaneously with the treatment were excluded from the study.

In order to prevent re-infection, other family members, who had head lice, but were not eligible for intervention were also treated with permethrin.

Drug preparation

Sumac clusters were bought from valid pharmaceuticals market) Buran of Kurdistan) and extra virgin olive oil was bought from Exir company.

The cover of the seeds was separated and placed in boiling water for 4 hours in the lab environment. Then, they passed the finer filters and were poured inside the crystallizer and it was placed on a water bath or set) Bain-Marie machine (for 24 hours. Finally, the dry extract obtained was diluted again with water. Besides, Soy lecithin, as an emulsifier, was gradually added to a very small amount of olive oil and mixed in the clockwise direction. Then, Sumac solution was gradually added to the mixture of Soy lecithin and olive oil. During the addition of the solution, it was rotated slowly clockwise to achieve a monotonic emulsion. The obtained emulsion contained (84.9%) water and (15.1%) oils) Olive oil and Soy lecithin). The solution was supplied as a spray to the patients.

Sample size

G * power V. 3.1.9.4 (reference number X) software was used to analyze the data. The first type error of 0.05, (95%) power of the test and the effect size of 0.18 for the statistical test of 2 domains for the sign test of 100 people were calculated for this study according to the following calculations [27].

Data collection method

Each patient filled out a questionnaire consisting of demographic information such as age, height, weight, BMI, level of education, number of baths per week, number of times for taking shower per a day, number of households, hair color, hair type, and head pruritus [28]. The severity of the complication and the degree of itching were determined and recorded 4 times (day zero and 4th, 10th, and 14th days after treatment). Any complications regarding the use of sumac solution and patient dissatisfaction were also questioned.

The diagnosis was done based on the direct observation of lice or nit. The head was divided into 5 regions by an applicator including the temporal regions and the areas behind the ears and neck [29]; each area was checked by magnifier for infection with lice and nit. Nits locating less than 1.4 inch of the scalp were considered clinically valuable and the rate of infection was graded according to the following division:

1. Zero (0-4 nit); 2. Grade 1 or mild (5-9 nit or live nymphs/lice); 3. Grade 2 or medium (10 to 25 nit or live nymphs/lice); 4. Grade 3 or severe (more than 25 nit or live nymphs/lice) [25, 30]. All examinations were performed 4 times; once before the treatment (day zero) and 3 times after the start of treatment (days 4, 10, 14). In order to eliminate the diagnostic errors and incorrect evaluation, all examinations were done by only one person (experienced physicians). Clinical symptoms such as pruritus or redness, scalp inflammation, and cervical lymphadenopathy were also evaluated.

Procedure

The study was conducted in the form of the before-after clinical trial. In the intervention group, each patient received 60ml of solution filled in spray plastic bottles for 3 continuous days, as well as 20ml daily, by a tiny
plastic brush teeth. The patients were asked to massage the area after applying the drug each time for 10 minutes on the head and, then, to wrap a towel around their hair and wait for 90 minutes. Then they were asked to comb by a fine brush then washing the hair with water and shampoo. Again, 60ml of the drug was given to patients to use in 3 continuous days on the 7th, 8th, and 9th days. Patients only used the drug that they were given. They were followed-up on the 4th, 10th, and 14th days of intervention. Patients were examined for lice, nit, and amount of itching or any complications. The information regarding the method of using the drug and the health instructions were given to the patients both in the written and verbal form.

Statistical analysis

All analyses of baseline and treatment effects were performed, using SPSS V. 22 software and inferential statistics tests (based on the scale of variables, McNemar, Wilcoxon, and the sign test).

3. Results

At the beginning of the study, the distribution of severity of head lice infection among 100 patients, who entered the intervention, was as follows: 1. (27%) of the patients had grade 1 infection; 2. (41%) of the patients had grade 2 infection; 3. (32%) of them had grade 3 infection. On the 10th day (when the administration of the second dose of the sumac solution was finished), the rate of infection for the patients (27 patients), who had grade 1 infection, was zero. Among 41 patients, who had grade 2 infection, 37 patients reached grade 0, and 2 patients reached grade 1, and the status of 2 patients did not change. Among 32 patients, who had grade 3 infection, 19 patients were fully treated, 5 patients reached grade 2, and 8 patients reached grade 1. The results of the Wilcoxon Signed Ranks test showed a significant difference in the severity of the infection before the treatment and 10 days after treatment (P<0.001). Table 1 presents the results of the comparison of the infection rate before the treatment and on the 10th day.

Finally, on the 14th day of the treatment, among 100 patients, (88%) of the patients reached grade zero (totally cured), (7%) of them reached grade 1, and (5%) of them reached grade 2. As a result, (88%) of the patients were fully treated and the rate of infection reduced to (12%), and (5%) of them reached grade 2 and (7%) of them reached grade 1.

The results of the Wilcoxon Signed Ranks test showed a significant difference in the severity of the infection before the treatment and 14 days after treatment (P<0.001). Table 2 presents the results of the comparison of the infection rate before the treatment and on the 14th day.

Table 1. Comparison of the grade of infection before the treatment and on the 10th day

<table>
<thead>
<tr>
<th>Grade</th>
<th>The Grade of Infection After Treatment (10th day)</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Zero (1-4)</td>
<td>One (5-9)</td>
</tr>
<tr>
<td>The grade of infection before treatment (first day)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>One (5-9)</td>
<td>27</td>
<td>0</td>
</tr>
<tr>
<td>Two (10-24)</td>
<td>37</td>
<td>2</td>
</tr>
<tr>
<td>Three (&gt;25)</td>
<td>19</td>
<td>8</td>
</tr>
<tr>
<td>Total</td>
<td>83</td>
<td>10</td>
</tr>
</tbody>
</table>

Table 2. Comparison of the grade of infection before the treatment and on the 14th day

<table>
<thead>
<tr>
<th>Grade</th>
<th>The Grade of Infection After Treatment (14th day)</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Zero (1-4)</td>
<td>One (5-9)</td>
</tr>
<tr>
<td>The grade of infection before treatment (first day)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>One (5-9)</td>
<td>27</td>
<td>0</td>
</tr>
<tr>
<td>Two (10-24)</td>
<td>38</td>
<td>3</td>
</tr>
<tr>
<td>Three (&gt;25)</td>
<td>23</td>
<td>4</td>
</tr>
<tr>
<td>Total</td>
<td>88</td>
<td>7</td>
</tr>
</tbody>
</table>
Surprisingly, itching reduced in the patients after the 4th day. Itching completely eliminated in (33%) of the patients; (27%) of them had only slight itching, (10%) of them had moderate itching, and (3%) of the patients had still high or very high itching. Table 3 shows the rate of itching before treatment and on the 4th day.

On the 10th day of the treatment, (86%) of the patients had no itching at all, (11%) of the patients had a little itching, and (1%) of the patients had medium, high, or very high itching. Finally, on the 14th day of the treatment, (89%) of the patients had no itching, (8%) of them had low pruritus, and (1%) of them had moderate itching. Only (2%) of the patients had high and very high pruritus. Table 4 presents the rate of itching before the treatment and on the 14th day.

<table>
<thead>
<tr>
<th>Rate</th>
<th>Before the treatment (the first day)</th>
<th>After the Treatment (The 4th day)</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0</td>
<td>Low</td>
<td>Medium</td>
</tr>
<tr>
<td>0</td>
<td>17</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Low</td>
<td>12</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>medium</td>
<td>8</td>
<td>18</td>
<td>1</td>
</tr>
<tr>
<td>high</td>
<td>4</td>
<td>7</td>
<td>5</td>
</tr>
<tr>
<td>Very high</td>
<td>9</td>
<td>2</td>
<td>4</td>
</tr>
<tr>
<td>Total</td>
<td>60</td>
<td>27</td>
<td>10</td>
</tr>
</tbody>
</table>

Table 3. Comparison of the rate of itching before the treatment and on the 4th day

<table>
<thead>
<tr>
<th>Itching Rate</th>
<th>Before the treatment (the first day)</th>
<th>After the Treatment (The 4th day)</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0</td>
<td>Low</td>
<td>Medium</td>
</tr>
<tr>
<td>0</td>
<td>17</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Low</td>
<td>12</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>medium</td>
<td>34</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>high</td>
<td>12</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td>Very high</td>
<td>14</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>Total</td>
<td>89</td>
<td>8</td>
<td>1</td>
</tr>
</tbody>
</table>

Table 4. Comparison of the rate of itching before the treatment and on the 14th day

4. Discussion

There was a significant difference in the severity of pediculosis before and after the treatment, and the rate of itching of the head became none or significantly less.

Seemingly, increasing the dose of the solution or the duration of using the solution would yield better results. The patients’ satisfaction of treatment was very high and significant, while the side effects of the solution were negligible and included only burning and pruritus in the first 1 to 2 minutes after the use of the drug in some patients. In the patient’s follow-up, a large number of subjects did not have pediculosis for several months, while in the case of using other drugs, they could not get rid of pediculosis at all, or they were quickly re-infected.

This indicates the strength of the drug in terms of its function, including the important effect of topical cleansing of the scalp and making the environment inappro-
appropriate for the continuation of the life of the insect. This event could result from the presence of tannin in sumac. Sumac contains a large number of tannins that have a tannery effect and cleansing effect that make the environment of scalp unpleasant for the life of the insect; it can remove the thick and sticky material underneath the scalp and clear the crust of substances, which accumulate in the pores of the skin and hair follicles. Therefore, most of the patients did not suffer from pediculosis for several months later.

Many studies have investigated the administration of herbal medicines for the treatment of head lice, but the use of sumac and olive oil, both of which had a long history in nutrition, seems to reduce the concerns about drug toxicity or complications, especially for long and frequent uses.

All patients, who entered the intervention, had used permethrin for at least 2 continuous periods and they did not positively respond to treatment. Most patients coexisted for a long time with lice and they did not get rid of the insect and nit at all and they had used other therapies including Lindane and Dimethicone lotion (4%) (Dilice), DDT, Gasoline, Oil, Mayonnaise, and Detergent. Today, pediculosis is widely spread and it has passed the economic, cultural, and health border; as it is widely seen in all societies, finding a suitable treatment is very necessary.

The limitations of the study include the lack of awareness of the patients, especially the students and their families, about the importance of treatment, follow-up, and the lack of cooperation between school and health care providers.

5. Conclusion

The results of this study demonstrate that Sumac solution can be a safe and effective solution for the treatment of head lice and can prevent drug resistance, cost increases, and confusion and frustration among the patients.

Ethical Considerations

Compliance with ethical guidelines

Patients who were consent to participate in the intervention filled out the form of ethics consent, and the parents of patients aged under 18 years old also filled out the consent form. The study protocol was approved by the Ethics Committee of Shahid Beheshti University of Medical Sciences (IR.SBMU.RETECH.REC.1397.125) and was registered in the Iranian Registry of Clinical Trials (IRCT20180712040446N1).

Funding

This article is based on PhD thesis Number 207 School of Traditional Medicine, Shahid Beheshti University of Medical Sciences, Tehran, Iran.

Author’s contributions

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Conflict of interest

The authors declared no conflict of interest.

Acknowledgments

The authors would like to acknowledge Shahid Beheshti center for research cooperation and financial support.

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