RESEARCH ARTICLE

Comparison of phenothrin mousse, phenothrin lotion, and wet-combing for treatment of head louse infestation in the UK: a pragmatic randomised, controlled, assessor blind trial [version 1; referees: 2 approved]

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Abstract

In this investigation of effectiveness of an alternative pediculicide dosage form, we recruited 228 children and 50 adult participants from Bedfordshire, UK, to a randomised, controlled, assessor blind trial comparing two insecticide products with mechanical removal of lice as a control group. Participants using insecticide were treated with either the investigative 0.5% phenothrin mousse, for 30 minutes, or 0.2% phenothrin lotion, for 2 hours as the reference product. Both treatments were applied only once, followed by shampoo washing. Those treated by wet-combing with conditioner were combed 4 times over 12 days. Parents/carers carried out the treatments to mimic normal consumer use. The outcome measure was the absence of lice, 14 days after treatment for the insecticides, and up to 14 days after completion of combing. Intention to treat analysis of the outcomes for 275 participants showed success for phenothrin mousse in 21/105 (20.0%), in 23/107 (21.5%) for phenothrin lotion, and in 12/63 (19.1%) for wet-combing. People receiving mousse were 1.07 (95% CI, 0.63 to 1.81) times more likely to still have lice after treatment compared with those treated with lotion. The group of participants who received the wet combing treatment were 1.13 (95% CI, 0.61 to 2.11) times more likely to still have lice after the treatment. None of the treatments was significantly (p < 0.05) more effective than any other. This study was carried out in an area where moderate resistance to phenothrin was demonstrated after the study by using a bioassay. Analysis of post treatment assessments found that failure of insecticides to kill louse eggs had influenced the outcome.

This article is included in the All trials matter collection.
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Competing interests: IFB is a consultant to several large, medium, and small companies that develop and market head louse treatment products, including medicinal products, medical devices, cosmeceuticals, and combs. CMB and PN are not aware of any competing interests.

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Introduction
During the early 1990s, a number of synthetic pyrethroid-based formulations for treating head louse infestation were introduced into the British market. The majority of these used \textit{d-}phenothrin as the active substance in a variety of dosage forms\(^1\). In 1997 a phenothrin mousse was developed based on a concept developed in Australia using natural pyrethrum\(^2\). The aim of the new product was designed to be more manageable during application and thus gain greater consumer acceptability than existing preparations. Laboratory studies of phenothrin lotion had indicated a high level of activity for the insecticide, so theoretically it could be incorporated into a formulation requiring a shorter application time\(^3\). However, at this time there was increasing evidence of insecticide resistance in several areas of the UK\(^4\), in parallel with a renewed consumer interest for treating head louse infestation by combing, either as the principal measure or as a component of conventional insecticide treatment.

At the time of the study the most widely promoted combing method was “Bug-Busting” (Community Hygiene Concern, London), which used a fine toothed plastic comb for wet combing with conditioner, repeated at 3–4 day intervals for 2 weeks. It suggested the first combing could remove all lice so only newly hatched nymphs would be found during subsequent combing sessions before they could mature and lay eggs\(^5\). Before our investigation only two studies of wet-combing had been conducted. In one Bug-Busting was half as effective as two applications of malathion lotion\(^6\). A second found it more effective than permethrin creme rinse, but the dropout rate from both treatments made the interpretation of the results difficult\(^7\).

We performed a pragmatic, observer blinded, three armed clinical trial analysed by intention to treat, comparing single applications of 0.5\% \textit{d-}phenothrin mousse or 0.2\% \textit{d-}phenothrin lotion with the Bug-Busting protocol of wet-combing with conditioner. The study was designed to evaluate the effectiveness of each treatment when in use by the public.

Methods

Participants
We recruited participants (children and adults) to the study from respondents to an information letter distributed through schools or via general practitioners associated with Bedfordshire Health. Prospective participants or their parents/caregivers telephoned the study co-ordinator to make an appointment for a home visit by a trained agency nurse. Most visits were within 24 hours unless requested at a different time. Nurses followed a standard approach to check for living lice using a plastic detection comb (Albyn of Stonehaven Ltd, Stonehaven, Scotland). If moving head lice were found, and the individual was 4 years or over, they were invited to join the study. Prospective participants were conducted through a standard consent procedure in which the content of information sheet was explained verbally and the parent/carer confirmed that they understood the function, processes, and commitments of the study before signing the consent form, which was witnessed by an independent adult. Participants were individually assigned a randomised treatment. All other household members were offered examination and, if found to be infested with head lice, were given the opportunity to join the study.

Prior to inclusion in the study, all participants provided baseline data including: age, gender, ethnicity, hair characteristics including length, thickness, degree of curl, and previous pediculicide use. Inclusion criteria required availability for up to 28 days to accommodate each of the possible treatment regimens and a suitable adult available to perform or assist in application of the treatment. Candidates excluded from this study were: pregnant or nursing mothers; anyone who had bleached, colour-treated or permanently waved their hair; used pediculicide, had been treated with antibiotics, or had participated in a clinical trial during the 4 weeks prior to this trial. In addition anyone with sensitivity to any pyrethroid insecticide or chrysanthemums; receiving treatment for asthma; suffering from a persistent skin disorder of the scalp (other than head lice); or had already participated in this study; was also excluded.

Household members with lice who either did not wish to participate or who failed to satisfy the inclusion criteria were given advice about appropriate treatment methods. Any participants found with lice after completion of the study period were supplied with 0.5\% malathion lotion (Suleo-M lotion, Seton Healthcare Group plc, Oldham, UK) in conformity with the Bedfordshire Health policy for treatment. No payment was offered for participation.

Ethics
Ethical approval was granted by both the South Bedfordshire and North Bedfordshire Local Research Ethics Committees of Bedfordshire Health, one of the conditions of which was that the data would be published in the public domain. The study was registered with the Current Controlled Trials database (ISRCTN73201839). See Supplementary files for the study protocol.

The study was conducted in compliance with Good Clinical Practices, and in conformity with the principles of the Declaration of Helsinki and European Standard, EN540: Clinical investigation of medical devices for human subjects. Written and witnessed informed consent was obtained from the participants and parents or guardians of children under 18 years of age.

Treatments
The nurses explained how to apply the treatment to the parent or carer and also gave a printed copy of these instructions. The parent then applied the treatment in the presence of the investigator. The nurses were instructed to answer any questions, and to note these, but not to intervene if any error was observed during treatment. This was intended to represent the clinical situation.

We supplied one group of participants with 0.5\% \textit{d-}phenothrin mousse in 50ml butane pressurised containers, with a camula fitting to allow direction of the mousse during delivery (Full Marks Mousse, Seton Healthcare Group plc, Oldham, UK). This also contained citrate buffered water, ethanol, and emulsifying wax. We supplied the second group with 0.2\% \textit{d-}phenothrin water/isopropanol lotion in 50ml glass bottles with a dropper aperture (Full Marks Lotion, Seton Healthcare Group plc). Carers applied the products to dry hair to saturate the hair and scalp. We made available as many containers of product necessary to comply with the instructions. People using mousse washed it from the hair with shampoo after...
30 minutes. Those treated with the lotion left it on the hair for 2 hours before shampooing. We supplied both groups with a non-medicated shampoo (L’Oréal Children’s Shampoo, L’Oréal (UK) Ltd, London, UK) for this. Both groups received a single application of treatment.

We supplied the third group of participants with a “Bug Buster” pack (Community Hygiene Concern, London, UK) for performing the wet-combing technique. We also supplied a bottle of the same non-medicated toiletry shampoo and four 60ml bottles of conditioner rinse (one for each treatment day) (L’Oréal Children’s Shampoo and L’Oreal Conditioner, L’Oréal (UK) Ltd), and a diary card. Parents/carers were instructed to wash the participant’s hair with the shampoo, massage in a generous amount of conditioner, and comb through the hair systematically from scalp to tip with the louse removal comb provided in the pack. They wiped the comb on a paper towel between strokes. Combing was repeated at regular intervals for 2 weeks (days 0, 4, 8, 12 of the study). We asked carers to fix any lice found during the combing onto the diary card using clear cellulose adhesive tape and to record how much time was spent combing.

Outcome measures
We made mousse and lotion treatment follow ups on days 4, 7, 10, and 14 and those for wet-combing on days 14, 21, and 28 after commencement of treatment. Day 14 was used as the point for measure of primary outcome for the insecticide groups. However, in the wet-combing group it was possible there could be viable eggs present on day 14 and therefore follow up examinations were also conducted on days 21 and 28 to detect any emerging nymphs or other lice missed at day 14. During each of the follow up assessment examinations, any lice found by detection combing were removed and fixed to the case record using clear cellulose adhesive tape. These were later examined in the laboratory to determine the gender or development stage of each insect.

Nurses collected the containers of mousse and lotion after treatment so that the quantities used could be measured. Bottles of conditioning rinse from the wet combing group were also collected for measurement after the final assessment on day 28, although some of these were mislaid by participants.

Sample size
We estimated sample sizes to show a difference between wet combing and phenothrin lotion treatment with 95% confidence, 90% power, and equivalence between the two phenothrin groups to within 20%. For this calculation we conservatively estimated that the phenothrin products would exhibit approximately 80% effectiveness and wet combing 50% success. Sample size calculations were made by the sponsor’s consultant statistician who estimated a minimum sample size of 104 participants in each phenothrin treated group and 58 participants treated with wet combing (266 evaluable participants) would satisfy this probability with greater than 90% power.

Randomisation and blinding
A computer generated list, prepared by the sponsor’s statistician, was used for randomisation of treatments, made up of balanced blocks of 133 treatment allocations with a relative frequency for each of the treatments of 52:52:29. Randomisation was by individual so that different members of a household could receive different treatments. Nurses involved in recruitment were supplied with envelopes in batches of ten and asked to issue them sequentially. Investigators, who were unaware of which treatment had been used, made follow up examinations using plastic detection combs to check for the presence of living lice. On day 4, 7, and 10 these assessors were a different group of agency nurses. On days 14, 21, and 28 the assessing investigators were from the Medical Entomology Centre (IFB and CMB).

Statistical analysis
We analysed for differences between groups based on the intention to treat (ITT) population and tested equivalence only using the per-protocol (PP) population. We calculated differences in cure rate using a chi-squared test and equivalence to within 20% based on the 95% confidence limits derived from the normal approximation to the binomial distribution. The initial analyses for the sponsor were performed by the consultant statistician (PN Lee Statistics and Computing Ltd, Sutton, UK) using bespoke software. Post hoc analyses performed by the investigators employed Epi-Info version 6, OXSTAT II version 1.11, and purpose built spreadsheet calculators. Differences between groups in baseline characteristics, safety, acceptability, and efficacy were tested using Fisher’s exact test for yes/no variables and the Mann-Whitney U test for ranked variables.

Tests for resistance
During the course of the study a high level of treatment failure was observed in the insecticide groups. We collected samples of lice at the final assessment from five participants from different parts of the study area for a bioassay evaluation for sensitivity to phenothrin. The insects were placed on treated or control filter papers using a method previously described for tests of permethrin sensitivity. Each treated filter disc was impregnated with 500 µl of 2% d-phenothrin solution, giving an insecticide deposition rate of 157 µg cm⁻². The mortality outcomes of the tests were compared, with a baseline sensitivity obtained using laboratory-reared, insecticide sensitive, body/clothing lice by means of log-probit analysis using LDP Line software.

Results
Participants
The study was conducted between June 1997 and March 1998, during which informed consent was obtained for 228 children and 50 adults to participate (Figure 1). Two people were excluded from further analysis as, upon inspection, no live lice had been found. The recruitment case record form for one other participant was lost so this case was also excluded from the study.

The 275 participants were randomly assigned to one of the treatment groups: 105 received phenothrin mousse, 107 phenothrin lotion, and 63 were allocated to the wet-combing treatment. Of these participants, 246 (89.5%) (100 mousse, 100 lotion, and 46 wet-combing) completed the trial with adequately complete follow-up data sets (Figure 1 and Dataset 1). From the original study group, 5 participants (2 treated with lotion and 3 wet-combing) were non-compliant and excluded from the per-protocol analyses. Non-compliance
involved additional combing or other unauthorised treatments (2 receiving lotion and 2 wet-combing) and one participant on wet-combing shaved his head. There were 24 other withdrawals: 2 people on wet-combing dropped out; 20 were lost to follow-up (5 from the mousse group, 5 from lotion, and 10 from wet-combing); and 2 people from the wet-combing group were not allocated Day 21 or Day 28 appointments in error following a communication failure between the study coordinators and the investigators conducting final assessments. Two of those lost to follow up were due to bereavement and the two drop outs chose not to continue in the study. The rate of protocol violation/withdrawal was significantly higher (p < 0.001) in the wet-combing group than the phenothrin-treated groups.

Of the 275 people known to satisfy the inclusion and exclusion criteria, 62 (23%) were male (Table 1). The percentage distribution of males was similar for the two phenothrin groups, 18% and 21% respectively, but higher in the wet-combing group (33%). This difference in proportion of males between wet-combing group and mousse-treated group was statistically significant (p < 0.05) and the difference between wet-combing group and the lotion-treated group was nearly significant (0.05 < p < 0.1).
Outcomes

Post-treatment examinations at day 14 showed that there were 20/107 successful treatments and 3 cases of reinfestation after cure (an overall success rate of 21.5%) using phenothrin lotion and 18/105 successes and 3 cases of reinfestation for phenothrin mousse (giving 20.0% overall success). This made mousse users 1.07 times more likely to have lice after completion of the treatment (95% confidence interval (CI) 0.63 to 1.81; odds ratio (OR) 1.10, 95% CI 0.56 to 2.13). In the case of wet-combing with conditioner there were 12/63 (19.1%) successful treatments and no cases of reinfestation. Participants treated with combing were, therefore, 1.13 times more likely to have lice (95% CI 0.61 to 2.11; OR 1.16, 95% CI 0.53 to 2.54) than if they had been treated with phenothrin lotion. People treated with wet-combing were also 1.05 times more likely to have lice than those participants receiving phenothrin mousse (95% CI 0.56 to 1.99; OR 1.06, 95% CI 0.48 to 2.34).

In both insecticide treated groups the majority of lice at post-treatment assessments were juveniles, of which 712 first were stage nymphs that could only have originated from eggs not killed by insecticide. However, it was not possible to properly analyse the full effect of oviducal failure due to participants being withdrawn early in the study on grounds of lack of efficacy. Nevertheless from the data available it was possible to determine that failure to kill louse eggs was a major contributing factor in the low rate of outcome success.

Intention to treat analysis found no statistical difference (p < 0.05) between the lotion and mousse. Similarly, no statistical difference was found between wet-combing and either of the insecticides. Success in curing the infestation was also not significantly associated with gender or hair type, thickness, or length, after adjustment for any randomisation anomalies. However, success rates were significantly (p < 0.01) higher in people who had previously used a head louse treatment successfully, by an estimated factor of 1.88 (95% CI 1.13 to 3.11). Success also significantly (p < 0.01) increased with age.

Analysis based on the per-protocol population, without taking into account the failure of randomisation, showed no significant difference in success rates between the two phenothrin groups (lotion 17.8%, mousse 13.7%), which showed equivalence to within 20% (mousse-lotion difference, -4.1%, 95% CI -6.1% to 14.4%).

Table 1. Demographic characteristics of the intention to treat population.

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Mousse</th>
<th>Lotion</th>
<th>Wet-comb</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of participants</td>
<td>105</td>
<td>107</td>
<td>63</td>
<td>275</td>
</tr>
<tr>
<td>Age</td>
<td>4–7</td>
<td>45 (42.9%)</td>
<td>42 (39.3%)</td>
<td>27 (42.9%)</td>
</tr>
<tr>
<td></td>
<td>8–12</td>
<td>36 (34.3%)</td>
<td>40 (37.4%)</td>
<td>24 (38.1%)</td>
</tr>
<tr>
<td></td>
<td>13–16</td>
<td>5 (4.8%)</td>
<td>6 (5.6%)</td>
<td>0 (0.0%)</td>
</tr>
<tr>
<td></td>
<td>&gt;17</td>
<td>19 (18.1%)</td>
<td>19 (17.8%)</td>
<td>12 (19.0%)</td>
</tr>
<tr>
<td>Median</td>
<td>9</td>
<td>9</td>
<td>8</td>
<td>9</td>
</tr>
<tr>
<td>Sex</td>
<td>Male</td>
<td>19 (18.1%) **</td>
<td>21 (19.6%) *</td>
<td>21 (33.3%) **</td>
</tr>
<tr>
<td>Previous treatment experience</td>
<td>85 (81.0%) **</td>
<td>98 (91.6%) **</td>
<td>49 (77.8%) **</td>
<td>232 (84.4%)</td>
</tr>
<tr>
<td>Previous treatment successful</td>
<td>23 (27.1%) §</td>
<td>46 (46.9%) §</td>
<td>14 (28.6%)</td>
<td>83 (35.8%)</td>
</tr>
<tr>
<td>Hair characteristics</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Length</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Above ears</td>
<td>18 (17.1%)</td>
<td>26 (24.3%)</td>
<td>23 (36.5%)</td>
<td>67 (24.4%)</td>
</tr>
<tr>
<td>Below shoulders</td>
<td>56 (53.3%) **</td>
<td>50 (46.7%)</td>
<td>24 (38.1%) **</td>
<td>130 (47.3%)</td>
</tr>
<tr>
<td>Thickness</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fine</td>
<td>39 (37.1%) **</td>
<td>43 (40.2%)</td>
<td>34 (54.0%) **</td>
<td>116 (42.2%)</td>
</tr>
<tr>
<td>Thick</td>
<td>66 (62.9%)</td>
<td>64 (59.8%)</td>
<td>29 (46.0%)</td>
<td>159 (57.8%)</td>
</tr>
<tr>
<td>Curl</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Straight</td>
<td>71 (67.6%)</td>
<td>73 (68.2%)</td>
<td>48 (76.2%)</td>
<td>192 (69.8%)</td>
</tr>
<tr>
<td>Wavy/curly</td>
<td>34 (32.4%)</td>
<td>34 (31.8%)</td>
<td>15 (23.8%)</td>
<td>83 (30.2%)</td>
</tr>
</tbody>
</table>

Levels of statistical variation between groups: Figures in bold type show the group exhibiting a statistical disparity indicating possible randomisation anomalies.
* Difference significant at p < 0.1; ** Difference significant at p < 0.05; § Difference significant at p < 0.01

Adverse events

We found several clearly defined treatment-related adverse events in people treated using the phenothrin products: 12 adverse events in 11 people using lotion (9 scalp irritation, 3 irritation of the respiratory system); 10 adverse events in six people treated with mousse (5 scalp irritation, 3 dry skin, 1 bullous reaction, 1 paesthesia of the scalp). There were no similarly defined adverse events for the wet-combing group but five carers reported children expressing signs of stress while being combed, one person reported discomfort.
during combing, and backache or arm/shoulder aches for three carers were also reported, but not formalised as reported adverse events. All events were considered mild and resolved rapidly except for two cases. The case of paraesthesia, which was classified as moderate, persisted for two days after treatment, and one case of dry skin persisted for some time after treatment but may have been an exacerbation of a pre-existing problem. Stinging of the hands and paraesthesia-like reactions were also reported by some of the carers while applying the phenothrin-based products. Paraesthesia has been reported from use of other pyrethroid preparations and would likely be exacerbated by the presence of alcohol in the product.

Tests for resistance
Lice from different participants showed marked differences when tested for sensitivity to phenothrin (Table 2). In all cases the lice were taken from people who had experienced treatment failure during the study so it was not surprising that the majority had insects that were resistant to the insecticide. However, one person had apparently also been reinfested with sensitive lice from a contact, as shown by the mixed sensitivity of the insects. All lice from another participant, treated using wet-combing, were phenothrin susceptible. Output data from the LDP Line analyses of the observations of head lice, in comparison with susceptible laboratory reared body lice, showed a resistance ratio (RR) of 54.74 when exposed to the insecticide. From the log-probit analyses the estimated time required to kill 50% of the insects (LT$_{50}$) was 502 minutes based on a mortality curve with a slope of 1.0096 ± 0.1324 (chi-squared 19.9681, p = 0.0005). In contrast the estimated LT$_{50}$ for the body lice was 24.74 minutes (slope = 5.1932 ± 0.5086, chi-squared = 11.6217, p = 0.0404). Log-probit analysis also suggested that the insecticide sensitive head lice (LT$_{50}$, 44 minutes; LT$_{95}$, 95 minutes) were approximately twice as tolerant of phenothrin as laboratory reared lice, although the number of insects involved was too small to provide a clear distinction.

Table 2. Sensitivity of head lice from study participants to d-phenothrin in vitro.

<table>
<thead>
<tr>
<th>Participant</th>
<th>Treatment</th>
<th>Number of lice</th>
<th>Time for insecticide effect</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>In study</td>
<td>In vitro</td>
<td>Total</td>
</tr>
<tr>
<td>18</td>
<td>Mousse</td>
<td>Phenothrin</td>
<td>7</td>
</tr>
<tr>
<td>181</td>
<td>Lotion</td>
<td>Phenothrin</td>
<td>6</td>
</tr>
<tr>
<td>271</td>
<td>Lotion</td>
<td>Phenothrin</td>
<td>11</td>
</tr>
<tr>
<td>273</td>
<td>Mousse</td>
<td>Phenothrin</td>
<td>37</td>
</tr>
<tr>
<td>278</td>
<td>Wet-comb</td>
<td>Phenothrin</td>
<td>24</td>
</tr>
<tr>
<td>Body lice</td>
<td></td>
<td>Phenothrin</td>
<td>60</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Control</td>
<td>23</td>
</tr>
</tbody>
</table>

* The time of death of these lice was sufficiently delayed that it may have been due to dehydration.
** Lice survived for longer than 540 minutes when provided with a blood meal.
Discussion

This was a pragmatic study designed to investigate the three treatments under conditions that mimicked normal use by parents and carers. The outcomes demonstrated that all three products were less than adequate to eliminate head louse infestation when used according to instructions.

In the case of the phenothrin mousse, a major limitation of the product in use was that the foam was too dry and insufficiently thermolabile. It did not break down to a fluid that could be readily spread through the hair. This problem probably arose because too little alcohol was included in the formulation so the foam generated from water and emulsifying wax did not break physically after application. Consequently, many of the parent carers probably failed to achieve an adequate or even coverage of the hair and scalp when applying it. In contrast, phenothrin lotion was too fluid, like other alcohol based products. This meant that in use it was easy to apply too little product because a small volume of the fluid made the hair look wet and, by implication, thoroughly coated. Also, this formulation did not contain any of the terpenes that had been shown to contribute much of the ovicidal activity shown by other alcoholic lotions. At that time the “Bug Buster” pack contained a two-part comb that was initially described as “unique, safe and well researched” and, “reliable even though its use may be time consuming”. Although when the product was shown to be relatively ineffective by independent investigators they were criticised by the pack suppliers for using a product that had been superseded by the time of publication. Our investigators found it relatively easy to find lice on heads that parents using the “Bug Buster” comb believed were louse free. Therefore, it can be concluded that the two-part comb (Figure 2) was not as effective as originally claimed and was probably not as easy to use as either the plastic detection comb we used or its replacement, which was similar to our detection comb.

For this study the sponsor engaged a general nursing agency to supply staff to perform the majority of study functions in place of trained investigators as a cost saving measure. We believe this was not only a false economy in terms of data quality but may also have resulted in a breakdown of Good Clinical Practice. We could not determine whether some of the nurses failed to understand their responsibilities as investigators, or simply could not follow instructions, but some of the apparent failures of randomisation could be attributed to poor practice. For example, people in the wet-comb group were significantly more likely to have short hair (p < 0.05) or fine hair (p < 0.05) than those in the mousse-treated group. These anomalies were attributed to a failure of randomisation at the point of enrolment by some nurses engaged in recruitment. Early in the study we drew this possibility the attention of the sponsor, as well as to the management of the nursing agency, who assured us that correct procedures were being followed. Nevertheless, a failure of randomisation was identified at analysis because we suspected a general reluctance on the part of carers and children to participate in the combing group. Boys with shorter hair were apparently seen as an easier option for wet-combing so it is possible the numbered envelopes were opened before they were specifically allocated to individuals in households were several members were participating. Therefore we adjusted for this anomaly using stratified chi-squared analyses, although ultimately it made little difference to the overall outcome analysis.

We detected another, less easily identified, anomaly in that people who had a previous experience of a successful treatment with an insecticide lotion were more likely to be recruited to the lotion group (lotion v mousse p < 0.01; lotion v wet combing p < 0.05) (Table 1), although there was no evidence this was due to a failure of randomisation at the point of allocation.

Another deviation arose because most participants that were found to have lice at post-treatment assessments on day 4 or later were not withdrawn and provided with rescue treatment by the agency nurses but allowed to remain in the study, in some cases, until assessed by us on day 14. As far as we were able to determine, this did not affect the outcomes, and was partly associated with the logistical difficulties of transporting large volumes of documentation between offices over a short period of time, but again suggests a lack of understanding of the requirements of the study.

In the study area the local policy had been to use malathion for head louse infestation during the previous few years, so we did not anticipate significant resistance to pyrethroids. However, the small sample of lice we collected during the later stages of the study demonstrated that resistance was present and probably existed over much of the study area. The level of treatment failure could not be fully explained by resistance because five years later we observed a 75% success rate using a phenothrin aqueous emulsion in a study area that overlapped with this geographically. Rather the problem may have rested with the preparations used because another study using the alcoholic phenothrin lotion, conducted in a different part of the country, obtained a similarly poor result with only 2/15 (13.3%) cures.

This study was conducted because the Medicines Control Agency (MCA), now the Medicines and Healthcare Products Regulatory Agency, did not consider clinical investigations of the mousse in India representative of conditions likely to be encountered in Britain, so the manufacturers were required to conduct a UK-based study for confirmation of efficacy. Surprisingly however, before we could complete the analysis of this study, the MCA issued a Marketing Authorization to the mousse for use in England and Wales.
Authorisation for the phenothrin mousse. As a result the product was launched without further assessment by the MCA. Not surprisingly, given the poor effectiveness observed in this clinical investigation, there was widespread anecdotal reporting of treatment failure by consumers after using the product. Nevertheless it remained in the UK market until 2009.

There are lessons that can be learned from this experience. The first is that bioassay tests conducted in a laboratory, whether using laboratory reared insects or even wild collected ones, can only be indicative of efficacy for a formulation and it is unknown for a treatment to perform poorly in vitro yet be effective in vivo. Of course, laboratory reared lice and other ectoparasites are usually poorly representative of the physiological characteristics of those found on their natural hosts, especially with regard to characteristics such as resistance. Even ex vivo screens, using insects recently collected from the wild, may be only partially representative and several replicate tests should be performed using insects from graphically separated locations to ensure that the outcome is not obtained either by chance or due to some happenstance of physiological difference in the insects from that location. Attempting to draw any kind of conclusion about efficacy from only two or three lice or a single replicate test is fraught with risk, although this appears to have been a common practice in some investigations.

The second lesson, one that was recognised by the Medicines Control Agency when they initially insisted that the phenothrin mousse should be clinically tested in the UK, is that a clinical investigation of a pediculicide in a country where treatments for head lice are not routinely used is not likely to be representative of the possible outcomes in the territory where the product is destined to be marketed. Such studies may be indicative of possible outcomes but basing strong claims about how a product will work in a developed country, where lice are regularly exposed to a range of chemical entities, on the results of studies conducted in a developing country is just as flawed as relying on in vitro data. However, in most developed countries there are products that appear to have been evaluated only in trials in developing countries.

The third, and to us, the most significant point is that the efficacy data for pediculicides must be a high priority for regulatory authorities before granting a Marketing Authorisation (MA). This is significant because the products are used on children and must be safe, clinically effective, and also cost effective. In this instance the product continued to be marketed after the final data were available, despite clearly showing its lack of effectiveness. This raises a question of how many other products could be on the market without evidence of efficacy. No doubt makers of such products rely on consumer feedback and complaints as a guideline as to whether their products are both acceptable and effective. However, in practice, most manufacturers receive relatively few complaints about efficacy and few pediculicides have been subjected to the kind of post marketing feedback and complaints as a guideline as to whether their products are both acceptable and effective. However, in practice, most manufacturers receive relatively few complaints about efficacy and few pediculicides have been subjected to the kind of post marketing surveillance applied to some other medications.

Even in the rather more litigious circumstances prevailing in the United States of America, there have been few who have gone as far as legal action to press claims of inefficacy. In overall terms it was found that the legal complaint process was hindered by the regulatory process, as argued by the defence attorneys acting for various drug companies in one class action stating “...The claims (of the Plaintiffs) stand in direct conflict with the Food Drug and Cosmetic Act, moreover, because the ‘defendants’ medications cannot be sold for the treatment of head lice and labeled to say that the medications are not effective when simultaneously federal law and regulations require the labeling to say that the products are effective.”

(http://www.law360.com/articles/36260/drug-makers-fight-class-action-over-lice-treatment) thereby turning the onus for verification of efficacy back to the competent authority, in this case the federal Food and Drug Administration (FDA), although in another plaint the Texas Supreme Court gave a per curiam ruling “...that the FDA contains no such ‘complex and interrelated federal scheme of law, remedy, and administration’ that would divest the state courts of jurisdiction...” (http://www.supreme.courts.state.tx.us/historical/2005/feb/031052.htm) suggesting that those courts could, if they so chose, declare products ineffective and presumably thereby place the FDA in an difficult position with respect to its approval of certain preparations. Therefore, only products for which adequate clinical studies have been conducted, and then those data placed in the public domain with appropriate periodic review to ensure resistance has not affected efficacy, can be considered effective. Simply relying on a competent authority MA is not adequate justification for continuing to sell a product when there are doubts about its effectiveness, as we highlighted in presenting the results from one of our recent investigations. Just because products or active substances may have been effective when first introduced does not mean that they remain so, as indicated by more recent clinical investigations using some so-called “standard of care” products as comparators, and both industry and regulators should be responsive to changes in circumstance.

Data availability
F1000Research: Dataset 1. Individual demographic data collected from participants and lice found by detection combing post-treatment, 10.5256/f1000research.2026.d31728

Participant consent
Written and witnessed informed consent was obtained from the participants and parents or guardians of children under 18 years of age.

Author contributions
All authors contributed to the original study design. PN was responsible for the ethics applications and liaison with the community health authorities, and was the Supervising Clinician throughout the study. IFB and CMB contributed to the logistics of participant management, conducted all final assessments on Days 14 to 28, and evaluated the collections of lice on the Case Record Forms for determination of treatment outcome. IFB conducted resistance testing of lice, analysis of those results, and post hoc analyses on the historical data to correct some errors and omissions. IFB was responsible for drafting the manuscript and all three authors read the draft and commented on its contents.

Competing interests
IFB is a consultant to several large, medium, and small companies that develop and market head louse treatment products, including...
medicinal products, medical devices, cosmeceuticals, and combs. CMB and PN are not aware of any competing interests.

Grant information
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The conduct of this study was supported financially by Seton Healthcare Group plc, which contributed to the original design of the study but played no role in the decision to publish the results, permission for which was withheld while the company was in existence as an independent commercial entity. All confidentiality agreements between the investigators and the sponsor in relation to this work have now expired. Reckitt Benckiser plc, the current beneficiary of the legacy Seton Healthcare assets for this category of product, has kindly granted approval for publication of the manuscript and the original study protocol. The study was supported logistically by Bedfordshire Health Authority and Bedfordshire Family Health Authority, Luton, both of which have now been replaced by other services with different responsibilities and jurisdictions during various reorganisations of the National Health Service in the UK. Our thanks go to Deirdre Power and Stanley Lindsay who were the Study co-ordinators on behalf of South Bedfordshire Community Healthcare Trust. All treatments were supervised and follow up assessments performed, except days 14 to 28, by staff from Independent Nursing Services Ltd, Wootton, Bedfordshire. Thanks also to all the schools and General Practitioners who circulated Information Sheets to families in their care, as prospective participants. The original statistical analyses were conducted by Peter Lee, PN Lee Statistics and Computing Ltd, Sutton, Surrey, on behalf of the sponsor, Seton Healthcare Group plc. Sara Gray and Barbara Temesi, both of Seton Healthcare Group plc, monitored the study documentation on behalf of the sponsor.

Supplementary material
Supplementary files: https://f1000researchdata.s3.amazonaws.com/ supplementary/2026/0086c068-7ac2-42a1-8135-946436252380.zip

References


Open Peer Review

Maria Inés Picollo
Centro de Investigaciones de Plagas e Insecticidas, Consejo Nacional de Investigaciones Científicas y Técnicas (CONICET), Buenos Aires, Argentina

This is an interesting clinical study to assess the comparative effectiveness of three head louse treatments when are used by the public. For this purpose, the treatments selected were 0.5% d-phenothrin mousse, 0.2% d-phenothrin lotion, and wet-combing with conditioner. The commercial products were applied according to manufacturer's instructions (30 minutes for 0.5% phenothrin mousse and 2 hours for 0.2% phenothrin lotion), and wet-combing according to the Bug-Busting Community Hygiene Concern, London (fine toothed plastic comb with conditioner, repeated at 3–4 day intervals for 2 weeks).

I consider that the manuscript is within the scope of the journal, the abstract is according to the objectives and results of the manuscript: The methodology, results and discussion are carefully written and adequate to the objectives of the study.

One of the relevant aspects of the work is that reinforces the need for the effectiveness of commercial products must be warranted by the health authorities, due to the importance of pediculosis in school-aged children both in developed and developing countries.

An unexpected and surprising result is the low effectiveness of the three methods evaluated. Although pyrethroid resistance and inadequate formulation of the active ingredients surely contribute to the low effectiveness of the phenothrin-based formulations (20% and 21.5% success for mousse and lotion respectively), the low effectiveness of wet-combing (19.15% success) has not been previously reported. This result is worrying considering that the use of a fine comb for removing lice and nits was proposed as a relevant tool both in the diagnosis of infestations and as part of an integrated control strategy of head lice.

I agree with the authors in their first conclusion about the limitations of laboratory tests: “bioassay tests conducted in a laboratory, whether using laboratory reared insects or even wild collected ones, can only be indicative of efficacy for a formulation and it is unknown for a treatment to perform poorly in vitro yet be effective in vivo. ….. even ex vivo screens, using insects recently collected from the wild, may be only partially representative and several replicate tests should be performed using insects from geographically separated locations to ensure that the outcome is not obtained either by chance or due to some happenstance of physiological difference in the insects from that location”. But as the authors surely know and probably should clarify in the paper, the main purpose of laboratory testing is the “comparative-evaluation” of the effectiveness of pure compounds or formulations. Thus, the parallel and
simultaneous evaluation of different compounds and/or formulations made in standardized conditions, is a reliable result of the comparative activity of the products evaluated.

Obviously laboratory tests are faster and cheaper than clinical trials, and represent an excellent tool for a first selection of the compound or product to be developed (a pure compound without pediculicide activity in laboratory tests, will not be effective in a clinical trial).

**Competing Interests:** No competing interests were disclosed.

I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

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Deon Canyon
Office of Public Health Studies, University of Hawaii, Honolulu, HI, USA

This research was conducted well, and the article was well written. There is one grammatical error in the Outcomes section (paragraph 2): "In both insecticide treated groups the majority of lice at post-treatment assessments were juveniles, of which 712 first were [1st, 2nd...] stage nymphs that could only have originated from eggs not killed by insecticide." Also, nymphs can transfer so this conclusion is not accurate.

"However, success rates were significantly (p < 0.01) higher in people who had previously used a head louse treatment successfully, by an estimated factor of 1.88 (95% CI 1.13 to 3.11)." So how many of the failures may have been due to inappropriately applying a treatment?

The title and abstract are appropriate for the article, and provide a suitable summary. The experiment appears to have been conducted properly, with appropriate controls and data measurements, and the analysis is also adequate. The conclusions drawn from the study are both sensible and balanced, and the competing interests of the authors have been sufficiently disclosed.

I'm just left wondering if any of the participant variables had an influence on the outcomes? If there was no effect, this should be stated.

**Competing Interests:** No competing interests were disclosed.

I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

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Author Response 22 Jul 2014

Ian Burgess, Medical Entomology Centre, UK
Thanks for finding the typographical error in Outcomes section (paragraph 2) : "In both insecticide treated groups the majority of lice at post-treatment assessments were juveniles, of which 712 first were [1st, 2nd...] stage nymphs that could only have originated from eggs not killed by insecticide." that eluded everyone else reading it.

It should have read, "In both insecticide treated groups the majority of lice at post-treatment assessments were juveniles, of which 712 were first stage nymphs that could only have originated from eggs not killed by insecticide." While I do not disagree that young nymphs can transfer, this is a less common event than transfer of third stage nymphs and adults by some considerable factor of difference. Consequently, since relatively fewer adults were found in general it seems reasonable to assume that most, if not all, newly hatched first stage nymphs originated from eggs not killed by treatment. Consequently, I suggest a possible change to the text to read, "In both insecticide treated groups the majority of lice at post-treatment assessments were juveniles, of which 712 were first stage nymphs the majority of which most likely originated from eggs not killed by insecticide.", which would satisfy all possibilities.

On the second point about how many treatment failures may have been due to inappropriate or incompetent application of treatment, the answer is we simply do not know. Firstly, common sense tells us that someone who has previously successfully negotiated the pitfalls of applying a head louse treatment is more likely to achieve a similar success on a subsequent occasion. Secondly, although the agency staff engaged by the sponsor were not experienced at performing head louse treatments so I do wonder how good they may have been (or not as the case may be) at detecting failures in application method. Thirdly, this is a risk you take when conducting pragmatic (real use by consumer) clinical studies. For most medications missing one dose is usually not the end of the world and does not usually affect the ultimate outcome. However, for a single dose application head louse study not applying the product properly is usually terminal from the perspective of efficacy, irrespective of how effective the product may or may not be, and in this case they were not as effective as the sponsors believed.

So, in respect of participant variables, I do not doubt they influenced the outcome. The difficulty remains in determining which variable had what effect. In a wholly investigator run study, whether pragmatic or with investigator applied treatment, at least some of the variables are accountable and can be analysed accordingly. However, in this case I don't think anyone could have done much to greatly affect the overall outcome.

**Competing Interests:** None

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**Discuss this Article**

**Version 1**

**Author Response 21 Jan 2015**

**Ian Burgess, Medical Entomology Centre, UK**

The way Ibarra et al. write their comment on our article it implies they believe that delay of publication of this trial was a vindictive act to undermine the current Bug Buster kit. Frankly it was nothing of that kind.
The original intention was to publish the results as soon as possible after completion of the study but appointment of a new Medical Director at Seton Healthcare resulted in delaying tactics on his part, presumably to avoid having to face the truth of the ineffectiveness of the product, which failed to deliver its active ingredient due to insufficient alcohol solvent. No permission was granted throughout the life of the product by which time we had other things to concentrate on and Seton Healthcare and its successor companies ceased to exist. Subsequently, it has been a tricky task finding someone who could authorise release of the information, hence the delay.

The statement “The white comb, with round-point teeth spanning 72 mm, which is illustrated in Figure 2 of Burgess et al. (2014) was only found in the 1996 Bug Buster Kit, where it was included for optional use to reduce a heavy infestation” appears to be a curious comment considering the staunch defence of the comb put forward by Ms Ibarra at the North Thames Head Lice Symposium. However, whether this was the only comb or just one of the combs in the pack at the time I cannot say because I never saw one. As stated in the text, all initial contacts with participants and guidelines for treatment were made by agency nurses, and the Bug Buster packs were issued as received from the supplier. Presumably had there been more than one comb in the pack it could have been used, yet all of the combing group participants or caregivers that I met referred only to the white two-part comb in relation to their descriptions of what they had done. Consequently, I can only leave readers to draw their own conclusions as to what may or may not have been available in those packs.

When it comes to the next point “Additionally, the authors’ description of the Bug Busting method omits the vital second stage combing of the hair after rinsing the conditioner off”, again, I cannot state exactly what was advised other than that the instructions supplied with the Bug Buster pack were given to participants’ caregivers. As it happens this “instruction” comes as a revelation to me because I had never encountered it until reading the comment, so perhaps it is a newer instruction than existed at that time.

Irrespective of what readers may conclude about Bug Busting now or in the 1990s, Bug Busting was not the primary interest of the study, which was in fact whether the phenothrin mousse could work to eliminate louse infestation. The trial showed that quite clearly it could not. Information about the Bug Busting arm was included in the manuscript solely because it was included in the original protocol and those participants need to be recorded in the public domain having freely donated their time and efforts to the study. For information, Bug Busting was only included in the study in the first place because at the time the study was being designed most General Practitioners had minimised their commitment to prescribing chemical treatments and were issuing generic instructions to their patients to perform “wet combing with conditioner”, as they interpreted what was described in the Department of Health leaflet that placed this physical means of treatment (including Bug Busting) at the forefront of louse management and which had been widely circulated at the time so that levels of GP prescribing had dropped to their pre-1993 levels.

I can understand the concerns about methodology voiced by Ibarra and her wider group of colleagues in relation to the method of randomisation. I also share them. However, in conducting this study and a number of others my hands have been tied. I suggest Ms Ibarra takes up the issue with the Medicines and Healthcare products Regulatory Agency, which has consistently declined to approve randomisation by household when asked and have requested/required studies that have been submitted to them for scientific advice to be randomised by individual on advice from their statistical and other consultants.

As to the pros and cons of Bug Busting, or any other combing means of louse elimination, I take no particular position. For some people it/they work and for others it/they don’t, in much the same way that many of the other types of treatment may or may not be successful. As anyone who has been engaged in this area of study for any time knows, there is no simple answer to the problem and all we can do is try things out to see if they work, and even then it is only truly possible to say with any conviction that such and
such a treatment worked for those people at that particular time. Try it again on the same people next month or next year and a different result is not impossible, and possibly quite likely. This is why I made the points that the regulatory agencies of all countries need to get their act together to make sure that approved products really do work\(^1\) and that the claims made for them are not just based on one or two observations, or for that matter on dubious information\(^7\). When that happens, perhaps this debate will come to an end.


**Competing Interests:** IFB is, as stated previously, a consultant to various companies that develop and market head louse treatment products and devices, including combs. He has no particular loyalties and has used a wide variety of methodologies in assisting families to eliminate their head louse infestations on a "horses for courses" basis.

Reader Comment 28 Nov 2014

**Joanna Ibarra**, Community Hygiene Concern, UK

**Modern Bug Busting wet combing does work**

Burgess et al. have delayed until 2014\(^1\) to report on a trial carried out from June 1997 to March 1998, which is some 16 years later. In the report certain parts of an early version of the Bug Buster Kit (1996) are described, but this kit was rapidly superseded by the gold standard Bug Buster Kit (1998 version)\(^2\) a fact that the authors do not state clearly in 2014.

The poor results of their trial with an apparently incomplete and definitely outdated version of the Bug Buster Kit (19.1% success rate) have confused referee Maria Inés Picollo into doubting the worth of wet combing and fine combs in general\(^1\). This doubt arises due to the failure of Burgess *et al.* (2014) to mention that Hill *et al.* (2005)\(^2\) found that the efficacy of the re-designed 1998 combs positively influences the effectiveness of Bug Busting wet combing. In this study the participants were followed up over a year. 57% of children allocated to the 1998 Bug Buster Kit were cured at 15 days, which was four times more effective than the insecticide products tested; moreover 97% of families continued to use their kit until they eradicated their existing infestation and for prophylactic purposes thereafter (Hill *et al.*, 2006)\(^3\).
As with formulated treatments, the results of trials of mechanical cures cannot be simply extrapolated to other combs and procedures. We maintain that there is no generic wet combing, and the evidence base pertains to a variety of comb designs (in plastic or metal) and methods of combing, which produce different results.

The white comb, with round-point teeth spanning 72 mm, which is illustrated in Figure 2 of Burgess et al. (2014) was only found in the 1996 Bug Buster Kit, where it was included for optional use to reduce a heavy infestation. But the principal comb in any Bug Buster kit has a short tooth span, and the 1996 grey 45 mm Bug Buster detector comb, is not specifically identified in their report. There is simply reference to “the louse removal comb provided”. Additionally, the authors’ description of the Bug Busting method omits the vital second stage combing of the hair after rinsing the conditioner off. This is always completed using a short span comb. The white and grey combs were replaced in 1998 with the much superior bright yellow maxi Bug Buster (62 mm tooth span) and the mini Bug Buster (40 mm tooth span) both with slim handles and a deeply bevelled leading edge on the teeth.

We are concerned that Burgess et al. (2014) highlight the white comb because when laid flat, it and the maxi Bug Buster have a similar outline, although their design is completely different in other respects. In 2014, this emphasis creates confusion amongst health providers who could easily assume that the poor findings of Burgess and his co-authors in the 1990s are applicable to the current gold standard Bug Buster Kit and discredit wet combing of any kind.

Studies in Gent, Belgium (De Maeseneer et al, 2000, Vander Stichele et al, 2002) and Copenhagen, Denmark (Olsen, 2002) used the yellow combs. A video demonstration of the gold standard Bug Buster Kit in use can be accessed on the home page at www.chc.org. See also Ibarra (2014) for comb illustrations (Figures 1 and 3).

We also share the methodological concerns identified previously by Vander Stichele and Lapeere (2008) about the type of trial protocol used by Burgess et al. (2014). Randomisation is by individual so different entrants in the same family can receive different treatments. This procedure means that failure to kill mature lice on day 0 with any of the products to be evaluated immediately puts other members of the family, including other trial entrants allocated to a different product, at risk of new infestation because these adult lice are especially inclined to move to a new head during close contact. The procedure is dubious for similar-looking products, but totally unsatisfactory when mechanical products are compared with formulated ones. In the 1997/8 trial it meant that those individuals who were allocated to wet combing might have become free of mature lice after the first combing session on day 0 by diligently following the full instructions supplied in each kit. This would greatly diminish their chances of passing on lice but still leave them vulnerable to catching new incoming lice. In contrast, failure to kill mature lice in entrants in the same family receiving formulated treatments on day 0 would permit them to continue to export lice. The randomisation procedure chosen by Burgess et al. in their 1997/8 trial may be cost saving at the expense of clear cut results. Entering one individual per family into a comparative trial of treatments for head lice is likely to work better.

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References


**Competing Interests:** JI, FF and CW work (either voluntarily or paid) for Community Hygiene Concern (CHC), a charitable organisation set up in 1988 to help parents and health providers manage pediculosis capitis successfully. CHC develops, produces and distributes the Bug Busting resources on a not-for-profit, primary purpose trading basis.