Face washing promotion for preventing active trachoma (Review)

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Face washing promotion for preventing active trachoma

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ABSTRACT

Background
Trachoma remains a major cause of avoidable blindness among underprivileged populations in many developing countries. It is estimated that about 146 million people have active trachoma and nearly six million people are blind due to complications associated with repeat infections.

Objectives
The objective of this review was to assess the effects of face washing on the prevalence of active trachoma in endemic communities.

Search strategy
We searched the Cochrane Central Register of Controlled Trials - CENTRAL (which contains the Cochrane Eyes and Vision Group Trials Register) in The Cochrane Library (Issue 3, 2007), MEDLINE (1966 to October 2007), EMBASE (1980 to October 2007), the reference lists of identified trials and the Science Citation Index. We also contacted investigators and experts in the field to identify additional trials.

Selection criteria
We included randomised or quasi-randomised controlled trials, comparing face washing with no treatment or face washing combined with antibiotics against antibiotics alone. Participants in the trials were people normally resident in endemic trachoma communities.

Data collection and analysis
Two review authors independently extracted data and assessed trial quality. Study authors were contacted for additional information. Two clinically heterogeneous trials are included, therefore a meta-analysis was considered inappropriate.

Main results
This review included two trials with data from a total of 2560 participants. Face washing combined with topical tetracycline was compared to topical tetracycline alone in three pairs of villages in one trial. The trial found a statistically significant effect for face washing combined with topical tetracycline in reducing 'severe' active trachoma compared to topical tetracycline alone. No statistically significant difference was observed between the intervention and control villages in reducing ('non-severe') active trachoma. The prevalence of clean faces was higher in the intervention villages than the control villages and this was statistically significant. Another trial compared eye washing to no treatment or to topical tetracycline alone or to a combination of eye washing and tetracycline drops in children with...
follicular trachoma. The trial found no statistically significant benefit of eye washing alone or in combination with tetracycline eye drops in reducing follicular trachoma amongst children with follicular trachoma.

Authors’ conclusions

There is some evidence that face washing combined with topical tetracycline can be effective in reducing severe trachoma and in increasing the prevalence of clean faces. Current evidence does not however support a beneficial effect of face washing alone or in combination with topical tetracycline in reducing active trachoma.

PLAIN LANGUAGE SUMMARY

Face washing promotion for reducing active trachoma

Trachoma is an infectious eye disease. Active infection usually begins in childhood and is characterised by eye discharges, redness and irritation. Poor facial hygiene can lead to the disease spreading from person to person through eye-seeking flies or contaminated fingers. Face washing is promoted as part of the World Health Organization ‘SAFE’ strategy to eliminate blindness around the world. The review authors identified two randomised controlled trials with a total of 2560 participants set in Australia and Tanzania. One trial had face washing in combination with tetracycline as the intervention and tetracycline ointment alone as the control. The second trial compared eye washing to no treatment or to topical tetracycline alone or to a combination of eye washing and tetracycline drops in children with follicular trachoma. Both trials reported on active trachoma as an outcome measure but only one trial reported on severe trachoma and percentage of clean faces. The trials included in this review evaluated the effect of face washing over a three to 12 month period. There is some evidence that face washing combined with topical tetracycline can be effective in reducing severe trachoma and in increasing the prevalence of clean faces.

BACKGROUND

Epidemiology

Trachoma is an infectious eye disease caused by the microorganism Chlamydia trachomatis. Trachoma remains a major cause of avoidable blindness among underprivileged populations in many areas of Africa, Asia and the Middle East, where poverty, overcrowding, poor personal and environmental hygiene favour transmission of the disease. It is estimated that about 146 million people have active trachoma and nearly six million people are blind due to complications associated with repeat infections (WHO 1997a). The organism causing trachoma is spread from person to person by close contact in overcrowded living conditions, or through contaminated fingers or cloths used by mothers to wipe away discharges on the faces of children (ICEH 1999). Flies, which are attracted to eye and nasal secretions on the faces of infected children, are also believed to be risk factors in the transmission of the organism (ICEH 1999; West 1991).

Presentation

In communities where trachoma is endemic, infection usually begins in childhood and repeat episodes of infection cause distortion of the eyelids (entropion), in-turned eyelashes (trichiasis), corneal abrasion and ultimately blindness due to corneal opacity. Active trachoma is more commonly observed in children (Taylor 1985; West 1991). It is characterised by redness and discharge associated with inflammatory thickening of the upper tarsal conjunctiva (mucous membrane lining the inner surface of the upper eyelids) and follicles (whitish elevations within the conjunctiva). A simplified grading system for the assessment of trachoma and its complications in endemic communities has been published (Thylefors 1987) and discussed in a Cochrane review of antibiotics for trachoma (Mabey 2005).

The role of face washing in trachoma control

Face washing is promoted by the World Health Organization (WHO) programme for the global elimination of trachoma as
part of the 'SAFE' strategy (WHO 1997b; WHO 1999). The SAFE strategy consists of surgery for trichiasis; antibiotics for infectious trachoma; facial cleanliness to reduce transmission; and environmental improvements (household sanitation and provision of clean water). The face washing component of this strategy aims to maintain clean faces in the community in order to reduce eye-seeking flies and person-to-person transmission of the trachoma organism. Face washing promotion as a community intervention can be combined with mass treatment with antibiotics in areas with high trachoma endemicity. Mass treatment with antibiotics aims to reduce the reservoir of Chlamydia trachomatis in the community while face washing aims to interrupt the cycle of infection and re-infection in the long term. The antibiotic and environmental arms of the SAFE strategy have been examined in other published Cochrane reviews (Mabey 2005; Rabiu 2007).

Rationale for a systematic review

The face washing principle appears simple and theoretically sound, but whether this intervention can reduce transmission of trachoma in practice is now a focus of debate (Bailey 2001). Some narrative reviews of the literature have suggested that facial cleanliness may be useful in preventing trachoma (Emerson 2000; Pruss 2000). However, most of the data were obtained from observational studies and the methodological quality of the few controlled trials included was not reported. In this review we aim to summarise systematically, research evidence from trials of face washing promotion for preventing active trachoma in endemic communities. In communities where water is scarce, the uptake and practice of face washing may not be as good as in communities where water is freely available. The potential influence of water availability on outcomes will be considered in this review.

OBJECTIVES

The objective of this review was to assess the effects of face washing promotion on the prevalence of active trachoma in endemic communities.

METHODS

Criteria for considering studies for this review

Types of studies

We included randomised and quasi-randomised controlled trials.

Types of participants

Participants in the trials were people normally resident in communities where trachoma is endemic.

Types of interventions

We considered the following interventions:

(1) face washing promotion versus no intervention;
(2) face washing promotion plus mass antibiotic treatment versus mass antibiotic treatment alone.

Face washing promotion can be delivered by any means appropriate to the local setting such as: radio or television; health education leaflets; community leaders; religious gatherings; role-play; drama in village halls; school teachers; women groups; music etc.

In trials where promotion of face washing was combined with mass antibiotic treatment, antibiotics considered include tetracycline ointment or capsules; azithromycin; or erythromycin, given at any dose or frequency.

Types of outcome measures

We considered the following outcomes.

(1) Number of participants with active trachoma (TF or TI) at 6, 12, or greater than 12 months post-treatment allocation (age group as reported in trials).

Active trachoma was defined using the Thylefors 1987 scale. On this scale, active trachoma is categorised as TF or TI. TF is trachoma follicular inflammation and is defined as the presence of five or more follicles, each of which is at least 0.5 mm in diameter, on the flat surface of the upper tarsal conjunctiva. TI is trachoma intense inflammation and is defined as the presence of marked inflammatory thickening of the upper tarsal conjunctiva that obscures more than half of the deep conjunctival vessels.

We planned to include trials that used other trachoma grading scales to assess active trachoma, provided the scales used can be related to the Thylefors 1987 scale.

(2) Number of participants with an unclean face at 6, 12, or greater than 12 months post treatment allocation (age group as reported in trials).

An unclean face was defined as the presence of eye or nasal discharge (WHO 2001) or any other definition used in trials.

(3) Number of participants with severe trachoma.

Severe trachoma was not exclusively specified as an outcome in the protocol for this review. However, we felt it was important to report it since one of the two trials that met the inclusion criteria defined and reported this outcome.

Search methods for identification of studies

Electronic searches
We identified trials from the Cochrane Central Register of Controlled Trials - CENTRAL (which includes the Eyes and Vision Group Trials Register) in The Cochrane Library, MEDLINE and EMBASE. There were no language or date restrictions in the searches for trials. The electronic databases were last searched on 16 October 2007. See: Appendices for details of search strategies for each database.

Searching other resources

Trachoma experts that were contacted for potentially relevant studies include Peach H and West S. Mabey D was a peer reviewer and she provided information on potentially relevant studies. Existing reviews were identified and their citations were checked for relevant trials. We used the Science Citation Index to search for references that cite the studies that are included in the review.

Data collection and analysis

Selection of studies

Two review authors independently screened titles and abstracts found by the electronic searches. We retrieved for further assessment hard copies of trials that were potentially relevant to the review. Those that met the selection criteria were assessed for methodological quality. Disagreements were resolved by discussion.

Assessment of methodological quality

Two review authors independently assessed included trials using the following criteria based on Section 6 of the Cochrane Handbook for Systematic Reviews of Interventions (Higgins 2006):

1) Concealment of allocation: A - Adequate; B - Unclear; C - Inadequate. For trials with unclear concealment of allocation, further information was sought from primary investigators.

2) Baseline comparability between intervention and control groups with respect to prevalence of active trachoma: A - Adequate if no substantial differences present; B - Unclear if not reported or not known whether substantial differences exist; C - Inadequate if substantial differences exist.

3) Comparability between intervention and control groups with respect to follow up: A - Adequate if no substantial differences in follow up rates; B - Unclear if not reported or not known; C - Inadequate if substantial differences exist in follow up rates.

4) Intention-to-treat analysis: A - Adequate if performed; B - Unclear if not known; C - Inadequate if not performed. Masking of participants and providers were not used to assess trial quality in this review. The nature of the intervention made it difficult to successfully apply masking. Posthoc we decided to use masking of outcome assessors as a parameter of quality.

Data collection

Two review authors independently extracted data onto a standardised data extraction form. We compared extracted data and reconciled differences. Disagreements were resolved by a third review author. Where studies reported the outcomes in different ways, primary investigators were contacted for further information to allow transformation of data.

Data analysis

Only two trials met the inclusion criteria for this review and these used different interventions and methods for outcome assessment. A meta-analysis was considered inappropriate and a narrative summary of results is presented. If additional studies become available in the future we will use the following methods: The specified outcomes are dichotomous therefore only relative risks will be calculated. Data will be combined in a meta-analysis if appropriate, using the random-effects model. If there are fewer than three studies and little evidence of heterogeneity a fixed-effect model will be used. In analysing cluster-randomised trials, if we encounter trials where the units of allocation and analysis are different (i.e. the unit of allocation was the community and the unit of analysis was individuals in the community) and this has not been accounted for in the analysis, we will contact primary investigators for additional data to develop estimates of intraclass correlation coefficients or design effect to calculate more appropriate confidence intervals. If a meta-analysis is not possible, a tabulated summary of results will be presented. We will not rely on statistical significance of a chi squared test to indicate heterogeneity but will consider this at all times during the review. The existence of heterogeneity may be apparent on visual examination of the forest plot. If present, heterogeneity will be explored using the following subgroups:

1) Communities with available water supply versus communities with scarce water supply. Water availability is defined in this review as the presence of a functional water source within 30 minutes walk or a distance of less than four kilometres from all households within the community (WHO 2001) or any other definition used in the trials.

2) Communities with intense active trachoma versus communities with less intense active trachoma. Intense active trachoma is defined in this review as communities with a baseline prevalence of TF or TI equal to or greater than 20%, while less intense is defined as communities with a prevalence of TF or TI less than 20% (WHO 1997b). If possible we will conduct a sensitivity analysis to investigate the influence of studies with quasi-random methods and those without concealment of allocation on the overall estimates of effect.

RESULTS
Description of studies

See: Characteristics of included studies; Characteristics of excluded studies.

Results of the search

The electronic searches generated 67 citations and abstracts. These were screened and the full text of two potentially relevant articles were retrieved for further assessment. One of these met the criteria for inclusion (West 1995). The other was not a randomised controlled trial and therefore not included in the review (Sutter 1983). A trachoma research expert drew our attention to a randomised trial that was not published in a journal (Peach 1987). In all two randomised trials are included in the review.

Updated searches

An updated search done in October 2007 identified 66 new reports of trials. The Trials Search Co-ordinator scanned the search results and removed any references which were not relevant to the scope of the review. The full text of three articles were checked for potential inclusion, however all were excluded. Edwards 2006 and Rubinstein 2006 were reports of health education promotion of face washing and Khandekar 2006 treated face washing and environmental sanitation interventions as one outcome.

Included studies

See Characteristics of included studies for further details.

Setting and participants

This review includes data from a total of 2560 participants in two trials. West 1995 was undertaken in Kongwa, Tanzania. In this trial a total of 1417 children, aged one to seven years from six villages, were randomised in three pairs to intervention or control. Peach 1987 was undertaken in the Northern Territory of Australia. In this trial 36 aboriginal communities were randomised to one of three intervention arms or one control arm. A total of 2530 children aged five to 14 were screened for follicular trachoma. A few more children above the age of 14 and some of preschool age were also screened. Of the total number of children screened in the participating communities, only 1143 children with follicular trachoma were recruited into the trial.

Interventions

In West 1995, 680 children from three villages were randomised to face washing promotion combined with tetracycline and 737 children from three villages were randomised to tetracycline ointment alone. Face washing promotion was community based and consisted of neighbourhood meetings to build consensus for increasing face washing and reinforcement activities such as school plays, seminars with the traditional healers and meetings with other village groups. Face washing promotion was carried out for one month during and after mass treatment with tetracycline. Tetracycline ointment was administered topically once daily for 30 days. In Peach 1987, 374 were randomised to tetracycline eye drops, 246 children were randomised to eye washing, 312 children were randomised to eye washing combined with tetracycline eye drops, and 211 children were randomised to the no treatment group. Children in the eye washing group had their eyes washed daily by school teachers for three months. Those in the tetracycline group had tetracycline eye drops applied daily for one week every month for three months. For the purpose of this review, data for the comparison between eye washing versus no treatment, and eye washing combined with eye drops versus eye drops alone are reported.

Outcome measures

In West 1995, outcomes reported include active trachoma, severe trachoma and clean faces. Trachoma was graded using the Thylefors 1987 scale. Severe trachoma was defined exclusively in the trial as 15 or more follicles, or the presence of inflammation that obscured all vessels of the tarsal plate. We extrapolated and extracted data from graphs presented in the report of the trial, as raw data were unavailable. These extrapolated data should be regarded as best approximations to the true figures. Our protocol specified ‘unclean faces’ as the outcome of interest, but this was reported as ‘clean faces’ in this trial. We have elected to present the outcome as reported in the trial as it would be difficult to transform the data without sufficient information from the trialists.

In Peach 1987, outcome was reported as the proportion of children with follicular trachoma who had follicles at three months after the intervention. The Aboriginal Health Workers simplified grading scheme was used to assess the presence of follicles as indicating active trachoma. Although this scale can be crudely compared to the TF grading on the Thylefors scale, it may have a lower specificity because of the tendency to classify participants with fewer than five follicles as having active trachoma.

Risk of bias in included studies

In West 1995 there was no information on how randomisation was done and whether allocation of villages to intervention or control was concealed. Baseline prevalences in active trachoma between comparison villages were not substantially different. Although 92% of the enrolled participants were followed up for one year, information regarding similarity of follow-up rates between comparison groups was not provided in the report. Information
on whether analysis of results was based on an intention to treat principle was not provided in the report. We note that this trial masked outcome assessment by taking photographs of tarsal plate read by an examiner who was not aware of the randomisation status of the villages.

In Peach 1987, details of how randomisation was done and concealment of allocation were not available in the report. Additional information from the author reveals that a random numbers table was used to allocate communities to the interventions or control group. The allocation was done after the initial screening by someone who was unaware of the prevalence of trachoma and unfamiliar with the communities, including their school teachers and health workers. It is unclear whether baseline prevalence of trachoma was similar among the comparison groups. Information on the number of communities randomised to each experimental group was not available in the report. However, during further correspondence the authors suggest that about nine communities were randomised to each arm. Almost 89% of enrolled participants were followed up for three months. All participants lost to follow up were assumed to have follicles at the end of the study and the intention to treat principle was applied in the analysis of results. Outcome was assessed by trachoma workers who were unaware of treatment allocation to the communities. Steps were taken to ensure that the outcome assessors did not learn which groups the communities were allocated to.

Table 1 gives the results of the assessment of methodological quality of the included trials.

<table>
<thead>
<tr>
<th>Trial ID</th>
<th>Method randomisation</th>
<th>Allocation conceal.</th>
<th>Baseline compar.</th>
<th>Attrition</th>
<th>Intention to treat</th>
</tr>
</thead>
<tbody>
<tr>
<td>Peach 1987</td>
<td>Unclear</td>
<td>Unclear</td>
<td>Adequate</td>
<td>Unclear</td>
<td>Unclear</td>
</tr>
<tr>
<td>West 1995</td>
<td>Adequate</td>
<td>? Adequate</td>
<td>Unclear</td>
<td>Adequate</td>
<td>Adequate</td>
</tr>
</tbody>
</table>

**Effects of interventions**

The two trials included were different in several respects, particularly with regard to types of intervention and definition of outcome measures. Therefore a meta-analysis was not considered appropriate. A narrative summary of the results is presented.

**Active trachoma (follicular or TF or TI)**

In West 1995, face washing combined with antibiotics was compared to antibiotics alone in three pairs of villages. In pair one, the percentage prevalence of active trachoma was lower in the village that received a combination of face washing and antibiotics than the village that received antibiotics alone at 12 months follow up (approximately 55% compared to 60%). In a second pair of villages, the percentage of active trachoma was also lower in the combination village than the antibiotic alone village (approximately 40% compared to 50%). However in a third pair of villages, the percentage of active trachoma in the combination village was higher than the antibiotics alone village (approximately 70% compared to 65%). The overall results for all the combination vil-
lages compared to the antibiotics alone villages suggest a reduction in the odds of any trachoma but this effect was not statistically significant (odds ratio (OR) 0.81, 95% confidence interval (CI) 0.42 to 1.59).

In Peach 1987, 191/246 children (77.6%) in the eye washing arm had follicles at three months compared to 160/211 (75.8%) in the no treatment arm. The difference was not statistically significant (p = 0.73). In the eye washing/eye drop combination arm, 215/312 (68.9%) had follicles at three months compared to 250/374 (66.8%) in the eye drop only arm. The difference was not statistically significant (p = 0.62). When a logit model was fitted to the data, taking age of participants, geographical location and trachoma outcome assessors into account, the results show that the odds of having follicular trachoma was higher in the eye drop only arm compared to the eye washing-eye drop combination (odds 1.17 to 1.00) but these odds were not significantly greater than 1 to 1. The odds of having follicular trachoma in the no treatment group compared to the eye washing group were similar (odds 1.02 to 1.00).

Severe trachoma

In West 1995 the three pairs of villages were also compared with respect to prevalence of severe trachoma. In pair one, the percentage prevalence of severe trachoma was lower in the village that received a combination of face washing and antibiotics than the village that received antibiotics alone at 12 months follow up (approximately 8% compared to 14%). In a second pair of villages, the percentage of active trachoma was also lower in the combination village than the antibiotic alone village (approximately 5% compared to 14%). However in a third pair of villages, the percentage of active trachoma in the combination village was slightly higher than the antibiotics alone village (approximately 10% compared to 8%). The overall results after adjustments for age and baseline trachoma status suggests a reduction in the odds of severe trachoma by the face washing antibiotic combination compared to antibiotic alone and this effect was statistically significant (OR 0.62, 95% CI 0.40 to 0.97). At six months follow up, there were no differences in the prevalence of severe trachoma between the intervention and control groups in the three pairs of villages. Peach 1987 did not report this outcome.

Clean faces

In West 1995 the percentage of children with clean faces was consistently higher in the face washing-antibiotic combination villages than the antibiotic alone villages. Total results showed an increase in the percentage of children with clean faces in the face washing/antibiotic combination villages from 18% at baseline to 33% at six months and 35% at 12 months follow up. There was a smaller increase in the percentage of children with clean faces in the antibiotic alone group (from 19% at baseline to 30% at six months and 26% at 12 months). The difference in the proportion of children with clean faces in the intervention villages compared to the control villages was statistically significant (p < 0.05). Peach 1987 did not report this outcome.

**DISCUSSION**

Although two trials are included in this review, a meta-analysis was not performed. This was because of notable clinical heterogeneity between the two trials, particularly with regard to intervention strategies and outcome definition. Although the report of the design and conduct of both trials suggests notable efforts by the investigators to strengthen the quality, lack of adequate information made it impossible objectively to assess the trials against some key quality parameters specified in the review (see section on methodological quality and additional Table 1). Outcomes were reported at three months in Peach 1987. Although the follow-up period fell short of what was specified in our protocol, we did not exclude the data from this trial in view of the paucity of randomised trials.

**Active trachoma**

It is unclear why face washing promotion combined with tetracycline had an effect in reducing active trachoma in two pairs of villages but no effect in a third pair in West 1995. Differences in baseline characteristics such as prevalence of trachoma, intensity of transmission, availability or access to water supplies between the third pair and the first two pair of villages may be important in explaining the differences in benefit. However, the overall results for the face washing/tetracycline combination villages compared to the tetracycline only villages suggest a modest beneficial effect of face washing in reducing active trachoma at 12 months, although this was not statistically significant. Peach 1987 suggests no benefit for face washing compared with no treatment. The raw data also show no benefit for the face washing/eye drops combination in comparison to eye drops alone. The age of participants varied and the authors observed a higher proportion of severe trachoma among older children. There were variations in the prevalence of trachoma in the different geographical locations from which the participating communities were drawn as well as slight differences in the diagnostic competence of outcome assessors. The authors hypothesised that community randomisation as done in the trial may not have adequately controlled for these factors hence the need to account for them in the logit model. After fitting the data to a logit model to control for perceived imbalances in the ages of participants, geographical location and outcome assessors, a marginal but not statistically significant benefit is suggested for the face washing/eye drops combination over the eye drops alone group. The report however does not state whether the analysis of results in a logit model was planned in advance or
simply informed by the apparent lack of effect suggested by the raw data.

The lack of effect of face washing in Peach 1987 can be explained by a number of factors. Firstly, the trachoma grading system used in the trial can potentially influence the results. Participants were recruited into the trial on the basis of whether follicles or papillae were present. Based on this definition, participants with follicles/papillae from causes other than active trachoma could have been included. For this group of people, treatment would appear to have no benefit if what is being treated is not trachoma. Furthermore, if participants had trachoma which was not intense, the effect of the face washing may not be readily apparent. Secondly, in analysing the results using the intention to treat principle, the authors assumed that the participants lost to follow up had follicular trachoma at the end of the study. If this assumption was inaccurate and there were more participants lost to follow up in a treatment group compared to control, as was the case with the eye washing arm (17% versus 10.4%), treatment might appear to be ineffective compared to control. However, a sensitivity analysis with the missing participants excluded from analysis did not alter the results. Thirdly, the intervention was administered for only three months. A longer intervention period and follow up may have significantly altered the results. Fourthly, in Peach 1987, face washing was applied to children with already established disease rather than the whole population at risk, and outcome was measured in this group of children. The face washing strategy aims to reduce active trachoma in endemic communities mainly by reducing the transmission of the disease. A better measure of effect would have been to evaluate the magnitude of the disease amongst the whole population or subset of the population rather than amongst persons with the disease, or to determine the number of new cases of disease since institution of the intervention. It is also unclear how much impact on transmission can be achieved by applying face washing only to individuals with the disease in endemic communities. The true impact of face washing on active trachoma in the communities might be better evaluated by a study design in which face washing is applied to whole populations at risk rather than only those with the disease.

Severe trachoma

As for active trachoma, benefits of face washing in reducing the prevalence of severe trachoma were apparent in the first and second pairs of villages in West 1995 at 12 months follow up. In the third pair, there appeared to be no benefit. The overall results for all the villages after adjusting for age and baseline trachoma status showed a benefit of face washing in reducing severe trachoma in the intervention villages compared to the control villages at 12 months follow up. It is probable that participants with severe active trachoma represent a subgroup with more intense transmission and therefore face washing, which aims to break transmission, would be more likely to show a stronger effect within this subgroup. On the other hand, the appropriateness of combining the results from the three pairs of villages is questionable, since presumably the villages were paired because of some differences between them. It is unclear why face washing showed no comparative benefit in the three pairs of villages at six months follow up. Apparent benefit at 12 months underscores the importance of a longer follow-up period to demonstrate impact of the intervention.

Clean faces

We note with interest that the percentage of participants with clean faces increased in both intervention and control groups over 12 months, even though the increase was higher in the intervention group. However, a statistically significant difference in the percentage of clean faces between the intervention and control groups at 12 months suggests a benefit of face washing promotion. Previous narrative reviews of the literature have reported possible beneficial effects of face washing in preventing active trachoma. The conclusions of these narrative reviews were based on data obtained from one trial and a number of observational studies. This review assessed the methodological quality of the trials included and found that some important quality parameters were not adequately addressed.

Authors’ Conclusions

Implications for practice

Evidence from one trial suggests that face washing can be effective in increasing facial cleanliness and in reducing severe trachoma, but its effect in reducing active trachoma is inconclusive. In another trial, there was no evidence of effect of face washing alone or in combination with tetracycline in reducing active trachoma in children with already established disease.

Implications for research

The trials included in this review evaluated the effect of face washing over a three to 12 month period. However, it is unclear whether this time period is long enough for a face washing promotional activity to demonstrate impact of the intervention. Therefore, future research should include longer follow-up periods and also address the questions of whether reinforcement activities are required over time to improve outcome. The reporting of the methodology of trials should be complete to enable reviewers and readers to assess the validity of their conclusions.
We are grateful to the editorial team of the Cochrane Eyes and Vision Group for executing the electronic searches. We also gratefully acknowledge the tremendous advice and co-operation we received from Professor Peach, the author for correspondence of one of the trials included in this review. We acknowledge the invaluable contribution of Denise Mabey who peer reviewed this review and also of Catey Bunce, Marie Diener-West and Roberta Scherer for their statistical and methodological advice.

REFERENCES

References to studies included in this review

Peach 1987 {published data only}

West 1995 {published data only}

References to studies excluded from this review

Edwards 2006 {published data only}

Khandekar 2006 {published data only}

Rubinstein 2006 {published data only}

Sutter 1983 {published data only}

Additional references

Bailey 2001

Emerson 2000

Glanville 2006

Higgins 2006

ICEH 1999

Mabey 2005

Pruss 2000

Rubinstein 2007

Taylor 1985

Thylefors 1987
**WHO 1997a**

**WHO 1997b**

**WHO 1999**

**WHO 2001**

* Indicates the major publication for the study
## Characteristics of included studies [ordered by study ID]

### Peach 1987

<table>
<thead>
<tr>
<th>Methods</th>
<th>36 aboriginal communities were randomised in stages to four experimental groups after stratification by geographical location. Method of randomisation: not stated. Unit of randomisation: communities, but individuals where analysed. Masking: outcome assessors masked but method of masking unclear. Analysis was by intention to treat principle (participants lost to follow up were assumed to have follicles at the end of the study).</th>
</tr>
</thead>
<tbody>
<tr>
<td>Participants</td>
<td>Country: Northern Territory of Australia. Participants: children aged five years and above drawn from 36 aboriginal communities. Age range: most participants were between 5 and 14 years, although a small proportion of children were older than 14 and a small proportion were pre-school. Total number of children randomised: 1143.</td>
</tr>
<tr>
<td>Interventions</td>
<td>Treatment: 1) Tetracycline eye drops daily for one week every month for 3 months (374 children randomised). 2) Eye washing daily for 3 months (246 children randomised). 3) Tetracycline eye drops plus eye washing (312 children randomised). Control: No treatment (211 children)</td>
</tr>
<tr>
<td>Outcomes</td>
<td>Follicular trachoma (proportion of children with follicular trachoma at 3 months).</td>
</tr>
</tbody>
</table>

### Risk of bias

<table>
<thead>
<tr>
<th>Item</th>
<th>Authors' judgement</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Allocation concealment?</td>
<td>Unclear</td>
<td>B - Unclear</td>
</tr>
</tbody>
</table>

### West 1995

<table>
<thead>
<tr>
<th>Methods</th>
<th>Six villages were randomised in three pairs to intervention or control. Method of randomisation: unclear. Masking: outcome assessors masked. Assessors examined photographs of tarsal plates for follicles.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Participants</td>
<td>Country: Kongwa, Tanzania. Participants: Children aged 1 to 7 years drawn from 6 trachoma endemic villages. Total number of children randomised: 1417.</td>
</tr>
<tr>
<td>Interventions</td>
<td>Treatment: Face washing promotion combined with mass tetracycline ointment (680 children).</td>
</tr>
</tbody>
</table>
Control: Mass tetracycline ointment only (737 children). Tetracycline ointment was administered topically once daily for 30 days.

Outcomes

1) Active trachoma.
2) Severe trachoma.
3) Clean face.

Notes

Risk of bias

<table>
<thead>
<tr>
<th>Item</th>
<th>Authors’ judgement</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Allocation concealment?</td>
<td>Unclear</td>
<td>B - Unclear</td>
</tr>
</tbody>
</table>

Characteristics of excluded studies  [ordered by study ID]

<table>
<thead>
<tr>
<th>Study</th>
<th>Reason for exclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Edwards 2006</td>
<td>Study intervention is health education promotion of face washing</td>
</tr>
<tr>
<td>Khandekar 2006</td>
<td>Unable to separate the effect of face washing from environmental sanitation interventions as both were indirectly examined as &quot;one intervention&quot;</td>
</tr>
<tr>
<td>Rubinstein 2006</td>
<td>Study intervention is health education promotion of face washing</td>
</tr>
<tr>
<td>Sutter 1983</td>
<td>Not a randomised controlled trial</td>
</tr>
</tbody>
</table>
This review has no analyses.

**APPENDICES**

**Appendix 1. CENTRAL search strategy used for Issue 3, 2007**

#1 MeSH descriptor Trachoma  
#2 MeSH descriptor Chlamydia trachomatis  
#3 (trachom*)  
#4 (tracom*)  
#5 (follicular near conjunctivitis)  
#6 (intense near conjunctivitis)  
#7 (#1 OR #2 OR #3 OR #4 OR #5 OR #6)  
#8 MeSH descriptor Hygiene  
#9 MeSH descriptor Face  
#10 face-wash or facewash*  
#11 face near wash*  
#12 facial near wash*  
#13 face near clean*  
#14 facial near clean*  
#15 face near hygien*  
#16 facial near hygien*  
#17 (#8 OR #9 OR #10 OR #11 OR #12 OR #13 OR #14 OR #15 OR #16)  
#18 (#7 AND #17)

**Appendix 2. MEDLINE search strategy used up to October 2007**

1 exp clinical trial/ [publication type]  
2 (randomized or randomised).ab,ti.  
3 placebo.ab,ti.  
4 dt.fs.  
5 randomly.ab,ti.  
6 trial.ab,ti.  
7 groups.ab,ti.  
8 or/1-7  
9 exp animals/  
10 exp humans/  
11 9 not (9 and 10)  
12 8 not 11  
13 exp trachoma/  
14 exp chlamydia-trachomatis/  
15 or/13-14  
16 trachoma$ .tw.  
17 (follicular adj3 conjunctivitis).tw.  
18 (intense adj3 conjunctivitis).tw.  
19 or/16-18  
20 15 or 19
The search filter for trials at the beginning of the strategy is from the published paper by Glanville et al (Glanville 2006).

**Appendix 3. EMBASE search strategy used up to October 2007**

1 exp randomized controlled trial/
2 exp randomization/
3 exp double blind procedure/
4 exp single blind procedure/
5 random$.tw.
6 or/1-5
7 (animal or animal experiment).sh.
8 human.sh.
9 7 and 8
10 7 not 9
11 6 not 10
12 exp clinical trial/
13 (clin$ adj3 trial$).tw.
14 ((singl$ or doubl$ or trebl$ or tripl$) adj3 (blind$ or mask$)).tw.
15 exp placebo/
16 placebo$.tw.
17 random$.tw.
18 exp experimental design/
19 exp crossover procedure/
20 exp control group/
21 exp latin square design/
22 or/12-21
23 22 not 10
24 23 not 11
25 exp comparative study/
26 exp evaluation/
27 exp prospective study/
28 (control$ or prospectiv$ or volunteer$).tw.
29 or/25-28
30 29 not 10
31 30 not (11 or 23)
32 11 or 24 or 31
33 exp trachoma/
34 exp chlamydia-trachomatis/
35 or/33-34
36 trac?oma$.tw.
37 (follicular adj3 conjunctivitis).tw.
38 (intense adj3 conjunctivitis).tw.
39 or/36-38
40 35 or 39
41 exp personal hygiene/
42 exp face/
43 or/41-42
44 ((face$ adj3 wash$) or clean$ or hygien$).tw.
45 ((facial adj3 wash$) or clean$ or hygien$).tw.
46 (face adj1 wash$).tw.
47 or/44-46
48 43 or 47
49 40 and 48
50 32 and 49

WHAT'S NEW
Last assessed as up-to-date: 15 October 2007.

<table>
<thead>
<tr>
<th>Date</th>
<th>Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>23 October 2008</td>
<td>Amended</td>
<td>Converted to new review format.</td>
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HISTORY
Protocol first published: Issue 2, 2002
Review first published: Issue 3, 2004

<table>
<thead>
<tr>
<th>Date</th>
<th>Type</th>
<th>Description</th>
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</thead>
<tbody>
<tr>
<td>20 February 2008</td>
<td>New search has been performed</td>
<td>Issue 2 2008: three new trials were identified in an updated search but were excluded.</td>
</tr>
<tr>
<td>31 March 2004</td>
<td>New citation required and conclusions have changed</td>
<td>Substantive amendment</td>
</tr>
</tbody>
</table>
CONTRIBUTIONS OF AUTHORS

Conceiving the review: HE
Coordinating the review: HE
Data collection for the review: HE
Screening search results: HE, MA
Organising retrieval of papers: HE
Screening retrieved papers against inclusion criteria: HE, MA, MR
Appraising quality of papers: HE, MA, MR
Abstracting data from papers: HE, MA, MR
Writing to authors of papers for additional information: HE
Resolving disagreements between reviewers: MR
Entering data into RevMan: HE
Interpretation of the data: HE, MR
Writing of the review: HE, MA, MR

DECLARATIONS OF INTEREST

None known.

SOURCES OF SUPPORT

Internal sources

• No sources of support supplied

External sources

• Effective Health Care Alliance Project, International Health Division, Liverpool School of Tropical Medicine, UK.
• National Eye Centre, Nigeria.

INDEX TERMS
Medical Subject Headings (MeSH)

*Chlamydia trachomatis; *Face; Anti-Bacterial Agents [administration & dosage]; Randomized Controlled Trials as Topic; Skin Care [*methods]; Tetracycline [administration & dosage]; Trachoma [*prevention & control]

MeSH check words

Humans