Topical preparations for the treatment of mild-to-moderate acne vulgaris: systematic review and network meta-analysis*

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Summary

Background Acne is very common and can have a substantial impact on wellbeing. Guidelines suggest first-line management with topical treatments, but there is little evidence regarding which treatments are most effective.

Objectives To identify the most effective and best tolerated topical treatments for acne using network meta-analysis.

Methods CENTRAL, MEDLINE, Embase and World Health Organization Trials Registry were searched from inception to June 2020 for randomized trials that included participants with mild/moderate acne. Primary outcomes were self-reported improvement in acne, and trial withdrawal. Secondary outcomes included change in lesion counts, Investigator’s Global Assessment, change in quality of life and total number of adverse events. Network meta-analysis was undertaken using a frequentist approach. Risk of bias was assessed using the Cochrane Risk of Bias Tool and confidence in evidence was assessed using CINeMA.

Results A total of 81 papers were included, reporting 40 trials with a total of 18 089 participants. Patient Global Assessment of Improvement was reported in 11 trials. Based on the pooled network estimates, compared with vehicle, benzoyl peroxide (BPO) was effective (35% vs. 26%) for improving self-reported acne. The combinations of BPO with adapalene (54% vs. 35%) or with clindamycin (49% vs. 35%) were ranked more effective than BPO alone. The withdrawal of participants from the trial was reported in 35 trials. The number of patients withdrawing owing to adverse events was low for all treatments. Rates of withdrawal were slightly higher for BPO with adapalene (2.6%) or clindamycin (2.7%) than BPO (1.8%) or adapalene alone (1.0%). Overall confidence in the evidence was low.

Conclusions Adapalene in combination with BPO may be the most effective treatment for acne but with a slightly higher incidence of withdrawal than monotherapy. Inconsistent reporting of trial results precluded firmer conclusions.

What is already known about this topic?

- Guidelines suggest a number of different topical preparations as first-line treatment for acne vulgaris.
Acne vulgaris (hereafter ‘acne’) is very common in both adolescents and adults.1 Acne can have significant impact on quality of life, including increased risk of depression.2 Guidelines differ in their recommendations and quality,3 but National Institute for Health and Care Excellence Clinical Knowledge Summary (NICE CKS) UK guidelines suggest that first-line treatment should be a single-agent topical treatment, followed by combination topical treatment.4 Guidelines in the USA, Canada and Europe are similar, recommending combination topical treatment as first-line therapy.5–7 Although topical preparations, such as benzoyl peroxide (BPO) and topical retinoids (e.g. adapalene) can be effective, there is uncertainty regarding the most appropriate strategy for initial and maintenance treatment.2 While the prescription of topical antibiotics as monotherapy in the UK is declining, topical antibiotics as monotherapy or in combination are still widely prescribed8 and contribute to antibiotic resistance.9,10

A 2014 James Lind Alliance Priority Setting Partnership for acne included the question ‘What is the best topical product for treating acne?’ in their top 10 priorities for future research.11 There are multiple topical acne treatments and it is not feasible to review and compare them all. However, it is reasonable to address the question set out in the Priority Setting Partnership by comparing treatments suggested in European guidelines as first-line topical preparations for mild and moderate acne that are prescribed in the UK.

Although these treatments are widely used, there are gaps in the evidence base regarding their effectiveness and tolerability. To date, there have been two Cochrane reviews that have assessed topical treatments for acne.12,13 However, these reviews were able to provide only limited head-to-head evidence for key treatments, including adapalene + BPO, which are widely used and recommended in guidelines.

The uncertainty in the evidence base regarding optimal choice of topical treatments for acne is important because (i) topical antibiotics, alone or in combination, may be used despite being no more effective than topical nonantibiotic treatments, (ii) uncertainty leads to potential delays in treating acne effectively, and (iii) patients may progress to other treatments if acne does not improve, e.g. long courses of oral antibiotics.

While traditional meta-analysis is limited to direct head-to-head comparisons, network meta-analysis techniques, sometimes also called multiple-treatments meta-analysis, can overcome this by using all available data to build a network of direct and indirect comparisons. It allows estimates of effectiveness of treatment in addition to estimates of incoherence (how well the whole network fits together).14

Materials and methods

Protocol and registration

The study was conducted and is reported in line with the PRISMA-NMA guideline15 and was preregistered on PROSPERO (CRD42019135570).

Public and patient involvement

Prior to undertaking this study, we convened a ‘patient panel’ of 10 people with current/former acne. We discussed the research question and how we might measure ‘effectiveness’ and ‘adverse events’. The patient panel felt strongly that a participant-reported outcome should be the primary measure; it was their own assessment of their acne that mattered most to them, not the assessment of a clinician. The patient panel also helped to decide on the scope of the review, stressing the importance of understanding whether prescribed topical medications actually worked. The panel saw little value in including medications not currently available to them in the UK. One member of the patient panel joined the study team and is a coauthor of this article.

Search strategy and information sources

We searched the Cochrane Central Register of Controlled Trials, MEDLINE and Embase, from inception to June 2020,
for relevant journal articles, conference abstracts and systematic reviews (Appendix S1; see Supporting Information). Our search was not limited by language. We also searched the World Health Organization International Clinical Trials Registry for relevant registered trials; we hand-searched references from included papers and relevant systematic reviews for additional relevant trials and we contacted experts and pharmaceutical companies to find any unpublished trials.

Study selection

We included randomized controlled trials but excluded split-face and split-body trials owing to concerns about contamination, quasirandomized trials and any nonrandomized designs.

Two reviewers independently screened all titles, abstracts and full papers, using the eligibility criteria below, with any disagreements resolved through discussion. We obtained and assessed full papers or conference abstracts for inclusion in the review only if they were written in English. However, we kept a record of papers not written in English whose title and abstract were potentially relevant for inclusion in future updates.

Eligibility criteria

Population

We included all trials where participants had mild-to-moderate acne (as defined by trial authors), regardless of age, sex, setting or previous treatments. We included trials in which there were mixed populations of acne severity, provided ≤ 50% of participants had severe acne. We excluded trials in which severity was not reported, or where it was unclear from the source material whether the trial was randomized.

We excluded trials in which all participants had truncal acne only, were diagnosed with rosacea, unusual forms of acne, chloracne, acne inversa, acne fulminans, neonatal acne, infantile acne, occupational acne, drug-induced acne and acne specifically associated with endocrinopathies, including polycystic ovary syndrome, had previously received oral isotretinoin, or were only using the trial treatment as maintenance therapy directly following another acne treatment.

Intervention

This review compares topical preparations for mild/moderate acne described in the NICE CKS or European guidelines. The list was refined by a panel of dermatologists, general practitioners and patients for relevance to clinical practice and patient needs. Treatment regimens available in the UK at any dose, formulation or duration were included. Preparations no longer manufactured or available in the UK, or studies comparing different strengths or dosages of the same preparation were excluded (Box 1).

| Box 1 List of included topical treatments |
|------------------------------------------|
| **Generic name** | **Examples of brand names** |
| Vehicle | |
| Azelaic acid | Skinoren® |
| Adapalene | Differin® |
| Adapalene + BPO | Epiduo® |
| BPO | Acneice® |
| Clindamycin | Dalacin T® |
| Clindamycin + BPO | Duac® |
| Clindamycin + zinc | Zinadclin® |
| Erythromycin + zinc | Zineryt® |
| Isotretinoin + erythromycin | Isotrexin® |
| Tretinoin | |
| Tretinoin + clindamycin | Treclin® |
| Tretinoin + erythromycin | Aknemycin Plus® |

BPO, benzoyl peroxide.

The comparator was placebo/vehicle or any treatment regimen, dose, or duration for the topical treatments listed in Box 1.

The primary outcomes were:

- proportion of participants self-reporting moderate or better global improvement in acne
- proportion of participants withdrawing from trial or cessation of trial medication owing to adverse events.

The secondary outcomes were:

- change in mean total lesion count from baseline as assessed by an investigator
- proportion of participants rated ‘clear’ or ‘almost clear’ on the Investigator’s Global Assessment (IGA) scale of acne severity
- proportion of participants rated as having at least a two-grade improvement from baseline on the IGA scale of acne severity
- change in quality of life from baseline (assessed using a validated instrument such as Skindex-16, Skindex-29 or Cardiff Acne Disability Index)
- reduction in Cutibacterium acnes strains
- total number of adverse events
- participant satisfaction with treatment.

Data collection and data items

A data extraction form was developed in Excel and piloted on two randomly selected papers to ensure consistency. Data available in graph format only were not extracted. Data extraction was performed by one reviewer and checked by a second reviewer.

All outcomes were reported in the medium term, defined as 5–16 weeks (with closest data point to 16 weeks used), with planned sensitivity analysis for short-term (2–4 weeks) and long-term (from 17 weeks to 12 months) outcomes. Trial arms that
reported different strengths or dosages of the same medication were pooled.

**Risk of bias in individual studies**

Risk of bias was assessed using the Cochrane Risk of Bias Tool, covering patient allocation sequence generation, allocation concealment, blinding and selective outcome reporting. 16

**Statistical analyses**

The network geometry has been presented graphically and describes the number of included interventions and the extent to which there are trials comparing different pairs of interventions. 17,18

The network meta-analysis was performed using a frequentist approach with a version of the R package netmeta, implemented in MetaInsight. 19 We anticipated heterogeneity between trials and therefore used random effects models and a common variance approach. 20 Equal heterogeneity across all comparators was assumed and a consistency model was adopted.

For continuous outcomes, the effects were summarized using mean difference if included trials used the same outcome metric or using standardized mean difference if trials reported different outcome metrics. Continuous outcomes were modelled using normal likelihood, and dichotomous outcomes were modelled using binomial likelihood models to produce odds ratios (ORs). A reduced weights approach was used to account for correlation between arms in multiarm...
Ranking of treatments was undertaken using the P-Score approach. We used the design-by-treatment test to evaluate global inconsistency, and node splitting was used to examine inconsistency between direct and indirect effects, with a $P$-value $< 0.05$ considered to be suggestive of conflicting evidence.

Confidence in evidence

The confidence in the evidence across trials was assessed using the Confidence in Network Meta-Analysis (CINEMA) approach and ratings were conducted in the CINEMA app.

CINEMA considers the following six domains: within-study bias, reporting bias, indirectness, imprecision, heterogeneity and incoherence. These domains are rated as ‘no concerns’, ‘some concerns’ or ‘major concerns’, with the exception of reporting bias, which is rated as ‘suspect’ or ‘undetected’. Judgements are then summarized across these six domains as ‘very low’, ‘low’, ‘moderate’ or ‘high’ confidence for each treatment comparison.

Comparisons were considered to have suspected risk of reporting bias if all or most of the comparisons were from industry-funded trials. Indirectness was downgraded for comparisons that were poorly connected in the network. For imprecision, the threshold was set at an odds ratio of 1.5 for binary comparisons and a difference of 10 for lesion counts based on discussion.

Results

Study selection and network structure

We identified 3717 references and, after removing duplicates, 2236 were screened by two reviewers for eligibility. We obtained 329 full texts and identified 133 eligible full texts reporting on 82 trials. An updated search in June 2020
identified a further 23 full texts, nine of which were eligible. We excluded 54 full texts, comprising 5126 participants, because the outcomes of interest could not be extracted. Of the trials identified by the original and updated searches, 81 full texts reporting on 40 trials including a total of 18 089 participants provided outcome data for meta-analysis (Figure 1).25–62

Figure 2 shows network plots for direct evidence between treatments. In all analyses, the main comparator was vehicle. For all outcomes, the most common treatment studied was BPO compared with vehicle, followed by adapalene and adapalene + BPO compared with vehicle. Fewer trials compared clindamycin + tretinoin, erythromycin + zinc or tretinoin, tretinoin alone or azelaic acid with any other treatment.

Trial characteristics and risk of bias

Key trial characteristics and risk of bias are detailed in Table S1 and Figures S1 and S2 (see Supporting Information). The mean sample size was 454 participants (SD 524). The average age was 19.77 years (SD 3.13) and 57.7% of participants were female. Overall, 50% of recruited participants were from North America, 29% were from Europe, 24% were from Asia, 5% were from South America and 3% were from Australia, but the ethnicity of these populations was poorly reported. Pharmaceutical companies sponsored 54% of trials and a further 38% did not report the funder.

Most trials had an unclear risk of bias for at least one domain owing to poor reporting and none had low risk of bias across all domains. While blinding of participants was generally well described in trials that included a vehicle, many trials were unclear in their description of the blinding of trial personnel. All trials were randomized, but the generation of the randomization sequence was poorly described in 30 trials.

Trial results

Table S2 sets out the pooled network analysis results and confidence ratings for all treatment comparisons. Figure 3 sets out all the pooled network comparisons relative to vehicle. Below, we consider the outcomes from the review for which sufficient data were available for network analysis. All treatment rankings and associated probabilities are set out in Tables S3–S6 (see Supporting Information).
Patient Global Assessment of Improvement

The proportion of participants who rated their acne as ‘improved or much improved’ was reported in 11 trials that included 6947 participants. Figure 3 shows that all treatments were significantly more effective than vehicle.

Table 1 sets out direct (no shading) and pooled (in grey) ORs and 95% confidence intervals (CIs) for comparisons. Compared with vehicle, adapalene + BPO had an OR of 3-65 (95% CI 2-58–5-15; moderate confidence) and network comparisons suggest that this treatment was significantly more effective than all other included treatments apart from clindamycin + BPO (OR 1-22; 95% CI 0-81–1-85; low confidence). Clindamycin + BPO was significantly more effective than BPO (OR 1-54; 95% CI 1-14–2-08; low confidence) or clindamycin alone (OR 1-91; 95% CI 1-36–2-68; moderate confidence).

Adverse events

The withdrawal of participants from the trial or participants stopping the trial medication was reported in 35 trials of 16735 participants. Results are set out in Table 2 and the rankings suggest that the lowest odds of withdrawal were in participants who used clindamycin. Clindamycin was associated with significantly lower odds of withdrawal than clindamycin + BPO (OR 2-17; 95% CI 1-25–3-70; very low confidence), BPO (OR 2-38; 95% CI 1-20–4-76; moderate confidence) or adapalene + BPO. The highest odds of withdrawal/discontinuation were for adapalene + BPO (OR 4-35; 95% CI 2-13–9-09; moderate confidence). Participants using adapalene + BPO had an OR of 2-56 (95% CI 1-41–4-76; moderate confidence) compared with adapalene alone, suggesting that the odds of withdrawal/discontinuation were three times higher with combination treatment than adapalene alone. Similarly, participants using adapalene + BPO had an OR of 2-22 (95% CI 0-94–5-26; moderate confidence) compared with those using tretinoin, and an OR of 1-85 (95% CI 1-08–3-13; moderate confidence) compared with those using BPO alone. However, the number of participants who withdrew owing to adverse events was low for all treatments (Table 3).

Total lesion counts

Mean change in total lesion counts was reported in 24 trials of 11717 participants (Table 4). The largest change was observed in those using adapalene + BPO with a difference of 20-96 lesions (95% CI −25-02 to −16-90; moderate confidence) compared with vehicle. Network comparisons suggest significant improvements with adapalene + BPO compared with all other treatments apart from erythromycin + tretinoin, where the CIs were very wide and confidence was very low. Compared with the second ranked treatment, clindamycin + BPO, there were −8-27 (95% CI −13-02 to −3-52; very low confidence) fewer lesions with adapalene + BPO. Clindamycin + BPO and BPO alone were more effective than clindamycin alone with low and moderate confidence, respectively.

Table 1 Direct and pooled comparisons for patient reported global improvement

| Treatment                  | OR (95% CI)          |
|----------------------------|----------------------|
| Adapalene                  | 3.65 (2.58–5.15)     |
| Adapalene + BPO            | 4.35 (2.13–9.09)     |
| Clindamycin + BPO          | 1.22 (0.81–1.85)     |
| Tretinoin                  | 2.17 (1.25–3.70)     |
| BPO                        | 2.38 (1.20–4.76)     |
| Vehicle                    | 1.00 (0.00–1.00)     |

Table 2 Direct and pooled comparisons and rankings for patient reported global improvement

| Treatment                  | OR (95% CI)          |
|----------------------------|----------------------|
| Adapalene                  | 3.65 (2.58–5.15)     |
| Adapalene + BPO            | 4.35 (2.13–9.09)     |
| Clindamycin + BPO          | 1.22 (0.81–1.85)     |
| Tretinoin                  | 2.17 (1.25–3.70)     |
| BPO                        | 2.38 (1.20–4.76)     |
| Vehicle                    | 1.00 (0.00–1.00)     |

Table 3 Adverse events

| Treatment                  | OR (95% CI)          |
|----------------------------|----------------------|
| Adapalene                  | 3.65 (2.58–5.15)     |
| Adapalene + BPO            | 4.35 (2.13–9.09)     |
| Clindamycin + BPO          | 1.22 (0.81–1.85)     |
| Tretinoin                  | 2.17 (1.25–3.70)     |
| BPO                        | 2.38 (1.20–4.76)     |
| Vehicle                    | 1.00 (0.00–1.00)     |

BPO, benzoyl peroxide. Comparisons are presented as odds ratio (95% confidence interval). Light grey shading indicates direct comparisons, dark grey shading indicates pooled comparisons, and black shading indicates treatment rankings.
Table 2: Direct and pooled comparisons and rankings for withdrawal owing to adverse events

|                  | Clindamycin | Clindamycin + zinc | Vehicle | Azelaic acid | Clindamycin + tretinoin | Adapalene | Erythromycin + zinc | Tretinoin | Clindamycin + BPO | BPO | Erythromycin + tretinoin | Adapalene + BPO |
|------------------|-------------|--------------------|---------|--------------|-------------------------|-----------|--------------------|-----------|-------------------|-----|---------------------|------------------|
| Clindamycin     | 1.0 (0.28–5.08) | 0.95 (0.37–2.44) | 1.00 (0.66–1.78) | 0.74 (0.26–2.09) | –                      | –         | 0.92 (0.37–2.27)  | 0.34 (0.18–0.63) | 0.67 (0.19–2.35) | –   | –                   | –                |
| Clindamycin + zinc | 0.85 (0.2–3.57) | 1.79 (0.36–9.09) | –       | –            | –                       | –         | –                  | –         | –                 | –   | –                   | –                |
| Vehicle         | 1.52 (0.79–2.86) | 1.79 (0.36–9.09) | –       | –            | –                       | –         | 0.89 (0.44–1.81)  | 0.48 (0.11–1.91) | 1.71 (0.61–5.03) | 0.64 (0.38–1.07) | 0.31 (0.03–3.10) | 0.37 (0.18–0.76) |
| Azelaic acid    | 1.14 (0.2–8.13)  | 1.54 (0.14–16.67) | 0.87 (0.13–5.88) | 1.11 (0.14–9.09) | –                       | 0.79 (0.05–13.50) | –      | –                  | –   | –                   | –                |
| Clindamycin + tretinoin | 1.45 (0.58–3.57) | 1.72 (0.31–9.09) | 0.96 (0.33–2.78) | 1.11 (0.14–9.09) | –                       | –         | 0.31 (0.03–3.20)  | 0.99 (0.14–7.16) | –      | –                   | –                |
| Adapalene       | 1.69 (0.82–3.57) | 2.12 (0.4–10)     | 1.12 (0.64–2)    | 1.3 (0.19–9.09)   | 1.18 (0.39–3.57)        | 0.48 (0.15–1.50) | 1.00 (0.14–7.32) | 0.73 (0.33–1.58) | –      | –                   | –                |
| Erythromycin + zinc | 2.08 (0.27–16.67) | 2.5 (0.2–33)      | 1.41 (0.12–7.17) | 1.61 (0.1–25)     | 1.45 (0.16–14.29)       | 1.23 (0.13–10)   | 9.94 (0.11–8.31) | 0.97 (0.13–7.09) | –      | –                   | –                |
| Tretinoin       | 2.12 (0.96–9)    | 2.31 (0.46–12.5)  | 1.12 (0.6–2.86)  | 1.32 (0.12–11.11) | 1.17 (0.46–4)            | 1.16 (0.53–2.56) | 0.94 (0.11–8.31) | –      | –                  | –   | –                   | –                |
| Clindamycin + BPO | 2.17 (0.75–3.7)  | 2.56 (0.54–12.5)  | 1.41 (0.76–2.7)  | 1.67 (0.24–11.11) | 1.49 (0.56–4)            | 1.28 (0.62–2.63) | 1.03 (0.14–7.69) | 1.09 (0.48–2.5)  | –      | –                   | –                |
| BPO             | 2.18 (0.72–4.76) | 2.86 (0.56–14.29) | 1.59 (0.98–2.56) | 1.82 (0.27–12.5)  | 1.64 (0.56–5)            | 1.41 (0.77–2.56) | 1.14 (0.14–9.09) | 1.2 (0.53–2.78)  | 1.11 (0.56–2.17) | –      | –                   | –                |
| Erythromycin + tretinoin | 5 (0.44–50)     | 5.88 (0.5–100)    | 1.23 (0.12–33.33) | 1.37 (0.19–100)   | 3.33 (0.27–50)           | 2.86 (0.27–33.33) | 2.33 (0.1–50)   | 2.44 (0.22–25)   | –      | –                   | –                |
| Adapalene + BPO | 4.15 (1.13–9.09) | 5.26 (1.03–25)    | 1.94 (1.69–9.5)  | 3.33 (0.49–25)    | 3.01 (1.9–9)             | 3.56 (1.4–6.26)   | 3.08 (1.6–6.67) | 2.32 (0.94–5.26) | –      | –                   | –                |

BPO, benzoyl peroxide. Comparisons are presented as odds ratio (95% confidence interval). Light grey shading indicates direct comparisons; dark grey shading indicates pooled comparisons; and black shading indicates treatment rankings.
There were 14 trials of 13,342 participants that evaluated improvement in the IGA to ‘clear’ or ‘almost clear’ (Table 5). All treatments were significantly more effective than vehicle apart from tretinoin (OR 0.83, 95% CI 0.46–1.52; low confidence). Adapalene + BPO was significantly more effective than all treatments apart from clindamycin + BPO, with an OR of improvement of 3.83 (95% CI 2.40–6.10; moderate confidence) compared with vehicle. Based on the pooled network estimate, adapalene + BPO was approximately twice as likely to lead to improvement than either BPO or adapalene, with low and moderate confidence, respectively.

Other outcomes and sensitivity analyses
There was insufficient data to undertake meta-analyses or network analyses for quality of life, patient satisfaction, C. acnes resistance and sensitivity analyses of outcomes in the short or long term.

Consistency
There was no evidence of global inconsistency. However, some analyses suggested local inconsistency between direct and indirect comparisons (Tables S7–S10; see Supporting Information). The number of trials where pairs of direct and indirect estimates could be compared was very low and in all instances CIs for estimates of differences were wide, but there was no evidence of systematic differences with respect to potential effect modifiers. Therefore, this apparent inconsistency may represent true differences between direct and indirect effects, with indirect estimates being more precise as they came from a network with larger trials.

Confidence in evidence
The grading of the comparisons with CINeMA (Tables S11–S14; see Supporting Information) showed mainly low to very low confidence ratings. This was due to concerns about reporting bias resulting from the involvement of industry in a large number of small trials and to concerns about within-study bias owing to poor reporting of the randomization and blinding procedures noted above. There were few concerns about transitivity (indirectness). Owing to the strict inclusion criteria, most trials included a homogeneous population of interest. There was also evidence of heterogeneity and imprecision, usually related to the low numbers of trials available for some comparisons in the network.

Discussion
This study compared the most commonly prescribed topical treatments for acne in the UK and found no convincing evidence that topical treatments containing antibiotics are more effective in treating acne than those that do not contain antibiotics. Adapalene + BPO appears to be ranked the most effective treatment on all included outcomes. It is also associated with a higher odds of withdrawal owing to adverse events, but the overall incidence of this outcome was low for all treatments.

Systematic reviews to date have not provided direct comparisons of some of the most commonly prescribed treatments. The recently published Cochrane review of BPO did not show statistically significant differences between BPO and other treatments, however, the study was not able to provide estimates for all other treatment comparisons. Similarly, the Cochrane review including azelaic acid was able to draw on only a limited number of direct trials to quantify differences between treatments.

This network analysis benefits from the additional power of indirect comparisons within the network. However, caution is still needed in interpreting these results. Findings presented here help to highlight gaps where further head-to-head trials are needed. The rankings we have reported are sensitive to inclusion criteria and may change as further evidence emerges. Moreover, the confidence in the evidence was low, with considerable uncertainty remaining about the true effect estimate owing to poor reporting of study methods and the substantial number of trials with industry involvement.

The use of oral antibiotics for acne is high and contributes to antibiotic resistance. Whereas resistance to topical antibiotics tends to be limited to the treated site, oral antibiotics can lead to resistance in commensal flora at all body sites. This study suggests that nonantibiotic treatments are effective as first-line treatment. Further research is needed to explore how these treatments compare with oral antibiotics used alone or in combination with topical treatments.

Although we looked at many outcomes that were important to our patient panel, the study was hampered by poor and inconsistent reporting of trial outcomes. For the participant-reported outcome, only 11 trials were included. The other 30 trials either did not report the outcome of interest (n = 26) or it was reported inconsistently between trials (n = 4). Efforts to harmonize the reporting of outcomes is needed, particularly as the outcomes most commonly reported, such as lesion counts, were not the ones that the patient panel felt were most meaningful.
Table 4 Direct and pooled comparisons and rankings for total lesion counts

|                  | Adapalene | Erythromycin | Azelaic acid | Clindamycin | Adapalene | Tretinoin | BPO | Erythromycin | Clindamycin | Clindamycin | Vehicle |
|-----------------|-----------|--------------|--------------|-------------|-----------|-----------|-----|--------------|-------------|-------------|---------|
| Adapalene+BPO   | 1. Adapalene+BPO | – | – | – | – | – | – | – | – | – | – |
| Adapalene+BPO   | – | – | – | – | – | – | – | – | – | – | – |
| Erythromycin+BPO| 2. Erythromycin+BPO | – | – | – | – | – | – | – | – | – | – |
| Azelaic acid    | 3. Azelaic acid | – | – | – | – | – | – | – | – | – | – |
| Clindamycin+BPO | 4. Clindamycin+BPO | – | – | – | – | – | – | – | – | – | – |
| Adapalene       | 5. Adapalene | – | – | – | – | – | – | – | – | – | – |
| Tretinoin       | 6. Tretinoin | – | – | – | – | – | – | – | – | – | – |
| BPO             | 7. BPO | – | – | – | – | – | – | – | – | – | – |
| Erythromycin+zinc| 8. Erythromycin+zinc | – | – | – | – | – | – | – | – | – | – |
| Clindamycin     | 9. Clindamycin | – | – | – | – | – | – | – | – | – | – |
| Clindamycin+tretinoin | 10. Clindamycin+tretinoin | – | – | – | – | – | – | – | – | – | – |
| Vehicle         | 11. Vehicle | – | – | – | – | – | – | – | – | – | – |

Comparisons are presented as odds ratio (95% confidence interval). Light grey shading indicates direct comparisons; dark grey shading indicates indirect comparisons; and black shading indicates treatment rankings.
For the purposes of this review, we considered total lesion counts. Members of our patient panel felt that this was more meaningful than the distinction between inflammatory and noninflammatory lesions. However, it is possible that the use of this global outcome disguises changes whereby certain phenotypes respond better to specific treatments.

Data on adverse events were particularly poorly reported and we were not able to assess this outcome. This makes it difficult to discuss relative risks and benefits of the different treatments in a meaningful way. Although we have been able to compare the likelihood of participants discontinuing the study, reasons were rarely reported. We were not able to compare adverse events that may concern patients starting a new treatment regimen, such as stinging, itching or peeling.

Blinding was reported in a number of trials and a suitable vehicle was used. However, BPO or retinoids can cause adverse events such as redness or peeling. This might have led to participants or clinicians guessing the allocation. It is hard to quantify the extent to which this may have occurred as it was not reported but, if this did occur, it would lower the overall quality of the reported evidence.

Transitivity is one of the key assumptions of network meta-analysis. In order to achieve a population that was as homogeneous as possible, we excluded full texts where the reported severity of acne was not clearly mild-to-moderate. Within the scope of the review, we did not have the resources to contact all authors of these excluded full texts to obtain clarification. It is possible that limiting the review in this way may have improved homogeneity but introduced a selection bias. Similarly, we did not have the resources to translate articles from other languages. We found 24 titles and abstracts in other languages that may potentially have been eligible. These represent a small proportion of the total titles and abstracts screened, but the inclusion of only English-language full texts may be a source of bias.

The medications in the network analysis account for about two-thirds of prescriptions in the UK in 2018,8 but there are notable gaps, with some treatments being poorly connected to the network and comparisons based on only a single trial. Data on azelaic acid were only available for the lesion count outcome and there were limited trials on combinations including erythromycin or erythromycin alone, which comprise a substantial proportion of topical prescriptions alone or in combination with other treatments.8,63

We were also unable to investigate different concentrations of included treatments in the scope of this review. The pooling of treatment strength into a single comparison may disguise differences in effectiveness of different formulations and strength and further research is needed to explore this topic. Moreover, ethnicity was too poorly reported to explore whether there were any differences with respect to different skin types or skin colours.

Based on evidence mainly graded as low to very low confidence, all topical treatments were more effective than vehicle, and adapalene + BPO was the most effective. Clinicians should evaluate this treatment option in consultation with patients as,
although withdrawal owing to adverse events was uncommon, treatment with adapalene + BPO also appeared to have a slightly higher odds of this outcome. Further work is needed to compare topical treatment with oral antibiotic treatments and to consider which treatments may be most cost-effective.

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Supporting Information

Additional Supporting Information may be found in the online version of this article at the publisher’s website:

Appendix S1 Acne topical treatments network meta-analysis search terms.

Table S1 Study characteristics.

Table S2 Summary of network pooled results and confidence in evidence.

Table S3 Treatment rankings with probability scores for patient-reported global improvements.

Table S4 Treatment rankings with probability scores for withdrawal due to adverse events.

Table S5 Treatment rankings with probability scores for total lesion counts.
Table S6  Treatment rankings with probability scores for Investigator’s Global Assessment.
Table S7  Inconsistency – Patient Global Assessment of Improvement.
Table S8  Inconsistency – withdrawal.
Table S9  Inconsistency – total lesion count.
Table S10 Inconsistency – Investigator’s Global Assessment.
Table S11 CINeMA – Patient Global Assessment of Improvement.
Table S12 CINeMA – withdrawal.
Table S13 CINeMA – total lesion count.
Table S14 CINeMA – Investigator’s Global Assessment.
Figure S1 Study risk of bias assessment.
Figure S2 Risk of bias summary.