Which outcomes are reported in cellulitis trials? Results of a review of outcomes included in cellulitis trials and a patient priority setting survey

Running head: Outcomes in cellulitis treatment and prevention trials

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Conflicts of interest: None known

What is already known about this topic?
- Cellulitis can have a significant impact on the physical and psycho-social wellbeing of patients.
- There is currently no consensus on what outcomes should be measured in cellulitis trials. This makes it difficult to summarise treatment effects in meta-analyses.

What does this study add?
- This review is the first to combine and compare clinical outcomes in randomised controlled trials (RCTs) assessing treatment and prevention of cellulitis with outcome themes deemed important to patients and health care professionals.
- We have highlighted the disparity in clinical outcomes published in these RCTs and the lack of patient reported outcomes.
- Following the COMET initiative, we suggest that homogenous outcomes should be sought and specified for future research in cellulitis.

Summary
Background: There is an emerging need to develop consistent outcomes in clinical trials to allow effective comparison of treatment effects. No systematic review has previously looked at the reporting of outcome measures used in randomised controlled trials (RCTs) on treatment and prevention of cellulitis (erysipelas).

Objectives: The primary aim of this review was to describe the breadth of outcomes reported from RCTs on cellulitis treatment and prevention. The secondary aim was to identify outcome themes from patient and health care professionals’ feedback from a cellulitis priority setting partnership (PSP).

Methods: We conducted a review of all outcome measures used in RCTs from two recent Cochrane reviews. Free text responses from a cellulitis priority setting survey were used to understand the perspectives of patients and healthcare professionals.

Results: Outcomes from 42 RCTs on treatment of cellulitis and six RCTs on prevention of cellulitis were reviewed. Only 28 trials stated their primary outcome. For trials assessing treatment of cellulitis, clinical response to treatment was categorised in 25 different ways. Five of these trials used an outcome that was in accordance with FDA guidance and only four trials incorporated either quality of life or patient satisfaction. For trials assessing prevention of cellulitis, recurrence was the key outcome measure. From the cellulitis PSP, prevention of recurrence, clinical features and long-term disease impact were the most important outcome themes for patients.

Conclusions: We have shown that in cellulitis treatment and prevention research, there is significant heterogeneity in clinical outcomes, inadequate focus on patient-reported outcomes, and a disparity between what is currently measured and what patients and healthcare professionals feel is important. We recommend that future cellulitis treatment trials consider the use of longer-term outcomes to capture recurrence and long-term morbidity, as well as short-term resolution of acute infection.

Keywords: cellulitis, erysipelas, outcomes, core outcome set, priority setting partnership

Introduction
Cellulitis is an acute, potentially serious bacterial infection of the cutaneous and subcutaneous tissue, usually occurring on the lower limb. The most common causative organism is group A streptococcus (streptococcus pyogenes) which causes a spreading area of erythema, tenderness and swelling. The illness can be complicated by ulceration, purulence and progression to systemic disease in severe cases. Erysipelas is characterised by a superficial, well-demarcated area of inflammation, whilst cellulitis often extends to the subcutaneous tissue. It is often difficult to distinguish between them clinically; therefore for the purpose of this review we have used cellulitis as the umbrella term for cellulitis, erysipelas and skin and soft tissue infection, which reflects current practice.

Cellulitis poses a significant health burden to the NHS: during 2014-15 in England, 114,190 patients were treated in hospital for cellulitis with a median length hospital stay of 6 days. In addition to short-term signs and symptoms, the disease can have a significant long-term impact on patient’s physical and mental health. In certain groups of patients there are high levels of recurrence of cellulitis arising from repeated damage to the lymphatic system and the presence of certain risk factors.

Despite the impact of the disease, there are few well-conducted trials looking into the treatment and prevention of cellulitis. It has been reported that clinical assessment of cellulitis many not capture the patient’s experience, and may be unreliable in reflecting real effects for patients.

Defining standard, measurable outcomes is essential for well-designed, robust clinical trials, to provide trial results can be reliably compared. Outcomes need to be meaningful and relevant for service-users, including patients and clinicians, in order to make a long-term difference to future practice.

It has been recognised that many outcomes in clinical trials lack sufficient validation and there is support for development of core outcome sets (COS) for trials to address this issue, as proposed by the COMET initiative. A COS is an agreed set of outcomes that should be measured and reported, as a minimum, in all clinical trials in a specific area of health care. The initial stage in developing a core outcome set is to identify what outcomes are available, and to establish which are most important and relevant for healthcare users.

The Food and Drug Authority (FDA) have attempted to address the lack of consistent outcomes for clinical trials of Acute Bacterial Skin and Soft tissue infections (ABSSIs). In 2013 they updated their guidance on endpoints in such trials which states the primary outcome should be: % reduction in lesion size, measured at 48 to 72 h compared to baseline. However, this recommendation is largely based on historical trial data dating to pre-antibiotic era and has undergone incomplete validation.

There have been no systematic reviews assessing outcome measures used in cellulitis trials. The main aim of this review was to describe all the outcome measures reported in randomised control trials (RCTs) on the treatment and prevention of cellulitis. The secondary aim was to identify outcome themes from patient and health care professionals’ feedback from a cellulitis priority setting partnership (PSP).

Methods
Study 1: Review of outcome measures in RCTs for the treatment and prevention of cellulitis

Search strategy

This review included all RCTs that were included in two Cochrane reviews: *Interventions for the prevention of recurrent erysipelas and cellulitis*<sup>5</sup>, published in June 2017, and the Cochrane update: *Interventions for the treatment of cellulitis and erysipelas* (awaiting publication but obtained from personal communication with authors). Studies that assessed treatment or prevention of cellulitis, erysipelas or skin and soft-tissue infection were included in this review. A search of COMET and PROSPERO was performed to ensure that no existing core outcome initiatives of thematically similar reviews were already registered.

Data regarding treatment complications, adverse events and side effects were not collected as this data is routinely collected in drug trials was not an aim of this review.

The study protocol for this review is available to view on the Centre of Evidence Based Dermatology website<sup>15</sup>

Data extraction and assessment of bias

Reported outcomes were extracted using a standardized template that was piloted prior to use. If the outcome measures were not clearly stated in the methods section of the trial report, but were described in the results section, then these were included.

Details of outcome measures were extracted from each paper by two independent researchers (ES and MP). Any disagreements were resolved by two further independent researchers.

Data extracted were: (i) demographics of trial (author and date of publication) (ii) trial type: treatment/ prevention of cellulitis, drug/ non-drug, (iii) whether primary outcome was stated (iv) total number of outcomes per trial (v) whether outcomes were in line with FDA guidance on skin infections (vi) additional comments on outcomes used in trials e.g. justification of outcomes (vii) Broad outcome domains: clinical, microbiological, biochemical, treatment-related, patient-focused, additional outcomes. For each outcome domain, further definitions were used for the specific outcomes measured e.g. cure/failure/response, length of hospital stay, quality of life. Additionally, for clinical outcome, the specific clinical features that were assessed as part of the outcome were identified e.g. erythema, swelling, warmth (viii) definition of each outcome (ix) how the outcome was assessed (x) scales used (xi) outcome assessor e.g. nurse/clinician/patient (xii) Timing/frequency of assessment.

Storage and analysis of all the data was undertaken by the lead researcher (ES) using Microsoft Excel, 2010, CEBD. Data are presented descriptively, and results reported separately for cellulitis treatment trials and prevention trials.

Study 2: Understanding patient and healthcare professionals’ perspectives

Using free text data collected during a Priority Setting<sup>14</sup> we sought to identify outcomes of importance to patients and healthcare professionals. Responses were submitted by 401 survey participants (171 patients/carers, 217 healthcare professionals, 13 other). Participants were asked the following: “What questions about the diagnosis, treatment or prevention of cellulitis would you like to see answered by research?” Data relevant to outcome measurement were extracted from the free text responses and used to identify key outcome themes using the word repetition technique<sup>16</sup>. 
Results

From the two Cochrane reviews, 48 trials were identified and included in the final analysis. This included 42 RCTs assessing acute treatment of cellulitis and six RCTs assessing prevention of cellulitis. No studies were excluded. The total number of outcomes measured per trial ranged from one to eight outcomes, across six different outcome domains (Figure 1). Only 28 (58%) trials stated the primary outcome. Two papers assessed non-pharmacological treatments for cellulitis; one reviewed the use of vibration therapy and the other an alternative therapy: sodium selenite. Six trials evaluated treatment strategies for skin and soft-tissue infections but did not analyse results for cellulitis separately. However, data from these papers were included in this review, to maximise capture of outcomes.

RCTs assessing treatment of cellulitis

Clinical outcomes

All trials assessed at least one clinical outcome. Of the RCTs assessing treatment of cellulitis, clinical response was categorised according to a range of definitions (Table 1) with the majority of papers classifying response according to “cure”, “failure” or “improvement”. The timing of the clinical assessment was variable. Some recent trials assessed clinical response at an early time-point, 2-3 days after treatment initiation. However, most trials assessed response at a “test-of-cure (TOC)” visit, which ranged from 2 to 42 days after the end of treatment (Table 1). A successful clinical response or “cure” was defined most commonly as either “complete resolution of presenting signs and symptoms”, or “resolution of the infection to an extent that no further antibiotic therapy was required” (Table 2). Four (10%) trials included “absence of recurrence” of infection as part of the definition of a successful clinical response.

There were 10 different signs and symptoms of infection used to describe clinical response across the trials (Figure 2). The most commonly assessed clinical feature was erythema. Of 34 trials that assessed erythema, six (18%) assessed this numerically, measuring the diameter or area of erythema, six (18%) graded the erythema on a severity scale (Figure 3); and 13 (38%) used “clinical assessment/ evaluation” without further detail of the methods used.

Sixteen (38%) of trials assessed at least one biochemical marker as part of clinical response to treatment. Of these, the most commonly assessed markers were white cell count, erythrocyte sedimentation rate and c-reactive protein.

Microbial outcomes

Twenty-four (57%) trials included a microbial outcome, assessed using either blood or wound culture. The majority of trials categorised microbial response according to complete/presumed eradication, persistence/presumed persistence and indeterminate/not evaluable (Table 4). Of the trials assessing a microbial response, the timing and frequency of assessment was either not stated or unclear in 18 (42%) trials.

Treatment-related outcomes

Sixteen (38%) trials assessed a treatment-related outcome. These included: duration of antibiotic treatment (13), number of doses of antibiotic (2) and serum antibiotic concentration (1). One trial used “number of doses of antibiotic until clinical response” as the primary outcome for the study.
Patient-focused outcomes

Four trials included patient-focused outcomes in their methodology. One trial assessed Quality of Life (QoL) using the EuroQol-5 dimensions (EQ-5D) questionnaire. Two assessed patient’s impression of improvement of cellulitis: one using the patient Global Impression of Improvement scale and one using predefined categorical statements: improved, stayed the same, worsened. One paper assessed patient satisfaction with treatment received, assessing each of the parameters: convenience, effectiveness and overall satisfaction on a scale of 0-4. Eleven (26%) trials assessed patient-reported pain (Table 3).

FDA guidance

Five (12%) trials assessing treatment of cellulitis that were published from 2011 onwards had their primary outcome in line with FDA guidance.

RCTs assessing prevention of cellulitis

The primary outcome of trials assessing prevention of cellulitis was either: number of episodes of recurrence or time to recurrence of cellulitis for all six RCTs. Three (50%) of the trials specified the clinical features to be used in assessing an episode of recurrence. Follow-up time varied from 3 months to 3 years. Two (33%) papers explained how patients were monitored during the follow-up period: via routine telephone calls, 3-monthly during the treatment phase and 6 monthly during the follow-up phase. It is worth noting that the PATCH I and II trials intended to collect data on the impact of cellulitis on QoL, however this was abandoned due to technical difficulties around timing of assessments in recurrent episodes.

Additional outcomes

Additional outcomes measured in all trials (assessing both treatment and prevention of cellulitis) included: length of hospital stay (ten trials) and cost-related outcomes (five trials).

Outcome themes from the cellulitis PSP

A total of 846 uncertainties were submitted during the PSP survey. These data were reviewed for outcome themes, and 254 (30%) responses included reference to outcomes of importance to participant. From these responses, 263 outcome themes were identified (some responses contained more than 1 outcome theme), of which 73 (28%) were from patients and 190 (72%) from healthcare professionals. Prevention of recurrence was the most frequently stated outcome theme of importance from both patients and healthcare professionals (Figure 4). “Treatment-focused” outcomes such as: length of acute and prophylactic antibiotic therapy and the use of objective markers to assess treatment response, were also key priorities for healthcare professionals. Assessment of clinical features (erythema, swelling and pain most commonly) and long-term morbidity/mortality were the next most important outcomes for patients. Surprisingly, only 2% of all outcome themes extracted focused on patient-orientated factors such as quality of life and treatment satisfaction.

Discussion
Main findings

1. Variability in clinical outcomes

This review highlights the significant variation of outcomes which are currently used in cellulitis trials, suggesting that efforts to streamline and develop a consensus-driven core outcome set would be valuable.

Clinical response to treatment was categorised in 25 different ways in the 42 treatment trials, and “cure”/ “success” or “resolution”, had 18 different definitions that were measured over a range of timescales, from day 2 to day 35 after the end of treatment period. The majority of trials defined “cure” according to complete or partial resolution of presenting signs and symptoms of infection. However, the specific signs and symptoms of infection assessed, such as erythema, swelling and warmth were often poorly defined. Up to 10 different signs and symptoms of infection were each used by at least one trial to assess treatment response. This reflects the general lack of consensus in outcome assessment for cellulitis.

Some of the more recent trials used “early clinical response” as the primary outcome, defined as: “cessation of lesion spread (measured as the product of the length x width of lesion) 48-72 hr after treatment initiation”. Assessing treatment response at this earlier time-point may increase reliability of the measurement of drug effect, because the outcome is not confounded by natural improvement of disease. Moreover, this endpoint is in line with the 2013 FDA guidance recommending early clinical response as the primary outcome.

2. Inadequate patient orientated outcomes

Patient-focused outcomes such as patient-reported pain, treatment satisfaction and quality of life were rarely reported. We know that cellulitis has a significant impact on physical and psychological health, as well as activities of daily living and quality of life, yet this significance is not reflected in the outcomes from the cellulitis trials published to date. Importantly, the survey results suggest that recurrence and long-term disease impact are important aspects to patients. To date, treatment trials have tended to be of short duration and focus on resolution of symptoms associated with the acute episode. This is not surprising given the cost and methodological complexities associated with longer-term trials, but may require greater consideration when planning future studies.

There were far more treatment trials than prevention trials, which again suggests an imbalance favoring short-term outcomes. More trials that address the prevention of long-term morbidity and recurrence are needed, so that treatment strategies can be developed that have a long-term beneficial impact for patients.

In contrast to patient responses, the survey results revealed the most important outcomes to healthcare professionals were: objective markers of disease response, length of antibiotic treatment and prophylaxis (in addition to recurrence). This is perhaps not surprising, given that healthcare professionals are responsible for both prescribing and objective assessment of treatment effect, rather than a subjective experience of disease.

3. Limited use of microbial outcomes

The use of blood or wound cultures in cellulitis management is debatable. Microbial outcomes are a poor indicator of treatment success as their yield is often low, even in the presence of infection, and can be adversely affected if patients have been pre-
treated with antibiotics. Furthermore, microbial cultures do not correlate well with severity of signs and symptoms or patient experience. This outcome measure is also of limited relevance to patients. However, the increasing incidence of antimicrobial resistance means that other indicators of resistance, such as the incidence and severity of other bacterial infections, may be important to collect.

Strengths and weaknesses

This review summarised the breadth of outcomes reported in published RCTs and analysed these alongside outcomes considered important by patients and healthcare professionals.

Since data on the views of patients and healthcare professionals were originally collected for another purpose (defining the research agenda for cellulitis research), they may not be as reflective of the results as if respondents had been asked specifically about research outcomes. Further qualitative work is required to replicate our findings and to full ascertain the perspectives of all key stakeholders.

The quality of included studies was not assessed as this was not the aim of this review, however many studies were poorly reported, making it difficult at times to establish a detailed understanding of the outcome measures used.

Due to time and resource limitations, included studies were identified from two recently updated Cochrane reviews. These were chosen because they are both up-to-date and sufficiently broad (covering any “treatment” and “prevention” of cellulitis) to gain the breadth of outcome data we aimed to capture. It is possible that more recent RCTs on cellulitis treatment and prevention have since been published.

Generalisability

The included trials were conducted in eight single countries and nine in multiple countries. This review captures a representative snapshot of existing outcomes measures used in cellulitis research. Others have highlighted similar variability in outcome measure instruments used in skin diseases such as eczema, vitiligo and acne.

Conclusion

Outcome measures for cellulitis should not only be consistent, to allow clinical trials to be adequately compared, but should be reflective of real impact in patient’s day to day lives. This review has highlighted that in cellulitis research currently there is a lack of consensus over what should be measured, how that should be measured and over what time-frame. Future research should work towards validating current outcomes with a view to developing a COS for cellulitis. Ideally this would include patient-reported outcomes and a long-term outcome measure, such as recurrence, to account for the chronicity of the disease that is important to many patients. Currently, researchers should seek to ensure outcomes are in line with FDA guidance specific to acute bacterial skin and soft tissue infections.

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45 Hepburn MJ, Dooley DP, Skidmore PJ et al. Comparison of short-course (5 days) and standard (10 days) treatment for uncomplicated cellulitis. *Archives of internal medicine* 2004; **164**: 1669-74.


Noel GJ, Strauss RS, Amsler K et al. Results of a double-blind, randomized trial of cefotibiprole treatment of complicated skin and skin structure infections caused by gram-positive bacteria. *Antimicrobial agents and chemotherapy* 2008; 52: 37-44.


Bucko AD, Hunt BJ, Kidd SL et al. Randomized, double-blind, multicenter comparison of oral cefditoren 200 or 400 mg BID with either cefuroxime 250 mg BID or cefadroxil 500 mg BID for the treatment of uncomplicated skin and skin-structure infections. *Clinical therapeutics* 2002; 24: 1134-47.


Results Tables

Table 1. Categorisation of clinical response and timing of assessment

<table>
<thead>
<tr>
<th>Categories used</th>
<th>Timing of assessment</th>
<th>Number of trials</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>Definition</th>
<th>Number of trials</th>
</tr>
</thead>
</table>
| Cure, failure | Variable<sup>28</sup>  
Day 25-35 (follow-up)<sup>29</sup>  
Day 7-14 after EOT (TOC)<sup>30</sup>  
Day 14-21 after EOT (TOC)<sup>31</sup>  
1 month after EOT<sup>32</sup> | 5 |
| Cure, improvement, failure, indeterminate | Unclear<sup>33</sup>  
Day 7 after EOT<sup>34</sup>  
Day 4-9 after EOT<sup>35</sup>  
EOT (variable)<sup>36</sup> | 4 |
| Cure, failure, indeterminate | Day 1 after EOT, Day 8-15 after EOT (TOC)<sup>37</sup> | 4 |
| Cure, improvement, failure | Day 11 after treatment initiation<sup>41</sup>  
EOT (variable)<sup>42</sup> | 3 |
| Success, failure | 48-72 hrs after treatment initiation, Day 14-15 (EOT)<sup>44</sup> | 3 |
| Relapse/ recurrence | Day 14, Day 28 after treatment initiation<sup>45</sup>  
Day 7 after treatment initiation<sup>46</sup> | 4 |
| Satisfactory, unsatisfactory | Unclear<sup>47, 48, 49</sup> | 3 |
| Cured, not cured | Day 10-14 after treatment initiation (TOC), 2-14 days after EOT (short-term follow-up)<sup>50</sup>  
Day 4-11 after EOT<sup>51</sup>  
Day 12 (EOT), Day 7-10 after EOT, Day 40 (1 month follow-up)<sup>52</sup> | 3 |
| Cure, failure, not evaluable | Day 7-14 after EOT (TOC)<sup>53, 54</sup>  
Day 28-35 after EOT (TOC)<sup>54</sup> | 2 |
| Cure + lesion regression | Day 14-17 (EOT)<sup>59</sup> | 1 |
| Cure, failure, relapse | Day 7-14 after EOT (TOC)<sup>55</sup> | 1 |
| Satisfactory (cure), unsatisfactory (failure/relapse) | Within 24 hrs last dose (EOT), Day 7-14 after EOT<sup>56</sup>  
1 month after EOT<sup>36</sup> | 1 |
| Cure, recurrence | 48-72 hr after treatment initiation<sup>57</sup> | 1 |
| Clinical success (cure, improvement), no improvement, failure | After 24 hrs of treatment<sup>58</sup> | 1 |
| Successful response, stable response, failure | 48-72 hr after treatment initiation<sup>46</sup> | 1 |
| Success (cure/ improvement), failure, indeterminate/ not assessable | Day 11 (EOT), Day 7-14 after EOT<sup>46</sup>  
Day 7-16 after EOT (TOC)<sup>59</sup>  
Day 7-14 after EOT<sup>50</sup> | 1 |
| Cure, improvement | Day 7-14 after EOT<sup>61</sup> | 1 |
| Success, improvement, failure | Day 3 of treatment<sup>61</sup> | 1 |
| Responder, non-responder, indeterminate | 48-72 hrs after treatment initiation<sup>62</sup> | 1 |
| Responder, failure | Day 11 (EOT)<sup>62</sup> | 1 |
| Satisfactory, failure, indeterminate, unevaluable | Unclear<sup>63</sup> | 1 |
| Cure, improvement, failure, relapse | Unclear<sup>64</sup> | 1 |
| Failure, recovery | Daily for 10 days of treatment<sup>65</sup> | 1 |

TOC= test of cure. EOT= End of Treatment

Table 2. Definition of "cure"/"success"/ "satisfactory" response across trials

<table>
<thead>
<tr>
<th>Definition</th>
<th>Number of trials</th>
</tr>
</thead>
<tbody>
<tr>
<td>Resolution of all signs and symptoms of the infection/ improvement to such an extent that no further antimicrobial therapy was necessary</td>
<td>1030,31,37,39,40,33-56,61</td>
</tr>
</tbody>
</table>
Complete resolution of presenting signs and symptoms

Early clinical response: cessation of lesion spread + afebrile ±

>20% reduction in lesion area (length x width of erythema, oedema, induration) ±

i) absence of fever ii) no increase in area of erythema plus absence of fever iii) no increase in area of erythema, no increase in area of swelling and the absence of fever iv) >20% reduction in area of erythema

Resolution of symptoms other than slight residual erythema/oedema

Clinical signs and symptoms resolved with no evidence of active infection at the time treatment was discontinued and no evidence of relapse

(i) Complete resolution of signs and symptoms of soft-tissue infection that was sufficient enough to result in either discontinuation of all antibiotic therapy or switch to the use of oral agents (ii) with no recurrence at the same site in 1 month

All signs and symptoms of the infection that were present before therapy were improved or had resolved and no new signs or symptoms of the infection were present at the post-treatment follow-up

Days until no advancement of the area of cellulitis

Days until no remaining flush/elevation of body temperature >37.8 degrees Celsius on that day

A score of 0 for erythema, oedema and pain and a normal temperature

Time to resolution of erythema, oedema, pain and temperature, without additional antibiotics

Body temperature <37.5 degrees, complete regression of local/general signs of severity and disappearance of the cutaneous plaque

Lesion size, defined as its length times its width, decreased from baseline, temperature was ≤ 37.6°C, fluctuance and localized heat/warmth were absent, and tenderness to palpation and swelling/induration were no worse than mild; for patients with a wound infection, the purulent drainage was to be improved and no worse than mild

Disappearance of warmth and tenderness at the site of infection, with substantial improvement in erythema and oedema, even with mild residual erythema, hyperpigmentation or oedema, not requiring further antibiotic therapy at day 14, and without symptom recurrence at day 28

Signs and symptoms disappeared or improved or signs and symptoms disappeared or improved during therapy but recurred

Improvement in signs and symptoms

Not defined

± Outcomes in line with FDA guidance: cessation of lesion spread (measured as the product of the length x width of lesion) 48-72 hr after treatment initiation

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<table>
<thead>
<tr>
<th>Table 3. Method of assessment of patient-reported pain</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Method of assessment</strong></td>
</tr>
<tr>
<td>0= absent, +/1 = moderate, ++/2 = marked/severe</td>
</tr>
<tr>
<td>100mm VAS scale</td>
</tr>
<tr>
<td>10cm VAS scale±</td>
</tr>
<tr>
<td>----------------</td>
</tr>
<tr>
<td>Faces rating scale±</td>
</tr>
<tr>
<td>Severity rating: none, mild, moderate, severe</td>
</tr>
<tr>
<td>Likert pain scale (0-10)</td>
</tr>
<tr>
<td>McGill pain score</td>
</tr>
<tr>
<td>Brief pain Inventory</td>
</tr>
</tbody>
</table>

0 = "no pain" and the 100 = "worst pain". ±10 point scale. 0 = "no pain" and 10 = "worst pain you can imagine"

<table>
<thead>
<tr>
<th>Table 4. Categorisation of microbial outcomes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Microbial response</td>
</tr>
<tr>
<td>----------------------</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th><strong>Complete eradication</strong></th>
<th>Absence of culturable material/pre-treatment pathogen from the original infection site by the end of therapy</th>
<th>17,50,31,33,35-37,39,41,46,51,53-55,59,63,64</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Not defined</td>
<td>35,60,61</td>
</tr>
<tr>
<td><strong>Presumed eradication</strong></td>
<td>Adequate source specimen not available to culture but patient was assessed as a clinical cure/responded</td>
<td>35,31,37,39,46,53,54,59</td>
</tr>
<tr>
<td></td>
<td>Not defined</td>
<td>35,60,61</td>
</tr>
<tr>
<td><strong>Partial eradication</strong></td>
<td>Absence of some, but not all pre-treatment pathogens at the end of therapy</td>
<td>23,31,37,46,33,39,46,53,55</td>
</tr>
<tr>
<td></td>
<td>Not defined</td>
<td>35,60,61</td>
</tr>
<tr>
<td><strong>Success</strong></td>
<td>Documented/presumed eradication of the baseline pathogen</td>
<td>14,31</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Cure</strong></td>
<td>The pathogen was eliminated with a clearing of inflammation, or the infection recurred with a new pathogen</td>
<td>14,31</td>
</tr>
<tr>
<td></td>
<td>Eradication of causative pathogen or no materials for culture due to clinical success</td>
<td>14,31</td>
</tr>
<tr>
<td><strong>Persistence</strong></td>
<td>Growth of the pre-treatment pathogen in a culture taken at the post-treatment visit</td>
<td>12,50,31,33,36,39,47,49,53,55,59,63,64</td>
</tr>
<tr>
<td></td>
<td>Absence of appropriate culture material in a clinical failure</td>
<td>13</td>
</tr>
<tr>
<td></td>
<td>Eradication of original pathogen with a post-baseline positive culture with a new pathogen requiring treatment</td>
<td>13</td>
</tr>
<tr>
<td></td>
<td>Not defined</td>
<td>34,35,37,46,50,51,54,61</td>
</tr>
<tr>
<td><strong>Presumed persistence</strong></td>
<td>No pathogen identified in a patient who was assessed as clinical failure</td>
<td>35,31,53</td>
</tr>
<tr>
<td></td>
<td>Not defined</td>
<td>46,50,54,61</td>
</tr>
<tr>
<td><strong>Superinfection/ new infection</strong></td>
<td>Emergence of a different pathogen organism during or at the end of therapy</td>
<td>35,33,36</td>
</tr>
<tr>
<td></td>
<td>Pathogen different from that isolated at baseline in presence of signs/symptoms of infection</td>
<td>31,53,59</td>
</tr>
<tr>
<td></td>
<td>Not defined</td>
<td>35,54,61</td>
</tr>
<tr>
<td><strong>Failure</strong></td>
<td>Lesion persisted with significant numbers of the original pathogen, or lesion improved but recurred with a culture that was positive with the same pathogen</td>
<td>13</td>
</tr>
<tr>
<td></td>
<td>Persistence/ presumed persistence of causative pathogen</td>
<td>24,43</td>
</tr>
<tr>
<td><strong>Recurrence</strong></td>
<td>Initial clearance of pre-treatment pathogen, same pathogen reappeared during the follow-up period</td>
<td>13</td>
</tr>
<tr>
<td><strong>Colonisation</strong></td>
<td>Pathogen isolated in absence of clinical signs/ symptoms of infection</td>
<td>13</td>
</tr>
<tr>
<td><strong>Indeterminate/ not evaluable</strong></td>
<td>Unevaluable results/ response not fitting any other category</td>
<td>11,31,34,37,39,40,43,53,54,59,61,64</td>
</tr>
</tbody>
</table>
Figures

Figure 1. Broad outcome domains used across trials

Explanation of outcome categories. Clinical: signs and symptoms of infection assessed as a measure of treatment effect, Microbial: growth or elimination of organisms at the site of infection or in blood cultures as a measure of treatment effect, Biochemical: blood markers measured before and after treatment, Treatment-related: e.g. number of doses/length of antibiotic therapy used as an outcome, Patient-focused: e.g. QoL effects in response to treatment.
Figure 2. Signs and symptoms of infection assessed in clinical response
Figure 3. Method of assessment of erythema

- Clinical assessment/evaluation
- Unclear
- Lesion surface area measured as product of maximal length x width
- Clinical score: 0 = absent, + = moderate, ++ = marked/severe
- Clinical score: none, mild, moderate or severe
- 10cm VAS scale
- Visual assessment after patient had been horizontal for 15 mins
- Demarcated edge marked with an indelible marker pen
- Maximum diameter measured using a standard tape measure

10cm VAS scale: ranging from skin normal colour, no inflammation at 0.0 to skin very red and inflamed at 10.0
Figure 4. Outcomes themes of importance extracted from participant feedback in the Cellulitis Priority-setting Partnership

- Recurrence
- Length of acute antibiotic treatment
- Length of antibiotic prophylaxis
- Objective markers to assess treatment response
- Clinical features
- Long-term morbidity/mortality
- Duration of infection
- Patient-focused outcome (preventing hospital admission/QoL)

Number of responses:

- Number healthcare-professional responders
- Number patient/carer responders