Antibiotics and antiseptics for surgical wounds healing by secondary intention

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ABSTRACT

Background
Following surgery, incisions are usually closed by fixing the edges together with sutures (stitches), staples, adhesives (glue) or clips. This process helps the cut edges heal together and is called 'healing by primary intention'. However, a minority of surgical wounds are not closed in this way. Where the risk of infection is high or there has been significant loss of tissue, wounds may be left open to heal by the growth of new tissue rather than by primary closure; this is known as 'healing by secondary intention'. There is a risk of infection in open wounds, which may impact on wound healing, and antiseptic or antibiotic treatments may be used with the aim of preventing or treating such infections. This review is one of a suite of Cochrane reviews investigating the evidence on antiseptics and antibiotics in different types of wounds. It aims to present current evidence related to the use of antiseptics and antibiotics for surgical wounds healing by secondary intention (SWHSI).

Objectives
To assess the effects of systemic and topical antibiotics, and topical antiseptics for the treatment of surgical wounds healing by secondary intention.

Search methods
In November 2015 we searched: The Cochrane Wounds Specialised Register; The Cochrane Central Register of Controlled Trials (CENTRAL) (The Cochrane Library); Ovid MEDLINE; Ovid MEDLINE (In-Process & Other Non-Indexed Citations); Ovid EMBASE and EBSCO CINAHL. We also searched three clinical trials registries and the references of included studies and relevant systematic reviews. There were no restrictions with respect to language, date of publication or study setting.

Selection criteria
Randomised controlled trials which enrolled adults with a surgical wound healing by secondary intention and assessed treatment with an antiseptic or antibiotic treatment. Studies enrolling people with skin graft donor sites were not included, neither were studies of wounds with a non-surgical origin which had subsequently undergone sharp or surgical debridement or other surgical treatments or wounds within the oral or aural cavities.
Data collection and analysis

Two review authors independently performed study selection, risk of bias assessment and data extraction.

Main results

Eleven studies with a total of 886 participants were included in the review. These evaluated a range of comparisons in a range of surgical wounds healing by secondary intention. In general studies were small and some did not present data or analyses that could be easily interpreted or related to clinical outcomes. These factors reduced the quality of the evidence.

Two comparisons compared different iodine preparations with no antiseptic treatment and found no clear evidence of effects for these treatments. The outcome data available were limited and what evidence there was low quality.

One study compared a zinc oxide mesh dressing with a plain mesh dressing. There was no clear evidence of a difference in time to wound healing between groups. There was some evidence of a difference in measures used to assess wound infection (wound with foul smell and number of participants prescribed antibiotics) which favoured the zinc oxide group. This was low quality evidence.

One study reported that sucralfate cream increased the likelihood of healing open wounds following haemorrhoidectomy compared to a petrolatum cream (RR: 1.50, 95% CI 1.13 to 1.99) over a three week period. This evidence was graded as being of moderate quality. The study also reported lower wound pain scores in the sucralfate group.

There was a reduction in time to healing of open wounds following haemorrhoidectomy when treated with Triclosan post-operatively compared with a standard sodium hypochlorite solution (mean difference -1.70 days, 95% CI -3.41 to 0.01). This was classed as low quality evidence.

There was moderate quality evidence that more open wounds resulting from excision of pyomyositis abscesses healed when treated with a honey-soaked gauze compared with a EUSOL-soaked gauze over three weeks' follow-up (RR: 1.58, 95% CI 1.03 to 2.42). There was also some evidence of a reduction in the mean length of hospital stay in the honey group. Evidence was taken from one small study that only had 43 participants.

There was moderate quality evidence that more Dermacym®-treated post-operative foot wounds in people with diabetes healed compared to those treated with iodine (RR 0.61, 95% CI 0.40 to 0.93). Again estimates came from one small study with 40 participants.

Authors’ conclusions

There is no robust evidence on the relative effectiveness of any antiseptic/antibiotic/anti-bacterial preparation evaluated to date for use on SWHSI. Where some evidence for possible treatment effects was reported, it stemmed from single studies with small participant numbers and was classed as moderate or low quality evidence. This means it is likely or very likely that further research will have an important impact on our confidence in the estimate of effect, and may change this estimate.

Plain Language Summary

Antibiotics and antiseptics for surgical wounds healing by secondary intention

What are surgical wounds healing by secondary intention?

These are surgical wounds which are left open to heal through the growth of new tissue, rather than being closed in the usual way with stitches or other methods which bring the wound edges together. This is usually done when there is a high risk of infection or a large amount of tissue has been lost from the wound. Wounds which are often treated in this way include chronic wounds in the cleft between the buttocks (pilonidal sinuses) and some types of abscesses.

Why use antibiotics and antiseptics to treat surgical wounds healing by secondary intention?

One reason for allowing a wound to heal by secondary intention after surgery is that the risk of infection in that wound is thought to be high. If a wound has already become infected, then antibiotics or antiseptics are used to kill or slow the growth of the microorganisms causing the infection and prevent it from getting worse or spreading. This may also help the wound to heal. Even where wounds are not clearly infected, they usually have populations of micro-organisms present. It is thought that they may heal better if these populations are reduced by antibacterial agents. However, the relationship between infection and micro-organism populations in wounds and wound healing is not very clear.
What we found

In November 2015 we searched for as many studies as possible that both had a randomised controlled design and looked at the use of an antibiotic or antiseptic in participants with surgical wounds healing by secondary intention. We found 11 studies which included a total of 886 participants. These all looked at different comparisons. Several different types of wounds were included. Studies looked at wounds after diabetic foot amputation, pilonidal sinus surgery, treatment of various types of abscess, surgery for haemorrhoids, complications after caesarean section and healing of openings created by operations such as colostomy.

Most studies compared a range of different types of antibacterial treatments to treatments without antibacterial activity, but four compared different antibacterial treatments. Although some of the trials suggested that one treatment may be better than another, this evidence was limited by the size of the studies and the ways they were carried out and reported. All of the studies had low numbers of participants and in some cases these numbers were very small. Many of the studies did not report important information about how they were carried out, so it was difficult to tell whether the results presented were likely to be true. More, better quality, research is needed to find out the effects of antimicrobial treatments on surgical wounds which are healing by secondary intention.

Assessed as up to date November 2015.

BACKGROUND

Description of the condition

Following surgery, incisions are usually closed by fixing the edges together with sutures (stitches), staples, adhesive glue or clips. This process helps the cut edges heal together and is called ‘healing by primary intention’.

However, a minority of surgical wounds are not closed in this way. Where the risk of infection is high or there has been significant loss of tissue, wounds may be left open to heal by the growth of new tissue rather than by primary closure; this is known as ‘healing by secondary intention’. This practice is commonly used following excision of pilonidal sinuses (chronic wounds which arise from hair follicles in the buttock cleft) or perianal or breast abscesses (although a Cochrane review did not find a clear benefit of open healing in pilonidal sinuses (AL-Khamis 2010)).

Wounds may also convert from healing by primary intention to healing by secondary intention when wound closure fails and dehiscence (full or partial separation of wound edges) occurs and cannot be rectified. A recent review using elements of systematic review methodology identified key risk factors for dehiscence as including obesity and wound infection, particularly in abdominal surgeries; however the methodology and quality of the fifteen included studies were highly variable and the review process was not well reported (Sandy-Hodgetts 2013). Dehisced wounds may be allowed to heal fully through secondary intention, or closed surgically after partial healing (delayed healing by primary intention). Delayed healing by primary intention may also be a planned approach.

Prevalence

The data on the incidence and prevalence of SWHSI (both planned SWHSI and those that occur following breakdown of a previously closed wound) are limited. Two published audits from cities in the north of England estimated that SWHSI comprise approximately 28% of all acute wounds receiving wound care provision (Srinivasaiah 2007; Vowden 2009). A more recent UK study which evaluated the prevalence of SWHSI over a two-week period in community, primary and secondary care settings found a prevalence of 0.41 per 1000 population in a total population of 590,585, almost half of which were planned to heal by secondary intention (Ashby 2014 [pers comm] - paper in preparation). An evaluation of the point prevalence of all types of complex wounds in a UK city (population 751,485) found a point prevalence of dehisced surgical wounds of 0.07 per 1000 population (Hall 2014).
The same study reported a prevalence of 0.03 per 1000 population for pilonidal sinus surgical wounds, while the categories of ‘other’ and ‘other surgical wound’ had respective prevalences of 0.11 and 0.14 per 1000 population. Published time-to-healing data for SWHSI are limited. Data from randomised controlled trials (RCTs) show median times to healing of between 54 and 68 days (ranges from 33 to 168 days) depending on aetiology, with pilonidal sinus excisions taking longer to heal than abdominal wounds (Berry 1996; Shackelford 2002; Viciano 2000).

Costs

Without good epidemiological data it is difficult to estimate the explicit costs to healthcare providers of managing SWHSI or the social costs including loss of income experienced by patients in work. The potential time to healing of these wounds, as well as possible requirements for further surgery and other complications including infection, indicate that costs for the broad category of SWHSI are likely to be substantial. The overwhelming majority of patients with SWHSI are treated with dressings of various types (AL-Khamis 2010 [pers comm]). Clearly this accounts for some proportion of the total UK National Health Service (NHS) spend of GBP 116 million on dressings and related wound care products (2008/9 prices) (MeReC 2010). However the majority of costs are likely to stem from the duration of hospital stays (although it is not clear that SWHSI necessitate longer hospital stays in all indications (AL-Khamis 2010)), and subsequent care requirements in the community. Wounds which become infected (develop a surgical site infection (SSI)) also incur increased cost - previously estimated at between GBP 814 and GBP 6626 per patient (Coello 2005; Plowman 2001). The prevalence of SSI was estimated at 5% of all surgeries in the UK and Ireland (Smyth 2008); this is likely to be conservative as many SSIs occur post-discharge from hospital (NICE 2008). These increased costs for SSI generally may apply equally to SWHSI.

Impact on patients and carers

In line with the paucity of epidemiological data, there is little evidence on the impact of SWHSI on patients’ quality of life. A recent qualitative study explored the impact of these wounds on 20 patients and carers. It found that the constant need for treatments and care and the diminished quality of life were comparable to those reported by patients with different types of ulcers; feelings of shock, anguish, frustration and powerlessness were highlighted. The possibility of wound infection was viewed as a constant and insidious threat, representing a major setback to healing (McCaughan 2014 [pers comm]- paper in preparation).

Wound infection

Open wounds such as SWHSI offer an ideal environment for microbial colonisation, and most wounds will contain some microorganisms although this will not necessarily lead to adverse events. An RCT in 54 patients with wounds with delayed healing by primary intention found 353 isolates of 44 different species; many wounds had a bacterial load greater than 1 x 10^7 g^-1 (previously considered a threshold predictive of infection (Robson 1968)) and healed without complication (Moues 2004). The isolates included Staphylococcus aureus and anaerobic species, often associated with infection and delayed healing (Madsen 1996).

Bacteria may be more resistant to bacteriocidal agents because they grow on the surface of a wound forming a film of cells, a ‘biofilm’. Recently opinion has shifted to the view that it is infection with enough or specific types, or both, of pathogenic micro-organisms and possibly resulting biofilms (Percival 2004; Wolcott 2008) that may lead to negative outcomes and potentially delayed healing (Bowler 2003; Davies 2007; Madsen 1996; Tiengove 1996). However, the impact of microbial colonisation on wound healing is not independent of the host response; the ability of the host to provide adequate immune response is likely to be as critical, if not more so, as the specifics of the flora in the wound in determining whether a wound heals.

The document ‘Wound Infection in Clinical Practice - An International Consensus’ (WUWHS 2008) outlines a scenario leading to wound infection where ‘bacteria multiply, healing is disrupted and wound tissues are damaged (local infection)’. The document also notes that ‘Bacteria may produce problems nearby (spreading infection) or cause systemic illness (systemic infection)’. Wound infection has also been seen as the far end of a continuum which starts with sterility (a brief period following surgery), moving to contamination (defined as the presence of microbes but little active growth and no clinical problems), then colonisation (considered as the normal status quo with growth in wound flora being managed by the host immune system and no damage to wound tissues) and finally to critical colonisation and then infection (Kingsley 2004). Critical colonisation is defined as a point between colonisation and infection where the ‘healthy’ balance of wound flora is no longer maintained by the host and the bacterial load or species present in the wound, or both shift away from a so-called safe level. (Kingsley 2004). Others have conceptualised critical colonisation as invasion of the wound surface by micro-organisms (AWMA 2011; Edwards 2004).

The classic clinical signs of infection include local pain, heat, redness, swelling and secretion of pus. The concept of critical colonisation lacks clear diagnostic criteria; it is generally described as being associated with delayed healing in the absence of overt signs of wound infection (Carville 2008; Cutting 2004) possibly with other symptoms such as increased exudate (though less than in infection) and hypergranulation/friable tissue (Cutting 2004; Gardner 2001) although associated evidence is limited. Although there is a widespread view that healing of complex wounds is likely to be retarded by critical colonisation or topical/
local infection, the empirical evidence to support this is extremely limited (Howell-Jones 2005). The literature on the impact of clinical colonisation or topical/local infection in SWHSI is even more limited and caution should be exercised in extrapolating from studies in other types of wounds healing by secondary intention; there are differences in the microbiology even between chronic wound types (Dowd 2008). The Australian Wound Management Association states that ‘the true extent of bacterial impairment of wound healing is unknown’ (AWMA 2011).

**Description of the intervention**

Where an antimicrobial intervention is considered clinically appropriate in the treatment of SWHSI there are two main approaches: an antibiotic may be administered systemically (orally, intravenously or intramuscularly) or a topical antibiotic or antiseptic may be applied. Systemic antibiotics affect the whole body while topical treatments affect only a specific area of the body. Antibiotics are substances that destroy or inhibit the growth of micro-organisms (Macpherson 2004). Systemic antibiotic treatments include groups of drugs which share similar modes of action such as penicillins, cephalosporins, aminoglycosides, macrolides and quinolones. Other antibiotics which do not belong to one of these main groups include clindamycin, metronidazole, trimethoprim and co-trimazole.

Topical antimicrobial interventions include both antibiotics and antiseptics. Antiseptics are thought to prevent the growth of pathogenic micro-organisms without damaging living tissue (Macpherson 2004). Topical applications broadly fall into two types: lotions used for wound irrigation or cleaning, or both, with a brief contact time (unless used as a pack/soak), and products which are in prolonged contact with the wound such as creams, ointments and impregnated dressings.

Agents used primarily for wound irrigation/cleaning are commonly based on povidone iodine, chlorhexidine and peroxide agents. Less commonly used are traditional agents such as gentian violet and hypochlorites. Longer contact creams and ointments include fusidic acid, mupirocin, neomycin sulphate and iodine (often as cadexomer iodine). The British National Formulary categorises antimicrobial dressings as honey-based, iodine-based, silver-based and other, which includes dressings impregnated with agents such as chlorhexidine or peroxides. Recommendations on dressing type are based primarily on the level of wound exudate which determines the dressing substrate as well as the antimicrobial agent (BNF 2014).

**Why it is important to do this review**

Whether systemic antibiotics, topical antibiotics or topical antiseptics can promote healing in SWHSI remains uncertain. An earlier systematic review of antimicrobial agents used for the treatment of all types of chronic wounds (O’Meara 2001) included three small trials assessing topical agents for pilonidal sinus excision wounds (but that were closed rather than left open) and a further three small trials of systemic antibiotics in the same indication. The reviewers were not able to generate definitive conclusions about the use of systemic or topical agents in these wounds because of methodological problems in the primary literature. A Health Technology Assessment (HTA) review from the same year assessed debridement for SWHSI (Lewis 2001). Some of these studies reported the use of chemical debriding agents with antiseptic properties. The review found insufficient evidence to support the use of any particular dressing.

A subsequent Cochrane systematic review (Vermeulen 2004) identified 13 RCTs of dressings and topical treatments for SWHSI, all of which were small and of poor quality. Six of these trials enrolled only patients following surgery for pilonidal sinuses, and five trials enrolled only patients with dehisced abdominal wounds. Interventions assessed included antiseptics such as EUSOL and povidone iodine. There was no evidence that the choice of dressing or topical treatment had any impact on wound healing; no data on clinical infection status were reported. This review is now over 10 years old.

The current review is one of a number of Cochrane reviews investigating the use of antibiotics and antiseptics in the treatment of different types of wounds, each of which updates elements of O’Meara 2001. While there will be some overlap between trials included in this review and in other Cochrane reviews of individual antimicrobial agents in wounds (Jull 2013; Storm-Versloot 2010) or dressings for particular types of SWHSI (Smith 2014), this re-
view will provide a single synthesis of the randomised evidence relating to all systemic and topical antimicrobials for SWHSI. Many options are available to health professionals who are considering using antimicrobial therapy for patients with SWHSI, either as a treatment for, or prophylaxis against, clinical infection. Evidence-based decision-making on the impact of antimicrobial agents on healing of these wounds can be challenging. Key decision problems include whether to use an antibacterial agent instead of standard care and whether different antimicrobial preparations have a differential impact on healing when directly compared.

OBJECTIVES

To assess the effects of systemic and topical antibiotics, and topical antiseptics for the treatment of surgical wounds healing by secondary intention.

METHODS

Criteria for considering studies for this review

Types of studies

We included published and unpublished randomised controlled trials (RCTs), including cluster RCTs, irrespective of language of report. Cross-over trials were only included if they reported outcome data at the end of the first treatment period, prior to cross-over. Quasi-randomised studies were excluded. RCTs reported only as abstracts were included only when available data were sufficient for reasonable data extraction, either from the abstract itself or from the study authors.

Types of participants

People aged at least 18 years with SWHSI. Studies enrolling people with skin graft donor sites were not included, as these sites are superficial wounds which require specific care. Studies of wounds with a non-surgical origin which have subsequently undergone sharp or surgical debridement or other surgical treatments were also excluded. We also excluded studies of wounds where the intention was delayed closure followed by healing with primary intent and studies of wounds which have been created in order to provide an entry or exit point such as percutaneous endoscopic gastrostomy (PEG) wounds or stoma. Wounds within the oral or aural cavities were also excluded. Studies which recruited people with a SWHSI alongside those with other types of wound were not included.

Types of interventions

The primary interventions of interest were topical antiseptic agents or antibiotic (antimicrobial) agents delivered either systemically or topically. We included any RCT in which the use of a topical or systemic antibiotic or a topical antiseptic is the only systematic difference between treatment groups. Systemic antibiotics may have been administered orally or by other routes (e.g. intravenously, intramuscularly). Both intervention and control regimens could consist of antibiotics or antiseptics administered singly or in combination; control regimens might also include placebo, another therapy, standard care or no treatment. Studies which evaluated intervention schedules including other therapies (e.g. pressure relieving devices, dressings) were included provided that these treatments were delivered in a standardised way across the trial arms. We excluded trials in which the presence or absence of a specific antibiotic/antiseptic was not the only systematic difference between trial arms.

We excluded studies evaluating antibiotic or antiseptic regimes that were part of pre-, intra- or post-operative surgical site infection (SSI) prevention practices. We also excluded physical and biological therapies sometimes purported to have incidental antibacterial properties such as heat therapy and larval therapy.

We anticipated that likely interventions would be antiseptic and antibiotic agents, which might include (but not be limited to) the following topical agents which can be available in the form of creams, sprays, ointments and impregnated into different types of dressings: chlorhexidine; povidone iodine; hydrogen peroxide and potassium permanganate; benzoyl peroxide; hypochlorites (e.g. EUSOL); gentian violet; mupirocin and fusidic acid; neomycin sulphate; peroxides; iodine; silver; and honey.

Systemic antibiotics could include: penicillins; cephalosporins; aminoglycosides; macrolides and quinolones; clindamycin; metronidazole; trimethoprim; and co-trimazole.

Types of outcome measures

We list primary and secondary outcome measures below. If a trial was otherwise eligible (correct study design, population and intervention/comparator) but did not report a listed outcome, then we contacted the study authors where possible in order to establish whether a relevant outcome was measured but not reported.

Primary outcomes

The primary efficacy outcome for this review was wound healing. Trialists use a range of different methods of measuring and reporting this outcome. RCTs which reported one or more of the following were considered to provide the most relevant and rigorous measures of wound healing:

- Time to complete wound healing (correctly analysed using survival, time-to-event approaches).
- Proportion of wounds completely healed during follow-up (frequency of complete healing).
We used authors’ definitions of complete wound healing; these were reported where possible. We reported outcome measures at the latest time point available (assumed to be length of follow-up if not specified) and the time point specified in the methods as being of primary interest (if this was different from latest time point available).

Where both the outcomes above were reported, we presented all data in a summary outcome table for reference but focused on reporting time to healing where possible. When time was analysed as a continuous measure (i.e. mean time to healing), but it was not clear whether all wounds healed, we documented the use of the outcome in the study, but did not plan to include the data in any meta-analysis.

We grouped outcome data for wound healing using the following categories; the review authors used their judgement as to whether statistical pooling within these time categories was appropriate.

- **Short term:** 8 weeks
- **Medium term:** > 8 weeks but \( \leq 24 \) weeks
- **Long term:** > 24 weeks

The primary safety outcome for the review was all reported adverse events. Where reported, we extracted data on all serious adverse events and all non-serious adverse events where a clear methodology for the collection of adverse event data was provided. This methodology should make it clear whether events were reported at the participant level or, where multiple events per person were reported, that an appropriate adjustment was made for data clustering.

We also treated wound infection as a primary safety outcome. This included changes in infection status as well as signs or symptoms of clinical infection. Study authors’ definitions of clinical infection were used. We did not include data on bacterial load, diversity, or the presence of individual species where it was not clear how the outcome related to infection.

**Secondary outcomes**

The following secondary outcomes were included:
- **Change (and rate of change) in wound size,** with adjustment for baseline size (we planned to contact study authors to request adjusted means when not presented). When change or rate of change in wound size was reported without adjustment for baseline size, we documented use of the outcome in the study, but did not extract, summarise or use the data in any meta-analysis.
- **Changes in bacterial (antibiotic) resistance.**
- **Health-related quality of life:** quality of life was included where it was reported using a validated scale such as the SF-36 (Ware 1992) or EQ-5D (EuroQoL Group 1990). Ideally reported data were adjusted for the baseline score. We did not include ad hoc measures of quality of life that were unlikely to be validated and would not be common to multiple trials.
- **Mean pain score** (including pain at dressing change) as a continuous outcome using a validated scale such as a visual analogue scale (VAS) or other recognised measurement instrument.
- **Number of wounds closed surgically** (or time to surgical closure).
- **Resource use** (when presented as mean values with standard deviation) including measures of resource use such as number of dressing changes, number of nurse visits, length of hospital stay, and need for other interventions including return to theatre.
- **Costs associated with resource use** (including estimates of cost-effectiveness).

**Search methods for identification of studies**

**Electronic searches**

We searched the following electronic databases to identify reports of relevant clinical trials:
- The Cochrane Wounds Specialised Register (searched 05 November 2015);
- The Cochrane Central Register of Controlled Trials (CENTRAL) (The Cochrane Library) (2015, Issue 10);
- Ovid MEDLINE (1946 to 05 November 2015);
- Ovid MEDLINE (In-Process & Other Non-Indexed Citations) (searched 05 November 2015);
- Ovid EMBASE (1974 to 05 November 2015);
- EBSCO CINAHL Plus (1937 to 05 November 2015).

The search strategies used for CENTRAL, Ovid MEDLINE, Ovid EMBASE and EBSCO CINAHL Plus can be found in Appendix 1. We combined the Ovid MEDLINE search with the Cochrane Highly Sensitive Search Strategy for identifying randomised trials in MEDLINE; sensitivity- and precision-maximising version (2008 revision) (Lefebvre 2011). We combined the EMBASE search with the Ovid EMBASE filter developed by the UK Cochrane Centre (Lefebvre 2011). We combined the CINAHL searches with the trial filters developed by the Scottish Intercollegiate Guidelines Network (SIGN 2011). There were no restrictions of the searches with respect to language, date of publication or study setting.

We also searched the following clinical trials registries:
- ClinicalTrials.gov (http://www.clinicaltrials.gov/);
- WHO International Clinical Trials Registry Platform (ICTRP) (http://apps.who.int/trialsearch/Default.aspx);

**Searching other resources**

We aimed to identify other potentially eligible trials or ancillary publications by searching the reference lists of retrieved included publications by searching the reference lists of retrieved included
trials, as well as relevant systematic reviews, meta-analyses and Health Technology Assessment reports.

Data collection and analysis

Selection of studies
Two review authors independently assessed the titles and abstracts of the citations retrieved by the searches for relevance. After this initial assessment, we obtained full-text copies of all studies considered to be potentially relevant. Two review authors independently checked the full papers for eligibility; we resolved disagreements by discussion and, where required, the input of a third review author. Where the eligibility of a study was unclear we attempted to contact study authors. We recorded all reasons for exclusion of studies for which we had obtained full copies. We completed a Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) flowchart to summarise this process (Liberati 2009). Where studies had been reported in multiple publications/reports, we obtained all publications. Whilst the study was included only once in the review, we extracted data from all reports to ensure all available, relevant data were obtained.

Data extraction and management
We extracted and summarised details of the eligible studies. Where possible we extracted data by treatment group for the pre-specified interventions and outcomes in this review. Two review authors independently extracted data; we resolved discrepancies through discussion or by consultation with a third review author. Where data were missing from reports, we attempted to contact the study authors and request this information.

Outcome data were collected for relevant time points as described in Types of outcome measures.
Where possible we extracted the following data:
- bibliographic data including date of completion/publication;
- country of origin;
- unit of randomisation (participant/wound);
- unit of analysis;
- trial design e.g. parallel, cluster;
- care setting;
- number of participants randomised to each trial arm and number included in final analysis;
- eligibility criteria and key baseline participant data including indication for and type of surgery; location(s) of wounds and whether planned healing by secondary intention;
- details of treatment regimen received by each group;
- duration of treatment;
- details of any co-interventions;
- primary and secondary outcome(s) (with definitions and, where applicable, time points);
- outcome data for primary and secondary outcomes (by group);
- duration of follow-up;
- number of withdrawals (by group) and number of withdrawals (by group) due to adverse events;
- publication status of study;
- source of funding for trial.

Assessment of risk of bias in included studies
Two review authors independently assessed included studies using the Cochrane tool for assessing risk of bias (Higgins 2011a). This tool addresses six specific domains: sequence generation, allocation concealment, blinding, incomplete data, selective outcome reporting and other issues - in this review we recorded issues with unit of analysis, for example where a cluster trial had been undertaken but analysed at the individual level in the study report. We assessed blinding of outcome assessment and completeness of outcome data for each of the review outcomes separately. We have presented our assessment of risk of bias using two 'Risk of bias' summary figures; one which is a summary of bias for each item across all studies, and a second which shows a cross-tabulation of each trial by all of the 'Risk of bias' items. We summarised a study's risk of selection bias, detection bias, attrition bias, reporting bias and other bias. For trials using cluster randomisation, we also planned to consider the risk of bias in relation to: recruitment bias, baseline imbalance, loss of clusters, incorrect analysis and comparability with individually randomised trials (Higgins 2011b) (Appendix 3).

Measures of treatment effect
For dichotomous outcomes, we calculated the risk ratio (RR) with 95% confidence intervals (CIs). For continuous outcome data, we used the mean difference (MD) with 95% CIs for trials that used the same assessment scale. We planned to report time-to-event data (e.g. time to complete wound healing) as hazard ratios (HRs) where possible, in accordance with the methods described in the Cochrane Handbook for Systematic Reviews of Interventions (Deeks 2011). If studies reporting time-to-event data (e.g. time to healing) did not report a HR, then, when feasible, we planned to estimate this using other reported outcomes, such as numbers of events, through the application of available statistical methods (Parmar 1998; Tierney 2007) although we did not implement such an approach here.

Unit of analysis issues
Where studies were randomised at the participant level and outcomes measured at the wound level, for example for wound healing, we treated the participant as the unit of analysis when the
number of wounds assessed appeared to be equal to the number of participants (e.g. one wound per person).

A possible unit of analysis issue we anticipated was that randomisation had been carried out at the participant level with the allocated treatment used on multiple wounds per participant (or perhaps only on some participants) but data were presented and analysed per wound (clustered data).

Where studies included some or all clustered data we planned to report this, noting whether data had been incorrectly treated as independent. We recorded this as part of the risk of bias assessment. We did not plan to undertake further calculation to adjust for clustering as part of this review.

Dealing with missing data

It is common to have data missing from trial reports. Excluding participants from the analysis post randomisation or ignoring participants who are lost to follow-up compromises the randomisation and potentially introduces bias into the trial. If it was thought that study authors might be able to provide some missing data, we planned to contact them; however, it is likely that data will often be missing because of loss to follow-up. In individual studies, when data on the proportion of wounds healed were presented, we assumed that randomly assigned participants not included in an analysis had an unhealed wound at the end of the follow-up period (i.e. they were considered in the denominator but not in the numerator). When a trial did not specify participant group numbers before dropout, we presented only complete case data. For time-to-healing analysis using survival analysis methods, dropouts should be accounted for as censored data. Hence all participants would contribute to the analysis. We acknowledge that such analysis assumes that dropouts are missing at random and there is no pattern of missingness. We have presented data for area change of wound and for all secondary dichotomous outcomes as a complete case analysis.

For continuous secondary outcome measures e.g. length of hospital stay we presented available data from the study reports/study authors and did not impute missing data. Where measures of variance were missing these were calculated (Higgins 2011a) or study authors contacted where possible. Where these measures of variation remained unavailable we excluded the study from any relevant meta-analyses.

Assessment of heterogeneity

Assessment of heterogeneity is a complex, multi-faceted process. Firstly, we planned to consider clinical and methodological heterogeneity: that is the degree to which the included studies varied in terms of participant, intervention, outcome and characteristics such as length of follow-up. We planned to supplement this assessment of clinical and methodological heterogeneity with information regarding statistical heterogeneity - assessed using the Chi^2 test (we considered a significance level of P < 0.10 to indicate statistically significant heterogeneity) in conjunction with the I^2 measure (Higgins 2003). I^2 examines the percentage of total variation across RCTs that is due to heterogeneity rather than chance (Higgins 2003). Very broadly, we considered that I^2 values of 25%, or less, meant a low level of heterogeneity (Higgins 2003), and values of more than 75% indicated very high heterogeneity (Deeks 2011). Where there was evidence of high heterogeneity we planned to explore this further if required: see Data synthesis.

Assessment of reporting biases

Reporting biases arise when the dissemination of research findings is influenced by the nature and direction of results. Publication bias is one of a number of possible causes of ‘small study effects’, that is, a tendency for estimates of the intervention effect to be more beneficial in smaller RCTs. Funnel plots allow a visual assessment of whether small study effects may be present in a meta-analysis. A funnel plot is a simple scatter plot of the intervention effect estimates from individual RCTs against some measure of each trial’s size or precision (Sterne 2011). Funnel plots are only informative when there is a substantial number of studies included in an analysis; we planned to present funnel plots for meta-analyses which included at least 10 RCTs using Review Manager 5.3 (RevMan) (RevMan 2014).

Data synthesis

We combined details of included studies in a narrative review according to the comparison between intervention and comparator, the population and the time point of the outcome measurement. If studies had appeared appropriately similar in terms of wound type and location, intervention type and antibacterial agent, duration of treatment and outcome assessment, we would have considered clinical and methodological heterogeneity and undertaken pooling.

In terms of meta-analytical approach, in the presence of clinical heterogeneity (review author judgement) or evidence of statistical heterogeneity, or both, we planned to use the random-effects model. We planned to use a fixed-effect approach only when clinical heterogeneity was thought to be minimal and statistical heterogeneity was estimated as non-statistically significant for the Chi^2 value and 0% for the I^2 assessment (Kontopantelis 2013). This approach was planned as it is recognised that statistical assessments can miss potentially important between-study heterogeneity in small samples, hence the preference for the more conservative random-effects model (Kontopantelis 2012). Where clinical heterogeneity was thought to be acceptable or of interest we potentially planned to meta-analyse even when statistical heterogeneity was high, with further exploration to interpret the causes behind this heterogeneity and using meta-regression for that purpose, if possible (Thompson 1999; Thompson 2002).
We presented data using forest plots where possible. For time-to-event data, we planned to plot (and, if appropriate, pool) estimates of HRs and 95% CIs as presented in the study reports using the generic inverse variance method in RevMan 5.3 (RevMan 2014). Where time to healing was analysed as a continuous measure but it was not clear if all wounds healed, we documented use of the outcome in the study but data were not summarised or used in any meta-analysis.

'Summary of findings' tables

We planned to present the main results of the review in ‘Summary of findings’ tables. These tables present key information concerning the quality of the evidence, the magnitude of the effects of the interventions examined and the sum of available data for the main outcomes (Schünemann 2011a). The ‘Summary of findings’ tables also include an overall grading of the evidence related to each of the main outcomes using the GRADE (Grades of Recommendation, Assessment, Development and Evaluation) approach. The GRADE approach defines the quality of a body of evidence as the extent to which one can be confident that an estimate of effect or association is close to the true quantity of specific interest. The quality of a body of evidence involves consideration of within-trial risk of bias (methodological quality), directness of evidence, heterogeneity, precision of effect estimates and risk of publication bias (Schünemann 2011b). We planned to present the following outcomes in the ‘Summary of findings’ tables.

- Time to complete wound healing when analysed using appropriate survival analysis methods.
- Proportion of wounds completely healed during the trial period.
- Changes in clinical infection status.

Where data were not pooled we have presented GRADE assessment without a 'Summary of findings’ table - which was the case for all comparisons in this review. In terms of the GRADE assessment, when making decisions for the risk of bias domain we downgraded only when studies had been classed at high risk of bias for one or more domains. We did not downgrade for unclear risk of bias assessments. In assessing the precision of effect estimates we followed GRADE guidance (GRADE 2013) and calculated an optimal information size (OIS) using conventional sample size calculation methods. We used the OIS, along with the size of 95% CIs in terms of whether they spanned estimates of benefit and harm, to assess for downgrading. We calculated the OIS based on GRADE guidance of using a relative risk reduction of between 20% and 30%. The OIS is summarised below but should not be treated as optimal sample sizes for any future research - within a GRADE assessment the OIS is used to assess the stability of CIs rather than being used to assess the appropriateness of a sample size to detect a difference per se.

- Time to wound healing - detect a reduction in time to healing of eight days from 28 days to 21 days (with 100 days recruitment and 100 days follow-up: 80% power; alpha 5%) = 388 participants
- Proportion of wounds healed - detect an increase in wound healing of 75% to 90% (80% power; alpha 5%) = 200 participants
- Changes in clinical infection status - detect a reduction in cases of wound infection from 14% to 10% 80% power; alpha 5%) = 2070 participants

Subgroup analysis and investigation of heterogeneity

If possible, we planned to assess heterogeneity across the following areas: where there was evidence of between-trial heterogeneity we would have conducted a subgroup analysis if feasible.

- Locations of wound types: abdominal wounds and other wound types.
- Planned and unplanned healing by secondary intention.

If possible we planned to perform analyses to explore the influence of risk of bias on effect size. We would have assessed the influence of removing from meta-analyses studies classed as having high and unclear risk of bias. These analyses would have included only studies assessed as having low risk of bias in all key domains, namely, adequate generation of the randomisation sequence, adequate allocation concealment and blinding of outcome assessor for the estimates of treatment effect.

Elements of this Methods section were based on the standard Cochrane Wounds Protocol Template.

RESULTS

Description of studies

Results of the search

The search for this review returned 3928 citations - 10 further citations were found from screening related systematic reviews and bibliographies. After screening we obtained full text 32 citations (relating to 28 studies). Of these studies 11 are included in the review: 13 were excluded. Four studies are awaiting assessment as they required further information before a decision on inclusion could be made and queries with study authors or translators are pending. See Figure 1.
Figure 1. Study flow diagram.

3928 records identified through database searching

10 additional records identified through other sources

3938 records available

3938 records screened

3906 records excluded

32 full-text articles (relating to 28 studies) assessed for eligibility

13 studies excluded, with reasons

4 studies awaiting assessment

11 studies included in qualitative synthesis

0 studies included in quantitative synthesis (meta-analysis)
**Included studies**

Eleven studies (886 participants) are included in this review with each making a different pairwise comparison of treatment - that is no two studies compared the same two treatments. Nine studies investigated use of antiseptic/antibacterial solutions on wounds post-operatively, delivering these via gauze or other inert dressings (Argen 2006; Brehant 2009; Okeniyi 2005; Piaggesi 2010) via a soak (Tosti 2014) or wash (Giannini 2014) or topically (Fernandez 2002; Gupta 2008; Schmidt 1991). Two studies investigated the use of antibiotics - one (Duong 2010) investigated trimethoprim-sulfamethoxazole post-operatively for people following surgery on skin abscesses while one (Khan 2014) investigated the use of antibiotics prior to haemorrhoidectomy surgery.

Included participants had a range of different wounds.

- Wound resulting from excision of pilonidal abscess/disease (Argen 2006; Fernandez 2002).
- Wound resulting from stoma orifice healing by secondary intention (Brehant 2009).
- Wound resulting from incision and drainage of a skin abscess (Duong 2010), pyomyositis abscesses (Okeniyi 2005) or hand abscess (Tosti 2014).
- Wound resulting from haemorrhoidectomy leaving open wounds (Giannini 2014; Gupta 2008; Khan 2014).
- Open post-surgical wounds on the foot/feet of people with diabetes (Piaggesi 2010).
- Wound complications following a caesarean section (c-section) resulting in an open wound (Schmidt 1991).

Studies had generally small sample sizes with a minimum of 37 participants (Fernandez 2002) and a maximum of 162 (Duong 2010) - the median sample size for the 11 included studies was 71 participants. The follow-up period of studies ranged from three weeks to six months and was unclear in one case (Schmidt 1991). Detailed data on each study are found in the Characteristics of included studies tables and an overview of included studies with their extracted outcome data can be found in Table 1.

**Excluded studies**

In total we excluded 13 studies from this review for the following reasons.

- Use of antibiotic/antiseptic was not the only difference between groups (n = 9).
- Not RCT (n = 1).
- No outcome relevant to review (n = 1).
- Not conducted in participants with SWHSI (n = 2).

**Risk of bias in included studies**

Risk of bias assessments are summarised in Figure 2 and Figure 3. Briefly, two studies were considered at low risk of selection bias (Argen 2006; Gupta 2008) the remaining studies were classed at unclear risk of bias. Four studies were classed at low risk of blinding for both staff and patient and blinded outcome assessment (Argen 2006; Duong 2010; Giannini 2014; Gupta 2008), two studies were classed as being at high risk of bias (Schmidt 1991; Tosti 2014); the remaining studies were classed as being at unclear risk of bias for blinding. Seven studies were classed as being at low risk of attrition bias (Argen 2006; Fernandez 2002; Giannini 2014; Gupta 2008, Okeniyi 2005; Piaggesi 2010; Tosti 2014), one study was classed at high risk of bias for attrition bias (Schmidt 1991); the remaining studies were at unclear risk of bias.
Figure 2. Risk of bias graph: review authors’ judgements about each risk of bias item presented as percentages across all included studies.
Figure 3. Risk of bias summary: review authors’ judgements about each risk of bias item for each included study.

<table>
<thead>
<tr>
<th>Study</th>
<th>Random sequence generation (selection bias)</th>
<th>Allocation concealment (selection bias)</th>
<th>Blinding of participants and personnel (performance bias)</th>
<th>Blinding of outcome assessment (detection bias)</th>
<th>Incomplete outcome data (attrition bias)</th>
<th>Selective reporting (reporting bias)</th>
<th>Other bias</th>
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<td>Tosti 2014</td>
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</tbody>
</table>
Effects of interventions

We have presented the effects of interventions by comparison below.

1. Antiseptic treatments compared with non-antimicrobial treatments

Comparison 1: Polyvidone iodine (Povidone iodine)-impregnated mesh compared with alginate mesh (1 trial; 71 participants)

This comparison includes one study which had six weeks' follow-up (Brehant 2009) and assessed treatment for wounds resulting from the closure of stoma orifice by secondary intention.

Primary outcomes

Time to wound healing Mean time-to-healing data presented but not all wounds healed. Data not presented further.

Proportion of wounds healed There was no clear evidence of a difference in number of wounds healed between groups. There were 25/34 (74%) wounds reported as healed in the iodine mesh group compared with 34/37 (92%) in the alginate mesh group: RR 0.80, 95% CI 0.64 to 1.00 (Analysis 1.1). GRADE assessment for proportion of wounds healed: Low quality evidence. Downgraded twice due to imprecision with 95% CIs for risk of wound healing in the iodine group ranging from 36% reduction in risk to 0% reduction, so confidence intervals include a RR of 1 and the estimate comes from a single small study with 71 participants (OIS = 200).

Wound infection There was no clear evidence of a difference in the number of wound infections between groups. In total 2/29 (7%) of participants had a wound infection in the iodine-impregnated group compared with 3/37 (8%) in the alginate mesh group. It was not clear from the report that the infection events occurred in different people but we have assumed this was the case: RR 0.85, 95% CI 0.15 to 4.76 (Analysis 1.2). GRADE assessment wound infection: Low quality evidence. Downgraded twice due to imprecision with 95% CIs for risk of wound infection in the iodine group ranging from a reduction in risk of 85% to an increase risk of 476% so crossing a RR of 1 and the estimate comes from a single small study with 71 participants (OIS = 2070).

Adverse events There were 10 events in the iodine mesh group and 17 events in the alginate mesh group. It was not clear whether each event occurred in a different person or not. We did not analyse the data further.

Secondary outcomes

Resource use (mean number of dressing changes to achieve healing): There was evidence of fewer dressing changes in the alginate mesh group with a mean of 17.9 (SD = 8) dressing changes in the iodine mesh group compared with a mean of 13.2 in the alginate mesh group (SD = 5). Mean difference = 4.70 more dressing changes in the iodine mesh group, 95% CI 1.66 to 7.74 (Analysis 1.3).

Costs associated with resource use (mean cost per participant - included nursing time for dressing changes and cost of dressing): The mean cost of the treatment in the iodine mesh group was EUR 287 compared with EUR 268 in the alginate mesh group. The authors did not present any measures of variance around these estimates, and we have not analysed the data further.

Comparison 2: 10% povidone iodine solution soak compared with no soak (1 trial; 100 participants)

This comparison included one study which had six weeks' follow-up (Tosti 2014) and assessed treatment for wounds resulting from the closure of stoma orifice by secondary intention.

Primary outcomes

None reported.

Secondary outcomes

Resource use Mean length of stay, number of readmissions and number of re-operations are reported in Table 1. It is not clear the number of participants that these events occurred in (thus if data are clustered). We have not analysed the data further here.

Comparison 3: Aloe vera compared with standard care (1 trial; 40 participants)

This comparison contains one study with unclear follow-up (Schmidt 1991) and evaluated treatment of open wounds following a c-section.

Primary outcomes

Time to wound healing The mean time to wound healing in the aloe vera group was 53 days (SD = 24) compared with 83 days (SD = 28) in the standard care group Table 1. There was large loss to follow-up in this group which impacts on knowledge of who...
Adverse events

The report notes that no adverse events were noted in either group, however data collection methods were not clear.

Secondary outcomes

None reported.

Comparison 4: Zinc oxide povidone mesh dressing compared with povidone mesh dressing (classed as placebo by study authors) (1 trial; 64 participants)

This comparison included one study which had 13 weeks' follow-up (Argen 2006) and assessed treatment for wounds resulting from excised pilonidal abscesses.

Primary outcomes

Time to wound healing

Median time to healing was described as 54 days (interquartile range (IQR) 42 days to 71 days) in the zinc oxide mesh group and 62 days (IQR 55 to 82 days) for the placebo mesh group. The study authors presented this as an unadjusted hazard ratio of 1.30, 95% CI 0.77 to 2.19 in the direction of the zinc oxide group. GRADE assessment: Low quality evidence - downgraded twice for imprecision: once as 95% CI were very imprecise ranging from a 23% reduction in hazard of healing in the zinc oxide group to a 219% increased hazard of healing (crossing RR 1) and once as data were from a single study with 64 participants (OIS = 388).

Wound infection

Study authors presented wound infection 'manifest on presence of foul smell' with 1/33 (3%) of participants classed as having an infected wound in the zinc oxide mesh group compared with 8/31 (26%) in the placebo group: RR 0.12, 95% CIs 0.02 to 0.89. Analysis 2.1. GRADE assessment for wound infection: Low quality evidence - downgraded once for indirectness of outcome which is a narrow definition of infection and once due to imprecision with data from a small single study with only 64 participants (OIS = 388).

Secondary outcomes

Pain

We have extracted and presented data in Table 1 for information, however the instrument used to measure pain scores presented (at seven days pre and post mesh removal) was not clearly described. It was also not clear if a high or low score corresponded to more (or less) pain. We have not presented data further here.

Comparison 5: Sucralfate cream compared with petrolatum cream (1 trial; 116 participants)

This comparison contained one trial with four weeks' follow-up (Gupta 2008) that evaluated treatments of participants with open wounds following haemorrhoidectomy.

Primary outcomes

Proportion of wounds healed

At four weeks 45/58 (78%) participants in the sucralate group had a healed wound compared with 30/58 (52%) in the control group RR: 1.50, 95% CI 1.13 to 1.99. Analysis 3.1. GRADE assessment for proportion of wounds healed: Moderate quality evidence. Downgraded once due to imprecision with the estimate coming from a small study (n = 116) which did not meet the OIS (OIS = 200).

Secondary outcomes

Pain

(measured using 10-point VAS scale 0 = no pain to 10 = very severe pain). Data are reported at seven days, 14 days and four weeks. We have presented all data in Table 1. As noted in our methods we have summarised the latest time point here. The mean pain score at four weeks in the sucralate group was 0.2 (SD = 0.3) compared with 1.4 (0.3). Mean difference = -1.20, 95% CI -1.28 to -01.12. Analysis 3.2.

Comparison 6: Trimethoprim-sulfamethoxazole compared with placebo (1 trial; 162 participants)

This comparison includes one study which had 90 days' follow-up (Duong 2010) and assessed treatment for wounds resulting from incision and drainage of a skin abscess.

Primary outcomes

Adverse events

In the trimethoprim group 14/73 (19%) participants had an adverse event attributed to medication compared with 9/76 in the placebo group. RR 2.43, 95% CI 0.99 to 5.98.
(Analysis 4.1). We have presented a complete case analysis. There were missing data in both groups.

Secondary outcomes

Wound recurrence (defined in this study as, "new lesions within 5 cm of original wound at 10 days"). At three months 13/46 (28%) participants had a new lesion in the trimethoprim group compared with 15/52 (29%) in the control. We have presented a complete case analysis. There was missing data in both groups. Mean difference: 0.98, 95% CI 0.52 to 1.84. Analysis 4.2.

Comparison 7: Metronidazole/Ceftriaxone prophylactic antibiotic use compared with standard care (1 trial; 100 participants)

This comparison contained one trial with six weeks’ to eight weeks’ follow-up (Khan 2014) that evaluated open wounds following haemorrhoidectomy.

Primary outcomes

Adverse events We have presented data for low grade fever in Table 1 but it is not clear how many participants had these events so we have not presented data further.

Secondary outcomes

None reported.

3. Antiseptic treatment compared with an alternative antiseptic treatment

Comparison 8: Rhizophora mangle bark extract compared with mercurochrome. (1 trial, 37 participants)

This comparison contained one trial with six weeks’ follow-up (Fernandez 2002) that evaluated open wounds resulting from excision of pilonidal cyst/fistula.

Primary outcomes

No data available for narrative review - see Table 1.

Secondary outcomes

None reported.

Comparison 9: Honey-soaked gauze compared with Edinburgh University Solution of Lime (EUSOL)-soaked gauze (1 trial; 43 participants)

This comparison contained one trial with three weeks’ follow-up (Okenyi 2005) that evaluated participants with open wounds resulting from excision of pyomyositis abscesses.

Primary outcomes

Proportion of wounds healed In the honey group 20/23 (87%) of wounds healed compared to 11/20 (55%) in the EUSOL group: RR: 1.58, 95% CI 1.03 to 2.42 (Analysis 5.1). GRADE assessment for proportion of wounds healed: Moderate quality evidence. Downgraded once due to imprecision with the estimate coming from a small study (n = 116) which did not meet the OIS (OIS = 200).

Adverse events No adverse events reported in either group.

Secondary outcomes

Resource use (mean length of hospital stay). The mean length of hospital stay in the honey group was 16.1 days (SD = 4.16) compared to 18.6 days (SD = 2.14) in the EUSOL group. Mean difference = -2.50 days, 95% CI -4.44 to -0.56.

Comparison 10: Gauze with povidone iodine compared with gauze with Dermacyn® Wound Care (1 trial; 40 participants)

This comparison contained one trial with 24 weeks’ follow-up (or until complete re-epithelialisation) (Piaggesi 2010) that evaluated diabetic participants with an infection on the foot requiring surgery resulting in an open wound.

Primary outcomes

Complete wound healing In the iodine group 11/20 (55%) of wounds healed during the six-month follow-up compared to 18/20 (90%) in the Dermacyn® group: RR: 0.61, 95% CI 0.40 to 0.93 (Analysis 6.1). GRADE assessment for proportion of wounds healed: Moderate quality evidence. Downgraded once due to imprecision with the estimate coming from a small study (n = 116) which did not meet the OIS (OIS = 200).

Adverse events We have presented data for number of events in Table 1 but it is not clear how many participants had these events so we have not presented data further.
Secondary outcomes
None reported.

Comparison 11: Triclosan compared with sodium hypochlorite (1 trial; 113 participants)
This comparison contained one trial with three weeks’ follow-up (Giannini 2014) that evaluated treatment of participants with open wound following haemorrhoidectomy.

Primary outcomes
Time to healing The mean time to healing in the triclosan group was 21.7 days (SD = 3.8) compared with 23.4 days (SD = 5.4) in the control group. We have presented summary data but it is not clear from the report that all wounds healed - and thus not clear that the mean is a valid measure of healing here. Mean difference = -1.70 days, 95% CI -3.41 to 0.01. Analysis 7.1. GRADE assessment for time to wound healing: Low quality evidence. Downgraded once due to imprecision with 95% CIs for a difference in mean time to healing ranging from a 3.4 day reduction in time to healing in the triclosan group to almost 0 days difference; the estimate was from a small study that did not meet the OIS (OIS = .388). The study was also downgraded once for indirectness of outcome as we could not be sure that all wounds had healed and thus that mean time to healing was a valid measure.

Adverse events: The study reported only adverse events related to bleeding/secretions from the wound. At 21 days’ follow-up adverse event scores in the triclosan group were 1.5 (SD 1.7) compared with 2.5 (SD 1.9) in the control group. Mean difference = -1.00, 95% CI -1.66 to -0.34 (Analysis 7.2).

Secondary outcomes
Table 1 reports data at one week, two weeks and three weeks for the following outcomes. Following our methods we have presented only three-week data here.

Pain (measured using 10-point VAS scale where lower score is better): At 21 days’ follow-up pain scores in the triclosan group were 1.4 (SD 1.6) compared with 2.6 (SD 1.8) in the control group. Mean difference = -1.20, 95% CI -1.83 to -0.57 (Analysis 7.3).

Discussion

Summary of main results
This review includes all available RCT evidence for assessments of antiseptic/antibiotic agents in the treatment of populations with surgical wounds healing by secondary intention. Eleven trials with a total of 886 participants were included.

Antiseptic treatments compared with non-anti-microbial treatments - four comparisons
Two comparisons compared iodine preparations with no antiseptic treatment and found no clear evidence of effects on wound healing or wound infection for these treatments. Available outcome data were limited and what evidence there was low quality. One trial reported fewer dressing changes in the povidone iodine group. One study compared a zinc oxide mesh dressing with a plain mesh dressing that was classified as placebo by study authors (Argen 2006). There was no clear evidence of a difference in time to wound healing between groups. There was some evidence of a difference in measures used to assess wound infection (wound with foul smell and number of participants prescribed antibiotics) which favoured the zinc oxide group. This was low quality evidence.

Antibiotic treatments compared with non-microbial treatments - three comparisons
This was classed as low quality evidence.

Antiseptic treatment compared with an alternative antiseptic treatment - four comparisons
There was a reduction in time to healing of open wounds following haemorrhoidectomy when treated with triclosan post-operatively compared with a standard sodium hypochlorite solution (mean difference -1.70 days, 95% CI -3.41 to 0.01) (Giannini 2014). This was classed as low quality evidence.

There was moderate quality evidence that more open wounds resulted from excision of pyomyositis abscesses healed when treated with a honey-soaked gauze compared with a EUSOL-soaked gauze over three weeks’ follow-up (Okenivy 2005). There was also some evidence of a reduction in the mean length of hospital stay in the honey group. Evidence was taken from one small study that only had 43 participants.

There was moderate quality evidence that more Dermacyn®-treated post-operative foot wounds in people with diabetes healed compared to those treated with iodine (Piaggesi 2010). Again estimates came from one small study with 40 participants.

Summary: There is no robust evidence on the relative effectiveness of any anti-microbial preparations evaluated to date for use on SWHSI. Where some evidence for possible treatment effects was reported, it stemmed from single studies with small participant numbers and was classified as moderate or low quality evidence. This means it is likely or very likely that further research will have an...
important impact on our confidence in the estimate of effect, and may change the estimate.

**Overall completeness and applicability of evidence**

Participants in the included trials had wounds which represented many of the types of surgery for which healing by secondary intention is commonly employed, and were therefore likely to be representative of clinical practice in terms of wounds which are intentionally healing in this way. However, only one trial appeared to include participants whose wounds were healing by secondary intention as a consequence of prior complications in healing by primary intention. The applicability of this review to patients with these types of wounds is therefore less clear.

**Quality of the evidence**

There was limited evidence available for this review. There were a small number of relevant studies eligible for inclusion and these evaluated a heterogeneous range of treatments. It is not clear how the treatments evaluated related to common practices in terms of antiseptic and antibiotic use for SWHSI. The studies themselves had small sample sizes. RCTs need to be adequately powered so that they are able to detect treatment effects of a specified size if they exist. This means that sample size calculations should be used to help estimate the optimum number of people recruited to a trial. Evidence from limited numbers of participants or where limited numbers of events occur, or both, can also lead to confidence intervals that are fragile (i.e. not stable). This can be assessed by comparing the data available for a comparison to the optimal information size. All the comparisons in this review were informed by evidence where numbers of participants/events were lower than the calculated optimal information size. It is also important that studies consider the outcome data that are being collected. Several studies presented data that could not easily be analysed here. Where time-to-healing data were presented it was as a mean time to healing. This is only valid if all participants have had the outcome of interest, otherwise the method cannot deal easily with participants who have not had the event, as mean time to event has to be imputed in these cases. Alternatively, people may only include data from those that had the event of interest in the mean calculation which places estimates at risk of bias. The correct approach is the use of survival (time to event methods). It is also important that trials clearly specify how outcome data are collected and defined. For example healing can be defined in a number of ways and this information was limited. We also note that given included studies were assessing antiseptics and antibiotics there was very limited presentation of wound infection data. Finally, studies should also clearly report how they planned to collect adverse event data and how this process was implemented in a standardised way across treatment arms. Overall the quality of outcome data presented was limited by both design and sample size.

Most studies were classed as being at low or unclear risk of bias. However, assessing risk of bias relies on good quality reporting of methods in study papers and this was limited in most of the eleven included studies. Key areas of good practice are the robust generation of a randomisation sequence, robust allocation concealment, and blinded outcome assessment where possible. All this information should be clearly included in the trial report, as trial authors should anticipate the inclusion of their trials in systematic reviews.

**Potential biases in the review process**

This review considered as much evidence as possible. It was not limited by language or publication status. We made attempts to identify studies which were not indexed on mainstream databases. We also contacted authors to try to ensure that limited reporting did not result in studies being excluded from the review. Although all of the included studies were published in English, we identified and assessed studies published in a range of languages. One of the excluded studies was published in Danish (Eldrup 1985) and three of the studies currently awaiting classification were published in other languages (Vasei 2008 (Farsi); Fillmann 2004 and Quilici 1998 (Portuguese)). It is possible that there may be unpublished data that we have been unable to identify. However, whilst we acknowledge the possibility of publication bias despite our attempts to locate unpublished studies, there were too few trials to test for its presence.

**Agreements and disagreements with other studies or reviews**

No other systematic review has specifically focused on antibacterial agents for surgical wounds healing by secondary intention. One previous Cochrane review (Vermeulen 2004) evaluated dressings and topical treatments for surgical wounds healing by secondary intention. Only two of the studies in our review were published before the Vermeulen review (Fernandez 2002; Schmidt 1991); Schmidt 1991 was included by the Vermeulen 2004 review but this did not identify Fernandez 2002. Our review identified and excluded several of the other trials included in Vermeulen's review (Cannavo 1998; Eldrup 1985; Viciano 2000; Walker 1991), principally because the use of an antibacterial agent was not the only systematic difference between trial arms. A review of debridement for surgical wounds healing by secondary intention (Lewis 2001) was completed before the publication of any of the included studies except Schmidt 1991. There is overlap in scope with a more general review of treatments for wounds healing by secondary intention (Bradley 1999) but this was also published before any of the included studies from our review.
except for Schmidt 1991 which is included in all three previous reviews as well as our review. As with the Vermeulen Cochrane review, our review identified and excluded some trials included in the older reviews (Viciano 2000 (included in Lewis 2001); Walker 1991; Williams 1981 (included in both Lewis 2001 and Bradley 1999)) because of additional differences between trial arms. There was only very limited overlap between our review and reviews of specific types of antimicrobials used to treat wounds in general: iodine (Vermeulen 2010); silver (Storm-Versloot 2010), aloe vera, (Dat 2012), and honey (Jull 2013).

A U T H O R S ’ C O N C L U S I O N S

Implications for practice
A comprehensive review of current evidence did not find convincing evidence in favour of the use of any particular antimicrobial treatment for surgical wounds healing by secondary intention for outcomes which matter (including wound healing and infection). Although some possible treatment effects were identified, the quality of the evidence varied from moderate to very low; all of the evidence was subject to some limitations. There was almost no evidence comparing different antimicrobial treatments. All of the evidence related to topical treatments.

Implications for research
Currently there is no consistent evidence of a difference in outcomes between open surgical wounds treated with antimicrobial and non-antimicrobial treatments or with different antimicrobial interventions. Any investment in future primary research on treatment choices for surgical wounds healing by secondary intention must maximise its value to decision-makers. Given the large number of treatment options, the design of future trials should be driven by high priority questions from patients and other decision makers. It is also important for research to ensure that the outcomes that are collected in research studies are those that matter to patients, carers and health professionals. Where trials are conducted, good practice guidelines must be followed in their design, implementation and reporting. Such trials should be adequately powered to detect differences in healing rates, should use appropriate statistical methods for time-to-event analyses and should include adequate follow-up (appropriate to the type of surgery undertaken) for all participants to heal.

A C K N O W L E D G E M E N T S
The authors are grateful to the following peer reviewers for their time and comments: Kurinchi Gurusamy, Elizabeth Matovinovic, Roy Buffery, Gill Worthy, Anna Joseph and AG Rhadika. The authors acknowledge the contribution of Megan Prictor and Denise Mitchell, copy editors. They would also like to thank Zhenmi Liu for her assistance with screening of papers for inclusion, and Ana Luiza C Martimbianco for her translation services.

R E F E R E N C E S

References to studies included in this review

Argen 2006  [published data only]

Brehant 2009  [published data only]

Duong 2010  [published data only]

Fernandez 2002  [published data only]

Giannini 2014  [published data only]

Gupta 2008  [published data only]

Khan 2014  [published data only]
References to studies excluded from this review

Cannavo 1998 [published data only]

Eldrup 1985 [published data only]
Eldrup J. Silastic foam dressing compared with mèche treatment in open treatment following excision of pilonidal cyst [Silastic foam dressing versus mèchebehandeling ved åben behandling efter excision af cystis pilonidalis]. *Ugeskr Laeger* 1985;147(5):408–9.

Goetze 2006 [published data only]

Hien 1988 [published data only]

Moore 2000 [published data only]

Murthy 2012 [published data only]

Nielsen 2012 [published data only]

Sozener 2011 [published data only]

Taylor 2011 [published data only]


Viciano 2000 [published data only]

Walker 1991 [published data only]

Williams 1981 [published data only]

Yang 2013 [published data only]

References to studies awaiting assessment

Fillmann 2004 [published data only]

Mirzabeygi 2011 [published data only]

Quilici 1998 [published data only]

Vasei 2008 [published data only]

Additional references

AL-Khamis 2010

Ashby 2014 [pers comm]

AWMA 2011

Berry 1996

BNF 2014

Bowler 2003

Bradley 1999

Carville 2008

Coello 2005

Cutting 2004

Dat 2012

Davies 2007

Deeks 2011

Dowd 2008

Edwards 2004

EuroQol Group 1990

Gardner 2001

GRADE 2013

Hahn 2005

Hall 2014

Higgins 2003

Higgins 2011a

Higgins 2011b

Howell-Jones 2005

Jull 2013

Kingsley 2004

Kontopantelis 2012

Kontopantelis 2013

Lefebvre 2011

Lewis 2001

Liberati 2009

Macpherson 2004

Madsen 1996

McCaugham 2014 [pers comm]

MeReC 2010

Moues 2004

NICE 2008

O’Meara 2001

Parmar 1998
Parmar MKB, Torri V, Stewart L. Extracting summary statistics to perform meta-analyses of the published

Percival 2004

Plowman 2001

RevMan 2014 [Computer program]

Robson 1968

Sandy-Hodgetts 2013

Schünemann 2011a

Schünemann 2011b

Shackelford 2002

SIGN 2011

Smith 2014

Smyth 2008

Srinivasaiash 2007

Sterne 2011

Storm-Versloot 2010

Thompson 1999

Thompson 2002

Tierney 2007

Trengove 1996

Vermeulen 2004

Vermeulen 2010

Vowden 2009
**Ware 1992**

**Wolcott 2008**

**WUWHS 2008**

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

**Argen 2006**

| Methods | Two-arm RCT  
| Multi-centred, conducted in two hospitals in Denmark  
| Patients were re-examined post-operative day 7, 30, 60, and 90 |
| Participants | 64 participants were randomised  
| Inclusion criteria: 18 years or older, operated on for the first time for pilonidal abscess or chronic pilonidal disease  
| Exclusion criteria: hypersensitivity to zinc, dementia, insufficient in Danish, or pregnant/lactating |
| Interventions | Group A: Zinc oxide (3%) polyvinylpyrrolidone (povidone) bound to a 50 cm-long cotton mesh of 2 (0.5 g) or 5 (1.25 g) cm widths. Median zinc oxide concentration of mesh reported as 33 mg/g (interquartile range 33 mg/g to 35 mg/g) (n = 33)  
| Group B: Polyvinylpyrrolidone (povidone) bound to a 50 cm-long cotton fine mesh of 2 (0.5 g) or 5 (1.25 g) cm widths (n = 31)  
| The dry meshes were applied to the wounds in at least four layers. Wounds were treated daily during the first seven post-operative days with either zinc oxide or placebo meshes and thereafter every second day  
| Perioperative prophylactic antibiotics were not given. The necessity of systemic antibiotic treatment post-operatively was judged on clinical signs of wound infection |
| Outcomes | Primary outcomes:  
| • Time to complete wound closure (defined as complete coverage of the wound with visible epithelium)  
| • Adverse events (Any serious or non-serious adverse events occurring until wound closure or day 90 were reported to the co-ordinating investigator. The investigator judged the adverse event as definitely not, improbable, probable, or most probably related to the intervention).  
| • Wound infection (defined by smell; the number of people given antibiotics post-operatively was also reported)  
| Secondary outcomes:  
| • Pain measured at day 7 post-op as pain intensity. The scale used was not reported although a VAS score measured as mm is mentioned. It is not clear if higher or low scores relate to more pain |
| Notes | Funding source: The Pharmacy Foundation of 1991, Denmark, and the Danish Medical Research Council (22-02- 0287)  
| Missing data were replaced by the mean value of both groups of respective covariates in the Cox model |

**Risk of bias**

| Bias | Authors' judgement | Support for judgement |
| Random sequence generation (selection bias) | Low risk | Quote: The patient allocation sequence was computer-generated 1:1 in variable block sizes of four or six stratified for center. Comment: Adequate. |
| Allocation concealment (selection bias) | Low risk | Quotes: Allocation concealment was performed using centrally packaged, consecutively numbered, identical packages containing zinc oxide or placebo meshes. The investigators were asked to use the next available number when a new patient entered the trial. Randomization codes were kept confidential until final assessments of the patients, data entry, statistical analyses, and main conclusions were completed. Comment: Adequate. |
| Blinding of participants and personnel (performance bias) | Low risk | Quotes: The zinc and placebo meshes were manufactured in Class 100,000 facilities, sterile, and indistinguishable in color, texture, and smell. In this randomized, double-blind, placebo-controlled multicentre trial, the patients were centrally randomized to receive either topical zinc oxide meshes or placebo meshes during pilonidal wound healing. Comment: Blinding occurred. |
| Blinding of outcome assessment (detection bias) | Low risk | **Outcome - wound healing** Quote: The wound was evaluated clinically with respect to complete wound closure by assessors blinded to treatment. Comment: Adequate. **Outcome - wound infection** Quote: In the present trial, we took all precautions to avoid bias including a priori sample-size calculation, adequate allocation concealment, placebo-controlled, blinded outcome assessment, a predefined primary outcome measure, and intention-to-treat analyses. Comment: Suggests that other outcomes were also assessed in blinded way. |
| Incomplete outcome data (attrition bias) | Low risk | Quotes: A 17-year-old female patient was included by mistake, but has been retained. |
in the trial for analyses. Three patients in the zinc oxide group and two patients in the placebo group withdrew from the study within 2 weeks after randomization for unknown reasons. Participants lost to follow-up were censored at the date of last contact to the investigators.

Comments: As close to ITT as possible. Limited loss of data.

<table>
<thead>
<tr>
<th>Selective reporting (reporting bias)</th>
<th>Low risk</th>
<th>None - protocol not obtained</th>
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</thead>
<tbody>
<tr>
<td>Other bias</td>
<td>Low risk</td>
<td>None noted</td>
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</tbody>
</table>

**Brehant 2009**

**Methods**
- Two-arm RCT
- Multi-centred, conducted in four hospitals in France
  - The patients were examined every day while they were in hospital. The volumetric measurement of the SO was carried out every 7 days for 1 month on days 0, 7, 14, 21, and 28. All the patients were reviewed during the sixth post-operative week for the end-of-study visit.

**Participants**
- 71 participants randomised
  - Inclusion criteria: patients having re-establishment of digestive continuity with closure of the stoma orifice by secondary intention
  - Exclusion criteria: patients aged 17 years or younger, emergency surgery, serious allergy to iodine or to one of the treatments used, a stoma affecting a digestive segment other than the small intestine or the colon, patient with dysthyroidia, patients requiring additional protection (pregnant or breast-feeding woman, subject under wardship or guardianship), and informed consent not signed.

**Interventions**
- **Group A**: polyvidone iodine-impregnated mesh (n = 34)
- **Group B**: alginate mesh (n = 37)
  - No manufacturer details for meshes given in paper. The length of the dressings was based on the volume of the whole cavity. No other local drainage was used. Dressings were changed every two days, with only one dressing whenever possible.
  - No co-interventions listed

**Outcomes**
- **Primary outcomes**:
  - Proportion of wounds healed (mean time to healing data was also reported but this has not been extracted as not all wounds healed).
  - Complications (a range of adverse events reported)
- **Secondary outcomes**:
  - Change in size (percentage filling of the SO cavity at day 28)
  - Surgical site infection (CDC definition)
  - Resource use (number of dressing change)
### Brehant 2009 (Continued)

<table>
<thead>
<tr>
<th>Notes</th>
<th>Funding source: not reported</th>
</tr>
</thead>
</table>

#### Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
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<th>Support for judgement</th>
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</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Unclear risk</td>
<td>No information reported</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Unclear risk</td>
<td>No information reported</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias)</td>
<td>Unclear risk</td>
<td>No information reported</td>
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<tr>
<td></td>
<td>All outcomes</td>
<td></td>
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<tr>
<td>Blinding of outcome assessment (detection bias)</td>
<td>Unclear risk</td>
<td>No information on who collected any outcome data reported</td>
</tr>
<tr>
<td></td>
<td>All outcomes</td>
<td></td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias)</td>
<td>Unclear risk</td>
<td>Quote: &quot;Five patients (all in the polyvidone iodine mesh group) were excluded from the analysis either because they refused to follow the protocol (n = 2), or because of the occurrence of an anastomotic fistula exteriorized through the old stoma orifice (n = 3). The statistical analysis therefore involved 66 patients&quot; Comment: Five participants in one group were excluded completely from analysis. Not clear what implications this has on attrition bias</td>
</tr>
<tr>
<td></td>
<td>All outcomes</td>
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<tr>
<td>Selective reporting (reporting bias)</td>
<td>Low risk</td>
<td>None noted - protocol not obtained</td>
</tr>
<tr>
<td>Other bias</td>
<td>Low risk</td>
<td>None noted</td>
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</tbody>
</table>

#### Duong 2010

| Methods | Two-arm RCT  
Undertaken in the emergency department of one hospital in the USA  
Follow-up took place at 10 to 14 days and 90 days |
| --- | --- |
| Participants | Participants were undergoing incision and drainage of a skin abscess  
161 Participants were randomised  
Inclusion criteria: aged 3 months to 18 years with a skin abscess; non-toxic with a temperature less than 38.4°C (101.1°F). Diagnostic criteria for skin abscess included the presence of all of the following features: (1) acute onset within 1 week, (2) fluctuance, (3) erythema, (4) induration, and (5) tenderness, with or without purulent drainage; |
patients or parent/guardian able to give informed consent
Exclusion criteria: known chronic health problems, such as diabetes; receiving immuno- suppressive medications, such as oral steroids for asthma; recent (within the last week) or current antibiotic usage; contraindication to trimethoprim-sulfamethoxazole, did not require surgery - for example for more superficial skin infections such as folliculitis
The surgery performed involved cleansing of the skin overlying all skin abscesses with 10% povidone iodine solution and then incision with a no. 11 blade, probing for loculations, and irrigation with normal saline solution. Decisions concerning the need for procedural sedation, the incision size, and 24-hour wound packing were physician dependent

| Interventions | Group A: trimethoprim-sulfamethoxazole (10 mg to 12 mg trimethoprim/kg/day divided into 2 doses, with a maximum dose of 160 mg trimethoprim/dose). Given for 10 days post-operatively. Only the liquid formulation of the antibiotic was used. The concentration of the antibiotic solution was 200 mg sulfamethoxazole/40 mg trimethoprim per 5 ml (n = 77)
Group B: placebo given for 10 days post-op (n = 84)
Patients and their parents were asked to monitor for any adverse effects of the medication and to call if they had any questions or concerns. At home, they were instructed to remove and discard the gauze packing, if used, 24 hours after it was placed in situ and to perform warm water soaks at least twice a day per standard of care. They were instructed to keep the wound clean and covered by a layer of gauze with taping around the edges and to avoid using topical antibiotic ointment/cream, hydrogen peroxide, alcohol, or Betadine to decrease the chance of confounding factors |

| Outcomes | Primary outcomes:
- Adverse events: clinical failure at day 10 (defined as presence of erythema, warmth, induration, fluctuance, tenderness, and drainage or worsening of symptoms requiring drainage, change in medication or hospital admission); adverse event attributed to medication
Secondary outcomes:
- Wound recurrence (defined as new lesion within 5 cm of original abscess site - new lesions could be folliculitis, furuncles, carbuncles, or abscesses. |

| Notes | Funding source: no clear information
Study was a non-inferiority study |

<table>
<thead>
<tr>
<th>Risk of bias</th>
<th>Bias</th>
<th>Authors’ judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Low risk</td>
<td>Quote: With a computer randomization program, subjects were then randomized in permuted blocks of 50 to receive a 10-day course of placebo or trimethoprim-sulfamethoxazole. The placebo consisted of a Maalox and tonic water combination that resembled the antibiotic in color, texture, and taste</td>
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</table>
### Duong 2010 (Continued)

<table>
<thead>
<tr>
<th>Source</th>
<th>Comments</th>
<th>Bias Factor</th>
<th>Risk</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Low risk</td>
<td></td>
<td>Quote: The medications were prepared, stored, and dispensed by the inpatient pharmacist who also generated the randomization sequence and assigned the participants to their groups. Comment: Unclear if the pharmacist knew what was being dispensed when allocating but assumed this was likely the case.</td>
<td></td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias)</td>
<td>Low risk</td>
<td>All outcomes</td>
<td>Quote: The patient, parents, and clinician who assessed the clinical outcome were blinded to group assignment. Comment: Adequate.</td>
<td></td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias)</td>
<td>Low risk</td>
<td>All outcomes</td>
<td>Quote: The patient, parents, and clinician who assessed the clinical outcome were blinded to group assignment. Comment: Adequate.</td>
<td></td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias)</td>
<td>Unclear risk</td>
<td>All outcomes</td>
<td>Quote: One hundred sixty-one subjects were enrolled, with 12 lost to follow-up. The patients who were lost to follow-up did not differ from the rest of study participants in terms of demographics or clinical presentation. Comment: Some lost to follow-up - but relatively small number - classed as low risk for adverse event data. There is more missing data for the wound recurrence outcome. Classed as unclear for this and unclear overall.</td>
<td></td>
</tr>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>Unclear risk</td>
<td></td>
<td>None noted - protocol not obtained</td>
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</tr>
<tr>
<td>Other bias</td>
<td>Unclear risk</td>
<td></td>
<td>None noted</td>
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</table>

### Fernandez 2002

<table>
<thead>
<tr>
<th>Source</th>
<th>Methods</th>
<th>Participants</th>
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<tbody>
<tr>
<td></td>
<td>Three-arm RCT</td>
<td>37 participants randomised</td>
</tr>
<tr>
<td></td>
<td>Undertaken in one centre in Cuba</td>
<td>Inclusion criteria: open wounds from surgical intervention of pilonidal cyst or pilonidal fistula</td>
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<tr>
<td></td>
<td>Duration of follow-up 6 weeks</td>
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</table>
### Interventions

<p>| | | |</p>
<table>
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<tr>
<th></th>
<th></th>
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<tbody>
<tr>
<td><strong>Group A</strong></td>
<td>Rhizophora mangle bark extract. Applied once a day (n = 12)</td>
<td></td>
</tr>
<tr>
<td><strong>Group B</strong></td>
<td>Rhizophora mangle bark extract. Applied twice a day (10 h to 12 h interval) (n = 12)</td>
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</tr>
<tr>
<td><strong>Group C</strong></td>
<td>Mercurochrome. Applied twice a day (10 h to 12 h interval) (n = 13)</td>
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</tbody>
</table>

### Outcomes

<table>
<thead>
<tr>
<th><strong>Primary outcomes:</strong></th>
<th></th>
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<tbody>
<tr>
<td>None</td>
<td></td>
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</tbody>
</table>

**Secondary outcomes:**

- Change in wound size (limited data for extraction)
- Adverse events (but methods not defined - very unclear and data not extracted)

### Notes

**Funding source:** not reported

### Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors’ judgement</th>
<th>Support for judgement</th>
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<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Low risk</td>
<td>Quote: The trial was single blind, random-ized and comparative. Three groups of patients were randomly created and assigned. Comment: Methodology unclearly reported in paper - authors of another review contacted the study authors and were informed that randomisation was performed with a random generated list.</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Unclear risk</td>
<td>Not details - methodology unclear</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias) All outcomes</td>
<td>Unclear risk</td>
<td><strong>All outcomes</strong> Quote: The trial was single blind, random-ized and comparative.. Comment: Unclear who was blinded from paper. Authors of another review contacted the study author and were able to establish that the patients were blinded.</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias) All outcomes</td>
<td>Unclear risk</td>
<td><strong>All outcomes</strong> Quote: The trial was single blind, random-ized and comparative.. Comment: Unclear whether blinded outcome assessment was conducted.</td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias) All outcomes</td>
<td>Low risk</td>
<td>Comment: Appears that all those randomised were included in analyses.</td>
</tr>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>Low risk</td>
<td>None noted - protocol not obtained</td>
</tr>
</tbody>
</table>
Giannini 2014

Methods

Two-arm RCT
Multi-centred study undertaken in Italy
Three weeks’ follow-up

Participants

113 participants randomised
Inclusion criteria: undergoing Milligan-Morgan haemorrhoidectomy for grade III or IV haemorrhoids
Exclusion criteria: patients with cancer, HIV, insulin-dependent diabetes, severe liver disease, Crohn’s disease, anal abscess/fistula, anal fissure, thrombosed haemorrhoids, anticoagulant treatment and pregnancy

Interventions

Group A: Triclosan (Proctocid, Uniderm Farmaceutici, Srl, Rome, Italy) (n = 55)
Group B: Sodium hypochlorite solution 1.15g/100ml) (n = 58)
In each group treatment consisted of anal wound washes using 10 ml of solutions diluted in a basin (no further details) three times a day and after defecation
Other drugs were discontinued during the treatment except stool softeners and painkillers. Painkiller tablets were administered as required

Outcomes

Primary outcomes:
• Wound healing (time to healing - healing not defined)
• Adverse events (bleeding/secretions - measured using a 10-point VAS scale - high and low not defined but seems lower scores refer to less bleeding/secretions)

Secondary outcomes:
• Pain (measured using a 10-point VAS scale - high and low not defined but seems lower scores refer to less pain)

Notes

Funding source: not reported but authors stated that there were no financial conflicts of interest

Risk of bias

Bias | Authors’ judgement | Support for judgement |
--- | --- | --- |
Random sequence generation (selection bias) | Unclear risk | Quote: Each centre participating the study [sic] received the randomisation code using 40 progressively numbered closed envelopes...the randomisation list was generated using random permuted blocks
Comment: No detail about method of sequence generation |
Allocation concealment (selection bias) | Low risk | Quote: See above and treatments were identical and participants and doctors blinded - assumes that those allocating were
## Giannini 2014 (Continued)

<table>
<thead>
<tr>
<th>Bias</th>
<th>Risk</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blinding of participants and personnel (performance bias)</td>
<td>Low</td>
<td>Adequate</td>
</tr>
<tr>
<td>All outcomes</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Quote:</strong> Triclosan and the hypochlorite solution were made indistinguishable (colour, texture fragrance) and were stored in similar bottles with the randomisation code. blinded to physician and patient <strong>Comment:</strong> Adequate</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias)</td>
<td>Low</td>
<td></td>
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<tr>
<td>All outcomes</td>
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<tr>
<td><strong>All outcomes</strong> Considered blinded. For wound healing it is noted that a photograph was sent to the co-ordinating centre. It is not explicit that the assessment was blinded but given the study methods we have inferred this</td>
<td></td>
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</tr>
<tr>
<td>Incomplete outcome data (attrition bias)</td>
<td>Low</td>
<td></td>
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<tr>
<td>All outcomes</td>
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</tr>
<tr>
<td><strong>Quote:</strong> There was one case of dropout in the Triclosan group <strong>Comment:</strong> Note one drop-out and flow chart presented suggests no data excluded from analysis. Considered low risk</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>Low</td>
<td></td>
</tr>
<tr>
<td><strong>No evidence from paper - protocol not obtained</strong></td>
<td></td>
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</tr>
<tr>
<td>Other bias</td>
<td>Low</td>
<td></td>
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<tr>
<td><strong>None noted</strong></td>
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</table>

## Gupta 2008

### Methods

- Two-arm RCT
- Undertaken in a single hospital in India
- Duration of follow-up 4 weeks

### Participants

- 116 participants undergoing haemorrhoidectomy resulting in an open wound
- Inclusion criteria: symptomatic and prolapsing haemorrhoids Grades III or IV and able to provide informed consent
- Exclusion criteria: associated fistula or fissure-in-ano, inflammatory bowel disease, dermatitis, or proctitis, patient unable to complete study documentation
- A single surgeon performed all the procedures, with the patient in the lithotomy position.
- Patients underwent a standard Milligan-Morgan haemorrhoidectomy

### Interventions

- **Group A:** Sucralfate cream - applied to the wounds three times daily (n = 58)
- **Group B:** Placebo (petrolatum cream) - applied to the wounds three times daily (n = 58)
- All patients were supplied with standard analgesic tablets containing a combination of tramadol hydrochloride 37.5 mg and acetaminophen 500 mg. The patients also received tablets containing 250 mg of metronidazole to be taken twice daily for seven days and 30 ml of lactulose
### Outcomes

**Primary outcomes:**
- Time to healing (time to 100% complete wound healing)
- Number of wounds completely healed (wound healing was defined as complete epithelial covering as observed by physical examination (per-rectal examination and anoscopy)

**Secondary outcomes:**
- Post operative pain (measured using 0 to 10 VAS scale with 0 = no pain)

### Notes

**Funding source:** not reported

### Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors’ judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Low risk</td>
<td>Quote: In this double-blind study, patients were prospectively randomized by computer-based sequential method into one of the two groups Comment: Adequate method</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Low risk</td>
<td>Quote: Randomization was performed at the completion of surgery; both creams looked the same and were impossible to distinguish Comment: Adequate method</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias) All outcomes</td>
<td>Low risk</td>
<td>See above quote and comment</td>
</tr>
</tbody>
</table>
| Blinding of outcome assessment (detection bias) All outcomes | Low risk           | **Pain assessment**

Quote: An independent observer, blinded to the postoperative prescription, collected and assessed the data

**Healing**

Quote: At the end of four weeks postoperatively, two surgeons, independent of the study group, examined the healing of patients' wounds

Comment: Adequate

**Incomplete outcome data (attrition bias) All outcomes**

| Low risk | Quote: It was found that two patients from the sucralfate group did not use the cream as advised. These patients reportedly used some other cream in addition to the sucralfate cream. Two patients from each group were lost to follow-up, |

Antibiotics and antiseptics for surgical wounds healing by secondary intention (Review)

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therefore, 110 patients (sucralfate n = 54, placebo n = 56) were available for final analysis. Comment: six lost to follow-up. Two participants appear to have been excluded necessarily. Remains limited loss of data though - classed as low risk

Selective reporting (reporting bias) | Low risk | None noted - protocol not obtained

Other bias | Low risk | None noted

Khan 2014

Methods
Two-arm RCT
Multi-centred study undertaken in Saudi Arabia and Pakistan
Duration of follow-up 6 weeks to 8 weeks or until complete healing occurred - with an assessment at 2 weeks

Participants
100 participants undergoing Milligan Morgan haemorrhoidectomy
Inclusion criteria: undergoing Milligan Morgan haemorrhoidectomy for grade III and IV haemorrhoids
Exclusion criteria: patients having some other concomitant perianal pathology warranting surgery, immunocompromised patients and patients who had recently (within last 2 weeks) taken any antibiotics, pregnant and lactating women were excluded from the study

Interventions
Group A: 500 mg I/V metronidazole and 1 g Ceftriaxone I/V before induction of anaesthesia (n = 50)
Group B: Standard care with no prophylactic antibiotic (n = 50)
All participants underwent standard Milligan Morgan haemorrhoidectomy by the same surgeon in each centre under regional (spinal) or general anaesthesia as per choice of the anaesthetist.
All the patients were given 75 mg I/M Diclofenac Sodium 12 hourly on operation day From first post-operative day and onwards patients were advised to take sitz bath in lukewarm water for minimum of 10 min, twice daily and also to apply the 0.2% GTN ointment with the finger tip after taking the sitz bath. Stool softeners were also advised to be taken (syrup lactulose 30 ml twice daily) from first post-operative day till complete wound healing. All the patients were advised to take tablet codeine phosphate and tablet paracetamol when required from first post-operative day and onwards

Outcomes
Primary outcomes:
- Complete wound healing (defined as complete epithelialisation of the wound).
Data not extracted as data unclear - presented as mean but referred to as complete healing - no units given). Study authors contacted to query, awaiting response
Secondary outcomes:
- Post-operative pain (measured using 0 to 100mm VAS scale with 0 = no pain).
Pain scores were taken daily - preferably after defecation. Data not presented continuously and not extracted - presented using % in stacked graphs only
- Adverse events (participants asked to report any complication occurring especially headache, fever and/or excessive purulent discharge from the wound)

**Notes**

**Funding source:** no funding was received for this study

Query sent to authors regarding outcome data

<table>
<thead>
<tr>
<th>Risk of bias</th>
<th>Authors’ judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
</table>
| Random sequence generation (selection bias) | Low risk          | Quote: Eligible patients were randomly assigned to one of the two treatment groups according to a computer generated list using Random Allocation Software
Comment: Adequate methodology                                                                 |
| Allocation concealment (selection bias) | Unclear risk       | Quote: The double blinding was ensured by not disclosing the treatment group to the patient and to the surgeon who examined the patients on each visit and filled the performa [sic]
Comment: Suggests that the control group were given a placebo - but this is not clear from the study report |
| Blinding of participants and personnel (performance bias) | Unclear risk       | Quote: The double blinding was ensured by not disclosing the treatment group to the patient and to the surgeon who examined the patients on each visit and filled the performa
Comment: Suggests that the control group were given a placebo - but this is not clear from the study report |
| Blinding of outcome assessment (detection bias) | Unclear risk       | **All outcomes:**
Quote: The double blinding was ensured by not disclosing the treatment group to the patient and to the surgeon who examined the patients on each visit and filled the performa
Comment: Suggests that the control group were given a placebo - but this is not clear from the study report |
<p>| Incomplete outcome data (attrition bias) | Unclear risk       | A total of 96 patients out of 100 patients (47 in group A and 49 in group B) completed the study. In group A 3 patients and in group B 1 patient did not complete the study |</p>
<table>
<thead>
<tr>
<th>Selective reporting (reporting bias)</th>
<th>Unclear risk</th>
<th>None noted - protocol not obtained. Presentation of data unclear to review authors</th>
</tr>
</thead>
<tbody>
<tr>
<td>Other bias</td>
<td>Unclear risk</td>
<td>None noted</td>
</tr>
</tbody>
</table>

### Okeniyi 2005

**Methods**

RCT

Undertaken in single hospital setting, Nigeria

Duration of follow-up 3 weeks

**Participants**

32 Nigerian children with 43 pyomyositis abscesses

10 patients had multiple pyomyositis

Inclusion criteria: Children with pyomyositis, included patients with multiple pyomyositis sites

**Interventions**

Group A: Honey-soaked gauze. Wounds were packed twice daily (n = 23 wounds)

Group B: Edinburgh University Solution of Lime (EUSOL)-soaked gauze. Wounds were packed twice daily (n = 20 wounds)

Co-interventions: following fresh surgical incisions and drainage, all subjects had 21-day course of ampicillin plus clavulanic acid (Ampiclox) and gentamicin

Number of participants in each group not reported

**Outcomes**

**Primary outcomes:**

- Complete wound healing (completion of epithelialization)
- Adverse events

**Secondary outcomes:**

- Resource use (length of hospital stay)

**Notes**

**Funding source:** no information given

### Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors’ judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Unclear risk</td>
<td>Quote: The dressing of the individual sites were randomly allocated to give insight into the rate of wound healing with honey and EUSOL devoid of confounding host factors</td>
</tr>
</tbody>
</table>
Okeniyi 2005  

| Blinding of outcome assessment (detection bias) | Unclear risk | Quote: Epithelialization was determined clinically in each case by the first investigator to avoid observer bias. Comment: Not clear if assessment was blinded |
| Incomplete outcome data (attrition bias) | Low risk | Appears to be no loss to follow up. No adverse events reported |
| Selective reporting (reporting bias) | Low risk | None noted - protocol not obtained |
| Other bias | Low risk | Quote: Among the subjects were 10 patients who had multiple pyomyositis. The dressing of the individual sites were randomly allocated to give insight into the rate of wound healing with honey and EUSOL devoid of confounding host factors. Comment: Seems that some participants had more than one wound. Clustering not alluded to in study report |

Piaggesi 2010

| Methods | Two-arm RCT Undertaken in a centre in Italy Duration of follow-up 6 months or to complete re-epithelization |
| Participants | 40 diabetic participants with an infection on the foot requiring surgery resulting in an open wound Inclusion criteria: a surgical lesion resulting from drainage or minor amputation, including trans-metatarsal amputations to treat an infected lesion distal to the ankle. Lesion grade 2B/3B Texas University grading score for diabetic foot ulcers, wider than 5 cm² and left open to heal by secondary intention; transcutaneous oxygen tension (TcPo2) value > 50 mm Hg distal to the ankle Exclusion criteria: bilateral lesions, having had a lesion in the same foot of duration longer than 6 months, HIV positive and any cause of immunodepression other than diabetes, local or systemic documented intolerance to povidone iodine, serum creatinine > 2 mg/dL, and life expectancy shorter than 1 year |
| Interventions | Group A: Gauze with povidone iodine diluted with 50% saline (n = 20) Group B: Gauze with Dermacyn® Wound Care a stable super-oxide solution with neutral pH (n = 20) Both groups were treated with a standardised clinical approach, comprising empiric systemic antibiotic therapy (piperacillin/tazobactam and metronidazole with the adjunct of teicoplanin when methicillin-resistant Staphylococcus aureus was present), prompt and aggressive surgical debridement, metabolic control, and stabilisation of the systemic condition of the patient All the patients were on insulin therapy and they monitored blood glucose; the adjust-
ment of antidiabetic therapy was part of the weekly control visit. Offloading was achieved using irremovable offloading devices and crutches, or alternatively, wheelchairs.

<table>
<thead>
<tr>
<th>Outcomes</th>
<th>Primary outcomes:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>• Time to healing; complete wound healing (not defined)</td>
</tr>
<tr>
<td></td>
<td>• Adverse events</td>
</tr>
<tr>
<td></td>
<td>Secondary outcomes:</td>
</tr>
<tr>
<td></td>
<td>• Number of re-interventions (any procedure carried out in operating theatre)</td>
</tr>
</tbody>
</table>

| Notes | Funding source: a non-restricted research grant from Oculus Innovative Sciences (Petaluma, CA), manufacturers of Dermacyn® Wound Care |

<table>
<thead>
<tr>
<th><strong>Risk of bias</strong></th>
<th><strong>Authors' judgement</strong></th>
<th><strong>Support for judgement</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Low risk</td>
<td>Quote: The patients were randomized into 2 groups: group A and group B by means of a computer-generated randomization code. Comment: Adequate methodology</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Unclear risk</td>
<td>No specific methodology noted</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias)</td>
<td>Unclear risk</td>
<td>Comment: No details but at discharge patients attended their own wounds, as iodine has a distinctive colour patients could probably not be blinded for the intervention</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias)</td>
<td>Low risk</td>
<td>Quote: Measurements and evaluations were performed by another diabetologist.. .. unaware of the allocation of the patients to the different groups. Comment: Adequate blinding</td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias)</td>
<td>Low risk</td>
<td>No drop outs noted</td>
</tr>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>Unclear risk</td>
<td>Comment: No reports on costs in the results, costs is mentioned as one of the endpoints in the methods section</td>
</tr>
<tr>
<td>Other bias</td>
<td>Unclear risk</td>
<td>None noted</td>
</tr>
</tbody>
</table>
Schmidt 1991

| Methods | Two-arm RCT  
|         | Single hospital centre in USA  
|         | Duration of follow-up unclear  
| Participants | 40 participants were randomised following wound complications following a c-section resulting in an open wound. Twenty-one participants had vertical incisions and 19 had transverse incisions  
|         | Inclusion criteria: women with surgical wounds that required healing by secondary intention after either caesarean section or laparotomy for gynaecological surgery. All incisions had opened spontaneously or had been drained to treat a seroma, haematoma or wound abscess before referral to clinic  
|         | Exclusion criteria: Patients with diabetes or cancer, requiring treatment with glucocorticoids or immunosuppressive drugs, had a history of abdominal irradiation, or a chronic debilitating disease  
| Interventions | Group A: standard treatment and aloe vera dermal gel (Carrington Laboratories, Irving, TX). The gel was applied with each dressing change to the granulation tissue in the wound bed at the level of the subcutaneous tissue and dermis (n = 20; 10 with vertical incision; 9 with transverse incision)  
|         | Group B: standard treatment only (n = 20; 11 with vertical incisions, 9 with transverse incisions)  
|         | Standard treatment: blunt debridement with gauze pad or sharp debridement of necrotic tissue. Wound then irrigated with high-volume, high-pressure irrigations using a 60 ml Luer Lock syringe with an 18-gauge angiocath and an irrigating volume of 1000 ml. A wet-to-dry dressing was applied, using a solution of equal parts of saline and sodium hypochlorite 0.025%. Wound care was performed every 8 hours initially and then every 12 hours after granulation tissue developed  
| Outcomes | Primary outcomes:  
|         | • Time to healing (defined as when wound was completely epithelialised)  
|         | • Adverse events  
|         | Secondary outcomes:  
|         | • None  
| Notes | Funding  
|         | source: no information given  

**Risk of bias**

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors’ judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
</table>
| Random sequence generation (selection bias) | Low risk | Quote: Randomised using a random number sequence generated by a computer program  
| | | Comment: Adequate methodology |
| Allocation concealment (selection bias) | Unclear risk | No information given |
**Schmidt 1991**  
*(Continued)*

<table>
<thead>
<tr>
<th>Blinding of participants and personnel (performance bias)</th>
<th>Unclear risk</th>
<th>No details</th>
</tr>
</thead>
<tbody>
<tr>
<td>All outcomes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias)</td>
<td>High risk</td>
<td>Quote The study was not blinded</td>
</tr>
<tr>
<td>All outcomes</td>
<td></td>
<td>Comment: Taken as not blinded for any outcome</td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias)</td>
<td>High risk</td>
<td>Comment: 19 participants (12 receiving standard treatment and 7 receiving aloe vera) were lost to follow up. No additional data was available for those lost to follow-up. Classed as high risk of attrition bias as large % of participants lost to follow-up</td>
</tr>
<tr>
<td>All outcomes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>Unclear risk</td>
<td>None noted - protocol not obtained</td>
</tr>
<tr>
<td>Other bias</td>
<td>Unclear risk</td>
<td>No defined follow up period</td>
</tr>
</tbody>
</table>

**Tosti 2014**

<table>
<thead>
<tr>
<th>Methods</th>
<th>Two-arm RCT</th>
</tr>
</thead>
<tbody>
<tr>
<td>Undertaken in one centre in the USA</td>
<td></td>
</tr>
<tr>
<td>Duration of follow-up 6 weeks</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Participants</th>
<th>100 participants undergoing surgery for all hand abscesses</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inclusion criteria: no further criteria listed</td>
<td></td>
</tr>
<tr>
<td>Exclusion criteria: purulence proximal to the wrist, absence of purulent fluid, chronic wound infections, and premature discharge from the hospital against medical advice (this final criterion seems to be a post-randomisation exclusion).</td>
<td></td>
</tr>
<tr>
<td>All patients received a thorough debridement in the operating room with excision of necrotic tissue. The wounds were irrigated with 3 L normal saline under a low-velocity irrigation system and then partially closed over a Penrose drain or an iodinated gauze wick. The wounds were then dressed with an iodine-petroleum gauze, sterile gauze, and an orthosis. Patients returned to the operating room if persistent or progressive erythema was noted or if purulent fluid was expressed from the wound. On the first post-operative day, the Penrose drain or gauze wicks were removed leaving open areas of the wound which were then exposed to the intervention or not, based on randomisation.</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Interventions</th>
<th>Group A: a 10% povidone iodine solution mixed with saline in a 1:1 ratio hand soak. Hands were submerged for 20 minutes, three times per day. After each soak wounds were redressed with sterile gauze and an orthosis. Soaks were stopped on discharge. (n = 50) Group B: no soaks, the wounds were treated with a daily dressing change with sterile gauze and an orthosis (n = 50) Empiric antibiotic coverage at time of admission (intravenous ampicillin-sulbactam if history of bite wound; otherwise vancomycin)</th>
</tr>
</thead>
</table>
Patients were given 10 days of oral antibiotics at discharge unless they had had positive blood cultures. In that case, they received 6 weeks of intravenous antibiotics and were observed by an infectious disease consultant.

### Outcomes

**Primary outcomes:**
- None

**Secondary outcomes:**
- Resource use (length of hospital stay; re-admissions and re-operations)

### Notes

Funding source: not reported

Authors were contacted and confirmed that these were all partially open wounds

### Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors' judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Low risk</td>
<td>Quote: when purulent fluid was confirmed at the time of debridement, patients were randomized via a computer-generated schedule. Comment: Computer-generated randomisation sequence - adequate</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Unclear risk</td>
<td>No information provided</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias) All outcomes</td>
<td>High risk</td>
<td>Quote: It was not possible to blind subjects and observers to the study protocol, which could have influenced the decision to perform repeated debridements. Comment: No blinding took place</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias)</td>
<td>High risk</td>
<td>Quote: It was not possible to blind subjects and observers to the study protocol, which could have influenced the decision to perform repeated debridements. Comment: No blinding took place</td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias)</td>
<td>Low risk</td>
<td>Quote: Two patients from each group were excluded from the final analysis for leaving the hospital against medical advice. Comment: Post-randomisation exclusions - but small number. Classed as low risk</td>
</tr>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>Low risk</td>
<td>None noted - protocol not obtained</td>
</tr>
<tr>
<td>Other bias</td>
<td>Low risk</td>
<td>None noted</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>Cannavo 1998</td>
<td>Use of antibiotic/antiseptic was not the only difference between groups</td>
</tr>
<tr>
<td>Eldrup 1985</td>
<td>Use of antibiotic/antiseptic was not the only difference between groups</td>
</tr>
<tr>
<td>Goetze 2006</td>
<td>Use of antibiotic/antiseptic was not the only difference between groups</td>
</tr>
<tr>
<td>Hien 1988</td>
<td>Not considered RCT based on available information, abstract only and no outcome data reported - unable to contact authors</td>
</tr>
<tr>
<td>Moore 2000</td>
<td>Use of antibiotic/antiseptic was not the only difference between groups</td>
</tr>
<tr>
<td>Murthy 2012</td>
<td>Not surgical wound healing by secondary intention</td>
</tr>
<tr>
<td>Nielsen 2012</td>
<td>Use of antibiotic/antiseptic was not the only difference between groups</td>
</tr>
<tr>
<td>Sozener 2011</td>
<td>No relevant outcome data for review collected - confirmed with author; purpose of study was to assess occurrence of fistula over year following surgery</td>
</tr>
<tr>
<td>Taylor 2011</td>
<td>Not surgical wounds healing by secondary intention</td>
</tr>
<tr>
<td>Viciano 2000</td>
<td>Use of antibiotic/antiseptic was not the only difference between groups</td>
</tr>
<tr>
<td>Walker 1991</td>
<td>Use of antibiotic/antiseptic was not the only difference between groups</td>
</tr>
<tr>
<td>Williams 1981</td>
<td>Use of antibiotic/antiseptic was not the only difference between groups</td>
</tr>
<tr>
<td>Yang 2013</td>
<td>Use of antibiotic/antiseptic was not the only difference between groups</td>
</tr>
</tbody>
</table>

### Characteristics of studies awaiting assessment  [ordered by study ID]

<table>
<thead>
<tr>
<th>Fillmann 2004</th>
</tr>
</thead>
<tbody>
<tr>
<td>Methods</td>
</tr>
<tr>
<td>Participants</td>
</tr>
<tr>
<td>Interventions</td>
</tr>
<tr>
<td>Outcomes</td>
</tr>
</tbody>
</table>
### Fillmann 2004 (Continued)

**Notes**  
Portuguese; identified in search update

### Mirzabeygi 2011

<table>
<thead>
<tr>
<th>Methods</th>
<th>RCT</th>
</tr>
</thead>
<tbody>
<tr>
<td>Participants</td>
<td>47 participants undergoing haemorrhoidectomy</td>
</tr>
<tr>
<td>Interventions</td>
<td>Metronidazole versus placebo</td>
</tr>
<tr>
<td>Outcomes</td>
<td>Post-operative pain</td>
</tr>
<tr>
<td>Notes</td>
<td>Abstract only; attempt to contact author unsuccessful</td>
</tr>
</tbody>
</table>

### Quilici 1998

<table>
<thead>
<tr>
<th>Methods</th>
<th>RCT</th>
</tr>
</thead>
<tbody>
<tr>
<td>Participants</td>
<td>40 participants undergoing haemorrhoidectomy</td>
</tr>
<tr>
<td>Interventions</td>
<td>Unguent containing neomycin sulfate antibiotic versus unguent containing (Xilodase(R)) plus hyaluronidase (50 UTR)</td>
</tr>
</tbody>
</table>
| Outcomes      | Wound closure time  
Surgical wound pain  
Local infection prevention  
Post-operative adverse effects |
| Notes         | English abstract; full text in Portuguese (not yet obtained) |

### Vasei 2008

<table>
<thead>
<tr>
<th>Methods</th>
<th>Randomized controlled clinical trial</th>
</tr>
</thead>
<tbody>
<tr>
<td>Participants</td>
<td>24 patients with surgical wound of pilonidal sinuses</td>
</tr>
<tr>
<td>Interventions</td>
<td>Honey dressing (n = 12) versus saline-soaked dressing (n = 12)</td>
</tr>
</tbody>
</table>
| Outcomes        | Complete wound healing  
Eradication of infection  
Adverse effects  
Resource use (length of hospital stay) |
| Notes           | English abstract; full text in Farsi (not yet obtained) |
### DATA AND ANALYSES

#### Comparison 1. Polyvidone iodine-impregnated mesh compared with alginate mesh

<table>
<thead>
<tr>
<th>Outcome or subgroup title</th>
<th>No. of studies</th>
<th>No. of participants</th>
<th>Statistical method</th>
<th>Effect size</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Proportion of wounds healed</td>
<td>1</td>
<td></td>
<td>Risk Ratio (M-H, Fixed, 95% CI)</td>
<td>Subtotals only</td>
</tr>
<tr>
<td>2 Wound infection</td>
<td>1</td>
<td></td>
<td>Risk Ratio (M-H, Fixed, 95% CI)</td>
<td>Subtotals only</td>
</tr>
<tr>
<td>3 Mean number of dressing changes</td>
<td>1</td>
<td></td>
<td>Mean Difference (IV, Fixed, 95% CI)</td>
<td>Subtotals only</td>
</tr>
</tbody>
</table>

#### Comparison 2. Zinc oxide mesh compared with placebo

<table>
<thead>
<tr>
<th>Outcome or subgroup title</th>
<th>No. of studies</th>
<th>No. of participants</th>
<th>Statistical method</th>
<th>Effect size</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Wound infection (based on presence of smell)</td>
<td>1</td>
<td></td>
<td>Risk Ratio (M-H, Fixed, 95% CI)</td>
<td>Totals not selected</td>
</tr>
<tr>
<td>2 Participants prescribed antibiotics</td>
<td>1</td>
<td></td>
<td>Odds Ratio (M-H, Fixed, 95% CI)</td>
<td>Totals not selected</td>
</tr>
</tbody>
</table>

#### Comparison 3. Sucralfate compared with petrolatum

<table>
<thead>
<tr>
<th>Outcome or subgroup title</th>
<th>No. of studies</th>
<th>No. of participants</th>
<th>Statistical method</th>
<th>Effect size</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Proportion of wounds healed</td>
<td>1</td>
<td></td>
<td>Risk Ratio (M-H, Fixed, 95% CI)</td>
<td>Subtotals only</td>
</tr>
<tr>
<td>2 Pain score (10-point VAS scale)</td>
<td>1</td>
<td></td>
<td>Mean Difference (IV, Fixed, 95% CI)</td>
<td>Subtotals only</td>
</tr>
</tbody>
</table>

#### Comparison 4. Trimethoprim-sulfamethoxazole compared with placebo

<table>
<thead>
<tr>
<th>Outcome or subgroup title</th>
<th>No. of studies</th>
<th>No. of participants</th>
<th>Statistical method</th>
<th>Effect size</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Adverse events (attributed to medication)</td>
<td>1</td>
<td></td>
<td>Risk Ratio (M-H, Fixed, 95% CI)</td>
<td>Subtotals only</td>
</tr>
<tr>
<td>2 Wound recurrence</td>
<td>1</td>
<td></td>
<td>Risk Ratio (M-H, Fixed, 95% CI)</td>
<td>Subtotals only</td>
</tr>
</tbody>
</table>
### Comparison 5. Honey compared with EUSOL

<table>
<thead>
<tr>
<th>Outcome or subgroup title</th>
<th>No. of studies</th>
<th>No. of participants</th>
<th>Statistical method</th>
<th>Effect size</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Proportion of wounds healed</td>
<td>1</td>
<td>Risk Ratio (M-H, Fixed, 95% CI)</td>
<td>Subtotals only</td>
<td></td>
</tr>
<tr>
<td>2 Mean length of hospital stay</td>
<td>1</td>
<td>Mean Difference (IV, Fixed, 95% CI)</td>
<td>Subtotals only</td>
<td></td>
</tr>
</tbody>
</table>

### Comparison 6. Iodine compared with Dermacyn®

<table>
<thead>
<tr>
<th>Outcome or subgroup title</th>
<th>No. of studies</th>
<th>No. of participants</th>
<th>Statistical method</th>
<th>Effect size</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Proportion of wounds healed</td>
<td>1</td>
<td>Risk Ratio (M-H, Fixed, 95% CI)</td>
<td>Subtotals only</td>
<td></td>
</tr>
</tbody>
</table>

### Comparison 7. Triclosan compared with sodium hypochlorite

<table>
<thead>
<tr>
<th>Outcome or subgroup title</th>
<th>No. of studies</th>
<th>No. of participants</th>
<th>Statistical method</th>
<th>Effect size</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Time to wound healing</td>
<td>1</td>
<td>Mean Difference (IV, Fixed, 95% CI)</td>
<td>Subtotals only</td>
<td></td>
</tr>
<tr>
<td>2 Adverse events</td>
<td>1</td>
<td>Mean Difference (IV, Fixed, 95% CI)</td>
<td>Subtotals only</td>
<td></td>
</tr>
<tr>
<td>(bleeding/secretions)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3 Pain</td>
<td>1</td>
<td>Mean Difference (IV, Fixed, 95% CI)</td>
<td>Subtotals only</td>
<td></td>
</tr>
</tbody>
</table>

### Analysis 1.1. Comparison 1 Polyvidone iodine-impregnated mesh compared with alginate mesh, Outcome 1 Proportion of wounds healed.

Review: Antibiotics and antiseptics for surgical wounds healing by secondary intention

Comparison: 1 Polyvidone iodine-impregnated mesh compared with alginate mesh

Outcome: 1 Proportion of wounds healed

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>polyvidone-iodine mesh</th>
<th>alginate mesh</th>
<th>Risk Ratio M-H,Fixed,95% CI</th>
<th>Weight</th>
<th>Risk Ratio M-H,Fixed,95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brehant 2009</td>
<td>25/34</td>
<td>34/37</td>
<td>-</td>
<td></td>
<td>0.80 [ 0.64, 1.00 ]</td>
</tr>
</tbody>
</table>

Test for subgroup differences: Not applicable

Antibiotics and antiseptics for surgical wounds healing by secondary intention (Review)
Copyright © 2016 The Cochrane Collaboration. Published by John Wiley & Sons, Ltd.
Analysis 1.2. Comparison 1 Polyvidone iodine-impregnated mesh compared with alginate mesh, Outcome 2 Wound infection.

Review: Antibiotics and antiseptics for surgical wounds healing by secondary intention

Comparison: 1 Polyvidone iodine-impregnated mesh compared with alginate mesh

Outcome: 2 Wound infection

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>polyvidone-iodine mesh n/N</th>
<th>alginate mesh n/N</th>
<th>Risk Ratio M-H,Fixed 95% CI</th>
<th>Weight M-H,Fixed 95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brehant 2009</td>
<td>2/29</td>
<td>3/37</td>
<td>0.85 [0.15, 4.76]</td>
<td></td>
</tr>
</tbody>
</table>

Test for subgroup differences: Not applicable
Analysis 1.3. Comparison 1 Polyvidone iodine-impregnated mesh compared with alginate mesh, Outcome 3 Mean number of dressing changes.

Review: Antibiotics and antiseptics for surgical wounds healing by secondary intention

Comparison: 1 Polyvidone iodine-impregnated mesh compared with alginate mesh

Outcome: 3 Mean number of dressing changes

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>Polyvidone-iodine mesh</th>
<th>Alginate mesh</th>
<th>Mean Difference</th>
<th>Weight</th>
<th>Mean Difference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brehant 2009</td>
<td>37</td>
<td>37</td>
<td>17.9 (8)</td>
<td>4.70</td>
<td>[1.66, 7.74]</td>
</tr>
</tbody>
</table>

Test for subgroup differences: Not applicable

Analysis 2.1. Comparison 2 Zinc oxide mesh compared with placebo, Outcome 1 Wound infection (based on presence of smell).

Review: Antibiotics and antiseptics for surgical wounds healing by secondary intention

Comparison: 2 Zinc oxide mesh compared with placebo

Outcome: 1 Wound infection (based on presence of smell)

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>Zinc oxide</th>
<th>Placebo</th>
<th>Risk Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n/N</td>
<td>n/N</td>
<td>M-H,Fixed,95% CI</td>
</tr>
<tr>
<td>Argen 2006</td>
<td>1/33</td>
<td>8/31</td>
<td>0.12</td>
</tr>
</tbody>
</table>

0.005 0.1 1 10 200

Favours Zinc-oxide  Favours placebo
### Analysis 2.2. Comparison 2 Zinc oxide mesh compared with placebo, Outcome 2 Participants prescribed antibiotics.

**Review:** Antibiotics and antiseptics for surgical wounds healing by secondary intention

**Comparison:** 2 Zinc oxide mesh compared with placebo

**Outcome:** 2 Participants prescribed antibiotics

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>Zinc oxide</th>
<th>Placebo</th>
<th>Odds Ratio</th>
<th>Odds Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n/N</td>
<td>n/N</td>
<td>M-H,Fixed,95% CI</td>
<td>M-H,Fixed,95% CI</td>
</tr>
<tr>
<td>Argen 2006</td>
<td>3/33</td>
<td>12/31</td>
<td><strong>0.16 [0.04, 0.64]</strong></td>
<td></td>
</tr>
</tbody>
</table>

0.01 0.1 1 10 100
Favours zinc Favours placebo

### Analysis 3.1. Comparison 3 Sucralfate compared with petrolatum, Outcome 1 Proportion of wounds healed.

**Review:** Antibiotics and antiseptics for surgical wounds healing by secondary intention

**Comparison:** 3 Sucralfate compared with petrolatum

**Outcome:** 1 Proportion of wounds healed

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>Sucralfate</th>
<th>Petrolatum</th>
<th>Risk Ratio</th>
<th>Weight</th>
<th>Risk Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n/N</td>
<td>n/N</td>
<td>M-H,Fixed,95% CI</td>
<td></td>
<td>M-H,Fixed,95% CI</td>
</tr>
<tr>
<td>Gupta 2008</td>
<td>45/58</td>
<td>30/58</td>
<td></td>
<td></td>
<td><strong>1.50 [1.13, 1.99]</strong></td>
</tr>
</tbody>
</table>

Test for subgroup differences: Not applicable

0.01 0.1 1 10 100
Favours sucralfate Favours petrolatum
Analysis 3.2. Comparison 3 Sucralfate compared with petrolatum, Outcome 2 Pain score (10-point VAS scale).

Review: Antibiotics and antiseptics for surgical wounds healing by secondary intention

Comparison: 3 Sucralfate compared with petrolatum

Outcome: 2 Pain score (10-point VAS scale)

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>Sucralfate</th>
<th>Petrolatum</th>
<th>Mean Difference</th>
<th>Weight</th>
<th>Mean Difference</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N Mean(SD)</td>
<td>N Mean(SD)</td>
<td>IV,Fixed,95% CI</td>
<td></td>
<td>IV,Fixed,95% CI</td>
</tr>
<tr>
<td>Gupta 2008</td>
<td>58 0.2 (0.1)</td>
<td>58 1.4 (0.3)</td>
<td>-1.20 [-1.28, -1.12 ]</td>
<td></td>
<td>-1.20 [-1.28, -1.12 ]</td>
</tr>
</tbody>
</table>

Test for subgroup differences: Not applicable

Analysis 4.1. Comparison 4 Trimethoprim-sulfamethoxazole compared with placebo, Outcome 1 Adverse events (attributed to medication).

Review: Antibiotics and antiseptics for surgical wounds healing by secondary intention

Comparison: 4 Trimethoprim-sulfamethoxazole compared with placebo

Outcome: 1 Adverse events (attributed to medication)

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>trimethoprim-sulfamethoxazole</th>
<th>Placebo</th>
<th>Risk Ratio</th>
<th>Weight</th>
<th>Risk Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n/N</td>
<td>n/N</td>
<td>M-H,Fixed,95% CI</td>
<td></td>
<td>M-H,Fixed,95% CI</td>
</tr>
<tr>
<td>Duong 2010</td>
<td>14/73</td>
<td>6/76</td>
<td>2.43 [ 0.99, 5.98 ]</td>
<td></td>
<td>2.43 [ 0.99, 5.98 ]</td>
</tr>
</tbody>
</table>

Test for subgroup differences: Not applicable
Analysis 4.2. Comparison 4 Trimethoprim-sulfamethoxazole compared with placebo, Outcome 2 Wound recurrence.

Review: Antibiotics and antiseptics for surgical wounds healing by secondary intention

Comparison: 4 Trimethoprim-sulfamethoxazole compared with placebo

Outcome: 2 Wound recurrence

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>trimethoprim-sulfamethoxazole</th>
<th>Placebo</th>
<th>Risk Ratio</th>
<th>Weight</th>
<th>Risk Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n/N</td>
<td>n/N</td>
<td>M-H,Fixed,95% CI</td>
<td></td>
<td>M-H,Fixed,95% CI</td>
</tr>
<tr>
<td>Duong 2010</td>
<td>13/46</td>
<td>15/52</td>
<td>0.98 [ 0.52, 1.84 ]</td>
<td>0.01</td>
<td>0.1 1 10 100</td>
</tr>
</tbody>
</table>

Test for subgroup differences: Not applicable

Analysis 5.1. Comparison 5 Honey compared with EUSOL, Outcome 1 Proportion of wounds healed.

Review: Antibiotics and antiseptics for surgical wounds healing by secondary intention

Comparison: 5 Honey compared with EUSOL

Outcome: 1 Proportion of wounds healed

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>Honey</th>
<th>EUSOL</th>
<th>Risk Ratio</th>
<th>Weight</th>
<th>Risk Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n/N</td>
<td>n/N</td>
<td>M-H,Fixed,95% CI</td>
<td></td>
<td>M-H,Fixed,95% CI</td>
</tr>
<tr>
<td>Okewenyi 2005</td>
<td>20/23</td>
<td>11/20</td>
<td>1.58 [ 1.03, 2.42 ]</td>
<td>0.01</td>
<td>0.1 1 10 100</td>
</tr>
</tbody>
</table>

Test for subgroup differences: Not applicable
**Analysis 5.2. Comparison 5 Honey compared with EUSOL, Outcome 2 Mean length of hospital stay.**

**Review:** Antibiotics and antiseptics for surgical wounds healing by secondary intention

**Comparison:** 5 Honey compared with EUSOL

**Outcome:** 2 Mean length of hospital stay

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>Honey</th>
<th>EUSOL</th>
<th>Mean Difference</th>
<th>Weight</th>
<th>Mean Difference</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N</td>
<td>Mean(SD)</td>
<td>N</td>
<td>Mean(SD)</td>
<td>IV,Fixed,95% CI</td>
</tr>
<tr>
<td>Okenyi 2005</td>
<td>23</td>
<td>16.1 (4.16)</td>
<td>20</td>
<td>18.6 (2.14)</td>
<td>-2.50 [-4.44, -0.56]</td>
</tr>
</tbody>
</table>

Test for subgroup differences: Not applicable

---

**Analysis 6.1. Comparison 6 Iodine compared with Dermacyn®, Outcome 1 Proportion of wounds healed.**

**Review:** Antibiotics and antiseptics for surgical wounds healing by secondary intention

**Comparison:** 6 Iodine compared with Dermacyn

**Outcome:** 1 Proportion of wounds healed

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>Iodine</th>
<th>Dermacyn</th>
<th>Risk Ratio</th>
<th>Weight</th>
<th>Risk Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n/N</td>
<td>n/N</td>
<td>M-H,Fixed,95% CI</td>
<td>M-H,Fixed,95% CI</td>
<td></td>
</tr>
<tr>
<td>Piaggi 2010</td>
<td>11/20</td>
<td>18/20</td>
<td>0.61 [0.40, 0.93]</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Test for subgroup differences: Not applicable
**Analysis 7.1. Comparison 7 Triclosan compared with sodium hypochlorite, Outcome 1 Time to wound healing.**

Review: Antibiotics and antiseptics for surgical wounds healing by secondary intention

Comparison: 7 Triclosan compared with sodium hypochlorite

Outcome: 1 Time to wound healing

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>Experimental</th>
<th>Control</th>
<th>Mean Difference</th>
<th>Weight</th>
<th>Mean Difference</th>
</tr>
</thead>
<tbody>
<tr>
<td>N</td>
<td>Mean(SD)</td>
<td>N</td>
<td>Mean(SD)</td>
<td>IV/Fixed,95% CI</td>
<td>IV/Fixed,95% CI</td>
</tr>
<tr>
<td>Giannini 2014</td>
<td>55</td>
<td>21.7 (3.8)</td>
<td>58</td>
<td>23.4 (5.4)</td>
<td>-1.70 [-3.41, 0.01]</td>
</tr>
</tbody>
</table>

Test for subgroup differences: Not applicable

**Analysis 7.2. Comparison 7 Triclosan compared with sodium hypochlorite, Outcome 2 Adverse events (bleeding/secretions).**

Review: Antibiotics and antiseptics for surgical wounds healing by secondary intention

Comparison: 7 Triclosan compared with sodium hypochlorite

Outcome: 2 Adverse events (bleeding/secretions)

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>Triclosan</th>
<th>Sodium hypo</th>
<th>Mean Difference</th>
<th>Weight</th>
<th>Mean Difference</th>
</tr>
</thead>
<tbody>
<tr>
<td>N</td>
<td>Mean(SD)</td>
<td>N</td>
<td>Mean(SD)</td>
<td>IV/Fixed,95% CI</td>
<td>IV/Fixed,95% CI</td>
</tr>
<tr>
<td>Gupta 2008</td>
<td>55</td>
<td>1.5 (1.7)</td>
<td>58</td>
<td>2.5 (1.9)</td>
<td>-1.00 [-1.66, -0.34]</td>
</tr>
</tbody>
</table>

Test for subgroup differences: Not applicable
Analysis 7.3. Comparison 7 Triclosan compared with sodium hypochlorite, Outcome 3 Pain.

Review: Antibiotics and antiseptics for surgical wounds healing by secondary intention

Comparison: 7 Triclosan compared with sodium hypochlorite

Outcome: 3 Pain

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>Triclosan</th>
<th>Sodium hypo</th>
<th>Mean Difference</th>
<th>Weight</th>
<th>Mean Difference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gupta 2008</td>
<td>55</td>
<td>58</td>
<td>-1.20</td>
<td></td>
<td>-1.20</td>
</tr>
</tbody>
</table>

Test for subgroup differences: Not applicable

ADDITIONAL TABLES

Table 1. Study outcomes

<table>
<thead>
<tr>
<th>Study</th>
<th>Comparison</th>
<th>Length of follow-up and time points of data presented</th>
<th>Time-to-healing data</th>
<th>Proportion of wounds completely healed</th>
<th>Adverse events</th>
<th>Change in ulcer size</th>
<th>Wound infection</th>
<th>Pain</th>
<th>Resource use</th>
<th>Cost</th>
<th>Wound recurrence</th>
</tr>
</thead>
<tbody>
<tr>
<td>Argen 2006</td>
<td>Group A: Zinc oxide povidone mesh (n = 33)</td>
<td>Follow-up 90 days for healing 7 days for pain Data presented at 90 days/13 weeks</td>
<td>Median time to healing: Group A: 54 days (IQR 42 to 71 days) Group B: 62 days (55 to 82 days) Unad-</td>
<td>n/a</td>
<td>n/a</td>
<td>n/a</td>
<td>Manifest by foul smell: Group A: 1/33 Group B: 8/31 Prescribed antibiotics during post-</td>
<td>Measured at 7 days as pain intensity. The scale used was not reported although</td>
<td>n/a</td>
<td>n/a</td>
<td>n/a</td>
</tr>
</tbody>
</table>
Table 1. Study outcomes  

<table>
<thead>
<tr>
<th>Study</th>
<th>Follow-up</th>
<th>Mean time-to-event data presented</th>
<th>Change in wound volume at 28 days</th>
<th>Defined according to the Centers for Disease Control and Prevention (CDC) - both</th>
<th>Mean number of dressing changes to achieve healing (SD - assumed from data in)</th>
<th>Mean cost per participant (nursing time for dressing changes and cost of dressing)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brehant 2009</td>
<td>Group A: povidone iodine-impregnated mesh (n = 34)</td>
<td>28 days (change in wound volume)</td>
<td>(no data on variation around point estimates)</td>
<td>-</td>
<td>n/a</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>Group B: Alginate mesh (n = 37)</td>
<td>6 weeks</td>
<td>Data presented as not all wounds healed during</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Table 1. Study outcomes (Continued)

<table>
<thead>
<tr>
<th>Author</th>
<th>Year</th>
<th>Group A</th>
<th>Group B</th>
<th>Follow-up</th>
<th>Number of participants</th>
<th>Number of participants with adverse events</th>
<th>Number of participants with treatment failures</th>
<th>Deep and superficial infections included</th>
</tr>
</thead>
<tbody>
<tr>
<td>Duong 2010</td>
<td></td>
<td>trimethoprim-sulfamethoxazole (n = 77)</td>
<td>placebo (n = 84)</td>
<td>10 days to 14 days and 90 days post-op</td>
<td>n/a</td>
<td>n/a</td>
<td>n/a</td>
<td>Group A: 2/29 Group B: 3/37 It was not clear from the paper whether the SSIs reported occurred in different people</td>
</tr>
</tbody>
</table>

Follow-up 6 weeks following follow-up. It is not clear how many people these events occurred in possible clustering.

Group A: 87% Group B: 91%

Deep and superficial infections included

Group A: 17.9 (8) Group B: 13.2 (5)

Euro Group A: 286.5 Group B: 268

New lesions within 5 cm of original wound at 10 days

Group A: 9/73 Group B: 19/76 At 3 months* Group A: 13/46 Group B: 15/52 *numerator figure calculated by re-
<table>
<thead>
<tr>
<th>Study</th>
<th>Group A</th>
<th>Group B</th>
<th>Group C</th>
<th>Follow-up</th>
<th>Pain</th>
<th>Complete wound healing</th>
<th>Complete wound healing</th>
<th>Pain</th>
<th>Complete wound healing</th>
<th>Complete wound healing</th>
<th>Pain</th>
<th>Change in ulcer size</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Fernández 2002</em></td>
<td>Rhi-zophora mangle bark extract. Applied once a day (n = 12)</td>
<td>Rhi-zophora mangle bark extract. Applied twice a day (10 h to 12-h interval) (n = 12)</td>
<td>Mer-curochrome. Applied twice a day (10 h to 12-h interval) (n = 13)</td>
<td>6 weeks Data presented at 3, 4, 5, 6 weeks following surgery</td>
<td>n/a</td>
<td>n/a</td>
<td>Very unclear methodology and limited data. Not extracted.</td>
<td>n/a</td>
<td>n/a</td>
<td>n/a</td>
<td>n/a</td>
<td>n/a</td>
</tr>
<tr>
<td><em>Gupta 2008</em></td>
<td>Sucralfate cream (n)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*Table 1. Study outcomes (Continued)*
Table 1. Study outcomes  

<table>
<thead>
<tr>
<th>Group A</th>
<th>Group B</th>
<th>Follow-up</th>
<th>Mean time to healing (SD)</th>
<th>Mean bleeding/secretions (SD)</th>
<th>Mean anal pain (SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>41</td>
<td>52</td>
<td>21 days</td>
<td>21.7 days (3.8)</td>
<td>6.24 (1.84)</td>
<td>6.4 (2.0)</td>
</tr>
<tr>
<td>45/58</td>
<td>30/58</td>
<td>23.4</td>
<td>6.77 (1.76)</td>
<td>6.77 (1.76)</td>
<td>7.2 (2.0)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(5.37)</td>
<td>4.03 (1.93)</td>
<td>4.03 (1.93)</td>
<td>3.7 (1.9)</td>
</tr>
</tbody>
</table>

*Figures calculated by review authors from % presented in study - also taking missing data into account.
Table 1. Study outcomes (Continued)

<table>
<thead>
<tr>
<th>Study</th>
<th>Group A</th>
<th>Group B</th>
</tr>
</thead>
<tbody>
<tr>
<td>Khan 2014</td>
<td>Group A: 500 mg I/V metronidazole and 1 g Ceftriaxone I/V (n = 50)</td>
<td>Group B: Standard care with no prophylactic antibiotic (n = 50)</td>
</tr>
<tr>
<td></td>
<td>Follow-up 6 weeks to 8 weeks or until complete healing Data presented at end of follow-up</td>
<td>n/a</td>
</tr>
<tr>
<td></td>
<td>Presented as mean (SD) for groups. Not clear what this mean figure related to - just referred to as complete healing - no units given</td>
<td>n/a</td>
</tr>
<tr>
<td></td>
<td>Low grade fever</td>
<td>n/a</td>
</tr>
<tr>
<td></td>
<td>Group A: 5</td>
<td>Group B: 4</td>
</tr>
<tr>
<td></td>
<td>Collected using VAS scales but not presented continuously as mean - presented only as % with scores ranging from 0 to 70. Unclear how calculated and not mean pain score. Not extracted</td>
<td>n/a</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Okeniyi 2005</th>
<th>Group A: Honey-soaked</th>
<th>Group B:</th>
<th>n/a</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Follow-up 21 days Data</td>
<td></td>
<td>n/a</td>
</tr>
<tr>
<td></td>
<td>No. of wounds with</td>
<td></td>
<td>n/a</td>
</tr>
<tr>
<td></td>
<td>Reported no</td>
<td></td>
<td>n/a</td>
</tr>
<tr>
<td></td>
<td>Mean Length of hosp-</td>
<td></td>
<td>n/a</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>n/a</td>
</tr>
<tr>
<td>Study Outcomes</td>
<td>Group A</td>
<td>Group B</td>
<td>Group A</td>
</tr>
<tr>
<td>----------------</td>
<td>---------</td>
<td>---------</td>
<td>---------</td>
</tr>
<tr>
<td>Presented at 21 days/3 weeks</td>
<td></td>
<td></td>
<td>complete epithelialisation at 21 days (%)</td>
</tr>
<tr>
<td>Adverse events in either group.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Data presented at 6 months or to complete re-epithelialisation</td>
</tr>
<tr>
<td></td>
<td></td>
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<td></td>
</tr>
</tbody>
</table>

**Table 1. Study outcomes (Continued)**

<table>
<thead>
<tr>
<th>Study Outcomes</th>
<th>Group A</th>
<th>Group B</th>
<th>Group A</th>
<th>Group B</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Piaggi 2010</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Follow-up 6 months or to complete re-epithelialisation</td>
<td></td>
<td></td>
<td>Time to healing (whilst Kaplan-Meier curve presented data noted to be mean and (SD)) Group A: 10.5 (5.9) Group B: 16.5 (7.1) As not</td>
<td>Complete wound healing</td>
</tr>
<tr>
<td></td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Study</td>
<td>Group A</td>
<td>Group B</td>
<td>Follow-up</td>
<td>Mean time in days to complete healing (SD)</td>
</tr>
<tr>
<td>-------</td>
<td>---------</td>
<td>---------</td>
<td>-----------</td>
<td>------------------------------------------</td>
</tr>
<tr>
<td>Schmidt 1991</td>
<td>Aloe vera and standard treatment (n = 20)</td>
<td>Standard treatment only (n = 20)</td>
<td>Unclear</td>
<td>Group A: 53 ± 24 Group B: 83 ± 28</td>
</tr>
<tr>
<td>Tosti 2014</td>
<td>10% povidone iodine solution mixed with saline (n = 20)</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Table 1. Study outcomes (Continued)

<table>
<thead>
<tr>
<th>Group</th>
<th>Readmissions*</th>
<th>Reoperations*</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>4</td>
<td>2</td>
</tr>
<tr>
<td>B</td>
<td>2</td>
<td>3</td>
</tr>
</tbody>
</table>

*Not clear if data presented per participant or per event - clustering possible

**Appendix 1. Search Strategies**

The Cochrane Central Register of Controlled Trials (CENTRAL) (The Cochrane Library)

#1 MeSH descriptor: [Anti-Infective Agents] explode all trees
#2 MeSH descriptor: [Penicillins] explode all trees
#3 MeSH descriptor: [Cephalosporins] explode all trees
#4 MeSH descriptor: [Aminoglycosides] explode all trees
#5 MeSH descriptor: [Quinolones] explode all trees
#6 MeSH descriptor: [Clindamycin] explode all trees
#7 MeSH descriptor: [Metronidazole] explode all trees
#8 MeSH descriptor: [Trimethoprim] explode all trees
#9 MeSH descriptor: [Mupirocin] explode all trees
#10 MeSH descriptor: [Neomycin] explode all trees
#11 MeSH descriptor: [Fusidic Acid] explode all trees
#12 MeSH descriptor: [Framycetin] explode all trees
#13 MeSH descriptor: [Polymyxins] explode all trees
#14 MeSH descriptor: [Chlorotetracycline] explode all trees
#15 (antibiotic* or antimicrobial* or antibacterial* or penicillin* or cephalosporin* or aminoglycoside* or quinolone* or clindamycin or metronidazole or trimethoprim or mupirocin or pseudomonic acid or neomycin or fusidic acid or framycetin or polymyxin*)
Antibiotics and antiseptics for surgical wounds healing by secondary intention (Review)

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Antibiotics and antiseptics for surgical wounds healing by secondary intention (Review)

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Appendix 2. 'Risk of bias' assessment (individually randomised controlled trials)

1. Was the allocation sequence randomly generated?
Low risk of bias:
The investigators describe a random component in the sequence generation process, such as referring to a random number table; using a computer random number generator; tossing a coin; shuffling cards or envelopes; throwing dice; drawing lots.
High risk of bias:
The investigators describe a non-random component in the sequence generation process. Usually, the description would involve some systematic, non-random approach, for example, sequence generated by odd or even date of birth; sequence generated by some rule based on date (or day) of admission; sequence generated by some rule based on hospital or clinic record number.
Unclear:
Insufficient information about the sequence generation process to permit judgement of low or high risk of bias.

2. Was the treatment allocation adequately concealed?
Low risk of bias:
Participants and investigators enrolling participants could not foresee assignment because one of the following, or an equivalent method, was used to conceal allocation: central allocation (including telephone, web-based and pharmacy-controlled randomisation); sequentially numbered drug containers of identical appearance; sequentially numbered, opaque, sealed envelopes.
High risk of bias:
Participants or investigators enrolling participants could possibly foresee assignments and thus introduce selection bias, such as allocation based on using an open random allocation schedule (e.g. a list of random numbers); assignment envelopes used without appropriate safeguards (e.g. if envelopes were unsealed or non-opaque or were not sequentially numbered); alternation or rotation; date of birth; case record number; any other explicitly uncontrolled procedure.

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Unclear:
Insufficient information to permit judgement of low or high risk of bias. This is usually the case if the method of concealment is not described or is not described in sufficient detail to allow a definitive judgement, for example, if the use of assignment envelopes is described, but it remains unclear whether envelopes were sequentially numbered, opaque and sealed.

3. Blinding (participants, personnel and outcome assessors) - was knowledge of the allocated interventions adequately prevented during the study?

Low risk of bias:
Any one of the following:
- No blinding, but the review authors judge that the outcome and the outcome measurement are not likely to be influenced by lack of blinding.
- Blinding of participants and key study personnel ensured, and unlikely that the blinding could have been broken.
- Either participants or some key study personnel were not blinded, but outcome assessment was blinded and the non-blinding of others is unlikely to introduce bias.

High risk of bias:
Any one of the following:
- No blinding or incomplete blinding, and the outcome or outcome measurement is likely to be influenced by lack of blinding.
- Blinding of key study participants and personnel attempted, but likely that the blinding could have been broken.
- Either participants or some key study personnel were not blinded, and the non-blinding of others is likely to introduce bias.

Unclear:
Any one of the following:
- Insufficient information to permit judgement of low or high risk of bias.
- The study did not address this outcome.

4. Were incomplete outcome data adequately addressed?

Low risk of bias:
Any one of the following:
- No missing outcome data.
- Reasons for missing outcome data unlikely to be related to true outcome (for survival data, censoring unlikely to be introducing bias).
- Missing outcome data balanced in numbers across intervention groups, with similar reasons for missing data across groups.
- For dichotomous outcome data, the proportion of missing outcomes compared with observed event risk not enough to have a clinically relevant impact on the intervention effect estimate.
- For continuous outcome data, plausible effect size (difference in means or standardised difference in means) among missing outcomes not enough to have a clinically relevant impact on observed effect size.
- Missing data have been imputed using appropriate methods.

High risk of bias:
Any one of the following:
- Reason for missing outcome data likely to be related to true outcome, with imbalance in numbers or reasons for missing data across intervention groups.
- For dichotomous outcome data, the proportion of missing outcomes compared with observed event risk enough to induce clinically relevant bias in intervention effect estimate.
- For continuous outcome data, plausible effect size (difference in means or standardised difference in means) among missing outcomes enough to induce clinically relevant bias in observed effect size.
- ‘As-treated’ analysis done with substantial departure of the intervention received from that assigned at randomisation.
- Potentially inappropriate application of simple imputation.

Unclear:
Any one of the following:
- Insufficient reporting of attrition/exclusions to permit judgement of low or high risk of bias (e.g. number randomised not stated, no reasons for missing data provided).
- The study did not address this outcome.

5. Are reports of the study free of the suggestion of selective outcome reporting?
Low risk of bias:
Any one of the following:
- The study protocol is available and all of the study’s prespecified (primary and secondary) outcomes that are of interest in the review have been reported in the prespecified way.
- The study protocol is not available but it is clear that the published reports include all expected outcomes, including those that were prespecified (convincing text of this nature may be uncommon).

High risk of bias:
Any one of the following:
- Not all of the study’s prespecified primary outcomes have been reported.
- One or more primary outcomes are reported using measurements, analysis methods or subsets of the data (e.g. subscales) that were not prespecified.
- One or more reported primary outcomes were not prespecified (unless clear justification for their reporting is provided, such as an unexpected adverse effect).
- One or more outcomes of interest in the review are reported incompletely so that they cannot be entered in a meta-analysis.
- The study report fails to include results for a key outcome that would be expected to have been reported for such a study.

Unclear:
Insufficient information to permit judgement of low or high risk of bias. It is likely that the majority of studies will fall into this category.

6. Other sources of potential bias
Low risk of bias:
The study appears to be free of other sources of bias.

High risk of bias:
There is at least one important risk of bias. For example, the study:
- had a potential source of bias related to the specific study design used; or
- has been claimed to have been fraudulent; or
- had some other problem.

Unclear:
There may be a risk of bias, but there is either:
- insufficient information to assess whether an important risk of bias exists; or
- insufficient rationale or evidence that an identified problem will introduce bias.

Appendix 3. 'Risk of bias' assessment (cluster randomised controlled trials)
In cluster randomised trials, particular biases to consider include: (i) recruitment bias; (ii) baseline imbalance; (iii) loss of clusters; (iv) incorrect analysis; and (v) comparability with individually randomised trials.

(i) Recruitment bias can occur when individuals are recruited to the trial after the clusters have been randomised, as the knowledge of whether each cluster is an ‘intervention’ or ‘control’ cluster could affect the types of participants recruited.

(ii) Cluster randomised trials often randomise all clusters at once, so lack of concealment of an allocation sequence should not usually be an issue. However, because small numbers of clusters are randomised, there is a possibility of chance baseline imbalance between the randomised groups, in terms of either the clusters or the individuals. Although not a form of bias as such, the risk of baseline differences can be reduced by using stratified or pair-matched randomisation of clusters. Reporting of the baseline comparability of clusters, or statistical adjustment for baseline characteristics, can help reduce concern about the effects of baseline imbalance.

(iii) Occasionally complete clusters are lost from a trial, and have to be omitted from the analysis. Just as for missing outcome data in individually randomised trials, this may lead to bias. In addition, missing outcomes for individuals within clusters may also lead to a risk of bias in cluster randomised trials.

(iv) Many cluster randomised trials are analysed by incorrect statistical methods, not taking the clustering into account. Such analyses create a ‘unit of analysis error’ and produce over-precise results (the standard error of the estimated intervention effect is too small) and P values that are too small. They do not lead to biased estimates of effect. However, if they remain uncorrected, they will receive too much weight in a meta-analysis.

(v) In a meta-analysis including both cluster and individually randomised trials, or including cluster randomised trials with different types of clusters, possible differences between the intervention effects being estimated need to be considered. For example, in a vaccine trial of infectious diseases, a vaccine applied to all individuals in a community would be expected to be more effective than if the vaccine...
was applied to only half of the people. Another example is provided by discussion of a Cochrane review of hip protectors (Hahn 2005). The cluster trials showed large positive effect whereas individually randomised trials did not show any clear benefit. One possibility is that there was a ‘herd effect’ in the cluster randomised trials (which were often performed in nursing homes, where compliance with using the protectors may have been enhanced). In general, such ‘contamination’ would lead to underestimates of effect. Thus, if an intervention effect is still demonstrated despite contamination in those trials that were not cluster randomised, a confident conclusion about the presence of an effect can be drawn. However, the size of the effect is likely to be underestimated. Contamination and ‘herd effects’ may be different for different types of cluster.

**Contributions of Authors**

Gill Norman: conceived and designed the review; checked the quality of data extraction; undertook and checked quality assessment; performed part of data analysis or interpretation; performed statistical analysis; completed the first draft of the review; approved the final version prior to submission.

Jo Dumville: conceived, designed and coordinated the review; extracted data; checked the quality of data extraction; analysed or interpreted data; undertook and checked quality assessment; performed statistical analysis; completed the first draft of the review; approved the final version prior to submission; secured funding; wrote to study authors / experts / companies; and is a guarantor of the review.

Devi Prasad Mohapatra: made an intellectual contribution to the review, advised on the review, and approved the final version of the review prior to submission.

Gemma Owens: extracted data; checked the quality of data extraction; undertook quality assessment; performed part of writing and editing the review; approved the final version prior to submission; advised on the review and wrote to study authors / experts / companies.

Emma Crosbie: checked the quality of statistical analysis; performed part of writing and editing the review; approved the final version prior to submission; and advised on the review.

**Contributions of editorial base:**

Julie Bruce: edited the review, advised on methodology, interpretation and content. Approved the final review prior to submission.

Sally Bell-Syer and Gill Rizzello: coordinated the editorial process. Advised on interpretation and content. Edited the review.

Rocio Rodriguez-Lopez: designed the search strategy. Reetu Child edited the search methods section and ran the searches.

**Declarations of Interest**

Gill Norman: my employment at the University of Manchester is funded by NIHR and focuses on high priority Cochrane reviews in the prevention and treatment of wounds.

Jo Dumville: nothing to declare.

Devi Prasad Mohapatra: nothing to declare.

Gemma Owens: is a recipient of an MCRC Clinical Research Training Fellowship and previously received a Wellbeing of Women Entry Level Scholarship.

Emma Crosbie: is a Scientific Editor for BJOG, has received funding from an NIHR Clinician Scientist Award, the HTA, Wellbeing of Women/the Wellcome Trust and Central Manchester University Hospitals NHS Foundation Trust. She is an employee of the University of Manchester.
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External sources
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DIFFERENCES BETWEEN PROTOCOL AND REVIEW

We have amended the 'Summary of findings' section in the methods. The changes we made included the use of optimal information size (OIS) to inform the GRADE assessment of imprecision.