# Table of Contents

**Abstract** .................................................................................................................................................................................. 1
**Plain Language Summary** .......................................................................................................................................................... 2
**Background** ................................................................................................................................................................................ 4
**Objectives** .................................................................................................................................................................................. 5
**Methods** .................................................................................................................................................................................. 5
**Results** .................................................................................................................................................................................... 7
  - Figure 1. ...................................................................................................................................................................................... 8
  - Figure 2. ...................................................................................................................................................................................... 9
**Discussion** ................................................................................................................................................................................ 18
**Authors' Conclusions** ............................................................................................................................................................. 19
**Acknowledgements** ............................................................................................................................................................... 19
**References** ............................................................................................................................................................................... 20
**Characteristics of Studies** .......................................................................................................................................................... 29
**Data and Analyses** ................................................................................................................................................................. 58
  - Analysis 1.1. Comparison 1 basic hand hygiene versus alcohol rub, Outcome 1 SSI. ................................................................. 59
  - Analysis 2.1. Comparison 2 chlorhexidine versus iodine, Outcome 1 CFUs. ............................................................................ 59
  - Analysis 3.1. Comparison 3 chlorhexidine versus iodine plus triclosan, Outcome 1 CFUs. ....................................................... 60
  - Analysis 4.1. Comparison 4 alcohol rub versus other alcohol rub, Outcome 1 CFUs. ................................................................. 61
  - Analysis 5.1. Comparison 5 scrub versus alcohol-only rub, Outcome 1 SSI. ........................................................................... 61
  - Analysis 6.1. Comparison 6 scrub versus alcohol rub, Outcome 1 SSI. .................................................................................. 62
  - Analysis 7.1. Comparison 7 scrub (chlorhexidine) versus alcohol rub + additional ingredient, Outcome 1 CFUs. ................. 62
  - Analysis 8.1. Comparison 8 scrub (povidone iodine) versus alcohol rub + additional ingredient, Outcome 1 CFUs. ........... 63
  - Analysis 9.1. Comparison 9 scrub (chlorhexidine) versus rub + additional ingredient, Outcome 1 CFUs. ............................ 63
  - Analysis 10.1. Comparison 10 scrub (chlorhexidine) versus alcohol rub + additional ingredient, Outcome 1 CFUs. ............ 64
  - Analysis 11.1. Comparison 11 duration - Kappstein (5 minutes versus 3 minutes), Outcome 1 CFUs immediately after antisepsis. ................................................................................................................................. 64
  - Analysis 12.1. Comparison 12 duration - 5 + 3 min versus 3 + 0.5 min with chlorhexidine), Outcome 1 CFUs. ...................... 65
  - Analysis 13.1. Comparison 13 duration - 5 + 3 min versus 3 + 0.5 minutes with iodine), Outcome 1 CFUs. ......................... 65
  - Analysis 14.1. Comparison 14 duration - 5 + 3.5 min versus 3 + 2.5 min chlorhexidine), Outcome 1 CFUs. ......................... 66
  - Analysis 15.1. Comparison 15 scrub versus scrub plus brush, Outcome 1 CFUS. ................................................................. 67
  - Analysis 16.1. Comparison 16 scrub versus scrub plus nail pick, Outcome 1 CFUs. ................................................................. 67
  - Analysis 17.1. Comparison 17 scrub plus brush versus scrub plus nail pick, Outcome 1 CFUs. ................................................. 67
**Additional Tables** ................................................................................................................................................................. 69
**Appendices** ............................................................................................................................................................................ 72
**Feedback** ................................................................................................................................................................................ 76
**What's New** ............................................................................................................................................................................ 77
**History** .................................................................................................................................................................................. 77
**Contributions of Authors** ....................................................................................................................................................... 77
**Declarations of Interest** .......................................................................................................................................................... 78
**Sources of Support** ............................................................................................................................................................... 78
**Differences between Protocol and Review** .................................................................................................................................. 78
**Index Terms** ............................................................................................................................................................................. 78

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*Surgical hand antisepsis to reduce surgical site infection (Review)*

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Surgical hand antisepsis to reduce surgical site infection

Judith Tanner1, Jo C Dumville2, Gill Norman2, Mathew Fortnam3

1School of Health Sciences, University of Nottingham, Nottingham, UK. 2School of Nursing, Midwifery and Social Work, University of Manchester, Manchester, UK. 3Cochrane Wounds Group, University of York, York, UK

Contact: Judith Tanner, School of Health Sciences, University of Nottingham, Queens Medical Centre, Nottingham, NG7 2HA, UK. judith.tanner@nottingham.ac.uk.

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ABSTRACT

Background
Medical professionals routinely carry out surgical hand antisepsis before undertaking invasive procedures to destroy transient microorganisms and inhibit the growth of resident micro-organisms. Antisepsis may reduce the risk of surgical site infections (SSIs) in patients.

Objectives
To assess the effects of surgical hand antisepsis on preventing surgical site infections (SSIs) in patients treated in any setting. The secondary objective is to determine the effects of surgical hand antisepsis on the numbers of colony-forming units (CFUs) of bacteria on the hands of the surgical team.

Search methods
In June 2015 for this update, we searched: The Cochrane Wounds Group Specialized Register; The Cochrane Central Register of Controlled Trials (CENTRAL) (The Cochrane Library); Ovid MEDLINE; Ovid MEDLINE (In-Process & Other Non-Indexed Citations) and EBSCO CINAHL. There were no restrictions with respect to language, date of publication or study setting.

Selection criteria
Randomised controlled trials comparing surgical hand antisepsis of varying duration, methods and antiseptic solutions.

Data collection and analysis
Three authors independently assessed studies for inclusion and trial quality and extracted data.

Main results
Fourteen trials were included in the updated review. Four trials reported the primary outcome, rates of SSIs, while 10 trials reported number of CFUs but not SSI rates. 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.

SSIs
One study randomised 3317 participants to basic hand hygiene (soap and water) versus an alcohol rub plus additional hydrogen peroxide. There was no clear evidence of a difference in the risk of SSI (risk ratio (RR) 0.97, 95% CI 0.77 to 1.23, moderate quality evidence downgraded for imprecision).
One study (500 participants) compared alcohol-only rub versus an aqueous scrub and found no clear evidence of a difference in the risk of SSI (RR 0.56, 95% CI 0.23 to 1.34, very low quality evidence downgraded for imprecision and risk of bias).

One study (4387 participants) compared alcohol rubs with additional active ingredients versus aqueous scrubs and found no clear evidence of a difference in SSI (RR 1.02, 95% CI 0.70 to 1.48, low quality evidence downgraded for imprecision and risk of bias).

One study (100 participants) compared an alcohol rub with an additional ingredient versus an aqueous scrub with a brush and found no evidence of a difference in SSI (RR 0.50, 95% CI 0.05 to 5.34, low quality evidence downgraded for imprecision).

**CFUs**

The review presents results for a number of comparisons; key findings include the following.

Four studies compared different aqueous scrubs in reducing CFUs on hands. Three studies found chlorhexidine gluconate scrubs resulted in fewer CFUs than povidone iodine scrubs immediately after scrubbing, 2 hours after the initial scrub and 2 hours after subsequent scrubbing. All evidence was low or very low quality, with downgrading typically for imprecision and indirectness of outcome. One trial comparing a chlorhexidine gluconate scrub versus a povidone iodine plus triclosan scrub found no clear evidence of a difference—this was very low quality evidence (downgraded for risk of bias, imprecision and indirectness of outcome).

Four studies compared aqueous scrubs versus alcohol rubs containing additional active ingredients and reported CFUs. In three comparisons there was evidence of fewer CFUs after using alcohol rubs with additional active ingredients (mild or very low quality evidence downgraded for imprecision and indirectness of outcome). Evidence from one study suggested that an aqueous scrub was more effective in reducing CFUs than an alcohol rub containing additional ingredients, but this was very low quality evidence downgraded for imprecision and indirectness of outcome.

Evidence for the effectiveness of different scrub durations varied. Four studies compared the effect of different durations of scrubs and rubs on the number of CFUs on hands. There was evidence that a 3 minute scrub reduced the number of CFUs compared with a 2 minute scrub (very low quality evidence downgraded for imprecision and indirectness of outcome). Data on other comparisons were not consistent, and interpretation was difficult. All further evidence was low or very low quality (typically downgraded for imprecision and indirectness).

One study compared the effectiveness of using nail brushes and nail picks under running water prior to a chlorhexidine scrub on the number of CFUs on hands. It was unclear whether there was a difference in the effectiveness of these different techniques in terms of the number of CFUs remaining on hands (very low quality evidence downgraded due to imprecision and indirectness).

**Authors’ conclusions**

There is no firm evidence that one type of hand antisepsis is better than another in reducing SSIs. Chlorhexidine gluconate scrubs may reduce the number of CFUs on hands compared with povidone iodine scrubs; however, the clinical relevance of this surrogate outcome is unclear. Alcohol rubs with additional antiseptic ingredients may reduce CFUs compared with aqueous scrubs. With regard to duration of hand antisepsis, a 3 minute initial scrub reduced CFUs on the hand compared with a 2 minute scrub, but this was very low quality evidence, and findings about a longer initial scrub and subsequent scrub durations are not consistent. It is unclear whether nail picks and brushes have a differential impact on the number of CFUs remaining on the hand. Generally, almost all evidence available to inform decisions about hand antisepsis approaches that were explored here were informed by low or very low quality evidence.

**Plain Language Summary**

Surgical hand antisepsis to reduce surgical site infection

**What are surgical site infections and who is at risk?**

The inadvertent transfer of micro-organisms such as bacteria to a patient’s wound site during surgery can result in a wound infection that is commonly called a surgical site infection (SSI). SSIs are one of the most common forms of health care-associated infections for surgical patients. Around 1 in 20 surgical patients develop an SSI in hospital, and this proportion rises when people go home. SSIs result in delayed wound healing, increased hospital stays, increased use of antibiotics, unnecessary pain and, in extreme cases, the death of the patient, so their prevention is a key aim for health services.

**Why use hand antisepsis prior to surgery?**

There are many different points in the care pathway where prevention of SSIs can take place. This includes antiseptic cleansing of the hands for those who are operating on the patient. Surgical hand antisepsis is the focus of this review. The two most common forms of hand antisepsis involve aqueous scrubs and alcohol rubs. Aqueous scrubs are water-based solutions containing antiseptic ingredients such as chlorhexidine gluconate or povidone iodine. Scrubbing involves wetting the hands and forearms with water, systematically applying an aqueous scrub solution using either hands or sponges, rinsing under running water and then repeating this process. Alcohol solutions containing additional active ingredients are used to perform an ‘alcohol rub’. Surgical teams systematically apply the alcohol rub solutions to their hands and allow it to evaporate. Alcohol is effective against a wide range of bacteria and other micro-organisms. Following hand
antisepsis, operating staff then put on gloves, which provide an important barrier between operating staff and the patient; however, because gloves can become perforated during surgery, it is necessary to have hands as germ-free as possible.

What we found

In June 2015 we searched for as many relevant studies that had a robust design (randomised controlled trials) as we could find and compared different types of hand antisepsis before surgery. We included 14 studies that compared a range of methods for performing surgical hand antisepsis. The two measures used to assess the effectiveness of treatments were the number of cases of SSIs in patients (presented in four included studies) and the number of viable bacteria or fungal cells (known as colony-forming units, or CFUs) on the hand of the person operating before surgery and after surgery (which is a way of counting the bacteria present on the skin surface). It is not clear whether the method of hand antisepsis influences the risk of SSI, as most of the studies were too small and had flaws. There was some evidence that hand antisepsis with chlorhexidine may reduce the number of bacteria on the hands of health professionals compared with povidone iodine. Importantly, we do not know what the number of CFUs on the hands tells us about the likelihood of patients developing SSIs. There was also some evidence that alcohol rubs with additional antiseptic ingredients may reduce CFUs compared with aqueous scrubs.

Up-to-date June 2015
The inadvertent transfer of micro-organisms to patients' wound sites during surgery can result in postoperative surgical site infections (SSIs). SSIs are one of the most common forms of healthcare-associated infections for surgical patients (NICE 2008). Around 5% of surgical patients develop an SSI (NICE 2008), though this incidence can double when surveillance includes active postdischarge follow-up (Leaper 2015). SSIs result in delayed wound healing, increased hospital stays, increased use of antibiotics, unnecessary pain and, in extreme cases, the death of the patient (Plowman 2000).

Micro-organisms that cause SSIs come from a variety of sources within the operating room, including the hands of the surgical team. Members of the surgical team wear sterile gloves to prevent transferring bacteria from their hands to patients. However, gloves can become perforated during surgery, so it is necessary to have hands as germ-free as possible. This is achieved by conducting surgical hand antisepsis immediately before donning sterile gloves prior to commencing surgical or invasive procedures. While handwashing removes transient micro-organisms, surgical hand antisepsis goes a step further to inhibit the growth of resident microorganisms, thereby minimising the risk of a patient developing an SSI (WHO 2009). This is achieved using antiseptic agents that kill and inhibit bacteria, fungi, protozoa and bacterial spores. An ideal antiseptic agent would be fast-acting, persistent (effective for a number of hours), cumulative (repeated exposure inhibits bacterial growth for a number of days), have a broad spectrum of activity and be safe to use. There are several individual components of surgical hand antisepsis, including the pre-wash, the application technique; the use of sponges, brushes or nail picks; the choice of antiseptic solution and the duration of the antisepsis.

Defining terms - scrub and rubs

Several different terms are used when describing surgical hand antisepsis. Antiseptic with running water and an aqueous solution is referred to as a surgical or a traditional scrub. Antiseptic with an alcohol solution is referred to as an alcohol rub or a waterless scrub. In this review, we understand surgical hand antisepsis to encompass both methods of surgical antisepsis: scrubbing and rubbing. The very first antisepsis of the day is referred to as the initial antisepsis. Scrubs or rubs performed thereafter but on the same day are referred to as subsequent antisepses.

Surgical hand antisepsis - current practice

The Association for Perioperative Practice (AfPP) recommends a pre-wash prior to the first antisepsis of the day, when hands are washed with soap or an antimicrobial solution under running water (AfPP 2011). The function of the pre-wash is to remove dirt (organic material). AfPP 2011 then recommends cleaning nails using a pick under running water. Clinicians can then perform antisepsis using either an antimicrobial solution with running water, referred to as a traditional scrub, or an alcoholic rub without water. AfPP 2011 suggests alcohol rubs are more effective in reducing bacteria on the skin but should not be used if there is visible dirt present. The AfPP does not cite any specific antimicrobial solution as being the most effective, but, like many other organisations, recommends that the solution chosen meets the ideal properties for an antimicrobial solution (ACORN 2012; AORN 2010; WHO 2009). These properties are identified by the Centers for Disease Control and Prevention (CDC) as the solution being:

- fast-acting;
- persistent (effective for a number of hours);
- cumulative (repeated exposure inhibits bacterial growth for a number of days);
- having a broad spectrum of activity; and
- safe to use (CDC 2002).

AfPP 2011 recommends a duration of 2 to 5 minutes (depending on manufacturers instructions) for a traditional scrub, but does not provide details on the recommended duration of an alcoholic rub. There is some discrepancy regarding aspects of hand antisepsis between different organisations. For example, the World Health Organization (WHO) and the Australian College of Operating Room Nurses (ACORN) only recommend a pre-wash if hands are visibly dirty (ACORN 2012; WHO 2009), and ACORN 2012 recommends that the first scrub of the day last 5 minutes while subsequent rubs last 3 minutes. Nail brushes no longer appear to be recommended as these can damage skin (ACORN 2012; AfPP 2011; AORN 2010; WHO 2009).

Guidelines for surgical antisepsis also cover topics such as rings, artificial nails and nail polish (AORN 2010; ACORN 2012; AfPP 2011; HIS 2001; Mangram 1999). The impact of these factors on SSI is the focus of another Cochrane review (Arrowsmith 2014). There are concerns that hand antisepsis causes skin damage to staff hands and that some products are more abrasive than others (Larson 1986b). This topic is outside the remit of this review.

Surgical hand antisepsis solutions

Solutions for hand antisepsis are either aqueous (water) based or alcohol based.

Aqueous scrubs

Aqueous scrubs are water based solutions containing active ingredients that are used during traditional handscrubs. The most common solutions contain chlorhexidine gluconate or povidone iodine (see below). Scrubbing involves wetting the hands and forearms with water, systematically applying an aqueous scrub solution using either hands or sponges, rinsing under running water and then repeating the process.

Alcohol rubs

Alcohol-based solutions are used to perform an 'alcohol rub'. Health professionals apply the solution to dry hands and then rub them together systematically before allowing the solution to evaporate. Alcohol rubs do not require water for their application. Some alcohol rub solutions contain additional active antiseptic agents.

Antiseptic agents

Alcohol

Alcohols have little or no residual effect, and the concentration rather than the type of alcohol is thought to be most important in determining its effectiveness (Larson 1995). Alcohol rubs are usually available in preparations of 60% to 90% strength and are effective against a wide range of gram-positive and gram negative bacteria, mycobacterium tuberculosis, and many fungi and viruses. The three main alcohols used are ethanol, isopropanol.
and n-propanol, and some rubs may contain a mixture of these. Compared with other common antiseptic products, alcohol is associated with the most rapid and greatest reduction in microbial counts (Lowbury 1974a), but it does not remove surface dirt as it does not contain surfactants or have a foaming action (Hobson 1998). Alcohol-based solutions usually (but not always) contain additional active ingredients to combine the rapid bacteriocidal effect of alcohol with more persistent chemical activity.

**Iodine and iodophors**

Iodine has mostly been replaced by iodophors, as iodine often causes irritation and discolouring of skin. Iodophors are composed of elemental iodine, iodide or triiodide, and a polymer carrier of high molecular weight (WHO 2009). Combining iodine with various polymers increases the solubility of iodine, promotes sustained release of iodine and reduces skin irritation. Iodophors are effective against a wide range of gram-positive and gram-negative bacteria, mycobacterium tuberculosis, fungi and viruses (Joress 1962). Iodophors contain iodine with a carrier such as polyvinylpyrrolidone (PVP). PVP, also known as povidone, is a polymer that detoxifies and prolongs the activities of drugs. PVP prolongs the activity of iodine by releasing it slowly. A combination of PVP and iodine, known as povidone iodine (PI), is less irritating than earlier solutions of iodine tincture (Joress 1962). Iodophors rapidly reduce transient and colonising bacteria but have little or no residual effect (Larson 1990).

**Chlorhexidine**

Chlorhexidine is a biguanide. It is effective against a wide range of gram-positive and gram-negative bacteria, lipophilic viruses and yeasts (Hibbard 2002a). It is not sporicidal. Although its immediate antimicrobial activity is slower than that of alcohols, it is more persistent because it binds to the outermost layer of skin, the stratum corneum (Larson 1990). Over time, repeated exposure can lead to a cumulative effect where both transient and resident organisms are reduced (Larson 1990). Chlorhexidine gluconate is effective in the presence of blood and other protein-rich biological materials (Hibbard 2002a).

**Quaternary ammonium compounds**

Quaternary ammonium compounds (QACs) are composed of a nitrogen atom linked to four alkyl groups. Alkyl benzalkonium chlorides are the most widely used as antiseptics, though other compounds include benzethonium chloride, cetrimide and cetylpromidium chloride. QACs are primarily bacteriocidal and fungicidal, although they are microbicidal against some organisms at high concentrations. They are more active against gram-positive bacteria than against gram-negative bacilli. QACs have relatively weak activity against mycobacteria and fungi and greater activity against lipophilic viruses. Their antimicrobial activity is adversely affected by the presence of organic material, and they are not compatible with anionic detergents. In 1994, the Food and Drug Administration (FDA) Tentative Final Monograph (TFM) tentatively classified benzalkonium chloride and benzethonium chloride as having insufficient data to classify as safe and effective for use as an antiseptic handwash (WHO 2009).

**Hexachlorophene**

Hexachlorophene is a halophenol compound. It is a slow-acting antiseptic that forms a film over the skin (Crowder 1967). The film retains bacteriostatic properties and is effective against gram-positive bacteria but is less effective with gram-negative bacteria and fungi (Crowder 1967). A report of toxicity in neonates led to restricted usage (Kimbrough 1973), and today, hexachlorophene has mostly been replaced by triclosan.

**Triclosan**

Triclosan (2,4,4′–trichloro-2′-hydroxydiphenyl ether) has been incorporated in detergents (0.4% to 1%) and alcohols (0.2% to 0.5%) used for hygienic and surgical hand antisepsis or preoperative skin disinfection. It inhibits staphylococci, coliforms, enterobacteria and a wide range of gram-negative intestinal and skin flora (Bartzokas 1983). Most strains of pseudomonas are resistant, and triclosan has only fair activity against mycobacterium tuberculosis and poor activity against fungi (Faoagali 1995).

**Chloroxylenol**

Chloroxylenol, also known as para-chloro-meta-xyleneol (PCMX), is a halogen-substituted phenolic compound. It is not as quick-acting as chlorhexidine or iodophors, and its residual activity is less pronounced than that observed with chlorhexidine gluconate (McDonnell 1999). In 1994, the FDA TFM tentatively classified chloroxylenol as having insufficient data to classify as safe and effective (WHO 2009).

**OBJECTIVES**

To assess the effects of surgical hand antisepsis on preventing surgical site infections (SSIs) in patients treated in any setting. The secondary objective is to determine the effects of surgical hand antisepsis on the number of bacteria colony-forming units (CFUs) present on the hands of the surgical team.

**METHODS**

Criteria for considering studies for this review

**Types of studies**

All published and unpublished randomised controlled trials (RCTs) of surgical hand antiseptic techniques were included. Controlled clinical trials were to be considered in the absence of RCTs. Two possible units of randomisation were considered: the scrub team or individual members of the scrub team.

**Types of participants**

All members of the scrub team or personnel working within the operating theatre or day case setting. The SSI outcome is measured in participants who have undergone surgery.

**Types of interventions**

This review included comparisons of the following with each other and/or placebo and/or no antiseptic:

- Surgical hand antiseptic;
- Aqueous scrub solutions;
- Alcohol rubs;
- Alcohol rubs containing additional active ingredients;
- Surgical hand antiseptic of different durations;
- Surgical hand antiseptic using different equipment (e.g. brush, sponge, nail pick).
Types of outcome measures

Primary outcomes
Occurrence of postoperative SSI, as defined by the CDC (Mangram 1999) or the study authors. We did not differentiate between superficial and deep-incisional infection.

Secondary outcomes
Number of bacterial CFUs found on the hands of the surgical team.

Search methods for identification of studies

Electronic searches
We describe the search methods of the original version and first update of this review in Appendix 1.

For this first update we searched:
The Cochrane Wounds Group Specialized Register (searched 10 June 2015);
The Cochrane Central Register of Controlled Trials (CENTRAL) (The Cochrane Library 2015, Issue 6);
Ovid MEDLINE (1946 to 9 June 2015);
Ovid MEDLINE - In-Process & Other Non-Indexed Citations (9 June 2015);
Ovid EMBASE (1974 to 9 June 2015);
EBSCO CINAHL (1982 to 10 June 2015).

The following search strategy was used in the Cochrane Central Register of Controlled Trials (CENTRAL):

#1 MeSH descriptor Surgical Wound Infection explode all trees
#2 MeSH descriptor Surgical Wound Dehiscence explode all trees
#3 surg* NEAR/5 infect*:ti,ab,kw
#4 surg* NEAR/5 wound*:ti,ab,kw
#5 surg* NEAR/5 site*:ti,ab,kw
#6 surg* NEAR/5 incision*:ti,ab,kw
#7 surg* NEAR/5 dehiscen*:ti,ab,kw
#8 ([post-operative or postoperative] NEAR/5 (wound NEXT infection*)):ti,ab,kw
#9 MeSH descriptor Preoperative Care explode all trees
#10 MeSH descriptor Perioperative Care explode all trees
#11 ([preoperative or pre-operative] NEXT care*:ti,ab,kw
#12 (#1 OR #2 OR #3 OR #4 OR #5 OR #6 OR #7 OR #8 OR #9 OR #10 OR #11)
#13 MeSH descriptor Skin explode all trees
#14 MeSH descriptor Antisepsis explode all trees
#15 (#13 AND #14)
#16 antiseptic*:ti,ab,kw
#17 MeSH descriptor Anti-Infective Agents, Local explode all trees
#18 MeSH descriptor Soaps explode all trees
#19 MeSH descriptor Povidone-Iodine explode all trees
#20 MeSH descriptor Iodophors explode all trees
#21 MeSH descriptor Chlorhexidine explode all trees
#22 MeSH descriptor Alcohols explode all trees
#23 MeSH descriptor Detergents explode all trees
#24 (iodophor* or povidone-iodine or betadine or chlorhexidine or triclosan or hexachlorophene or benzalkonium or alcohol or alcohols or antiseptic* or soap* or detergent*):ti,ab,kw
#25 MeSH descriptor Disinfection explode all trees
#26 MeSH descriptor Disinfectants explode all trees
#27 (#25 OR #26)
#28 (#13 AND #27)
#29 (skin NEAR/5 disinfect*):ti,ab,kw
#30 (#15 OR #16 OR #17 OR #18 OR #19 OR #20 OR #21 OR #22 OR #23 OR #24 OR #28 OR #29)
#31 MeSH descriptor Handwashing explode all trees
#32 MeSH descriptor Hand explode all trees
#33 ("hand" or "hands" or handwash* or (hand NEXT wash*) or (surgical NEXT scrub*)):ti,ab,kw
#34 (#31 OR #32 OR #33)
#35 (#12 AND #30 AND #34)

The search strategies for Ovid MEDLINE, Ovid EMBASE and EBSCO CINAHL are available in Appendix 2, Appendix 3 and Appendix 4, respectively. 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 terms developed by the UK Cochrane Centre (Lefebvre 2011). We combined the CINAHL search with the trial filters developed by the Scottish Intercollegiate Guidelines Network (SIGN) (SIGN 2011). There were no restrictions with respect to language, date of publication or study setting.

Searching other resources
We handsearched the bibliographies of all retrieved and relevant publications identified by these strategies for further studies.

Data collection and analysis

Selection of studies
For the original review, three authors independently assessed the titles and abstracts of potentially relevant studies identified through the search strategy, retrieving the full text of all studies that potentially met the criteria. If it was unclear from the title or abstract whether a study met the criteria or there was a disagreement over the eligibility, we retrieved the full text of the study. The three authors then decided independently whether or not to include the studies. There were no disagreements among authors regarding which studies to include.

For the update, two authors independently assessed all titles and abstracts using the same methods, seeking assistance from translators where necessary. Again, there were no disagreements among authors about which studies to include.

Data extraction and management
We piloted a standardised data extraction form, and two authors independently used the finalised version to extract the following data from studies:

Trial data extracted
- Duration of surgical antisepsis
- Antiseptic solution used
- Equipment used (e.g. brush, sponge, nail pick)
- Role of the person carrying out the hand antisepsis, for example, scrub nurse or surgeon
- Scrub history of the person scrubbing, for example, initial or subsequent scrub
• Surgical specialty, for example, orthopaedics, ophthalmics, urology, etc.
• Type of surgical procedure: elective or emergency
• Duration of surgical procedure
• Surgical glove material
• Size of groups
• Method of SSI detection
• Duration of follow-up

**Trial outcomes**

- Number of SSIs
- Number of CFUs (bacteria) on hands of surgical team

**Assessment of risk of bias in included studies**

Two review authors independently assessed the included studies using the Cochrane tool for assessing risk of bias (Higgins 2011). This tool addresses six specific domains: sequence generation, allocation concealment, blinding, incomplete data, selective outcome reporting and other issues (Appendix 5). 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. For this review, we anticipated that blinding of participants (surgical staff) may not be possible. For this reason the assessment of the risk of detection bias focused on whether trials reported blinded outcome assessment (because wound infection can be a subjective outcome, it can be at high risk of measurement bias when outcome assessment is not blinded).

We 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.

**Data synthesis**

Data were entered into Review Manager software (RevMan 2014). Continuous outcomes (i.e. CFUs) were reported as mean differences (MDs) with 95% confidence intervals (CI). Dichotomous outcomes (i.e. SSIs) were presented as risks ratio (RR) with 95% CI. We reported findings narratively and considered pooling of data after exploring clinical and statistical heterogeneity. We examined clinical heterogeneity by looking at the type of intervention, the participant population and the type of surgery. For assessment of statistical and related heterogeneity we used I^2 values (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, may mean a low level of heterogeneity Higgins 2003, and values of more than 75%, or more, indicate very high heterogeneity.

**Handling of data where the appropriateness of the analysis reported in the paper was unclear**

Where the trial had a cross-over design or was cluster-randomised but the analysis did not appear to take this into account, we reported the available raw data (e.g. mean values) as well as the effect estimate calculated in the paper and discussed the likely effect of an incorrect analysis on the effect estimate.

**‘Summary of findings’ tables**

In the update, in line with current Cochrane methods, we planned to present the main results of the review in ‘Summary of findings’ tables where we had pooled data. 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 for each comparison.

- SSI events
- Number of CFUs

Where we did not pool data, we decided to conduct the GRADE assessment for each comparison and present this narratively within the Results section without the presentation of separate ‘Summary of findings’ tables.

**RESULTS**

**Description of studies**

Also see Characteristics of included studies and Characteristics of excluded studies.

**Results of the search**

The search for this update took place in June 2015 and yielded 274 abstracts. We obtained 18 of these as full-text records for further assessment, subsequently excluding 14 (see Excluded studies) and including four (Al-Naami 2009; Nthumba 2010; Tanner 2009; Vergara-Fernandez 2010). The addition of these four new studies to the 10 studies in the previous version of the review brought the total number of studies included in this update to 14.

Over the life of the review, we have made attempts to contact seven authors to obtain further information (Gupta 2007, Hajipour 2006; Herruzo 2000; Kappstein 1993; Pereira 1997; Pietsch 2001; Sensoz 2003). Five authors responded (Hajipour 2006; Herruzo 2000; Kappstein 1993; Pereira 1997; Sensoz 2003). We included Hajipour 2006, Herruzo 2000, Kappstein 1993, Pereira 1997 and Pietsch 2001 in the review. We also included Gupta 2007, although we have not carried out any independent analyses on their findings. The update identified one study (from a bibliographic search) that is awaiting assessment pending further information (Chen 2012).

**Included studies**

We present an overview of included studies and comparisons in Table 1.
We identified and included 14 eligible trials in this review. Four trials reported the primary outcome, namely SSI (Al-Naami 2009; Nthomba 2010; Parienti 2002; Vergara-Fernandez 2010). The remaining 10 trials reported the number of CFUs (on the hands of the surgical team), which is a surrogate outcome that is thought to give an impression of the likelihood of infection. All 14 trials took place in operating departments, with 8 studies involving surgery and 6 testing interventions in surgical staff but without surgery taking place. Of the included studies, four compared different durations of scrubs or rubs (Kappstein 1993; Pereira 1990; Pereira 1997; Wheelock 1997).

Some of the included studies had complicated designs: Parienti 2002 was an equivalence, cluster, cross-over trial where the unit of randomisation was the surgical service. Each surgical service carried out one intervention for one month and then switched to the alternative intervention the following month. Nthumba 2010 also used a cluster, cross-over design, whereby operating theatres were allocated to an intervention with cross-over every two months. We also considered Hajipour 2006 to be a cluster trial, as it randomised four surgeons to different antisepsis methods which they used prior to surgery on multiple participants. Seven other studies (Gupta 2007, Herruzo 2000, Kappstein 1993, Pereira 1990; Pereira 1997; Pietsch 2001; Wheelock 1997) were also cross-over trials.

**Definition of scrub procedure**

Six trials gave detailed protocols for their antisepsis techniques (Furukawa 2005; Parienti 2002; Pereira 1990; Pereira 1997; Tanner 2009; Wheelock 1997). Authors of eight trials reported using a brush or sponge (Gupta 2007; Furukawa 2005; Herruzo 2000; Parienti 2002; Pereira 1990; Pereira 1997; Vergara-Fernandez 2010), while Tanner 2009 compared a nail brush and nail pick. Parienti 2002 and Wheelock 1997 stated that antisepsis protocols met with national guidelines. Seven of the trials employed a 'supervisor' to observe compliance with the antisepsis protocol (Furukawa 2005; Nthumba 2010; Parienti 2002; Pereira 1990; Pereira 1997; Tanner 2009; Wheelock 1997). Three trials presented minimal details of the antisepsis protocol (Gupta 2007; Hajipour 2006; Nthumba 2010), and the remaining five trials did not comment on antisepsis techniques (Al-Naami 2009; Herruzo 2000; Kappstein 1993; Pietsch 2001; Vergara-Fernandez 2010).

**Excluded studies**

See Characteristics of excluded studies

**Risk of bias in included studies**

We summarise our ‘Risk of bias’ assessment in Figure 1, Figure 2 and in the ‘Risk of bias’ assessment tables. Overall, most studies were at unclear risk or high risk of bias for one of the following: selection bias, detection bias and attrition bias. Only two studies were at low risk of bias for all these (Nthumba 2010; Tanner 2009).

**Figure 1.** Risk of bias graph: review authors’ judgements about each risk of bias item presented as percentages across all included studies.
Figure 2. Risk of bias summary: review authors' judgements about each risk of bias item for each included study.
Surgical hand antisepsis to reduce surgical site infection (Review)

We included four studies to be at low risk of detection bias, as they blinded outcome assessors (Al-Naami 2009; Nthumba 2010; Hajipour 2006; Tanner 2009). We considered two studies to be at high risk of detection bias (Parienti 2002; Wheelock 1997).

We also judged two studies to be at high risk of attrition bias. Pereira 1997 reported that 9/32 randomised members of staff failed to complete with no reasons given for withdrawal. Al-Naami 2009 excluded 100 of 600 patients from the analysis for reasons such as their condition was revised on subsequent histopathological examination, they had incomplete forms, or they "failed follow-up".

We judged several studies to be at unclear risk of other bias due to uncertainty as to whether a cluster-randomised or cross-over design had been taken into account in the analysis. Parienti 2002 and Hajipour 2006 were at unclear risk of other bias as a cluster design was detected but it did not seem that clustering had been taken into account in the analyses. Herruzo 2000, Kappstein 1993, Pereira 1990, Pereira 1997; Wheelock 1997 were at unclear risk because it appeared that the crossover designs had not been taken into consideration in the analyses.

Effects of interventions

We included 14 trials in this review. In total, the trials evaluated nine basic comparisons related to the type (i.e. scrub or rub, active ingredients), duration and tools used for surgical hand antisepsis.

- **Comparison 1**: basic hand hygiene versus alcohol rub containing additional active ingredients (Nthumba 2010).
- **Comparison 2**: different aqueous scrub solutions: chlorhexidine gluconate versus povidone iodine (Furukawa 2005; Herruzo 2000; Pereira 1990; Pereira 1997).
- **Comparison 3**: comparison of different alcohol rubs containing additional active ingredients (Gupta 2007; Pereira 1997).
- **Comparison 4**: aqueous scrubs versus alcohol-only rubs (Al-Naami 2009).
- **Comparison 5**: aqueous scrubs versus alcohol rubs containing additional active ingredients (Gupta 2007; Herruzo 2000; Parienti 2002; Pietsch 2001; Vergara-Fernandez 2010).
- **Comparison 6**: duration of surgical antisepsis (Kappstein 1993; Pereira 1990; Pereira 1997; Wheelock 1997).
- **Comparison 7**: surgical hand antisepsis using a nail pick versus surgical hand antisepsis not using a nail pick (Tanner 2009).
- **Comparison 8**: surgical hand antisepsis using a brush versus surgical hand antisepsis not using a brush (Tanner 2009).
- **Comparison 9**: surgical hand antisepsis using a nail pick versus surgical hand antisepsis using a brush (Tanner 2009).

**Comparison 1**: basic hand hygiene versus alcohol rub containing additional active ingredients (1 study)

Basic hand hygiene (soap and water) compared with 75% isopropyl alcohol plus 0.125% hydrogen peroxide.
(Hibiclens) or 7.5% povidone iodine (Betadine) using a 3 minute initial and 30 second subsequent scrub. Control of the order of interventions was through a Latin square design. Investigators took hand bacterial samples immediately after the initial scrub, 2 hours after the initial scrub and 2 hours after the subsequent scrub. Although the study had a cross-over design, it did not appear that this was taken into consideration in the analysis, thus 95% CIs may be overestimated.

Furukawa 2005 compared 4% chlorhexidine gluconate (Hibiscrub) with 7.5% povidone iodine (Isodine) using a 3 minute scrub. Twenty-two operating room nurses were randomised to one of the two intervention groups. Each nurse took part only once. The nurses did not take part in any actual surgery.

Herruzo 2000 randomised 154 members of surgical teams and compared three intervention groups: relevant to this comparison was a 3 minute scrub of either aqueous chlorhexidine gluconate 4% or aqueous povidone iodine 7.5%. The study had a cross-over design, which appears to have been accounted for. The study also reports repeated measures which does not appear to have been accounted for in the analysis.

Pereira 1997 compared 4% chlorhexidine gluconate (Hibiclens) with 5% povidone iodine plus 1% triclosan (Microshield PVP) using a 3 minute initial and 2.5 minute subsequent scrub. Twenty-three operating room nurses were randomised to carry out each of five interventions for one week each. The order of interventions was controlled through a Latin square design. Participants did not take part in any actual surgery. Investigators took hand bacterial samples immediately after the first antisepsis, 2 hours after the first antisepsis and 2 hours after the subsequent antisepsis. Although the study had a cross-over design, it did not appear that trialists took this into consideration in the analysis, thus the 95% CI may be overestimated.

Outcomes

Surgical site infection

Not reported

Number of colony forming units (CFUs)

Chlorhexidine gluconate compared with povidone iodine

Pereira 1990 compared 4% chlorhexidine gluconate (Hibiclens) with 7.5% povidone iodine (Betadine) using a 5 minute initial and 3 minute subsequent scrub. There was some evidence that scrubbing with chlorhexidine might be more effective than with povidone iodine in reducing the number of CFUs on the hand immediately after scrubbing (MD − 0.34, 95% CI − 0.64 to − 0.04;), 2 hours after the initial scrub (MD − 0.75, 95% CI − 1.06 to − 0.44) and 2 hours after the subsequent scrub (MD − 1.10, 95% CI − 1.42 to − 0.78; Analysis 2.1).

Because the analysis did not account for the effects of paired data resulting from the cross-over design, the study may have overestimated the uncertainty of the effect estimate; the correct 95% CIs for the estimate may be narrower than those reported here. However, the study also used a repeated measures design which the analysis did not account for; this could lead to an underestimation of the uncertainty. The interaction of these two factors makes the true confidence intervals unclear, so we have downgraded the evidence twice due to imprecision.

Very low quality evidence due to imprecision and indirectness of outcome. Downgraded twice for imprecision due to two analytical issues which mean precision estimates may change upon correct analysis of data, and downgraded twice for indirectness as CFU is a surrogate outcome and because the intervention was used in the absence of surgery being conducted. A GRADE assessment of very low quality evidence means any estimate of effect is very uncertain.

Pereira 1990 compared 4% chlorhexidine gluconate (Hibiclens) with 7.5% povidone iodine (Betadine) using a 3 minute initial and 30 second subsequent scrub. There was evidence that chlorhexidine may be more effective than povidone iodine in reducing the number of CFUs immediately post scrubbing (MD − 0.17, 95% CI − 0.28 to − 0.06), 2 hours after the initial scrub (MD − 0.41, 95% CI − 0.72 to − 0.10) and 2 hours after the subsequent scrub (MD − 0.65, 95% CI − 0.93 to − 0.37 Analysis 2.1).

Very low quality evidence due to imprecision and indirectness of outcome (as above).

In Furukawa 2005, there were fewer CFUs in the chlorhexidine gluconate group after scrubbing (MD − 2.40, 95% CI − 3.26 to − 1.54; Analysis 2.1).

Low quality evidence: downgraded twice for indirectness as CFU is a surrogate outcome and because the intervention was used in the absence of surgery being conducted.

In Herruzo 2000, there was evidence that a 3 minute aqueous scrub using chlorhexidine gluconate was more effective in reducing CFUs on hands than a 3 minute aqueous scrub using povidone iodine, both immediately after antisepsis (MD − 48.00, 95% CI − 50.57 to − 45.4) and at the end of a surgical procedure (MD − 132.0, 95% CI − 141.20 to − 122.80; Analysis 2.1). Because the analysis did not account for the repeated measures, the study may have underestimated uncertainty around the effect estimate: the true confidence intervals for the estimate may be wider than those reported here.

Low quality evidence; downgraded once as precision estimates may change upon correct analysis of data and downgraded once due to indirectness as CFU is a surrogate outcome.

We considered pooling the three studies (four comparisons) (Furukawa 2005; Herruzo 2000; Pereira 1990); however, in light of the high degree of statistical heterogeneity for each possible comparison (ranging from 100% to 56%) and the differences in interventions noted, we did not undertake this.

Chlorhexidine gluconate compared with povidone iodine plus triclosan

Pereira 1997 reported no evidence of a difference in CFUs immediately after the first antisepsis (MD − 0.38, 95% CI − 0.84 to 0.08) (Analysis 3.1) and 2 hours after the first antisepsis (MD − 0.38, 95% CI − 0.90 to 0.14) (Analysis 3.2). The trial found a difference in favour of chlorhexidine 2 hours after the subsequent antisepsis (MD − 0.69, 95% CI − 1.13 to − 0.25) (Analysis 3.1).

Because the analysis did not account for the effects of paired data resulting from the cross-over design or for the effect of using repeated measures, the true confidence intervals for the estimate are uncertain.
Comparison 2 Summary: Comparison of different aqueous scrub solutions - chlorhexidine gluconate compared with povidone iodine

Data from four trials (five comparisons) of chlorhexidine-containing aqueous scrub solutions with povidone iodine containing solutions (all having initial longer duration of use followed by subsequent use) suggest that chlorhexidine containing agents may reduce the numbers of CFU on the hands to a greater extent than povidone iodine containing solutions. However, overall this is low or very low quality evidence, and the number of CFUs is a surrogate outcome for SSI. Some of the studies included appeared to have been incorrectly analysed and one was at high risk of attrition bias. No trials reported SSI events so there is no evidence to link the number of CFUs to clinical outcomes.

Comparison 3: comparison of different alcohol rubs containing additional active ingredients (2 trials)

Two trials compared alcohol rubs containing additional active ingredients (Gupta 2007; Pereira 1997).

Gupta 2007 compared three 2 ml aliquots of 1% chlorhexidine gluconate in 61% ethyl alcohol (Avagard) against a 3 minute application of zinc pyrithione in 70% ethyl alcohol (Triseptin). The 61% alcohol solution is a waterless product, and the 70% alcohol solution is a water aed product which requires rinsing with water. Eighteen operating room staff used each product for five consecutive days. Testing was carried out immediately before and after antisepcy on day one, and at the end of days two and five.

Pereira 1997 compared 0.5% chlorhexidine gluconate in isopropanol compared with 0.5% chlorhexidine gluconate in ethanol. The alcohol rubs were used immediately after an aqueous scrub (with 4% chlorhexidine gluconate) and also as the subsequent antiseptic agent. The active ingredient in both alcohol rubs was the same (i.e. 0.5% chlorhexidine gluconate), and both preparations had 70% strength alcohol, the only difference being the alcohol (isopropanol versus ethanol). Scrubs lasted for 2 minutes, and the initial and subsequent applications of alcohol rubs lasted for 30 seconds.

Outcomes

Surgical site infection

Not reported

Number of colony forming units (CFUs)

Pereira 1997 did not find a difference in the number of CFUs between the isopropanol- and ethanol-based rubs immediately after the first antisepcy (MD 0.00, 95% CI – 0.57 to 0.57), 2 hours after the first antisepcy (MD 0.07, 95% CI -0.45 to 0.59) or 2 hours after the subsequent antisepcy (MD 0.11, 95% CI – 0.49 to 0.71); see Analysis 4.1. Differences between groups were consistently small and imprecise. As previously noted, the true confidence intervals for the effect estimates is uncertain because of the issues with the analysis.
Comparison 5: aqueous scrubs versus alcohol rubs containing additional active ingredients (6 trials)

Six studies compared traditional scrubs with alcohol rubs containing additional active ingredients (Gupta 2007; Hajipour 2006; Herruzo 2000; Parienti 2002; Pietsch 2001; Vergara-Fernandez 2010). The six trials used different antiseptic solutions, therefore it was not appropriate to perform a meta-analysis. Each trial is considered separately.

Parienti 2002 compared a 5 minute scrub using either 4% povidone iodine (Betadine) or 4% chlorhexidine gluconate (Hibiscrub) versus a 5 minute handrub using 75% propanol-1, propanol-2 with meccetromin ethylsulphate (Sterillium). Participants in the aqueous scrub group could choose between chlorhexidine gluconate or povidone iodine solutions. Participants in the handrubbing group carried out a single handwash for 1 minute with non-antiseptic soap at the start of each day. The entire scrub team in each of six hospitals took part. The trial included 4387 consecutive patients undergoing clean and clean-contaminated surgery and assessed SSI at 30 d using the CDC definition. The study was an equivalence, cluster cross-over trial but did not appear to have accounted for the clustering in the analysis.

Vergara-Fernandez 2010 compared an aqueous scrub with 4% chlorhexidine gluconate (which involved the use of a brush and a sponge) versus a handrub using 61% ethyl alcohol plus 1% chlorhexidine gluconate. Mean duration of the aqueous scrub was 3.9 minutes (SD 1.07), and mean duration of the alcohol rub was 2.0 minutes (SD 0.47). This trial took place in a single institution and involved 400 staff operating on 100 patients undergoing clean and clean-contaminated surgery, who were randomised to the two hand antisepsis groups. Investigators assessed SSI at one month using the CDC definition. Twenty per cent of the included staff had hand samples sent for microbiological examination, and samples were assessed as positive or negative for hand cultures.

Herruzo 2000 compared three intervention groups: chlorhexidine gluconate scrub versus povidone iodine scrub versus an alcohol rub with N-duopropenide. Each scrub or rub lasted 3 minutes. We successfully contacted Herruzo 2000 for additional information regarding sample size. 154 members of the surgical team were randomised for 55 operations. Investigators measured CFUs before antisepsis, immediately after antisepsis and at the end of the surgical procedure.

Pietsch 2001 compared scrubbing using 4% chlorhexidine gluconate (Hibiscrub) with hand rubbing using an alcoholic solution of 45% propanol-2, 30% propanol-1 plus 0.2% ethylhexadecyldimethyl ammonium ethylsulphate (Sterillium). Seventy-five surgeons from one hospital participated in this randomised cross-over trial, using one product for four weeks then switching to the alternative product following a rest week. CFUs were measured before antisepsis, immediately after antisepsis and after the surgical procedure.

Hajipour 2006 compared a 3 minute 4% chlorhexidine gluconate scrub versus a 3 minute chlorhexidine in alcohol rub (Hydrex). We contacted the trial authors, who provided additional study details. Following an aqueous chlorhexidine scrub at the start of each day, four surgeons were randomised to one or other intervention and were evaluated repeatedly in that condition. Testing was carried out using the finger press method at the end of each surgical procedure.

Gupta 2007 compared 7.5% povidone iodine aqueous scrub against two alcohol rubs: the N-duopropenide solution of 1% chlorhexidine gluconate in 61% ethyl alcohol (Avagard) and a 3 minute application of zinc pyrithione in 70% ethyl alcohol (Triseptin). The paper does not provide further details regarding the application of the products. Eighteen operating room staff used each of the three products for five consecutive days. Testing was carried out immediately before and after antisepsis on day one, and at the end of days two and five.

Outcomes

Surgical site infection (SSI)

Aqueous povidone iodine or chlorhexidine gluconate versus 75% propanol-1, propanol-2 plus meccetromin ethylsulphate

Parienti 2002 collected data for 30 days following surgery. There was no clear evidence of a difference in the rates of SSI between aqueous scrub and alcohol rub: 2.5% (53/2135) of participants developed an SSI in the scrub group compared with 2.4% (55/2252) in the handrub group (RR 1.02, 95% CI 0.70 to 1.48) (Analysis 6.1).

Low quality evidence downgraded once due to risk of detection bias and once due to imprecision.

Aqueous chlorhexidine gluconate compared with 61% ethyl alcohol plus 1% chlorhexidine gluconate

Vergara-Fernandez 2010 found no clear evidence of a difference in SSI rates between groups. In total 2% of participants (1/50) had a SSI in the aqueous scrub group compared with 4% (2/50) in the alcohol handrub group (RR 0.50, 95% CI 0.05 to 5.34). The study was small and the resulting 95% CI intervals wide, ranging from a 95% reduction in risk of SSI to a 400% increased risk of SSI (Analysis 6.1).

Low quality evidence downgraded twice due to imprecision.

Number of colony-forming units

Aqueous chlorhexidine gluconate versus N-duopropenide

Herruzo 2000 reported CFU data (log$_{10}$) after antisepsis and after surgery (Analysis 7.1). We were unable to produce an estimate of treatment effect for the review. Using bivariate analysis, Herruzo 2000 reports that N-duopropenide is more effective than chlorhexidine in reducing the number of CFUs on participants’ hands immediately after antisepsis (P value < 0.01) and at the end of a surgical procedure (P value < 0.01); the paper did not provide any further information on estimates.

Very low quality evidence downgraded twice as precision estimates are not available and once due to indirectness of outcome.

Aqueous povidone iodine versus N-duopropenide

Herruzo 2000 reported CFU data (log$_{10}$) after antisepsis and after surgery (Analysis 8.1). We were unable to produce an estimate of treatment effect for the review. Using bivariate analysis, Herruzo 2000 reports that N-duopropenide was statistically significantly more effective than povidone iodine in reducing the number of CFUs on participants hands immediately after antisepsis (P value <0.01) and at the end of a surgical procedure (P value <0.01).
Very low quality evidence downgraded twice as precision estimates are not available and once due to indirectness of outcome.

Aqueous chlorhexidine gluconate versus 45% propanol-2, 30% propanol-1 plus 0.2% ethylhexadecyldimethyl ammonium ethylsulphate (Sterillium)

Pietsch 2001 reported that rubbing using 45% propanol-2, 30% propanol-1 plus 0.2% ethylhexadecyldimethyl ammonium ethylsulphate (Sterillium) was more effective in reducing CFUs on participants’ hands than scrubbing using 4% chlorhexidine gluconate, both immediately after antisepsis (MD − 1.27, 95% CI − 1.23 to − 1.31) and at the end of the surgical procedure (MD − 1.07, 95% CI − 1.03 to − 1.11); Analysis 9.1.

Moderate quality evidence downgraded once due to indirectness of outcome.

Aqueous 4% chlorhexidine gluconate versus 0.5% chlorhexidine gluconate in 70% alcohol

Hajipour 2006 reported finding fewer CFUs following an aqueous scrub than after an alcohol rub (MD − 135.60, 95% CI − 153.39 to − 117.81; Analysis 10.1).

This study had a cluster design, in which each surgeon constituted a cluster, giving two clusters in each trial arm. However, the analysis did not take this into consideration, so the reliability of the effect estimate reported by the authors is uncertain and could be wider than reported.

Very low quality evidence downgraded twice due to potential imprecision as re-analysis of data could increase the confidence intervals and change study conclusions and once due to indirectness of outcome.

Aqueous povidone iodine versus 61% ethyl alcohol and 70% ethyl alcohol

Gupta 2007 did not present sufficient raw data for us to be able to conduct independent statistical analysis, so we contacted the author to request additional data. In the interim we present Gupta 2007’s own analysis. When CFUs were compared collectively from all the sample times, Gupta 2007 reports ‘no statistically significant difference’ between the solutions (P = 0.21). It must be noted that this analysis has not been independently verified and it is unclear if investigators adjusted this analysis to account for the cross-over design.

Very low quality evidence downgraded twice for imprecision as CIs not available and twice for indirectness as we were unable to assess the actual results and the analysis undertaken and because the outcome is a surrogate outcome.

Comparison 5 summary: Aqueous scrubs compared with alcohol rubs containing additional active ingredients

It is unclear if there is a difference in numbers of SSIs between aqueous scrubs and alcohol rubs. The CFU outcome data were varied with two studies finding in favour of the alcohol rubs (moderate and very low quality evidence), one favouring the scrub arm (very low quality evidence) and one study reporting a ‘non-statistically significant difference’ with no other data (very low quality evidence).

Comparison 6: duration of surgical antisepsis (4 trials)

Four trials compared surgical antisepsis of different durations but used different antiseptic agents, which prevented us from pooling results (Kappstein 1993; Pereira 1990; Pereira 1997; Wheelock 1997).

Wheelock 1997 randomised 25 operating room nurses and surgical technologists to either a 2 minute or a 3 minute scrub. After carrying out the trial scrub, and following a one-week washout period in which they remained to undertake scrubbing as part of their usual work, the participants switched to the other intervention. Though the intention of the trial authors was for participants to use aqueous 4% chlorhexidine gluconate (Hibiclens), participants with a history of skin irritation (15/25 participants) used either 2% chlorhexidine gluconate or parachlorometaxylenol (PCMX). CFUs were measured 1 hour after the surgical scrub.

Kappstein 1993 compared a five minute rub with a three minute rub using alcoholic disinfectant. The disinfectant is not identified. Both rubs followed 1 minute handwashes using soap and water. Twenty-four surgeons carried out each of three intervention groups once in a random order. Samples were taken before and immediately after antisepsis.

Pereira 1990 compared a 5 minute initial and 3 minute subsequent scrub with a 3 minute initial and 30 second subsequent scrub using chlorhexidine gluconate. Thirty-four participants were randomly assigned to one of four groups, and each group was assigned to one of four interventions, each lasting one week.

Pereira 1990 also compared a 5 minute initial and 3 minute subsequent scrub with a 3 minute initial and 30 second subsequent scrub using povidone iodine.

Pereira 1997 compared a 5 minute initial and a 3.5 minute subsequent scrub with a 3 minute initial and a 2.5 minute subsequent scrub using 4% chlorhexidine gluconate. Twenty-three operating room nurses were randomised to carry out each of five interventions for one week each.

Outcomes

Surgical site infection (SSI)
Not reported

Number of colony forming units (CFUs)

Three minute scrub versus two minute scrub

Wheelock 1997 presented paired data, which the review authors re-analysed but did not present. There were fewer CFUs on hands immediately after a 3 minute scrub compared with a 2 minute scrub (MD − 0.29, 95% CI − 0.06 to − 0.52).

Very low quality evidence downgraded once due to risk of detection bias and twice due to indirectness, once due to indirectness of outcome and once because the intervention was used in the absence of surgery being conducted.

Five minute rub versus three minute rub (Analysis 11.1)

Kappstein 1993 favoured the 3 minute scrub over the 5 minute scrub when assessed immediately after antisepsis (MD 0.26, 95% CI 0.14 to 0.38; Analysis 11.1). Because the use of a cross-over design...
was not accounted for in the analysis the true confidence intervals may be narrower than those reported here.

**Low quality evidence: downgraded once due to imprecision and once due to indirectness.**

Five minute initial and three minute subsequent scrub versus three minute initial and 30 second subsequent scrub using chlorhexidine (Analysis 12.1)

Pereira 1990 reported no clear difference between groups in the number of CFUs immediately after the initial scrub (MD = −0.19, 95% CI −0.51 to 0.13) or 2 hours after the initial scrub (MD = −0.23, 95% CI −0.52 to 0.06). There was evidence of a difference in CFUs favouring the 5 minute arm 2 hours after the subsequent scrub (MD = −0.58, 95% CI −0.92 to −0.24); see Analysis 12.1.

**Very low quality evidence down graded twice due to imprecision and once due to indirectness of outcome.**

Five minute initial and three minute subsequent scrub compared with a three minute initial and 30 second subsequent scrub using povidone iodine (Analysis 13.1)

In Pereira 1990, there was no clear difference in the number of CFUs at any time point, whether immediately after the initial scrub (MD 0.02, 95% CI −0.28 to 0.32), 2 hours after the initial scrub (MD 0.12, 95% CI −0.21 to 0.45) or 2 hours after the subsequent scrub (MD −0.13, 95% CI −0.37 to 0.11); see Analysis 13.1.

**Very low quality evidence downgraded due to imprecision and indirectness of outcome.**

Five minute initial and three and 30 second subsequent scrub compared with a three minute initial and two and a half minute subsequent scrub using chlorhexidine (Analysis 14.1)

In Pereira 1997, there was no clear difference in the number of CFUs at any time point reported, whether immediately after the initial antisepsis (MD 0.08, 95% CI −0.44 to 0.60), 2 hours after the initial antisepsis (MD −0.19, 95% CI −0.78 to 0.40) or 2 hours after subsequent antisepsis (MD −0.17, 95% CI −0.71 to 0.37); see Analysis 14.1.

**Very low quality evidence due to risk of bias of attrition bias; imprecision and indirectness of outcome.**

**Comparison 6 summary: duration of surgical antisepsis**

Outcome data were only available for CFUs. One study reported evidence of fewer CFUs on hands after using a 3 minute rather than 2 minute chlorhexidine scrub. Another study reported fewer CFUs after a 3 minute alcohol rub compared with a 5 minute alcohol rub; evidence was low quality in both cases. One study reported that 3 minute subsequent scrubs with aqueous chlorhexidine (following initial scrubs) were more effective in reducing the number of CFUs on hands than 30 second subsequent scrubs; this difference was not observed with povidone iodine treatments used in the same way; estimates from this study was classed as being of very low quality. Other comparisons reported no clear differences in number of CFUs.

**Comparison 7: surgical hand antisepsis using a nail pick versus surgical hand antisepsis not using a nail pick (1 trial)**

One three-arm trial compared the effect of surgical hand antisepsis using a nail pick with surgical hand antisepsis alone (Tanner 2009). The study randomised 164 participants to one of three groups. All groups scrubbed with two measured doses of 2 ml aqueous chlorhexidine gluconate 4% (Hibiscrub) for 1 minute per dose; the total scrub time, which was observed and timed, was 2 minutes. One group performed only this surgical hand antisepsis. A second group used a disposable nail pick to clean their nails under running water before the hand antisepsis procedure. The third group used a disposable nail brush to clean their nails under running water before the hand antisepsis procedure. Participants then undertook circulating duties in the operating room for one hour but did not participate in any surgeries.

**Outcomes**

**Surgical site infection**

Not reported

**Number of colony forming units (CFUs)**

There was no clear evidence of a difference between nail pick and no nail pick in the number of CFU detected after one hour on the dominant hands of participants (MD 0.13, 95% CI −0.14 to 0.40; Analysis 15.1).

**Very low quality evidence - downgraded once due to imprecision and once due to indirectness of outcome and further again for indirectness as no surgery was performed.**

**Comparison 8 surgical hand antiseptic using a brush versus surgical hand antiseptic not using a brush (1 trial)**

In the same three-arm trial (Tanner 2009) compared the effect of surgical hand antisepsis and using a brush with surgical hand antisepsis alone. Participants were allocated to groups as described above. There was no clear evidence of a difference between using and not using a brush during hand antisepsis on the number of CFUs (MD 0.24, 95% CI −0.04 to 0.51; Analysis 16.1).

**Very low quality evidence; downgraded once due to imprecision and once due to indirectness of outcome and further again for indirectness as no surgery was performed.**

**Comparison 9: surgical hand antisepsis using a nail pick versus surgical hand antisepsis using a brush (1 trial)**

In the same three-arm trial (Tanner 2009) compared the effect of surgical hand antisepsis using a nail pick with surgical hand antisepsis using a brush. There was no difference in the number of CFUs detected after 1 hour on the dominant hands of participants who used a brush before hand antisepsis compared with those who used a nail pick before hand antisepsis (MD 0.10, 95% CI −0.17 to 0.37; Analysis 17.1).

**Very low quality evidence - downgraded once due to imprecision and once due to indirectness of outcome and further again for indirectness as no surgery was performed.**

**Summary of Comparisons 7 to 9: Surgical hand antisepsis using a nail pick and brush**

There was no clear evidence of a difference in CFUs when a nail pick was compared with a brush or with no pick or brush.

**Brief overview of findings**
## Comparison 1: basic hand hygiene versus alcohol rub containing additional active ingredients

| Evidence | SSI | GRADE ASSESSMENT | CFUs (on hands) | GRADE ASSESSMENT |
|----------|-----|-----------------|----------------|-----------------|
| 1 study  | There was no clear evidence of a difference between treatments: RR 0.97 (95% CI 0.77 to 1.23) (Nthumba 2010) | Moderate quality evidence | Not reported | — |

## Comparison 2: different aqueous scrub solutions: chlorhexidine gluconate versus povidone iodine

| Evidence | SSI | GRADE ASSESSMENT | CFUs (on hands) | GRADE ASSESSMENT |
|----------|-----|-----------------|----------------|-----------------|
| 4 studies | Not reported | — | In 4 comparisons (3 studies: Furukawa 2005; Herruzo 2000; Pereira 1990), there was evidence of lower CFU counts immediately following scrubs with chlorhexidine, and in Pereira 1990, also after subsequent scrubbing. In 1 comparison (1 study) there was no evidence of a difference in the CFU count between an aqueous scrub of chlorhexidine gluconate and an aqueous scrub of povidone iodine plus triclosan (Pereira 1997). | Low quality evidence for Furukawa 2005 and Herruzo 2000 and very low quality evidence for Pereira 1990 and Pereira 1997 | — |

## Comparison 3: comparison of different alcohol rubs containing additional active ingredients

| Evidence | SSI | GRADE ASSESSMENT | CFUs (on hands) | GRADE ASSESSMENT |
|----------|-----|-----------------|----------------|-----------------|
| 2 studies | Not reported | — | 1 study reported small mean difference values with imprecision around estimates at the 3 time points reported (Pereira 1997). We could not verify the findings of 1 study, which study authors reported as not statistically significant (Gupta 2007). | Very low quality evidence | — |

## Comparison 4: aqueous scrubs versus alcohol-only rubs

| Evidence | SSI | GRADE ASSESSMENT | CFUs (on hands) | GRADE ASSESSMENT |
|----------|-----|-----------------|----------------|-----------------|
| 1 study  | RR 0.56 (95% CI 0.23 to 1.34) (Al-Naami 2009) | Very low quality evidence | Not reported | — |

The estimates was imprecise, and it was not possible to rule out an effect in either direction.

## Comparison 5: aqueous scrubs versus alcohol rubs containing additional active ingredients

| Evidence | SSI | GRADE ASSESSMENT | CFUs (on hands) | GRADE ASSESSMENT |
|----------|-----|-----------------|----------------|-----------------|
| 6 studies | 2 studies reported SSI data: RR 1.02, 95% CI 0.70 to 1.48 (Parienti 2002) RR 0.56, 95% CI 0.05 to 5.34 (Vergara-Fernandez 2010) | Low quality evidence | Reported for 5 sub-comparisons based on product type | Moderate quality evidence for Pietsch 2001, and very low quality for Hajipour 2006, Herruzo 2000 and Gupta 2007 | — |
Both estimates were imprecise, and it was not possible to rule out an effect in either direction.

One sub-comparison (1 study) suggested no evidence of a difference between interventions (Gupta 2007).

| Comparison 6: duration of surgical antisepsis | 4 studies | Not reported | 3 min versus 2 min scrub: evidence of fewer CFUs in 3 min scrub group immediately after scrubbing (MD 0.29, 95% CI 0.06 to 0.52; Wheelock 1997) | Low quality evidence for Kappstein 1993 and very low quality evidence for Pereira 1990, Pereira 1997 and Wheelock 1997 |
|---------------------------------------------|----------|--------------|--------------------------------------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------|
|                                            |          |              | 5 min scrub versus 3 min scrub: evidence of fewer CFUs immediately after scrubbing in the 3 min group (MD 0.26, 95% CI 0.14 to 0.38; Kappstein 1993) |
|                                            |          |              | 5 min initial and 3 min subsequent scrub versus a 3 min initial and 30 s subsequent scrub: Chlorhexidine scrub: evidence of fewer CFUs in the longer scrub duration group after the subsequent scrubs (Pereira 1990) |
|                                            |          |              | Povidone iodine scrub: no evidence of a difference in CFUs between groups at any stage (Pereira 1990) |
|                                            |          |              | 5 min initial and 3.5 min subsequent scrub versus a 3 min initial and 2.5 min subsequent scrub using chlorhexidine: no evidence of a difference in CFUs at any stage (Pereira 1997) |
| Comparison 7: surgical hand antisepsis using a nail pick versus surgical hand antisepsis not using a nail pick | 1 study | Not reported | Mean difference in CFUs 1 h after antisepsis was 0.13 (95% CI −0.14 to 0.40; Tanner 2009). No clear evidence of a difference. | Very low quality evidence |
| Comparison 8: surgical hand antisepsis using a brush versus surgical hand antisepsis not using a brush | 1 study | Not reported | Mean difference in CFUs 1 h after antisepsis was 0.24 (95% CI −0.04 to 0.51; Tanner 2009). No clear evidence of a difference. | Very low quality evidence |
| Comparison 9: surgical hand antisepsis using a nail pick versus surgical hand antisepsis using a brush | 1 study | Not reported | Mean difference in CFUs 1 h after antisepsis was 0.10 (95% CI −0.17 to 0.37; Tanner 2009). No clear evidence of a difference. | Very low quality evidence |
**DISCUSSION**

**Summary of main results**

Fourteen trials met the inclusion criteria and were included in this review. Only four trials reported risk of surgical site infection (SSI), the primary outcome for this review (Al-Naami 2009; Nthumba 2010; Parienti 2002; Vergara-Fernandez 2010). These results were inconclusive, and it is not clear whether any method of hand antisepsis is any more effective than another in reducing subsequent SSI. Most of the evidence was of low or very low quality, mainly due to risk of bias and imprecision. Most of the studies were too small to detect a difference in SSI rates even if one existed.

The remaining 10 trials measured the number of CFUs on hands. Where trials were cluster and/or crossover trials they were often not analysed correctly, which may impact on the precision of estimates.

In four comparisons of aqueous scrub solutions containing chlorhexidine versus those containing povidone iodine, (three studies) there was evidence that chlorhexidine-glucanote containing scrubs reduce numbers of CFU on the hands more than povidone iodine (Furukawa 2005; Herruzo 2000; Pereira 1990). The evidence was of low or very low quality. High levels of heterogeneity precluded meta-analysis.

Results of studies comparing alcohol rubs with aqueous scrubs were mixed. In three comparisons (two studies) alcohol rubs containing additional active ingredients were found to be more effective than aqueous scrubs in reducing the number of CFUs on hands (moderate and very low quality evidence) (Herruzo 2000; Pietsch 2001). Gupta 2007 reported no difference between povidone iodine scrubs and ethyl alcohol based rubs of 61% and 70% (very low quality evidence). Hajipour 2006 found chlorhexidine aqueous scrubs more effective than chlorhexidine in alcohol rubs (very low quality evidence).

In terms of the duration of hand antisepsis, there was evidence that a 3 minute scrub reduced the number of CFUs compared to a 2 minute scrub (Wheelock 1997) (very low quality evidence). Data on other comparisons were not consistent, and interpretation is difficult.

One three-arm trial assessed the use of a disposable nail pick or disposable nail brush under running water prior to scrubbing with chlorhexidine and found no clear difference in the number of CFUs (very low quality evidence) (Tanner 2009).

Nthumba 2010, which found no difference in SSIs when comparing surgical hand antisepsis with basic hand hygiene (moderate quality evidence), has interesting implications for practice, which warrant further research.

**Quality of the evidence**

Surgical hand antisepsis is carried out to reduce the number of bacteria on the hands of the surgical team so that in the event of a breach in glove barrier protection, the risk of transferring infections to patients is reduced. Therefore, the most appropriate outcome measure for a study of surgical hand antisepsis is postoperative SSI. Only four studies used this measure (Al-Naami 2009; Nthumba 2010; Parienti 2002; Vergara-Fernandez 2010), while the remaining 10 trials measured CFUs. It is widely assumed that interventions reducing CFUs may impact on incidence of SSIs but there is little evidence to support this assertion. There was considerable variation in the timing of outcome assessment for measuring CFUs as well as some differences in the methods used to obtain samples.

All of the studies included in this review were underpowered in terms of small sample sizes, a limited number of outcome events or both. RCTs need to be adequately powered so that they are able to detect treatment effects of a specified size if they exist, and sample size calculations should be used to help estimate the number of people recruited to a trial. Evidence from limited numbers of participants and/or where limited numbers of events occur can also lead to confidence intervals that are fragile (that is not stable) which is also likely to have been a risk for all comparisons here.

A key issue with the included studies was the incorrect or unclear analysis and presentation of cross-over and cluster data. When these methods are used, correct planning, conduct and reporting are vital to maximise the value of the research. Cross-over studies involve an individual receiving more than one of the treatments of interest, with the randomisation consisting of the sequence in which these are received. The analysis requires statistical tests that take the paired nature of these data into account. If investigators do not use these approaches, they may overestimate standard errors, leading to confidence intervals which are less precise than they might be - that is, there is a risk of a type II error (concluding no significant difference when there is one). Cluster trials, where allocation of treatment is made at a group level and the unit of assessment is at the individual level, require careful planning and analysis to avoid producing artificially small standard errors which can lead to type I errors - that is, finding a statistical significant difference when one does not exist.

**Potential biases in the review process**

We attempted to overcome potential publication bias through rigorous searching. We had two studies translated, one from German and one from Spanish (Kappstein 1993; Vergara-Fernandez 2010). Five other trials also took place in countries where English is not the first language: Al-Naami 2009 in Saudi Arabia, Furukawa 2005 in Japan, Herruzo 2000 in Spain, Nthumba 2010 in Kenya and Pietsch 2001 in Germany.

We attempted to contact seven trial authors for additional information (Gupta 2007; Hajipour 2006; Herruzo 2000; Kappstein 1993; Pereira 1997; Pietsch 2001; Sensoz 2003). Five authors responded (Hajipour 2006; Herruzo 2000; Kappstein 1993; Pereira 1997; Sensoz 2003). We included six of these trials in the review (Gupta 2007; Hajipour 2006; Herruzo 2000; Kappstein 1993; Pereira 1997; Pietsch 2001). For the update we excluded Sensoz 2003, as there were no indications from the publication that participants were randomised, and we were unable to obtain information to the contrary. For the update we attempted to contact one author (Chen 2012); this trial is awaiting assessment.

Four trials acknowledged the role of commercial companies in supplying antiseptic products (Parienti 2002; Pereira 1999; Pereira 1997; Vergara-Fernandez 2010), and one trial was led by a research employee of a commercial company (Pietsch 2001). Tanner 2009 reported funding from a commercial company. As the lead author of
this review was also an author on an included study (Tanner 2009), two other authors independently extracted data from it and drafted related review text to minimise any bias.

Agreements and disagreements with other studies or reviews

This updated review differs from the earlier review as it includes an additional four studies (Al-Naami 2009; Nthumba 2010; Tanner 2009; Vergara-Fernandez 2010). However the addition of these new studies did not result in a change to the findings of the earlier review.

We identified two other systematic reviews of surgical hand antisepsis (Goncalves 2012; Hsieh 2006). We discussed Hsieh 2006 in the earlier version of this Cochrane review, but Goncalves 2012, published in 2012, is new to this update. Goncalves 2012 included 23 studies and two systematic reviews (the previous version of this Cochrane review and Hsieh 2006), focusing on traditional scrubs versus alcohol rubs. It included randomised trials, non-randomised trials, cohort studies, case-control studies, descriptive studies and case reports. Goncalves 2012 included six randomised studies which fell within the search dates of this update review but are excluded from this review (Larson 1990; Hobson 1998; Mulberry 2001; Larson 2001b; Marchetti 2003; Rotter 2006). Goncalves 2012 also included one unpublished study that we were unable to retrieve (Sigler 2001). However, we did include the three of the same randomised studies (Pietsch 2001; Parienti 2002; Gupta 2007). Goncalves 2012 concluded that alcohol preparations can replace traditional scrubs using aqueous chlorhexidine or iodine.

Hsieh 2006 was published in 2006 and includes three trials (Bryce 2001; Larson 2001b; Wheelock 1997). This Cochrane review update also included Wheelock 1997 but excluded Bryce 2001 and Larson 2001b, as they were not randomised. Hsieh 2006 failed to identify six of the eight trials published before 2006 that were included in this Cochrane review (Furukawa 2005; Herruzo 2000; Kappstein 1993; Pereira 1990; Pereira 1997; Pietsch 2001), and they excluded Parienti 2002 because it measured SSI rather than CFUs. Hsieh 2006 draws two conclusions: that surgical handrubs using alcohol-based products are more effective than 6 minute scrubs using 4% chlorhexidine, and that there is no evidence that a 2 minute scrub is more effective than a 3 minute scrub using 4% chlorhexidine.

Guidelines for hand antisepsis have changed slightly since the first version of this Cochrane review was published. The main changes are the increasing status of alcohol rubs, which are now considered to have equal standing to traditional scrubs and are acceptable alternatives (AIPP 2011; ACRN 2012); and the use of brushes which has been downgraded from not necessary to not recommended (AIPP 2011; ACRN 2012).

The conclusions one can draw from the evidence in this review are limited for the following reasons. Ten of the 14 trials included in this review measured CFUs rather than SSI (Furukawa 2005; Gupta 2007; Hajipour 2006; Herruzo 2000; Kappstein 1993; Pereira 1990; Pereira 1997; Pietsch 2001; Tanner 2009; Wheelock 1997). Two of these trials used the finger press method rather than the recognised standard glove juice test (Hajipour 2006; Herruzo 2000). Sample sizes are small in most of the trials. In three trials, participants in the scrub group had a choice of antiseptic solutions (Al-Naami 2009; Parienti 2002; Wheelock 1997), and details of the randomisation are poor in many of the trials.

A U T H O R S' C O N C L U S I O N S

Implications for practice

Uncertainty remains about the optimal method of hand antisepsis for minimising SSI outcomes: only four studies reported this outcome, and they were heterogeneous, at high risk of bias and mainly greatly underpowered. For the secondary (and surrogate) outcome of the number of CFUs on the hands, there is some generally low or very low quality evidence that quality chlorhexidine aqueous scrubs may reduce CFUs compared with povidone iodine aqueous scrubs and that an alcohol rub with added ingredient reduced the number of CFUs on the hand compared with aqueous scrubs (different types) immediately after antisepsis and at the end of the surgical procedure. The evidence regarding the impact of using a nail pick or brush on CFUs is unclear. In terms of duration of antisepsis, scrubbing for 3 minutes rather than 2 minutes may lead to fewer CFUs on the hand (low or very low quality evidence). Other data are more difficult to interpret.

Implications for research

Trials should attempt to measure the impact of hand antisepsis on SSIs. Investigators should also adhere to a recognised definition of SSI, which includes a 30 day follow-up. Trials should use adequate sample sizes based on a priori sample size calculations and take account of any data cross-over and/or clustering. The following trials might address important clinical questions - these and related uncertainties should be prioritised in conjunction with health professionals and policy makers. Future studies should be adequately designed and powered.

- Antisepsis compared with standard handwashing approaches.
- Relative effectiveness of aqueous scrubs versus alcohol rubs (outcome SSI).
- Optimum duration of aqueous scrubs (outcome SSI).
- Optimum duration of alcohol rubs (outcome SSI).
- Brushes, nail picks and sponges compared with no brushes, nail picks or sponges (outcome SSI).

A C K N O W L E D G E M E N T S

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CHARACTERISTICS OF STUDIES

Characteristics of included studies [ordered by study ID]

Al-Naami 2009
Methods
- Randomised controlled trial (described as an equivalence study)
  - Generation of random number sequence: no details given
  - Allocation concealment: no details given
  - Blinding: blinding of outcome assessors; no other details given
  - A priori sample calculations: yes
  - Antisepsis protocol: yes (partial)
  - Withdrawals: incomplete details given
  - Intention-to-treat analysis: no
  - Clear inclusion or exclusion criteria: no

Participants
- 1 hospital General Surgery division; surgeons (no further information on personnel)
- 600 patients initially randomised from general surgery; data reported on 500
- General Surgery: abdominal (e.g. cholecystectomy) and other; mixture of clean and clean-contaminated operations

Interventions
- Group 1 - traditional surgeons' handscrub for 3-5 min using 7.5% povidone iodine (Betadine) or 4% chlorhexidine gluconate (Hibiscrub) (228 patients)
- Group 2 - As with group 1 for first case; subsequent antisepsis with alcohol handrub with 62% ethanol (Purrel) 10 ml, allowed to dry (272 patients)

Outcomes
- Surgical site infection defined as any one or more of the following: symptoms and signs (pain, swelling, redness, tenderness, indurations, purulent discharge, opened wound) occurring within 30 d from surgery (examinations before discharge, at 1 week, at 1 month, and C/S results); no further detail supplied.

Notes
- All patients had standardised skin preparation. 76% of patients had prophylactic antibiotics (indicated for specified surgeries); no difference between arms.

Risk of bias

| Bias                              | Authors' judgement | Support for judgement                                                                 |
|----------------------------------|--------------------|---------------------------------------------------------------------------------------|
| Random sequence generation (selection bias) | Unclear risk       | No information on sequence generation                                                 |
|                                  |                    | Quote: "Participants were randomised to either a routine hand scrub or an alcohol hand-rub upon selecting a sealed envelope for each case." No further information. |
| Allocation concealment (selection bias) | Unclear risk       | No further information; not clear if envelopes were opaque or sequentially numbered   |
| Blinding (performance bias and detection bias) All outcomes - Blinding participants | High risk          | Clear differences between the procedures employed mean blinding of participants was not possible |
### Al-Naami 2009 (Continued)

| Characteristics | Risk of Bias | Description |
|-----------------|--------------|-------------|
| Blinding (performance bias and detection bias) All outcomes - Blinding care givers | Unclear risk | No information reported |
| Blinding (performance bias and detection bias) All outcomes - Blinding outcome assessors | Low risk | Assessment of wound appearance and swab by personnel unaware of allocation. Quote: "Surgeons who examined surgical sites were unaware of the groups' allocation"; "A swab was sent for C/S from any suspected SSI. Health care personnel taking swabs and interpreting results of C/S were also unaware of how hand disinfection for each group had been allocated." |
| Incomplete outcome data (attrition bias) All outcomes | High risk | 100 (1 in 6) participants originally randomised were excluded from analysis. The reasons for this are not fully explored. Quote: "Initially an equal number of cases (300 patients in each group) were randomised to each method. However, more cases were further excluded from each group as they turned out to be non-eligible for inclusion after the original randomization (e.g. acute or chronic cholecystitis on histopathological examination), incomplete forms, failed follow-ups, etc." |
| Selective reporting (reporting bias) | Low risk | Authors reported all specified objectives. Quote: "The objective of this study is to determine the equal efficacy of alcohol-based hand-rub as compared to traditional surgical scrub in the prevention of SSI as the primary outcome measure; the compliance of surgical staff and skin tolerance as the secondary outcome measure; also keeping in mind cost effectiveness and the potential change in surgical practice at least in our institution" |
| Other bias | Low risk | No other sources of bias detected |

### Furukawa 2005

| Characteristics | Description |
|-----------------|-------------|
| Methods | Randomised controlled trial Generation of random number sequence: no details given Allocation concealment: no details given Blinding: no details given A priori sample calculations: no Antiseptic protocol: yes Withdrawals: no details given Intention-to-treat analysis: no Clear inclusion or exclusion criteria: no |
| Participants | 22 operating room nurses Baseline comparability: baseline bacterial counts |
| Interventions | Group 1 - 3 min scrub using aqueous chlorhexidine gluconate Group 2 - 3 min scrub using aqueous povidone iodine |
| Outcomes | Outcome measure: CFUs on participants' hands Method of testing: glove juice method Timing of testing: before antiseptic and after antiseptic (no information regarding how long after antiseptic testing was conducted) |
| Notes | Participants did not take part in any surgical procedures |
**Furukawa 2005** (Continued)

**Risk of bias**

| Bias                                           | Authors' judgement | Support for judgement                                                                                                                                 |
|------------------------------------------------|--------------------|------------------------------------------------------------------------------------------------------------------------------------------------------|
| Random sequence generation (selection bias)   | Unclear risk       | Quote: "Twenty-two operating room nurses were randomly divided into two groups as follows: the PVI group (n = 11) and the CHG group (n = 11). All the nurses were examined for bacterial contamination of their hands before and after surgical handwashing".
Comment: Evidence of randomisation, however not enough evidence to suggest truly randomised sequence generation |
| Allocation concealment (selection bias)       | Unclear risk       | Quote: "Twenty-two operating room nurses were randomly divided into two groups as follows: the PVI group (n = 11) and the CHG group (n = 11). All the nurses were examined for bacterial contamination of their hands before and after surgical handwashing".
Comment: No evidence that appropriate allocation concealment took place |
| Blinding (performance bias and detection bias) | Unclear risk       | Quote: "All the nurses were examined for bacterial contamination of their hands before and after surgical handwashing"
Comment: No evidence of blinding of participants or personnel to blinding to intervention |
| Blinding (performance bias and detection bias) | Unclear risk       | Quote: "All the nurses were examined for bacterial contamination of their hands before and after surgical handwashing"
Comment: No evidence of blinding of participants or personnel to blinding to intervention |
| Blinding (performance bias and detection bias) | Unclear risk       | Tap water outcome
Quote: "We took water samples from 4 faucets just before hand washing and 1 ml of each sample was injected onto Brain-Heart infusion bouillon plates (Eiken K. K) and cultured at 37°Celsius for 48 hours for bacterial detection."
Comment: Samples seemingly taken by personnel, however unclear as to the role bias would play in the quantitative study of bacterial colonies in water faucets.
Hand and fingers outcome
Quote: "The samples were collected and pre-treated according to the Glove Juice method. In detail, the sample liquid was taken from the right glove just before hand washing and from the left glove after hand washing". This was then "cultured at 37°Celsius for 48 hours, thereafter the number of bacterial colonies was counted".
Comment: Unclear as to whether those obtaining the samples were blinded to the intervention. It is likely that they were not blinded; however the overall judgement is unclear. |
### Furukawa 2005 (Continued)

| **Incomplete outcome data** (attrition bias) | **Low risk** | No direct quotes, however no losses to follow-up encountered |
| **Selective reporting (reporting bias)** | **Low risk** | Quote: no direct quotes. |
| **Other bias** | **Low risk** | No other sources of bias detected |

### Gupta 2007

| **Methods** | Randomised cross-over controlled trial |
| **Population** | 18 operating room staff working in ophthalmic, podiatric and general surgery |
| **Interventions** | Group 1 - brush application of 7.5% povidone iodine aqueous scrub |
| **Outcomes** | Outcome measure: CFUs on participants' hands |

### Risk of bias

| **Bias** | **Authors' judgement** | **Support for judgement** |
| **Random sequence generation (selection bias)** | Unclear risk | Quote: Participants "were assigned to at random to one of three groups. Each group used one of the three solutions for five consecutive days. The following week, each group used a different scrub solution, such that all participants used each product over the study duration" |
| **Allocation concealment (selection bias)** | Unclear risk | Quote: Participants "were assigned to at random to one of three groups. Each group used one of the three solutions for five consecutive days. The following week, each group used a different scrub solution, such that all participants used each product over the study duration" |
### Gupta 2007 (Continued)

| Bias Category | Risk Type | Risk Level | Comment |
|---------------|-----------|------------|---------|
| Blinding (performance bias and detection bias) | All outcomes - Blinding participants | High risk | Quote: “Participants could not be blinded to the three solutions due to differences in their nature and method of application”.
Comment: Study describes that blinding of participants was not possible. There is no information given as to whether the investigators were blinded, but it is likely there were similarly unblinded to the intervention for the reasons given for the participants above. The judgement for participant blinding is therefore high risk. |
| Blinding (performance bias and detection bias) | All outcomes - Blinding care givers | Unclear risk | Quote: “Participants could not be blinded to the three solutions due to differences in their nature and method of application”.
Comment: Study describes that blinding of participants was not possible. There is no information given as to whether the investigators were blinded, but it is likely there were similarly unblinded to the intervention for the reasons given for the participants above. The judgement for blinding of caregivers here is unclear. |
| Blinding (performance bias and detection bias) | All outcomes - Blinding outcome assessors | Unclear risk | Bacterial sampling
Quote: “The subject introduced their hands into this bag and the investigator massaged their hands externally with emphasis on web spaces and subungual areas”
Comment: no attempt made at blinding method of obtaining bacterial sample, which would advise a high risk of bias decision
Microbial assay
Quote: "Samples were sent to the microbiology laboratory immediately after collection in a blinded manner"
Comment: There was adequate evidence that approach microbial testing was blinded.
Therefore, overall assessment of the risk of bias for outcome assessment is unclear. |
| Incomplete outcome data | (attrition bias) | Low risk | Quote: “Eighteen surgical staff members participated in this study. All members used each of the three scrub solutions over the duration of the study. . . Two volunteers developed a skin rash and a burning sensation on their hands within and a few minutes after their first scrub with the ABWA product. They also reported a metallic taste in their mouth and palpitations. Although none of their symptoms were severe and resolved shortly thereafter without any medical intervention, they were removed from the study. All the remaining staff volunteers completed the study”.
Comment: small loss to follow-up and full explanations given as to the reasons for dropout |
| Selective reporting (reporting bias) | Low risk | No direct quotes, although the efficacy of the product in terms of reduction of log reduction in bacterial counts and product preference by participants are both included in the Results in full and comprehensive manner, as outlined in the Methods. |
| Other bias | Unclear risk | Cross-over design, unclear if accounted for in analysis |
## Hajipour 2006

### Methods

| Bias                               | Authors' judgement | Support for judgement                                                                 |
|------------------------------------|--------------------|----------------------------------------------------------------------------------------|
| Random sequence generation (selection bias) | Low risk           | Quote: “Orthopaedic surgeons were allocated to one of two different handwashing protocols using a randomisation table” |
|                                    |                    | Comment: evidence of random sequence generation, therefore judged as low risk           |
| Allocation concealment (selection bias) | Unclear risk      | Quote: "[T]he surgeon was randomised to wash for 5 min with either chlorhexidine or alcohol gel" |
|                                    |                    | Comment: no evidence that there was an attempt at allocation concealment                |
| Blinding (performance bias and detection bias) | Unclear risk | Quote: “The hand-washing protocol dictated that all surgeons should wash for 5 min with chlorhexidine for their first case with thorough cleaning under the fingernails. Thereafter, the surgeon was randomised to wash for 5 min with either the chlorhexidine or alcohol gel. Alcohol was allowed to dry on the hands prior to double gloving”. |
|                                    |                    | Comment: no evidence that participants or personnel were blinded to intervention        |
|                                    |                    |                                                                                         |

### Participants

- 4 surgeons working in a trauma surgery
- Baseline comparability: surgeon's grade, order of patient on the operating list, duration of surgery

### Interventions

| Group 1 | Group 2 |
|---------|---------|
| 3 min scrub using aqueous chlorhexidine gluconate | 3 min application of 0.5% chlorhexidine gluconate in 70% alcohol |
| All surgeons washed with chlorhexidine (no further detail) for 5 min for first procedure with thorough cleaning under fingernails. |

### Outcomes

- Outcome measure: CFUs on participants' hands
- Method of testing: finger press testing with agar plates
- Timing of testing: at the end of the surgical procedure

### Notes

- The 4 surgeons, who were not blinded, were randomised once and tested 53 times
Comment: no evidence that participants or personnel were blinded to intervention

| Risk of bias                                      | Low risk                                                                 | Quote: “The number of bacterial colonies present after 24 h and 48 h of incubation were recorded for each agar plate by a microbiologist blinded to the washing protocol used” |
|--------------------------------------------------|--------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Blinding (performance bias and detection bias)   | Low risk                                                                 | Comment: adequate blinding of assessment outcome                                                                                                                                               |
| All outcomes - Blinding outcome assessors        |                                                                           |                                                                                                                                                                                                 |
| Incomplete outcome data (attrition bias)         | Low risk                                                                 | Quote: “Overall, 41 procedures and 82 episodes of handwashes were included in the study. Two episodes were discarded due to contamination at the time of glove removal. There was no incidence of outer glove perforation during this study” |
| All outcomes                                     |                                                                           | Comment: good evidence to suggest losses to follow-up were accounted for and there was minimal effect of attrition bias                                                                     |
| Selective reporting (reporting bias)             | Low risk                                                                 | No direct quotes, but the assessment variable of bacterial colonisation after different methods of handwashing (which was outlined in the methodology) is accounted for in the results. |
| Other bias                                       | Unclear risk                                                             | This appears to be a clustered randomised trial; it does not seem that clustering was taken into account in the analysis.                                                                      |

### Herruzo 2000

| Methods                                           | Randomised cross-over controlled trial                                   |
|---------------------------------------------------|-------------------------------------------------------------------------|
| Generation of random number sequence: no details given |                                                                 |
| Allocation concealment: no details given          |                                                                 |
| Blinding: no details given                        |                                                                 |
| A priori sample calculations: no                  |                                                                 |
| Antisepsis protocol: minimal details              |                                                                 |
| Withdrawals: no details given                     |                                                                 |
| Intention-to-treat analysis: no                   |                                                                 |
| Clear inclusion or exclusion criteria: no         |                                                                 |

| Participants                                      | 154 members of the surgical teams working in plastic surgery and traumatology |
|---------------------------------------------------|--------------------------------------------------------------------------------|
| Baseline comparability: baseline bacterial counts  |                                                                              |

| Interventions                                     | Group 1 - 3 min scrub using aqueous chlorhexidine gluconate, n = 50          |
|---------------------------------------------------|------------------------------------------------------------------------------|
|                                                   | Group 2 - 3 min scrub using aqueous povidone iodine, n = 49                  |
|                                                   | Group 3 - 3 min rub with N-duopropenide, n = 55                              |

| Outcomes                                          | Outcome measure: CFUs on participants' hands                                |
|---------------------------------------------------|------------------------------------------------------------------------------|
|                                                   | Method of testing: finger press testing with agar plates                     |
|                                                   | Timing of testing: before antiseptic, immediately after antiseptic and at the end of the surgical procedure |

| Notes                                              | —                                                                           |

### Risk of bias

**Surgical hand antisepsis to reduce surgical site infection (Review)**

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| Bias                                           | Authors' judgement | Support for judgement |
|------------------------------------------------|--------------------|------------------------|
| Random sequence generation (selection bias)   | Unclear risk       | Quote: "Healthy volunteers washed with one of the three products for 1 week for 3 consecutive weeks. The order of the washings was randomised"  
Comment: indication that a randomisation process was undertaken, however not clear how the sequence was generated |
| Allocation concealment (selection bias)       | Unclear risk       | Quote: "Healthy volunteers washed with one of the three products for 1 week for 3 consecutive weeks. The order of the washings was randomised"  
Comment: no evidence of allocation concealment given |
| Blinding (performance bias and detection bias) All outcomes - Blinding participants | High risk          | Quote: "The effect of standard surgical washings with 7.5% iodine povidone or 4% chlorhexidine (both with scrubbing for 3 min following a standard technique) was compared with the effect of washing (without scrubbing with 2.3% N-duopropenide in 60° isopropanol with dermoprotective substance). The latter solution was poured over the hands, which were then rubbed together, and when it began to dry, it was reapplied over 3 min."  
Comment: The study design here is a cross-over design, reducing potential effects of bias; however, there is no evidence that participants or personnel were blinded to the intervention given, and the conditions differed sufficiently that blinding would not have been possible. |
| Blinding (performance bias and detection bias) All outcomes - Blinding care givers | High risk          | Quote: "The effect of standard surgical washings with 7.5% iodine povidone or 4% chlorhexidine (both with scrubbing for 3 min following a standard technique) was compared with the effect of washing (without scrubbing with 2.3% N-duopropenide in 60° isopropanol with dermoprotective substance). The latter solution was poured over the hands, which were then rubbed together, and when it began to dry, it was reapplied over 3 min."  
Comment: The study design here is a cross-over design, reducing potential effects of bias; however, there is no evidence that participants or personnel were blinded to the intervention given, and the conditions differed sufficiently that blinding would not have been possible. |
| Blinding (performance bias and detection bias) All outcomes - Blinding outcome assessors | Unclear risk       | Quote: "We used fingertips to sample hand bacteria as the standard European Norm (pr EN 1500) except we had not artificially contaminated the hands: five fingertips are rubbed for 1 min on a Petri dish containing 10 ml of TBS plus an antiseptic neutralize". These were cultured for "48 h at 37°C, and then the CFU/hand (the five fingertips), were counted and transformed into a decimal logarithm".  
Comment: Despite the assessment being quantitative, it is not clear whether those who obtained the bacterial samples were independent of the study or blinded to the intervention. |
| Incomplete outcome data (attrition bias)       | Low risk           | No direct quotes, although no losses to follow-up recorded in Results  
Comment: no obvious source of attrition bias |
| Selective reporting (reporting bias)           | Low risk           | No direct quotes  
Comment: tabular format of results incorporates the assessment outcomes outlined (CFUs) between the groups, suggesting a low risk |
| Other bias                                     | Unclear risk       | Although the trial had a cross-over design, it did not appear that this was reflected in the analysis. |
### Kappstein 1993

#### Methods

- Randomised cross-over trial (participants took part in each of 3 groups)
- Generation of random number sequence: no details given
- Allocation concealment: no details given
- Blinding: no details given
- A priori sample calculations: yes
- Antiseptic protocol: no
- Withdrawals: no details given
- Intention-to-treat analysis: no
- Clear inclusion or exclusion criteria: no

**Participants**

- 24 surgeons
- Baseline comparability: baseline bacterial counts

**Interventions**

- Group 1: 1 min wash with soap and water followed by 5 min rub with an alcoholic disinfectant
- Group 2: 1 min wash with soap and water followed by 3 min rub with an alcoholic disinfectant
- Group 3: 1 min wash with chlorhexidine soap followed by two min of rubbing with 0.5% chlorhexidine in isopropanol

**Outcomes**

- Outcome measure: CFUs on participants' hands
- Method of testing: glove juice method
- Timing of testing: before antisepsis and immediately after antisepsis

**Notes**

- —

#### Risk of bias

| Bias | Authors' judgement | Support for judgement |
|------|--------------------|-----------------------|
| Random sequence generation (selection bias) | Unclear risk | Just states “random order” with no further information |
| Allocation concealment (selection bias) | Unclear risk | As with sequence generation; no further information |
| Blinding (performance bias and detection bias) | Unclear risk | Standard technique compared with shorter techniques precludes blinding of personnel but no further information |
| Blinding (performance bias and detection bias) | Unclear risk | Standard technique compared with shorter techniques precludes blinding of personnel but no further information |
| Blinding (performance bias and detection bias) | Unclear risk | No information on blinding of assessment reported |
| Incomplete outcome data (attrition bias) | Low risk | No evidence of dropouts/loss of data |
| Selective reporting (reporting bias) | Low risk | All specified outcomes reported |
### Kappstein 1993 (Continued)

| Other bias | Unclear risk | Although the trial had a cross-over design, it did not appear that this was reflected in the analysis. |

### Nthumba 2010

| Methods | Cluster-randomised controlled cross-over trial |
|---------|-----------------------------------------------|
|         | Generation of random number sequence: appropriate |
|         | Allocation concealment: no details given but considered unlikely to be an issue in cluster-randomisation |
|         | Blinding: outcome assessors |
|         | A priori sample calculations: yes |
|         | Antisepsis protocol: yes |
|         | Withdrawals: details of patients without postdischarge surveillance |
|         | Intention-to-treat analysis: no |
|         | Clear inclusion or exclusion criteria: yes |

| Participants | 66 surgeons and trainees; 3317 patients |

| Interventions | Group 1 - plain soap and water: 4-5 min clean running water and plain soap; sterile cotton hand towel dry. 5 clusters (n = 1682 patients) |
|               | Group 2 - As group 1 before first procedure of day and subsequently in case of visible soiling, then alcohol-based handrub (75% isopropyl alcohol, 1.45% glycerol, 0.125% hydrogen peroxide) for 3 min and kept wet (7-10 ml per preparation) |

| Outcomes | SSI (defined using modified US Centers for Disease Control and Prevention definitions for nosocomial infection) detected by tours of hospital wards; reviews in outpatient clinic; telephone contact: diagnosis established jointly by study collaborators |

| Notes | — |

### Risk of bias

| Bias | Authors’ judgement | Support for judgement |
|------|--------------------|-----------------------|
| Random sequence generation (selection bias) | Low risk | Surgical hand preparation procedures were assigned randomly to the 6 participating operating theatres by tossing a coin, with a cross-over every 2 months. There was no indication of baseline imbalance for important variables such as type of surgery, contamination level of the surgery or use of antibiotic prophylaxis. |
| Allocation concealment (selection bias) | Low risk | Allocation was decided by toss of a coin. Not clear who undertook this process and if it was concealed from the sites. Given that this was a cluster trial with cross-over, the potential for bias stemming from allocation concealment was limited. As the Cochrane Handbook for Systematic Reviews of Interventions notes, “Cluster-randomised trials often randomise all clusters at once, so lack of concealment of an allocation sequence should not usually be an issue.” |
| Blinding (performance bias and detection bias) All outcomes - Blinding participants | High risk | Personnel were aware of allocation as procedures differed in obvious ways. Compliance by surgical teams was determined by observation of practices. A trained observer who did not belong to the surgical team checked whether each sink had the correct hand preparation and whether all surgeons (including visiting staff) followed the recommended hand preparation procedures. |
### Nthumba 2010 (Continued)

| Blinding (performance bias and detection bias) | Unclear risk | No information reported |
|-----------------------------------------------|--------------|-------------------------|
| All outcomes - Blinding care givers           |              |                         |
| Blinding (performance bias and detection bias) | Low risk     | Diagnosis, documentation and determination of SSI by personnel without knowledge of allocation. |
| All outcomes - Blinding outcome assessors     |              | *"SSI was diagnosed ... and documented by a trained nurse who visited the surgical wards three to four times each week during the 30 days after surgery. Patients discharged before this were reviewed in the outpatient clinic or contacted by telephone ... The nurse was blinded to the method of surgical hand preparation. The diagnosis of SSI was established jointly by the study collaborators; differences in SSI ascertainment were resolved by consensus without knowledge of the study allocation of the patient."*
| Incomplete outcome data (attrition bias)      | Low risk     | 3722 patients underwent a surgical procedure in the operating theatre, and 3317 were included in the 10 study clusters. Postdischarge surveillance data could not be obtained for 184 patients (5.5%). |
| All outcomes                                  |              |                         |
| Selective reporting (reporting bias)          | Low risk     | The primary objective of this cluster-randomised, cross-over trial was to compare the efficacy of plain soap and water with alcohol-based handrub, using SSI rates as the main outcome measure. The feasibility and affordability of the local production of an alcohol-based handrub was also investigated, together with an assessment of its acceptability among healthcare workers. All specified outcomes were reported. |
| Other bias                                    | Low risk     | There was no evidence of other sources of bias. Clustered nature of the data was taken into account in the trial. |

### Parienti 2002

| Methods                                     | Cluster-randomised controlled equivalence trial |
|---------------------------------------------|-----------------------------------------------|
| Generation of random number sequence:      | random number tables                            |
| Allocation concealment:                    | no details given                                |
| Blinding: discussed but only conducted       | during postdischarge follow-up                  |
| A priori sample calculations:               | yes                                            |
| Antisepsis protocol:                        | yes                                            |
| Withdrawals: 51 patients lost during       | follow-up                                       |
| Intention-to-treat analysis:                | yes                                            |
| Clear inclusion or exclusion criteria:      | yes                                            |
| Participants                                | Surgical teams within 6 hospitals were randomised. 4387 patients undergoing clean and clean-contaminated surgery were included in the study. Baseline comparability: details of surgical procedures, duration of surgery, patients' ASA classifications |
| Interventions                               | Group 1 - 5 min scrub using either 4% povidone iodine or 4% chlorhexidine gluconate |
|                                             | Group 2 - 5 min handrub with alcohol solution containing 75% propanol-1, propanol-2 with mecetronium ethylsulphate |
| Outcomes                                    | Outcome measure: SSIs in patients at 30 d using CDC definition |
|                                             | Method of testing: observation by surgeon or infectious disease specialist, case note review, telephone interview |
|                                             | Timing of testing: 30 d follow-up               |
**Risk of bias**

| Bias | Authors' judgement | Support for judgement |
|------|-------------------|----------------------|
| Random sequence generation (selection bias) | Low risk | Quote: "Each participating surgical service was assigned a 2-digit random number by using a random number table. Surgical services corresponding to the 3 higher numbers were assigned to hand-rubbing with AAS and the remaining 3 services were assigned to traditional hand-scrubbing". Comment: adequate evidence of random sequence generation |
| Allocation concealment (selection bias) | Unclear risk | Quote: "Each participating surgical service was assigned a 2-digit random number by using a random number table. Surgical services corresponding to the 3 higher numbers were assigned to handrubbing with AAS and the remaining 3 services were assigned to traditional hand-scrubbing" Comment: no evidence of allocation concealment |
| Blinding (performance bias and detection bias) | High risk | Quote: "[O]bservers of the clinical outcome could not be blinded to the hand antisepsis protocol." The study also mentions that "compliance observers did not belong to the operating department team but were usually present in the surgical suite. To avoid a Hawthorne effect the surgical teams were not informed of the timing of the evaluations". Comment: Although the effect of blinding has been considered, in the comparison of different scrubbing protocols it would be difficult to blind the participant or personnel. The risk of bias is still high in this instance, however. |
| Blinding (performance bias and detection bias) | High risk | Quote: "[O]bservers of the clinical outcome could not be blinded to the hand antisepsis protocol." The study also mentions that "compliance observers did not belong to the operating department team but were usually present in the surgical suite. To avoid a Hawthorne effect the surgical teams were not informed of the timing of the evaluations". Comment: Although the effect of blinding has been considered, in the comparison of different scrubbing protocols it would be difficult to blind the participant or personnel. The risk of bias is still high in this instance, however. |
| Blinding (performance bias and detection bias) | High risk | Surgical site infection outcome Quote: "According to CDC guidelines, all SSIs had to be confirmed by the surgeon or the physician in charge on the patient. Thus, observers of the clinical outcome could not be blinded to the hand antisepsis protocol" Comment: As the surgeon was the participant in this case. it is clear that this could constitute a high risk of bias. Tolerance and compliance outcome Quote: "[T]he surgical personnel (77 subjects) were asked to estimate the effect of the 2 protocols on their skin. We used 2 10 cm visual analogue scales, at month 0 and after 3 crossovers; 0 cm representing absence of an tolerance problem and 10 cm representing maximal dryness with chapped hands and desquamation or maximal irritation with erythema, burning sensation, and abrasion." Comment: The surgeon was the (unblinded) participant who reported the variables, so the results may be affected by performance bias. |
### Parienti 2002 (Continued)

| Incomplete outcome data (attrition bias) | Low risk | Quote: "During the study period ... 4823 consecutive patients underwent surgery. Among these, 385 patients underwent contaminated or dirty-contaminated surgery, and 51 were lost to follow up at 30 days (17 in the hand-rubbing group). The remaining 4387 patients (68.5% of whom underwent clean surgery) were considered for analysis". |
| Selective reporting (reporting bias) | Low risk | Quote: "Thirty-day surgical site infections rates were the primary end point; operating department teams’ tolerance of and compliance with hand antisepsis were secondary end points". |
| Other bias | Unclear risk | This appears to be a clustered cross-over study; it does not seem that clustering was taken into account in the analysis. |

### Pereira 1990

| Methods | Randomised controlled cross-over trial (Latin square design - participants took part in each of 4 interventions) |
| Allocation concealment: no details given |
| Blinding: no details given |
| A priori sample calculations: no |
| Antisepsis protocol: yes |
| Withdrawals: 2 participants withdrew |
| Intention-to-treat analysis: no |
| Clear inclusion or exclusion criteria: no |

| Participants | 34 anaesthetic, recovery and ward nurses |
| Baseline comparability: gender, age, ethnicity, hand dominance, baseline bacterial counts |

| Interventions | Group 1 - 5 min initial scrub and 3 min subsequent scrub using chlorhexidine |
| Group 2 - 3 min initial and 30 s subsequent scrub using chlorhexidine |
| Group 3 - 5 min initial and 3 min subsequent scrub using povidone iodine |
| Group 4 - 3 min initial and 30 s subsequent scrub using povidone iodine |

| Outcomes | Outcome measure: CFUs on participants' hands |
| Method of testing: glove juice method |
| Timing of testing: before antiseptics, immediately after antiseptics, 2 h after initial antiseptic, 2 h after subsequent antiseptic |

| Notes | Participants did not take part in any surgical procedures |

### Risk of bias

| Bias | Authors' judgement | Support for judgement |
| Random sequence generation (selection bias) | Unclear risk | Quote: "Subjects were randomly assigned to one of four groups, and each group was assigned to one of the four scrub regimens each week. Control on the treatment order was achieved through a Latin square design, as described by Winder." |
### Pereira 1990 (Continued)

| Bias Type                                    | Risk | Comment:                                                                 | Direct Quote                                                                 | Comment:                                                                                              |
|----------------------------------------------|------|--------------------------------------------------------------------------|------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------------|
| **Allocation concealment** (selection bias)  | Unclear risk | adequate evidence of an appropriate study design, but on balance not enough evidence of truly random sequence generation | "Subjects were randomly assigned to one of four groups, and each group was assigned to one of the four scrub regimens each week. Control on the treatment order was achieved through a Latin square design, as described by Winder." | inadequate evidence of truly random sequence generation                                                                                                   |
| **Blinding (performance bias and detection bias)** | Unclear risk | no indication that allocation to each group was concealed to the personnel | "Subjects were randomly assigned to one of four groups, and each group was assigned to one of the four scrub regimens each week . . . Subjects were supervised by the investigator while they scrubbed on all test occasions.‖ | inadequate evidence of truly random sequence generation                                                                                                 |
| **Blinding (performance bias and detection bias)** | Unclear risk | no evidence to suggest that there was appropriate blinding of participants or personnel during the study | "Subjects were randomly assigned to one of four groups, and each group was assigned to one of the four scrub regimens each week . . . Subjects were supervised by the investigator while they scrubbed on all test occasions.‖ | inadequate evidence of truly random sequence generation                                                                                                 |
| **Blinding (performance bias and detection bias)** | Unclear risk | no evidence to suggest that there was appropriate blinding of participants or personnel during the study | "Subjects were randomly assigned to one of four groups, and each group was assigned to one of the four scrub regimens each week . . . Subjects were supervised by the investigator while they scrubbed on all test occasions.‖ | inadequate evidence of truly random sequence generation                                                                                                 |
| **Bacterial contamination**                  | Unclear risk | indication that those collecting the samples, administering the fluid or those performing the microbial assays were in any way blinded to the intervention or protocol | "While the glove was still on the hand, a sample of the fluid was taken . . . [S]amples were collected on four occasions for each condition: (1) immediately before scrubbing (both hands), (2) immediately after the initial surgical scrub (non-dominant hand only) (3) 2 hours after the initial surgical scrub, immediately before the consecutive scrub (dominant hand) and (4) 2 hours after one consecutive surgical scrub (dominant hand).‖ | inadequate evidence of truly random sequence generation                                                                                                 |
| **Incomplete outcome data** (attrition bias) | Low risk | evidence that losses to follow-up were small and fully accounted for | "Thirty-six subjects were recruited, but two subjects withdrew from the experiment before completing all four treatments (scrubs) because of skin reactions, including erythema, burning sensations and local swelling" | incomplete evidence of truly random sequence generation                                                                                                  |
| **Selective reporting (reporting bias)**     | Low risk | results of each 'scrub' are displayed fully at baseline and subsequent time intervals in the Results as laid out in the Methodology. | No direct quotes, but the results of each 'scrub' are displayed fully at baseline and subsequent time intervals in the Results as laid out in the Methodology. | selective reporting of study results                                                                                                                          |
| **Other bias**                               | Unclear risk | trial had a cross-over design, it did not appear that this was reflected in the analysis. | Although the trial had a cross-over design, it did not appear that this was reflected in the analysis. | inadequate evidence of truly random sequence generation                                                                                                   |
### Pereira 1997

**Methods**
- Randomised controlled trial cross-over (Latin square design - participants took part in each of 5 interventions)
- Generation of random number sequence: no details given
- Allocation concealment: no details given
- Blinding: no details given
- A priori sample calculations: no
- Antisepsis protocol: yes
- Withdrawals: yes details provided
- Intention-to-treat analysis: no
- Clear inclusion or exclusion criteria: no

**Participants**
- 23 operating room nurses
- Baseline comparability: age, gender, skin condition, baseline bacterial counts

**Interventions**
- Group 1 - 5 min initial and 3.5 min subsequent scrub using 4% chlorhexidine
- Group 2 - 3 min initial and 2.5 min subsequent scrub using 4% chlorhexidine
- Group 3 - 3 min initial and 2.5 min subsequent scrub using povidone iodine with triclosan
- Group 4 - 3 min initial scrub using 4% chlorhexidine followed by a 30 s application of isopropanol 70% and chlorhexidine 0.5%, and subsequent scrubs using 30 s application of isopropanol 70% and chlorhexidine 0.5%
- Group 5 - 2 min initial scrub using 4% chlorhexidine followed by a 30 s application of ethanol 70% and chlorhexidine 0.5%, and subsequent scrubs using 30 s application of ethanol 70% and chlorhexidine 0.5%

**Outcomes**
- Outcome measure: CFUs on participants' hands
- Method of testing: glove juice method
- Timing of testing: before antisepsis, immediately after antisepsis, 2 h after initial antisepsis, 2 h after subsequent antisepsis

**Notes**
- Participants did not take part in any surgical procedures

### Risk of bias

| Bias                                           | Authors' judgement | Support for judgement |
|------------------------------------------------|--------------------|-----------------------|
| Random sequence generation (selection bias)    | Unclear risk       | Quote: "Subjects who agreed to participate in the study followed each of the scrub protocols in turn, the order controlled by the use of a Latin square design. That is, every nurse was required to complete every protocol but not in the same sequence." |
| Allocation concealment (selection bias)        | Unclear risk       | Comment: clear that the study design does reduce selection bias; however not clear whether there is enough evidence to suggest random sequence was generated |
| Blinding (performance bias and detection bias) | High risk          | Quote: "Subjects who agreed to participate in the study . . . followed each scrub protocol each day for one week (referred to as the 'test week') with a week of normal activities between each test week. They were issued with a supply of the specific antiseptic to be used in excess to their requirements for scrubbing so that the appropriate antiseptic could be used exclusively dur- |

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**Surgical hand antisepsis to reduce surgical site infection (Review)**

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ing the test week. Subjects were assessed before commencing each scrub pro-
tocol and at the end of the test week to determine changes in the number of
colony forming units (cfu) after scrubbing and changes in the condition of the
hands.”

Comment: Although study design allows for repeated testing, there is no ev-
idence that there was effective blinding of participants and personnel to the
different interventions, and differences between the conditions would have
been clearly apparent to those taking part.

### Blinding (performance bias and detection bias)

| All outcomes - Blinding care givers | High risk |
|------------------------------------|-----------|

Quote: "Subjects who agreed to participate in the study . . . followed each
scrub protocol each day for one week (referred to as the ‘test week’) with a
week of normal activities between each test week. They were issued with a
supply of the specific antiseptic to be used in excess to their requirements for
scrubbing so that the appropriate antiseptic could be used exclusively dur-
ing the test week. Subjects were assessed before commencing each scrub pro-
tocol and at the end of the test week to determine changes in the number of
colony forming units (cfu) after scrubbing and changes in the condition of the
hands.”

Comment: Although study design allows for repeated testing, there is no ev-
idence that there was effective blinding of participants and personnel to the
different interventions, and differences between the conditions would have
been clearly apparent to those taking part.

### Blinding (performance bias and detection bias)

| All outcomes - Blinding outcome assessors | Unclear risk |
|------------------------------------------|--------------|

Sampling method

Quote: "A glove juice sampling method to estimate the numbers of CFU
present on the hands”

Comment: no indication that those performing the sampling or performing the
microbial assays were independent to the study

Skin condition

Quote: "Larson’s Weekly Skin Assessment Rating Scale was used to rate the
condition of the hands with respect to appearance, integrity, moisture and
sensation. Participants rated themselves on a weekly basis. An independent
rater, blind to the protocol being followed by the subject, also rated the sub-
ject’s dominant hand each week.”

Comment: evidence that reasonable measures were undertaken to blind out-
come assessment. Overall assessment unclear for outcome assessment blind-
ing.

### Incomplete outcome data

| All outcomes | High risk |
|--------------|-----------|

Quote: "Of the 32 participants recruited, 23 completed all five scrub protocols
(scrubs) because it was discovered [that several participants] were allergic or
sensitive to one of the more antiseptics. One felt that her skin was already too
dry to be able to participate further. One person could not complete the study
because she was unexpectedly off work for an extended period and the other
two gave no reason for withdrawing.”

Comment: fairly high rates of losses to follow-up and no information given as
to why some participants did not complete the study. Although it seems that
the researchers acted appropriately here, it is difficult to give a judgement oth-
er than high risk
Pereira 1997 (Continued)

Selective reporting (reporting bias) Low risk
No direct quotes, but reasonable evidence to suggest that both bacterial contamination and skin condition have been appropriately and fully tested and reported upon effectively.

Other bias Unclear risk
Although the trial had a cross-over design, it did not appear that this was reflected in the analysis.

Pletsch 2001

Methods
Randomised cross-over trial
Generation of random number sequence: no details given
Allocation concealment: no details given
Blinding: no details given
A priori sample calculations: no
Antiseptics protocol: no
Withdrawals: no details given
Intention-to-treat analysis: no
Clear inclusion or exclusion criteria: no

Participants
75 surgeons
Baseline comparability: baseline bacterial counts

Interventions
Group 1 - surgical scrub using 4% chlorhexidine (details of the duration are not given)
Group 2 - alcohol rub using Sterillium (45% w/w of propan-2-01, 30% w/w of propan-1-01 and 0.2% w/w of ethylhexadecyl(dimethyl ammonium ethylsulfate))(details of the duration are not given)

Outcomes
Outcome measure: CFUs on participants’ hands
Method of testing: glove juice method
Timing of testing: before antisepsis, immediately after antisepsis and after surgical procedure completed

Notes
—

Risk of bias

| Bias                               | Authors’ judgement | Support for judgement                                                                 |
|------------------------------------|--------------------|--------------------------------------------------------------------------------------|
| Random sequence generation (selection bias) | Unclear risk       | Dermal tolerance study                                                               |
|                                    |                    | Quote: “The first study was performed with a panel of 60 volunteers divided in two subgroups of 30 persons each.” |
|                                    |                    | Comment: no indication of how the subgroups were divided                              |
|                                    |                    | Surgical hand disinfection study                                                     |
|                                    |                    | Quote: “In a randomised cross-over clinical trial the antimicrobial efficacy of Sterillium for the surgical hand-rub was tested against Hibiscrub in the Kantonsspital Basel over a period of 11 weeks … two weeks were needed for recruitment, four weeks to conduct the Sterillium arm of the study, one week’s interval and then four weeks for the Hibiscrub part.” |
|                                    |                    | Comment: no further information given as to how participants were recruited and how they were assigned and allocated to which intervention |
|                                    |                    | Hygienic hand disinfection study                                                     |
Quote: "The antimicrobial efficacy of each product was compared with that of 60% (v/v) 2-propanol on artificially contaminated hands (E. coli K12, NCTC 10538) using a cross-over design with 15 volunteers."

Comment: no indication of if a randomised sequence was used and what method was implemented

| Allocation concealment (selection bias) | Unclear risk | Dermal tolerance |
|-----------------------------------------|--------------|------------------|
|                                        |              | Quote: "The first study was performed with a panel of 60 volunteers divided in two subgroups of 30 persons each" |
|                                        |              | Comment: No indication of how allocation was concealed |

Surgical hand disinfection

Quote: "In a randomised cross-over clinical trial the antimicrobial efficacy of Sterillium for the surgical hand-rub was tested against Hibiscrub in the Kantonsspital Basel over a period of 11 weeks . . . two weeks were needed for recruitment, four weeks to conduct the Sterillium arm of the study, one week's interval and then four weeks for the Hibiscrub part."

Comment: no indication if allocation was concealed

Hygienic hand disinfection

Quote: "The antimicrobial efficacy of each product was compared with that of 60% (v/v) 2-propanol on artificially contaminated hands (E. coli K12, NCTC 10538) using a cross-over design with 15 volunteers."

Comment: no indication if allocation was concealed

| Blinding (performance bias and detection bias) | Unclear risk | Dermal tolerance |
|-----------------------------------------------|--------------|------------------|
| All outcomes - Blinding participants          |              | Quote: "The first panel started with Hibiscrub, the second with Sterillium. Both products were used over seven weeks after one week of preconditioning. After an interval of four weeks the second run started with a single crossover of products."
|                                               |              | Comment: although cross-over trial used, no indication that the participants or personnel were blinded to the intervention |

Surgical hand disinfection

Quote: "Organisms were recovered by the glove juice method. Plastic bags with the sampling fluid were placed on the subjects hands. The bag on each hand was secured and massaged for 1 min in a uniform manner by a laboratory technician."

Comment: no indication whether participants were blinded to intervention and unclear whether technician was blinded to which intervention had been used

Hygienic hand disinfection

Quote: "The antimicrobial efficacy of each product was compared with that of 60% (v/v) 2-propanol on artificially contaminated hands (E. coli K12, NCTC 10538) using a cross-over design with 15 volunteers. The hands were first washed for 1 min with soft soap, dried with paper towels immersed in the contamination fluid up to the mid-metacarpals for 5 s with fingers spread and then allowed to dry for 3 min"
### Pietsch 2001 (Continued)

| Blinding (performance bias and detection bias) All outcomes - Blinding care givers | Unclear risk | Dermal tolerance |
|---|---|---|
| **Comment:** unclear as to whether participants or personnel were blinded to the interventions |

#### Surgical hand disinfection

*Quote:* "The first panel started with Hibiscrub, the second with Sterilium. Both products were used over seven weeks after one week of preconditioning. After an interval of four weeks the second run started with a single crossover of products".

*Comment:* although cross-over design used, no indication that the participants or personnel were blinded to the intervention

#### Hygienic hand disinfection

*Quote:* "The antimicrobial efficacy of each product was compared with that of 60% (v/v) 2-propanol on artificially contaminated hands (*E. coli* K 12, NCTC 10538) using a cross-over design with 15 volunteers. The hands were first washed for 1 min with soft soap, dried with paper towels immersed in the contamination fluid up to the mid-metacarpals for 5 s with fingers spread and then allowed to dry for 3 min."

*Comment:* unclear as to whether participants or personnel were blinded to the interventions

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#### Blinding (performance bias and detection bias) All outcomes - Blinding outcome assessors

| Unclear risk | Dermal tolerance |
|---|---|
| **Comment:** unclear as to whether participants or personnel were blinded to the interventions |

#### Surgical hand disinfection

*Quote:* "Organisms were recovered by the glove juice method. Plastic bags with the sampling fluid were placed on the subjects hands. The bag on each hand was secured and massaged for 1 min in a uniform manner by a laboratory technician"

*Comment:* no indication whether participants were blinded to intervention and unclear whether technician was blinded to which intervention had been used

#### Hygienic hand disinfection

*Quote:* "The following parameters were measured before and after application: Clinical assessment by observation of the hands and the forearms of the volunteers by a dermatologist. The number of volunteers which dropped out of the study because of skin damage as decided by a dermatologist"

*Comment:* no indication that the dermatologist was independent to the study

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#### Surgical hand disinfection

*Quote:* "The bag on each hand was secured and massaged for 1 min in a uniform manner by a laboratory technician"

*Comment:* no indication that the laboratory technician was blinded to the intervention

#### Hygienic hand disinfection

*Quote:* "Control microbial counts were obtained by rubbing the fingertips for 1 minute in a Petri dish containing a liquid broth using a separate dish for each hand. Either 3 ml of the hand gel or two aliquots of 3 ml of the reference alcohol were applied to the hands. The rub-in period was 30 s for the hand gels and 60 s for the reference alcohol as prescribed by EN 1500."

*Comment:* no mention of whether those supervising the process or the overseeing the microbial assays were independent of the study
Incomplete outcome data (attrition bias)

**Dermal tolerance**

*Quote:* "A dramatic finding is the very high number of subjects dropping out of the Hibiscrub group. Altogether 15 persons gave up using the Hibiscrub for reasons related to the use of the product, but there was only one person who discontinued Sterillium."

*Comment:* This total represented a large proportion of the total participants used. Despite attributing the high rates of dropout to reasons related to the product, the risk of attrition bias is fairly high in this case.

**Surgical hand disinfection**

No direct quotes, but no indication given as to the total number of participants used or whether there was any observed loss to follow-up. Therefore the judgement remains unclear.

**Hygienic hand disinfection**

*Quote:* "The antimicrobial efficacy of each product was compared with that of 60% (v/v) 2-propanol on artificially contaminated hands (E. coli K12, NCTC 10538) using a cross-over design with 15 volunteers."

*Comment:* No losses to follow-up were discussed within the results; however, there is no evidence to suggest no losses to follow-up occurred. Overall judgement therefore remains unclear for outcome assessment.

**Selective reporting (reporting bias)**

**Dermal tolerance**

No direct quotes; however, the form used to tabulate the results gives no quantitative figures for variables that were predominantly quantitative in nature. For example "D-squames", "Electrical capacity", "Trans epidermal water loss" are expressed as either "−− very poor", "− poor", "+ good" or "++ very good". It is unclear why the outcomes have been reported in this way.

**Surgical hand disinfection**

No direct quotes, although the results indicate the microbial concentration in the sampling fluid before and after treatment, outlined in the methodology and expressed as logarithm.

**Hygienic hand disinfection**

No direct quotes, although all the hand disinfectants that were discussed in the methodology as appropriate for testing are discussed and tabulated in an appropriate manner. Overall judgement of low risk.

**Other bias**

**Low risk**

No other sources of bias detected; cross-over design taken into account in analysis.

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**Methods**

**Parallel group randomised controlled trial. Unit of randomisation and analysis is individual staff member.**

- Generation of random number sequence: computerised generation
- Allocation concealment: sealed opaque envelopes
- Blinding: blinded outcome assessment
- A priori sample calculations: yes
- Antisepsis protocol: yes
Participants 164 nurses, operating department practitioners and healthcare assistants

Interventions
Group 1 - chlorhexidine (aqueous chlorhexidine gluconate 4% (Hibiscrub)) alone. Total application time of 2 min, n = 54
Group 2 - chlorhexidine as group 1 plus cleaning nails with disposable nail pick (before scrub under running water), n = 54
Group 3 - chlorhexidine as above plus cleaning nails with disposable nail brush (before scrub under running water), n = 54
All groups were observed and timed when scrubbing. Total antiseptic application time in each group was 2 min (measured dose of 2 x 2 ml for 1 min/dose).

Outcomes Primary outcome: number of CFUs on dominant hand
Method of testing: modified glove juice method (sterile Gammex Powder Free, Ansell glove)

Notes No surgical procedures were performed; all staff performed circulating duties within the operating theatre for 1 h.

Risk of bias

| Bias                                      | Authors’ judgement | Support for judgement |
|-------------------------------------------|--------------------|-----------------------|
| Random sequence generation (selection bias) | Low risk           | An appropriate method of generating the randomisation sequence was reported: "Randomisation was in random size blocks in multiples of three and was generated by a statistician using a computer software package." |
| Allocation concealment (selection bias)   | Low risk           | A recognised method for ensuring allocation concealment was reported: "Group allocation details were placed inside sequentially numbered sealed opaque envelopes by an individual independent from the study. The envelopes were opened by participants after baseline bacterial counts had been taken and immediately before the scrub intervention was performed. The researcher conducting the baseline sample was unaware of each participant's group allocation." |
| Blinding (performance bias and detection bias) All outcomes - Blinding participants | High risk | Scrub protocols differed such that all participants were aware of their allocation |
| Blinding (performance bias and detection bias) All outcomes - Blinding care givers | High risk | For personnel (not caregivers): "The researcher conducting the baseline sample was unaware of each participant’s group allocation. As the researcher observed the participants' scrubbing they were therefore aware of each participant’s group allocation when conducting the post-scrub sample.” |
| Blinding (performance bias and detection bias) All outcomes - Blinding outcome assessors | Low risk | CFU assessment was performed in a blinded manner: "Laboratory staff estimating the bacterial counts were unaware of group allocation status.” |
| Incomplete outcome data (attrition bias) All outcomes | Low risk | Data missing for only 2 participants; reasons given: "164 operating department staff took part in the study. No participants dropped out, but the laboratory
### Tanner 2009 (Continued)

Results for two people were spoiled. The findings on 162 participants are presented.

**Selective reporting (reporting bias)**  
**Low risk**  
All specified outcomes reported: “The primary purpose of the trial was to compare any two of the trial groups by measuring the difference in post-intervention CFU per hand. A secondary purpose was to identify any relationships between participants and baseline CFU counts.”

**Other bias**  
**Low risk**  
No evidence of other sources of bias

### Vergara-Fernandez 2010

**Methods**  
Parallel group randomised controlled trial. Unit of randomisation and analysis is patient  
Generation of random number sequence: not reported  
Allocation concealment: sealed envelopes, no further information  
Blinding: not reported  
A priori sample calculations: no  
Antisepsis protocol: no  
Withdrawals: no patient withdrawals for evaluation of SSI; CFUs only assessed for 20% of staff but all included in analysis.  
Intention-to-treat analysis: yes  
Clear inclusion or exclusion criteria: types of surgery only

**Participants**  
400 staff classified as surgeons, "instrumentalists" and helpers  
100 patients undergoing clean or clean-contaminated surgery

**Interventions**  
Group 1 - aqueous scrub with 4% chlorhexidine gluconate with brush and sterile water. Mean duration of scrub 3.9 (SD 1.07) min.  
Group 2 - alcohol rub with 61% ethanol, 1% chlorhexidine gluconate. Mean duration 2.0 (SD 0.47) min.

**Outcomes**  
SSI after 1 month (CDC criteria; method of diagnosis not further reported)  
CFUs on hands (20% of personnel only): reports number of personnel with positive cultures (no further detail)

**Notes**  
Only 20% of the 400 enrolled staff were assessed for bacteria on hands; these were classified as having or not having a positive culture. No data on number of CFUs on hands were reported.

### Risk of bias

| Bias                          | Authors' judgement | Support for judgement |
|-------------------------------|--------------------|-----------------------|
| Random sequence generation    | Unclear risk       | Just says "used closed envelopes" |
| Allocation concealment        | Unclear risk       | Just says "used closed envelopes" |
| Blinding (performance bias and detection bias) | Unclear risk       | No information given but interventions clearly differed |
### Vergara-Fernandez 2010 (Continued)

| Bias | Authors' judgement | Support for judgement |
|------|--------------------|-----------------------|
| Blinding (performance bias and detection bias) All outcomes - Blinding care givers | Unclear risk | No information given but interventions clearly differed |
| Blinding (performance bias and detection bias) All outcomes - Blinding outcome assessors | Unclear risk | No information on blinding of outcome assessors |
| Incomplete outcome data (attrition bias) All outcomes | Low risk | Data on all patients reported for SSI |
| | | Data on 20% of personnel collected for CFUs; results reported for all those collected |
| Selective reporting (reporting bias) | Low risk | All specified outcomes reported |
| Other bias | Low risk | No other sources of bias detected |

### Wheelock 1997

| Bias  | Authors' judgement | Support for judgement |
|-------|--------------------|-----------------------|
| Methods | Randomised cross-over trial | Generation of random number sequence: no details given Allocation concealment: no details given Blinding: no details given (dermal tolerance) A priori sample calculations: yes Antisepsis protocol: yes Withdrawals: no details given Intention-to-treat analysis: no Clear inclusion or exclusion criteria: no |
| Participants | 25 operating theatre nurses and surgical technologists Baseline comparability: age, gender, hand size, role, length of perioperative experience |
| Interventions | Group 1 - 3 min surgical scrub using either 4% chlorhexidine, 2% chlorhexidine or parachlorometaxylenol Group 2 - 2 min surgical scrub using either 4% chlorhexidine, 2%chlorhexidine or parachlorometaxylenol |
| Outcomes | Outcome measure: CFUs on participants' hands Method of testing: glove juice method Timing of testing: 1 h after antisepsis |
| Notes | Participants did not take part in any surgical procedures. |

### Risk of bias

| Bias | Authors' judgement | Support for judgement |
|------|--------------------|-----------------------|
| Random sequence generation (selection bias) | Unclear risk | Quote "We randomly assigned subjects to one of two study groups (i.e., two-minute, three-minute surgical hand scrub times)." Comment: no indication as whether a truly randomised sequence was generated |
Wheelock 1997 (Continued)

| Allocation concealment (selection bias) | Unclear risk | Quote "We randomly assigned subjects to one of two study groups (i.e., two-minute, three-minute surgical hand scrub times)."
Comment: no indication as to whether allocation was concealed to the participants or personnel. The study was a cross-over trial, therefore all the participants were likely to have undertaken the same interventions; however, the role of bias is unclear in this case.

| Blinding (performance bias and detection bias) | High risk | No direct quotes given, but no mention as to whether the subjects or the personnel were blinded to the intervention. It is very likely that the personnel were not blinded as they would be able to calculate the time spent handwashing.

| Blinding (performance bias and detection bias) | High risk | No direct quotes given, but no mention as to whether the subjects or the personnel were blinded to the intervention. It is very likely that the personnel were not blinded as they would be able to calculate the time spent handwashing.

| Blinding (performance bias and detection bias) | High risk | No direct quotes given, but investigators were responsible for measurement of time and for preparation of solution for bacterial culture, as well as the sampling and measurement of log CFU counts. Although this is an objective measure, it is reasonable to suggest that as the investigators were not blinded to the intervention then the risk of bias here could be considered as high.

| Incomplete outcome data (attrition bias) | Low risk | Quote: "Each subject was able to complete both surgical hand scrub trials, which resulted in 300 agar plates for incubation and enumeration."
Comment: adequate evidence of no loss to follow-up

| Selective reporting (reporting bias) | Low risk | The main outcome variable was the log counts of bacterial colonies found on participants’ hands after washing their hands for a specified time. This was fully represented in table format in the results section.

| Other bias | Unclear risk | Although the trial had a cross-over design, it did not appear that this was reflected in the analysis

AAS: aqueous alcohol solution; ABWA: alcohol-based water-aided; ASA: American Society of Anesthesiologists; CDC: Centers for Disease Control; CFU: colony forming units; CHG: chlorhexidine gluconate; C/S: culture and sensitivity; PVI: povidone iodine; SSI: surgical site infection; v/v: volume/volume per cent.

Characteristics of excluded studies [ordered by study ID]

| Study         | Reason for exclusion               |
|---------------|------------------------------------|
| Adjoussou 2009| Not a randomised controlled trial  |
| Aly 1983      | A laboratory-based study           |
| Aly 1988      | A laboratory-based study           |
| Aly 1998      | Evaluated patient skin preparations|
| Arata 1993    | Evaluated patient skin preparations|
| Ayliffe 1984  | A discussion paper                 |
| Study            | Reason for exclusion                                      |
|------------------|-----------------------------------------------------------|
| Ayliffe 1988     | A laboratory and ward study - not hand antisepsis         |
| Ayliffe 1990     | Hand hygiene study                                       |
| Babb 1991        | Study carried out on volunteers, not scrub staff in an operating theatre |
| Bansal 2002      | Wound irrigation study                                   |
|Barsanti 2009     | Infection in ITU                                          |
| Bartzokas 1983   | A laboratory-based study                                  |
| Bearman 2010     | Gloving study                                             |
| Beeuwkes 1986    | Participants were not randomised                          |
| Bendig 1990      | A laboratory-based study                                  |
| Bernam 2004      | Not a randomised controlled trial                         |
| Bibbo 2005       | Evaluated patient skin preparations                       |
| Blomgren 1983    | Body exhaust suit study                                  |
| Borer 2001       | Infection surveillance study                              |
| Boyce 2000a      | An editorial                                              |
| Boyce 2000b      | Evaluated skin condition rather than SSIs or CFUs        |
| Braumann 2008    | Study of dressings                                        |
| Breeze 1994      | Discussion paper                                          |
| Brooks 2001      | Evaluated patient skin preparations                       |
| Bruckner 2009    | Study of dressings                                        |
| Bryce 2001       | Not a randomised controlled trial. Participants used product A for 2 weeks then swapped to product B for the following 2 weeks. |
| Caelli 2000      | Study of MRSA decolonisation                              |
| Carro 2007       | Not a randomised controlled trial                         |
| Cheng 2001       | Literature review                                         |
| Coelho 1984      | Not relevant to this review                               |
| Cremieux 1989    | A laboratory-based study                                  |
| Crowder 1967     | Study was not randomised. Participants performed antisepsis using their usual solution. There were no comparison groups |
| Culligan 2005    | Randomised controlled trial comparing antiseptic solutions on patients' skin |
| Study                      | Reason for exclusion                                                                 |
|----------------------------|----------------------------------------------------------------------------------------|
| Curti 1974                 | Chemical agents                                                                       |
| Da Cunha 2011              | Not a randomised controlled trial                                                     |
| Dahl 1990                  | Chlorhexidine scrub which was left on the surgeons arm was compared with a surgeons arm where the chlorhexidine scrub was rinsed off. Did not meet the objectives of this review |
| Das 2005                   | Study on hygienic hand washing                                                        |
| De Castro Peraza 2010      | Study of gloves in surgery                                                            |
| Demir 2009                 | Staff survey                                                                           |
| Deshmukh 1998              | Participants were randomised to 2 groups. Group 1 participants were tested after 1 hour and group 2 participants were tested after 2 hours. Participants in both groups used product A 1 day and product B the next day |
| Dineen 1969                | Participants' hands were covered with bacterial inoculum. A laboratory-based study     |
| Dineen 1978                | A laboratory-based study                                                               |
| Dohmen 2006                | Antibiotic prophylactic study                                                         |
| Durani 2008                | Non-systematic literature review                                                      |
| Elenbaas 1982              | Dog bite study                                                                         |
| Ellenhorn 2005             | Evaluated patient skin preparations                                                    |
| Faoagali 1995              | A laboratory style study using non clinical hospital staff                             |
| Ford 2005                  | Antimicrobial suture study                                                             |
| Grabsch 2004               | Not randomised                                                                         |
| Grinbaum 1995              | A retrospective study                                                                  |
| Gruendemann 2001           | Discussion paper                                                                      |
| Guilhermetti 2001          | MRSA decontamination study                                                            |
| Hagen 1995                 | Evaluated patient skin preparations                                                    |
| Harnoss                    | Study carried out on volunteers, not scrub staff in an operating theatre               |
| Heeg 1986                  | Experimental and clinical conditions                                                  |
| Heeg 2001                  | Measured the impact of hand care products on alcohol rubs                              |
| Heeg 2008                  | Testing methods not suitable; not randomised                                           |
| Hibbard 2002a              | A laboratory-based study                                                               |
| Hibbard 2002b              | A laboratory-based study                                                               |
| Study            | Reason for exclusion                                                                 |
|------------------|--------------------------------------------------------------------------------------|
| Hingst 1992      | A laboratory-based study                                                             |
| Hobson 1998      | A laboratory-based study                                                             |
| Hubner 2006      | A laboratory-based study                                                             |
| Incoll 2009      | Skin preparation of patient                                                          |
| Jeng 1998        | A laboratory-based study                                                             |
| Jeng 2001        | A study of skin antiseptics used on patients skin                                     |
| Jones 2000       | A laboratory-based study and participants were not randomised                        |
| Joress 1962      | No comparison group was used in the first part of the trial. Comparison groups were used in the second part of the trial, but solutions were applied to the forearm rather than as surgical scrubs |
| Kampf 2005       | A laboratory-based study                                                             |
| Kargi 2008       | Anaesthetic agents study                                                             |
| Keser 2005       | Not a randomised controlled trial                                                   |
| Kikuchi 1999     | Measured condition of skin on hands of participants; did not compare CFUs or SSIs   |
| Kjellander 1960  | Not randomised                                                                       |
| Kong 1994        | Not relevant topic.                                                                  |
| Kramer 2007      | Not randomised                                                                       |
| Kramer 2008      | Descriptive paper of scrubbing methods                                               |
| Larson 1984      | Study focused on handwashing rather than hand antisepsis                             |
| Larson 1986a     | A laboratory-based study                                                             |
| Larson 1986b     | A laboratory-based study                                                             |
| Larson 1990      | A laboratory-based study                                                             |
| Larson 1993      | A laboratory-based study                                                             |
| Larson 2001a     | Study of handwashing in intensive care                                               |
| Larson 2001b     | Not randomised to appropriate groups. 5 participants were randomised to a reference group at the beginning of the study. The participants randomised to the intervention group used an alcohol rub for 3 weeks and then a surgical scrub for 3 weeks |
| Lehmann 1985     | Analgesics study                                                                     |
| Lepor 2009       | Patient skin prep study                                                              |
| Lilly 1978       | A laboratory-based study                                                             |
| Study             | Reason for exclusion                                                                 |
|------------------|---------------------------------------------------------------------------------------|
| Lio 2009         | Study on topical agents                                                               |
| Llanos 2006      | Wound management                                                                      |
| Loeb 1997        | Study carried out on volunteers, not scrub staff in an operating theatre              |
| Lowbury 1974a    | Not relevant to this review                                                          |
| Lowbury 1974b    | A laboratory-based study                                                             |
| Lung 2004        | A literature review                                                                  |
| Magann 1993      | Evaluated patient skin preparations                                                  |
| Magera 2007      | Patient skin preparation study                                                       |
| Marchetti 2003   | Laboratory-based study                                                               |
| Marra 2008       | Hand hygiene study                                                                   |
| Mathias 2000     | A discussion paper                                                                    |
| Mathias 2002     | A discussion paper                                                                    |
| McBride 1973     | A laboratory-based study                                                             |
| Meers 1978       | Not relevant topic                                                                   |
| Minakuchi 1993   | A study of handwashing rather than hand antisepsis.                                   |
| Misterka 1991    | Study of dressings                                                                    |
| Moralejo 2003    | Not a randomised controlled trial; summary of previously published studies          |
| Mulberry 2001    | A laboratory-based study                                                             |
| Murie 1980       | Cross-over trial but without any randomisation                                         |
| Nakano 2008      | Wound irrigation study                                                                |
| Nowak 1982       | Bowel prep study                                                                      |
| O'Shaughnessy 1991 | All participants carried out intervention 1 on day 1, intervention 2 on day 2 and intervention 3 on day 3. |
| Olson 2012       | Study carried out on volunteers, not scrub staff in an operating theatre              |
| Ortiz 2012       | Assessed intra-operative rescrubbing                                                  |
| Parienti 2004    | Central venous catheter study                                                        |
| Paulson 1994     | A laboratory-based study                                                              |
| Study          | Reason for exclusion                                           |
|---------------|----------------------------------------------------------------|
| Paulson 1999  | A laboratory-based study                                      |
| Peterson 1978 | A laboratory-based study                                      |
| Phimolsarnit 1986 | Not randomised                                                |
| Poon 1998     | Not randomised                                                |
| Rehork 1991   | Study carried out on volunteers, not scrub staff in an operating theatre |
| Reid 1991     | Study of wound management                                     |
| Reverdy 1984  | A laboratory-based study                                      |
| Rotter 1980   | A laboratory-based study                                      |
| Rotter 1984   | Study of handwashing rather than hand antisepsis              |
| Rotter 1986   | A laboratory-based study                                      |
| Rotter 1998   | A laboratory-based study                                      |
| Rotter 2005   | Explores hand hygiene rather than hand antisepsis             |
| Rotter 2006   | Laboratory-based study                                        |
| Sattar 2000   | A laboratory-based study                                      |
| Scheibel 1991 | Study of clean air systems                                    |
| Scott 1991    | Evaluated user satisfaction                                   |
| Sensoz 2003   | No evidence that the study was randomised                     |
| Serra 2005    | Study of dressings                                            |
| Shirahatti 1993 | Evaluated patient skin preparations                          |
| Simor 2007    | MRSA decolonisation                                           |
| Springer 2002 | Discussion paper                                              |
| Stahl 2007    | Lab based study of effect of saline in antimicrobial skin preparations |
| Starr 2005    | Evaluated patient skin preparations                           |
| Stevenson 2003 | Study of antibiotics                                        |
| Sullivan 2008 | Patient skin preparation study                                |
| Tanner 2008   | Earlier systematic review                                     |
| Thiele 2008   | Literature review                                             |
| Tucci 1977    | Not a randomised controlled trial; no control group           |
## Characteristics of studies awaiting assessment [ordered by study ID]

| Study           | Reason for exclusion                                                                 |
|-----------------|--------------------------------------------------------------------------------------|
| Valente 2003    | Wound irrigation study                                                               |
| Vogt 2006       | Evaluated iodine based wound dressings                                               |
| Voss 1997       | Looked at compliance with various handwashing methods                                 |
| Vossinakis 2004 | Study of local anaesthetic                                                            |
| Walwaikar 2002  | Each intervention group contained a scrub solution, a patient prep solution and a follow-up wound cleansing product. It was not possible to look at the effect of the scrub solution on its own |
| Waterman 2006   | Study of glove juice and rings                                                        |
| Webster 1989    | Study of handwashing in a neo-natal unit                                              |
| Wernze 1975     | Anaesthetics study                                                                    |
| Whittaker 2005  | Study of antibiotics in hand injuries                                                |
| Yeung 2007      | Hand hygiene literature review                                                       |
| Yuldashehkan 2008 | Not a randomised controlled trial                                                   |
| Zaragoza 1999   | A study of handwashing, not hand antisepsin.                                          |

**MRSA**: methicillin-resistant Staphylococcus aureus

## Data and Analyses

**Comparison 1. basic hand hygiene versus alcohol rub**

| Outcome or subgroup title | No. of studies | No. of participants | Statistical method                          | Effect size  |
|---------------------------|----------------|---------------------|---------------------------------------------|--------------|
| 1 SSI                     | 1              | 3133                | Risk Ratio (M-H, Fixed, 95% CI)             | 0.97 [0.77, 1.23] |
### Analysis 1.1. Comparison 1 basic hand hygiene versus alcohol rub, Outcome 1 SSI.

| Study or subgroup | Scrub (soap and water) | Alcohol rub | Risk Ratio | Weight | Risk Ratio |
|-------------------|------------------------|-------------|------------|--------|------------|
|                   | n/N                    | n/N         | M-H, Fixed, 95% CI |        | M-H, Fixed, 95% CI |
| Nthumba 2010      | 128/1596               | 127/1537    | 100%       | 0.97[0.77,1.23] |

Total (95% CI): 1596/1537

Total events: 128 (Scrub (soap and water)), 127 (Alcohol rub)

Heterogeneity: Not applicable

Test for overall effect: Z =0.25(P=0.8)

Favours soap and water

### Comparison 2. chlorhexidine versus iodine

| Outcome or subgroup title | No. of studies | No. of participants | Statistical method | Effect size |
|---------------------------|----------------|---------------------|--------------------|-------------|
| 1 CFUs                    | 3              |                     | Mean Difference (IV, Fixed, 95% CI) | Totals not selected |
| 1.1 CFUs immediately after antisepsis | 3 | | Mean Difference (IV, Fixed, 95% CI) | 0.0 [0.0, 0.0] |
| 1.2 CFUs 2 h after initial antisepsis | 1 | | Mean Difference (IV, Fixed, 95% CI) | 0.0 [0.0, 0.0] |
| 1.3 CFUs 2 h after subsequent antisepsis | 1 | | Mean Difference (IV, Fixed, 95% CI) | 0.0 [0.0, 0.0] |
| 1.4 CFUs after surgical procedure | 1 | | Mean Difference (IV, Fixed, 95% CI) | 0.0 [0.0, 0.0] |

### Analysis 2.1. Comparison 2 chlorhexidine versus iodine, Outcome 1 CFUs.

| Study or subgroup | Chlorhexidine | Iodine | Mean Difference | Mean Difference |
|-------------------|---------------|--------|----------------|----------------|
|                   | N Mean(SD)    | N      | Fixed, 95% CI  | Fixed, 95% CI  |
| 2.1.1 CFUs immediately after antisepsis | | | | |
| Furukawa 2005     | 11 0.1 (0.4) | 11 2.5 (1.4) | -2.4[3.26,-1.54] |
| Herruzo 2000      | 50 18 (6)     | 49 66 (7) | -48[50.57,-45.43] |
| Pereira 1990      | 34 4 (0.7)    | 34 4.3 (0.6) | -0.34[0.64,-0.04] |
| Pereira 1990      | 34 4.1 (0.2) | 34 4.3 (0.2) | -0.17[-0.28,-0.06] |

| 2.1.2 CFUs 2 h after initial antisepsis | | | | |
| Pereira 1990      | 34 3.6 (0.6) | 34 4.4 (0.7) | -0.75[-1.06,-0.44] |
| Pereira 1990      | 34 3.8 (0.6) | 34 4.2 (0.7) | -0.41[-0.72,-0.1] |

| 2.1.3 CFUs 2 h after subsequent antisepsis | | | | |
| Pereira 1990      | 34 3.4 (0.8) | 34 4.5 (0.5) | -1.1[-1.42,-0.78] |

Favours Chlorhex. -1 -0.5 0 0.5 1 Favours Iodine
| Study or subgroup | Chlorhexidine | Iodine | Mean Difference | Mean Difference |
|------------------|---------------|--------|-----------------|-----------------|
|                  | N  Mean(SD)   | N      | Mean(SD)        | Fixed, 95% CI   |
| Pereira 1990     | 34 4 (0.6)    | 34 4.7 (0.5) | -0.65 [-0.93, -0.37] |

### 2.1.4 CFUs after surgical procedure

|                  |          |        |               |               |
|------------------|----------|--------|---------------|---------------|
| Herruzo 2000     | 50 37 (11) | 49 169 (31) | -132 [-141.2, -122.8] |

Favours Chlorhex. -1 -0.5 0 0.5 1 Favours Iodine

**Comparison 3. chlorhexidine versus iodine plus triclosan**

| Outcome or subgroup title | No. of studies | No. of participants | Statistical method | Effect size |
|---------------------------|----------------|---------------------|--------------------|-------------|
| 1 CFUs                    | 1              |                     | Mean Difference (IV, Fixed, 95% CI) | Totals not selected |
| 1.1 CFUs immediately after antisepsis | 1          |                     | Mean Difference (IV, Fixed, 95% CI) | 0.0 [0.0, 0.0] |
| 1.2 CFUs 2 h after initial antisepsis | 1           |                     | Mean Difference (IV, Fixed, 95% CI) | 0.0 [0.0, 0.0] |
| 1.3 CFUs 2 h after subsequent antisepsis | 1           |                     | Mean Difference (IV, Fixed, 95% CI) | 0.0 [0.0, 0.0] |

### Analysis 3.1. Comparison 3 chlorhexidine versus iodine plus triclosan, Outcome 1 CFUs.

| Study or subgroup                  | Chlorhexidine | Iodine triclosan | Mean Difference | Mean Difference |
|-----------------------------------|---------------|------------------|-----------------|-----------------|
|                                  | N  Mean(SD)   | N  Mean(SD)      | Fixed, 95% CI   | Fixed, 95% CI   |
| 3.1.1 CFUs immediately after antisepsis | Pereira 1997 | 23 4.4 (0.7) | 23 4.8 (0.9) | -0.38 [-0.84, 0.08] |
| 3.1.2 CFUs 2 h after initial antisepsis | Pereira 1997 | 23 4.1 (1) | 23 4.4 (0.8) | -0.38 [-0.9, 0.14] |
| 3.1.3 CFUs 2 h after subsequent antisepsis | Pereira 1997 | 23 4.1 (0.9) | 23 4.8 (0.6) | -0.69 [-1.13, -0.25] |

Favours Chlorhex -2 -1 0 1 2 Favours Iodine triclosan

### Comparison 4. alcohol rub versus other alcohol rub

| Outcome or subgroup title | No. of studies | No. of participants | Statistical method | Effect size |
|---------------------------|----------------|---------------------|--------------------|-------------|
| 1 CFUs                    | 1              |                     | Mean Difference (IV, Fixed, 95% CI) | Totals not selected |
### Analysis 4.1. Comparison 4 alcohol rub versus other alcohol rub, Outcome 1 CFUs.

| Study or subgroup | Isopropanol | Ethanol | Mean Difference | Mean Difference |
|-------------------|-------------|---------|----------------|----------------|
| N                 | Mean(SD)    | N       | Mean(SD)       | Fixed, 95% CI  |
| 4.1.1 Immediately after antisepsis |             |         | 0.0 [-0.57, 0.57] |               |
| 23                | 4 (1)       | 23      | 4 (1)          |                |
| Pereira 1997      |             |         |                |                |
| 4.1.2 2 h after initial antisepsis | 23                | 3.5 (0.8) | 23 | 3.5 (1) | 0.07 [-0.45, 0.59] |
| Pereira 1997      | 23          | 3.5 (1) |                |                |
| 4.1.3 2 h after subsequent antisepsis | 23                | 3.9 (1) | 23 | 3.8 (1.1) | 0.11 [-0.49, 0.71] |
| Pereira 1997      | 23          | 3.9 (1) |                |                |

| Favours Isopropanol | -2 | -1 | 0 | 1 | 2 |
|----------------------|----|----|---|---|---|
| Favours Ethanol      |    |    |   |   |   |

### Comparison 5. scrub versus alcohol-only rub

| Outcome or subgroup title | No. of studies | No. of participants | Statistical method | Effect size |
|---------------------------|----------------|---------------------|-------------------|-------------|
| 1 SSI                     | 1              | 500                 | Risk Ratio (M-H, Fixed, 95% CI) | 0.56 [0.23, 1.34] |

### Analysis 5.1. Comparison 5 scrub versus alcohol-only rub, Outcome 1 SSI.

| Study or subgroup | Alcohol rub | Aqueous scrub | Risk Ratio | Weight | Risk Ratio |
|-------------------|-------------|---------------|------------|--------|------------|
| n/N               | n/N         | M-H, Fixed, 95% CI |            |        | M-H, Fixed, 95% CI |
| Al-Naami 2009     | 8/272       | 12/228        | 0.56[0.23,1.34] | 100%   |            |
| Total (95% CI)    | 272         | 228           | 0.56[0.23,1.34] | 100%   |            |

Total events: 8 (Alcohol rub), 12 (Aqueous scrub)
Heterogeneity: Not applicable
Test for overall effect: Z=1.3(P=0.19)
### Comparison 6. scrub versus alcohol rub

| Outcome or subgroup title | No. of studies | No. of participants | Statistical method | Effect size |
|--------------------------|----------------|---------------------|--------------------|-------------|
| 1 SSI                    | 2              |                     | Risk Ratio (M-H, Fixed, 95% CI) | Subtotals only |

**Analysis 6.1. Comparison 6 scrub versus alcohol rub, Outcome 1 SSI.**

| Study or subgroup | 4% chlorhexidine | Alcohol rub | Risk Ratio | Weight | Risk Ratio |
|-------------------|------------------|-------------|------------|--------|------------|
| Parienti 2002     | 53/2135          | 55/2252     | M-H, Fixed, 95% CI | 0% | 1.02[0.7,1.48] |
| Vergara-Fernandez 2010 | 1/50             | 2/50        | M-H, Fixed, 95% CI | 0% | 0.5[0.05,5.34] |

Favours chlorhexidine

| Study or subgroup |  |  |  |  |  |
|-------------------|---|---|---|---|---|
|                   | 0.1| 0.1| 1| 10| 100|
|                   |  |
|                   | Favours chlorhexidine | Favours alcohol rub |

### Comparison 7. scrub (chlorhexidine) versus alcohol rub + additional ingredient

| Outcome or subgroup title | No. of studies | No. of participants | Statistical method | Effect size |
|---------------------------|----------------|---------------------|--------------------|-------------|
| 1 CFUs                    | 1              |                     | Mean Difference (IV, Fixed, 95% CI) | Totals not selected |
| 1.1 Immediately after antisepsis | 1              |                     | Mean Difference (IV, Fixed, 95% CI) | 0.0 [0.0, 0.0] |
| 1.2 After surgical procedure | 1              |                     | Mean Difference (IV, Fixed, 95% CI) | 0.0 [0.0, 0.0] |

**Analysis 7.1. Comparison 7 scrub (chlorhexidine) versus alcohol rub + additional ingredient, Outcome 1 CFUs.**

| Study or subgroup | Chlorhexidine Scrub | N duopropenide Rub | Mean Difference Fixed, 95% CI | Mean Difference Fixed, 95% CI |
|-------------------|---------------------|--------------------|-----------------------------|-----------------------------|
| 7.1.1 Immediately after antisepsis | 50 | 55 | 1 (0) | Not estimable |
| Herruzo 2000      | 50 | 18 (6) | 37 (11) | 55 | 1 (0) | Not estimable |

Favours Scrub

| Study or subgroup |  |  |  |  |  |
|-------------------|---|---|---|---|---|
|                   | -10| -5| 0 | 5 | 10|
|                   | Favours Scrub | Favours Rub |

### Comparison 8. scrub (povidone iodine) versus alcohol rub + additional ingredient

| Outcome or subgroup title | No. of studies | No. of participants | Statistical method | Effect size |
|---------------------------|----------------|---------------------|--------------------|-------------|
| 1 CFUs                    | 1              |                     | Mean Difference (IV, Fixed, 95% CI) | Totals not selected |
| Outcome or subgroup title | No. of studies | No. of participants | Statistical method | Effect size |
|---------------------------|--------------|-------------------|-------------------|-------------|
| 1.1 Immediately after antisepsis | 0 | | Mean Difference (IV, Fixed, 95% CI) | 0.0 [0.0, 0.0] |
| 1.2 After surgical procedure | 1 | | Mean Difference (IV, Fixed, 95% CI) | 0.0 [0.0, 0.0] |

### Analysis 8.1. Comparison 8 scrub (povidone iodine) versus alcohol rub + additional ingredient, Outcome 1 CFUs.

#### 8.1.1 Immediately after antisepsis

| Study or subgroup | Iodine Scrub | Duopropenide Rub | Mean Difference |
|------------------|--------------|------------------|-----------------|
| Herruzo 2000     | 49 Mean(31)  | 55 Mean(1)      | Not estimable   |
| Herruzo 2000     | 49 Mean(7)   | 55 Mean(0)      | Not estimable   |

### Analysis 9.1. Comparison 9 scrub (chlorhexidine) versus rub + additional ingredient, Outcome 1 CFUs.

#### 9.1.1 Immediately after antisepsis

| Study or subgroup | Rub | Scrub | Mean Difference |
|------------------|-----|-------|-----------------|
| Pietsch 2001     | 75 Mean(0.1) | 75 Mean(0.1) | -1.27 [-1.31, -1.23] |

### Comparison 9. scrub (chlorhexidine) versus rub + additional ingredient

| Outcome or subgroup title | No. of studies | No. of participants | Statistical method | Effect size |
|---------------------------|--------------|-------------------|-------------------|-------------|
| 1 CFUs                    | 1            |                  | Mean Difference (IV, Fixed, 95% CI) | Totals not selected |
| 1.1 Immediately after antisepsis | 1 | | Mean Difference (IV, Fixed, 95% CI) | 0.0 [0.0, 0.0] |
| 1.2 After surgical procedure | 1 | | Mean Difference (IV, Fixed, 95% CI) | 0.0 [0.0, 0.0] |
### Comparison 10. scrub (chlorhexidine) versus alcohol rub + additional ingredient

| Outcome or subgroup title | No. of studies | No. of participants | Statistical method | Effect size |
|---------------------------|----------------|---------------------|--------------------|-------------|
| 1 CFUs                    | 1              | 53                  | Mean Difference (IV, Fixed, 95% CI) | -135.6 [-153.39, -117.81] |

#### Analysis 10.1. Comparison 10 scrub (chlorhexidine) versus alcohol rub + additional ingredient, Outcome 1 CFUs.

| Study or subgroup | Chlorhexine Scrub N Mean(SD) | Alcohol Rub N Mean(SD) | Mean Difference Fixed, 95% CI | Weight | Mean Difference Fixed, 95% CI |
|-------------------|-------------------------------|------------------------|-----------------------------|--------|-------------------------------|
| Hajipour 2006     | 25 3.1 (6.1)                  | 28 138.7 [47.6]        | -135.6 [-153.39, -117.81]   | 100%   | 100%                          |
| Total ***         | 25                            | 28                     | -135.6 [-153.39, -117.81]   | 100%   | 100%                          |

Heterogeneity: Not applicable
Test for overall effect: Z=14.94(P<0.0001)

### Comparison 11. duration - Kappstein (5 minutes versus 3 minutes)

| Outcome or subgroup title | No. of studies | No. of participants | Statistical method | Effect size |
|---------------------------|----------------|---------------------|--------------------|-------------|
| 1 CFUs immediately after antisepsis | 1              | 48                  | Mean Difference (IV, Fixed, 95% CI) | 0.26 [0.14, 0.38] |

#### Analysis 11.1. Comparison 11 duration - Kappstein (5 minutes versus 3 minutes), Outcome 1 CFUs immediately after antisepsis.

| Study or subgroup | 5 minutes N Mean(SD) | 3 minutes N Mean(SD) | Mean Difference Fixed, 95% CI | Weight | Mean Difference Fixed, 95% CI |
|-------------------|----------------------|----------------------|-----------------------------|--------|-------------------------------|
| Kappstein 1993    | 24 4.8 (0.2)         | 24 4.6 (0.2)         | 0.26 [0.14, 0.38]           | 100%   | 100%                          |
| Total ***         | 24                   | 24                   | 0.26 [0.14, 0.38]           | 100%   | 100%                          |

Heterogeneity: Not applicable
Test for overall effect: Z=4.19(P<0.0001)

### Comparison 12. duration - 5 + 3 min versus 3 + 0.5 min with chlorhexidine

| Outcome or subgroup title | No. of studies | No. of participants | Statistical method | Effect size |
|---------------------------|----------------|---------------------|--------------------|-------------|
| 1 CFUs                    | 1              |                     | Mean Difference (IV, Fixed, 95% CI) | Totals not selected |
| Outcome or subgroup title | No. of studies | No. of participants | Statistical method | Effect size |
|--------------------------|---------------|---------------------|--------------------|------------|
| 1.1 Immediately after antisepsis | 1 | | Mean Difference (IV, Fixed, 95% CI) | 0.0 [0.0, 0.0] |
| 1.2 2 h after initial antisepsis | 1 | | Mean Difference (IV, Fixed, 95% CI) | 0.0 [0.0, 0.0] |
| 1.3 2 h after subsequent antisepsis | 1 | | Mean Difference (IV, Fixed, 95% CI) | 0.0 [0.0, 0.0] |

**Analysis 12.1. Comparison 12 duration - 5 + 3 min versus 3 + 0.5 min with chlorhexidine), Outcome 1 CFUs.**

| Study or subgroup | 5 minutes | 3 minutes | Mean Difference | Mean Difference |
|-------------------|-----------|-----------|----------------|----------------|
|                    | N Mean(SD)| N Mean(SD)| (Fixed, 95% CI) | (Fixed, 95% CI) |
| 12.1.1 Immediately after antisepsis | | | | |
| Pereira 1990 | 34 4 (0.7) | 34 4.2 (0.6) | -0.19 [-0.51, 0.13] | |
| 12.1.2 2 h after initial antisepsis | | | | |
| Pereira 1990 | 34 3.6 (0.6) | 34 3.8 (0.6) | -0.23 [-0.52, 0.06] | |
| 12.1.3 2 h after subsequent antisepsis | | | | |
| Pereira 1990 | 34 3.4 (0.8) | 34 4 (0.6) | -0.58 [-0.92, -0.24] | |

**Comparison 13. duration - 5 + 3 min versus 3 + 0.5 minutes with iodine)**

| Outcome or subgroup title | No. of studies | No. of participants | Statistical method | Effect size |
|--------------------------|---------------|---------------------|--------------------|------------|
| 1 CFUs | 1 | | Mean Difference (IV, Fixed, 95% CI) | Totals not selected |
| 1.1 Immediately after antisepsis | 1 | | Mean Difference (IV, Fixed, 95% CI) | 0.0 [0.0, 0.0] |
| 1.2 After initial antisepsis | 1 | | Mean Difference (IV, Fixed, 95% CI) | 0.0 [0.0, 0.0] |
| 1.3 2 h after subsequent antisepsis | 1 | | Mean Difference (IV, Fixed, 95% CI) | 0.0 [0.0, 0.0] |

**Analysis 13.1. Comparison 13 duration - 5 + 3 min versus 3 + 0.5 minutes with iodine), Outcome 1 CFUs.**

| Study or subgroup | 5 minutes | 3 minutes | Mean Difference | Mean Difference |
|-------------------|-----------|-----------|----------------|----------------|
|                    | N Mean(SD)| N Mean(SD)| (Fixed, 95% CI) | (Fixed, 95% CI) |
| 13.1.1 Immediately after antisepsis | | | | |
| Pereira 1990 | 34 4.3 (0.6) | 34 4.3 (0.7) | 0.02 [-0.28, 0.32] | |

**Surgical hand antisepsis to reduce surgical site infection (Review)**

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### Study or subgroup

| 5 minutes | 3 minutes | Mean Difference | Mean Difference |
|-----------|-----------|-----------------|-----------------|
| N Mean(SD) | N Mean(SD) | Fixed, 95% CI | Fixed, 95% CI |

#### 13.1.2 After initial antisepsis

| Pereira 1990 |
|-------------|
| 34          |
| 4.4 (0.7)   |
| 4.2 (0.7)   |
| 0.12 [-0.21, 0.45] |

#### 13.1.3 2 h after subsequent antisepsis

| Pereira 1990 |
|-------------|
| 34          |
| 4.5 (0.5)   |
| 4.7 (0.5)   |
| -0.13 [-0.37, 0.11] |

### Comparison 14. duration - 5 + 3.5 min versus 3 + 2.5 min chlorhexidine)

| Outcome or subgroup title | No. of studies | No. of participants | Statistical method | Effect size |
|---------------------------|----------------|---------------------|-------------------|-------------|
| 1 CFUs                    | 1              | 108                 | Mean Difference (IV, Fixed, 95% CI) | 0.24 [-0.03, 0.51] |
| 1.1 Immediately after antisepsis | 1           | 4.5 (1)           | Mean Difference (IV, Fixed, 95% CI) | 0.08 [-0.44, 0.6] |
| 1.2 2 h after initial antisepsis | 1           | 3.9 (1.1)         | Mean Difference (IV, Fixed, 95% CI) | -0.19 [-0.78, 0.4] |
| 1.3 2 h after subsequent antisepsis | 1           | 4 (1)             | Mean Difference (IV, Fixed, 95% CI) | -0.17 [-0.71, 0.37] |

### Analysis 14.1. Comparison 14 duration - 5 + 3.5 min versus 3 + 2.5 min chlorhexidine), Outcome 1 CFUs.

| Study or subgroup    | 5 minutes Mean(SD) | 3 minutes Mean(SD) | Mean Difference Fixed, 95% CI | Mean Difference Fixed, 95% CI |
|----------------------|--------------------|--------------------|-------------------------------|-------------------------------|
| 14.1.1 Immediately after antisepsis Pereira 1997 | 23 | 4.5 (1) | 23 | 4.4 (0.7) | 0.08 [-0.44, 0.6] |
| 14.1.2 2 h after initial antisepsis Pereira 1997 | 23 | 3.9 (1.1) | 23 | 4.1 (0.9) | -0.19 [-0.78, 0.4] |
| 14.1.3 2 h after subsequent antisepsis Pereira 1997 | 23 | 4 (1) | 23 | 4.1 (0.9) | -0.17 [-0.71, 0.37] |

### Comparison 15. scrub versus scrub plus brush

| Outcome or subgroup title | No. of studies | No. of participants | Statistical method | Effect size |
|---------------------------|----------------|---------------------|-------------------|-------------|
| 1 CFUS                    | 1              | 108                 | Mean Difference (IV, Fixed, 95% CI) | 0.24 [-0.03, 0.51] |
### Analysis 15.1. Comparison 15 scrub versus scrub plus brush, Outcome 1 CFUs.

| Study or subgroup | Experimental | Control | Weight | Mean Difference | Mean Difference |
|-------------------|--------------|---------|--------|-----------------|----------------|
|                   | N Mean(SD)   | N Mean(SD) |        |                 |                |
| Tanner 2009       | 54 3.1 (0.7) | 54 2.9 (0.7) | 100%  | 0.24 [0.03, 0.51] |                |
| **Total *** **    | 54           | 54      | 100%  | **0.24 [0.03, 0.51]** |                |

Heterogeneity: Tau^2 = 0; Chi^2 = 0, df=0 (P=0.0001); I^2 = 100%
Test for overall effect: Z=1.77 (P=0.08)

Favours scrub plus brush: -100, -50, 0, 50, 100
Favours scrub only: -100, -50, 0, 50, 100

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### Comparison 16. scrub versus scrub plus nail pick

| Outcome or subgroup title | No. of studies | No. of participants | Statistical method | Effect size |
|---------------------------|----------------|---------------------|--------------------|-------------|
| 1 CFUs                    | 1              | 108                 | Mean Difference (IV, Fixed, 95% CI) | 0.13 [-0.14, 0.40] |

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### Analysis 16.1. Comparison 16 scrub versus scrub plus nail pick, Outcome 1 CFUs.

| Study or subgroup | Experimental | Control | Weight | Mean Difference | Mean Difference |
|-------------------|--------------|---------|--------|-----------------|----------------|
|                   | N Mean(SD)   | N Mean(SD) |        |                 |                |
| Tanner 2009       | 54 3 (0.7)   | 54 2.9 (0.7) | 100%  | 0.13 [-0.14, 0.4] |                |
| **Total *** **    | 54           | 54      | 100%  | **0.13 [-0.14, 0.4]** |                |

Heterogeneity: Not applicable
Test for overall effect: Z=0.95 (P=0.34)

Favours scrub plus nail pick: -100, -50, 0, 50, 100
Favours scrub alone: -100, -50, 0, 50, 100

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### Comparison 17. scrub plus brush versus scrub plus nail pick

| Outcome or subgroup title | No. of studies | No. of participants | Statistical method | Effect size |
|---------------------------|----------------|---------------------|--------------------|-------------|
| 1 CFUs                    | 1              | 108                 | Mean Difference (IV, Fixed, 95% CI) | 0.11 [-0.16, 0.38] |

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### Analysis 17.1. Comparison 17 scrub plus brush versus scrub plus nail pick, Outcome 1 CFUs.

| Study or subgroup | Scrub plus brush | Scrub plus nail pick | Weight | Mean Difference | Mean Difference |
|-------------------|------------------|----------------------|--------|-----------------|----------------|
|                   | N Mean(SD)       | N Mean(SD)           |        |                 |                |
| Tanner 2009       | 54 3.1 (0.7)     | 54 3 (0.7)           | 100%  | 0.11 [-0.16, 0.38] |                |
| **Total *** **    | 54               | 54                   | 100%  | **0.11 [-0.16, 0.38]** |                |

Heterogeneity: Tau^2 = 0; Chi^2 = 0, df=0 (P=0.0001); I^2 = 100%

Favours scrub plus brush: -100, -50, 0, 50, 100
Favours scrub plus nail pick: -100, -50, 0, 50, 100
| Study or subgroup | Scrub plus brush | Scrub plus nail pick | Mean Difference | Weight | Mean Difference |
|------------------|-----------------|---------------------|-----------------|--------|----------------|
|                  | N   | Mean(SD) | N   | Mean(SD) | Fixed, 95% CI | Fixed, 95% CI |
| Test for overall effect: Z=0.81 (P=0.42) | | | | | |
| Favours scrub brush | -100 | -50 | 0 | 50 | 100 | Favours scrub plus nail pick |

Surgical hand antisepsis to reduce surgical site infection (Review)
### Table 1. Overview of included studies

| Study       | 1          | 2          | 3          | 4          | 5          | Country            | Trial involved surgery                                                                 | SSI | CFU |
|-------------|------------|------------|------------|------------|------------|--------------------|----------------------------------------------------------------------------------------|-----|-----|
| Al-Naami 2009 | Aqueous scrub | Alcohol rub | NA         | NA         | NA         | Saudi Arabia       | Clean and clean-contaminated operations. Mainly abdominal.                             | ✓   | #   |
| Furukawa 2005 | Aqueous scrub | Aqueous scrub | NA         | NA         | NA         | Japan              | #                                                                                       | #   | ✓   |
| Gupta 2007   | Aqueous scrub | Alcohol rub + active ingredient | NA         | NA         | NA         | USA                | Ophthalmic, podiatric and general surgery                                               | ✓   | #   |
| Hajipour 2006 | Aqueous scrub | Alcohol rub + active ingredient | alcohol rub + active ingredient | NA         | NA         | UK                 | Trauma                                                                                 | #   | ✓   |
| Herruzo 2000 | Aqueous scrub | Aqueous scrub | NA         | NA         | NA         | Spain              | Plastic surgery and traumatology                                                        | ✓   | #   |

- **Additional Tables**
- Cochrane Library: Trusted evidence. Informed decisions. Better health.
| Study             | Participants | Methodology | Antisepsis 1 | Antisepsis 2 | Antisepsis 3 | Country | Guidelines | Compliance | Notes |
|-------------------|--------------|-------------|--------------|--------------|--------------|---------|------------|------------|-------|
| Kappstein 1993*   | n = 24 surgeons | Aqueous scrub 1 (duration 1) | Aqueous scrub 2 (duration 2) | NA | NA | NA | Germany | # | # | ✓ | Before antisepsis and immediately after antisepsis; glove juice method |
| Nthumba 2010      | n = 66 surgical staff and 3317 patients | Alcohol rub + active ingredient | Standard hand hygiene | NA | NA | NA | Kenya | ✓ | ✓ | # | Clean and clean-contaminated operations. Mixed surgery types. |
| Parienti 2002     | n = 4387 patients | Aqueous scrub | Alcohol rub + active ingredient | NA | NA | NA | France | ✓ | ✓ | # | Mix of procedures; CDC guidelines |
| Pereira 1990a     | n = 34 nurses | Aqueous scrub 1 Duration 1 | Aqueous scrub 2 Duration 1 | Aqueous scrub 1 Duration 2 | Aqueous scrub 2 Duration 2 | NA | Australia | # | # | ✓ | Immediately after antisepsis, 2 hours after initial antisepsis; 2 hours after subsequent antisepsis; glove juice method |
| Pereira 1997      | n = 34 operating room nurses | Aqueous scrub 1 (duration 1) | Aqueous scrub 2 (duration 2) | Aqueous scrub 3 (duration 2) | Alcohol rub + active ingredient 1 | Alcohol rub + active ingredient 2 | Australia | # | # | ✓ | Immediately after antisepsis, 2 hours after initial antisepsis, 2 hours after subsequent antisepsis; glove juice method |
| Pietsch 2001      | n = 75 surgeons | Aqueous scrub | Alcohol rub + active ingredient | NA | NA | NA | Germany | ✓ | # | ✓ | Immediately after antisepsis and after surgical procedure completed; |

Table 1. Overview of included studies (Continued)
Table 1. Overview of included studies (Continued)

| Study            | Method                      | Comparison 1 | Comparison 2 | Country | Demo | San | Time/Hand Juicy Method |
|------------------|-----------------------------|--------------|--------------|---------|------|----|------------------------|
| Tanner 2009      | Aqueous scrub               | Aqueous scrub + nail pick | NA          | NA      | UK   | # | # | glove juice method     |
| n= 164 staff     | Aqueous scrub + nail brush  | NA           | NA           | 1 hour After antisepsis; |
|                  |                             |              |              | modified glove juice method |
| Vergara-Fernandez 2010 | Aqueous scrub | Alcohol rub + active ingredient | NA          | NA      | Mexico | ✓ | ✓ | Only 20% of the 400 enrolled staff were assessed for bacteria on hands; data not included |
| n = 100 patients | Aqueous scrub + nail brush  | NA           | NA           | Clean and clean-contaminated operations. Mixed surgery types. |
|                   |                             |              |              | CDC guidelines |
| Wheelock 1997     | Aqueous scrub 1 (duration 1) | Aqueous scrub 2 (duration 2) | NA          | NA      | USA  | # | # | glove juice method     |
| n = 25 operating theatre nurses and surgical technologists | NA | NA | NA | 1 hour after antisepsis; |
|                  |                             |              |              | glove juice method |

NA: not applicable
APPENDICES

Appendix 1. Search methods used in previous versions

Original review (2007)

- Cochrane Wounds Group Specialised Register (Searched 12 June 2007);
- The Cochrane Central Register of Controlled Trials (CENTRAL) (The Cochrane Library 2007, Issue 2);
- Ovid MEDLINE (2005 to May Week 5 2007);
- Ovid EMBASE (2005 to 2007 Week 23);
- Ovid CINAHL (2005 to June Week 2 2007);
- ZETOC database of conference proceedings was searched from 1993 to 2005.

The following search strategy was used for searching CENTRAL:

1 MeSH descriptor Surgical Wound Infection explode all trees
2 surgical NEAR infection*
3 surgical NEAR wound*
4 (post-operative or postoperative) NEAR (wound NEXT infection*)
5 MeSH descriptor Preoperative Care explode all trees
6 MeSH descriptor Perioperative Care explode all trees
7 preoperative or pre-operative
8 (#1 OR #2 OR #3 OR #4 OR #5 OR #6 OR #7)
9 MeSH descriptor Skin explode all trees
10 MeSH descriptor Antisepsis explode all trees
11 (#9 AND #10)
12 antiseptic
13 MeSH descriptor Iodine explode all trees
14 MeSH descriptor Iodophors explode all trees
15 MeSH descriptor Povidone-Iodine explode all trees
16 MeSH descriptor Chlorhexidine explode all trees
17 MeSH descriptor Alcohols explode all trees
18 MeSH descriptor Soaps explode all trees
19 MeSH descriptor Detergents explode all trees
20 MeSH descriptor Disinfection explode all trees
21 iodophor* or povidone-iodine or betadine or chlorhexidine or alcohol or alcohols or antiseptic* or soap* or detergent* or disinfect*
22 (#11 OR #12 OR #13 OR #14 OR #15 OR #16 OR #17 OR #18 OR #19 OR #20 OR #21)
23 MeSH descriptor Handwashing explode all trees
24 hand or hands or handwash* or surgical scrub*)
25 (#23 OR #24)
26 (#8 AND #22 AND #25)

Appendix 2. Ovid MEDLINE search strategy

1 exp Surgical Wound Infection/
2 exp Surgical Wound Dehiscence/
3 (surg* adj5 infect*).tw.
4 (surg* adj5 wound*).tw.
5 (surg* adj5 site*).tw.
6 (surg* adj5 incision*).tw.
7 (surg* adj5 dehiscen*).tw.
8 ((post-operative or postoperative) adj5 wound infection*).tw.
9 exp Preoperative Care/
10 exp Perioperative Care/
11 ((preoperative or pre-operative) adj care).tw.
12 or/1-11
13 exp Skin/
14 exp Antisepsis/
15 and/13-14
Surgical hand antisepsis to reduce surgical site infection (Review)

Appendix 3. Ovid EMBASE search strategy

1 exp Surgical Wound Infection/
2 exp Surgical Wound Dehiscence/
3 (surg* adj5 infect*).tw.
4 (surg* adj5 wound*).tw.
5 (surg* adj5 site*).tw.
6 (surg* adj5 incision*).tw.
7 (surg* adj5 dehiscen*).tw.
8 ((post-operative or postoperative) adj5 wound infection*).tw.
9 exp Preoperative Care/
10 exp Perioperative Care/
11 ((preoperative or pre-operative) adj care).tw.
12 or/1-11
13 exp Skin/
14 exp Antisepsis/
15 and/13-14
16 skin antisept*.tw.
17 exp Anti-Infective Agents, Local/
18 exp Iodophors/
19 exp Povidone-Iodine/
20 exp Chlorhexidine/
21 exp Alcohols/
22 exp Soaps/
23 (iodophor* or povidone-iodine or betadine or chlorhexidine or triclosan or hexachlorophene or benzalkonium or alcohol or alcohols or antiseptic* or soap*).tw.
24 exp Disinfectants/
25 13 and 24
26 (skin adj5 disinfect*).tw.
27 exp Detergents/
28 13 and 27
29 (skin adj5 detergent*).tw.
30 or/15-23,25-26,28-29
31 exp Handwashing/
32 exp Hand/
33 (hand or hands or handwash* or surgical scrub*).tw.
34 or/31-33
35 12 and 30 and 34

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Appendix 4. EBSCO CINAHL search strategy

S32 S12 and S25 and S31
S31 S26 or S27 or S28 or S29 or S30
S30 TI (surgical scrub*) or AB (surgical scrub*)
S29 TI (hand or hands or handwash*) or AB (hand or hands or handwash*)
S28 (MH "Surgical Scrubbing")
S27 (MH "Hand+")
S26 (MH "Handwashing+")
S25 S13 or S14 or S15 or S16 or S17 or S18 or S19 or S20 or S23 or S24
S24 TI skin N5 disinfect* or AB skin N5 disinfect*
S23 S21 and S22
S22 (MH "Skin+")
S21 (MH "Disinfectants")
S20 TI (iodophor* or povidone-iodine or betadine or chlorhexidine or triclosan or hexachlorophene or benzalkonium or alcohol or alcohols or antiseptic* or soap* or detergent*) or AB (iodophor* or povidone-iodine or betadine or chlorhexidine or triclosan or hexachlorophene or benzalkonium or alcohol or alcohols or antiseptic* or soap* or detergent*)
S19 (MH "Detergents+")
S18 (MH "Soaps")
S17 (MH "Alcohols+")
S16 (MH "Chlorhexidine")
S15 (MH "Povidone-Iodine")
S14 (MH "Iodine")
S13 TI antisepsis or AB antisepsis
S12 S1 or S2 or S3 or S4 or S5 or S6 or S7 or S8 or S9 or S10 or S11
S11 TI (preoperative care or pre-operative care) or AB (preoperative care or pre-operative care)
S10 (MH "Perioperative Care+")
S9 (MH "Preoperative Care+")
S8 TI (postoperative* N5 wound infection* OR post-operative* N5 wound infection* OR post-operative* N5 wound infection*) or AB (postoperative* N5 wound infection* OR post-operative* N5 wound infection*)
S7 TI surg* N5 dehiscen* or AB surg* N5 dehiscen*
S6 TI surg* N5 incision* or AB surg* N5 incision*
S5 TI surg* N5 site* or AB surg* N5 site*
S4 TI surg* N5 wound* or AB surg* N5 wound*
S3 TI surg* N5 infection* or AB surg* N5 infection*
S2 (MH "Surgical Wound Dehiscence")
S1 (MH "Surgical Wound Infection")

Appendix 5. Risk of Bias assessment

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 of 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 available to permit a 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 were 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 unsealed procedure.

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
Either of the following.
- Insufficient information available to permit a 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**

Either 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**

Either 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 available 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.

**FEEDBACK**

Enquiry about status of any ongoing trials, 11 August 2008

**Summary**

The Authors conclusions’ include suggestions for trials that are needed. Ecolab Ltd would like to know if any of the trials have or are being carried out?
Submitter has modified conflict of interest statement: I certify that I have affiliations with an organisation or entity with a financial interest in the subject matter of my feedback.

Reply
Tanner conducted a randomised controlled trial in April to June 2008 comparing nail brushes and nail picks with 164 operating room staff. This study found no difference in the number of colony forming units on the hands of the scrub staff one hour after they had scrubbed with antiseptic solution and a nail pick, antiseptic solution and a nail brush or antiseptic solution alone. The authors of this review have not been informed of other studies in this field.

New searches have been conducted for this review which is currently being updated.

Contributors
Author of feedback: Alex Haworth Occupation Business Development Manager, Ecolab Ltd.

Review author: Judith Tanner

WHAT'S NEW

| Date              | Event                                           | Description                                                                 |
|-------------------|-------------------------------------------------|------------------------------------------------------------------------------|
| 11 January 2016   | New citation required but conclusions have not changed | First update. New search. Four new trials. Conclusions unchanged.             |
| 30 June 2015      | New search has been performed                   | Risk of bias updated, Grade assessment undertaken. Revision of Results to streamline text and introduction of new supporting table to replace text. |

HISTORY
Protocol first published: Issue 3, 2003
Review first published: Issue 1, 2008

| Date              | Event                                           | Description                                                                 |
|-------------------|-------------------------------------------------|------------------------------------------------------------------------------|
| 26 August 2008    | Feedback has been incorporated                   | Response to enquiry regarding the status of ongoing trials.                  |
| 8 August 2008     | Amended                                         | Converted to new review format.                                              |
| 4 October 2007    | New citation required and conclusions have changed | Substantive amendment                                                       |

CONTRIBUTIONS OF AUTHORS

Judith Tanner: conceived and designed the review; performed part of the data analysis and interpretation, writing and editing; made an intellectual contribution to and advised on the review; approved the final version before submission; performed previous work that was the foundation for the current review and wrote to study authors/experts/companies.

Jo Dumville: extracted data; analysed and interpreted data and checked quality assessment; performed statistical analysis; completed the first draft of the review, approved the final version prior to submission and is the guarantor of the review.

Gill Norman: checked quality of data extraction and analysed and interpreted data; checked quality assessment; checked quality of statistical analysis; performed part of writing and editing the review; and approved the final version before submission.

Matthew Fortnam: undertook quality assessment and approved the final version before submission.
Contributions of the editorial base

For this update:

Joan Webster and Nicky Cullum (Editors) edited the protocol, advised on methodology, interpretation and content; approved the final review update prior to submission.

Sally Bell-Syer: co-ordinated the editorial process; advised on content; Gill Rizzello: edited the review update.

Rocio Rodriguez: designed the search strategy, ran the searches. Reetu Child edited the search methods section.

DECLARATION OF INTEREST

Judith Tanner:

Judith Tanner gave a lecture at two study days in 2013 for which her employer was paid an honorarium from Molnlycke HealthCare. Judith Tanner had no control of the funds. Molnlycke Healthcare produce a surgical hand antiseptic solution.

Judith Tanner gave a lecture at a study day in 2015 for which she was paid an honorarium from Molnlycke HealthCare. This money was later forwarded to her employer and Judith Tanner has no control of the funds. Molnlycke Healthcare produce a surgical hand antiseptic solution.

Judith Tanner gave a lecture at a study day in 2015 for which she was paid an honorarium from Smith and Nephew. Smith and Nephew do not make any products relevant to this review.

Jo Dumville: none known.

Gill Norman: my employment at the University of Manchester is funded by NIHR Cochrane programme Grant 13/89/08- High Priority Cochrane Reviews in Wound Prevention and Treatment.

Matthew Fortnam: none known.

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• School of Nursing, Midwifery and Social Work, University of Manchester, UK.

External sources

• National Institute for Health Research (NIHR), UK, UK.

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DIFFERENCES BETWEEN PROTOCOL AND REVIEW

In this update we removed rates of septicaemia from the outcomes.

INDEX TERMS

Medical Subject Headings (MeSH)

*General Surgery; Anti-Infective Agents, Local [*administration & dosage]; Antisepsis [*methods]; Colony Count, Microbial; Hand [*microbiology]; Hand Disinfection [*methods]; Randomized Controlled Trials as Topic; Surgical Wound Infection [epidemiology] [*prevention & control]

MeSH check words

Humans