ABSTRACT

Objective: To determine whether there still remains a significant advantage in the use of sutures to staples for orthopaedic skin closure in adult patients.

Design: Systematic Review/ Meta-Analysis.

Data sources: MEDLINE-OVID, EMBASE-OVID, CINAHL and Cochrane Library. Grey and unpublished literature was also explored by searching: International Clinical Trial Registry, Grey Matters BIOSIS Previews, Networked Digital Library of Theses and Dissertations, ClinicalTrials.gov, UK Clinical Trials Gateway, UK Clinical Research Network Study Portfolio, Open Grey, Grey Literature Report, and Web of Science.

Selection criteria: Articles were from any country, written in English and published after 1950. We included all randomised control trials and observational studies comparing adults (≥18 years) undergoing orthopaedic surgery who either received staples or sutures for skin closure. The primary outcome was the incidence of surgical site infection. Secondary outcomes included closure time, inflammation, length of stay, pain, abscess formation, necrosis, discharge, wound dehiscence, allergic reaction and health-related quality of life.

Results: 13 studies were included in our cumulative meta-analysis conducted using Review Manager V.5.0. The risk ratio was computed as a measure of the treatment effect taking into account heterogeneity. Random-effect models were applied. There was no significant difference in infection comparing sutures to staples. The cumulative relative risk was 1.06 (0.46 to 2.44). In addition, there was no difference in infection comparing sutures to staples in hip and knee surgery, respectively. Lastly, except for closure time, there was no significant difference in secondary outcomes comparing sutures to staples.

Conclusions: Except for closure time, there was no significant difference in superficial infection and secondary outcomes comparing sutures to staples was found. Given that there may in fact be no difference in effect between the two skin closure and the methodological limitations of included studies, authors should begin to consider the economic and logistic implications of using staples or sutures for skin closure.

PROSPERO registration number: CRD42015017481.

INTRODUCTION

In the context of orthopaedic surgery, surgical site infection (SSI)—defined as the occurrence of wound infection following surgery—are frequent postoperative complications that represent 20% of all nosocomial infections. SSIs are clinically classified as the occurrence of infection affecting either the superficial or deep incision sites within 30 days postoperatively or within 1 year if an implant is left inside the patient.1 Orthopaedic SSIs are frequent postoperative complications that represent 20% of all nosocomial infections. In addition, orthopaedic SSIs have been shown to extend postoperative hospital stay, double hospital readmission rates and increase annual healthcare costs up to 300%.1–4 Unfortunately, orthopaedic SSIs also inflict a tremendous burden on the patient, often increasing physical limitations and reducing postoperative quality of life.4,5

In response to the critical need to reduce the incidence of SSIs, a wealth of research has been published identifying effective preoperative, intraoperative and postoperative strategies to reduce the incidence of SSIs.1 Preoperative strategies include increasing host immune status, prophylactic antibiotic administration and proper sterilisation of the patient and all surgical staff.6 During surgery, it is imperative that the surgical environment is properly ventilated, all surgical equipment are sterilised, and the use of proper surgical technique is employed.5 Postoperatively,
incisions should be properly wrapped with sterile dressing and proper sterile techniques should be used during dressing changeover.5

With the increased pressure on orthopaedic surgeons to promote rapid wound healing, reduce postoperative length of stay (LOS) and complications, the method of skin closure during surgery has become increasingly important.7–9 The optimal goal of skin closure, ‘is to promote rapid skin healing and an acceptable cosmetic result while minimising the risk of dehiscence or infection’.7 In orthopaedic surgery, the most common skin closure methods are the use of staples or sutures.2 10 11 Yet, there seems to be no consensus in the literature as to which closure method is superior, with some studies reporting no difference and others reporting a higher wound complication rate following the use of staples.9 12–15

A systematic review and meta-analysis (SRMA), published in 2009, compared the efficacy of sutures to staples after orthopaedic surgery.7 They reported that the risk for postoperative infection was over three times greater in patients who receive staples for skin closure compared with those who receive sutures.7 Yet, due to small sample sizes, noticeable heterogeneity in effect size across outcomes, and lack of methodological rigour within studies, called for additional research in this area.7–10 12 15–17

Consequently, in response to the paucity of information and continuing discrepancy of results, numerous observational studies and RCTs have been published within the past 6 years comparing methods of skin closure after orthopaedic surgery. Therefore, the primary aim of our study is to perform an updated SRMA, and to investigate whether there still remains an advantage in the use of sutures to staples for orthopaedic skin closure.

METHODS
Search strategy
We searched MEDLINE-ovid, EMBASE-ovid, CINAHL and Cochrane Library from 1950 to January 2015 (see online supplementary appendix A for detailed search strategy pertaining to each databases). Grey and unpublished literature was also explored by searching: International Clinical Trial Registry, Grey Matters, BIOSIS Previews, Networked Digital Library of Theses and Dissertations, ClinicalTrials.gov, UK Clinical Trials Gateway, UK Clinical Research Network Study Portfolio, Open Grey, Grey Literature Report, and Web of Science. The following databases were searched within the Web of Science platform: Web of Science Core Collection, BIOSIS Previews, KCI-Korean Journal Database, MEDLINE, SciELO Citation Index, and Zoological Record. Conference Proceedings were indexed within the Web of Science Core Collection database search. No language or date of coverage restrictions was placed on all searches. Lastly, on completion of levels 1, 2 and 3 screening, forward and backward citation searches were done to assure no studies were mistakenly omitted from our search strategy. A review protocol was submitted and approved with PROSPERO on 10 March 2015, and approved on 12 March 2015 (registration number: CRD42015017481). The search strategy was designed to accommodate for database and platform-specific terminology, and syntax. A detailed search strategy including MeSH terms, keywords and Boolean operators can be found within the online supplementary appendix.

Population, Intervention, Comparator, Outcomes and Study Design
The Population, Intervention, Comparator, Outcomes and Study Design (PICOS) were adopted from Smith et al.7 Articles were from any country, written in English and published after 1950. We included all randomised control trials and observational studies (not an editorial, opinion, case report or a review article) comparing adults (≥18 years) undergoing orthopaedic surgery (trauma or elective) who either received staples or sutures (excluding skin adhesives, barbed sutures and surgical zipper) for skin closure. The primary outcome was the incidence of SSI after orthopaedic surgery. Secondary outcomes included closure time, inflammation, LOS, patient satisfaction, pain, abscess formation, necrosis, discharge, wound dehiscence, allergic reaction and health-related quality of life (HRQoL). This study has been reported according to the PRISMA guidelines.18

Study selection
A total of 947 studies were identified and imported into EPPI V.4.0 Reference Manager. As per the prespecified search strategy, 280 studies were retrieved through relevant database searches and an additional 667 studies retrieved from grey literature searches and various clinical trial registries. After removing 216 duplicates, 731 articles underwent title screening. Studies were excluded if they were not human (n=4), not a research study (n=11), and if they did not look at orthopaedic surgery (n=84) or skin closure (n=511). On completion of level 1 screening, 610 studies were excluded and 121 were included for level 2 (abstract) screening. Studies were excluded if they did not directly compare sutures to staples for skin closure (n=102). Nineteen studies were eligible for full-text screening. Articles were excluded if: they were not English (n=1), the results were not published (n=1), the studies had not yet completed (n=2; International Clinical Trial Registry: (1) NCT02040312001 and (2) NTR3946) and if one could not differentiate results from orthopaedic and non-orthopaedic groups (n=1). Contact with authors were attempted to obtain English manuscripts, and any unpublished results or interim data. Two independent reviewers (RK and SDM) assessed articles for eligibility; κ statistics were calculated to assess reviewer agreement on the completion of levels 1 and 2 screening.
Additionally, all disagreements were corrected through negotiated consensus. The agreement between reviewers for levels 1 and 2 screening was: 0.50 and 0.85, respectively. A detailed PRISMA flow chart outlining the number of studies screened for eligibility, and reasons for exclusion are provided in figure 1. Specific screening questions are provided in the online supplementary appendix.

**Data extraction**
Relevant quantitative and qualitative data necessary for adequate study comparison and analysis was abstracted by a reviewer (RK). Information on study objective, design, operation type, closure material, sample size, age (mean), sex ratio, time to removal, follow-up, infection, closure time, inflammation, LOS, patient satisfaction, pain, abscess formation, necrosis, discharge, wound dehiscence, allergic reaction and HRQoL was collected. All outcomes were prespecified within PROSPERO.

**Risk of bias assessment**
Studies were critically appraised for adequate randomisation, allocation concealment, baseline similarity, blinding, follow-up completion and intention-to-treat (ITT) analysis. Subsequently, using the Grading of Recommendations, Assessment, Development, and Evaluation (GRADE) guidelines, included articles were used to assign an overall grade of either high, moderate, low quality of evidence for infection.19–22

**Summary measures and statistical analysis**
Statistical analysis was conducted using Review Manager V.5.0.23 Dichotomous outcomes were reported as risk ratios and continuous outcomes as mean differences (MDs). All analyses were conducted using the Mantel-Haenszel random-effects model. Heterogeneity was assessed using the $\chi^2$ and $I^2$. All results were reported at a significance level of 0.05, and 95% CI was provided. Stratification of infection based on surgery type has been prespecified, and a funnel plot was calculated to assess publication bias. In addition, ITT analysis was performed in studies in which results were reported ‘as per protocol’.

**RESULTS**

**Study selection**
Thirteen studies were included in our cumulative meta-analysis (table 1).8–10 12 13–17 24–29 All articles were published between 1990 and 2015. The cumulative sample size of all studies was 563 patients within the suture group, and 692 patients within the staple group.

Ten studies were randomised controlled trials and three were observational studies.

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**Figure 1** PRISMA flow diagram.33

- Records identified through database searching (n = 280)
- Additional records identified through other sources (n = 667)
- Records after duplicates removed (n = 216)
- Records screened (n = 731)
- Records excluded (n = 712)
- Full-text articles assessed for eligibility (n = 19)
- Studies included in qualitative synthesis (n = 13)
- Studies included in quantitative synthesis (meta-analysis) (n = 13)

<table>
<thead>
<tr>
<th>Full-text articles excluded, with reasons (n = 6)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Not English: 1</td>
</tr>
<tr>
<td>2. Could not differentiate between orthopaedic and non-orthopaedic patients: 2</td>
</tr>
<tr>
<td>3. Study completed but published results not found: 1</td>
</tr>
<tr>
<td>4. Study had not been completed and interim data not available: 2</td>
</tr>
<tr>
<td>Study</td>
</tr>
<tr>
<td>----------------------------</td>
</tr>
<tr>
<td>Stockley and Elson</td>
</tr>
<tr>
<td>Clay and Southwood</td>
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<tr>
<td>Liew and Haw</td>
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<tr>
<td>Graham et al</td>
</tr>
<tr>
<td>Murphy et al</td>
</tr>
<tr>
<td>Shetty et al (calculations used intention to treat)</td>
</tr>
<tr>
<td>Khan et al</td>
</tr>
<tr>
<td>Singh et al</td>
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<tr>
<td>Eggers et al</td>
</tr>
<tr>
<td>Newman et al</td>
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<tr>
<td>Patel et al</td>
</tr>
<tr>
<td>Slade Shantz et al</td>
</tr>
<tr>
<td>Wyles et al</td>
</tr>
</tbody>
</table>

Baseline characteristics of all included studies. All missing information are omitted in black.

ITT: sample size when considering intention to treat.

ORIF, fraction open reduction, internal fixation; PC, prospective cohort; RC, retrospective cohort; RCT, randomised controlled trial; THR, total hip arthroplasty; TKR, total knee arthroplasty.
Study characteristics

Seven studies provided data on the mean age of patients within each group. Using sample size weights, the overall mean age among the staple and suture groups were 68 and 66 years, respectively. Six studies provided data on the distribution of sex within each group. Among the 216 patients who received staples, there were 66 males and 150 females. Additionally, among the 214 patients within the suture group, there were 68 males and 146 females. The majority of studies focused on lower extremity orthopaedic operations. Surgeries ranged from total knee arthroplasty; total hip arthroplasty; hip fracture surgery; knee fracture surgery; and miscellaneous ankle, tibia, patella, olecranon and humerus fracture corrections. Of the seven articles that included more than one surgery type, only three studies stratified the results based on procedure. Six studies did not report suture and staple removal time. In the 12 papers that provided the relevant data, the overall time to removal was consistent across all studies and ranged from 10 to 16 days. One study did not report suture and staple follow-up time. Among the rest, there was considerable variation in postoperative follow-up, ranging from 3 to 365 days. There was some variation in suture material. The most common materials used were Nylon and Vicryl. A summary of all baseline characteristics abstracted from included studies can be found in Table 1.

Risk of bias assessment

A risk of bias assessment of all 13 studies showed considerable variation in study quality (Table 2). Two studies were retrospective chart reviews, 1 study was a prospective cohort and 10 studies were randomised controlled trials. Of the 10 RCTs, 2 study did not appropriately randomise patients, 5 studies did not perform or clearly report allocation concealment, and 5 studies did not provide baseline characteristics. In addition, seven studies did not blind patients and six studies did not perform or did not clearly report ITT analysis. However, among all included articles, only six did not complete or adequately report follow-up. Despite the low quality, all studies were included in the analysis. Lastly, with respect to our primary outcome of infection, an overall grade of ‘very low quality’ was assigned due to the considerable...
amount of imprecision, inconsistency and indirectness among the evidence (table 3).

**Primary outcome: infection**
All studies provided information on infection among patients receiving sutures and staples for skin closure (table 4). Wound infection comparing sutures to staples was insignificant, with a cumulative relative risk (RR) of 1.06 (95% CI 0.45 to 2.44; figure 2).

**Secondary outcomes**
1. **Dehiscence**: Seven studies provided data on wound dehiscence. The cumulative risk ratio was insignificant, with a RR of 0.96 (0.32 to 2.84; table 4 and figure 3).
2. **Inflammation, discharge, necrosis, allergic reaction, abscess formation**: There was no significant difference in effect comparing sutures to staples across all of these outcomes (table 4).
3. **Pain**: Two studies measured pain. Slade Shantz et al. assessed pain using the 100 mm pain scale, and Stockley and Elson noted 31 cases of reported pain among removal of staples as compared to seven cases of reported pain within the suture group. Although no inferential statistics was used, it seems that staples may be more painful to remove than sutures.
4. **Closure time**: Four studies provided information on closure time. However, only within two studies was mean closure time provided or calculable. The cumulative MD was significant, with a MD of 5.84 (4.52 to 7.15) comparing suture and staple groups. Qualitatively, Liew and Haw noted that wound closure was approximately 7 times faster with staples compared with sutures. No inferential statistics was used. Lastly, Khan et al. noted that closure time was faster using skin clips compared with sutures. Data were reported as the median and

### Table 3 Grade assessment for overall quality of evidence with respect to infection

<table>
<thead>
<tr>
<th>Number of participants</th>
<th>Risk of bias</th>
<th>Inconsistency</th>
<th>Indirectness</th>
<th>Imprecision</th>
<th>Publication bias</th>
<th>Overall quality of evidence</th>
</tr>
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<tbody>
<tr>
<td>(studies)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Infection (follow-up: range 365 days)</td>
<td>Serious&lt;sup&gt;1-6&lt;/sup&gt;</td>
<td>Not serious&lt;sup&gt;7&lt;/sup&gt;</td>
<td>Not serious&lt;sup&gt;8&lt;/sup&gt;</td>
<td>Serious&lt;sup&gt;9&lt;/sup&gt;</td>
<td>Publication bias strongly suspected&lt;sup&gt;10&lt;/sup&gt;</td>
<td>Very low</td>
</tr>
<tr>
<td>(10 RCTs)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>761</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Infection (follow-up: range 365 to)</td>
<td>Serious&lt;sup&gt;1&lt;/sup&gt;</td>
<td>Serious&lt;sup&gt;7&lt;/sup&gt;</td>
<td>Not serious&lt;sup&gt;8&lt;/sup&gt;</td>
<td>Serious&lt;sup&gt;9&lt;/sup&gt;</td>
<td>Publication bias strongly suspected&lt;sup&gt;10&lt;/sup&gt;</td>
<td>Very low</td>
</tr>
<tr>
<td>(3 observational studies)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>484</td>
<td></td>
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</table>

GRADE evaluation for overall quality of evidence. Included studies are separated into clinical trials and observational studies. Overall, included studies are of very low quality. Explanations for risk of bias, inconsistency, indirectness, imprecision and publication bias are provided below.

1. Singh et al., Newman et al., Patel et al. were observational studies (no randomisation, allocation concealment, blinding).
2. Clayer and Southwood and Stockley et al. did not truly randomise (randomised based on unit number and date of birth).
3. Of the randomised trials, six studies did not report or conceal allocation.
4. Of the randomised trials, seven studies did not blind (patient or assessor) to the treatment (staples or sutures).
5. For seven studies, completion of follow-up for all patients was not clearly reported or did not occur.
6. For two studies infection was not the primary outcome.
7. I<sup>2</sup> value <75%.
8. All studies included in the meta-analysis directly compare sutures to staples.
9. Very few events occurred in suture and staple treatment arms resulting in very large confidence intervals.
10. One study did not publish results and was excluded from our systematic review/meta-analysis.

GRADE, Grading of Recommendations, Assessment, Development, and Evaluation; RCT, randomised controlled trial.

### Table 4 Primary and secondary outcomes comparing suture to staples for skin closure after orthopaedic surgery

<table>
<thead>
<tr>
<th>Outcome</th>
<th>Incidence</th>
<th>Relative risk (95% CI)</th>
<th>p Value</th>
<th>Heterogeneity</th>
<th>Mean difference (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Sutures</td>
<td>Staples</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Infection&lt;sup&gt;8-10, 12, 15-17, 24-29&lt;/sup&gt;</td>
<td>171563</td>
<td>21692</td>
<td>1.06 (0.46 to 2.44)</td>
<td>0.89</td>
<td>18</td>
</tr>
<tr>
<td>Dehiscence&lt;sup&gt;8, 10, 12, 16, 26, 28, 29&lt;/sup&gt;</td>
<td>6308</td>
<td>21441</td>
<td>0.96 (0.32 to 2.84)</td>
<td>0.94</td>
<td>0</td>
</tr>
<tr>
<td>Inflammation&lt;sup&gt;8, 16&lt;/sup&gt;</td>
<td>359</td>
<td>22172</td>
<td>0.22 (0.00 to 12.07)</td>
<td>0.46</td>
<td>84</td>
</tr>
<tr>
<td>Discharge&lt;sup&gt;8, 9, 16&lt;/sup&gt;</td>
<td>8123</td>
<td>17135</td>
<td>0.66 (0.14 to 3.23)</td>
<td>0.61</td>
<td>58</td>
</tr>
<tr>
<td>Necrosis&lt;sup&gt;15, 16&lt;/sup&gt;</td>
<td>1192</td>
<td>3197</td>
<td>0.51 (0.07 to 3.88)</td>
<td>0.52</td>
<td>0</td>
</tr>
<tr>
<td>Allergic reaction&lt;sup&gt;9, 15, 26&lt;/sup&gt;</td>
<td>21226</td>
<td>1211</td>
<td>1.37 (0.22, 8.60)</td>
<td>0.74</td>
<td>0</td>
</tr>
<tr>
<td>Abscess&lt;sup&gt;15, 29&lt;/sup&gt;</td>
<td>21115</td>
<td>2244</td>
<td>1.86 (0.22 to 15.71)</td>
<td>0.57</td>
<td>0</td>
</tr>
<tr>
<td><strong>Closure time</strong>&lt;sup&gt;17, 28&lt;/sup&gt; (min)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>5.84 (4.52 to 7.15)</td>
</tr>
</tbody>
</table>

Primary and secondary outcomes comparing sutures to staples for skin closure after orthopaedic surgery. All outcomes (95% CI) are calculated using the Mental-Haeszel random-effects model. Heterogeneity is calculated using the I<sup>2</sup> statistic. χ<sup>2</sup> significance level dictates whether the heterogeneity between studies is significant (*p<0.05).
IQR and thus, could not be used for the cumulative
effect estimate.

5. Length of stay: Two studies reported LOS.9  8 However,
Eggers et al28 9 and Khan et al28 9 reported LOS as mean
(CI) and median (IQR) days, respectively. Normality
was not assumed, and the data was not combined.
Qualitatively, there is no noticeable difference among
patients who receive staples or sutures.

6. HRQoL: Only one study administered the Short
Form-36, a generic HRQoL questionnaire within
suture and staple groups.28 There was no difference
in HRQoL across all subscales.

Sensitivity analysis
Three randomised controlled trials reported results as
per protocol. We adjusted the results according to ITT
and performed a sensitivity analysis to examine the
overall RR of infection. Accounting for lost to follow-up,
the RR of infection remained insignificant (table 5,
figures 4 and 5).

Subgroup analysis
1. Hip surgery: Six studies provided data comparing infec-
tion between patients receiving sutures to staples after
hip surgery.8  10  12  15  29 The cumulative sample size
was 164 and 245 patients in the suture and staple
group, respectively. The cumulative risk ratio was insig-
ificant, with a RR of 0.48 (0.10 to 2.45; figure 6).

2. Knee surgery: Seven studies provided data comparing infec-
tion between patients receiving sutures to staples
after knee surgery.9  15  25–29 The cumulative sample
size was 239 and 263 patients in the suture and staple
group, respectively. The cumulative risk ratio was insign-
ificant, with a RR of 1.38 (0.42 to 4.52; figure 7).

Publication bias
Publication bias was assessed using a funnel plot for our
primary outcome of infection (figure 8). The funnel for
studies was not fully symmetrical. Included studies were
scattered throughout the bottom and top right corner of
the plot. Yet, there seems to be some evidence for
missing studies as seen by the empty space in the top left
corner. Nevertheless, publication bias can be suspected
but cannot be concluded based on this plot. The reason
for difficulty in interpretation is mainly due to the small
group of studies and high heterogeneity and small effect
size. In addition, publication bias can be one of many
possible explanations for our funnel plot asymmetry.

DISCUSSION
As compared with the systematic review and meta-analysis
published in 2009,7 we found no significant difference in
infection among patients who receive staples and sutures
for skin closure. Although there was no significant

Figure 2  Forest plot of the cumulative risk ratio (95% CI) of wound infection comparing patients who received staples or sutures
for wound closure after orthopaedic surgery. Analysis conducted with Mantel-Haenszel random-effects model.

Figure 3  Forest plot of the cumulative risk ratio (95% CI) of wound dehiscence comparing patients who received staples or sutures
for wound closure after orthopaedic surgery. Analysis conducted with Mantel-Haenszel random-effects model.
heterogeneity across all studies (p > 0.05), there was some noticeable variation in magnitude and direction of effect (figure 1). In addition, our study also showed no significant difference in dehiscence, inflammation, discharge, necrosis, allergic reaction, and abscess formation. Three studies did not report results as per ITT. Sensitivity analysis was conducted following ITT analysis. Two scenarios were created accounting for two logical extreme cases. Scenario 1 assumes all lost to follow-up within the suture group had infection, and patients who received staples developed no infection. Conversely, scenario 2 assumes all patients who were lost to follow-up within the staple cohort received infection and no patient experienced infection in the suture group. The risk of infection comparing sutures to staples within scenarios 1 and 2 was insignificant, further strengthening our conclusion of no significant difference in infection risk between sutures and staples. However, it is important to note that the magnitude of effect was large within both scenarios (figures 4 and 5; table 5), suggesting that our sensitivity analysis may have been underpowered to detect a significant difference.

Subgroup analysis was also performed comparing the effect of sutures and staples on infection, stratified by surgery type. Limitations on available data allowed us to only analyse the effect of infection comparing hip and knee surgery only. There was no significant difference in the incidence of infection across both surgery types. However, the use of staples was potentially harmful, with a RR of 1.38 (0.42 to 4.52). However, caution must be given to the interpretation of results in this particular group due to the significant heterogeneity in magnitude and direction of effect between studies. With respect to patient-centred outcomes, only two studies measured patient-reported pain. Unfortunately, the results from both studies could not be combined quantitatively. However, based on qualitative analysis, there seems to be a trend that staples cause more pain on removal than sutures. This trend is consistent with non-orthopaedic surgeries.50–52

Table 5 Sensitivity analysis

<table>
<thead>
<tr>
<th>Study</th>
<th>Initially randomised sample size</th>
<th>Lost to follow-up</th>
<th>Deaths</th>
<th>Per protocol sample size</th>
<th>Per protocol Relative risk</th>
<th>Favours sutures (worst case staples)</th>
<th>Favours staples (worst case sutures)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Slade Shantz et al17</td>
<td>97</td>
<td>28</td>
<td>0</td>
<td>69</td>
<td>1.14 (0.07 to 17.96)</td>
<td>0.06 (0.01 to 0.47)</td>
<td>27.80 (3.87 to 200)</td>
</tr>
<tr>
<td>Liew and Haw24</td>
<td>23</td>
<td>4</td>
<td>1</td>
<td>19</td>
<td>2.85 (0.12 to 65.74)</td>
<td>0.25 (0.03 to 2.07)</td>
<td>5.0 (0.25 to 98.75)</td>
</tr>
<tr>
<td>Shetty et al10</td>
<td>55</td>
<td>8</td>
<td>1</td>
<td>47</td>
<td>0.10 (0.01 to 1.84)</td>
<td>0.09 (0.01 to 1.61)</td>
<td>(no loss to follow-up)</td>
</tr>
</tbody>
</table>

Sensitivity analysis for the relative risk of infection. Adjusted studies were those who did not follow ITT protocol and had lost to follow-up. Adjusted relative risks were calculated for two scenarios. Scenario 1: all lost to follow-up had infection in suture group, and none in staple group. Scenario 2: all lost to follow-up had infection in staple group, and none in the suture group. Original randomised treatment sizes were used for adjusted relative risk calculations. ITT, intention to treat.
comparing sutures to staples was 5.84 (4.52 to 7.15). The results do suggest that staples may be slightly faster to use compared with sutures for skin closure. Slade Shantz et al.17 reported an estimated 7 min time savings with the use of staples for skin closure. If an average orthopaedic surgeon were to perform five surgeries per day, the use of staples could save an estimated 35 min of operative time potentially reducing overscheduling and hospital costs.17 28 Liew and Haw24 also noted that the use of staples for skin closure was approximately 7 times faster as compared with sutures. However, the potential utility of staples for skin closure may still be a false economy driven by surgeon perception and bias.7 The use of staples has always been noted as a ‘more expensive method’ for skin closure as compared with using sutures.15 16 In addition, the number dressing changes required for patients who undergo skin stapling is much higher than those who undergo suturing.7 Consequently, the savings in intraoperative costs could be negated by dressing changes and staple/applicator costs. To our knowledge, no formal cost-benefit analysis has been done comparing the use of sutures and staples for orthopaedic surgery. Other economic considerations include difference in LOS comparing sutures to staples. A shorter LOS can result in shorter postoperative costs, inpatient care, physical therapy and pharmaceuticals.28 Additionally, Eggers et al.28 noted that the level of wound cosmesis might also affect LOS, such that a shorter LOS may be partly due to a higher emotional satisfaction predicated on a more cosmetically appealing incision.

In our study, there seems to be no noticeable difference in LOS comparing sutures to staples. However, it is important to note that only two studies analysed LOS, and our conclusions are solely speculative. A major limitation of our SRMA is the substantially low quality of evidence and considerable heterogeneity across studies. Of the 13 studies included, 7 studies did not provide baseline characteristics.10 15 16 24–26 29 Thus, we could not conclusively establish that patients between groups were similar across all confounders for infection such as age, body mass index and comorbidity burden.7 Another potential limitation is that we included two studies in which comparing wound complications...
between sutures and staples was not the primary objective. One study which was published in the year 2000 was captured by the previous systematic review/meta-analysis. The authors noted that the study was excluded because comparing the incidence of wound complications was not the primary objective. However, for the purposes of our SRMA, the two studies were included because they effectively randomised patients to suture and staple treatments. In addition, information on postoperative infection was reported within both studies.

In addition, no information was available evaluating specific suture techniques, or surgeon skill. Murphy et al. suggested that poor surgical technique and accuracy of suture or staple closure can have an effect on wound healing. However, it is unclear whether suturing technique is confounder with respect to post-operative wound complications. Second, the majority of studies did not specify whether infections were superficial or deep. Superficial infections typically require the administration of antibiotics while deep infections require more severe courses of treatment.

The strengths of our study are twofold. First, our SRMA was completed with high methodological rigour. All of our outcomes and subgroup analysis were prespecified and our study was registered with PROSPERO prior to data extraction. Second, in contrast to Smith et al., we found no significant difference comparing sutures to staples across all post-operative complications and subgroups; suggesting that there may be no clinically relevant difference in effect between the two skin closure methods within this population.

Future studies should focus on conducting high-quality randomised control trials. Detailed baseline characteristics, and a detailed patient recruitment flow chart should be presented. In addition, authors should remove themselves from ‘as per protocol’ analysis and convey results according to the ITT principle. Second, our study failed to capture two large randomised controlled trials because they had not yet been completed. Thus, future SRMAs should be conducted comparing the use of sutures to staples for skin closure.

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**REFERENCES**