Prevention of surgical site infection in lower limb skin lesion excisions with single dose oral antibiotic prophylaxis: a prospective randomised placebo-controlled double-blind trial

Samuel C Smith,1 Clare F Heal,2 Petra G Buttner3

ABSTRACT
Objectives: To determine the effectiveness of a single perioperative prophylactic 2 g dose of cephalexin in preventing surgical site infection (SSI) following excision of skin lesions from the lower limb.

Design: Prospective double-blinded placebo-controlled trial testing for difference in infection rates.

Setting: Primary care in regional North Queensland, Australia.

Participants: 52 patients undergoing lower limb skin lesion excision.

Interventions: 2 g dose of cephalexin 30–60 min before excision.

Main outcome measures: Incidence of SSI.

Results: Incidence of SSI was 12.5% (95% CI 2.7% to 32.4%) in the cephalexin group compared with 35.7% (95% CI 18.6% to 55.9%) in the placebo group (p=0.064). This represented an absolute reduction of 23.21% (95% CI –0.39% to 46.82%), relative reduction of 65.00% (95% CI –12.70% to 89.13%) and number-needed-to-treat of 4.3.

Conclusions: Administration of a single 2 g dose of cephalexin 30–60 min before skin lesion excision from the lower limb may produce a reduction in the incidence of infection; however, this study was underpowered to statistically determine this.

Trial registration number: ACTRN12611000595910.

INTRODUCTION

Skin cancer causes a significant burden of disease in many developed countries.1, 2 The majority of skin cancers are treated by surgical excision3, 4 which is increasingly being performed in outpatient and primary care settings.5, 6 As the majority of skin cancer surgery takes place in general practice in Australia,5 it is important to study infection rates in this setting. Skin lesion excisions form a large proportion of a typical Australian general practitioner’s (GPs) workload, and this proportion is even greater for Queensland GPs, given that this State has the highest incidence of skin cancer.5 General practice dermatological surgery may differ from a hospital setting, with most procedures taking place in treatment rooms rather than in formal operating theatres.

Surgical site infection (SSI) is one of the few complications of this relatively minor surgery. These infections often require antibiotics and repeat consultations to assess wound healing. They can potentially lead to significant bacteraemic complications and impair cosmetic outcome.7 The acceptable rate of infection following clean minor surgery (class 1) is less than 5%.8–12 This is reflected in skin lesion excisions, with a rate of between 1% and 3% in most studies.13–18 The exceptions are studies conducted by the present authors in 2005 and 2009 which reported infection rates of 8.6% and 11.7%.19 20 The reason for this higher infection rate remains unclear, but might be related to tropical humidity. Even within cohorts with a low overall incidence of infection, some procedures may be at higher risk because of the body site, pathology or patient factors and infection rates may be greater than 5% in these high-risk groups. Previously identified risk factors include
Excision from the region of the lower leg, ear and nose, other predetermined significant medical conditions. A body site map was used to record the excision site. At the end of the study, practice nurses were asked to re-examine computer records to fill in any missing data. The principal researcher visited participating GPs and practice nurses prior to the start of and regularly during the trial to ensure that data collection was standardised.

**Intervention and control**

The intervention arm of the trial consisted of a 2 g dose of cephalaxin administered 30–60 min before an excision. The cephalaxin (Keflex 500 mg capsules) was compounded into generic gel capsules (SurgiPack Empty Gelatine Capsules, Size ‘00’, 0.95 mL capacity). Rice flour was used to fill the placebo capsules as it had an identical appearance to the cephalaxin powder and no foreseeable adverse effects. Each dose consisted of four capsules filled with either cephalaxin or rice flour.

**Randomisation and blinding**

A computer-generated random number table was used to randomise a consecutive sequence of numbers into two groups—intervention and control—in permuted blocks of 20 with a 1:1 ratio. These numbers were used to label medicine jars containing the corresponding capsules. The content of the individual jar was known only to the principal researcher who had no direct involvement with participants or in data collection. All others involved in the trial were blinded to the capsule identity.

**Participant recruitment and trial protocol**

Eligible participants were identified by a practice nurse or doctor at the time of their pre-excision appointment. All patients provided signed consent before enrolling in the study. Participants were given a numbered medicine jar along with written and verbal instructions to take the four capsules 30–60 min before their excision appointment.

**Surgical wound management protocol**

Workshops were run at the participating practices to develop guidelines to ensure that excision management was standardised. The excision was performed using an aseptic technique and skin preparation with 2% chlorhexidine solution. Wounds were dressed with non-waterproof adhesive fabric dressings. Patients were provided with verbal and written instructions regarding wound care, including the need for wounds to be kept dry and covered for the first 2 days and for topical antiseptics or antibiotics not to be used. Time to removal of sutures (ROS) was 12–16 days at the treating doctor’s discretion.

**Outcome measures**

Wounds were assessed for infection by the practice nurse or doctor at the time of ROS, or sooner if patients re-presented earlier with a perceived infection. Our definition of SSI was based on the presence of any of the criteria shown in box 2. These were adapted from the

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**Box 1 Exclusion criteria for participant recruitment**

- <18 years old or not capable of providing informed consent.
- Declined to participate.
- Currently taking antibiotics or treating clinician feels they are clinically indicated for antibiotic treatment following excision.
- Repair of lacerations or lesions considered as contaminated/infected prior to surgery.
- Excision not utilising primary closure (eg, shave biopsy or curette).
- Excision of a sebaceous cyst.
- Patient unable to return for suture removal.
- Penicillin or cephalosporin allergy.
Centre for Disease Control and Prevention (CDC) definition for superficial SSI. Information was also collected on wound swabs performed, prescription of antibiotics and number of follow-up visits. Patients were asked about any adverse effects which could be attributed to the trial medication.

Sample size

Our sample size was calculated on the basis of previous studies which predicted a baseline infection rate of 15% from the lower limb. We decided that an absolute reduction in the SSI rate of 10% or more would be clinically significant. To reach this conclusion with statistical confidence (power 80%, α error 5%), it was calculated that a sample size of 282 excisions would be required.

The trial was conducted as part of an honours research project which ran over a 2-year period. The decision was made to cease data collection once the desired sample size of 282 was collected or by May 2012 to allow time for completion of the honours thesis.

Statistical methods

Data were entered and stored in a Microsoft Access database and statistical analysis and tests conducted in IBM SPSS Statistics V.20. Numeric data were summarised using median values and IQRs. Categorical variables were reported as proportions with Clopper-Pearson 95% CIs calculated for incidence of infection.

Fisher’s exact test was used to test the primary hypothesis; absolute and relative risk reductions were calculated with 95% CIs as well as the number-needed-to-treat (NNT). Comparison of other infection-related parameters, such as prescription of antibiotics and number of additional visits, was conducted with Fisher’s exact test and Mann-Whitney U tests, respectively.

No provisions were made for an interim analysis as it was felt that there was low risk of potential harm.

Ethics approval and trial registration

The trial was registered on the Australian New Zealand Clinical Trials Registry (ACTRN: 12611000595910) prior to its start.

RESULTS

Of a total of 78 excisions from the lower limb conducted during the collection period, 26 cases were excluded from the study (figure 1). Fifty-two cases were subsequently randomised and underwent skin lesion excision. Three participants forgot to take the trial medication prior to their excision, and one participant treated his wound with a topical antiseptic. Follow-up was completed in all 52 (100%) participants randomised. Six doctors across the two practices were involved, each recruiting between 3 and 18 patients.

Baseline characteristics

Comorbid conditions (excluding diabetes) were more common among those cases randomised to the cephalexin group compared with placebo (87.5% vs 67.9%) and antihypertensive use was also higher (70.8% vs 49.2%).

Box 2 Criteria for presence of a surgical site infection

- Purulent discharge.
- Erythema >1 cm from the wound margin.
- Localised swelling.
- Patient reports increased tenderness at the wound site.
- Patient reports increased heat at the wound site.

*Presence of any one or more of these criteria was deemed to represent the presence of a surgical site infection.
The median excision size was 6 mm smaller in the cephalexin group (table 1).

Outcome measures and estimates

The primary intention-to-treat analysis was conducted on all 52 cases which were randomised and in which the excision was performed (table 2). A secondary per protocol analysis was also performed that examined only cases where patients took their allocated trial medication and did not violate trial protocol (table 3).

In the intention-to-treat analysis, the incidence of SSI in the intervention group was 12.5% (3/24) compared with 35.7% (10/28) in the control group. This approached statistical significance (p=0.064), producing an absolute reduction of 23.2% (95% CI −0.39% to 46.82%), relative reduction of 65.00% (95% CI −12.70% to 89.13%) and an NNT of 4.3.

There were also clinically relevant reductions in antibiotic prescriptions and follow-up visits. These were also not of statistical significance (table 2).

In the per protocol analysis, the difference in incidence of infection was statistically significant (8.7% (2/23) vs 36% (9/25); p=0.039). The relative risk reduction for an SSI produced by prophylaxis was 75.85% (95% CI −0.30% to 94.18%), resulting in an absolute risk reduction of 27.30% (95% CI −3.50% to 51.10%) and an NNT of 3.7.

No adverse effects attributed to the trial medication were reported by participants.

DISCUSSION

In this trial, a 2 g dose of cephalexin 30–60 min before skin lesion excision from the lower limb was associated with a relative reduction in infection rate of 65% and an absolute reduction of 23.2% from a baseline infection rate of 35.7–12.5%. Based on these findings, the NNT with prophylactic cephalexin was 4.31 in order to prevent one infection following excision from the lower limb. Additionally, prophylaxis resulted in a reduction in the number of therapeutic antibiotic courses required and number of follow-up visits.
Limitations

Although these findings met our predetermined level for clinical relevance, an absolute reduction of infection rate by 10%, they were not statistically significant (p=0.064). Furthermore, a slower than predicted recruitment rate, and time constraints of the honours research project, resulted in the trial being terminated well before the calculated sample size was achieved. As such, this trial was unable to address the proposed hypothesis and no definitive conclusions can be drawn from it on the actual efficacy of antibiotic prophylaxis.

Various characteristics influence SSI and although information on as many variables as possible was recorded, it is difficult to ensure that all baseline data are comparable. Several characteristics did vary notably between the prophylaxis and placebo groups in this study. Given that the study was underpowered and did not achieve statistical significance, it was decided not to pursue adjustment for these imbalances using multiple logistic regression. Future studies, appropriately powered to address this hypothesis, will need to adjust for such baseline difference as well as consider cluster analysis based on individual GPs to limit the confounding effect on variable training, experience and outcomes among practitioners.

Even when using guidelines, the diagnosis of infection is still subjective and there may be interobserver and intraobserver variation. The 1992 CDC definition of SSI that we chose to use has limitations. However, it is the most widely implemented standard definition of wound infection and is the closest to a gold standard available. We have no evidence to support intrapractice and interpractice reproducibility of measurement and recording procedures. This is also an area which future studies may consider addressing.

Generalisability

The baseline infection rate for lower limb excisions in this trial was considerably higher than that reported in other studies, including those previously conducted in Mackay. The underlying reason for this is uncertain, but may be related to the majority of the excisions being from below the knee, a site that has previously been identified as being at an even higher risk of infection. Furthermore, the infection rate in studies conducted in Mackay has consistently been higher than that reported in other centres. This further adds to the argument that climate/environmental factors may be important determinants of SSI following skin lesion excision. Owing to the higher than expected infection rate, there

<table>
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<tr>
<th>Table 2</th>
<th>Primary and secondary outcome measures intention-to-treat analysis*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Cephalexin (n=24)</td>
</tr>
<tr>
<td>Number of infections</td>
<td>3</td>
</tr>
<tr>
<td>Infection rate (95% CI)</td>
<td>12.5 (2.7 to 32.4)</td>
</tr>
<tr>
<td>Prescription of antibiotics</td>
<td>3</td>
</tr>
<tr>
<td>Number of additional follow-up visits required</td>
<td>Median 0 [0–0]</td>
</tr>
<tr>
<td>Total (Range)</td>
<td>7 (0–3)</td>
</tr>
<tr>
<td>Additional total amount of antibiotics (excluding prophylaxis) (mg)†</td>
<td>Median 0 [0–0]</td>
</tr>
<tr>
<td>Total (Range)</td>
<td>27 000 (0–12 000)</td>
</tr>
</tbody>
</table>

*Data are median (IQR) (range) or number (%) unless otherwise stated; 95% CI for infection rate are the Clopper-Pearson interval; p values are Fisher’s exact test or the Mann-Whitney U test for categorical and continuous variables, respectively.
†Cephalexin group: cephalexin (15 000 mg), flucloxacillin (12 000 mg), placebo group: cephalexin (100 000 mg), flucloxacillin (24 000 mg); ciproflaxacin (14 000 mg), amoxicillin/clavulonic acid (8750 mg).
NA, not applicable.

<table>
<thead>
<tr>
<th>Table 3</th>
<th>Primary and secondary outcome measures per protocol analysis*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Cephalexin (n=23)</td>
</tr>
<tr>
<td>Number of infections</td>
<td>2</td>
</tr>
<tr>
<td>Infection rate (95% CI)</td>
<td>8.7 (1.1 to 28.0)</td>
</tr>
<tr>
<td>Prescription of antibiotics</td>
<td>2 (8.7%)</td>
</tr>
<tr>
<td>Number of additional follow-up visits required</td>
<td>Median 0 [0–0]</td>
</tr>
<tr>
<td>Total</td>
<td>5</td>
</tr>
</tbody>
</table>

95% CI for infection rates are the Clopper-Pearson interval.
*Data are median [IQR] or number (%) unless otherwise stated; 95% CI for infection rate are the Clopper-Pearson interval; p values are Fisher’s exact test or the Mann-Whitney U test for categorical and continuous variables, respectively.
NA, not available.
may be limitations to extrapolating these results to other centres, where the baseline infection is lower. However, similar relative reductions have been produced with prophylactic regimens, including single dose parenteral antibiotics, in settings with lower baseline infection rates (table 4).

Antibiotic prophylaxis is probably prescribed excessively or inappropriately for dermatological surgery and is thought to be best reserved for high-risk patients such as excisions from the lower limb. There are no data available on the current prescribing habits of Australian GPs regarding oral prophylaxis for minor excisions. The decision to prescribe antibiotic prophylaxis is complicated; in addition to efficacy, antibiotic costs, adverse effects and resistance must also be taken into account.

The role of antibiotic prophylaxis in the prevention of SSI following skin lesion excision remains controversial. Randomised trials have demonstrated that topical antibiotic prophylaxis is ineffective and should not be used. Infusion of antibiotic mixed with local anaesthetic, and systemic prophylactic regimes, have produced significant reductions in infection rates. The limited available literature has demonstrated comparable relative reductions in SSI to this study (table 4). Two of these studies examined the use of oral prophylaxis while the other two considerably larger studies examined various regimens of intramuscular cefazolin. With the exception of Amland et al and one arm of the 1994 study by Bencini et al, all of these studies examined short courses of prophylaxis rather than single doses. Although variations in the study populations and protocols make comparison difficult, the reductions achieved in this study appear to be at least similar to those achieved with short courses of prophylaxis and parenteral methods of delivery.

### CONCLUSION

In this study, a 2 g dose of cephalixin 30–60 min before skin lesion excisions from the lower limb was associated with a reduced risk of SSI from 35.7% to 12.5%.

### Table 4: Studies of systemic antibiotic prophylaxis for skin lesion excision

<table>
<thead>
<tr>
<th>Study, country</th>
<th>Setting and sample size</th>
<th>Intervention and control</th>
<th>Wound type</th>
<th>Infection rate (relative reduction)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amland et al.,28 Norway</td>
<td>Plastic surgery unit, 60 patients</td>
<td>Control: placebo Intervention: single dose 1000 mg orally azithromycin the night before surgery</td>
<td>Skin cancer surgery</td>
<td>Control: 16.0% Azithromycin: 5.7% (0.36)</td>
</tr>
<tr>
<td>Bencini et al.,29 Italy</td>
<td>Outpatient dermatologic surgery unit, 527 patients</td>
<td>A: no prophylaxis B: 1 g intramuscular cefazolin 12 hourly for 48 h before and after surgery C: 1 g intramuscular cefazolin 12 hourly from 2 h before surgery to 24 h after D: single 1 g intramuscular dose cefazolin 2 h before surgery</td>
<td>Excisions from contamination-prone areas (groin, axillae, interdigital spaces of feet)</td>
<td>A: 12% B: 4.6% (0.38)* C: 0.77% (0.064)* D: 2% (0.17)*</td>
</tr>
<tr>
<td>Czarnecki et al.,30 Australia</td>
<td>Outpatient dermatology clinic, 97 lesions</td>
<td>Intervention 1: 500 mg orally cephalexin three times a day, starting 2 days before and continuing 24 h postoperatively Intervention 2: topical mupirocin three times a day from 2 days before and continuing 24 h postoperatively Control: cetrimide-chlorhexidine cream 2 days before and continuing 24 h postoperatively</td>
<td>Ulcerated skin cancers which cultured positive for <em>Staphylococcus aureus</em> or Gram-negative bacteria</td>
<td>Control: 21.7% Intervention 1: 2.7% (0.12)* Intervention 2: 0% (0)*</td>
</tr>
<tr>
<td>Bencini et al.,31 Italy</td>
<td>Outpatient dermatologic surgery unit, 2165 patients</td>
<td>A: no antibiotic B: 1 g intramuscular cefazolin 12 hourly immediately after surgery continuing for 3 days C: 250 mg cefazolin powder applied locally during surgery D: 1 g cefazolin intramuscular 12 hourly starting 2 days before and continuing 2 days after surgery</td>
<td>Excision of skin lesions</td>
<td>A: 4.3% B: 1.5% (0.35)* C: 0.9% (0.21)* D: 0.2% (0.047)*</td>
</tr>
</tbody>
</table>

*Statistically significant reduction compared to control.
producing an absolute reduction of 23.2%, a relative reduction of 65% and an NNT of 4.3. This reduction, however, was not statistically significant (p = 0.064), and the trial was underpowered to address the hypothesis. As such, no definitive conclusion can be drawn on the efficacy of oral prophylaxis for lower limb skin lesion excision from this trial. Given the potential benefits, further investigation is warranted.

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Contributors SCS established and oversaw the study design and implementation, compiled the data and conducted the primary analysis. CFH conceived the study idea, identified suitable practices and assisted in study design and analysis. PGB assisted in sample size calculation and study analysis. All authors contributed to manuscript production.

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Provenance and peer review Not commissioned; externally peer reviewed.

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