Neuropathic agents in the management of pruritus in burn injuries: a systematic review and meta-analysis

Christopher McGovern1,2, Tara Quasim1,2, Kathryn Puxty1,2, Martin Shaw1,3, Wijnand Ng4, Charlotte Gilhooly1,2, Nikolaos Arkoulis5,6, Michael Basler7, Alan Macfarlane1,7, Lia Paton2

ABSTRACT

Objectives Pruritus is a common and often distressing complication after a burn injury. The purpose of this review is to explore the efficacy of drugs classically used to treat neuropathic pain in the management of pruritus after burn injury.

Methods A systematic literature search of medical databases was conducted to find studies investigating drugs listed in the National Institute for Health and Care Excellence (NICE) guideline (CG173, "neuropathic pain in adults") for the management of pruritus after burn injury in patients of any age. Controlled studies were stratified by the drug class studied and their risk of bias before conducting meta-analysis. A narrative review of case series or observational studies was presented. Severity of pruritus at any time point, with all quantitative and qualitative measures, was included.

Results Fifteen studies were included in the final analysis, 10 investigated the use of gabapentinoids, 4 studied doxepin, and 1 local anesthetic agents. Meta-analysis of three randomized controlled trials (RCTs) demonstrated that the use of gabapentinoids was associated with an improvement in mean VAS (Visual Analog Scale) 0–10 scores of 2.96 (95% confidence interval (95% CI) 1.20 to 4.73, p<0.001) when compared with placebo or antihistamines. A meta-analysis of four RCTs investigating topical doxepin showed an improvement in mean VAS scores of 1.82 (95% CI 0.55 to 3.09, p<0.001). However, when excluding two studies found to be at high risk of bias, no such improvement was found (−0.32, 95% CI −1.64 to −0.99, p=0.83).

Conclusion This study suggests that gabapentinoids are beneficial in the management of burn-related pruritus. There is a lack of evidence to suggest that doxepin is an effective treatment. Topical local anesthetic agents may be safe and beneficial, but studies are scarce.

Level of evidence Systematic review, level II.

INTRODUCTION

Burn injuries are acute traumatic insults. However, even with successful treatment, it is increasingly recognized that such injuries have far reaching and long lasting consequences.1 After even relatively minor burns, patients experience a persisting inflammatory response and immune dysfunction,2 leading to an increased risk of cardiovascular disease,3 cancer,4 infections,5 and early death.6 Although seemingly minor in comparison, pruritus is an additional long-term consequence that is common and significantly affects the quality of life of patients surviving burns.

There is a high prevalence of sensory disorders such as chronic pain, paresthesia, and pruritus in burns patients.7 The incidence of pruritus has been reported to be 93% at hospital discharge,8 67% to 73% at 2 years,9 9 and 44% at 4 years to 10 years.8 9 Factors such as deep dermal injury, greater total body surface area (TBSA) burned, an increased number of surgical interventions, female gender, and symptoms of post-traumatic stress disorder increase the risk.9

Achieving meaningful control of pruritus symptoms can be difficult and only a paucity of clinical trials have evaluated interventions.10–12 Histamine produced both from mast cell degranulation and as a by-product of collagen formation is thought to be a major contributor to the development of pruritus. A survey performed in the UK showed that over 90% of burns units used antihistamines as the first-line treatment.13 However, the involvement of various other peripherally acting pruritogens and the pathophysiological changes that occur more centrally mean that antihistamine monotherapy is often inadequate, especially in chronic pruritus.14

Although the neuronal pathways involved in the perception of pain have been extensively explored, the equivalent neuroanatomical basis for pruritus remains incompletely understood. A subset of afferent slow conducting C-fibers are activated by pruritogens including histamine, acetylcholine, calcitonin gene-related peptide, bradykinin, leukotrienes, prostaglandins, and various cytokines.15 Pain and pruritus share a similar neurophysiological basis, thought to be a consequence of evolutionary changes15 and after activation, these C-fibers conduct impulses in a similar manner to the pain pathway via the dorsal root ganglion, spinothalamic tract, thalamus, and then to various higher centers including the somatosensory cortex.14 15 Similarities have been drawn between chronic pruritus and neuropathic pain. Clinical features such as hyperkinesia and allokinesis mirror the hyperalgesia and allodynia seen in neuropathic pain secondary to peripheral and central sensitization.14 16–17 Such pathological processes are reflected in one classification of pruritus, as pruritogenic, neuropathic, neurogenic, and psychogenic.16

The objective of this narrative, systematic review is to evaluate the effectiveness of agents used in neuropathic pain, as detailed by the National
Institute of Clinical Excellence (NICE),18 in the management of pruritus after a burn injury.

**METHODS**

**Registration**

This review was registered on the PROSPERO Register of Systematic Reviews, ID number CRD420201647777. The PRISMA (Preferred Reporting Items for Systematic Review and Meta-Analyses) guidelines for the conduct of systematic reviews were followed throughout.19

**Eligibility criteria**

Articles were included that investigated the management of pruritus in patients of any age that had sustained a burn injury with the use of neuropathic agents that are listed in the NICE guideline (CG173) “neuropathic pain in adults: pharmacological management in non-specialist settings”.18 Given the likelihood of several studies being observational in nature, no restrictions were made regarding the use of a control group. Animal studies, human volunteer studies, literature reviews, and conference abstracts were excluded, otherwise no restrictions on the type of study were made.

**Search strategy**

Three databases, MEDLINE, EMBASE and the Cochrane Central Register of Controlled Trials (CENTRAL), were searched with no time period or language restrictions (last accessed January 7, 2021). The search strategies for each database can be found in the online supplemental material.

**Study selection**

After amalgamation of search results from the three sources and removal of duplicates, two authors (CM, WN) independently conducted a title review, abstract review, and then full article review to select articles for inclusion. Any disagreement between the two reviewers was resolved by a third reviewer (LP). The references of all titles included in the data analysis were screened for further articles to be included.

**Table 1** Randomized controlled trials of gabapentinoids

| Study            | Year | Setting/design | Inclusion criteria | Patient number | Age (years) | Intervention groups | Gabapentinoid dosing regimen | Outcomes | Follow-up |
|------------------|------|----------------|--------------------|----------------|-------------|---------------------|-------------------------------|----------|-----------|
| Zheng et al17     | 2015 | Single center RCT | >5% TBSA, second degree burn, >80% healed or healed within 3 months | 58 | 18 to 60 | 1. Gabapentin 2. Cetirizine 3. Placebo | Gabapentin 300 mg twice daily | Pruritus VAS | 28 days |
| Ahuja et al19     | 2013 | Single center RCT | >5% TBSA, 80% epithelialized or healed within 1 month | 80 | 18 to 60 | Four groups: 1. Antihistamine 2. Pregabalin and antihistamine 3. Placebo 4. Pregabalin | Pregabalin varied with severity of pruritus: Mild—75 mg twice daily Moderate—75 mg three times a day Severe—150 mg twice daily | Pruritus VAS | 28 days |
| Ahuja et al19     | 2011 | Single center RCT | >5% TBSA, 80% epithelialized or healed within 1 month | 60 | 12 to 70 | Three groups: 1. Cetirizine 2. Gabapentin 3. Combination cetirizine and gabapentin | Gabapentin varied with severity of pruritus: Mild—300 mg once daily Moderate—300 mg twice daily Severe—300 mg three times a day | Pruritus VAS | 28 days |
| Gray et al20      | 2011 | Single center RCT | Admitted to burn unit, >5% TBSA | 90 | 18 to 65 | 1. Pregabalin 2. Placebo | Pregabalin 75 mg twice daily, titrated up to 150 mg twice daily or 300 mg twice daily based on clinical response | NPS (includes itch score 0 to 10) Procedural pain score 0 to 10 4-point side effect scale | 28 days |

NPS, Neuropathic Pain Scale; QOL, quality of life; RCT, randomized controlled trial; TBSA, total body surface area.

**Data extraction**

Data were extracted by CM using a predefined spreadsheet which included study design, patient demographics, interventions, and outcomes. In the event of missing data, study investigators were contacted.

**Outcomes measured**

The outcome of interest was the severity of pruritus at any time point. No specific restrictions were used, with all quantitative severity scales, qualitative measures, and questionnaire methods of assessment included. For inclusion in meta-analysis, any quantitative scales were converted to an 11-point continuous scale and the mean difference between groups reported.

**Risk of bias assessment**

Each included study was assessed independently by CM and WN using a specific risk of bias tool. The RoB2 (the updated Cochrane risk of bias tool) was used for randomized controlled trials (RCTs), and ROBINS-I tool for non-randomized studies. The quality of evidence for the outcomes of interest were assessed using the GRADE (Grading of Recommendations, Assessment, Development and Evaluations) system.20

**Data synthesis**

Studies were categorized based on the intervention studied, specifically the neuropathic agent of interest, and are presented in tables for each class of agent. In controlled studies using the same drug or class of drug (eg, gabapentinoids) and comparable outcome measures (eg, a continuous variable such as visual analogue scale), results were collated using a random-effects meta-analysis, and two-sided p values and 95% confidence intervals were calculated. Heterogeneity was expressed as an I² statistic for studies included in meta-analyses. No specific sensitivity analyses were performed; however, studies were stratified by their risk of bias and meta-analyses were conducted separately for those at high and low/moderate risk and then combined. For agents where only case series or observational studies were available, a narrative review of the study findings was undertaken.
Table 2  Observational studies of gabapentinoids

| Study            | Year   | Setting/design  | Inclusion criteria                                                                 | Patient number | Age (years) | Intervention groups                                                                 | Dosing regimen                                                                 | Outcomes                          | Follow-up |
|------------------|--------|-----------------|-------------------------------------------------------------------------------------|----------------|-------------|-------------------------------------------------------------------------------------|--------------------------------------------------------------------------------|----------------------------------|-----------|
| Kneib et al⁵⁶    | 2019   | Single center retrospective cohort study | Admitted to burn unit, complained of itch                                             | 411            | >14         | Use of neuropathic pain protocol. Analyzed by group: 1. Gabapentin at <72 hours 2. Gabapentin >72 hours 3. No gabapentin | Gabapentin 300 mg once daily increased every 2 days if NRS5 to max 900 mg three times a day | Morphine equivalent dose Total gabapentin use Pain and itch NRS Short Form-12 survey | 24 months |
| Neusvendijk et al⁴⁹ | 2018   | Single center prospective observational study | In burn ward or outpatient clinic, complained of itch                                | 413            | <13         | Not protocolized. Mixture of no treatment, gabapentin, and antihistamine           | Gabapentin 5 mg/kg                | Itch Man Scale 0 to 4               | 1 week to 3 months |
| Kaul et al⁵⁰     | 2018   | Single center retrospective observational study | Pruritus or neuropathic pain on gabapentin                                           | 136            | 0 to 20     | Gabapentin only or in combination with pregabalin                                  | Various gabapentin doses used from 50 mg three times a day to 1200 mg three times a day, Pregabalin 50 mg to 100 mg three times a day added if inadequate response to gabapentin. | Retrospective review to describe effective dose of drugs | Unclear |
| Zachariah et al⁴⁷ | 2012   | Single center prospective observational study | Burn 6 weeks to 2 years old, pruritus with failure of cetirizine and emollients       | 23             | 4 to 60     | Gabapentin added if cetirizine inadequate                                            | Adults—gabapentin 100 mg twice daily, increased to max 300 mg three times a day Children—5 mg/kg twice daily to max 5 mg/kg three times a day | Itch severity scale (7 to 21 points) | 6 months |
| Goutos et al⁴⁴   | 2010   | Single center prospective observational study | Within 72 hours of injury, admitted to burn unit, sense of itch and urge to scratch | 91             | Adults and children                                                                 | Two consecutive protocols. Gabapentin introduced early or late, compared with pre-protocol data. | Adult 300 mg once daily titrated to 300 mg three times a day over 3 days Children 5 mg/kg once daily titrated to 5 mg/kg three times a day over 3 days | Itch Man Scale 0 to 4             | Unclear. Only inpatient data. |
| Mendham et al⁴⁷  | 2004   | Single center prospective observational study | Itching burn wound, admitted to burns unit                                            | 35             | Children    | Gabapentin (no comparator or control)                                               | 5 mg/kg three times a day to max 5 mg/kg twice daily plus 10 mg/kg nocte | Staff or parent reporting of itch reduction | Unclear. Some followed up for 18 months |

NRS, Numeric Rating Scale.

Numeric data were extracted from graphs if the required data were not included elsewhere in an article using Graphgrabber (V.2.0.2, Quintessa Ltd, Oxfordshire, United Kingdom). Meta-analysis was conducted using the software package Revman (V.5.4.1, Copenhagen: The Nordic Cochrane Centre, The Cochrane Collaboration).

RESULTS
Study inclusion
The literature search returned 5469 articles after removal of duplicates. The PRISMA flowchart (see online supplemental figure S1) details the selection process. One article required translation from Chinese.

Three main classes of neuropathic agents were investigated in the articles included in the final analysis:
- Gabapentinoids (gabapentin and pregabalin).
- Topical doxepin.
- Topical local anesthetic agents.

Gabapentinoids
Ten articles²¹⁻⁴⁰ investigated gabapentinoids in the management of pruritus after burn injury. Four studies were RCTs (table 1) and six were observational studies with varying methodology (table 2).

Randomized controlled trials
Three RCTs provided sufficient data to perform meta-analyses. Two studies²¹ ³⁰ included groups comparing gabapentin to a control arm given an antihistamine, cetirizine. Both studies demonstrated a mean reduction in pruritus severity, measured on 0 to 10 VAS (Visual Analog Scale) of around 6 points in the gabapentin group over the 28-day trial period. Those treated with cetirizine had a reduction of 3.9 and 3.5. The meta-analysis demonstrated a greater reduction in VAS score of 2.19 (95% CI 1.74 to 2.63) with gabapentin compared with cetirizine (figure 1).

Studies by Gray et al²⁴ and Zheng et al³⁰ included cohorts treated with a placebo. Gabapentinoids differed between studies, with Gray et al using pregabalin 150 mg 600 mg daily and Zheng et al using gabapentin 600 mg daily. Meta-analysis demonstrated an improvement in 0 to 10 itch severity score of 3.63 (95% CI 1.20 to 8.46) when a gabapentinoid was used in comparison to placebo (figure 1).

Combination of the above subgroup meta-analyses demonstrated an improvement in VAS of 2.96 (95% CI 1.20 to 4.73) when gabapentinoids are compared with control.

Although pruritus was not reported as a primary outcome in the study by Gray et al, it did report elements of the NPS (Neuropathic Pain Scale) including a 0 to 10 scale of pruritus

McGovern C, et al. Trauma Surg Acute Care Open 2021;6:e000810. doi:10.1136/tsaco-2021-000810
Treatment with gabapentin alone for pruritus, the measure of this outcome relied on adequate documentation in the patient medical notes and there was no comparator or control group.

Nieuwendijk et al. investigated the incidence, severity, and risk factors associated with pruritus in pediatric burn injury, then went on to describe pharmacotherapies used with 17.9% having received gabapentin. Unfortunately, as the study was principally designed to explore factors associated with pruritus, no conclusions could be drawn on the effectiveness of the pharmacological therapy.

Kneib et al. conducted a retrospective cohort study investigating the use of a neuropathic pain and pruritus protocol. Patients were started on incremental doses of gabapentin if itch scores remained greater than 4 (on 0 to 10 NRS) despite initial treatment with cetirizine. Comparison was made between various groups including pre-protocol and post-protocol introduction as well as patients that received gabapentin early (<72 hours), late (>72 hours), or not at all. There was no difference in itch severity odds ratios between any group at discharge through to 24 months.

Topical doxepin

Doxepin is a tricyclic antidepressant agent, but due to its potent antihistaminergic activity, is used topically to treat pruritus in eczema. Four studies investigated the use of topical doxepin on pruritic burn scars in adult patients (table 3). Given significant differences in both the results of these studies and the risk of bias assessments, meta-analyses are presented on studies at high risk and low/moderate risk of bias separately and then combined (figure 2).

Demling et al. performed two single center RCTs comparing topical doxepin to standard care. The results of both trials showed a marked improvement in itch VAS scores at all time points compared with standard care. The results of a meta-analysis including these studies demonstrated an improvement in mean VAS score of 3.10 (95% CI 2.73 to 3.47). Both of these studies were found to be at high risk of bias from a lack of blinding and unclear randomization methods (see online supplemental table S1). Additionally, the control arm of “standard care” involved titration of oral antihistamines that all participants were already taking prior to enrollment rather than introduction of another therapy or placebo.

Kwa et al. conducted two multicenter, blinded RCTs investigating the use of doxepin cream. The first study showed no difference in itch intensity at any time point between doxepin

| Study or Subgroup | Gabapentin vs antihistamine | Control | Mean | SD | Total | Mean | SD | Total | Weight | Mean Difference IV, Random, 95% CI | Mean Difference IV, Random, 95% CI |
|------------------|-----------------------------|---------|------|----|-------|------|----|-------|--------|----------------------------------|----------------------------------|
| 8.1.1 Gabapentin vs antihistamine | | | | | | | | | | 8.1.2 Gabapentin/Pregabalin vs placebo | |
| Ahuja 2011 | 6 | 0.55 | 20 | 3.9 | 0.98 | 20 | 26.5% | 2.10 [1.61, 2.59] | 2.60 [1.52, 3.69] | |
| Zheng 2015 | 6.1 | 1.54 | 20 | 3.5 | 1.83 | 20 | 24.9% | 2.19 [1.74, 2.63] | |
| Subtotal (95% CI) | 40 | 40 | 51.1% | 2.19 [1.74, 2.63] | |
| Heterogeneity: Tau² = 0.00; Chi² = 0.68; df = 1 (P = 0.41); I² = 0% | |
| Test for overall effect: Z = 9.87 (P < 0.00001) | |

Figure 1 Forest plot showing the reduction in mean VAS (Visual Analog Scale) in each treatment arm, comparing gabapentinoids with controls. 95% CI, 95% confidence interval.

severity. The effect of pregabalin on pruritus appeared much smaller than that demonstrated by the other studies included in the meta-analysis, with an improvement in mean scores of 1.17 (95% CI 0.10 to 2.24).

A further RCT was not included in the meta-analysis. Although the percentage changes in mean VAS scores were reported, the analysis did not include sufficient information regarding the distribution of the sample data, such as SD, to allow inclusion. This trial did, however, demonstrate a 78.9% fall in mean pruritus VAS scores in the group given pregabalin in comparison to 33.3% in the placebo group when focusing on patients with the most severe initial VAS scores (9 to 10). The placebo and antihistamine groups in this study suffered high dropout rates, however, potentially reflecting inadequate symptom relief.

Both RCTs conducted by Ahuja et al. also included groups given a gabapentinoid and antihistamine in combination but found no additional benefit when compared with gabapentinoid alone.

### Observational studies

Of the six observational studies (table 2), three were considered to be at serious risk of bias, two at moderate risk, and one to be of low risk (see online supplemental table S2). Sources of possible bias were primarily outcome measurements, which were often generated by research staff rather than patient reported.

Mendham et al. reported improvements in itch intensity when gabapentin was used in children with persisting itch despite treatment with antihistamines. Unfortunately, this outcome was measured solely on subjective reporting by nursing staff, parents, and children.

Goutos et al. investigated the use of two antipruritic protocols, with early or late introduction of gabapentin as part of incremental pharmacotherapy. In 41.3% of patients given gabapentin as the first-line agent, satisfactory itch control was achieved, in comparison to just 10% when cetirizine was used first line.

Zachariah et al. reported improved mean itch severity scores when gabapentin was given to pediatric and adult patients complaining of pruritus with inadequate relief on antihistamines and emollients. On a scale ranging from 7 to 21, mean itch severity scores fell from 13.35 to 8.36 within 1 month of treatment and this effect was sustained for the 6-month follow-up period.

Kaul et al. conducted a retrospective review of drug prescribing data of 136 mainly pediatric patients given gabapentin, pregabalin, or a combination of both for pruritus or neuropathic pain. Although 91.4% of patients had an adequate response to treatment with gabapentin alone for pruritus, the measure of this outcome relied on adequate documentation in the patient medical notes and there was no comparator or control group.

Nieuwendijk et al. investigated the incidence, severity, and risk factors associated with pruritus in pediatric burn injury, then went on to describe pharmacotherapies used with 17.9% having received gabapentin. Unfortunately, as the study was principally designed to explore factors associated with pruritus, no conclusions could be drawn on the effectiveness of the pharmacological therapy.

Kneib et al. conducted a retrospective cohort study investigating the use of a neuropathic pain and pruritus protocol. Patients were started on incremental doses of gabapentin if itch scores remained greater than 4 (on 0 to 10 NRS) despite initial treatment with cetirizine. Comparison was made between various groups including pre-protocol and post-protocol introduction as well as patients that received gabapentin early (<72 hours), late (>72 hours), or not at all. There was no difference in itch severity odds ratios between any group at discharge through to 24 months.

Topical doxepin

Doxepin is a tricyclic antidepressant agent, but due to its potent antihistaminergic activity, is used topically to treat pruritus in eczema. Four studies investigated the use of topical doxepin on pruritic burn scars in adult patients (table 3). Given significant differences in both the results of these studies and the risk of bias assessments, meta-analyses are presented on studies at high risk and low/moderate risk of bias separately and then combined (figure 2).

Demling et al. performed two single center RCTs comparing topical doxepin to standard care. The results of both trials showed a marked improvement in itch VAS scores at all time points compared with standard care. The results of a meta-analysis including these studies demonstrated an improvement in mean VAS score of 3.10 (95% CI 2.73 to 3.47). Both of these studies were found to be at high risk of bias from a lack of blinding and unclear randomization methods (see online supplemental table S1). Additionally, the control arm of “standard care” involved titration of oral antihistamines that all participants were already taking prior to enrollment rather than introduction of another therapy or placebo.

Kwa et al. conducted two multicenter, blinded RCTs investigating the use of doxepin cream. The first study showed no difference in itch intensity at any time point between doxepin

| Study or Subgroup | Gabapentin vs antihistamine | Control | Mean | SD | Total | Mean | SD | Total | Weight | Mean Difference IV, Random, 95% CI | Mean Difference IV, Random, 95% CI |
|------------------|-----------------------------|---------|------|----|-------|------|----|-------|--------|----------------------------------|----------------------------------|
| 8.1.1 Gabapentin vs antihistamine | | | | | | | | | | 8.1.2 Gabapentin/Pregabalin vs placebo | |
| Ahuja 2011 | 6 | 0.55 | 20 | 3.9 | 0.98 | 20 | 26.5% | 2.10 [1.61, 2.59] | 2.60 [1.52, 3.69] | |
| Zheng 2015 | 6.1 | 1.54 | 20 | 3.5 | 1.83 | 20 | 24.9% | 2.19 [1.74, 2.63] | |
| Subtotal (95% CI) | 40 | 40 | 51.1% | 2.19 [1.74, 2.63] | |
| Heterogeneity: Tau² = 0.00; Chi² = 0.68; df = 1 (P = 0.41); I² = 0% | |
| Test for overall effect: Z = 9.87 (P < 0.00001) | |
cream and the control group. Due to difficulty recruiting to the trial and a high dropout rate, this study was underpowered. A second study33 addressed these recruitment issues using a cross-over study design comparing doxepin against placebo without the inclusion of an antihistamine. Again, this demonstrated no difference in itch intensity between groups. A meta-analysis of both articles by Kwa et al (figure 2) using outcome data at 14 days showed no difference in changes in VAS scores in comparison to placebo or antihistamines, with a mean VAS change of \(-0.32\) (95% CI \(-1.64\) to 0.99).

All four studies investigating doxepin have been included in the final meta-analysis (figure 2) with no adjustment made for the risk of bias assessment. This demonstrated a reduction in mean VAS of 1.82 (95% CI 0.55 to 3.09). This result should be interpreted with caution as, when assessed using the GRADE tool, was found to be of very low quality, principally due to the high risk of bias (see online supplemental table S3).

### Topical local anesthetics

One study35 investigated the use of a topical local anesthetic agent in the management of pruritus after burn injury in children 1 to 5 years old. EMLA cream, a mixture of prilocaine and lidocaine, was applied to healed partial thickness burns with persisting pruritus in five patients. The main purpose of this study was to assess the safety and pharmacokinetics of this therapy. This study was performed over 3 days, with the first 2 days acting as a control for the treatment being implemented on day 3. There was an improvement in itch intensity as measured by a VAS and number of pruritic episodes. Owing to the young age of the children, outcome measures were made by parents, nursing staff, and the study investigators, potentially introducing an element of bias. This study suggested that the use of such topical local anesthetic agents was safe and may have potential benefit.

### DISCUSSION

This systematic review identified 15 studies investigating the use of various drugs often used to manage neuropathic pain to treat pruritus after burn injury. The analysis has demonstrated that gabapentin is effective in treating pruritus after a burn injury, resulting in an improvement of around 2 points on a VAS when compared with antihistamines. When compared with placebo, gabapentinoids were also beneficial, although the confidence intervals in the meta-analysis were wide. The drugs used in each of the two included studies also differed, as did the indication for their initiation. Zheng et al30 investigated gabapentin in the management of pruritus, whereas Gray et al24 investigated the use of pregabalin in patients with neuropathic pain, demonstrating a much smaller improvement in pruritic symptoms. This perhaps reflects the patient selection in this study whereby pain was the cardinal symptom, rather than pruritus. Given these limitations, it is therefore not possible to conclude whether this improvement is reproducible among the class of gabapentinoids or only evident with gabapentin.

Gabapentinoids are now used for a wide variety of indications. Although structurally similar to the inhibitory neurotransmitter GABA (gamma aminobutyric acid) found throughout the brain, they produce their analgesic effect through a different mechanism:

### Table 3 Studies of topical doxepin

| Study          | Year | Setting/design          | Inclusion criteria                        | Patient number | Intervention          | Control                        | Outcomes                  | Follow-up |
|----------------|------|-------------------------|-------------------------------------------|----------------|-----------------------|--------------------------------|---------------------------|-----------|
| Kwa et al32    | 2020 | Multicenter cross-over RCT | Healed burn, itch VAS≥3, pruritic area <10% | 27             | Doxepin cream         | Placebo cream                  | Pruritus VAS               | BIQ at week 2 and week 5 | 5 weeks   |
| Kwa et al32    | 2019 | Multicenter RCT         | Healed burn, itch VAS≥3                   | 31             | Doxepin cream and placebo tablet | Placebo cream and antihistamine tablet | Pruritus VAS               | BIQ QOL SF-36 Somnolence Erythema | 12 weeks |
| Demling et al34| 2003 | Single center RCT       | Healed, <35%TBSA partial thickness burn, pruritic area <20% | 31             | Doxepin               | Standard care                  | Pruritus VAS               | Erythema (Vancouver Scar Scale) | 12 weeks |
| Demling et al31| 2002 | Single center RCT       | Healed burn, pruritic area <15%           | 41             | Doxepin               | Standard care                  | Pruritus VAS               | Erythema (Vancouver Scar Scale) | 12 weeks |

BIQ, Burn Itch Questionnaire; QOL SF-36, Quality of Life Short Form-36 questionnaire; VAS, Visual Analog Scale.

Figure 2  Forest plot showing the reduction in mean VAS (Visual Analog Scale) in each treatment arm, comparing topical doxepin with controls. 95% CI, 95% confidence interval.
central nervous system they do not act on GABA receptors and their benefit in the management of pain and pruritus is likely to be due to action at voltage-gated calcium channels and NMDA (N-methyl D-aspartate) receptors within the spinal cord and brain, inhibiting the release of excitatory neurotransmitters.

Previous studies have demonstrated the benefit of gabapentinoi

drugs.59–61 In the context of individuals suffering a burn injury, such conditions63–67 but the evidence for benefit in neuropathic pain is only small numbers being included in the meta-analyses and a narrative review being conducted for the remaining studies.

Although the drugs we investigated are classically used to manage neuropathic pain, we elected not to focus on this specific pain condition, but rather pruritus given the increasing use of such drugs in other pruritic disorders and the recognized clinical and pathophysiological similarities between neuropathic pain and pruritus. Although some studies have demonstrated a reduction in morphine consumption and improved acute pain scores with the use of gabapentin,21–23 there is a lack of evidence studying their specific use in neuropathic pain in patients with burns.

The VAS or other numerical scoring systems were commonly used to report pruritus outcomes. However, other methods included itch episodes, breakthrough doses of antihistamines, and the itch severity scale. Difficulties were encountered due to these multiple reporting methods. Similar issues have been highlighted in other systematic reviews, often from pain management literature, highlighting the difficulties in standardizing and validating such outcome measures.

This systematic review also limited the interventions being investigated to those drugs listed by NICE for the management of neuropathic pain. Despite this wide inclusion criteria including multiple drugs and drugs classes, the literature search did not return any information on therapies such as other tricyclic antidepressants, including amitriptyline, or selective serotonin reuptake inhibitors that have been used to manage pruritus in other conditions.76–77 Other therapies beyond pharmacological management may be of benefit in pruritus, namely, psychotherapy such as cognitive behavioral therapy, transcutaneous electrical nerve stimulation, and acupuncture. These were not addressed in this review.

CONCLUSIONS

Gabapentin appears effective in the management of pruritus associated with burn injury. Topical lidocaine may be a safe and effective option for managing pruritus in small surface area healed burns. Topical doxepin, although used to manage pruritus in eczema, does not appear to be effective in burn injuries.

Acknowledgements The authors would like to thank the burn team at Glasgow Royal Infirmary for their advice on guidance, especially Mr Stuart Watson, Ms Eleanor Roberston, Katrina Dalgarno, Gillian Calder, and Ellen Meland.

Contributors CM preformed literature searches, study review and selection, conducted data collection and analysis, and drafted and revised the article. CM is the guarantor. WN performed study reviews and data collection. UP supervised literature review and study selection. TQ and KP revised the manuscript and supervised all aspects of the study. MS supervised statistical analysis and revised the manuscript. AM, NA, CG, and MB contributed to the planning of the study and revised the manuscript. CM is the

Competing interests None declared.

Patient consent for publication Not required.

Provenance and peer review Not commissioned; externally peer reviewed.

All data relevant to the study are included in the article

Open access This is an open access article distributed in accordance with the Creative Commons Attribution Non Commercial (CC BY-NC 4.0) license, which permits others to distribute, remix, adapt, build upon this work non-commercially, and license their derivative works on different terms, provided the original work is properly cited, appropriate credit is given, any changes made indicated, and the use is non-commercial. See: http://creativecommons.org/licenses/by-nc/4.0/.

ORCID iD

Christopher McGovern http://orcid.org/0000-0002-1587-1242
REFERENCES

1 Barnett LW, Fev WS, Waltham JC, Wood FM, Fear MW. Understanding acute burn injury as a chronic disease. Burns Trauma 2019;7:23.

2 Jeschke MG, Gauglitz GG, Kulp GA, Finnerty CC, Williams FN, Kraft R, Suman OE, Milic RP, Herndon DN. Long-term persistence of the pathophysiologic response to severe burn injury. Plast Reconstr Surg 2015;136:621–30.

3 O’Halloran E, Shah A, Dembo L, Hool L, Viola H, Grey C, Boyd J, O’Neill T, Wood F, Duke J, et al. The impact of non-severe burn injury on cardiac function and long-term cardiovascular pathology. Sci Rep 2016;6:34650.

4 Duke JM, Bauer J, Fear MW, Rea S, Wood FM, Boyd J. Burn injury, gender and cancer risk: population-based cohort study using data from Scotland and Western Australia. BMJ Open 2014;4:e003845.

5 Duke JM, Randall SM, Wood FM, Boyd JH, Fear MW. Burns and long-term infection disease morbidity: a population-based study. Burns 2017;43:5035-47(19):3049-1-273:81.

6 Duke JM, Rea S, Boyd JH, Randall SM, Wood FM. Mortality after burn injury in children: a 33-year population-based study. Pediatrics 2015;135:e903–10.

7 Malefant A, Forget R, Papillon J, Amzel R, Frigon YJ, Choiomère M. Prevalence and characteristics of chronic sensory problems in burn patients. Pain 1996;67:493–500.

8 Carragher GJ, Martinez EM, McMullen KS, Fauerbach JA, Holavanahalli RK, Herndon W, Wicman SA, Engwah LH, Gibson NS. Pruritus in adult burn survivors: postburn prevalence and risk factors associated with increased intensity. J Burn Care Res 2013;34:94–101.

9 Van Loey NEE, Bremer M, Faber AW, Middeckle E, Nieuwenhuis MK. Itching following burns: epidemiology and predictors. Br J Dermatol 2008;158:95–100.

10 Bell PI, Gabriel E. Evidence-based review for the treatment of post-burn pruritus. J Burn Care Res 2009;30:55–61.

11 Zachariah JR, Rao AL, Prabha R, Gupta AK, Paul MK, Lamba S. Post burn pruritus–a review of current treatment options. Burns 2012;38:621–9.

12 Morgan M, Deurs JR, Frøsig-Jørgensen M, Lewis RJ, Cabot PJ, Gray PD, Vetter I. Burn pain: a systematic and critical review of epidemiology, pathophysiology, and treatment. Pain Med 2018;19:708–34.

13 Bell I, MacDams T, Morgan R. Pruritus in burns: a descriptive study. J Burn Care Rehabil 1988;9:305–8.

14 Goutos I. Neuropathic mechanisms in the pathophysiology of burns pruritus: Redefining directions for therapy and research. J Burn Care Res 2013;34:82–93.10.1097/BCR.0b013e32828e4444.

15 Paus R, Schmelz M, Bird T, Steinhoff M. Frontiers in pruritus research: scratching and itchy skin (“alloknesis”) produced by intracutaneous injection of histamine. Somatosens Mot Res 1999;16:291–8.

16 Aghajanyan K, Amin A, Rosenberg M, Rosenberg L, Meyer WJ. Use of gabapentin and pregabalin in the treatment of antipruritic protocols in acute burns. The emerging value of gabapentin in the management of post burn pruritus. Burns 2011;37:203–7.

17 Goutos I, Aghajanyan K, Amin A, Rosenberg M, Rosenberg L, Meyer WJ. Use of gabapentin and their combination in the relief of post-burn pruritus: a pilot study. J Burn Care Rehabil 2001;22:235–42.

18 Rose MA, Kam PCA. Gabapentin: pharmacology and its use in pain management. Anaesthesia 2002;57:451–62.

19 Gottrup H, Juhl G, Kristensen AD, Lai R, Chi BH, Brown J, Bach FW, Jensen TS. Chronic oral gabapentin reduces elements of central sensitization in human experimental hyperalgesia. Anesthesiology 2004;101:1400–8.

20 Chincholkar M. Analogic mechanisms of gabapentinoids and effects in experimental pain models: a narrative review. Br J Anaesth 2018;120:1315–34.

21 Latremoileire A, Woof CJ. Central sensitization: a generator of pain hypersensitivity by central neural plasticity. J Pain 2009;10:695–92.

22 Apers G, Palik A, Ziersch E, Fuhrmann A, Alivans P. The use of pregabalin in the treatment of uracemic pruritus in haemodialysis patients. J Ren Care 2010;36:180–5.

23 Bueller HA, Bernhard JD, Dubroff LM. Gabapentin treatment for brachioradial pruritus. J Eur Acad Dermatol Venereol 1999;13:227–8.

24 Yusudian PD, Wilson NJE. Efficacy of gabapentin in the management of pruritus of unknown origin. Arch Dermatol 2005;141:1507–9.

25 Clarke H, Bonin RP, Orse BA, Engelsakis M, Wijersundera DN, Katz J. The prevention of chronic post-surgical pain using gabapentin and pregabalin: a combined systematic review and meta-analysis. Anesth Analg 2012;115:428–50.

26 Singh D, Kennedy DH. The use of gabapentin for the treatment of postoperative neuralgia. Clin Ther 2003;25:852–89.

27 Cooper TE, Derry S, Wilfen PJ, Moore RA. Gabapentin for fibromyalgia pain in adults. Cochrane Database Syst Rev 2017;2017:CD012188.

28 Vernet M, Lauzier F, Zychanski R, Perron C, Savard X, Pinard-A LA, Leblanc G, Cossi M-J, Nieuve X, Turgeon AE, et al. Perioperative use of gabapentinoids for the management of postoperative acute pain: a systematic review and meta-analysis. Anesthesiology 2020;133:265–79.

29 KY H, Gan TJ, Habib AS. Gabapentin and pregabalin postoperative pain – a systematic review of randomized controlled trials. Pain 2006;126:91–101.

30 Han Y, Xiao XJ, Jiang HQ, Ma LX, Ma XL. The use of gabapentin in the management of postoperative pain after total knee arthroplasty: a PRISMA-compliant meta-analysis of randomized controlled trials. Medicine 2016;95:e3883.

31 Zhai L, Song Z, Liu K. The effect of gabapentin on acute postoperative pain in patients undergoing total knee arthroplasty: a meta-analysis. Medicine 2016;95:e3673.

32 Tonnare N, Veluchamy A, Zhou Y, Fletcher EH, Moir E, Hebert HL, Donnan PT, Watson L, Coulon LA, Smith BH. Trends in gabapentin and pregabalin prescribing, co-prescribing of opioids and benzodiazepines, and associated deaths in Scotland. Br J Anaesth 2020;125:0007-0912(20)30373-1:159–67.

33 Montastry L, Foo SY, Renoux C. Trends in first gabapentin and pregabalin prescriptions in primary care in the United Kingdom, 1993-2017. JAMA Intern Med 2018;178:292–4.

34 Evoy KE, Morrison MD, Saklad SR. Abuse and misuse of pregabalin and gabapentin. Drugs 2017;77:402–26.

35 Molero Y, Larsson H, D’Onofrio BM, Sharp DJ, Fazel S. Associations between gabapentinoids and suicidal behaviour, unintentional overdoses, injuries, road traffic incidents, and violent crime: population based cohort study in Sweden. BMJ 2019;19:12147.

36 Smith RV, Haverus JR, Walsh SL. Gabapentin misuse, abuse and diversion: a systematic review. Addiction 2016;111:141.

37 Bonnet I, Schebaum N. How addictive are gabapentin and pregabalin? A systematic review. Eur Neuropsychopharmacol 2017;27:50924-977X(17)30897-1:1185–215.

38 Medicines and Healthcare Products Regulatory Agency. Pregabalin (Lycrata), gabapentin (Neurontin) and risk of abuse and dependence: new scheduling

McGovern C, et al. Trauma Surg Acute Care Open 2021;6:e000810. doi:10.1136/tsaco-2021-000810
requirements from 1 April 2019. https://www.gov.uk/drug-safety-update/pregabalin-lyrica-gabapentin-neurontin-and-risk-of-abuse-and-dependence-new-scheduling-requirements-from-1-april-risk-of-abuse-and-dependence.

59 Vetrievevel TP, Randall SM, Wood FM, Rea S, Boyd JH, Duke JM. A population-based comparison study of the mental health of patients with intentional and unintentional burns. Burns Trauma 2018;6:31.

60 Davis CS, Esposito TJ, Palladino-Davis AG, Rythlik K, Schermer CR, Gamelli RL, Kovacs EJ. Implications of alcohol intoxication at the time of burn and smoke inhalation injury: an epidemiologic and clinical analysis. J Burn Care Res 2013;34:120–6.

61 Klifto KM, Quiroga L, Hultman CS. Substance use and inhalation injury in adult burn patients: retrospective study of the impact on outcomes. Burns Trauma 2019;7:15.

62 Fasar JT, Berlin JA, Strom BL. Clinically important changes in acute pain outcome measures. J Pain Symptom Manage 2003;25:406–11.

63 Demant DT, Lund K, Finnerup NB, Vollert J, Maier C, Segerdahl MS, Jensen TS, Sindrup SH. Pain relief with lidocaine 5% patch in localized peripheral neuropathic pain in relation to pain phenotype: a randomised, double-blind, and placebo-controlled, phenotype panel study. Pain 2015;156:2234–44.

64 Derry S, Wiffen PJ, Moore RA, Quinlan J. Topical lidocaine for neuropathic pain in adults. Cochrane Database Syst Rev 2017:10.

65 Fiorelli A, Pace C, Cascone R, Carlucci A, De Ruberto E, Izzo AC, Passavanti B, Chiiodini P, Pota V, Ausilio C, et al. Preventive skin analgesia with lidocaine patch for management of post-thoracotomy pain: results of a randomized, double blind, placebo-controlled study. Thorac Cancer 2019;10:631–41.

66 Saber AA, Elgamal MH, Rao AJ, Iravri EA, Martinez RL. Early experience with lidocaine patch for postoperative pain control after laparoscopic ventral hernia repair. Int J Surg 2009;7:36–8.

67 Khanna M, Peters C, Singh JR. Treating pain with the lidocaine patch 5% after total knee arthroplasty. PM&R 2012;4:51934-1482(12)00285-7:642–6.

68 Inan S, Dun NJ, Cowan A. Inhibitory effect of lidocaine on pain and itch using formalin-induced noceception and 5′-guanidinonaltrindole-induced scratching models in mice: behavioral and neuroanatomical evidence. Eur J Pharmacol 2009;616:141–6.

69 Allenby CF, Johnstone RS, Chatfield S, Pike LC, Tidy G. PERINAL—a new no-touch spray to relieve the symptoms of pruritus ani. Int J Colorectal Dis 1993;8:184–7.

70 Layton AM, Cotterill JA. Natalgia paraesthetica—report of three cases and their treatment. Clin Exp Dermatol 1991;16:197–8.

71 Rimar S, Alavi C, Sedighinejad A, Toloie M, Kavoosi S, Kouchakinejad L. Effect of gabapentin on morphine consumption and pain after surgical debridement of burn wounds: a double-blind randomized clinical trial study. Arch Trauma Res 2012;1:38–43.

72 Cuignet O, Pirson J, Soudon O, Zizi M. Effects of gabapentin on morphine consumption and pain in severely burned patients. Burns 2007;33:81–6.

73 Gray P, Williams B, Crammond T. Successful use of gabapentin in acute pain management following burn injury: a case series. Pain Med 2008;9:371–6.

74 McClenaghan F. The concurrent use of gabapentin and opioid analgesia in burns patients. Arch Trauma Res 2012;1:81–2.

75 Malhotra A, Mackey S. Outcomes in pain medicine: a brief review. Pain Ther 2012;1:5.

76 Kouwenhoven TA, van de Kerkhof PCM, Kamsteeg M. Use of oral antidepressants in patients with chronic pruritus: a systematic review. J Am Acad Dermatol 2017;77:1068–73.

77 Magazin M, Daze RP, Okeson N. Treatment refractory brachioradial pruritus treated with topical amitriptyline and ketamine. Cureus 2019;11:e5117.