Review Article

Pain Management of Pediatric Musculoskeletal Injury in the Emergency Department: A Systematic Review

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Background. Pain management for children with musculoskeletal injuries is suboptimal and, in the absence of clear evidence-based guidelines, varies significantly. Objective. To systematically review the most effective pain management for children presenting to the emergency department with musculoskeletal injuries. Methods. Electronic databases were searched systematically for randomized controlled trials of pharmacological and nonpharmacological interventions for children aged 0–18 years, with musculoskeletal injury, in the emergency department. The primary outcome was the risk ratio for successful reduction in pain scores. Results. Of 34 studies reviewed, 8 met inclusion criteria and provided data on 1169 children from 3 to 18 years old. Analgesics used greatly varied, making comparisons difficult. Only two studies compared the same analgesics with similar routes of administration. Two serious adverse events occurred without fatalities. All studies showed similar pain reduction between groups except one study that favoured ibuprofen when compared to acetaminophen. Conclusions. Due to heterogeneity of medications and routes of administration in the articles reviewed, an optimal analgesic cannot be recommended for all pain categories. Larger trials are required for further evaluation of analgesics, especially trials combining a nonopioid with an opioid agent or with a nonpharmacological intervention.

1. Introduction

We have close to 20 years of evidence showing that analgesia for pediatric patients in the emergency department (ED) is suboptimal [1–6]. A recent medical record review of children presenting to an ED with acute musculoskeletal injury (MSK-I) confirmed that only 35% received an analgesic of any kind [7]. This is surprising, given the robust evidence that children experience both short- and long-term consequences when pain is undertreated [8–11].

Policy statements of pain and pediatric and emergency societies endorse the appropriate treatment of children’s pain as a key part of ED clinical care [12–15]. Children with MSK-I (i.e., simple fracture or severe sprain) in the ED will likely experience maximal pain during the period between the occurrence of the fracture and the point of immobilization of the limb in a cast. During this period, the child will often undergo manipulation of the injured limb by a nurse, physician, medical trainee, and an X-ray technician [7,12–16]. There is little doubt that sufficient analgesia would improve the child’s experience [16]. Yet, there is no consensus as to what constitutes the best analgesic or combination of analgesics for this type of injury.
To our knowledge, there are no systematic reviews of interventions in pain relief for children presenting to the ED with a MSK-I. Our objectives were (a) to review the available randomized controlled trials related to the ED pain management of MSK-I in children and (b) to quantify the occurrence of any serious adverse events related to the use of the analgesics among the included studies. We aimed to assess the efficacy of the various interventions used and, thereby, identify which one(s) to recommend in the ED for addressing pediatric MSK-I related pain.

2. Methods

2.1. Study Design. We conducted a systematic review of randomized controlled trials of nonpharmacological (e.g., distraction and music), physical (e.g., splinting), and pharmacological (e.g., opioids, nonopioids, and nonsteroidal anti-inflammatory medications) interventions in children and adolescents under 19 years of age presenting to the ED with a musculoskeletal injury. Studies of both adult and pediatric participants were included if the authors were able to provide pediatric-specific data. We excluded studies that compared therapies for fracture reduction, pain management in patients with osteogenesis imperfecta, and pain management for postdischarge pain. Substudies of previously reported trials, crossover studies, and abstracts or conference proceedings of trials without sufficient information were excluded. The primary outcome was the risk ratio for successful reduction in pain scores. The secondary outcome was the occurrence of serious adverse events.

2.2. Search Strategy. We performed a systematic search for published and unpublished articles in accordance with the Preferred Reporting Items for Systematic Review and Meta-Analysis (PRISMA) guidelines [17] using the following databases: Cochrane Central Register of Controlled trials (CENTRAL) (1984–April 2015), MEDLINE (1984–April 2015) using the Ovid interface, PubMed (1975–2015), Embase (1988–April 2015), and ProQuest (1980–April 2015). The search strategy is described in the Appendix. In order to identify unpublished and ongoing trials, we searched numerous international trials registers and Google Scholar. Key journals and conference proceedings from major meetings were hand-searched from 2000 to 2015. Lastly, we contacted authors for further information and checked reference lists of all included trials. We restricted our search to articles written in either French or English.

2.3. Study Selection. Three review authors (SLM, SA, and SG) independently screened titles and abstracts using a standardized form with predefined eligibility criteria. Full-text articles were independently assessed by each of the three authors to determine whether or not they met inclusion criteria. Discrepancies were resolved by consensus. Selected studies were evaluated for methodological quality and appropriateness. Studies rejected at this stage were noted with reasons for exclusion. Authors of studies were contacted, if necessary, for clarification or missing data. Review authors were not blinded to author, institution, journal, or results of a study prior to its assessment.

2.4. Data Collection and Analysis. Data for each included study were extracted and recorded independently by three review authors (SLM, SA, and SG) on a standardized data extraction form. Any differences were resolved by discussion between review authors until consensus was reached. The data was entered into Review Manager (RevMan) software (version 5.1. Copenhagen: The Nordic Cochrane Centre, The Cochrane Collaboration, 2011). We collected data for study eligibility, randomization technique, interventions, age, type of intervention, dose, route, and cointerventions. For the primary and secondary outcomes, we collected pain scores, mean pain score difference, and occurrence of serious (life-threatening) adverse events. Any missing data were sought through communication with the study authors.

Risk ratios were calculated to evaluate treatment effects according to the studies retained for this review. For continuous outcomes, results were summarized using a mean difference and 95% confidence interval (CI). We measured the efficacy of the interventions, but not the number needed to treat to benefit (NNT) or number needed to treat to harm (NNH). The decision to pool the data was based on comparability of the analgesic agents and their dosing among studies.

2.5. Assessment of Methodological Quality. Studies that met inclusion criteria were graded independently for methodological rigour using the Cochrane Collaboration’s Risk of Bias assessment tool [18] by three reviewers (SLM, AD, and NP). Studies were scored as high, low, or unclear risk of bias based on random sequence generation, allocation concealment, blinding of participants and personnel, blinding of outcome assessment, incomplete outcome data, selective reporting, and other biases. Discrepancies were resolved through discussion.

2.6. Dealing with Missing Data. All authors were contacted for missing data. When data remained missing, the studies were not included and are indicated in the flow diagram (Figure 1) with “no raw data available” as the reason of exclusion.

3. Results

3.1. Study Selection. The PRISMA Study Flow Diagram (Figure 1) shows the flow of studies for this review. Of the 34 studies of potential relevance identified, 26 were excluded: nine were not randomized controlled trials [16, 19–26]; one was an adult-only study [27]; one included only two children over a total of 67 patients with no specific pediatric data available [28]; one was concerned with fracture reduction [29]; one was conducted with healthy volunteers [30]; six had data collected at the participant’s home after discharge from the ED [31–36]. The remaining seven studies were excluded despite adequate methodology, because we were unable to obtain the raw data necessary to calculate their risk ratio,
1852 records identified through database search
138 records identified through other sources

1542 records after duplicates removed
1542 records screened
1508 records excluded

26 full text articles excluded:
(i) Not a RCT (n = 9)
(ii) Uncomparable outcome (n = 6)
(iii) No raw data available (n = 7)
(iv) Ineligible population (n = 4)

34 full text articles assessed for eligibility

8 studies included for analyses

Figure 1: PRISMA Study Flow Diagram.
Table 1: Characteristics of included studies.

| Author            | Participants                                      | Intervention*                                                                 | Main outcome                                                                                                                                                                                                 | Findings                                                                                                                                                                                                                                                                 |
|-------------------|---------------------------------------------------|-------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Borland et al. 2007 [44] | N = 67 children with acute long-bone fractures  
Age: 7 to 15 years | Group 1: n = 33, IN fentanyl 1.4 μg/kg  
Group 2: n = 34, IV morphine 0.1 mg/kg | Difference in MPS greater than 13 mm on the VAS at 5 minutes after analgesic administration | No significant differences in pain scores between groups at all study points                                                                                                      |
| Clark et al. 2007 [45] | N = 300 children with acute musculoskeletal pain  
Age: 6 to 17 years | Group 1: n = 100, acetaminophen 15 mg/kg  
Group 2: n = 100, ibuprofen 10 mg/kg  
Group 3: n = 100, codeine 1 mg/kg | Difference in MPS greater than 15 mm on the VAS at 60 minutes after analgesic administration | (i) Ibuprofen group showed significantly greater reduction in VAS but only 52% of participants in this group received adequate analgesia (VAS < 30 mm) |
| Friday et al. 2009 [46] | N = 66 children with fractures or dislocations  
Age: 5 to 17 years | Group 1: n = 34, ibuprofen 10 mg/kg, (max 400 mg)  
Group 2: n = 32, acetaminophen + codeine 15 mg/kg and 1 mg/kg (max 60 mg codeine) | Difference in mean pain score greater than 1 face on the WBS at 15 and 30 minutes after analgesic administration | Equivalent analgesic effectiveness of both agents at 40 minutes                                                                 |
| Furyk et al. 2009 [47] | N = 73 children with suspected limb fractures  
Age: 4 to 13 years | Group 1: n = 36, nebulized fentanyl 4 μg/kg  
Group 2: n = 37, IV morphine 0.1 mg/kg | Difference in mean pain score greater than 2 cm on the CAS at 40 minutes after analgesic administration | (i) Significantly decreased pain scores in both groups at all study time points  
(ii) No significant differences in pain scores between groups                                                                 |
| Graudins et al. 2015 [48] | N = 73 children with limb injury; weighing < 50 kg  
Age: 3 to 13 years | Group 1: n = 36, IN ketamine 1 mg/kg + ibuprofen 10 mg/kg  
Group 2: n = 37, IN fentanyl 1.5 μg/kg + ibuprofen 10 mg/kg | Median reduction in pain 30 min after analgesic administration  
FPS-R for children aged 3 to 6 and VAS for children 7 years and older | Similar pain reduction in both groups                                                                 |
| Kendall et al. 2001 [49] | N = 384 children with a clinical fracture of an upper or lower limb in the ED  
Age: 3 to 16 years | Group 1: n = 191, IN diamorphine spray 0.1 mg/kg  
Group 2: n = 193, IM morphine 0.2 mg/kg | Pain score on WBS at 30 minutes after analgesic administration | (i) Both medications significantly reduced pain  
(ii) Onset of pain relief was faster in the IN spray group than IM (lower pain scores at 5, 10, and 20 minutes)  
(iii) No difference between groups at 30 minutes                                                                 |
| Le May et al. 2013 [50] | N = 81 children with limb trauma  
Age: 6 to 18 years | Group 1: n = 41, ibuprofen 10 mg/kg + codeine 1 mg/kg (max 60 mg)  
Group 2: n = 40, ibuprofen 10 mg/kg (max 600 mg) + placebo | Difference in MPS of 20 mm on the VAS at 90 minutes after analgesic administration | No significant differences in mean pain scores between groups at all study time points  
(i) Significant reduction in mean pain scores in both groups  
(ii) No significant differences in pain scores between groups at 120 minutes                                   |
| Neri et al. 2013 [51] | N = 125 children with suspected fracture or dislocation  
Age: 4 to 17 years | Group 1: n = 60, ketorolac (SL) 0.5 mg/kg  
Group 2: n = 65, tramadol (SL) 2 mg/kg | McGrath scale for children up to 6 years and VAS for those older than 6 years. Primary outcome was MPS < 5/10 on a 0–10 VAS at 120 minutes | (i) Significant reduction in mean pain scores in both groups  
(ii) No significant differences in pain scores between groups at 120 minutes  
(iii) Rescue dose of paracetamol-codeine administered in 2/60 children in the ketorolac group versus 8/65 in the tramadol group (not significant) |

*Note: medications were given orally, except where otherwise indicated.  
CAS: Color Analog Scale; IM: intramuscular; IN: intranasal; IV: intravenous; MPS: Mean Pain Score; RCT: Randomized Controlled Trial; SL: sublingual; VAS: Visual Analog Scale; WBS: Wong-Baker faces pain scale.
medication used among the eight studies precluded the use of meta-analysis techniques. Three studies [45, 48, 50] provided the data necessary for the calculation of their RR in their respective article. For the remaining five studies [44, 46, 47, 49, 51], data was provided upon request by each of the corresponding author or coauthors. In all of the included studies, there were no "true" control groups (group receiving no analgesia), as it would be considered ethically unacceptable to refrain from treating a child's pain.

We were able to combine two studies, as they both compared fentanyl (intranasal and nebulized) with morphine (intravenous) [44, 47]. No statistically significant heterogeneity ($I^2 = 0\%$) was found and the RR was not significant (RR: 1.13; 95% CI: 0.93–1.38; $P = 0.22$) (Figure 2). The six other studies could not be pooled, as they studied different analgesics. The study by Friday et al. [46] compared the combination of acetaminophen-codeine to ibuprofen. The RR was not significant towards either group (RR: 1.30; 95% CI: 0.96–1.76; $P = 0.09$) ($n = 200$ children). Clark et al. [45] compared three different analgesics each one to the other, yielding three separate comparisons. In this study, ibuprofen was not considered as a control but having the same weight as acetaminophen and codeine. The RR of the first comparison between codeine and ibuprofen was not significant for either analgesic (RR: 0.90; 95% CI: 0.63–1.28; $P = 0.56$) ($n = 200$ children). Le May et al. [50] compared a combination of ibuprofen-codeine to a combination of ibuprofen-placebo of codeine. The RR was not in favour of either combination (RR: 0.76; 95% CI: 0.45–1.29; $P = 0.31$) ($n = 62$ children). The study by Graudins et al. [48] compared intranasal fentanyl/oral ibuprofen to intranasal ketamine/oral ibuprofen. The RR was not in favour of either combination (RR: 0.96; 95% CI: 0.77–1.22; $P = 0.76$) ($n = 68$). The study by Kendall et al. [49] compared intranasal diamorphine with intramuscular morphine. The RR was not in favour of either analgesic (RR: 1.14; 95% CI: 0.81–1.59; $P = 0.45$) ($n = 384$). Finally, the study by Neri et al. [51] compared sublingual ketorolac to sublingual tramadol. No statistically significant difference was observed between analgesics (RR: 1.10; 95% CI: 0.99–1.22; $P = 0.06$) ($n = 125$).

The minimally clinically significant difference (MCSD) in analgesic effectiveness varied across studies depending on the scale used. One study [47] used a mean pain score difference $>1.0$ for the WBS. On a 0 to 100 mm VAS, one study [44] used $>13$ mm and two studies [46, 48] used $>20$ mm, while one study [45] considered $<30$ mm as the pain score indicating adequate pain relief on that scale. Another study [50] considered 1.5 (standard deviation: 2.0) as a MCSD on a VAS that ranged from 0 to 10. Finally, on the same scale, another study [51] considered $<5/10$ as an adequate pain relief.

3.6. Serious Adverse Events. Only two serious adverse events were reported in the eight studies. Clark et al. [45] noted one important error in the preparation of the study medication

![Figure 2: Risk of bias.](image-url)
(codeine), which led to the administration of fivefold higher dose to the patient. No adverse outcome was observed for this patient as the error was quickly identified and the patient received proper treatment (oral charcoal and monitoring in the ED). In the group receiving IM morphine, Kendall et al. [49] reported one episode of nausea and vomiting that they considered a serious adverse event of moderate intensity. The patient was admitted for a short period of observation but recovered spontaneously. No other serious adverse events occurred among the remaining 1167 participants, including the 783 (66.98%) patients who received opioid.

4. Discussion

A child’s risk of sustaining a fracture before the age of 16 years ranges from 27 to 42%, [52–54] making MSK-I very common presentation to the ED [2, 55]. Unfortunately, the injury itself, as well as its assessment and treatment (i.e., radiographic tests and cast application), can cause significant pain and distress to a child. Children presenting to the ED with either a fracture or severe sprain often suffer from moderate-to-severe pain [33, 45, 56]. Current pain management, however, is recognized as suboptimal [7] and may be due in part to lack of clarity as to which medication to prescribe [57, 58]. The main objective of this systematic review was to identify the most effective interventions for reducing pain related to acute MSK-I. To our knowledge, this is the first systematic review of this subject. All of the studies retained were level I evidence: prospective randomized controlled trials with control groups (standard care or equivalent standard care). This review included eight randomized controlled trials [44–51] (1169 participants) assessing analgesics via different routes: oral, sublingual, intravenous, intramuscular, and intranasal.

4.1. Oral Pain Medications. One study demonstrated that ibuprofen was significantly more effective than acetaminophen [45] to relieve pain due to a simple fracture or severe sprain. Of note, only 52% of patients receiving ibuprofen achieved adequate pain relief, defined as <30 mm on a 100 mm Visual Analog Scale.

When codeine, alone, was compared to ibuprofen or acetaminophen [45], it did not provide better relief of pain. Moreover, even when codeine was combined with acetaminophen [46] or ibuprofen [50], these dual therapy combinations were considered equivalent to ibuprofen (10 mg/kg) alone [46]. The poor clinical performance for codeine can likely be explained by its genetic polymorphism-influenced metabolism, which can reduce its clinical effectiveness and influence its side effect profile [59–61]. Codeine, a prodrug, requires metabolizing to release morphine, its active analgesic component. The enzyme responsible for conversion of the prodrug to its active form is cytochrome P450-2D6. Approximately 50% of the North American Caucasian population has at least one reduced functioning allele for CYP2D6; this decreases the effectiveness of codeine for each affected individual [62–65]. Codeine has fallen out of favour as an analgesic for children because of this high interindividual variability and its association with drug-related deaths due to ultrarapid metabolizing [59, 66]. Currently, Health Canada recommends that no codeine-containing products should be used in any child less than 12 years of age [67]. In an effort to identify a reasonable agent to replace codeine, a recent study has compared oral morphine (0.5 mg/kg) to ibuprofen (10 mg/kg) for fracture pain in the post-ED setting [35]. Authors found no significant difference in analgesic efficacy between orally administered morphine and ibuprofen but noted that morphine was associated with a significantly greater number of side effects (56% versus 31%).

4.2. Intranasal Pain Medications. Three trials studied the use of intranasal fentanyl [44, 47, 48]. Two trials demonstrated that intranasal fentanyl was no different in efficacy compared to intravenous morphine to treat children with fractures [44, 47]. Larger sample sizes, with increased power, may show a significant difference between treatments, in the future. A third study compared intranasal fentanyl to intranasal ketamine (a dissociative anaesthetic) and demonstrated that the two were comparable in effectiveness for pain reduction, when combined with ibuprofen [48]. Similarly, another study [49] compared intranasal diamorphine (a semisynthetic moderate potency derivative of morphine) to intramuscular morphine and determined that they were associated with similar pain reduction in children with moderate-to-severe pain. Recent work has confirmed that intranasal diamorphine demonstrates no serious adverse events in over 200 children [68]. But another study [69] has shown an adverse event rate of 26.5%, which is in keeping with most other opioid medications. Intranasal pain medications have great intuitive appeal for clinicians, as they can be administered more quickly and without a potentially painful intravenous insertion or intramuscular injection. It is widely accepted that all intramuscular analgesic injections should be avoided in children when other routes are available [70]. Still, there remains limited evidence to definitively support intranasal pain medication use in children, at this time.

4.3. Sublingual Pain Medications. Sublingual medications have the appeal of rapid onset, can be used in children as young as six years of age, and preclude the need to swallow pills. A single study compared sublingual tramadol (a synthetic analogue of codeine and a weak u-opioid receptor agonist) to sublingual ketorolac (a nonsteroidal anti-inflammatory drug) [51]. While this study failed to demonstrate that either drug was more effective than the other, the results trended strongly towards ketorolac being more clinically effective. A larger trial would likely be able to clarify this. As with ibuprofen and codeine, a nonsteroidal anti-inflammatory medication appears to be outperforming weak opioid, even via the sublingual route.

4.4. Nonpharmacological Techniques. Few studies have addressed the efficacy of nonpharmacological interventions (e.g., splinting, cold, and distraction techniques) to relieve children’s pain related to MSK-I. We did not find a single trial specifically studying the analgesic effect of splinting injured limbs. We identified one nonpharmacological intervention study in our search [25] but it could not be included in the present review as it was not a randomized controlled
trial. In this study, Tanabe et al. [25] demonstrated that all groups (standard care, standard care plus 10 mg/kg ibuprofen, and standard care plus distraction) achieved a statistically significant reduction in pain at 30 minutes; the largest decrease in pain ratings was for those who received standard care plus distraction in the form of music and/or toys. A recently updated Cochrane Database systematic review of psychological interventions for procedural pain has confirmed that there is strong evidence for the use of distraction, in this setting [71]. Given that there is minimal risk involved in the use of distraction, it stands to reason that its benefits as an adjuvant to pharmacological therapy are both feasible and likely beneficial. Recognizing that implementing trials that study the efficacy of nonpharmacological interventions are extremely challenging and that nonpharmacological interventions, alone, may not be adequate, their utility could be studied in the context of an adjuvant to pharmacological therapy.

4.5. Perspectives on Future Studies. Current research would suggest that ibuprofen would be the recommended first choice for mild-to-moderate acute musculoskeletal pain. However, it may not be adequate, when used alone, for moderate-to-severe pain. The intranasal route for pain medication administration has recently become quite popular, with four of our eight included studies exploring this modality. While intranasal fentanyl appears to be promising, a large scale randomized controlled trial would be required to confirm its efficacy, as it is rapidly gaining popularity within the clinical realm, despite limited evidence. Further, because of its short-acting effect, it would probably require more than one dose to provide proper relief for a MSK-I. Moreover, once clinical efficacy and adverse effect profiles are better understood, further studies should also focus on the cost-effectiveness of these medications. While the cost of an individual dose of an oral medication (ibuprofen $0.38 for 400 mg suspension dose and acetaminophen $0.42 for 500 mg suspension dose) versus intranasal fentanyl ($0.45 for a 100 mcg ampoule) is comparable, the single patient-use atomizer required for intranasal delivery can be costly ($3.50–$5.00 per atomizer).

Finally, we would also recommend that future studies on pain management of MSK-I with children adopt a standardized primary outcome to assess pain management efficacy. Studies retained for this review had various primary outcomes of MPS from 13 to less than 30 mm on a 0–100 mm VAS as well as less than 5 over 10 on a 0–10 VAS and a difference of 2 cm on a 0–10 cm CAS. This wide range of primary outcomes makes it hard to properly compare results between studies. Further, when comparing analgesics in a trial, one should make sure that their respective delay for onset and peak of action are similar as well as setting a study time measure for the primary outcome accordingly. For instance, in Friday’s trial [46], measure of the primary outcome was set at 40 minutes after analgesic administration. Yet, all of the study medications for this trial were oral analgesics (ibuprofen, acetaminophen, and codeine) requiring at least 60 minutes to reach their peak of action. It was then kind of expected that the authors mentioned that they did not find any significant difference between groups at 40 minutes. They concluded by stating that ibuprofen was comparable in efficacy to the combination of acetaminophen and codeine, which is misleading. In fact, their results showed a positive trend in pain reduction in the group who received acetaminophen and codeine at 60 minutes.

4.6. Limitations. There were some limitations inherent to this systematic review. Firstly, we were unable to pool results and generate a summary statistic due to the disparity in the analgesics chosen and variations in the way intergroup differences in efficacy were reported. Secondly, the absence of response from some authors and the nonavailability of the data from several corresponding authors of studies meeting inclusion criteria prevented us from presenting all available studies regarding pain management of children presenting to the ED with MSK-I. Finally, the search was limited to articles published in either English or French, and no grey literature was searched for non-English language articles. The search was limited to randomized controlled trials and therefore excluded quasi-experimental studies and other designs.

5. Conclusion

In conclusion, based on our review of the currently available evidence, no one specific analgesic agent or intervention has been clearly identified as the optimal choice in the ED for all scenarios of pain management related to pediatric MSK-I. The ideal analgesic agent(s) for moderate-to-severe pain and the utility of nonpharmacological interventions in the pediatric population have yet to be determined. Our review results underscore the need for larger trials with a standardized primary outcome, to generate strong evidence for pain treatment of children presenting to the ED with a MSK-I. Finally, there is an urgent need to definitively address the issue of safety of commonly used analgesic agents in children, with special emphasis on opioid medications.

Appendix

Ovid Search Strategy. Database: EBM Reviews-Cochrane Central Register of Controlled Trials, Embase, Ovid MEDLINE(R)

(1) PAIN/
(2) pain$ or discomfort.ti,ab.
(3) Exp ANALGESICS
(4) analges$.ti,ab.
(5) OR/(1)–(4)
(6) FRACTURES, BONE/
(7) FRACTURES, CLOSED/
(8) Exp RADIUS FRACTURES
(9) HUMERAL FRACTURES/
(10) SHOULDER FRACTURES/
(11) ((simple or buckle or greenstick) adj6 fracture$).ti,ab.
Musculoskeletal injury (MSK-I) in children is very common and almost universally painful. Pain management of MSK-I in the emergency department (ED) varies greatly between hospitals. This paper is a systematic review of the current pain management for children presenting to the ED with MSK-I. Eight randomized controlled trials met inclusion criteria and provided data on 1,169 children. Due to heterogeneity of medications among the studies retained, an optimal analgesic cannot be presently recommended. Current research would suggest ibuprofen as first choice for mild-to-moderate musculoskeletal pain. However, there is no evidence for its superiority when used alone for moderate-to-severe pain.

**Competing Interests**

The authors declare that they have no competing interests.

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