Efficacy and cultural appropriateness of psychosocial interventions for Aboriginal and Torres Strait Islander paediatric burn patients and caregivers: a systematic review.

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Abstract

Background: Paediatric burns are highly painful and traumatising injuries that are overrepresented among Aboriginal and Torres Strait Islander people. Paediatric burn patients’ pain remains poorly managed by pharmacological interventions, leading to increased anxiety, distress and trauma in patients and their caregivers. Non-pharmacological psychosocial interventions have been suggested as effective in reducing pain and psychological morbidities among paediatric burn patients and their caregivers; however, the degree of effectiveness and appropriateness for Aboriginal and Torres Strait Islander people is unclear.

Methods: A non-date restricted systematic review was conducted through four databases. Studies published in English assessing psychosocial interventions on paediatric burn patients’ physical pain along with theirs and/or their caregiver’s anxiety, distress, or trauma symptoms were identified and included in this review. Included studies were assessed for their ability to reduce one of the outcomes of interests and for their reflection of Aboriginal and Torres Strait Islander peoples’ perspectives of health.

Results: Of the 3,178 identified references, 17 were eligible. These include distraction based techniques (n=8), hypnosis/familiar imagery (n=2), therapeutic approaches (n=4), and patient preparation/procedural control (n=3). Distraction techniques incorporating procedural preparation reduced pain, while discharge preparation and increased ‘patient control’ reduced patient and caregiver anxiety; and internet based Cognitive Behaviour Therapy reduced short-term but not long-term post-traumatic stress symptoms. No interventions reflected Aboriginal and Torres Strait Islander peoples’ perspectives of health; and few targeted caregivers
and focused on reducing their symptoms.

Conclusions: The development and assessment of psychosocial interventions to appropriately meet the needs of Aboriginal and Torres Strait Islander paediatric burn patients is required.

Background

Burn injuries cause severe pain [1–5] and can result in psychological trauma [2–4], anxiety [3, 5] and distress [1, 3]. These uniquely challenging injuries affect Aboriginal and Torres Strait Islander people at higher rates than non-Indigenous Australian people. This is highlighted by the Burns Registry of Australia and New Zealand’s most recent report that between 2017–2018 Aboriginal and Torres Strait Islander people were hospitalised for burn injuries three times more often than non-Indigenous people, and experienced significantly larger burns covering 10–49% of their Total Body Surface Area (TBSA) [6]. Paediatric specific data indicates similar discrepancies with Aboriginal and Torres Strait Islander children and adolescents experiencing 2.4 times higher rate of hospitalisation from burn injuries than non-Indigenous Australian children between 2011–2013 [7].

A unique challenge of these injuries lies within the persistent and debilitating base level of pain that is further intensified by regular procedures undergone for months to years following the initial injury [8, 9]. The complex nature of burn related pain often results in poor management despite the administration of standard doses of analgesia [8–13] and is particularly difficult to monitor among paediatric burn patients who are less able to articulate the intensity of their own pain [14]. This is further complicated for Aboriginal and Torres Strait Islander people who may not report their pain at all [15, 16] or verbally express their pain differently to non-
Indigenous Australians [17]. This is particularly concerning as poorly managed pain during hospitalisation strongly predicts burn patients’ psychosocial adjustment and overall wellbeing up to two years following treatment and hospital discharge [18]. More specifically, the pain and discomfort experienced by burn patients is associated with increased distress and anxiety [19]. This, in turn, increases the risk of developing other psychological morbidities such as acute stress and post-traumatic stress disorders (PTSD) [19]. The impact of burn related pain and distress is further exacerbated for paediatric burn patients who have a limited understanding of their injury and treatment [20], restricted agency in their care [21], and reduced ability to cope with the unpredictability of a hospital setting [22]. The struggles faced by paediatric burn patients is also greatly felt by their caregivers [20] who often experience overwhelming feelings of guilt, worry, panic, and anxiety whilst struggling with drastic shifts in their parenting role and ability to assist their child [23]. A review of empirical data highlights that 10–20% of paediatric burn patients and 4–42% of their caregivers reportedly experience symptoms of PTSD following the burn injury [24]. This highlights the need to effectively treat paediatric burn patients’ and their caregivers’ anxiety, distress, and psychological trauma. This is further emphasised by the finding that early onset of such psychological morbidities have a high rate of relapsing later in life [25].

The use of non-pharmacological, psychosocial interventions in conjunction with pharmacological analgesia have been suggested for reducing burn related pain and consequential psychological morbidities [26]. Interventions incorporating Gate Control Theory [21] techniques into change of dressing (COD) procedures are suggested as particularly effective in distracting the patient and reducing their ability to concentrate on painful stimuli [27]. Following this theory, virtual reality
has shown particularly favourable results on pain management among paediatric burn patients alone [28], and combined with young adult burn patients [29, 30]. Likewise, music therapy has shown promising effects in reducing anxiety and distress among paediatric burn patients alone [31–33], and combined with adults [34, 35]. Other psychosocial interventions utilising cognitive approaches and behavioural strategies have demonstrated similar effects and suitability for use among a wide age range of children and adolescents. More specifically, cognitive approaches including imagery, preparation techniques, information sharing, and coping strategies are suggested as particularly suitable for older children and adolescents [14]. While behavioural strategies including breathing exercises, desensitisation, and positive reinforcement are suggested as particularly suitable for younger children [14]. Several studies have presented the usefulness of such psychosocial interventions, however, no comprehensive comparison or systematic review has been conducted to assess their rigour, effectiveness, or appropriateness in meeting the needs of Aboriginal and Torres Strait Islander paediatric burn patients and their caregivers. This study assessed the effectiveness of any psychosocial intervention in reducing pain and psychological trauma, distress, and/or anxiety among paediatric burn patients and their caregivers generally. Alongside this assessment, we systematically evaluated the appropriateness and applicability of such interventions for use among Aboriginal and Torres Strait Islander families to inform the necessity and directions for future developments of culturally appropriate interventions.

Methods

Protocol and registration
Details of the protocol for this systematic review were registered on PROSPERO, the international prospective register of systematic reviews (CRD42018073451) [36].

Eligibility criteria

The below eligibility criteria were applied. Associated search terms were developed in consultation with experts from the University of Queensland library [see Additional file 1].

Table 1: Eligibility criteria

| Inclusion                                                                 | Exclusion                                                                 |
|---------------------------------------------------------------------------|----------------------------------------------------------------------------|
| 1. Studies focused on unintentional pediatric burn injuries.              | 1. Focus on non-burn injuries/illnesses or intentional burn injuries.      |
| 2. Injured children < 18 years receiving treatment at time of study and/or | 2. Injured adults > 18 years and/or injured children < 18 years post burns care. |
| their caregivers.                                                          | 3. Assessment of physical interventions i.e. dressings, physical therapy, massage etc. |
| 3. Assessment of psychosocial interventions*                              | 4. Studies with no clear comparison group.                                 |
| 4. Randomised control trials (RCT) or non-randomised control trials (NRCT) | 5. Assessment of any other outcome variable, or studies measuring only pain. |
| with clear comparison groups.                                              |                                                                            |
| 5. Assessing patient pain and theirs and/or caregiver’s anxiety, distress, |                                                                            |
| and/or trauma symptoms.                                                    |                                                                            |
| 6. Studies published in English with no date restrictions.                |                                                                            |

*Defined here as any intervention designed primarily to improve psychosocial wellbeing rather than physiological aspects of health.

Information sources

The Cumulative Index to Nursing and Allied Health Literature (CINAHL), Embase, MEDLINE, PubMed, and PsycINFO databases were systematically searched up to May 2018. Database alerts were established at this time and resulting references added as they became available. Reference lists of included manuscripts were hand-screened to identify additional articles not previously captured.

Study selection

Duplicates were removed prior to the lead author double screening all title and abstract references in the online systematic review software package Covidence, Australia and Argentina (v1086 e0dda871) [37]. An additional reviewer screened 10% of title and abstract references in Excel to verify accuracy with 91% agreement.

The lead author screened all full text references and two additional authors screened 50% each of full text references. Conflicts were resolved via consensus among all authors.

Assessment of cultural components, study quality and risk of bias.

Cultural components were extracted via a form developed in line with Milroy et al.’s
the Dance of Life [38], a multi-dimensional model reflecting the interconnectedness and complex layering of Australian Aboriginal people’s perspectives of health and wellbeing [see Additional file 2]. In accordance with the model, data was extracted on five core aspects of health: physical (4 items), psychological (3 items), social (4 items), spiritual (3 items), and cultural wellbeing (3 items). All items was graded zero (not present) or one (present) with relevant details extracted. The cultural components data form is included as supplementary materials.

Risk of bias was assessed against the Cochrane Collaboration’s tool for assessing risk of bias [39] (nine items), and an additional three items from the Cochrane suggested risk of bias criteria for EPOC reviews [40] to assess potential intervention contamination, and baseline outcome and characteristic similarities. All risk of bias items were graded zero (high/unclear bias) or one (low risk of bias). Study quality was evaluated by the Downs & Black Checklist [41] (28 items) to assess reporting, external validity, internal validity, and study power. Power was graded based on the smallest sample group (0 = n<1, 1 = n1-n2, 2 = n3-n4, 3 = n5-n6, 4 = n7-n8, 5 = n>8), and all other items graded zero (no) or one (yes); possible maximum score of 31.

Hooper et al.’s classifications [58] were adapted to provide overall quality rankings of poor (≤ 14), fair (15–19), good (20–25), or excellent (26–31). Assessment and data extraction were conducted by the lead author for all studies, and independently by two co-authors for 50% of studies.

Data synthesis

Meta-analysis was not appropriate due to high heterogeneity among studies. Therefore, data was synthesised narratively, presented in text by outcome (i.e. pain, anxiety, distress, and trauma), and tabulated by intervention type.
Results

Study selection

Database searches returned 3,638 abstracts published prior to 18th November 2019. Duplicates were removed and the remaining 1,937 abstracts underwent title and abstract screening, resulting in 1,821 exclusions and 116 inclusions for full text revision. Exhaustive attempts to obtain full manuscripts was successful for 100 abstracts. Of the remaining 16, five were not able to be acquired and 11 were unavailable in English. Full manuscripts were double screened by the lead author and two co-authors, resulting in a further 82 exclusions. References of the remaining 18 studies were hand-screened, identifying an additional ten abstracts of interest, two of which were included for data extraction [see Additional file 3]. Data extraction was not possible for three studies despite repeated attempts to contact study authors to obtain missing data and further information. Data was extracted from the final 17 eligible studies and narratively synthesised. Exclusion rates are outlined in the PRISMA flow diagram (Fig. 1.).

Main results

The study and participant characteristics of included interventions are presented in Tables 2 and 3 respectively. The key findings of included interventions are presented by intervention type in Table 4, and narratively synthesised by outcome below.

|          | N   | Age M (SD) | Age Range | Male n (%) | TBSA% M (SD) | Ethnicity n (%) |
|----------|-----|------------|-----------|------------|--------------|-----------------|
|          |     |            |           |            |              | Reported Not reported First Nation |
| Blakeney | I   | 32         | 14 (1.8)  | 12-17      | 9 (28)       | 36.8 (25.1)     | 32 (100) 0 (0) 2 (6)* |
|          | C   | 32         | 14.2 (1.9)|            | 17 (53)      | 44.2 (20.6)     | 32 (100) 0 (0) 0 (0)* |
| Brown    | I   | 47         | 8.3 (2.5) | 4-13       | 27 (57.5)    | 1.9 (2.2)       | 47 (100) 0 (0) 0 (0) |
|          | C   | 52         | 8.2 (2.7) |            | 33 (63.5)    | 1.9 (2.1)       | 52 (100) 0 (0) 0 (0) |
| Burns-   | I   | 15         | 7.8 (2.3) | 4-12       | 8 (53)       | 9.2             | 15 (100) 0 (0) 0 (0) |
| Nader | C  | 15 | 7.1 (2.8) | 11 (73) | 6.4 (7.5) | 15 (100) | 0 (0) | 0 (0) |
|-------|----|----|-----------|---------|-----------|---------|-------|-------|
| Chester | I  | 29 | 8.6 (3.4) | 16 (59) | 1.2 (2.0)^ | 0 (0) | 29 (100) | 0 (0)^** |
|       | C  | 35 | 7.1 (2.7) | 22 (63) | 1.0 (2.0)^ | 0 (0) | 35 (100) | 0 (0)^** |
| Elliott | I  | 4  | 8.5 (3.5) | 4 (100) | 21.5 (15.0) | 3 (75) | 1 (25) | 0 (0) |
|       | C  | 4  | 6.7 (2.1) | 4 (100) | 32 (24.3) | 1 (25) | 3 (75) | 0 (0) |
| Foertsch | I  | 13 | 5.8 | 12 (52) | 11.4 | 0 (0) | 13 (100) | 0 (0) |
|       | C  | 10 | 5.8 | 0 (0) | 0 (0) | 0 (0) | 10 (100) | 0 (0) |
| Hyland | I  | 50 | 2.3 (1.5–4.5)^ | 25 (50) | 0.8 (0.5–2.0)^ | 0 (0) | 50 (100) | 0 (0) |
|       | C  | 50 | 2.2 (1.6–3.9)^ | 27 (54) | 0.5 (0.5–2.0) | 0 (0) | 50 (100) | 0 (0) |
| Jeffs | VR | 8  | 14.3 (2.0) | 3 (38) | 7.4 (8.5) | 6 (75) | 2 (25) | 0 (0) |
|       | PD | 10 | 12.6 (2.1) | 8 (80) | 3.4 (3.3) | 9 (90) | 1 (10) | 0 (0) |
|       | C  | 10 | 13.9 (2.8) | 8 (80) | 4.7 (6.9) | 10 (100) | 0 (0) | 0 (0) |
| Kavanagh | I  | 4  | 6.25 (4.4) | 4 (100) | 22.7 (9.1) | 0 (0) | 4 (100) | 0 (0) |
|       | C  | 5  | 7.1 (3.8) | 2.5–11.5 | 3 (60) | 37.5 (27.1) | 5 (100) | 0 (0) |
| Kipping | I  | 20 | 12.6 (1.3) | 13 (65) | 5.1 (6.3) | 0 (0) | 20 (100) | 0 (0) |
|       | C  | 21 | 13.5 (1.8) | 15 (71) | 4.7 (4.5) | 0 (0) | 21 (100) | 0 (0) |
| Miller^a | VGD | 20 | 6.6 (2.5) | 12 (60) | 2.6 (1.4) | 0 (0) | 20 (100) | 0 (0) |
|       | MMMD-D | 20 | 6.6 (2.6) | 13 (65) | 2.8 (1.9) | 0 (0) | 20 (100) | 0 (0) |
|       | MMMD-PP | 20 | 5.5 (2.1) | 14 (70) | 4.3 (4.2) | 0 (0) | 20 (100) | 0 (0) |
|       | C  | 20 | 6.1 (2.1) | 8 (40) | 2.5 (1.4) | 0 (0) | 20 (100) | 0 (0) |
| Miller^b | Patient | 12 | 3.0^ | 6 (50) | 2.8 (1.0) | 20 (100) | 0 (0) | 0 (0) |
|       | Caregiver | 9  | 3.0^ | 3 (33) | 12 (100) | 0 (0) | 0 (0) | 0 (0) |
| Moore | Patient | 12 | 3.0^ | 6 (50) | 2.8 (1.0) | 20 (100) | 0 (0) | 0 (0) |
|       | Caregiver | 9  | 3.0^ | 1 (8) | 12 (100) | 0 (0) | 0 (0) | 0 (0) |
| Quay | I  | 26 | 5.3 | 0.7–15 | 23 (1–81) | 0 (0) | 26 (100) | 0 (0) |
|       | C  | 24 | 5.3 | 0 (0) | 24 (100) | 0 (0) | 0 (0) | 0 (0) |
| Sveen | Patient | 26 | 5.3 (3.5) | 13 (50) | 8.5 (7.0) | 0 (0) | 26 (100) | 0 (0) |
|       | Caregivers | 31 | 36.4 (6.6) | 14 (61) | 9.9 (7.0) | 0 (0) | 23 (100) | 0 (0) |
|       | Caregivers | 31 | 38.3 (5.5) | 9 (29) | N/A | 0 (0) | 31 (100) | 0 (0) |
| Van der Heijden | I  | 71 | 2.0 (13.1–4.1)^ | 37 (52) | 7 (4–13)^ | 0 (0) | 71 (100) | 0 (0) |
|       | C  | 64 | 1.7 (1.3–2.9)^ | 32 (50) | 10.5 (5–15)^ | 0 (0) | 64 (100) | 0 (0) |
| Whitehe | I  | 8  | - | 6–16 | 5 (36) | 0 (0) | 8 (100) | 0 (0) |
|       | C  | 6  | - | 0 (0) | 6 (100) | 0 (0) | 0 (0) | 0 (0) |

I: Intervention, VR: Virtual reality intervention, PD: Passive distraction intervention, VGD: Video game distraction intervention, MMMD-D: Multi-modal Device-Distraction intervention, MMMD-PP: Multi-modal Device-Procedural Preparation intervention, C: Control; M: Mean, SD: Standard deviation, ^Median (IQR), *Native American, **Aboriginal and Torres Strait Islander/South Sea Islander.

Table 3
Study characteristics
| First author, year. Location. Design. | Intervention | Control | Outcome: measures\(^{\text{assessor}}\) | Measurement time points |
|--------------------------------------|-------------|---------|------------------------------------------|------------------------|
| **Blakeney, 2005. USA. RCT. 42**     | n = 32^\text{\(^3\)} (9) 4-day group social skills workshop based on Changing Faces REACH OUT, and ‘usual’ treatment. | n = 32^\text{\(^{10}\)} (31) ‘Usual’ treatment, follow-up psychological appointments upon request only. | Anxiety/depression: CBCL\(^{(C)}\). | Pre-intervention & 1 year post-intervention: CBCL. |
| **Brown, 2014. Australia. RCT. 45**  | n = 47^\text{\(^{12}\)} (23) Ditto™ PP pre-COD & distraction interactive story/game during COD. | n = 52^\text{\(^{12}\)} (26) Standard distraction during COD: TV, videos, books, toys, caregiver soothing. | Pain: FPS-R\(^{(pt.)}\), HR\(^{(N)}\). Pain & distress: FLACC\(^{(N)}\). Anxiety: VAS-A\(^{(pt. > 8\ yrs)}\). Trauma: CTSO\(^{(pt. > 6\ yrs)}\). | Pre-randomisation: FPS-R, HR, FLACC, VAS-A. Pre-removal: FPS-R, FLACC, VAS-A. Post-removal: FPS-R, HR, FLACC, VAS-A, CTSO\(^{(1\text{st} COD)}\). Post-application: FPS-R, HR, FLACC, VAS-A. During removal & application: HR. 3-months post-re-epithelisation: CTSO. |
| **Burns-Nader, 2017. USA. RCT. 48** | n = 15^\text{\(^{0}\)} (0) Tablet distraction game, CLT support during 2nd and/or 3rd COD. | n = 15^\text{\(^{0}\)} (0) Standard distraction, CLT support during 2nd and/or 3rd COD. | Pain: FACES\(^{(pt.)}\), nurse’s pain reports\(^{(N)}\). Anxiety: CEMS\(^{(Pt.)}\). | Prior & during hydrotherapy: CEMS. Post-hydrotherapy: FACES, nurse’s pain reports, CEMS. |
| **Chester, 2018. Australia. RCT. 44** | n = 29^\text{\(^{0}\)} (0) Hypnosis pre & during COD: guided imagery, breathing, muscle relaxation, permissive & direct hypnotic suggestions. | n = 35^\text{\(^{0}\)} (0) Standard interventions pre & during COD: parent presence, books, TV, electronic games, DVDs, toys, bubbles, music, Ditto™ PP & distraction. | Pain: FPS-R\(^{(pt.)}\), FLACC\(^{(N)}\), NRS\(^{(C)}\), HR\(^{(R)}\). Anxiety: VAS-A\(^{(pt. > 8\ yrs, \ C < 8\ yrs)}\). Trauma: CPSS\(^{(pt. > 7\ yrs)}\). YCPC\(^{(C < 7\ yrs)}\). | Pre & post-procedure: FPS-R, FLACC, NRS, VAS-A. During procedure: FPS-R, FLACC, NRS. Pre-medications, post-application: HR. 3-months post-injury: CPSS, YCPC. |
| **Elliott, 1983. USA. NRCT. 55**     | n = 4^\text{\(^{0}\)} (0) Stress management during COD: Distraction, breathing, emotive imagery, pain reinterpretation. | n = 4^\text{\(^{0}\)} (0) SC during COD. | Pain & distress: BTDS\(^{(MS)}\). | Removal, first 15 min of hydrotherapy, during physical therapy, dressing re-application: BTDS. |
| **Foertsch, 1998. USA. RCT. 56**     | n = 13^\text{Total \(1\)} (43) Familiar imagery during COD: focus on childhood memory/experience. | n = 10^\text{Total \(1\)} (43) Social support during COD: researcher conversation & encouragement. | Pain & anxiety: FACES\(^{(Pt. ’s 3–9\ yrs)}\), VAS\(^{(Pt. ’s 9–12\ yrs)}\). Distress: OSBD\(^{(R)}\). | Baseline & 15-second intervals during procedure: OSBD. Post-procedure: FACES, VAS. |

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**First author, year. Location. Design.**

**Intervention**

**Control**

**Outcome: measures\(^{\text{assessor}}\)**

**Measurement time points**
| First author, year, Location, Design. | Intervention | Control | Outcome: measures (assessor) | Measurement time points |
|---------------------------------------|--------------|---------|-----------------------------|------------------------|
| Jeffs, 2014. USA. RCT.                | n = 8^0 (0) 1VR, 3D interactive program pre & during COD. | n = 10^0 (0) PD; PD pre & during COD: movie | n = 10^0 (0) 'Typical' care during COD: standard nurse communication. | Procedural pain and anxiety questionnaire, CFS. |
| Kavanagh, 1983. USA. NRCT.            | n = 4^0 (0) Max. procedure 'predictability' - specific nurse attire, 'patient control' of 'appropriate' aspects of procedure. | n = 5^0 (0) Min. 'predictability', medical staff control over procedure. | Pain: Nurse reports 0–6 scale(N). Anxiety: CBI(N & C of pt.'s 1–3 yrs). | 2–3 times daily: Nurse reports. Weekly: CBI. |
| Kipping, 2012. Australia. RCT.        | n = 20^0 (0) Off-the shelf VR pre & during COD. | n = 21^0 (0) Standard distraction during COD: TV, stories, music, caregiver. | Pain: VAS(Pt. & C). Pain & Distress: FLACC(N). | Baseline, retrospective post-removal, & application: VAS, FLACC. |
| Miller, 2010. Australia. RCT.         | n = 20^2 (10) MMD−PP: MMD−PP pre-COD, standard distraction during COD. | n = 20^3 (15) MMD−D: MMD-D interactive story/game during COD. | n = 20^4 (20) MVD: VGD during COD. | Pain: FACES(Pt.), VAS(C). Pain & distress: FLACC(N). Pre & post-removal, pre & post-application: FACES, VAS, FLACC. |
| Miller, 2011. Australia. RCT.         | n = 20^0 (0) MMD−PP, MMD-D interactive story/game pre & during COD. | n = 20^0 (0) Standard PP & distraction pre & during COD. | n = 20^0 (0) Standard PP & distraction during COD: toys, TV, & nurse/caregiver interactions. | Pain: FACES(Pt.), VAS(C). Pre & post-removal, pre & post-application: FACES, VAS, FLACC. |
| Moore, 2015. USA. NRCT.               | n = 12^0 (0) CBT MP pre-COD: standard medical equipment & puppets. | n = 9^0 (0) SC during COD: Standard PP, clinical staff verbal explanations. | n = 20^0 (0) MMD−PP, MMD−D interactive story/game & puppets. | Pain: FPS(Pt.), Pain & distress: FLACC(R). Pre & post-procedure: FPS, FLACC, STAI-CH. Post-removal: FPS. |
| Ouay, 1983. USA. RCT.                 | n = 26^0 (0) Discharge preparation weekly by nurse, written information, procedural rehearsal 3 days pre-discharge. | n = 24^0 (0) Routine instructions 3 days pre-discharge. | n = 24^0 (0) Routine instructions 3 days pre-discharge. | 1-day pre-discharge, 1st follow-up visit: STAI-CH. |
| Sveen, 2017. Sweden. RCT.             | n = 31^16 (52) Internet based CBT & ACT support program. | n = 31^3 (10) SC during COD. | n = 31^3 (10) SC during COD. | Posttraumatic stress: IES-R, PSI-SF, PSS(C). Pre-procedure, post-procedure, 3mths post-injury, 12mths post-injury: IES-R, PSI-SF, PSS. |
| Van der Heijden, 2018. South Africa    | n = 71^3 (4) 3–5 minute MT. | n = 64^3 (4) SC during COD. | n = 64^3 (4) SC during COD. | Pain: COMFORT-R(R), FACES(PL). Pre-procedure, hallway, entering |
| Author (year) | Outcomes | Results |
|--------------|----------|---------|
| *PP & distraction* Brown [45] | Pain | • FPS-R scores lower in Ditto™ than control at post-application of 2nd COD (MD=-1.51 [CI:-2.89, -0.13] \( p = 0.032 \)). • HR lowered across 3 COBs in Ditto™ group (MD=-4.89 [CI:-9.69, -0.09], \( p = 0.046 \)). |
|            | Pain & distress | • FLACC scores not reported. |
|            | Anxiety | • VAS-A scores lower in Ditto™ than control at pre-removal (MD=-1.79 [CI:-3.59, -0.01] \( p = 0.51 \)). |
| *Burns-Nader* [48] | Pain | • Intervention group did not affect CTSQ scores 1 week post-injury (MD not reported [CI:-1.49, 0.87] \( p = 0.602 \)) or 3 months post re-epithelialisation (MD not reported [CI: -1.26, 2.00] \( p = 0.651 \)). • Nurse’s pain reports lower in tablet group (M = 3.73, SD = 0.88) than control (M = 2.93, SD = 1.03), \( p = 0.030 \). |
|            | Anxiety | • CEMS scores higher in control (\( p = 0.001 \)) and after (\( p = 0.002 \)) hydrotherapy. • CEMS scores remained higher in control post-procedure (\( p < 0.050 \)), tablet group returned to baseline levels (\( p = 0.57 \)). |
| *Miller* [46] | Pain | • FACES scores sig. differed at pre & post-removal, and pre & post-application of all 3 COBs (\( p = \leq 0.001 \) all time points). |
• FACES scores lowered across 3 COD’s in:
  o MMD-D at pre-removal (p = ≤ 0.001), post-removal (p = 0.005), and pre-application (p = 0.004).
  o MMD-PP at pre-removal (p = 0.044).
  o VGD at post-application (p = 0.030).

• FACES scores lowered in:
  o MMD-PP more than VGD and control at pre-removal (both p = ≤ 0.01), post-removal (both p = < 0.001), pre-application (both p = < 0.001), and post-application (both p = < 0.001).
  o MMD-D more than control at pre-application (p = ≤ 0.05), post-removal (p = < 0.001), and VGD at post-removal (p = < 0.001) and post-application (p = < 0.001).

• VAS scores sig. differed at pre-removal of 2nd & 3rd COD; and post-removal, pre & post-application of all 3 CODs (p = ≤ 0.001).

• VAS scores increased across 3 CODs in VGD compared to MMD-PP at post-removal & application (both p = < 0.001), and MMD-D at post-removal (p = ≤ 0.05) and post-application (p = ≤ 0.001).

• VAS scores lowered across 3 COD’s in:
  o MMD-D at pre & post-removal, and pre-application (all p = ≤ 0.001), and post-application (p = 0.002).
  o MMD-PP at pre-removal (p = 0.035), and post-application (p = 0.009).
  o Control at pre-removal (p = 0.034).

• VAS scores lowered in MMD-PP and MMD-D more than control at post-removal (both p = < 0.001) and post-application (both p = < 0.001).

Pain & distress

• FLACC scores sig. differed at pre-removal of 2nd & 3rd COD (p = ≤ 0.001); post-removal at 1st (p = 0.003), 2nd & 3rd CODs (p ≤ 0.001); pre-application of 1st (p = 0.010), 2nd & 3rd CODs (p ≤ 0.001); and post-application all 3 CODs (p ≤ 0.001).

• FLACC scores lowered across 3 COD’s in:
  o MMD-D at post-removal (p = 0.008), pre-application (p = 0.047), and post-application (p = 0.018).
  o Control at pre-removal (p = ≤ 0.001).

• FLACC scores lowered in:
  o MMD-PP more than control at post-removal (p = ≤ 0.050) and post-application (p = < 0.001); and VGD at post-removal (p = ≤ 0.050) and post-application (p = ≤ 0.001).
  o MMD-D more than control at
post-removal (p = < 0.01), and post-application (p = < 0.001); and VGD at post-removal (p = ≤ 0.01) and post-application (p = < 0.001).

Millerb [47] Pain • FACES scores lower in MMD than control at pre-removal (p = 0.004); post-removal, pre & post-application (all p = < 0.001), 30% reduction.
• VAS scores lower in MMD than control at pre-removal (p = 0.018), post-removal (p = 0.010), pre-application (p = 0.001), post-application (p = < 0.001), 30% reduction.
• MMD combined sig. lowered pre-removal FACES (p = 0.009) and VAS scores (p = 0.035) compared to MMD-D.
• HR lowered in MMD at removal and application (both p = 0.040).

Pain & distress • FLACC scores lower in MMD than control at post-removal (p = < 0.001), pre-application (p = 0.021), post-application (p = < 0.001), 50% reduction at removal.
• MMD combined borderline less effective than MMD-D in reducing post-removal FLACC scores (p = 0.050).

Jefts [49] Pain • APPT-WGRS pre-procedure scores highest in VR, then SC and PD (p = 0.041).
• APPT-WGRS procedure scores lower in VR than PD (MD = 23.7 mm [CI:2.4, 45.0] p = 0.029), and SC (MD = 9.7 mm [CI:-9.5, 28.9] not sig. p = 0.320).
• Male patients reported less procedural pain (p = < .001).

Anxiety • Intervention group did not affect state (p = 0.060) or trait anxiety (p = 0.710).

Kipping [50] Pain • Intervention group did not affect patient VAS scores at dressing removal (p = 0.160) or application (p = 0.400).
• Intervention group did not affect caregiver VAS scores at dressing removal (p = 0.710) or application (p = 0.750).

Pain & distress • FLACC scores lower in VR (M = 2.9, SD = 2.4) than control (M = 4.7, SD = 2.5) at dressing removal (p = 0.020), but not application (p = 0.230).

Van der Heijden [51] Pain • Intervention group did not affect COMFORT-B scores before or after intervention (SMD = 0.04 [CI:-0.30, 0.38] p = 0.990).
• FACES scores lower in MT than SC (p = 0.050); N = MT13, SC5.

Distress • Intervention group did not affect OSBD-r scores before or after intervention (SMD = 0.11 [CI:-0.23, 0.45] p = 0.530).
| Intervention | Pain | Distress | Anxiety | Trauma | Pain & distress | Anxiety | Trauma | Pain & anxiety |
|--------------|------|----------|---------|--------|----------------|---------|--------|--------------|
| Whitehead-Pleaux [52] | Intervention group did not affect FPS-R scores (p = 0.200). | Intervention group did not affect FPS-R scores (p = 0.181), or after procedure (p = 0.345). | HR MD from before to after procedure greatest in control (p = 0.003) | NAPI scores higher in MT than control during procedure (p = .020). | Fear Thermometer scores higher in MT than control before (p = 0.043), and during procedures (p = 0.002), but not after (p = 0.228). | FLACC scores not reported. | CPSS impairment severity scores lower in hypnotherapy than SC (MD = 0.46 [CI: -0.01, 0.92] p = 0.050). | CPSS impairment severity scores lower in hypnotherapy than SC (MD = 0.46 [CI: -0.01, 0.92] p = 0.050). | CPSS impairment severity scores lower in hypnotherapy than SC (MD = 0.46 [CI: -0.01, 0.92] p = 0.050). |
| Chester [47] | Intervention group did not affect overall FPS-R scores before, during or after any procedure (p = > 0.100). | FPS-R scores lower in in patients < 8 years at 3rd COD (MD = 4.71 [CI: 0.33, 9.09] p = 0.04); relevant sample 3 per group. | HR lower in hypnotherapy than SC at pre-removal (MD=-15.20 [CI:-27.20, -3.20] p = 0.010) and post-application of 3rd COD (MD=-15.49 [CI:-28.25, -2.53] p = 0.020). | Patients > 8 years VAS-A scores lower in hypnotherapy than SC at pre-removal of 2nd COD (MD=-0.80 [CI:-1.50, -0.10] p = 0.030). | VAS-A scores for patients <8 years lower in hypnotherapy than SC at pre-removal of 2nd (MD=-1.37 [CI:-2.57, -0.16] p = 0.030) and 3rd CODs (MD=-2.07 [CI:-3.64, -0.49] p = 0.010). | Patient CPSS impairment severity scores lower in hypnotherapy than SC (MD = 0.75 [CI:0.05, 1.45] p = 0.040). | Patient CPSS impairment severity scores lower in hypnotherapy than SC (MD = 0.75 [CI:0.05, 1.45] p = 0.040). | Patient CPSS impairment severity scores lower in hypnotherapy than SC (MD = 0.75 [CI:0.05, 1.45] p = 0.040). |
| Hypnosis & guided imagery | Intervention group did not affect overall FPS-R scores before, during or after any procedure (p = > 0.100). | FPS-R scores lower in in patients < 8 years at 3rd COD (MD = 4.71 [CI: 0.33, 9.09] p = 0.04); relevant sample 3 per group. | HR lower in hypnotherapy than SC at pre-removal (MD=-15.20 [CI:-27.20, -3.20] p = 0.010) and post-application of 3rd COD (MD=-15.49 [CI:-28.25, -2.53] p = 0.020). | Patients > 8 years VAS-A scores lower in hypnotherapy than SC at pre-removal of 2nd COD (MD=-0.80 [CI:-1.50, -0.10] p = 0.030). | VAS-A scores for patients <8 years lower in hypnotherapy than SC at pre-removal of 2nd (MD=-1.37 [CI:-2.57, -0.16] p = 0.030) and 3rd CODs (MD=-2.07 [CI:-3.64, -0.49] p = 0.010). | Patient CPSS impairment severity scores lower in hypnotherapy than SC (MD = 0.75 [CI:0.05, 1.45] p = 0.040). | Patient CPSS impairment severity scores lower in hypnotherapy than SC (MD = 0.75 [CI:0.05, 1.45] p = 0.040). | Patient CPSS impairment severity scores lower in hypnotherapy than SC (MD = 0.75 [CI:0.05, 1.45] p = 0.040). |
| Foertsch [59] | Intervention group did not affect OSBD scores between groups (F_{1,9}=0.18, p = > 0.500), or across 4 CODs (exact | FACES and VAS scores not analysed due to patient difficulty in comprehending tools. | | | | | | |

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- Cry behaviors correlated with verbal resistance at 2nd (r[22] = 0.77, p = < 0.001), 3rd (r[22] = 0.56, p = < 0.050), and 4th CODs (r[22] = 0.49, p = < 0.050); with emotional support at 1st (r[23] = 0.58, p = < 0.050), and 2nd (r[22] = 0.88, p = < 0.010); and with verbal pain at 1st (r[23] = 0.52, p = < 0.050), and 2nd CODs (r[22] = 0.83, p = < 0.010).
- Female patients displayed higher verbal resistance at baseline (t[21]=-2.40, p = 0.020); and cry behaviors at 2nd -4th COD (t[20]=-2.26, p = 0.030).

**Therapeutic approaches**

**Blakeney [45]**  
**Anxiety/ distress**  
- CBCL anxious and depressed scores sig. lowered from pre-intervention to 1 year post-intervention in intervention group (t=-2.5, p = .017) and control (t=-2.4, p = .026); however not between groups (p = > 0.300).

**Elliott [58]**  
**Pain & distress**  
- Group comparisons not possible.
- BTDS scores reduced in Intervention group by 25–52% (mean = 36.7%) from baseline to post-intervention.
- BTDS scores consistently increased for intervention group in therapist absence.
- Patient’s preferred: relaxation, emotive imagery, distraction, imagery of pleasant scenery, and earning tangible reinforcement for coping techniques.

**Hyland [57]**  
**Pain**  
- CLT group received fewer additional analgesic medication during procedure than SC (n = 6, 12% vs n = 9, 18%).
- Average CHEOPS scores lower in CLT (Mdn = 5.3, IQR: 4.5–6.7) than SC (Mdn = 6.0, IQR: 5.4–7.6), (CI: 0.1, 1.2, p = 0.020).
- Nursing staff observed higher pre-procedure pain in CLT than SC (Mdn = 1.0, IQR: 0.0–2.0 vs. Mdn = 0.5, IQR: 0.0–1.0).
- Intervention group did not affect nursing staff observations of procedural pain (Mdn = 2.0 for both groups).
- FACES scores not reported

**Pain & anxiety**  
- CLT caregivers observed higher patient pre-procedure pain than SC (Mdn = 3.5, IQR: 0.0–4.0 vs. Mdn = 3.0, IQR: 0.0–5.0).
- CLT caregivers observed lower patient procedural pain than SC (Mdn = 2.0, IQR: 0.0–4.0 vs. Mdn = 3.0, IQR: 1.0–7.0).
- Intervention group had no affect caregiver observations of patient pre-procedure anxiety (Mdn = 2.0, IQR: 1.0–5.0 vs. Mdn = 2.0, IQR: 0.0–5.0).
- CLT caregivers observed less patient procedural anxiety than
### Anxiety

- Intervention group did not affect average CFS scores (CI: 0-0.2, \( p = 0.300 \)).
- CLT caregivers had higher anxiety than SC at pre-procedure (Mdn = 7.0, IQR: 5.0-8.0 vs. Mdn = 6.0, IQR: 4.0-8.0), and during procedure (Mdn = 5.0, IQR: 1.0-7.0 vs. Mdn = 3.5, IQR: 2.0-7.5).
- Nursing staff observed higher patient pre-procedure anxiety in CLT than SC (Mdn = 2.0, IQR: 0.0-4.0 vs. Mdn = 1.5, IQR: 0.0-3.0).
- Intervention group did not affect nursing staff observations of patient procedural anxiety (Mdn = 2.0 for both groups).

### Sveen [61]

#### Posttraumatic stress

- IES-R scores lower in intervention than control at 6 weeks post-randomization (\( \beta = -11.5 \ [SE: 3.88] \ p = 0.003 \)) and 3mths post-intervention (\( \beta = -7.89 \ [SE: 3.38] \ p = 0.020 \)).
- Intervention group did not affect IES-R scores at baseline or 12mths post-intervention.
- Intervention group did not affect caregivers PSI-SF or PSS scores at any time point during CODs.
- Caregivers perceived the intervention as informative and meaningful but time consuming.

### Preparation & ‘patient control’

#### Kavanagh [56]

#### Pain

- Intervention group required less analgesic pain medication in 1st 2 weeks of hospitalisation (\( p = < 0.010 \)).
- Intervention group received more analgesic medication between CODs (\( p = < 0.025 \)).
- Nurse reports not reported.

### Moore [46]

#### Pain

- Intervention group did not affect FPS scores from baseline, during, or post-procedure (\( p = 0.717 \)).

#### Pain & distress

- FLACC scores lower in MP than SC during CODs (0.5 vs 2 respectively), not sig. (\( p = 0.165 \)).

#### Anxiety

- Intervention group did not affect caregivers state anxiety from baseline to post-procedure (\( p = 0.421 \)).

### Quay [60]

#### Anxiety

- Caregivers able to rehearse treatments and share concerns about returning home.
- STAI-CH scores decreased in intervention caregivers of patients with > 30% TBSA burns at discharge (\( p = < 0.050 \) and
1st follow-up (p = < 0.050).
• Intervention group did not affect STAI-CH scores for any patient’s or caregiver of patients with < 30% TBSA burns at discharge or 1st follow-up visit.

APPT-WGRS: Adolescent Paediatric Pain Tool, word graphic rating scale, BTDS: Burn-Treatment Distress Scale, CBCL: Children’s Behavior Checklist, CEMS: Children’s Emotional Manifestation Scale, CFS: Children’s Fear Scale, CHEOPS: Children’s Hospital of Eastern Ontario Pain Scale, CI: 95% confidence interval, CLT: Child Life Therapy, COD: Change of dressing, COMFORT-B: COMFORT-behavioral scale, CPSS: Child PTSD Symptom Scale, CTSQ: Child Trauma Screening Questionnaire, FACES: Wong-Baker FACES pain rating scale, FLACC: Faces Legs Arms Cry Consolability, FPS: Faces Pain Scale, FPS-R: Faces Pain Scale-Revised, HR: Heart rate, IES-R: Impact of Event Scale-Revised, IQR: Interquartile range, M: Mean, MD: Mean difference, Mdn: Median, MMD: Multi-modal Device, MMD-D: Multi-modal Device Distraction, MMD-PP: Multi-modal Device Procedural Preparation, MP: Medical play, MT: Music therapy, NAPI: Nursing Assessment of Pain Index, NRS: Numeric Rating Scale, OSBD: Observational Scale of Behavioral Distress, OSBD-r: Observational Scale of Behavioral Distress-revised, PD: Passive distraction, PSI-SF: Parenting Stress Index Short Form, PSS: Perceived Stress Scale, SC: Standard care, SD: Standard deviation, SMD: Standardised mean difference, STAI-CH: Spielberger State-Trait Anxiety Inventory for Children, TBSA: Total Body Surface Area, VAS: Visual Analogue Scale, VAS-A: Visual Analog Scale-Anxiety, VGD: Video game distraction, VR: Virtual reality, YCPC: Young Child PTSD Checklist.

Table 2: Participant characteristics

Table 3: Study characteristics

Table 4: Key results of included studies.

Cultural components

None of the 17 included studies incorporated Australian Aboriginal cultural components as presented in the Dance of Life’s model of Australian Aboriginal people’s perspectives of health and wellbeing. Blakeney et al.’s social skills psychoeducation was the only study to include Native American children (n = 2, 3%); however, this study did not assess intervention effect by ethnicity or incorporate any cultural components. Therefore, its ability to meet the needs of First Nations people (respectfully used here in reference to Indigenous peoples globally) could not be ascertained [42]. Moore et al. acknowledged the lack of cultural diversity within their study on medical play; however, this was brief and specific to the inclusion of African American and Hispanic families [43]. Similarly, Chester et al.’s study on hypnotherapy briefly acknowledged the lack of representation of Aboriginal and/or Torres Strait Islander children; however, did not elaborate on any potential implications or outline the ethnic/cultural diversity of
included children [44].

Pain

Distraction based interventions had variable effects on patient pain. The Multi-Modal Device (MMD) and Ditto™ devices reduced patient self-reported pain [45–47], caregiver observations of pain [46, 47], and nurse observations of pain and distress [46, 47] when the procedural preparation story “Bobby get’s a burn” and interactive distraction games were provided together [45, 47], and separately [46]. The benefits of MMD distraction increased with repeated use and was borderline more effective in reducing nurse observations of pain and distress behaviours when used alone than in combination with procedural preparation [46, 47]. However, less interactive video game distraction was found to reduce self-reported pain and increase caregiver observations of pain over time compared to Multi-Modal Device – Procedure Preparation (MMD-PP), and Multi-Modal Device – Distraction (MMD-D) [46]; or reduce nurse observations of pain and distress but not affect self-reported pain compared to standard distraction [48].

Three-dimensional virtual reality increased self-reported pre-procedural pain and reduced self-reported procedural pain more effectively than passive distractions [49]. While off-the-shelf virtual reality increased nursing staff’s observations of pain and distress behaviours and had no effect on patients’ or caregivers’ reports of pain [50]. Music therapy reduced self-reported pain compared to standard care when provided immediately following COD [51, 52]; however, did not affect patients’ self-reported pain when provided during COD [51, 52]. Likewise, ‘medical play’ prior to COD commencement did not affect patients’ self-reported pain; however, reduced nursing staff observations of pain and distress behaviour at insignificant levels [43].

In contrast, the use of Child Life Therapy (CLT) reduced patients’ pain as observed
by caregivers and an independent assessor, and increased nursing staff’s observations of pre-procedural pain [53]. Similarly, hypnotherapy reduced pain levels at the third COD as self-reported by patients less than 8 years of age and caregivers [44]. Stress management during COD reduced self-reported pain and distress from baseline to post-intervention; however increased similar to the control group when the therapist was absent [54]. Further, patients that received increased ‘patient control’ and ‘predictability’ required less analgesic medication during the first two weeks of hospitalisations and more in between CODs [55].

Distress

The MMD and Ditto™ devices’ procedural preparation and distraction also reduced patient self-reported distress [45, 47], compared to standard care which increased self-reported distress [45]. Similarly, stress and pain management during COD reduced patient self-reported distress but only in the presence of a therapist [54]. In contrast, the use of familiar imagery did not reduce patients’ self-reported distress or investigators’ observations of distress behaviours [56]. Likewise, music therapy during and following COD did not reduce nurse observations of distress [51, 52] but rather increased observations of distress when performed during COD procedures [52].

Anxiety

Preparation for dressing procedures or hospital discharge reduced patients’ and caregivers’ anxiety [43, 55, 57]. Similarly, increased ‘predictability’ and ‘patient control’ during CODs reduced patient’s anxiety during the first two weeks of hospitalisation; however, not significantly [55]. Hospital discharge preparation reduced anxiety among caregivers of children with burns affecting ≥ 30% of their TBSA, but did not impact other caregivers’ or patients’ state anxiety during CODs
[57]. Blakeney et al. found that providing psychoeducational programs had similar effects as standard care in reducing patient’s self-reported anxiety/depression scores from pre-intervention to 1 year post-intervention [42]. Hypnotherapy lowered patients’ pre-removal anxiety as reported by patients aged older than 8 years at second COD, and caregivers for patients aged less than 8 years at second and third COD [44]. Child Life Therapy (CLT) reduced caregivers’ observations of patients’ procedural anxiety; however increased both caregiver anxiety at pre and during procedures, and nurse observations of anxiety at pre-procedure [53]. The Ditto™ device lowered self-reported anxiety at pre-removal for patient’s > 8 years [45]. Likewise, tablet based electronic game distraction reduced anxiety during and after COD procedures compared to standard distraction [48]. In contrast, virtual reality did not reduce patient anxiety during COD [45, 49, 52]; and music therapy during CODs increased self-reported anxiety before and during procedure [52].

Trauma

Few studies measured psychological trauma symptoms and those that did had mixed results [44, 45, 58]. Sveen et al.’s online self-help program reduced patient posttraumatic stress scores six weeks post-baseline and three months post-intervention; however symptoms returned to baseline levels 12 months post-intervention [58]. The online self-help program also did not reduce caregivers’ actual or perceived stress at any time [58]. Hypnotherapy significantly lowered patient self-reported trauma impairment severity compared to standard care; however, also increased caregiver’s observations of trauma symptoms in children aged < 7 years at three months post-injury [44]. The Ditto™ device did not reduce children’s stress or trauma symptoms one week following injury or three months following wound healing [45].
Risk of bias

Risk of bias assessment is presented in Fig. 2. Allocation blinding was often not possible for participants [46–49, 54], investigators [42, 46–48, 52–56], and outcome or data assessors; however, only two studies attempted to reduce such bias with counter-rationales [54, 55]. Non-randomised Control Trials (NRCT) assigned participants based on attendances’ month [55], weekday [43], or unspecified time [54]. Selective reporting potential biased the impression of efficacy in some studies. This was present in Hyland et al. who did not present Wong Baker FACES pain scores [53]; Brown et al. and Chester et al. who did not report nurses’ FLACC measures [44, 45]; and Kavanagh who did not present Nurse reports of patient pain[55]. Missing data was acknowledged by Foertsch et al. [56] and Hyland et al. [53] but was not adequately addressed. Elliott & Olson reported minimal results with no group comparison due to heterogeneity in patient age, time of data collection, and length of hospitalisation [54].

Intervention and control groups differed at baseline. Blakeney et al.’s intervention group contained all (n = 2) Native American participants and more females than control [42]. Elliott & Olson’s ‘baseline’ participants were younger (M = 6.75yrs) with greater burn TBSA (M = 32%) than intervention (Age M = 8.50yrs and TBSA% M = 21.50) [54]. Moore et al.’s groups significantly differed in burn location (p = 0.02) [43]. Kavanagh’s intervention group were all male, younger, with smaller burn TBSA, and shorter hospitalisation than control [55]. Whitehead-Pleaux et al.’s control had significantly lower baseline distress behaviours (p = 0.02) and self-reported anxiety (p = 0.04); however, anxiety analysis included unexplained extra participant [52].

Downs & Black quality assessment

Studies were classified as excellent [44, 47, 49, 50], good [43, 45, 46, 48, 51, 53,
56, 58], fair [42, 52, 57], or poor quality [54, 55]. Three studies had small samples of six or less per group [52, 54, 55]. Some studies did not make clear if participants were recruited from the same population [42, 54], or over the same time period [42, 48, 52-54]; and only four studies adequately described adverse events [44, 47, 49, 50]. Two studies did not outline differences in follow-up lengths or perform adequate adjustment [54, 57]. Studies did not outline losses to follow-up [49, 52, 55]; or describe the characteristics of the source population [42-58] or those identified as lost to follow-up [42, 45, 46, 51, 54, 56-58]. Distribution of confounders were not outlined [52, 54, 55, 57] or clear imbalances inadequately addressed [42, 43]. Two studies performed seemingly unplanned analysis by burn TBSA [57] and patient age [51]; and data dredging were unclear for two studies [42, 55]. Two studies did not present random variability estimates [54, 55], and three did not present actual p values [54, 56, 57]. Furthermore, Kavanagh did not clearly describe outcome measures and stated some participants did not receive both aspects of the intervention [55].

Discussion

This systematic review highlights a gap in understanding the effectiveness of psychosocial interventions for Aboriginal and Torres Strait Islander paediatric burn patients’ and their caregivers. Previous systematic reviews have not accounted for this specific group [59–63], assessed the effects of all psychosocial interventions available [59, 61], or considered psychological trauma as a primary outcome [59–62]. To the authors’ knowledge, this systematic review is the first to assess the effectiveness of psychosocial interventions in reducing pain and/or anxiety, distress, and trauma symptoms among paediatric burn patients and their caregivers as well
as their relevance to First Nations peoples.

The appropriateness of the included interventions for Aboriginal and Torres Strait Islander people could not be determined due to the lack of cultural components and First Nations participants. The limited representation of First Nations people is disconcerting given all but one study [58] were conducted in countries with strong First Nations presence; including the United States of America (n = 9, 53%), Australia (n = 6, 35%), and South Africa (n = 1, 7%). The circumstance surrounding the omittance of First Nations people from most studies is unclear; however, some potential reasons may include inaccurate ethnicity records as exemplified by the categorisation of ‘lighter skin’ vs ‘darker skin’ by Brown et al. [45], or potential bias in intervention design appeal or accessibility for First Nations people.

Similarly, only two studies provided interventions to caregivers and focused on their symptoms [57, 58]. The limited focus on caregivers is also concerning given the strong evidence of psychological implications of paediatric burn injuries on caregivers [20, 23, 64]. Furthermore, sufficient psychosocial support, education and involvement of caregivers in CODs can build caregivers’ competency and aid with their coping during their child’s burn treatment [65, 66].

The relatively small number of included studies (n = 17) demonstrates the limited scope of formally assessed psychosocial interventions with clear comparison groups. We acknowledge the difficulty of applying rigour and standardisation to psychosocial interventions due to their need for flexibility and adaptability; attributes often key to their responsiveness to individual needs. Standardisation is further compounded by the multi-factorial nature of ‘standard’ burns care, resulting in varied COD approaches [45], length of time from pain medication to dressing removal [43], days of hospitalisation [57], informal procedural preparation [46], and
'standard' distraction during COD [44]. However, some studies lacked standardisation of elements outside of ‘standard care’ that were provided to controls including additional investigator verbal support [56] and active distraction during CODs [55].

The generalisability of included studies was difficult to determine due to poor reporting of source population [42-58], and participants’ characteristics [54]. Small sample sizes of ≤ 20 per group impeded generalisability and power of ten studies [43, 46-50, 52, 54-56]. Further, some studies excluded families involved with child protection services or prior Suspected Child Abuse and Neglect reports [44, 45, 50]; patients diagnosed with an impairment or Autism Spectrum Disorder [45]; and patients receiving initial CODs in theatre [44], requiring skin grafts or other diagnosed medical conditions [45]. Other studies restricted inclusion by verbal communicability, inadvertently limiting representation of younger children [52] and non-English speaking families [45, 50]. And other studies demonstrated bias towards families with higher education [43] [58] and ‘socio-economic status’ [45], or with married/cohabitating caregivers and low family conflicts or symptoms of PTSD [58]. In contrast, Kavanagh’s intervention participants all reported a history of ‘family disorganisation’ or psychopathological symptoms, confounding factors to intervention success [55]. And of particular concern, two included interventions were not tested among females due to gender imbalances between groups or lack of female participants [54, 55]. Potential gender differences in intervention experience and primary outcomes was not considered by any of the included studies.

This review was limited to studies available in English and with Randomised Control Trial (RCT) and NRCT designs, resulting in the exclusion of seven otherwise relevant studies. These excluded studies were generally reflective of this review’s findings;
however, three reported on interventions not captured here [67–69]. One study found art therapy effective in allowing paediatric burn patient’s to express their traumas [69], another found art and play therapy combined reduced patient anxiety [67], and the other found group therapy reduced caregiver anxiety [68]. It is highly recommended that the results of these excluded studies be considered when developing future interventions.

Authors of the included studies often criticised quantitative measures for being relatively subjective [51, 53] and heavily reliant on self-reports and structured assessments [51]. Studies reported the Fear Thermometer [52] and Wong-Baker FACES scale [56] to be confusing and difficult for young children to understand, despite being validated for use among children. Likewise, FLACC behavioural pain scale was heavily criticised as inappropriate for children aged 9 + years whom are less likely to display observable signs of distress and pain [56], and its inability to capture pain and distress differences between groups [43]. Similarly, the Achenbach Child Behavioral Checklist [42], State-Trait Anxiety Inventory [57], and anxiety tool used by Kavanagh [55] were reportedly too general to capture paediatric burn patients’ anxiety during COD.

Despite these challenges, the included interventions demonstrated that procedural distraction via hand-held devices are effective in reducing patient’s pain [45–48]; however, less effective when only games were available [48] opposed to procedural preparation stories as offered by the Ditto™ [45] and MMD [46, 47]. This indicates that procedural information provided at any time during procedures can reduce patient pain. This is supported by Damanhuri et al.’s finding that active distraction incorporating additional COD information and encouragement was far more effective than passive distraction such as music therapy or games [14]. However, this review
also found that distraction techniques did not reduce paediatric patient’s anxiety [45, 49, 52], or trauma symptoms [45] regardless of incorporation of procedural preparation. In contrast, therapeutic approaches were effective in reducing psychological morbidities among patients and caregivers. In particular, CLT reduced caregiver and independent assessor’s observations of patient pain, and caregiver’s observations of patient anxiety [53]. An online self-help program was the only intervention found to effectively reduce patient trauma symptoms [58]. Similarly, incorporating stress and pain management into CODs reduced patient distress but was not sustained in the therapist’s absence [54]. The results of the included studies should be interpreted with consideration of potential bias due to difficulties in intervention allocation blinding [42–58].

Conclusion

This review returned a limited number of interventions that effectively reduced paediatric burn patient and caregiver psychological morbidities. The scarcity of work on reducing psychological trauma symptoms is particularly disconcerting given the volume of work emphasising the highly traumatising nature of burn injuries for both patients and families [2–4]. This highlights a need for additional work to better support and prepare caregivers for their vital role in providing security and comfort to their children during procedures. Of main concern to this review, the well-documented overrepresentation of Aboriginal and Torres Strait Islander paediatric burn patients was not reflected in the included studies nor were their perspectives on health and wellbeing. This lack of representation highlights the urgency for psychosocial interventions to be developed in partnership with and assessed among Aboriginal and Torres Strait Islander families. Finally, it is suggested that the effects
of the included psychosocial interventions be further explored within broader healthcare settings and contexts; in particular, distraction featuring procedural information, CLT, stress and pain management, discharge preparation, and online self-help programs.

Abbreviations

ACT
Acceptance and Commitment Therapy

APPT-WGRS
Adolescent Paediatric Pain Tool, word graphic rating scale

BTDS
Burn-Treatment Distress Scale

C
Control

CBCL
Children’s Behavior Checklist

CBI
Children’s Behavior Inventory

CBT
Cognitive Behavioral Therapy

CEMS
Children’s Emotional Manifestation Scale

CFS
Children’s Fear Scale

CHEOPS
Children’s Hospital of Eastern Ontario Pain Scale

CI
95% confidence interval

CINAHL
Cumulative Index to Nursing and Allied Health Literature

CLT
Child Life Therapy
COD
Change of dressing
COMFORT-B
COMFORT-behavioral scale
CPSS
Child PTSD Symptom Scale
CTSQ
Child Trauma Screening Questionnaire
FACES
Wong-Baker FACES pain rating scale
FLACC
Faces Legs Arms Cry Consolability
FPS
Faces Pain Scale
FPS-R
Faces Pain Scale-Revised
HR
Heart rate
I
Intervention
IVR
Virtual reality intervention
IPD
Passive distraction intervention
IVGD
Video game distraction intervention
IMMD−D
Multi-modal Device-Distraction intervention
IMMD−PP
Multi-modal Device-Procedural Preparation intervention
IES-R
Impact of Event Scale-Revised
IQR
Interquartile range
M
Mean
MD
Mean difference
Mdn
Median
MMD
Multi-modal Device
MMD-D
Multi-modal Device Distraction
MMD-PP
Multi-modal Device Procedural Preparation
MP
Medical play
MT
Music therapy
NAPI
Nursing Assessment of Pain Index
NRCT
Non-randomised Control Trial
NRS
Numeric Rating Scale
OSBD
Observational Scale of Behavioral Distress
OSBD-r
Observational Scale of Behavioral Distress-revised
PD
Passive distraction
PP
Declarations
**Ethics approval and consent to participate**

Not applicable.

**Consent for publication**

Not applicable.

**Availability of data and material**

Not applicable.

**Competing interests**

The authors declare that they have no competing interests.

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**Authors’ contributions**

HW, BG, KH, and KC conceptualised the review. HW conducted all search strategies and hand screening of references. HW double screened all references for title and abstract; CR screened 10% of references for title and abstract. HW screened and extracted data for all full text references; BG and KH screened and extracted data...
for 50% each of full text references. HW performed the narrative data synthesis. HW led and BG, KH, KC, and RK contributed to the data interpretation. All authors contributed to the drafting and revision of the manuscript, approved the final version, and agree to be accountable for all aspects of the work.

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Figures
Figure 1

PRISMA flow chart of inclusion/exclusion rates
Figure 2
Risk of Bias
Supplementary Files

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