Measuring musculoskeletal pain in infants, children and adolescents

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ABSTRACT:
Accurate, reliable and timely assessment of pain is critical for effective management of musculoskeletal pain conditions. The assessment of pain in infants, children and adolescents with and without cognitive impairment can be particularly challenging for healthcare clinicians for a number of reasons including factors related to the consultation (e.g. heterogeneous patient population, time constraints), the healthcare clinician (e.g. awareness / knowledge of available pain scales), standardised assessment scales (e.g. availability, psychometric properties and application of each scale) the patient (e.g. developmental stage, ability to communicate) and the context in which the interaction takes place (e.g. familiarity with the setting, physiological and psychological state). As a result, pain is frequently not assessed or measured during the consultation, and in many instances, is underestimated and undertreated in this population. The purpose of this paper is to provide healthcare clinicians with an overview of scales that can be used to measure pain in infants, children and adolescents. Specifically, the paper reviews the various approaches to measure pain intensity, identifies factors that can influence the pain experience, expression and assessment in infants, children and adolescents, provides age appropriate suggestions for measuring pain intensity in patients with and without cognitive impairment, and identifies ways to assess the impact of pain using multidimensional pain scales.
Musculoskeletal (MSK) pain is a common experience for children and adolescents. As many as half of all children and adolescents will report experiencing MSK pain at least once a month and as many as a third will report persistent or recurrent MSK pain. Musculoskeletal pain is known to have a substantial impact on the everyday life of children and adolescents, for example, in those who report MSK pain; ~40% report interference with daily activities and sports participation, 20% report missing school/work, 20-30% take medication, and over half seek healthcare which is associated with significant healthcare costs especially for those who experience persistent pain. Critical to the effective management of MSK pain by healthcare clinicians is the accurate, reliable and timely assessment of pain and the routine implementation of age appropriate, valid and reliable measurement of pain is a pivotal component of evidence-based medicine. Specifically, the valid and reliable measurement of pain is helpful in understanding a person’s pain experience, can help identify appropriate treatment options, and assist in monitoring change in a person’s pain condition, minimising potential adverse physiological and psychological consequences of unrelieved or inadequately managed pain.

Fundamental differences exist between infants, children, adolescents and adult’s pain experience and expression which highlight the need to assess and interpret pain in a way that is specific to each age group. For example, while the definition of pain is universal “a distressing experience associated with actual or potential tissue damage with sensory, emotional, cognitive and social components”, the way in which these components interact with environmental, developmental, sociocultural and contextual factors means that the conceptualisation, understanding and communication of pain is distinctly different for infants, children and adolescents and adults. For example, pain vocabulary emerges from
months (e.g. ‘ouch’, ‘ow’, ‘hurt’,) and continues to develop until it can reliably be used from around 5 years of age. Similarly, it is from this time that a child begins to develop an understanding of the causes and consequence of pain and the ability to control the expression of pain. In the absence of intellectual or cognitive deficit, a child’s age serves as a reasonable and easily measured proxy for development, and should be used to guide the way in which pain is measured in children and adolescents.

The assessment and measurement of pain in infants, children and adolescents can be a challenge for healthcare clinicians. Reasons for this include factors related to the consultation (e.g. heterogeneous patient population, time constraints), the healthcare clinician (e.g. awareness / knowledge of available pain scales), standardised assessment scales (e.g. availability, psychometric properties and application of each scale) the patient (e.g. developmental stage, ability to communicate) and the context in which the interaction takes place (e.g. familiarity with the setting, physiological and psychological state). As a result pain is frequently not formally assessed or measured during the consultation with more informal questioning used e.g. questions such as ‘are you ok?’ or ‘feeling better?’ and the presence of pain validated though observation of behavioural cues such as crying or grimacing. Even in populations who are at a higher risk of experiencing MSK pain e.g. children with cerebral palsy there is data to suggest that pain is assessed using validated tools in less than 10% of encounters. The inconsistent assessment, measurement and documentation of pain means that in in many instances pain is underestimated and undertreated in this population. Reports of hospital audit data suggests that a third of children experience moderate to severe pain during their hospital admission and documentation of pain assessment is varied (occurring between 12-100% of the time) across hospital settings and between clinicians but all too frequently the assessment, measurement and documentation of pain does not meet hospital or professional guidelines. Encouragingly, clinical practice has been found to be
responsive to knowledge translation and implementation strategies aimed at improving the
assessment, measurement and documentation of pain in children and adolescents. Further work is needed to understand the frequency at which the assessment and
measurement of pain occurs in other healthcare settings such as primary care and
community facilities.

Pain is recognised as a core outcome domain by a number of national and international
initiatives such as the Pediatric Initiative on Methods, Measurement and Pain Assessment in
Clinical Trials Consensus Group (PedIMMPACT) and the Society of Pediatric Pain Assessment Task Force and NIH Tool Box. These initiatives promote the use of
evidence based measures of pain intensity and impact in clinical practice and research.
Recommendations are the results of formal, collaborative processes and methodologies that
combine empirical evidence, expert opinion and clinical utility. The purpose of this paper is to
provide healthcare clinicians with an overview of scales that can be used to measure
musculoskeletal pain in infants, children and adolescents in a way that is quick, accurate and
reliable. However, few pain scales have specifically been evaluated for this purpose and
suggestions are based on scales that have been evaluated to measure procedural (e.g.
immunisation pain) and non-specific pain (e.g. MSK pain) with the emphasis being on clinical
utility of scales. Using the social communication model of pain as a framework the paper
specifically considers factors that influence the pain experience and expression, reviews the
various approaches that can be used to assess and measure pain; provides age appropriate
suggestions for measuring pain intensity in patients with and without cognitive impairment,
and identifies ways to assess the impact of pain using multidimensional pain scales. This
paper incorporates and extends the work of the previous collaborations outlined above by
identifying and integrating evidence from more recent publications into the measurement of
pain in infants, children and adolescents. Scales designed for use in the intensive care
setting (e.g. to assess pain in the intubated patient) or those that have only been evaluated
to assess post-operative pain are not reported in this paper unless specifically stated. We also acknowledge that there are aspects of pain beyond pain intensity, such as affective and evaluative dimensions but we do not address them comprehensively in this article.

The social communication model of pain

It is often helpful to use a theoretical model as a framework for understanding the many aspects and processes involved in the experience of pain. This review has chosen the social communication model of pain (see Figure 1) as a framework from which to examine, understand and approach the assessment and management of pain in individuals of all ages and for those with cognitive impairments. This comprehensive model highlights that at each stage there are a number of factors (biological, psychological and social factors) related to the individual in pain and clinician that influence how pain is experienced, expressed and interpreted and the effectiveness of this communication. The model also considers the social context (interpersonal context) in which the communication of pain occurs, and importantly recognises that the communication of pain may differ whether it is self-initiated by the person in pain or elicited by an observer's question (or repeat questioning e.g. after treatment when a socially desirable response may be provided). This model places assessment and measurement at the heart of understanding a person's pain experience and highlights the importance of how that information is obtained. The remainder of this paper will consider the measurement and assessment of pain in children and adolescents with a particular focus on the social communication model as a guide.

As outlined in the above social communication model, a number of factors related to the individual in pain, clinician and context can influence each stage. In Figure 2 below we have outlined some of the key factors within these stages that are relevant to the assessment and measurement of pain in children, for example the cognitive ability of the child to understand concepts such as pain severity or intensity, the burden that a child may feel when
questioned by a clinician, the potential influences of parents who are present within the consultation.

Approaches to measuring pain intensity

There are three main approaches to measuring pain intensity: physiological (how the body reacts), observations of behaviour (how the child reacts) and self-report (what the child says). It is important to note that the choice of approach will depend on the age and abilities of the child and that the different approaches are not interchangeable and typically only correlate poor to moderately. Ideally, information from each approach is used simultaneously to provide a detailed understanding of the pain experience with consideration of both automatic processing and higher cognitive function (see Figure 1).

Physiological indicators (e.g. increased heart rate, blood pressure, sweating) are associated with generalised (non-specific) stress reaction and are more strongly associated with distress and anxiety than self-report pain measures. For this reason, physiological indicators should not be used in isolation to estimate presence, quality or intensity of pain. Further, these indicators habituate over time and are therefore not appropriate for use in acute pain that is continuous or in those with chronic pain.

Observational measures involve observing an individual’s non-verbal behaviour (e.g. crying, facial expression, torso and limb movements) and interactions (e.g. social, appetite). The behavioural response to pain is recognised to be more of an automatic and reflexive response to actual or potential tissue damage. Parents and carers can often provide specific and helpful information about typical and idiosyncratic pain-related behaviours that reflect different quality or intensity of pain in their child, this information can then be used inform the selection of appropriate pain management or prevention strategies. As cognitive skills and function increase with age as does one’s ability control (e.g. suppress, exaggerate or feign) behaviour and where possible observational measures should be used to complement
self-report measures of pain (see Figure 1). Observational measures are particularly useful for assessing pain in:

- Children aged less than 4 years who do not have the language skills necessary to communicate pain, or lack the comprehension necessary for self-report measures.
- Patients with cognitive or communication impairments e.g. cerebral palsy.
- Situations where valid self-report is not possible (e.g. extreme distress) or the credibility of the self-report is in doubt.53

**Self-report** is the most direct and reliable approach for measuring pain in those who can self-report.129, 139 The ability of a child to understand and report the presence and intensity of pain requires cognitive skills including receptive language and understanding, knowledge and memory of pain, executive function (e.g. cognitive flexibility, working memory), and the ability to understand and estimate magnitudes and symbolic processing.63, 150, 151, 161 These skills begin to emerge as early as 3 years of age and gradually develop to enable the accurate and reliable self-reporting of pain intensity by children aged 5 years (on average) or older. While screening tasks (e.g. counting, comprehension and seriation task) are available these are time consuming and do not predict a child’s ability to accurately and reliably self-report pain beyond age alone.154, 161 The association between a child’s age and cognitive skills highlights the need to measure pain using different pain scales in children of different ages e.g. using a more simplified scale (with fewer response options) for younger children.151, 161

**Proxy-report;** while input from parents / guardians has a place in the assessment of pain in children, clinicians should be mindful of over-reliance on this information. Numerous studies have shown discrepancies between parental and child report of pain. Typically, studies in the general population of healthy children show that parents under-report pain in their children.29, 66, 76 In contrast, studies in children with painful health conditions typically report better
concordance, although with a tendency for parents to over-estimate pain severity, compared to the child’s report.\textsuperscript{32, 148}

\textbf{AGE APPROPRIATE SCALES TO MEASURE PAIN INTENSITY}

Single-item scales of pain intensity are most commonly used to measure pain as they are quick, simple to administer and closely correlated with the impact of pain on the individual (e.g. activity limitations, health care seeking, medication use).\textsuperscript{140} Tables 2-4 outline available scales of pain intensity including a general description of the scale, age range, psychometric properties, and practical considerations for use. These tables synthesise evidence from several systematic reviews,\textsuperscript{23, 31, 34, 36, 40, 59, 74, 80, 129, 141, 153} practice guidelines\textsuperscript{110} and peer-reviewed articles. The evidence outlined in Tables 2-4 was used to identify an appropriate pain intensity scale for each age-group, the suggested scales are summarised in Table 5. The scales that have been suggested for use are based on authors’ judgements along with consideration of the psychometric properties of the scale, type of pain (e.g. procedural versus non-specific pain), population, and context in which the scale has been evaluated.

\textbf{Infant (3 years or younger)}

Observations of behaviour are most commonly used in this age-group, most typically: crying, facial expression, verbalisation, torso and leg movements. It is important to note that no scale has been comprehensively evaluated to assess pain in children aged 3 years or younger in primary care settings, or for children with chronic or persistent pain.\textsuperscript{34, 153} The majority of observational scales have been developed to measure post-operative pain in the hospital setting, but a number of these scales have since been used to assess brief pain associated with medical procedures e.g. venepuncture, immunisations. In the absence of more robust evidence, the scales that have been validated to assess procedural pain are
reported in Table 2. Scales that only evaluate post-operative pain or incorporate physiological measures (e.g. blood pressure, oxygen levels), have not been included as this information is not readily available or feasible for use by many healthcare clinicians.

**Suggested scale:** The Face, Legs, Activity, Cry and Consolability (FLACC) Scale.

The FLACC scale was originally designed and validated for use in infants and children aged 2 months to 7 years to measure post-operative pain. Since original development, the scale has been used to measure acute and procedural pain in emergency departments, immunisation centres, and various clinical settings (e.g. radiology, ambulatory, dental) and in research. The FLACC scale is suggested to measure pain in infants aged 3 years or younger on the basis that is has been validated to measure acute procedural pain in a variety of settings (e.g. outpatient paediatric clinic, emergency department, immunization clinic), and in the absence of a scale that has been more comprehensively evaluated.

**Description:** A score of 0, 1 or 2 is given for each of the five items. Descriptions of typical behaviours are provided for each item, e.g. Legs: “Normal or relaxed position” (score 0); “Uneasy, restless, tense” (score 1); “Kicking, or leg drawn up” (score 2). Item scores are summed to provide a total score from 0-10. Table 2 contains additional details about the FLACC, and other observational scales to measure pain intensity in children who are unable to self-report.

**Preschool child (3-5 years)**

Age is the strongest predictor of a child’s ability to understand and use self-report pain scales. It is noted however, that rate of development is varied. While preschool aged children (3-5 years) are generally less likely to be able to understand self-reported pain scales than older children, some will be able to. Experience of pain and prior use of a scale appears to influence a child’s ability to use a pain scale reliably, emphasising the need to
measure pain consistently. It is suggested that pain intensity is captured though self-report in children of this age (if deemed appropriate), supplemented by information from parents / guardians of the child, and observation of behaviour (FLACC scale).

Suggested scale: Pieces of hurt tool (as able), supplemented by parent / guardian report and observation.

The pieces of hurt tool was designed and validated for use in children aged 4 to 6 years to measure procedural pain at immunisation clinics. The scale has since been used to measure acute (e.g. post-operative), procedural, chronic and recurrent pain in hospitalised children as young as 3 years. The Pieces of hurt tool is the suggested pain scale for infants aged 3 to 5 years.

Description: The child is asked ‘Does it hurt?’, if the child says ‘no’ then zero is recorded. If the child responds ‘yes’ the child is presented with 4 tokens (e.g. poker chips) and it is explained that each token represents a ‘piece of hurt’ (one token is a little bit of hurt, two is a bit more, to four tokens the most hurt you could ever have). The child is then asked ‘how many pieces of hurt they have right now?’ The number of tokens is then recorded; additional details are reported in Table 3.

Child (6-11 years)

Face scales are consistently preferred by children over numerical, analogue, or word descriptor scales. Several versions are available that use either line drawings or photographs (e.g. Faces pain scale – Revised, Wong-Baker, Oucher). While the scales perform similarly they are not interchangeable due to the different anchors used by each of the scales, highlighting the importance of using the same scale consistently. The main limitation of face scales is that pain intensity, a sensory component of pain, is being
measured using faces that express the effective / emotional dimension of pain. The type of
face anchors used by scales (e.g. smiling vs. neutral) has been found to influence children’s
responses.  

Suggested scale: Faces Pain Scale-Revised (FPS-R) scale.

The FPS-R was adapted from the original Faces Pain Scale and validated in children 4 to
12 years undergoing a painful procedure (ear piercing), and in an inpatient clinical
population. The FPS-R is the suggested pain scale for children aged 6 to 11 years based
on considerable evidence in support of its reliability and validity in this age-group. The
primary strength of the FPS-R compared to other face scales are the gender neutral face
anchors that do not convey the emotional / affective dimension of pain (e.g. smiling,
crying). 

Description: Faces Pain Scale-Revised (FPS-R) is a set of 6 line-drawn faces with depictions
of increasing levels of pain from left to right. Children are asked to specify which face best
illustrates the amount of pain they are experiencing at that time. Each face is assigned an
increasing score from left to right, either 0-5, or 0-10 (increments of 2). Electronic versions of
the FPS-R have also been developed and validated e.g. the Sydney Animated Facial
Expressions (SAFE) scale and Painometer App, additional details are reported in
Table 3.

Adolescent (12-18 years)

Suggested scale: Verbal Numerical Rating Scale (NRS)-11.

The NRS-11 is one of the most commonly used scales to measure pain intensity in both
clinical and research settings, despite only recently undergoing appropriate psychometric
evaluation in children and adolescents. The scale has been validated to measure acute, procedural pain, as well as chronic pain in a wide range of settings including school children receiving immunisations, outpatient pain clinics, emergency departments. The NRS-11 is suggested to measure acute pain in children aged 12 to 18 years due to its simplicity, validity, reliability, and brevity as a pain assessment tool.

Description: The child is asked “On a scale of 0 to 10, where 0 is no pain and 10 is the worst possible pain, tell me what number best represents your pain?” The individual responds with a number that reflects their pain. The NRS-11 has also been adapted and validated for use to capture pain intensity for both acute and chronic conditions by SMS and online, additional details are reported in Table 3.

Children and adolescents with cognitive impairment

Children and adolescents with cognitive impairments (e.g. cerebral palsy) experience more significant and frequent pain than children without cognitive impairment, and are less likely to receive adequate pain management, indicating the need for specific and appropriate pain assessment measures. Pain behaviours displayed by children with cognitive impairment are not always comparable to those of children without cognitive impairment, although pain expression has been found to be consistent, observable and reflective of the presence and severity of pain. Thus pain measurement tools should be adaptable to reflect individual pain related behaviours, but ideally also contain standardised items that enable its use in any setting. Very few scales are available to assess pain in children with cognitive impairment and as seen in Table 4, these have only been tested in post-operative, residential care, or school settings. No scale has been tested to measure brief procedural, chronic, or recurrent pain in children and adolescents with cognitive impairment.
Suggested scale: Revised-Face, Legs, Activity, Cry and Consolability (r-FLACC) Scale

The r-FLACC was adapted from the FLACC scale\(^9\) for use in children and adolescents with cognitive impairment.\(^8\) The r-FLACC is suggested for children and adolescents with cognitive impairment, based on evidence demonstrating valid and reliable measurement of post-operative pain in a hospital setting, the ability of the scale to be individualised, and evidence of its clinical utility. This suggestion is made in the absence of any other more comprehensively evaluated scale.

**Description:** The r-FLACC is essentially the same as the FLACC scale but also enables behaviours that are unique to the individual to be described for each of the five behaviours (face, movement of the body and legs, cry and consolability). Identifying pain behaviours that are unique to the individual requires input from a family member or carer.

Assessing the broader impact of pain using multidimensional pain scales

While this paper has focused predominantly on the measurement of pain intensity it is acknowledged that pain experience is complex and contains other dimensions including affective (e.g. unpleasantness) and evaluative dimensions (e.g. appraisal of pain) as well as the impact pain has on everyday life including an individual’s physical, social, emotional functioning and ability to fulfil their ‘role’. The social communication model presented at the start of this paper can still be used to conceptualise the communication of these other dimensions of pain albeit using broader, multidimensional pain scales. Multidimensional pain scales are particularly useful for assessing recurrent and chronic pain as they can capture various dimensions of the pain experience (including pain duration, frequency, location, nature of pain, aggravating and easing factors) and how pain impacts everyday life e.g. interference with daily activity, or participation in school and sport. This fills a well-accepted need to differentiate between low intensity, transient pain and more persistent pain that has substantial impact on life.\(^9\) Multidimensional scales differ with respect to the factors
assessed e.g. psychosocial factors, situational factors, nature of disability and period of time. Some of the most commonly-used multidimensional pain scales for use in children and adolescents with chronic or recurring pain are outlined in Table 5. No specific scale has been suggested, as the choice will depend on the purpose of measurement and the health condition being measured e.g. region- or condition-specific scales. Common to many of the scales is identification of impact on school absenteeism, interference with sports participation, interference with activities of daily living, medication use, and healthcare utilisation. These being acknowledged important indicators of pain impact in paediatric populations.99

Another important dimension of pain assessment is fluctuation over time. In addition to the scales outlined in Table 5, pain diaries are often used to capture information about pain (e.g. pain intensity, frequency and location), and its effect on behaviour over time. The information collected in a pain diary may provide a more accurate and reliable measure of pain if completed on a regular basis, by minimising recall bias. Recent advances in information and communication technology (e.g. Internet, smartphones) have permitted the development of electronic methods such as e-diaries. Advantages of this electronic approach over traditional paper-based techniques include: minimizing errors in data transfer and transcription, ability to capture time-stamped data, ease of data sharing, increased compliance, and heightened patient satisfaction78. Recently, Stinson and colleagues have validated a number of web-based and smart-phone-based multidimensional electronic pain assessment tools (e-Ouch, Standardized Universal Pain Evaluation for Rheumatology providers (SUPER-KIDZ) and Pain-QuILT™ (freely available).77, 78, 132 These tools can be used in a variety of clinical settings to monitor MSK in children in real time.77, 78

GOOD PRACTICE POINTS
Pain in children has been inadequately assessed, underestimated, and undertreated for many years. This situation can only be rectified through the improved communication of pain between the individual in pain and the observer at every relevant clinical contact, until this becomes a part of routine care. Central to the social communication model of pain framework are the various intra- and interpersonal factors that may influence the expression, assessment and management of pain. Consideration of these factors and the context in which the communication occurs is vital for effective, accurate and reliable communication of pain, early detection of pain and timely management (including reassurance and advice). Most importantly the improved communication of pain has been found to improve patient outcomes (e.g. reduction in mean pain scores, improved satisfaction) and reduce in health care costs (e.g. reduction in length of stay).

An individual’s pain experience and expression is determined by a range of biopsychosocial factors that are specific to their developmental stage. The clarity of the pain expression can be optimised through the consistent use of valid and reliable pain scales that are age appropriate and meet the cognitive and communication capabilities of the individual, where available. For infants this means using observational scales of behaviour, for children integrating both observational and self-report scales until later childhood and teenage years where self-report can primarily be used. In all cases, patient self-report of pain should be considered and interpreted alongside knowledge of the context, and supplemented with information gained from observation of their behavior and input from parents / guardians and carers, when appropriate. A child’s ability to use a pain scale accurately and reliably increases with their familiarity of the scale, highlighting the need to introduce and educate children on the use of pain scales early in their life course. By providing children with the vocabulary and skills necessary to express their pain the clarity of the pain expression can be improved and potentially reduce errors in the interpretation by the observer.
Effective pain assessment and management by clinicians can be enhanced through the consistent use of standardised pain scales within and across health care settings and the accurate and timely documentation of assessment findings. Advantages of this approach include improved continuity of care for the individual and the ability to generate consistent data for longitudinal comparison of pain overtime. Evidence from studies on global perceived effect scales show that patient reported outcomes taken overtime provide a more accurate understanding of changes in a person’s health status compared to recall of improvement / deterioration. The minimally clinically important change has been determined for a number of scales (reported in Table 3 where available) which can assist clinician’s determine theeffectiveness of an intervention and provide insight into the meaningfulness of the change for the individual. Areas in need of further development however is the assessment and measurement of pain that is the result of other mechanisms e.g. neuropathic and central pain, for which there are currently no scales that have been validated for use by children and adolescents. Undoubtedly preferable to the timely assessment and management of pain are practices that can help minimise or prevent the experience of pain. In many instances pain can be anticipated (e.g. procedural, vaccinations, post-operative pain), and proactively managed by clinicians and parents / carers using both pharmacological (e.g. topical analgesics) and non-pharmacological interventions (e.g. distraction). Initiatives such as the ‘It doesn’t have to hurt’ online video is one such example of providing clinicians, parents / guardians with effective, evidence based information that can positively influence a child’s experience of painful situations.

DISCUSSION

To date little research has been done to evaluate the use of measures designed for assessment of pain in infants, children and adolescents outside the hospital setting. This
constitutes an important evidence gap given that primary care, community outpatient and rehabilitation clinics have frequent contact with infants, children and adolescents with pain. While further work is needed, this paper provides clinicians with a pragmatic, evidence-based overview of scales that can be used to measure pain intensity in infants, children, and adolescents, with and without cognitive impairment, and assess the impact of pain using multi-dimensional pain scales.
<table>
<thead>
<tr>
<th>Term</th>
<th>Operational definition</th>
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<tbody>
<tr>
<td><strong>Reliability</strong></td>
<td>The reproducibility of a measure over different occasions and is concerned with minimizing sources of random error so that measures are reproducible. In general, acceptable reliability coefficients for research and clinical purposes are ≥0.7 and ≥0.9 respectively.</td>
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<td>Test-retest</td>
<td>The agreement between observations with the same individuals on at least two occasions.</td>
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<td>Inter-rater reliability</td>
<td>The agreement between different raters/observers of an observational measure of pain.</td>
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<td><strong>Validity</strong></td>
<td>Used to assess whether the scale is measuring what it is intending to measure.</td>
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<td>Face validity</td>
<td>Whether the pain scale includes appropriate items that appear to measure what it is proposing to measure.</td>
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<td>Content validity</td>
<td>The assessment of whether the items in the pain measure include the appropriate information and content.</td>
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<tr>
<td>Criterion</td>
<td>Includes concurrent validity and predictive validity. In concurrent validity, a new pain measure is correlated with a gold standard measure which is administered at the same time. In general, correlations between the new measure and the gold standard should be at least r ≥ 0.3–0.5. The magnitude of the coefficients are hypothesis dependent but should not be too high as to make the new measure redundant. In predictive validity, the correlation of the measure to the criterion variable is determined later.</td>
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<tr>
<td><strong>Construct</strong></td>
<td>Determines the validity of abstract variables that cannot be directly observed, such as pain. These constructs are assessed by their relationships with other variables.</td>
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<td>Convergent validity</td>
<td>Evaluates how well items on a pain scale correlate with other measures of the same construct or related variables. In general, correlations between the measure and another measure of the same construct should be r ≥ 0.3–0.5; however, the magnitude of the coefficients are hypothesis dependent.</td>
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<tr>
<td>Discriminant validity</td>
<td>Evaluates how items on a pain scale correlate with other measures that are unrelated. In general, correlations between the measure and another unrelated measure should be r &lt; .3; however, the magnitude of the coefficients are hypothesis dependent.</td>
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<td><strong>Responsiveness</strong></td>
<td>Measures whether the measure is able to identify changes in pain over time where change is expected (e.g. after analgesia). (COSMIN taxonomy)</td>
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<td>Interpretability</td>
<td>The meaningfulness of the scores obtained from a pain measure.</td>
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<td>Feasibility</td>
<td>How easily a pain measure can be scored and interpreted.</td>
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<td>Minimally clinically important change</td>
<td>The smallest difference in score in the domain of interest which patients perceive as beneficial and which would mandate, in the absence of troublesome side effects and excessive cost, a change in the patient’s management.</td>
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<tr>
<td>Pain scales</td>
<td>Description</td>
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<tr>
<td>Face, Arms, Legs, Activity Cry and Consolability Scale (FLACC)</td>
<td>5, 34, 46, 90, 91, 135</td>
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<tr>
<td>Child facial coding system (CFCS)</td>
<td>17, 31, 52</td>
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<tr>
<td>Children’s Hospital Of Eastern Ontario Pain Scale (CHEOPS)</td>
<td>34, 35, 39</td>
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<tr>
<td>Modified behaviour pain scale (MBPS)</td>
<td>35, 136</td>
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**Table 2: Observational scales used to measure pain in infants and children 3 years or younger**
movement scored on a 0-3 scale and cry 0-4 scale, range 0-10

<table>
<thead>
<tr>
<th>Neonatal Infant Pain Scale (NIPS)</th>
<th>6 item scale (facial expression, breathing patterns, cry, arm movement, arousal, leg movement). All items 0 (absent/relaxed) to 1 (change from normal) except cry; 0 to 2</th>
<th>I: 0-2 months</th>
<th>✓</th>
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<td>- Scored on a commonly understood 0-10 scale</td>
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<td></td>
<td>- Validated for acute procedural pain e.g. immunisations</td>
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<td>- Multidimensional scale taking into consideration factors such as arousal</td>
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<td></td>
<td>- Evaluated for infants only, limited applicability</td>
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</tbody>
</table>

✓ Evidence available; – no supporting evidence
I: intended age range; S: studied age range.
<table>
<thead>
<tr>
<th>Pain scale</th>
<th>Description</th>
<th>Age range</th>
<th>Reliability</th>
<th>Validity</th>
<th>Responsiveness</th>
<th>Practical considerations</th>
<th>Strengths</th>
<th>Limitations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pieces of hurt tool</td>
<td>The child is asked “Did it hurt?”; if the child responds yes they are given four chips - ‘pieces of hurt’. The child is told; ‘These are pieces of hurt- one chip is a little bit of hurt, and four chips are the most hurt you could ever have. Do you have one, two, three or four pieces of hurt?’ The number of chips is recorded.</td>
<td>I: 4-7 years S: 3-18 years</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>- Tangible quantity of pain is easier for younger children to understand - Developmentally appropriate, yes/no followed by limited response options - Developed in English. Validated in Jordanian and Thai - Validated recurrent or persistent pain - Can use any items</td>
<td>- Little testing in younger children - Potential bias toward higher pain scores especially in younger children - Requires the ability to count and estimate quantities using numbers - Infection risk, storage, and availability of tokens</td>
<td></td>
</tr>
<tr>
<td>Faces pain scale - revised</td>
<td>6 line drawn faces aligned horizontally from an expression of “no pain” (left) to “most pain possible” (right). The child points to the face that shows their pain. Standardised instructions are used. The original Faces Pain Scale had 7 faces scored on 0-6 scale. Revised to make compatible with other 0-10 scales.</td>
<td>I: 4-12 years S: 3-18 years</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>- Scored on a commonly understood 0-10 scale - Gender neutral drawings of faces - Developed in English. Translations: &gt; 35 languages, validated in French, Thai and Catalan. - Minimally clinically important change: 2/10 (1 face), or 25% - Electronic version available</td>
<td>- Reduced accuracy with decreasing age - Response bias in those under 5 years of age - Skewed pain intensity ratings towards ‘no pain’, may underestimate pain intensity</td>
<td></td>
</tr>
<tr>
<td>Verbal Numerical rating Scale-11</td>
<td>The child is asked “On a scale of 0 to 10, where 0 is no pain and 10 is the worst possible pain, tell me what</td>
<td>I: 8-18 years</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>- Scored on a commonly understood 0-10 scale</td>
<td>- Requires the ability to count, estimate quantities using numbers, recall pain</td>
<td>-</td>
</tr>
</tbody>
</table>
number best represents your pain?" The individual responds with a number that reflects their pain.

<table>
<thead>
<tr>
<th>Scale Type</th>
<th>Age</th>
<th>Description</th>
<th>优点</th>
<th>缺点</th>
</tr>
</thead>
<tbody>
<tr>
<td>Coloured analogue scale (CAS)</td>
<td>I: 5-17 years, S: 3-18 years</td>
<td>Visual analogue scale with a mechanical device with a plastic slider over a 143 mm long tetragon varying from narrow (10 mm) and white labelled 'no pain', to wide (30 mm) and dark red at the end labelled 'most pain'. Range 0-10.</td>
<td>- Adaptable e.g. usual, strongest, lowest pain, at rest / activity - Developed in English. Translations: Spanish, French - suitable for 8-20 years, physical disabilities (e.g. cerebral palsy, neuromuscular disease), chronic pain - electronic version available&lt;sup&gt;114, 115&lt;/sup&gt; - Minimally clinically important change 1/10</td>
<td>Requires users to have the CAS device available - Infection risk</td>
</tr>
<tr>
<td>OUCHER (NRS and photographic)</td>
<td>I: 3-7 years, S: 3-18 years</td>
<td>0-10 numerical rating scale aligned vertically next to 6 photographs ranging from &quot;no hurt at all&quot; (at the bottom - 0) to the &quot;biggest hurt you could ever have&quot; (at the top - 10). &lt;sup&gt;(Prior to 2009 scoring was 0-100) Boucher manual&lt;/sup&gt;</td>
<td>- Various versions (Caucasian, Hispanic, Asian, boy/girl) - Child can choose photographic or VAS - photographic scale (other face scales are line drawings) - Scored on a commonly understood 0-10 scale - Evaluated in specific populations e.g. sickle cell - Little testing in younger children - Printed OUCHER scale required - infection risk - Photographs resemble acute pain only - The NRS left of the photographs may be confusing for young children (3-7 years)&lt;sup&gt;104&lt;/sup&gt;</td>
<td>- Little testing in younger children</td>
</tr>
<tr>
<td>Visual Analogue Scale (VAS)</td>
<td>I: 2-17 years, S: 3-18 years</td>
<td>10cm vertical/horizontal line with anchors (e.g. 'No pain', 'Worst possible pain'). The child marks along the line to</td>
<td>- Evaluated in several populations e.g. sickle cell, juvenile chronic arthritis - Variability in line length, demarcations, orientation (and anchors e.g. worst pain, strongest pain)</td>
<td>- Variability in the time period (e.g. past week, current pain) and anchors used (e.g. worst possible pain, strongest pain) - Further testing required in younger children (6-8 years).</td>
</tr>
<tr>
<td>Scale Type</td>
<td>Description</td>
<td>Intended Age Range</td>
<td>Studied Age Range</td>
<td>Evidence Available</td>
</tr>
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</table>
| Wong-Baker Faces Pain Scale | 6 line drawn faces aligned horizontally from a smiling “no hurt” face (left) to crying “hurts worst” face (right), range 0-5 (or 0-10 if each face is 2 points). | I: 3-18 years | S: 9 months – 18 years | ✓ ✓ ✓ ✓ ✓ - ✓ ✓ ✓ ✓ ✓ ✓ | - Developed in English. Validated for use in English-speaking populations. Validated for use in Jordanian. - Scored on a commonly understood 0-10 scale that is suitable for 8-20 years with physical disabilities, specific conditions e.g. sickle cell. - Younger children tend to use the extremes and may provide higher pain intensity rating than other face scales. - May not perform as a continuous scale; pain ratings cluster around words which reduces sensitivity. |}
| Word descriptor scales / Word Graphic Scale | 5-6 words to describe pain from 'no pain' to 'worst pain'. A word is selected to best describe their pain. Each word has a number for scoring. Range from 0 (no pain) to 5 (worst pain). | I: 8-17 years | S: 3-18 years | ✓ - ✓ ✓ ✓ - ✓ - ✓ - ✓ | - Developed in English. Validated for use in English-speaking populations. - Incorporates into multidimensional scales e.g. Adolescent Pediatric Pain Tool. - Requires ability to comprehend pain intensity terms. - May not perform as a continuous scale; pain ratings cluster around words which reduces sensitivity. |

I: intended age range; S: studied age range. ✓ Evidence available; – no supporting evidence.
<table>
<thead>
<tr>
<th>Pain scale</th>
<th>Description</th>
<th>Age range</th>
<th>Reliability</th>
<th>Validity</th>
<th>Responsiveness</th>
<th>Feasibility</th>
<th>Interpretability</th>
<th>Training</th>
<th>User preference</th>
<th>Acceptable to carers</th>
<th>Strengths</th>
<th>Limitations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Revised Face, Arms, Legs, Activity Cry and Consolability Scale (rFLACC)</td>
<td>5 item scale measures facial expression, leg movements, activity, cry and consolability. Each item is scored 0-2, total range 0-10. Needs input from parent/guardian to identify ‘baseline’ behaviours. Includes open-ended descriptor for individual pain behaviours.</td>
<td>I: 4-19 years</td>
<td>✓ ✓</td>
<td>✓ ✓</td>
<td>✓ ✓</td>
<td>✓ ✓ ✓ ✓ ✓ ✓</td>
<td>✓ ✓ ✓ ✓ ✓</td>
<td>✓ ✓ ✓ ✓ ✓</td>
<td>✓ ✓ ✓ ✓ ✓</td>
<td>simple to use, score and interpret - Scored on a commonly understood 0-10 scale - Individualised by parent/carer - Developed in English. Validated in Danish</td>
<td>- evaluated in post-operative settings only - underlying motor impairments e.g. spasticity may affect assessment - Observation time not established; 5 min used in previous testing. - Ambiguity around amount and timing of some items e.g. 'jaw clench', consolability.</td>
<td></td>
</tr>
<tr>
<td>Individualised numerical rating scale (INRS)</td>
<td>Carers provide word descriptors to be used as anchors for their child’s pain behaviours from 0 (no pain) to 10 (worst possible pain).</td>
<td>I: 6 – 18 years</td>
<td>✓ ✓</td>
<td>✓ ✓</td>
<td>✓ ✓</td>
<td>✓ ✓ ✓ ✓ ✓ ✓</td>
<td>✓ ✓ ✓ ✓ ✓</td>
<td>✓ ✓ ✓ ✓ ✓</td>
<td>✓ ✓ ✓ ✓ ✓</td>
<td>- Uniquely created for each individual according using information from carer - 1 minute observation time</td>
<td>- Minimal psychometric evaluation, not assessed in procedures e.g. injections. - Requires parent-nurse collaboration</td>
<td></td>
</tr>
<tr>
<td>Non-communicating children pain checklist – revised (NCCPC-R)</td>
<td>30 items (6 sub-scales: vocal, social, facial, activity, body and limb, physiological). Frequency of each behaviour; 0=not at all to 3=very. Range 0-90. Post-operative version (NCCPC-PV - 27-items, total range 0-81), does not include eat/sleeping items.</td>
<td>I: 3-18 years</td>
<td>✓ -</td>
<td>- -</td>
<td>✓ ✓</td>
<td>✓ ✓ ✓ ✓ ✓ ✓</td>
<td>✓ ✓ ✓ ✓ ✓</td>
<td>✓ ✓ ✓ ✓ ✓</td>
<td>✓ ✓ ✓ ✓ ✓</td>
<td>- Intended for use by parents but can be used by anyone involved in child’s care. - Has been used in observational studies</td>
<td>- Psychometric properties not assessed in procedures e.g. injections. - 10 minute observation time before scoring - Validation study used 2 hour observation period.</td>
<td></td>
</tr>
<tr>
<td>Pediatric Pain Profile (PPP)</td>
<td>20 items rated on 4 point Likert scales (0=not at all to 3=a great deal). Used by an observer familiar with the</td>
<td>I: 1-18 years</td>
<td>✓ ✓ ✓ ✓</td>
<td>✓ ✓</td>
<td>✓ ✓</td>
<td>✓ ✓ ✓ ✓ ✓ ✓</td>
<td>✓ ✓ ✓ ✓ ✓</td>
<td>✓ ✓ ✓ ✓ ✓</td>
<td>✓ ✓ ✓ ✓ ✓</td>
<td>- Developed for use in residential (during normal morning routine in a home, residential care, residential</td>
<td>- Psychometric properties not assessed in procedures e.g. injections.</td>
<td></td>
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</tbody>
</table>
** All of these scales are observational scales which have been evaluated to assess post-operative pain. None of the above scales have been evaluated to assess procedural type pain.

I: intended age range; S: studied age range. ✓Evidence available; – no supporting evidence
## Table 5: Multidimensional pain scales that measure the impact of pain in infants, children and adolescents

<table>
<thead>
<tr>
<th>Pain scale</th>
<th>Description</th>
<th>Age range</th>
<th>Reliability</th>
<th>Validity</th>
<th>Practical considerations</th>
<th>Strengths</th>
<th>Limitations</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Adolescent pediatric pain tool (APPT)</strong> 44, 64, 117, 118</td>
<td>Pain intensity measured on 0-100mm VAS, body chart and 67-word descriptors to express sensory (37 words), evaluative (8), affective (11) and temporal (11) qualities 5 pain subscale scores: number of pain sites (from body chart), pain intensity score, number of temporal descriptors (%), total pain quality and temporal descriptors.</td>
<td>I: 8-18 years S: 8-18 years</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>- Self-report</td>
<td>- Evaluated as single components (e.g. body chart and word graphic scale has been evaluated in school, medical and surgical pain setting), but not entire scale - evaluation required in complex, recurrent, chronic pain states - few descriptors represent neuropathic pain - requires equipment and overlay to score - Children must read or understand English. Some children required assistance</td>
</tr>
<tr>
<td><strong>Bath Adolescent Pain Questionnaire (BAPQ) 41, 42, 155</strong></td>
<td>61 items in 7 domains: social functioning, physical functioning, physical functioning, depression, general anxiety, pain specific anxiety, family functioning and development. Each item rated on 5 point scale (never=0 to always=4) except the development subscale rated from 0 (“very behind”) to 4 (“very ahead”). Range 0-244</td>
<td>I: 11-18 years S: 10-18 years</td>
<td>✓</td>
<td>-</td>
<td>✓</td>
<td>- Initially developed and tested in outpatient Rheumatology and multidisciplinary pain clinic. - Used to assess chronic pain from perspective of adolescent or parent - Validated in secondary care e.g. Rheumatology / pain management clinics.</td>
<td>- Questionnaire length - Complicated scoring with reverse scoring for some items - Total sum score not clinically useful</td>
</tr>
<tr>
<td><strong>Childhood Health Assessment Questionnaire (CHAQ) 19, 70, 94, 111, 122</strong></td>
<td>Includes: Disability and Discomfort, in the last week. Disability incl. 30 items, eight subscales: dressing, grooming, arising, eating, walking, reaching, grip, activities. Each item scored 0=no difficulty to 3=unable to do it (or ‘not applicable’ if beyond</td>
<td>I: 1-19 years</td>
<td>✓ ✓</td>
<td>✓ ✓</td>
<td>✓ ✓ ✓</td>
<td>- evaluated in Juvenile Idiopathic Arthritis (JIA), cerebral palsy, inflammatory myopathies. - core outcome measure for clinical trials in JIA - Self-report and parent proxy versions</td>
<td>- relatively insensitive to important short term changes in children with JIA - evidence of a floor effect</td>
</tr>
</tbody>
</table>
| Test                                                                 | Description                                                                                                                                                                                                 | I: 8 – 18 years | Notes                                                                                                                                                                                                 | Validated in 32 countries. | Tested in specific populations. Requires further evaluation.
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<tbody>
<tr>
<td>Child Activity Limitations Interview (CALI / CALI-21) 100, 101</td>
<td>8 activities selected from list of 21 options found difficult or bothersome due to pain. Importance and difficulty over the last 4 weeks is rated on 5-point scale from 0=not important/difficult to 4=extremely important. Ratings are summed; total score 0-32. CALI-21: participants report on limitations for all 21 activities. Ratings summed; total score 0-84.</td>
<td>I: 8 – 18 years</td>
<td>Developed in healthy children and validated in chronic or recurrent pain e.g. headache, JIA, sickle cell. - Developed and tested in specific populations. Requires further evaluation. Validated in 32 countries.</td>
<td>- 10 mins to complete</td>
<td></td>
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</table>
| Functional disability Inventory (FDI) 30, 70, 157                   | 15 items assessing everyday activities in the past 2 week e.g. walking up stairs. Each item scored 0= no trouble to 4= Impossible. Scores summed; total score 0-60.                                                                 | I: 8 – 17 years | Evaluated in a variety of conditions e.g. abdominal pain, acute minor illnesses, back pain. Recommended by PedIMMPACT90 - Interview or self-report                                       | - Developed and validated in mixed population with chronic pain e.g. fibromyalgia, rheumatoid arthritis, headache - Self report and parent scale - Validated in German. Based on Multidimensional Pain Inventory | Requires further evaluation.
<p>| Pain experience Questionnaire (PEQ) 56                            | 15-item questionnaire, four subscales: pain severity, pain-related interference, social support and affective distress. Each item scored on 7 point Likert scale from 'not at all' to 'very much'.                                                                 | I: 7 – 18 years | - Developed and validated in mixed population with chronic pain e.g. fibromyalgia, rheumatoid arthritis, headache - Self report and parent scale - Validated in German. Based on Multidimensional Pain Inventory | - Requires further evaluation | - Requires further evaluation - Analysis of real time data collection is complex potentially limiting its clinical implementation - E-ouch recorded on a personal handheld device as opposed to an app or online - Pre-specified reporting times used in testing |
| <strong>Pain-Quilt</strong> | Web-based tool for tracking pain (quality, intensity, location) using time-stamped records. Pain quality involves choosing from a validated library of pain icons, e.g. a matchstick for ‘burning pain’. Pain intensity; 0-10 NRS from ‘no pain’ to ‘worst pain imaginable’. Pain location by dragging-and-dropping pain icons onto a virtual body-map. | I: 12 – 18 years | ✓ | ✓ | ✓ | ✓ | ✓ | ✓ | ✓ | - Ability to self-report complex sensations associated with chronic pain e.g. ‘burning’ - Real time data entry - Users able to record information faster than other electronic pain assessment tools (median completion 1.4 min) - Available at: <a href="http://www.painquilt.mcmaster.ca">www.painquilt.mcmaster.ca</a> | - Requires further evaluation - Developed and tested in an adult population and utility tested in adolescent populations |
| <strong>Pediatric pain assessment tool (PPAT)</strong> | 32 word descriptors, 10 cm VAS: present and worst pain, pain coping strategies, and influence of pain on daily activities. Modelled on McGill Pain questionnaire, Pediatric Pain Questionnaire. | I: 5 – 17 years | ✓ | - | ✓ | ✓ | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | Used JIA, cancer, surgery - child and parent versions - Children 7-16 years had no problems using scale - Developed in Dutch. Validated in Jordanian. Has been administered to Arab-American children. | - Younger children may require assistance - PPAT has been evaluated as single components, not the entire scale |
| <strong>Promis Pediatric Pain Interference Scale (PPIS)</strong> | Pain interference bank contains 13 questions, (8 in the short form). All questions use 7 day recall scored on a 5 point Likert scale with anchors 0=‘never’ to 4=‘almost always’. Raw score totals can be converted to a T-score using reference tables | I: 8 – 18 years | - | - | ✓ | ✓ | ✓ | ✓ | - | - | - | - | - | - | - | - | - | - | - | - | - | Developed for wide range of conditions e.g. arthritis, rehabilitation, asthma. - developed using Item Response Theory, users can select appropriate items from a ‘bank’ of questions - Preliminary validation in chronic MSK pain receiving multidisciplinary treatment - Child (8-17 years) and parent proxy (children 5-17 years) versions. - Developed in English. Validated in German and Spanish | - Requires further evaluation - Most accurate way to score the scale is through the online assessment centre - Scale can be scored manually however require reference tables to convert to a T-score. |
| <strong>Teen Nordic Musculoskeletal Screening Questionnaire (TNMQ-S)</strong> | 27-item questionnaire, dichotomous responses. The presence of MSK symptoms, their impact on school attendance, and on sports and leisure activity participation over the past 6 months. | I: 6 – 18 years | ✓ | - | - | ✓ | - | - | - | - | - | - | - | ✓ | - | - | - | - | - | - | - | Musculoskeletal symptom screening tool - Developed and tested in French. | - Requires further evaluation |
| <strong>Varni-Thompson</strong> | Assesses chronic pain intensity, location, sensory, evaluative and | I: 4 – 18 | ✓ | ✓ | ✓ | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | Developed and tested JIA | - Administered by interview |</p>
<table>
<thead>
<tr>
<th><strong>Pediatric Pain Questionnaire (PPQ)</strong></th>
<th>affective qualities. Pain intensity measured on 10 cm VAS, body chart (location and number of pain sites), 46 word descriptors to assess sensory, evaluative and affective qualities of pain. Modelled on the McGill Pain questionnaire.</th>
<th>year</th>
<th>applicable age range</th>
<th>I: 9-11 years</th>
<th>S: 6 – 16 years</th>
<th>- Assesses chronic pain from perspective of adolescent, parent or clinician - in various populations (juvenile rheumatoid arthritis, sickle-cell, fibromyalgia) - Adapted for children with cerebral palsy - Developed in English. Validated: Norwegian. Translations: Danish, Portuguese, Swedish, French.</th>
<th>10-15 mins to complete. &lt;5 mins to score. - younger children likely require assistance</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Young spine questionnaire</strong></td>
<td>For each region of the spine (cervical, thoracic, lumbar); pain presence, frequency and intensity (Revised Faces Pain Scale). Also includes function at school, recreation, treatment and family history of pain. No summary score.</td>
<td>I: 9-11 years</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>- Iterative development with children aged 9-11 years. - Self report - Developed in Danish. Translated to English.</td>
</tr>
</tbody>
</table>

I: intended age range; S: studied age range. ✓ Evidence available; – no supporting evidence
Table 5: Summary of the scales that have been suggested to measure pain intensity for each age group

<table>
<thead>
<tr>
<th>Age range</th>
<th>Recommended scale</th>
<th>Type of scale</th>
<th>Psychometric properties</th>
<th>Strengths</th>
<th>Limitations</th>
</tr>
</thead>
</table>
| Infants (3 years or younger) | Face, Arms, Legs, Activity Cry and Consolability Scale (FLACC) | Observational | Reliability: Yes Validity: Yes Responsiveness: Yes User preference: Yes Patient preference: Yes | - Validated for acute procedural pain e.g. immunisations and post-operative pain  
- Scored on a commonly understood 0-10 scale  
- Recommended for use by PedIMMPACT  |
|                          |                                                             |               |                                                                                        | - Additional studies needed to evaluate generalisability.                                                                                      |                                                                                                |
| Preschool child (3-5 years) | Pieces of hurt tool (Hester's Poker Chip Tool) | Self-report   | Reliability: Yes Validity: Yes Responsiveness: Yes Assessor preference: - Patient preference: Yes | - Developmentally appropriate scale i.e. yes/no followed by limited response options  
- Validated recurrent or persistent pain  
- Validated in Jordanian and Thai  
- Can use any items  
- Recommended for use by PedIMMPACT |
|                          |                                                             |               |                                                                                        | - Little testing in younger children  
- Potential bias toward higher pain scores especially in younger children  
- Requires the ability to count and estimate quantities using numbers  
- Infection risk, storage, and availability of tokens |                                                                                                |
- Gender neutral drawings of faces  
- Translations: > 35 languages, validated in French, Thai and Catalan  
- Minimally clinically important change: 2/10 (1 face), or 25%  
- Electronic version available; Sydney Animated Facial Expressions Scale (SAFE), Painometer App  
- Recommended for use by PedIMMPACT  |
|                          |                                                             |               |                                                                                        | - Reduced accuracy with decreasing age  
- Response bias in those under 5 years of age  
- Skewed pain intensity ratings towards 'no pain', may underestimate pain intensity |                                                                                                |
- Adaptable e.g. usual, strongest, lowest pain, at rest / activity  
- Translations: Spanish, French  
- suitable for 8-20 yrs, physical disabilities (e.g. cerebral palsy, neuromuscular disease), chronic pain  
- electronic version available114, 115  
- Minimally clinically important change 1/10  |
|                          |                                                             |               |                                                                                        | - Requires the ability to count, estimate quantities using numbers, recall pain  
- Variability in the time period (e.g. past week, current pain) and anchors used (e.g. worst possible pain, strongest pain)  
- Further testing required in younger children (6-8 years). |                                                                                                |
<table>
<thead>
<tr>
<th>Cognitive Impairment (4-19 years)</th>
<th>Revised Face, Arms, Legs, Activity Cry and Consolability Scale (rFLACC)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>28, 36, 43, 85, 103-105, 149</td>
</tr>
<tr>
<td>Observational</td>
<td>Reliability: Yes</td>
</tr>
<tr>
<td></td>
<td>Validity: Yes</td>
</tr>
<tr>
<td></td>
<td>Responsiveness: Yes</td>
</tr>
<tr>
<td></td>
<td>User preference: Yes</td>
</tr>
<tr>
<td></td>
<td>Carer preference: Yes</td>
</tr>
<tr>
<td></td>
<td>- simple to use, score and interpret</td>
</tr>
<tr>
<td></td>
<td>- Scored on a commonly understood 0-10 scale</td>
</tr>
<tr>
<td></td>
<td>- Individualised by parent/carer</td>
</tr>
<tr>
<td></td>
<td>- Validated in Danish</td>
</tr>
<tr>
<td></td>
<td>- evaluated in post-operative settings only</td>
</tr>
<tr>
<td></td>
<td>- underlying motor impairments e.g. spasticity may affect assessment</td>
</tr>
<tr>
<td></td>
<td>- Observation time not established; 5 min used in previous testing.</td>
</tr>
<tr>
<td></td>
<td>- Ambiguity of some items e.g. 'jaw clench' and consolability (amount and timing).</td>
</tr>
</tbody>
</table>
Figure 1: The social communication model of pain

* Pain: a distressing experience associated with actual or potential tissue damage with sensory, emotional, cognitive and social components (Williams, 2016).
† Pain experience and expression can be influenced by a number of individual, social and environmental factors, examples of each of these factors are outlined in Figure 2. The expression of pain may verbal, self-reported or observation of behaviour and this may be self-initiated by the person in pain or elicited by an observer’s question.
‡ Pain assessment can be influenced by a number of clinician factors and practical considerations, examples of these are outlined in Figure 2.
§ Pain management can include pharmacological, conservative (e.g. distraction, relaxation, exercise, education) and environmental interventions.
Pain management can be influenced by a clinician / parent or carers appraisal of the situation, their own knowledge / understanding and beliefs and contextual factors e.g. setting.

Adapted from T Hadjistavropoulos and KD Craig, 2002, Fig 1, p 555 and KD Craig, 2009, Figure 1, p24.33, 53
Figure 2: Factors (biological, psychological and social factors) related to the individual in pain, clinician and context that influence how pain is experienced, expressed and assessed.\textsuperscript{58, 78, 112, 150, 153, 160}
References:


79. Lauridsen H, Hestbaek L. Development of the young spine questionnaire. *BMC Musculoskelet Disord*. 2013;14:


