Section 2.0: Clinical Practice Points
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Conflict of Interest Declaration
The contributing committee members have no conflict of interest to declare.

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**How to Cite This Document**

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Section 2.0 Clinical practice points

Section 3.0 Pediatric chronic pain assessment tools

Section 4.0 Implementation supports manual
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This Chronic Pain Assessment Toolbox for Children with Disabilities is a compendium of resources based upon high quality research evidence intended to support the assessment of chronic pain in pediatric disability clinical practice. Evidence to Care spearheaded this project using an integrated knowledge translation strategy by engaging an inter-professional group from Holland Bloorview Kids Rehabilitation Hospital.

The current document presents the recommendations sourced from the Registered Nurses’ Association of Ontario (RNAO) Best Practice Guideline (BPG) for the Assessment and Management of Pain. This BPG was selected from a structured review and is available for download from the RNAO website. The accompanying evidence-informed practice points were developed through consultation with an inter-professional group of childhood disability experts from Holland Bloorview Kids Rehabilitation Hospital. The practice points are tailored to children with cerebral palsy (CP), but have vast applicability to children with other disabilities with chronic pain concerns.

For additional resources, including implementation supports, please visit the Evidence to Care website or contact etc@hollandbloorview.ca.
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Section 1a: Conducting a Pain Assessment – The Importance of Assessing Pain

<table>
<thead>
<tr>
<th>Registered Nurses’ Association of Ontario Recommendation</th>
<th>Evidence Rating</th>
</tr>
</thead>
<tbody>
<tr>
<td>Recommendation 1.1&lt;sup&gt;1&lt;/sup&gt;</td>
<td></td>
</tr>
<tr>
<td>Screen for the presence, or risk of, any type of pain:</td>
<td>Ib</td>
</tr>
<tr>
<td>• On admission or visit with a healthcare professional;</td>
<td></td>
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<tr>
<td>• After a change in medical status; and</td>
<td></td>
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<tr>
<td>• Prior to, during and after a procedure.</td>
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Holland Bloorview Cerebral Palsy and Chronic Pain: Practice Points

1. Chronic pain in children with cerebral palsy (CP) is under-recognized and under-treated, and is one of the most important factors negatively impacting a child’s ability to participate in activities of daily living. The presence of cognitive, communication, and motor impairments can make accurate assessment difficult.<sup>3-6</sup>

Following the points will facilitate communication between the inter-professional healthcare team who are treating the child, to better the management of the child’s pain.

2. Be conscientious for any indication of pain by asking the child, family or caregiver about the presence of pain or discomfort.<sup>7</sup>

3. Pain assessment should be considered the fifth vital sign in clinical care.<sup>7-9</sup> It is essential that children with CP are routinely monitored, assessed, and reassessed for pain and that their experience of pain is well documented.<sup>10,11</sup> Screening for pain should be routine and should consider the:

- Child’s level of function (e.g. Gross Motor Function Classification System [GMFCS] level);
- CP classification (e.g. type of hypertonia such as spastic or dystonic CP);
- Comorbidities (e.g. constipation, gastroesophageal reflux disease, fractures, hip subluxation);
- Setting (e.g. home, school, hospital); and
- Intervention (e.g. therapy, Botox injections).

4. Evaluation of chronic pain in children with CP should involve a comprehensive, child and family-centred health approach, whereby all aspects (e.g. physical, psychological, social) are considered as a whole and not independently in the assessment of the pain (e.g. biopsychosocial circumstances, culture, [family] financial stress, mental health...
### Section 1 b: Conducting a Pain Assessment – Assessment Parameters, the Role of Self-Report and Use of Pain Tools

<table>
<thead>
<tr>
<th>Registered Nurses’ Association of Ontario Recommendation</th>
<th>Evidence Rating</th>
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</thead>
<tbody>
<tr>
<td>Recommendation 1.2¹</td>
<td>Ib</td>
</tr>
<tr>
<td>Perform a comprehensive pain assessment on persons screened having the presence, or risk of, any type of pain using a systematic approach and appropriate, validated tools.</td>
<td></td>
</tr>
<tr>
<td>Recommendation 1.3¹</td>
<td>III</td>
</tr>
<tr>
<td>Perform a comprehensive pain assessment on persons unable to self-report using a validated tool.</td>
<td></td>
</tr>
<tr>
<td>Recommendation 1.4¹</td>
<td>III</td>
</tr>
</tbody>
</table>

Chronic Pain Assessment Toolbox for Children with Disabilities

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<table>
<thead>
<tr>
<th>Pain Assessment</th>
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<tbody>
<tr>
<td><strong>Recommendation 3.3</strong></td>
</tr>
<tr>
<td>Teach the person, their family and caregivers about the pain management strategies in their plan of care and address known concerns and misbeliefs.</td>
</tr>
</tbody>
</table>

**Holland Bloorview Cerebral Palsy and Chronic Pain: Practice Points**

<table>
<thead>
<tr>
<th>Pain Assessment</th>
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<tbody>
<tr>
<td><strong>Pain Assessment</strong></td>
</tr>
<tr>
<td>1. When assessing pain in children with CP, it is important to recognize that there is no single objective assessment strategy and that a multidimensional approach (e.g. assessments of behaviour, communication needs, and functional abilities) should be taken.</td>
</tr>
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<thead>
<tr>
<th>Pain Assessment</th>
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<tbody>
<tr>
<td><strong>The Role of Self-Report</strong></td>
</tr>
<tr>
<td>2. Children with CP may communicate in different ways. In addition to verbal self-report, consider self-report through yes/no responses, use of communication devices, facial expressions, gestures, eye contact/eye pointing, and others.</td>
</tr>
</tbody>
</table>
2.0 Clinical Practice Points

**Involving Family and Caregivers**
3. Family and caregivers are integral to the pain assessment process. Their participation in the assessment of their child’s pain allows them to:
   - Better understand their child’s cause of and response to pain;
   - Provide an opportunity for reflection of ongoing, chronic pain;\(^{11,14}\)
   - Draw upon their knowledge regarding the child’s communication patterns, mood, and behaviour;\(^{11}\) and
   - Become comfortable and confident with pain assessment and monitoring strategies by learning how to use the pain assessment tools and document pain.

4. In situations where the child with CP is not able to self-report pain, pain assessment may include an observational or proxy tool as well as behavioural and physiological indicators (e.g. behavior appearance, activity level, and vital signs). Whenever possible, standardized measures should be used when assessing pain.

**Use of Pain Tools**
5. It is important to recognize that no individual tool can be broadly recommended for pain assessment in all children across all contexts. To ensure the most effective assessment tool is chosen for a child with CP and chronic pain, it is important to consider the:
   - Child’s individual condition and circumstances (e.g. age, GMFCS level, developmental level, presence of developmental disability, communication ability, setting);
   - Purpose of the pain assessment (e.g. presence of pain, benefit of treatment, coping); and
   - Selection of an alternative tool if the tool that has been chosen is not providing the required information.

The International Classification of Functioning, Disability and Health Model can be used to direct a comprehensive pain assessment.\(^{15}\)

**Body Functions and Structure**

**Pain History**
- Conduct a complete medical and pain history, including onset, intensity, quality, location, duration, variability, predictability, exacerbating, and alleviating factors.\(^{16}\)
- Ask about the presence of regular or persistent crying, groaning, facial grimacing, loss of interest in food, difficulty sleeping, changes in appearance, and activity level.\(^{8,13}\)
- Document previous pain experiences (e.g. surgeries, history of unmanaged pain,
### 2.0 Clinical Practice Points

Painful procedures during previous hospitalization) and identify any past or current treatments for pain and associated timelines (e.g. use of medication for >72 hours), level of effectiveness, and any adverse effects.\(^1\)\(^,\)\(^8\)

**Pain Evaluation**
- Basic vital signs and growth parameters should be obtained during at least the first evaluation. Additional symptoms such as motor (e.g. increased tone, muscle spasms), sensory (e.g. changes in heart rate, blood pressure, respiratory rate) and autonomic changes (e.g. numbness, hyperesthesia) as measures of pain should be noted.\(^17\)
- Include an observation of the child’s general appearance, posture, gait, and cognitive state with the focus on, but not limited to the affected area.\(^16\),\(^18\) Note muscle spasms, trigger points, and areas of somatic sensitivity to touch and functional range of motion.\(^18\),\(^19\) Serial examinations may be required.\(^16\)

**General Medical History**
Conduct a general medical history by including documentation of primary and secondary diagnosis, past medical history, allergies/sensitivities, risk factors (e.g. airways stability, disease process), medications, review of systems, and impact of interventions (medication, therapy, procedures).\(^17\),\(^20\)

**Activity and Participation**
- Assess the impact of pain on daily life activities: sleep, eating, school, social and physical activities, and family and peer interaction.\(^8\),\(^16\)

**Environmental Factors**
- Engage parents and caregivers to better understand the cause and response to the child’s pain.\(^11\),\(^14\)
- Review past pain problems and pain experiences, as well as, current treatments (e.g. medications, home remedies, alternative treatments) with parents and caregivers.\(^16\),\(^18\)
- Assess environmental factors that may impact the child’s pain experience (e.g. school environment, home environment, extracurricular activities).

**Personal Factors**
- Consider the individual’s developmental stage (i.e. chronological vs. mental age).
- Consider that language, ethnic, and cultural factors may influence the expression and assessment of pain.\(^7\)
- Assess the child’s emotional functioning, coping responses to stress and pain, and the ability to talk about pain.\(^7\),\(^16\)
- Include information about the presence of psychological symptoms (e.g. anxiety, fears, depression or anger), psychiatric disorders, and personality traits/states.\(^17\),\(^21\)
2.0 Clinical Practice Points

Section 2: Pain Reassessment

<table>
<thead>
<tr>
<th>Registered Nurses’ Association of Ontario Recommendation</th>
<th>Evidence Rating</th>
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</thead>
<tbody>
<tr>
<td>Recommendation 4.1&lt;sup&gt;1&lt;/sup&gt; &lt;br/&gt;Reassess the person’s response to the pain management interventions consistently using the same re-evaluation tool. The frequency of reassessments will be determined by: &lt;br/&gt;• presence of pain; &lt;br/&gt;• pain intensity; &lt;br/&gt;• stability of the person’s medical condition; &lt;br/&gt;• type of pain e.g. acute versus persistent; and &lt;br/&gt;• practice setting.</td>
<td>IIb</td>
</tr>
<tr>
<td>Recommendation 3.3&lt;sup&gt;1&lt;/sup&gt; &lt;br/&gt;Teach the person, their family and caregivers about the pain management strategies in their plan of care and address known concerns and misbeliefs.</td>
<td>Ib</td>
</tr>
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Holland Bloorview Cerebral Palsy and Chronic Pain: Practice Points

Reassess Pain
1. Ongoing pain assessment is an integral component of comprehensive pain management strategy for children with CP.<sup>8, 19</sup>
2. When re-examining the child, be aware of alternative organic causes of pain (e.g. dental pain, pressure ulcers) or psychological (e.g. stress) or psychiatric comorbidities (e.g. depression, anxiety).<sup>22</sup> As part of pain reassessment, consider repeating the components of section 1b, practice point 5 (where necessary) and additionally review the: <br/>• Effects of pain on activities of daily living (ADLs), sleep, and mood; <br/>• Adverse effects of medications for pain treatment (e.g., nausea, constipation); <br/>• Level of sedation; and <br/>• Strategies used to relieve pain, both pharmacological, and non-pharmacological.<sup>1</sup> 
3. If a child has persistent pain despite the clinician’s best efforts to relieve it, involve an inter-professional team to consult on alternative pain reduction strategies.

Involve Family and Caregivers
4. Provide instructions to the child, their family and caregivers on: <br/>• How to use a pain log or diary (provide a tool); and <br/>• How to communicate unrelieved pain to their clinician and support them in efforts to advocate for themselves or for their child.<sup>1</sup>
### 2.0 Clinical Practice Points

#### Section 3: Documenting Pain

<table>
<thead>
<tr>
<th>Registered Nurses’ Association of Ontario Recommendation</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Recommendation 4.2&lt;sup&gt;1&lt;/sup&gt; Communicate and document the person’s responses to the pain management plan.</td>
<td>IIb</td>
</tr>
<tr>
<td>Recommendation 1.5&lt;sup&gt;1&lt;/sup&gt; Document the person’s pain characteristics.</td>
<td>IIa</td>
</tr>
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</table>

#### Holland Bloorview Cerebral Palsy and Chronic Pain: Practice Points

1. Document the comprehensive pain assessment including pain history, evaluation, and the impact of the pain on the child and their parent, caregiver and family. Communicate the following information to the members of the inter-professional team:
   - Parameters of pain obtained through the use of a structured assessment tool;
   - Relief or lack of relief obtained from treatment methods and related adverse effects; and
   - Child, family or caregiver’s goals for pain treatment and the effect of pain on the child.<sup>1</sup>

2. Document the pain assessment tool(s) used effectively to capture the child’s experience of pain so that there is consistency within the inter-professional team when assessing and reassessing pain.
**Interpretation of Evidence: Levels of Evidence**

The following table presents the level of evidence and corresponding rationale, adapted by the Registered Nurses’ Association of Ontario and displayed within the Assessment and Manage of Pain, Third Edition¹ (page 11).

<table>
<thead>
<tr>
<th>Level of Evidence</th>
<th>Rationale</th>
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<tbody>
<tr>
<td>Ia</td>
<td>Evidence obtained from meta-analysis or systematic reviews of randomized controlled trials.</td>
</tr>
<tr>
<td>Ib</td>
<td>Evidence obtained from at least one randomized controlled trial.</td>
</tr>
<tr>
<td>IIa</td>
<td>Evidence obtained from at least one well-designed controlled study without randomization.</td>
</tr>
<tr>
<td>IIb</td>
<td>Evidence obtained from at least one other type of well-designed quasi-experimental study without randomization.</td>
</tr>
<tr>
<td>III</td>
<td>Evidence obtained from well-designed non-experimental descriptive studies, such as comparative studies, correlation studies and case studies.</td>
</tr>
<tr>
<td>IV</td>
<td>Evidence obtained from expert committee reports or opinions and/or clinical experience of respected authorities.</td>
</tr>
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</table>
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References


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Holland Bloorview

Kids Rehabilitation Hospital

Teaching and Learning Institute

Accelerate knowledge

Evidence to Care

Inspire our people

Transform care

Lead the system