 TOOLKIT FOR PAIN ASSESSMENT

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About RHSCIR

The RICK HANSEN SPINAL CORD INJURY REGISTRY (RHSCIR) is a pan-Canadian prospective observational registry located at 31 major Canadian acute care and rehabilitation facilities. Across Canada, RHSCIR is collecting comprehensive SCI data for the purpose of improving spinal cord injury (SCI) care and clinical outcomes. Using standardized research protocols and data collection forms, RHSCIR tracks the experiences and outcomes of people with traumatic SCI during their journey from acute care to rehabilitation to community reintegration. Details about participants’ spinal cord injuries including extent of injury and level of paralysis, recovery, and success of various treatments are among the data recorded.

The data collected in RHSCIR contains powerful information that will help track the effectiveness of specific treatments, practices or programs for improving functional outcomes and quality of life after SCI. RHSCIR promotes, encourages and supports the pursuit of excellence in all areas of SCI health care management.

This network of 31 facilities has a new goal of standardizing the basic assessment of pain across facilities in their patients who have sustained a SCI. This minimum amount of standardized pain assessment information is not meant to replace current clinical practice, but is being standardized and will be added to RHSCIR in order to support tracking and reporting of this important information across the facilities. The ultimate goals are to support the use of this important data in informing facilities of their facility or program level trends, providing information on how your facility compares to national trends, while capturing data to answer valuable clinical and research questions at a national level.

To learn more about RHSCIR, please visit www.rickhanseninstitute.org.
WHY IS THIS INFORMATION IMPORTANT?

The majority of people with a spinal cord injury (SCI) will experience pain during their lifetime. Pain negatively affects health-related quality of life, social participation and everyday function (e.g., sleep, exercise, and employment) as well as psychological well-being. Pain is likely to increase over time in 47% of persons with SCI, and decrease in only 7% (1). Given the prevalence and negative effects of pain, it is important to gather information to better understand how to manage it. This information can then be used to identify/develop needed resources (e.g., staffing, intervention programs). In addition, such information can serve as the basis for patient education as part of learning about self-management and directing care.

All the information collected will be added to RHSCIR providing validation and support of evidence-based practice. Much of this information is aligned with the International SCI Pain Basic Data set (Version 2.0), which is an important source of expert-derived recommendations developed by experts in SCI-related pain.

Benefits to Clinicians and Patients

Collection and reporting of this data can benefit clinicians and patients by:

- Provides information on the more common pain types that are most troublesome to patients
- Makes you aware of what to ask for and what to look for in new patients
- Contributes to the discipline by furthering understanding of:
  - Risk factors for worst pain experienced.
  - Average pain severity for worst pain experienced.
  - The most common location of worst pain experienced.
  - Impact of pain on daily functioning.
  - Frequency with which different treatment options are used.
  - Level of satisfaction with treatment strategies.

Should a patient ask why the information is being collected, it may be useful to respond in a similar manner by highlighting some key points (e.g. "As a patient, members of your health care team will be asking you many different questions about pain at multiple time points after your injury. Since there are a variety of types of pain caused by different things that can occur after a spinal cord injury, these questions are important as they will assist your health care team in determining how best to treat your specific pain."

As clinicians, you will be collecting pain data from your patients throughout their stay at your facility. Section 3 outlines the minimum amount of pain assessment data recommended to collect (i.e. Clinical Data Collection Forms). Collecting this data consistently on your patients will allow you to objectively track and analyze any changes and help modify, enhance, and direct future care and treatment plans. In addition, community follow-up interviews with the patient will be completed by a RHSCIR coordinator or designate.
Benefits to the Program

In order to optimize the efficiency and outcome of pain management in individuals with SCI on your program, it is important to collect, analyze, and review the pain assessment information in a standardized manner. The collection of this standardized pain assessment data by both clinicians and RHSCIR staff at your facility, will allow you to access the type of data outlined below that will support both program level management and evaluation as well as give you the ability to compare your program with similar programs across Canada.

Collection and reporting of this data can benefit your program by:

- Setting staffing levels, determining what type of staff are required (e.g., RN, OT, PT, research) and how many.
- Determining required equipment.
- Larger system planning.
- Providing metrics to support quality improvement initiatives.
- Providing comparisons to national data.
- Reporting metrics to program managers and facility administrators.
- Meeting requirements for the new Accreditation Canada SCI Acute Care and Rehabilitation Standards.
- Providing insight into understanding and ensuring that inpatient programs are meeting the needs of patients (i.e. patient satisfaction), and setting the stage for life in the community.
- Identifying gaps in the community.

What Happens Once I Collect the Data?

Providing invaluable data to RHSCIR: Once you collect the data, your facility’s RHSCIR coordinator will abstract this information and input it along with any other relevant clinical information into the registry database. The national RHSCIR team has developed and implemented a number of practices to ensure patient confidentiality is maintained and strict privacy policies and procedures are adhered to.

Providing a baseline for management of SCI across Canada: The de-identified data from your facility and other facilities participating from your site will be reported back on a quarterly basis and will provide information on the frequency of pain treatment (worst pain, general pain) including things like treatment type, satisfaction with pain management, pain interference, location, intensity, as well as the frequency, intensity and location of neuropathic pain, etc.

To access your site’s data reports, visit Supporting Clinical Initiatives in SCI (SCI²) resource site at http://sci2.rickhanseninstitute.org. Please see your local RHSCIR coordinator, or designated representative, to receive this log in information.

You can also access the SCI² site by visiting www.rickhanseninstitute.org.
Please take note of the following instructions to complete the Clinical Data Collection.

**Time**
- Estimated time required: 20 minutes

**Equipment (if applicable)**
- Clinician Pain Assessment form
- Pain assessment tool e.g., DN4, LANSS (optional)
Please complete this form within 7 days of admission and 7 days prior to discharge from facility.

1. Date pain assessment completed: 
   Enter as much of the date as is known.

2. Clinician Name/Signature: ________________________________

3. Is participant receiving any treatment for pain? (e.g., medications, recreational drugs, physical therapies, psychological treatment, etc.)
   □ Yes
   □ No

4. a) Has the participant had any pain in the last 7 days including today?
   □ Yes
   □ No
   □ Unknown

   b) How many different pain problems does the participant have?
   □ 1
   □ 2
   □ 3
   □ 4
   □ ≥5

   c) For participant’s pain, please indicate worst pain location and characteristics below:

<table>
<thead>
<tr>
<th>Pain Location/Sites:</th>
<th>Right</th>
<th>Midline</th>
<th>Left</th>
</tr>
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<tbody>
<tr>
<td>Head</td>
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<td>Lower back</td>
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<td>Buttocks</td>
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<td>Hips</td>
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<td>Anus</td>
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<td>Upper legs/thighs</td>
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<td>Lower legs/feet</td>
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<td>Knee</td>
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<td>Ankle</td>
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<td>Foot/Toes</td>
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<tr>
<td>Unknown location/site</td>
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</tbody>
</table>

□ Pain as bad as you can imagine
□ Unknown

□ None

□ Other (specify): ______________________

□ Leeds Assessment of Neuropathic Symptoms and Signs (LANSS)
□ Douleur Neuropathique en 4 Questions (DN4)
□ Other (specify): ______________________

□ Pain Intensity:
□ 0 No pain
□ 1
□ 2
□ 3
□ 4
□ 5
□ 6
□ 7
□ 8
□ 9
□ 10 Pain as bad as you can imagine
□ Unknown

□ Which pain assessment tool was used?
□ No therapy

□ 0
□ 1
□ 2
□ 3
□ 4
□ 5
□ 6
□ 7
□ 8
□ 9
□ 10 Pain as bad as you can imagine
□ Unknown

□ Date of Onset (Date this pain started): ________/____/____
Timing of Assessment

**Admission:** A comprehensive Clinician Pain Assessment should be performed as soon as possible after admission, but within seven days at a minimum. If you are unable to complete the comprehensive assessment within seven days (e.g. the patient is too critically ill, or access is not possible due to surgery, etc.), please complete the pain assessment and data collection form as soon as possible.

**Discharge:** A comprehensive Clinician Pain Assessment should be performed within seven days prior to discharge. Ideally, please complete as close to discharge as possible.

Step 1: Schedule the Comprehensive Clinician Pain Assessment

- Schedule the comprehensive Clinician Pain Assessment with the patient.
- Inform the patient of this procedure and obtain verbal consent from the patient.

Step 2: Collect Supplies

- Clinician Pain Assessment form.
- Pain assessment tool e.g. DN4, LANSS (optional).

*Insert the date of assessment and your name on the form.*

Step 3: Review Patient Information

- Review available information on the patient e.g. history, chart notes, clinician assessments to obtain background information about the patient and their pain.

Step 4: Patient Interview

- Ask the patient the following questions, from the Clinical Pain Assessment form:
  3. *Is the participant receiving any treatment for pain?*
  4. *Has the participant had any pain in the last 7 days including today?*

- If the patient identifies the presence of pain, document their worst pain and complete the following sections on the form pertaining **only to their worst pain:**
  - Pain location/sites
  - Type of pain (nociceptive vs. neuropathic)
  - Date of onset
  - Which pain assessment tool was used (if any)
  - Pain intensity
DEFINITIONS OF SPINAL CORD PAIN TYPES

NOCICEPTIVE PAIN TYPES

Musculoskeletal: refers to pain occurring in a region where there is preserved sensation above, at or below the neurological level of injury and which is believed to be arising from musculoskeletal structures. The presence of this type of pain is suggested by pain descriptors such as dull or aching, pain related to movement, tenderness of musculoskeletal structures on palpation, response to anti-inflammatory or opioid medications and evidence of skeletal pathology on imaging consistent with the pain presentation. Examples include mechanical pain, spinal fractures, muscular injury, shoulder overuse syndromes and muscle spasm.

Visceral: refers to pain usually located in the thorax, abdomen, or pelvis and believed to be generated in visceral structures. The presence of this type of pain is suggested by characteristics such as dull, tender, or cramping and a relationship to visceral pathology or dysfunction (e.g. infection or obstruction). Examples include urinary tract infection, ureteric calculus and bowel impaction. Note: Failure to find evidence of visceral pathology or failure to respond to treatment directed at visceral pathology may indicate the presence of neuropathic pain.

Other: refers to nociceptive pains that may be present but do not fall into the musculoskeletal or visceral categories. Examples include pain associated with ulceration of the skin and headache.

NEUROPATHIC PAIN TYPES

At-level: refers to neuropathic pain presenting in a segmental pattern. A necessary condition for this to occur is that there is a lesion or disease affecting the spinal cord or nerve roots. At-level neuropathic pain is perceived anywhere within the dermatome of the level of neurological injury and three dermatomes below this level. At-level neuropathic pain is often characterized as hot-burning, tingling, pricking, pins and needles, squeezing, cold, electric, or shooting. Sensory changes such as allodynia, hypoalgesia, or hyperalgesia within the pain distribution are often found.

Below-level: refers to neuropathic pain that is present more than three dermatomes below the dermatome of the neurological level of injury; it may be in addition be perceived up to the dermatome representing the neurological level of injury and the three dermatomes just below this. A necessary condition for this to occur is that there is a lesion or disease affecting the spinal cord and that the pain is believed to arise as a result of this damage. Below-level pain is often characterized as hot-burning, tingling, pricking, pins and needles, squeezing, cold, electric, or shooting; it usually has a regional distribution. Sensory changes such as allodynia, hypoalgesia, or hyperalgesia may be present.

Other: refers to neuropathic pains that are present above, at or below the neurological level of injury but are not directly related to the SCI. Examples include postherpetic neuralgia, pain associated with diabetic neuropathy, central post stroke pain, and compressive mononeuropathies.

Other pain: refers to pain that occurs when there is no identifiable noxious stimulus nor any detectable inflammation or damage to the nervous system responsible for the pain and the pain is thought to be unrelated to the underlying SCI, both temporally and mechanistically. It is unclear what causes the pain to develop or persist. Examples include Complex Regional Pain Syndrome type I, interstitial cystitis pain, irritable bowel syndrome pain and fibromyalgia.

Note: Source of definitions from the International Spinal Cord Injury Pain Basic Data Set (v2.0)(2).
Training Resources

The International SCI Pain Basic Data Set version 2.0 has Training Cases to help you understand the types of pain patients express and how to chart it: http://www.iscos.org.uk/sitefiles/2013%2006%2011_International%20SCI%20Pain%20Basic%20Data%20Set_Version%202%200.pdf.


Questions or comments regarding this guideline?
Email clinical@rickhanseninstitute.org.
REFERENCES & ADDITIONAL RESOURCES

References


Additional Resources

Pain BC has many resources for health care providers and their patients:
http://www.painbc.ca/resources-for-health-care-providers.

Pain resource centre, developed by the Canadian Pain Coalition is available here:

Other resources are available on the SCI² site, including examples of assessment and charting forms.


In addition to the clinically collected data outlined in this toolkit, there will be additional patient self-reported data collected by your facility’s RHSCIR team as outlined in the following pages. In addition, there will also be data collected by RHSCIR from those participants who consent to the community follow-up in RHSCIR. This includes subjective information on neuropathic pain, pain treatment(s), and pain interference and is collected upon discharge to the community and during subsequent RHSCIR community follow-up interviews (data collection points). This portion of data does not require any clinician time, but is collected by registry personnel and will also be made available to your facility.

These questions are collected by the RHSCIR team when the participant is discharged into the community (i.e. home, extended care, etc.) as well as at one year, two years, five years, and then every five years from their date of injury.
# Pain Questionnaire

These questions ask your opinion about any pain you may be experiencing and how it may interfere with your daily living. Also, you will be asked to describe what you do to manage it.

1. a) Are you currently using or receiving any treatment for a pain problem?
   - ☐ Yes (e.g., medications, recreational drugs, physical therapies, psychological treatment, etc.)
   - ☐ No
   - ☐ Unknown

   b) If Yes, What treatments do you use to manage your pain? (check ALL that apply)
   - ☐ Complementary (e.g., biofeedback, acupuncture, hypnosis)
   - ☐ Medical and procedural or neuromodulation (e.g., nerve blocks, injections, implanted stimulators, intrathecal pump, TENS)
   - ☐ Non-prescription medications (e.g., non-prescription pain killers such as Tylenol®)
   - ☐ Non-traditional (e.g., naturopathy, homeopathy, herbal remedies)
   - ☐ Psychotherapeutic (e.g., psychotherapy, cognitive behavioural therapy, relaxation, stress management, psycho-education, support group)
   - ☐ Physical therapies (e.g., physiotherapy, massage, chiropractic)
   - ☐ Recreational drugs (e.g., marijuana)
   - ☐ Prescription medications (e.g., morphine, codeine)
   - ☐ Other (specify): ____________________

2. Overall, how satisfied are you with the management of your pain?

   0 1 2 3 4 5 6 7 8 9 10
   - ☐ Unknown
   - Not at all satisfied
   - Completely satisfied

3. a) Have you had any pain during the last 7 days, including today?
   - ☐ Yes
   - ☐ No (skip to question 4)

   b) If YES, in the LAST WEEK:

   i) In general, how much has pain interfered with your day to day activities in the last week?

   0 1 2 3 4 5 6 7 8 9 10
   - ☐ Unknown
   - No interference
   - Extreme interference

   ii) In general, how much has pain interfered with your overall mood in the past week?

   0 1 2 3 4 5 6 7 8 9 10
   - ☐ Unknown
   - No interference
   - Extreme interference

   iii) In general, how much has pain interfered with your ability to get a good night’s sleep?

   0 1 2 3 4 5 6 7 8 9 10
   - ☐ Unknown
   - No interference
   - Extreme interference
4. a) Since your injury, have you experienced neuropathic pain? (Pain that is often ongoing and intense that occurs spontaneously or by light touching and is characterized by feelings of burning, shooting, tingling, etc.)
   - Everyday
   - Few times a week
   - Few times a month
   - Few times a year
   - Once a year
   - Never *(Pain information complete, no other questions answered)*
   - Don’t know *(Pain information complete, no other questions answered)*

b) Location(s) of your neuropathic pain (check ALL that apply):
   - Head
   - Neck and/or shoulders
   - Arms and/or hands
   - Torso (chest, abdomen, pelvis, and/or genitals)
   - Back (upper and/or lower back)
   - Hips, buttocks, and/or anus
   - Upper legs/thighs
   - Lower legs or feet

c) Average pain intensity of your neuropathic pain in the past week:

0 1 2 3 4 5 6 7 8 9 10
No pain

Pain as bad as you can imagine

   d) Have you received some form of treatment for the neuropathic pain?
   - Yes
   - No
   - Unknown

e) When you had neuropathic pain, to what extent did it limit your activities?
   - Not at all
   - Very little
   - To some extent
   - To a great extent
   - Completely
   - Don’t know