Canadian C-spine rule and the National Emergency X-Radiography Utilization Study (NEXUS) for detecting clinically important cervical spine injury following blunt trauma
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Canadian C-spine rule and the National Emergency X-Radiography Utilization Study (NEXUS) for detecting clinically important cervical spine injury following blunt trauma (Protocol)

Saragiotto BT, Maher CG, Lin CWC, Verhagen AP, Goergen S, Michaleff ZA

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# Table of Contents

- **Header** ...................................................................................................................... 1
- **Abstract** ...................................................................................................................... 1
- **Background** .................................................................................................................. 1
  - Figure 1. .......................................................................................................................... 3
- **Objectives** .................................................................................................................... 5
- **Methods** ....................................................................................................................... 5
- **References** ................................................................................................................... 7
- **Additional Tables** ........................................................................................................ 8
- **Appendices** .................................................................................................................. 10
- **What's New** .................................................................................................................. 11
- **History** ......................................................................................................................... 11
- **Contributions of Authors** ............................................................................................ 11
- ** Declarations of Interest** .............................................................................................. 11
- **Sources of Support** ...................................................................................................... 12
ABSTRACT

This is a protocol for a Cochrane Review (Diagnostic test accuracy). The objectives are as follows:

To describe and compare the diagnostic accuracy of the Canadian C-spine rule and the National Emergency X-Radiography Utilization Study (NEXUS) to screen for clinically important cervical spine injury (CSI) in patients following blunt trauma.

BACKGROUND

Cervical spine injury (CSI) represents approximately 3.5% of the cases of trauma presenting to emergency departments around the world (Hasler 2012; Milby 2008; Niska 2010). Approximately 2% of cervical spine injuries will be clinically important injuries such as fracture or dislocation and require specialist intervention (e.g. immobilisation or surgical intervention). Due to the potentially catastrophic consequences of a delayed or missed diagnosis of clinically important CSI (Davis 1993), diagnostic imaging is undertaken for the great majority of patients with CSI. As the prevalence of clinically important CSI is only 2% of total CSI, a mandatory imaging policy would lead to the large majority of patients with CSI receiving imaging that confers no net health benefit (Stiell 1997).

Clinical decision rules can assist clinicians to rule out clinically important CSI by identifying those patients with a lower likelihood of a clinically important CSI and therefore do not require imaging (Motor Accidents Authority 2014; TRACsa 2008). The use of validated tools to improve clinical assessment of pre-test risk has the potential to minimise costs, resource utilisation, length of stay in emergency departments, and unnecessary exposure to radiation (Griffith 2011). Use of such tools can facilitate shared decision
making with patients and provide them with reassurance about the reasoning behind a clinical decision to defer or not perform imaging at all.

The National Emergency X-Radiography Utilization Study (NEXUS) criteria and the Canadian C-spine rule are two clinical decision rules developed to help clinicians risk-stratify patients with cervical spine trauma to determine if they need imaging to rule out clinically important CSI. These clinical decision rules have been externally validated internationally and their use in routine clinical practice is recommended by international guidelines (Hoffman 2000; NICE Guidance 2007; The College of Emergency Medicine 2010; TRACsa 2008; Stiell 2001). For safe and effective screening, the rule must have a high-sensitivity rate, indicating that a clinically important CSI will not be missed. Other attributes of a high-performing clinical decision rule are selection criteria that allow its application to a broad range of patients at risk of the target condition and unambiguous definition/description of the items within the rule. These attributes work together to increase the likelihood of the correct use of the rule for the largest number of patients in a variety of clinical settings.

Despite evidence indicating higher sensitivity and specificity of the Canadian C-spine rule than NEXUS for patients meeting the inclusion/exclusion criteria for either of these rules (Michaleff 2012; Stiell 2003), the current imaging and other clinical practice guidelines do not reflect the differences in both performance as well as patient selection criteria for the two clinical decision rules (ACEM / RANZCR Guidelines 2012; ACR Appropriateness Criteria; Diagnostic Imaging Pathways).

**Target condition being diagnosed**

Clinically important CSI is defined as fracture, dislocation or mechanical instability of the cervical spine which requires specialist intervention (e.g. immobilisation or surgical intervention) (Stiell 1999; Stiell 2001). Blunt trauma results from an impact to the body in the absence of any penetrating trauma, examples of blunt trauma include motor vehicle accident, fall or assault.

The prevalence of CSI in trauma patients is estimated as less than 4%, of which only about half of these will have a fracture or dislocation, of which about 25% will report neurological deficits (Hasler 2012; Milby 2008; Niska 2010). Clinically important CSI requires specific treatment (e.g. operative stabilisation) to prevent secondary injury to the spinal cord, which would lead to significant disability, morbidity or mortality (Davis 1993; Goergen 2015).

**Index test(s)**

The NEXUS criteria (Hoffman 1998) (Table 1) and the Canadian C-spine rule (Stiell 2001) (Figure 1) are two clinical decision rules designed to be used for patients with CSI following blunt trauma. Both decision rules were developed for patients with cervical trauma in whom clinically important CSI is a concern; however the Canadian C-spine rule specifies its use for patients who are aged 16 years or older, alert (as indicated by a score of 15 on the Glasgow Coma Scale), and in a stable condition (Stiell 2001), whereas NEXUS has one inclusion criterion (suspected clinically important CSI following trauma) and one exclusion criterion (penetrating trauma) (Hoffman 2000). For both rules, patients with a negative result have an acceptably low risk of clinically important CSI and imaging is not required. A positive test result is interpreted quite differently as both rules have only modest specificity (< 50%), meaning that most positive test results are false positives (Michaleff 2012). This means that many patients who test positive and are imaged will be subsequently shown not to have clinically important CSI.
Figure 1. The Canadian C-Spine Rule.

Adult Canadian C-Spine Rule
for alert (GCS = 15) and stable trauma patients

**Inclusion Criteria:**
- Adults (defined as ≥13 years of age), AND
- Acute trauma to the head or neck, AND
- Stable (i.e. normal vital signs as per Revised Trauma Score), AND
- Alert (GCS = 15), AND
- Injury within previous 48 hours, AND EITHER
- Neck pain; OR
- No neck pain but meet the following criteria:
  - Visible injury above the clavicles; AND
  - Non-ambulatory; AND
  - Dangerous mechanism of injury*

**Exclusion Criteria:**
- Trivial injuries (e.g. Simple facial lacerations) and did not fulfill the “at risk” inclusion criteria;
- Penetrating trauma;
- Present with acute paralysis;
- Known vertebral disease (e.g. ankylosing spondylitis, rheumatoid arthritis, spinal stenosis, or previous cervical surgery) as determined by the examining physician;
- Returned to ED for reassessment of same injury;
- Pregnancy.

**Algorithm:**
- NO to ANY
- Exclude
- YES to ANY
- NO to ALL
- YES to ANY
- 1. Any high-risk factor that mandates radiography?
- Age ≥ 65
- or
- Dangerous mechanism
- or
- Paraesthesia in extremities
- NO to ALL
- YES to ANY
- 2. Any low-risk factor that allows safe assessment or range of movement?
- Simple rear-end MVC
- or
- Sitting position in ED
- or
- Ambulatory at any time
- or
- Delays onset of neck pain
- or
- Absence of midline/c-spine tenderness
- NO to ALL
- YES to ANY
- 3. Able to actively rotate neck?
- 45° left and right
- IMAGING NOT RECOMMENDED

*Dangerous mechanism:
- Fall from ≥2 stairs / 5 stairs
- Avulsion head to head, eg diving
- MVC high speed (~100 km/h), rollover, ejection
- Motorised recreational vehicle
- Bicycles collision

Simple rear-end MVC excludes:
- Pushed into oncoming traffic
- Hit by bus / large truck
- Rollover
- Hit by high speed vehicle

Delayed:
- Not immediate onset of neck pain

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The Canadian C-spine rule consists of a) three high-risk criteria, the presence of any one of which results in a recommendation for imaging (age ≥ 65 years, dangerous mechanism, or paraesthesia in extremities); b) five low-risk criteria the presence of any one of which allows the collar to be removed and cervical spine rotation assessed (simple rear-end motor vehicle collision, siting position in the emergency department, ambulatory at any time, delayed onset of neck pain, or absence of midline cervical-spine tenderness); and c) an assessment of the ability of patients to rotate the neck 45 degrees left and right (Stiell 2003) (Figure 1). The five criteria of the NEXUS are: 1) no posterior midline cervical tenderness; 2) no evidence of intoxication; 3) normal level of alertness (i.e. the patient is alert and oriented to person, place, time, and event); 4) no focal neurological deficit; and 5) no painful distracting injuries (e.g. long-bone fracture) (Hoffman 1998; Hoffman 2000) (Table 1). Patients who meet all five of the NEXUS criteria can have safely clinically important CSI excluded without the use of imaging. A positive test result for either rule suggests that the patient is more likely to have a clinically important CSI and in these patients recommends the use of imaging investigations. Neither rule however specifies the type of imaging investigation that should be used (i.e. computed tomography (CT) or plain radiography) and both are silent on whether plain radiography should be followed by CT if the patient is deemed, by some criterion (e.g. neurological abnormality or injury mechanism) to be at high risk.

**Clinical pathway**

When patients with CSI arrive at the emergency department, they typically undergo a series of assessments to rule out clinically important CSI. Firstly, the initial assessment includes the Glasgow Coma Scale score, history taking, which consists of medical history (i.e. history of spine surgery, previous neck pain), demographic details, mechanism of injury, presence, onset and progression of symptoms (i.e. pain, paraesthesia) (Ackland 2012; Goergen 2015). Then, examination of physical and neurological status is performed, which includes motor function (i.e. tone, strength, reflexes) and sensation (i.e. paraesthesia) (Ackland 2012; Goergen 2015). Further assessments would include the presence of bony tenderness in the midline, and the patient's ability to rotate the cervical spine 45 degrees to each side (Goergen 2015). Finally, most patients undergo imaging, usually radiography or CT (Como 2009; Daffner 2007). A semi-rigid cervical collar is often recommended until imaging can be conducted (Ackland 2012). In practice, the choice of using CT versus plain radiography for the diagnosis of cervical spine injury is subjective and depends on many factors, such as hospital policies and protocol, the availability of imaging equipment e.g. CT; the severity of the trauma and age of the patient. In children, radiography tends to be performed first and only when it is abnormal is CT considered (Daffner 2007), and magnetic resonance imaging (MRI) may be performed instead of CT in certain situations (e.g. persistent neck pain with normal plain radiography and no neurological signs or symptoms) because of suspicion of ligamentous injury, although there is no evidence to support the clinical utility of MRI in this situation as demonstration of ligamentous oedema or even disruption does not necessarily change planned treatment (collar immobilisation). In adults who attend emergency departments with what is thought to be clinically important cervical spine trauma (this is mostly based on clinical suspicion without consistent application or understanding of decision rules), CT is usually performed first as it is more sensitive and specific than plain radiography in the demonstration of fractures and subluxation or dislocation (Parizel 2010). When CT is not available or when pre-test suspicion is low some people will carry out plain radiography despite its inferior performance (Holmes 2005). The original NEXUS and Canadian C-spine rule studies did not indicate what proportion of patients had CT versus radiography, nor did they mandate CT if imaging was indicated. However, there is evidence that supports CT being more accurate than plain radiography; plain radiography often fails to show the cervicothoracic junction adequately (Holmes 2005; Parizel 2010). Appropriate application of evidence-based clinical decision rules for patients with CSI can help to focus clinical examination and history taking, improve rates of positive imaging, and reduce unnecessary use of imaging that carries with it financial costs for the health system and opportunity costs for other patients who need to access imaging in resource-constrained environments such as public emergency departments, especially at night and on weekends.

**Rationale**

Clinically important CSI following blunt trauma accounts for approximately 2% of the cases of cervical spine injury in emergency departments; however, most patients undergo diagnostic imaging (Stiell 1997). The NEXUS criteria and the Canadian C-spine rule are two clinical decision rules available to assist emergency clinicians to evaluate the need for imaging in patients with CSI. These rules have the potential to rule out a clinically important cervical spine injuries and therefore reduce the number of unnecessary imaging in these patients. A review conducted in 2012 (Michaelf 2012) concluded that the Canadian C-spine rule has better diagnostic accuracy than the NEXUS criteria; however we are aware of new studies published since the search date of this review (Goode 2014; Griffith 2013; Griffith 2014; Matteucci 2015; Morrison 2014). Therefore, this new Cochrane review will be an update of the previous systematic review (Michaelf 2012).
OBJECTIVES
To describe and compare the diagnostic accuracy of the Canadian C-spine rule and the National Emergency X-Radiography Utilization Study (NEXUS) to screen for clinically important cervical spine injury (CSI) in patients following blunt trauma.

METHODS
Criteria for considering studies for this review

Types of studies
We will consider prospective cohort or cross-sectional studies if they compare the results of the Canadian C-spine rule or NEXUS with an appropriate reference standard. Only studies that enrolled clinical populations where there is diagnostic uncertainty will be included in the review. We will only include results from full-text reports that include sufficient raw data to allow reconstruction of contingency tables. If studies have been published as an abstract or conference proceeding, full-text publications will be sought, or alternatively we will contact authors for their data where possible. We will also contact the authors for data in case we are not able to construct the 2 x 2 tables from the reports. Studies published in languages other than English will be included if translations can be obtained.

Participants
We will include studies that assess the diagnostic accuracy of the Canadian C-spine rule or NEXUS in adults presenting with cervical spine injury after blunt trauma. Clinically important CSI is defined as fracture, dislocation or mechanical instability of the cervical spine which requires specialist intervention (e.g. immobilisation or surgical intervention) (Stiell 1999; Stiell 2001). Blunt trauma results from an impact to the body in the absence of any penetrating trauma; examples of blunt trauma include motor vehicle accident, fall or assault.

We will include studies carried out in all settings (i.e. hospital emergency departments and general practice) and will include studies in which medically trained and qualified individuals as well as nurse practitioners and allied health professionals have undertaken assessment of participants using either of the two clinical decision rules).

Index tests
We will include studies that evaluated the diagnostic accuracy of either the Canadian C-spine rule (Stiell 2001) or NEXUS rule (Hoffman 1998), or both to rule out a potentially serious CSI. Both instruments are clinical decision rules commonly used to decide whether or not diagnostic imaging is needed to prove or rule out CSI after blunt trauma. Since they have slightly different inclusion criteria (i.e. the C-spine rule is not applicable for those with Glasgow Coma Scale less than 15 and NEXUS recommends imaging for those patients), we will assess the diagnostic accuracy of each of the rules when applied in accordance with their derivation.

Target conditions
Clinically important cervical spinal injury (CSI) after blunt trauma.

Reference standards
We will include studies if the diagnostic rule results were compared to results of diagnostic imaging procedures such as plain radiographs, and computed tomography (CT) to confirm the presence of cervical spine fracture, dislocation or mechanical instability. Imaging is regarded as a valid reference standard. For the development of both tools, plain radiography was used as a reference standard, unless CT or magnetic resonance (MRI) was performed (Hoffman 2000; Stiell 2001).

We will also include studies that image some patients and clinically followed the remainder as a reference standard but will conduct a sensitivity analysis (if possible) to investigate the effect of the use of this imperfect reference standard on review results. An example of this approach is the 14-day proxy method, where the assessing clinician elects to image patients based upon their clinical judgment and the remaining patients are contacted by a registered nurse 14 days after discharge and asked questions about pain and return to function (Vandemheen 1999). A positive response to these questions results in patients being asked to return to hospital for imaging investigations.

Search methods for identification of studies

Electronic searches
We will search the following databases from their inception.
- MEDLINE (OvidSP)
- Embase (OvidSP)
- Cumulative Index to Nursing and Allied Health Literature (CINAHL) (EBSCO)
- Latin American and Caribbean Health Sciences Literature (LILACS) (Bireme)

The search strategy for the MEDLINE database is presented on Appendix 1. We will also search ClinicalTrials.gov and the World Health Organization's International Clinical Trials Registry Platform (WHO ICTRP) for trials registry and protocols.

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Searching other resources
We will also search the reference lists of included studies and previous relevant reviews for potentially relevant studies.

Data collection and analysis

Selection of studies
Two review authors will independently screen titles and abstracts from the search results. The full-text publications of the potentially eligible studies will be retrieved and independently assessed for inclusion. Disagreements will be resolved by consensus or arbitration by a third review author. No language restrictions will be applied and translations will be sought where possible.

Data extraction and management
Two review authors will independently perform the data extraction. Disagreements will be resolved by consensus or arbitration by a third review author when required. We will extract the following data: characteristics of studies (country, recruitment modality, source of funding, risk of bias), study design, characteristics of participants (age, gender, duration of symptoms, number of participants, including number receiving the index test and reference standard), type of index tests and reference standards, including the methods of execution; experience, expertise, and training of the assessors, and the frequency of true positives, true negatives, false positives and false negatives for the index to the reference test in order to create a 2 x 2 table for each included study. We will contact the authors in case of incomplete or missing data.

Assessment of methodological quality
The methodological quality of included studies will be assessed using a modified version of the Quality Assessment of Diagnostic Accuracy Studies tool (QUADAS-2) (Whiting 2011; Slaar 2017; Wade 2013). The QUADAS-2 tool assesses methodological quality based on four domains: (1) patient selection, (2) index test, (3) reference standard, and (4) flow and timing. The developers of QUADAS-2 recommend that the tool can be tailored for each specific review by adding or omitting signalling questions to assist judgements. The signalling questions related to risk of bias and applicability that will be used in this review are described in Table 2. Studies judged as ‘no’ or ‘unclear’ for one or more domains are considered as having concerns regarding applicability (Whiting 2011). Two review authors will independently assess the methodological quality of the included studies. Disagreements will be resolved by discussion and, if necessary, arbitrated by a third review author.

Statistical analysis and data synthesis
We will generate diagnostic 2 x 2 contingency tables to record true positives, true negatives, false positives and false negatives for each study, and calculate sensitivity, specificity and the 95% confidence interval (CI) for each index test. To estimate the summary sensitivity and specificity we will perform a meta-analysis using the bivariate logistic model (Reitsma 2005). We expect that all studies share the same criteria for test positivity for each of the two tests (positivity is defined with the same threshold across studies); thus no issues of multiple thresholds reported are expected to rise. For studies that directly compared both tests, we will perform direct comparison, but comparison will not be limited to direct comparisons. If data are sufficient and adequate, we will indirectly compare the two tests in relation to their sensitivities or specificities. Test comparison will be performed by adding covariates for different types of index tests into the model, and testing the significance (significance level = 0.05) of the parameters of covariate’s. Analyses will be performed using STATA and Review Manager.

Investigations of heterogeneity
We will investigate factors that may contribute to heterogeneous results in a subgroup analysis, such as healthcare setting (e.g. emergency room versus primary care), different health professionals (e.g. medical specialists versus nurses), age of patient population, and the influence of delayed verification. We will also perform a sensitivity analysis for study quality (i.e., QUADAS-2) to investigate methodological heterogeneity. We will use forest plots and sensitivities and specificities plotted using an HSROC curve for visual examination of heterogeneity between studies, and add covariates (i.e. settings, health professional, age, delayed verification, and QUADAS items) to investigate the heterogeneity between studies in the meta-analysis.

Sensitivity analyses
As reported above we will conduct both sensitivity and subgroup analyses to evaluate the impact of methodological quality, healthcare setting and healthcare professional, population (e.g. age group) and reference standard (e.g. 14 day proxy) has on the performance of the index tests.
REFERENCES

Additional references

ACEM / RANZCR Guidelines 2012

Ackland 2012

ACR Appropriateness Criteria

Como 2009

Daffner 2007

Davis 1993

Diagnostic Imaging Pathways

Goergen 2015

Goode 2014

Griffith 2011

Griffith 2013

Griffith 2014

Hasler 2012

Hoffman 1998

Hoffman 2000

Holmes 2005

Matteucci 2015

Milby 2008

Morrison 2014

Motor Accidents Authority 2014
NICE Guidance 2007

Niska 2010

Parizel 2010

Reitsma 2005

Slaar 2017

Stiell 1997

Stiell 1999

Stiell 2001

Stiell 2003

The College of Emergency Medicine 2010

TRACs 2008

Vandemheen 1999

Wade 2013

Whiting 2011

References to other published versions of this review
Michaleff 2012

* Indicates the major publication for the study

ADDITIONAL TABLES

Table 1. The NEXUS Criteria

<table>
<thead>
<tr>
<th>Criteria</th>
<th>Explanations</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. No posterior midline cervical tenderness</td>
<td>Midline posterior bony cervical spine tenderness is present if the patient complains of pain on palpation of the posterior midline neck from the nuchal ridge to the prominence of the first thoracic vertebra, or if the patient reports pain with direct palpation of any</td>
</tr>
</tbody>
</table>
Table 1. The NEXUS Criteria (Continued)

<table>
<thead>
<tr>
<th>Criteria</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>2. No evidence of intoxication</td>
<td>Patients should be considered intoxicated if they have either of the following: (a) a recent history, by the patient or an observed intoxication or intoxicating ingestion; or (b) evidence of intoxication on physical examination, such as odour of alcohol, slurred speech, ataxia, dysmetria, or other cerebellar findings, or any behavior consistent with intoxication. Patients may also be considered to be intoxicated if tests of bodily secretions are positive for drugs that affect level of alertness, including a blood alcohol level greater than 0.08 mg/dL.</td>
</tr>
<tr>
<td>3. Normal level of alertness</td>
<td>An altered level of alertness can include any of the following: (a) Glasgow Coma Scale score of 14 or less; (b) disorientation to person, place, time, or events; (c) inability to remember 3 objects at 5 minutes; (d) delayed or inappropriate response to external stimuli; or (e) other</td>
</tr>
<tr>
<td>4. No focal neurological deficit</td>
<td>Any focal neurological complaint (by history) or finding (on motor or sensory examination)</td>
</tr>
<tr>
<td>5. No painful distracting injuries</td>
<td>No precise definition for distracting painful injury is possible. This includes any condition thought by the clinician to be producing pain sufficient to distract the patient from a second (neck) injury. Examples may include, but are not limited to, the following: (a) a long bone fracture; (b) a visceral injury requiring surgical consultation; (c) a large laceration, degloving injury, or crush injury; (d) large burns; or (e) any other injury producing acute functional impairment. Physicians may also classify any injury as distracting if it is thought to have the potential to impair the patient’s ability to appreciate other injuries</td>
</tr>
</tbody>
</table>

*If all of these criteria are met, imaging is not required in order to exclude clinically important CSI

Table 2. Assessment of methodological quality: modified version of QUADAS-2

<table>
<thead>
<tr>
<th>Quality domain</th>
<th>Risk of bias</th>
<th>Signalling questions</th>
<th>Applicability</th>
<th>Signalling questions</th>
</tr>
</thead>
<tbody>
<tr>
<td>1: Patient selection</td>
<td>Could the selection of patients have introduced bias? (high/low/unclear)</td>
<td>1) Was a consecutive or random sample of patients enrolled? (yes/no/unclear)</td>
<td>Are there concerns that the included patients and settings do not match the review question? (high/low/unclear)</td>
<td>Were all the patients recruited from the same clinical setting?</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2) Did the study avoid inappropriate exclusions? (yes/no/unclear)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2: Index test</td>
<td>Could the interpretation of the index test have introduced bias? (high/low/unclear)</td>
<td>1) Were the index test results interpreted without knowledge of the results of the reference standard? (yes/no/unclear)</td>
<td>Are there concerns that the index test, its conduct, or the interpretation differ from the review question? (high/low/unclear)</td>
<td>1) Did the study provide a clear definition of what was considered to be a “positive” result for the index test? (high/low/unclear)</td>
</tr>
</tbody>
</table>
Table 2. Assessment of methodological quality: modified version of QUADAS-2  (Continued)

<table>
<thead>
<tr>
<th>3: Reference Standard</th>
<th>4: Flow and timing</th>
</tr>
</thead>
<tbody>
<tr>
<td>Could the interpretation of the reference standard have introduced bias? (high/low/unclear)</td>
<td>Could the patient flow have introduced bias? (low/high/unclear)</td>
</tr>
<tr>
<td>1) Is the reference standard likely to correctly classify the target condition?  2) Were the reference standard results interpreted without knowledge of the results of the index test?</td>
<td>1) Was there an appropriate interval between index test and reference standard (e.g. short enough to be reasonably sure that the target condition did not change between the two tests)?  2) Did all patients receive the same reference standard?  3) Were all patients include in the analysis?  4) Were withdrawals from the study clearly reported?</td>
</tr>
<tr>
<td>Are there concerns that the target condition as defined by the preference standard does not match the review question? (high/low/unclear)</td>
<td>-</td>
</tr>
<tr>
<td>2) Was the index test implemented by the same type of health professional in all patients? (e.g. profession, level of training)</td>
<td>-</td>
</tr>
</tbody>
</table>
Appendix 1. MEDLINE search strategy

1. (NEXUS or CCR).mp.
3. (Canadian c-spine or Canadian cervical spine).mp.
4. ((Cervical spine or c-spine) adj5 clear$).mp.
5. (cervical adj5 (trauma$ or injur$ or fracture$ or sublux$ or dislocat$ or avuls$ or instab$)).mp.
6. (Spinal cord injury without radiographic abnormality or SCIWO A).mp.
7. exp Cervical Vertebrae/
8. exp Neck Injuries/
9. exp Spinal Injuries/
10. exp Spinal Cord Injuries/
11. spinal fractures/
12. 1 or 2 or 3 or 4 [Index tests]
13. 5 or 6 or 7 or 8 or 9 or 10 or 11 [target condition]
14. 12 and 13
DECLARATIONS OF INTEREST

None known.

SOURCES OF SUPPORT

Internal sources

• National Health and Medical Research Council of Australia, Australia.

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External sources

• No sources of support supplied