Canadian Practice Patterns of Venous Thromboembolism Prophylaxis for Adults with Spinal Cord Injury

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Abstract

Context: According to current reviews and guidelines, venous thromboembolism (VTE) prophylaxis in spinal cord injury (SCI) includes low molecular weight heparin (LMWH) in combination with mechanical prophylaxis. The level of evidence for VTE prophylaxis is varied among the recommendations. National best practice consensus is extremely important in the care of patients especially in situations when the evidence are vague and varied.

Objective: To ascertain practice patterns of VTE prophylaxis by Canadian SCI rehabilitation physiatrists in adults admitted to a rehabilitation unit for spinal cord injury rehabilitation.

Method: An invitation to participate in this project was distributed to Canadian SCI Rehabilitation physiatrists through the ‘SCI Hallways’, a private online forum for consultation between Canadian physiatrists.

Results: A total of 10 physiatrists from 8 of 13 Canadian academic rehabilitation programs participated. All participants stated that their practice involved using a form of mechanical VTE prophylaxis and LMWH for 8 to 12 weeks.

Conclusion: Use of VTE prophylaxis for SCI is consistent among Canadian physiatrist and matches guidelines for VTE prophylaxis in spinal cord injury.

Keywords: Thromboembolism; Spinal cord injury; Prophylaxis

Introduction

Venous thromboembolism (VTE) is the third most common cause of death in people with spinal cord injury (SCI) [1]. VTE encompasses both deep vein thrombosis (DVT) and pulmonary embolism (PE). The incidence of DVT among people with SCI ranges from 49% to 100%, with the most significant risk occurring in the first two weeks post-injury [2]. Up to 50% of DVTs can lead to a PE, which is potentially life threatening. Therefore, it is vital for clinicians to implement effective prophylaxis to prevent this potentially fatal complication.

Many studies have attempted to clarify the ideal form of prophylaxis and the level of evidence for the recommendations are varied [3-5]. Current guidelines and recommendations stem from three main sources: Antithrombotic Therapy and Prevention of Thrombosis 9th edn. by the American College of Chest Physicians in 2012 (this is not specific for SCI), Spinal Cord Injury Rehabilitation Evidence (SCIRE) systematic review in the journal Archives of Physical Medicine and Rehabilitation from 2009, and the Paralyzed Veteran’s Association Guidelines from 1999 [3-5].

These three publications recommend two methods of VTE prophylaxis in people with SCI: mechanical and pharmacological [3-5] (See Table 1 for a summary of the recommendations from each publication).

The care of SCI patients, after their acute injury and associated surgeries, are usually under the responsibility of physiatrists. Canada is a geographically large country with multiple major rehabilitation centers spread throughout. Because of the large geographical distance between rehabilitation centers, and the variations in the level of evidence within VTE prophylaxis recommendations, a survey was conducted to assess the current Canadian practices for VTE prophylaxis for people with SCI.

<table>
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<tr>
<th>ACP 2012</th>
<th>SCIRE 2009</th>
<th>PVA 1999</th>
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Pharmacological Prophylaxis

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<tr>
<th>ID #</th>
<th>Pharmacologic Prophylaxis</th>
<th>Mechanical Prophylaxis</th>
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<tbody>
<tr>
<td>1</td>
<td>Enoxaparin</td>
<td>8-12 weeks</td>
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<tr>
<td>2</td>
<td>Dalteparin</td>
<td>8-12 weeks</td>
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<td>3</td>
<td>Dalteparin</td>
<td>8-12 weeks</td>
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<tr>
<td>4</td>
<td>Dalteparin</td>
<td>8-12 weeks</td>
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Methods

The question, “What are you using for VTE and PE prophylaxis at your center?” was posted on a private online forum called “SCI Hallways” whose members consist exclusively of Canadian physiatrists engaging in SCI rehabilitation. Over the course of two weeks, various Canadian physiatrists posted their responses on the forum. The majority of the responses consisted of the specific name of the pharmacological and mechanical prophylaxis, but given the nature of using an online forum, five participants gave vague responses. These participants posted that they used an unspecified LMWH, an unspecified mechanical form of prophylaxis and or just posted "same as above". These five participants were contacted directly via email for clarifications.

Given that there are only a handful of physiatrists involved in inpatient SCI rehabilitation, it can be difficult to organize a face-to-face meeting to discuss local practice patterns. The “SCI Hallways” is an online forum used to bridge geographical distance, allowing Canadian clinicians who treat people with SCI to engage in discussion in a convenient way. In addition to aiding in facilitating a national consensus, it has also been used regularly to assist physicians in clinical decision making and with research. Most academic Canadian physiatrists engaging in SCI rehabilitation have access to, and have previously participated on this forum. Responses can be accessed by members at any time.

Results

There are a total of thirteen academic centers across Canada with physicians engaging in SCI rehabilitation [6]. Some centers have dedicated SCI rehabilitation wards, while others admit patients needing SCI rehabilitation into a general rehabilitation ward. Ten physicians from eight centers participated in this practice pattern assessment. (See Figure 2 for a flow chart of the summary of participation and data acquisition).

Table 1: Summary of recommendations from the three sets of recommendations for VTE prophylaxis in adults with SCI. Each journal used a different grading system for the level of evidence. In the recommendations from the American College of Chest Physicians, Grade 2C corresponds to weak recommendations based on low-quality evidence. In the recommendations from SCIRE, Level 1a evidence corresponds to data from 2 randomized control trials and Level 4 corresponds to data from pre-post study and case series. In the recommendations from PVA, Level I correspond to data from large randomized trials with definitive results, and Level II corresponds to data from small randomized trials with uncertain results.

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<th>Pharmacologic Prophylaxis</th>
<th>Mechanical Prophylaxis</th>
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<tr>
<td></td>
<td>LMWH or UH (Grade 2C)</td>
<td>LMWH or UH (Level II)</td>
</tr>
<tr>
<td>Duration</td>
<td>3 months</td>
<td>8-12 Weeks</td>
</tr>
<tr>
<td>Mechanical Prophylaxis</td>
<td>Preferably IPCD (Grade 2C)</td>
<td>IPCD or TEDS (Level I)</td>
</tr>
<tr>
<td>Duration</td>
<td>Unspecified</td>
<td>2 Weeks</td>
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</table>

Table 2: Summary of participation in survey and data acquisition. Response rate was 69% of all Canadian academic rehabilitation centers.

All physicians that participated use mechanical prophylaxis. One participant did not specify the type they use. Of the other nine, one used intermittent pneumatic compression devices (IPCD) until starting LMWH and then switched to thrombo-embolic deterrent stockings (TEDS), while the other eight used TEDS with IPCD in the acute setting for up to 2 weeks and then transitioned to TEDS only (Table 2). Ideally, both IPCD and TED stockings are worn around the clock. However, certain activities cannot be done with these on, such as during maintenance of hygiene and skin integrity assessments.

All physicians that participated use pharmacological prophylaxis with low molecular weight heparin (LMWH). Six used dalteparin exclusively, one enoxaparin exclusively, and three did not specify (Table 2).
Table 2: Tabulated results from respondents with names substituted with identification numbers. LMWH stands for low molecular weight heparin. IPCD stands for intermittent pneumatic compression devices. TEDS stands for Thrombo-Embolic Deterrent Stockings also known as Thrombo-Embolic Deterrent Hose.

Discussion

Relevance of guidelines

The practice patterns of Canadian SCI rehabilitation physicians reflect recommendations that state that there is evidence, although inconsistent, for both mechanical and pharmacological prophylaxis for patients with SCI [3-5]. The inconsistency may be because of the difference between the inclusion and exclusion criteria used by the authors to generate their conclusions. (See Figures 2 and 3 for comparison of patient population for pharmacologic and mechanical prophylaxis used in the three papers).

The SCIRE recommendations published in 2009 is the most recent of the three publications and reviewed studies exclusively in the SCI patient population. The SCIRE recommendations suggest that there is strong evidence for LMWH, but poor evidence for mechanical prophylaxis [4]. The guideline from American College of Chest Physicians in 2012 is not directed at only patients with SCI, but at all non-orthopedic major trauma patients (including traumatic brain injury and spinal surgery) [5]. Additionally, the guideline from PVA (Paralyzed Veteran's Association) in 1999 is based on research from the 1990s [3]. From our correspondence with the PVA, they have currently stopped distributing their guideline and are in the process of producing an updated guideline from more current literature.

The three guidelines and recommendations derive their conclusion from a variety of sources with minor overlap between them. For pharmacological prophylaxis, 4 studies were included in all three papers. These 4 studies were all published on or prior to 1990 [7-10]. The recommendations from the American College of Chest Physicians are derived from a total of 19 studies with 11 of the 19 studies done in the non-SCI population [5]. The recommendations from SCIRE were

<table>
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<tr>
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<th>Drug Type</th>
<th>Duration</th>
<th>Prophylaxis Type</th>
<th>Duration</th>
<th>LMWH Prophylaxis</th>
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<tr>
<td>5</td>
<td>Dalteparin</td>
<td>8-12 weeks</td>
<td>IPCD + TEDS for 2 weeks</td>
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<td>TED Stocking for duration of LMWH</td>
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<tr>
<td>6</td>
<td>Dalteparin</td>
<td>8-12 weeks</td>
<td>IPCD + TEDS for 2 weeks</td>
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<td>TED Stocking for duration of LMWH</td>
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<tr>
<td>7</td>
<td>Dalteparin</td>
<td>8-12 weeks</td>
<td>IPCD + TEDS for 2 weeks</td>
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<td>TED Stocking for duration of LMWH</td>
</tr>
<tr>
<td>8</td>
<td>Unspecified LMWH</td>
<td>8-12 weeks</td>
<td>IPCD + TEDS for 2 weeks</td>
<td></td>
<td>TED Stocking for duration of LMWH</td>
</tr>
<tr>
<td>9</td>
<td>Enoxaparin or Dalteparin</td>
<td>8-12 weeks</td>
<td>IPCD only until LMWH started</td>
<td></td>
<td>TED Stocking for duration of LMWH</td>
</tr>
<tr>
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<td>Unspecified LMWH</td>
<td>8-12 weeks</td>
<td>Unspecified</td>
<td></td>
<td>Unspecified</td>
</tr>
</tbody>
</table>

Figure 2: Graph of number of publications versus source comparing patient population for pharmacological prophylaxis. SCIRE derived their recommendations exclusively from SCI patients. American College of Chest Physicians included 4 studies on orthopedic patients, and 7 on other mixed surgical patients. PVA included 1 study on major trauma patients.

Figure 3: Graph of number of publications versus source comparing patient population for mechanical prophylaxis of VTE. The SCIRE recommendations published in 2009 is the most recent of the three publications and reviewed studies exclusively in the SCI patient population. The SCIRE recommendations suggest that there is strong evidence for LMWH, but poor evidence for mechanical prophylaxis [4]. The guideline from American College of Chest Physicians in 2012 is not directed at only patients with SCI, but at all non-orthopedic major trauma patients (including traumatic brain injury and spinal surgery) [5]. Additionally, the guideline from PVA (Paralyzed Veteran's Association) in 1999 is based on research from the 1990s [3]. From our correspondence with the PVA, they have currently stopped distributing their guideline and are in the process of producing an updated guideline from more current literature.
derived from 13 studies and the recommendations from PVA were derived from 6 studies with the newest study from 1995 [3,4].

There is even less overlap between the sources used by the major three recommendations for mechanical prophylaxis [3-5]. There are zero studies that all three recommendations cited [3-5]. The only overlap of sources occurs in the SCIRE and PVA guidelines for two studies with one done in 1982 and the other in 1992 [11,12]. Additionally, the primary sources all evaluated a vast number of interventions (including range of motion exercises, massage, bandages, rotating tables, electrical stimulation, venous foot pump, compression stockings, intermittent compression devices) and various patient populations (including SCI, general surgery, orthopedic surgery, neurosurgery, cardiac surgery and abdominal surgery) [10-39].

**Mechanical prophylaxis**

The options for mechanical prophylaxis includes IPCDs and TED stockings [3-5]. IPCDs are inflatable cuffs placed around the lower limbs that are intermittently inflated via an electrically powered pneumatic pump [40]. TED stockings are compression stockings with a decreasing gradient of pressure in a distal to proximal direction [40]. IPCDs simulate the actions of lower limb muscles to intermittently force the blood in the deep veins back to the heart, and TED stockings increase the velocity of blood flow and prevents the activation of extrinsic coagulation pathway from the venous distention exposing subendothelial tissue 40. One of the biggest advantages of mechanical prophylaxis is the cost effectiveness and the infrequent and minor adverse events (consisting of foot abrasions, superficial thrombophlebitis, and subjective warmth) [41].

The SCIRE review stated that there is Level 4 evidence (weak) for all forms of mechanical prophylaxis [4]. SCIRE evidence is based on two studies which although showing a positive effect for mechanical prophylaxis, were weak (a case series in 2001 and a pre-post study in 1999) [4]. The guideline from American College of Chest Physicians in 2012 stated that the evidence is grade 2C (recommendations based on low-quality evidence) [5]. American College of Chest Physician’s guideline is derived from multiple studies from major trauma patients and all were described as having various limitations [5]. The guideline from PVA in 1999 stated that the evidence is level I (data from large randomized trials with definitive results) [3]. The guidelines from PVA were based on studies from the 1970s and 1980s, including studies involving orthopedists, neurosurgeons, cardiac surgery and minor surgeons, and outdated forms of mechanical prophylaxis (range of motion exercises, massage, electrical stimuli, and venous foot pump) [3].

The three recommendations had varying grades of evidence for mechanical prophylaxis [3-5]. However, the options, especially TED stockings, have mild and infrequent side effects [40]. In our study, all participants stated that SCI patients used TED stockings and IPCDs during the initial 2 weeks and continued with the TED stockings for the duration of needing LMWH.

**Pharmacological Prophylaxis**

The SCIRE review states that there is Level 1a evidence (strong) for LMWH being superior to unfractionated heparin (UH) and that all LMWH are equally efficacious 4. However, guidelines from American College of Chest Physicians in 2012 and from PVA in 1999 state that the evidence for pharmacological prophylaxis is not strong: Grade 2C (weak recommendations based on low-quality evidence) and Level II (data from small randomized trials with uncertain results, although this statement is relatively outdated), respectively [3,5] (Table 1). The majority of studies analyzed by the American College of Chest Physicians were in non-SCI patients and all the studies by PVA were done prior to 1995 (Table 2).

As reflected by the data, the respondents all use a variety of LMWHs. There was a variety of responses with the majority (60%) using Dalteparin. We suspect that because there is no data in literature suggesting one type of LMWH is superior to another, the rationale for choosing the specific LMWH is based on the hospital’s pharmacy formulary. Interestingly, one clinician stated that he/she choose prophylaxis based on previous experiences, and used Dalteparin for non-traumatic SCI patients and Enoxaparin for traumatic SCI patients.

**Limitations of the Study**

Limitations of this study include a possible reporting bias due to the transparent nature of the study design, and a selection bias due to this study being voluntary.

Additionally, the guidelines for the timeframe for VTE prophylaxis are vague or unspecified. The only recommendations for the timeline come from PVA and they suggest a total of 8-12 weeks [3]. This range means that there are four weeks of potential variability that are up to the clinician's own discretion and interpretation of a patient's risk for VTE.

Finally, only physiatrists participated in this study and the trauma team, surgeons, and the intensivists were not involved. Because physiatrists are predominantly involved after a patient has been stabilized in the surgical and trauma units, they are not intimately involved in the decision making process for VTE prophylaxis during a patient's acute admission.

**Conclusion**

Use of pharmacological VTE prophylaxis for SCI is consistent among Canadian physiatry physicians and matches current guidelines for VTE prophylaxis in spinal cord injury. Although there is inconsistency in the level of evidence for mechanical VTE prophylaxis in current recommendations, these physicians all use mechanical VTE prophylaxis for SCI.

**Acknowledgments**

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**References**


6. Program descriptions. CarMS.


