Supporting chronic pain management across provincial and territorial health systems in Canada: Findings from two stakeholder dialogues

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Accessibility
Supporting chronic pain management across provincial and territorial health systems in Canada: Findings from two stakeholder dialogues

Michael G Wilson PhD1,2,3, John N Lavis MD PhD1,2,3,4,5, Moriah E Ellen MBA PhD6,7

BACKGROUND: Chronic pain is a serious health problem given its prevalence, associated disability, impact on quality of life, and the costs associated with the extensive use of health care services by individuals living with it.

OBJECTIVE: To summarize the research evidence and elicit health system policymakers', stakeholders' and researchers' tacit knowledge and views about improving chronic pain management in Canada and engaging provincial and territorial health system decision makers in supporting comprehensive chronic pain management in Canada.

METHODS: For these two topics, the global and local research evidence regarding each of the two problems were synthesized in evidence briefs. Three options were generated for addressing each problem, and implementation considerations were assessed. A stakeholder dialogue regarding each topic was convened (with 29 participants in total) and the deliberations were synthesized.

RESULTS: To inform the first stakeholder dialogue, the authors found that systematic reviews supported the use of evidence-based tools for strengthening chronic pain management, including patient education, self-management supports, interventions to implement guidelines and multidisciplinary approaches to pain management. While research evidence about patient registries/treatment-monitoring systems is limited, many dialogue participants argued that a registry/system is needed. Many saw a registry as a precondition for moving forward with other options, including creating a national network of chronic pain centres with a co-ordinating 'hub' to provide chronic pain-related decision support and a cross-payer, cross-discipline model of patient-centred primary health care-based chronic pain management. For the second dialogue, systematic reviews indicated that traditional media can be used to positively influence individual health-related behaviours, and that multistakeholder partnerships can contribute to increasing attention devoted to issues on policy agendas. Dialogue participants emphasized the need to mobilize behind an effort to build a national network that would bring together existing organizations and committed individuals.

CONCLUSIONS: Developing a national network and, thereafter, a national pain strategy are important initiatives that garnered broad-based support during the dialogues. Efforts toward achieving this goal have been made since convening the dialogues.

Key Words: Canada; Chronic pain; Dialogue; Health systems; Pain management

Chronic pain is a serious health problem given its prevalence, associated disability, impact on quality of life, and the costs associated with extensive use of health care services (1-5). Approximately one in five Canadian adults experience chronic pain (5,6), but it often goes unrecognized and/or is undertreated (7,8). Prevalence increases with age, with some estimates indicating that as many as 65% of community-dwelling older adults and 80% of those living in long-term care facilities experience chronic pain (5,6). Furthermore, quality of life for
individuals with chronic pain has been found to be lower than for those with most other chronic diseases (1). The financial impact of chronic pain in terms of health care expenditures and productivity costs has been estimated to be $56 to $60 billion per year in Canada (9).

To support and provide better care for Canadians affected by chronic pain, there is a need to identify how provinces and territories across the country may individually, as well as collectively, strengthen chronic pain management. One approach to addressing health system issues such as chronic pain is to convene stakeholder dialogues, in which the overriding objective is to support evidence-informed policymaking by pairing the best available research evidence with a robust deliberative process that gives voice to the tacit knowledge and real world views and experiences of those involved in and/or affected by the issue. In general, stakeholder dialogues convene health system stakeholders (eg, government officials, professional and community leaders, patients/citizens/groups representing them and researchers) for deliberations with the goal of supporting participants to champion creative efforts to address a pressing health system problem within their respective constituencies (10). Specifically, dialogues provide stakeholders with the opportunity to bring their tacit knowledge and their own views and experiences to bear on a pressing health system problem, options that would address it and consideration of implementation issues. Each dialogue was informed by an evidence or issue brief that mobilized the best available research evidence about each of these components (an issue brief uses the same approach as an evidence brief, but draws on findings from a previously conducted synthesis of the evidence) (10).

To foster these efforts, the Community Alliances for Health Research and Knowledge Translation on Pain partnered with and provided funding to the McMaster Health Forum (www.mcmaster-healthforum.org) in 2009. It was to act as a neutral convenor for a stakeholder dialogue focused on strengthening chronic pain management in Canada. This dialogue was followed by another in April 2011 (again with funding from CAHR-pain to the McMaster Health Forum) to build on a key finding from the first dialogue – the need to more systematically engage health system decision-makers in supporting comprehensive chronic pain management in provincial and territorial health systems in Canada. Specifically, the challenges of engaging policymakers in the first dialogue led to broad-based consensus among participants that long-term sustainable action is constrained by a lack of attention devoted to chronic pain by health system decision-makers. In the present article, we present the key findings from the two stakeholder dialogues and the briefs that were prepared to inform them (11,12). We also provide an overview of examples of progress that have been made since the dialogues were convened to draw attention to the types of actions that have been or are being taken to address this pressing health system issue.

METHODS

The two stakeholder dialogues were convened on December 9, 2009 and April 11, 2011, in Hamilton, Ontario. Briefs sent to participants in advance were prepared by the McMaster Health Forum working in collaboration with an interdisciplinary steering committee. The methods used for preparing the briefs and convening the dialogues are described below. A detailed article describing evidence briefs and stakeholder dialogues is also available for those interested in more detail about the approach (10).

Preparing the evidence and issue briefs

Each of the briefs was prepared through four steps. First, a steering committee comprising representatives from partner organizations and stakeholder groups was convened. The role of the steering committee was to engage with the McMaster Health Forum to provide guidance and expert advice across all stages of the process. In collaboration with the steering committee, terms of reference were developed for each of the briefs. These provided a preliminary outline framing the problem, three options for addressing it and implementation considerations. In the second step, key informant interviews were conducted (nine for the evidence brief and 11 for the issue brief) with policymakers, managers (eg, from health regions, health care institutions and community-based organizations), stakeholders (eg, from interest groups, provider associations or other stakeholder groups) and researchers, who were actively engaged in the issue of chronic pain. The terms of reference were iteratively revised based on feedback from the key informants and the steering committee and then used to guide the writing of each brief. The key informants were also asked to identify literature that would be relevant to preparing the briefs.

Third, for each brief, relevant research evidence regarding the problem, options and implementation considerations was identified, selected, appraised and synthesized. Whenever possible, research evidence was drawn from systematic reviews and, occasionally, from single studies when reviews were not identified. Published literature was identified by searching PubMed using the health services research search filters for appropriateness, process assessments, outcomes assessments and qualitative research. In addition, grey literature was searched for by reviewing the websites of a number of Canadian and international organizations (13-21). To identify research evidence about the three options in each of the briefs, Health Systems Evidence (www.healthsystemsevidence.org) was searched. Health Systems Evidence is a continuously updated database, which in January 2015 contained >4200 systematic reviews and >2200 economic evaluations of health service delivery, including consideration of financial and governance arrangements within health systems. Health Systems Evidence identifies documents from several sources including the Cochrane Database of Systematic Reviews (for systematic reviews of effects) and the Centre for Reviews and Dissemination (for systematic reviews of effects and economic evaluations) (22). The reviews and economic evaluations were identified by searching Health Systems Evidence for chronic pain in the title and abstract and by searching topic categories addressing features of each of the options.

The searches were reviewed for relevance by the lead author of each brief (JNL for the first brief and MGW for the second brief). For each systematic review, the focus of the review, key findings, the last year the literature was searched, the methodological quality (based on AMSTAR [23] ratings that are provided for all reviews contained in Health Systems Evidence), the proportion of included studies that were conducted in Canada and the proportion of included studies focused on chronic pain were extracted. For any reviews that had not been previously quality appraised using AMSTAR, two reviewers independently completed an assessment.

Fourth, key findings in the form of an evidence brief (for the first dialogue) and an issue brief (for the second dialogue) were synthesized. Specifically, the briefs were drafted in such a way as to present concisely and in accessible language the global and local research evidence. The final version of the briefs consisted of a one-page summary of key messages followed by a more detailed description of: the problem; three options (including the benefits, harms and costs of the options as well as key elements of and stakeholder experiences with them); and possible barriers to implementation of the options at the levels of individuals, providers, organizations and systems. A merit review process was then undertaken for each brief with a small number of policymakers, stakeholders and researchers to ensure each brief’s system relevance and scientific rigour.

Convening the stakeholder dialogues

Working collaboratively with the steering committee, health system stakeholders were identified (ie, government officials, professional and community leaders, groups representing people living with chronic pain and other stakeholders, as well as researchers). Participants were invited who had the ability to: bring unique views and experiences to bear on the challenge and learn from the research evidence and from others’ views and experiences; and champion within their respective constituencies actions that would address the challenge creatively. Participants were identified by reviewing government directories and the websites of relevant organizations and from suggestions provided

Table 1. Many options were available to address the issues. To provide a level playing field in terms of background information and time, we did not need to be devoted to reviewing the detailed contents of the brief. The goal was not to aim for consensus but rather to provide a space where diverging opinions could be shared and discussed and to identify where synergistic efforts among stakeholders could address the problem might be possible. In addition, each dialogue followed the Chatham House rule (ie, information used during the meeting may be used, but neither the identity nor the affiliation of participants were to be revealed). Finally, the dialogues were not recorded but notes were taken by the facilitator and students assisting with each dialogue. These notes were used to draft summaries of each dialogue that highlighted the key themes that emerged during each deliberation, points of disagreement or general consensus, and the types of action that participants thought could be taken following the dialogue (the identities of participants were kept confidential in the dialogue summaries).

### RESULTS

**Topic 1: Supporting chronic pain management across provincial and territorial health systems in Canada**

The first stakeholder dialogue addressed the issue of chronic pain management across Canada. We present below a summary of the key findings from the evidence brief and the key themes of the deliberations. For those interested in additional information, the evidence brief (12) and dialogue summary (24) are available on the McMaster Health Forum website (www.mcmasterhealthforum.org).

**Key findings from the evidence brief:** The challenge of strengthening chronic pain management in provincial and territorial health systems can be understood by considering four sets of inter-related issues, outlined along with a summary of contributing factors in Table 1. Many options were available to address the issues. To promote discussion about the pros and cons of potentially viable options, we selected three, which we outline in Table 2, along with a summary of key findings. Finally, we identified implementation barriers at the level of individuals, care providers and systems, as well as possible strategies to address the barriers, which we outline in Table 3.

**Summary of dialogue 1:** The dialogue brought together a diverse group of 13 stakeholders (two policymakers/managers, three health care provider association/group representatives, four researchers and four other stakeholders) from across Canada. The group was smaller than our target size of 18 to 22, largely due to the difficulty
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### TABLE 2
Three options for better supporting chronic pain management*

<table>
<thead>
<tr>
<th>Option</th>
<th>Option focus and elements</th>
<th>Summary of key findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Create a model patient registry/treatment-monitoring system in a single jurisdiction</td>
<td>• This option focuses on: o identifying what services are being offered to whom (i.e., what types of patients), by whom (e.g., what disciplines), and how frequently (with appropriate attention to privacy concerns); o identifying both under- and over-utilization; o monitoring efforts to improve service delivery and evaluating their impacts; and o publicly reporting opportunities for improvement • To further understand this option, it is useful to consider it according to four key health system elements: o patient registries; o treatment-monitoring systems; o privacy issues pertaining to patient registries/treatment-monitoring systems; and o public reporting of aggregated data</td>
<td>• Substantial uncertainty exists regarding this option’s benefits and potential harms</td>
</tr>
<tr>
<td>2. Create a national network of centres with a coordinating ‘hub’ to provide chronic pain-related decision-support</td>
<td>• The function/focus of a coordinating hub would be to: o analyze data about treatment patterns (which can include the analysis of data from a patient registry/treatment-monitoring system, which was the focus of the preceding option); o synthesize research evidence; o develop and disseminate resources and tools to support self-management; o develop and disseminate clinical practice guidelines and other resources and tools to support providers and organizations in prevention, early identification, and ongoing treatment; o offer support to undergraduate professional training programs; o offer continuing professional development and other strategies to support evidence-based care (both for single disciplines and multidisciplinary teams); and o monitor efforts to improve care (across the full range of payers and the full continuum of care, including primary health care, postsurgical care, etc) and evaluate their impacts • This option can be further understood by considering the effectiveness of the tools and resources the hub would provide, including: o tools to support self-management (e.g., education for people living with chronic pain, decision aids, personal health records, peer support and telephone support); o interventions to support the dissemination of clinical practice guidelines; o support for undergraduate professional training programs; o continuing professional development and other strategies to support evidence-based care; and o monitoring efforts to improve care and evaluate their impacts</td>
<td>• No relevant reviews were identified about privacy issues pertaining to a patient registry/treatment-monitoring system • Synthesized research evidence is available to support the use of a range of evidence-based tools and resources: o Two high-quality reviews and one medium-quality review focused on patient education and showed favourable results in terms of pain reduction (however, all reviews are more than five years old). o Other reviews identified some benefits and no harms with respect to other self-management supports (e.g., patient education, decision aids, personal health records, peer support and telephone support) (36–46), interventions to support the implementation of clinical practice guidelines (multi-faceted interventions were found to generally be most effective) (12), and continuing professional development to support evidence-based care (12)</td>
</tr>
<tr>
<td>3. Broker and support the implementation of a cross-payer, cross-discipline model of patient-centred primary health care-based chronic pain management</td>
<td>• This option focuses on rewarding: o quality, such as by re-balancing fee schedules away from procedures and toward payment for the time demands associated with assessment, management, support, and dealing with payers and employers, and by accrediting chronic pain ‘specialist’ providers or centres; and o efficiency, such as by engaging the most cost-effective providers and by providing tiered support from telecommunications to in-person interactions, and through tiered referrals from primary health care to accredited regional multidisciplinary pain clinics • This system redesign has a number of health system elements that each need to be considered: o cross-payer models of patient-centred primary health care-based chronic pain management; o cross-discipline models of patient-centred primary health care-based chronic pain management that address the full spectrum of comprehensive care (e.g., prevention, early intervention, treatment, management and rehabilitation); o rewards for quality and efficiency in primary health care; o fee schedules that consider the time demands associated with primary and secondary prevention, treatment, management and rehabilitation, as well as dealing with payers and employers; o accrediting chronic pain ‘specialist’ providers or centres; o engaging the most cost-effective providers; and o providing tiered support from telecommunications to in-person interactions and through tiered referrals from primary health care to accredited regional multidisciplinary chronic pain management centres</td>
<td>• No reviews were identified that relate directly to the concept of a hub. • Several reviews relate to cross-discipline models of care. o Three medium-quality reviews and one high-quality review that relate to multidisciplinary approaches to pain management found medium to strong evidence for improvements in patient function (47) o Another recent (2009) high-quality review showed no difference in patient outcomes between those receiving multidisciplinary rehabilitation and those in control groups (43) • No relevant reviews were identified about cross-payer models of patient-centred primary health care</td>
</tr>
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</table>

*The findings in this table are based on what was available at the time of publication of the evidence brief (December 2009)
Participants generally agreed that effective chronic pain management dialogue participants argued that "no recognition of chronic pain as a significant stigma was associated with chronic pain and highlighted the need for a follow-up dialogue about how to engage policymakers/managers from across the country. This approach could be informed by the work done in Australia, which was the first country in the world to develop a national strategy and framework for the community-based pain services.

Option 1 (create a model patient/registry/treatment-monitoring system in a single jurisdiction)
- Operational challenge in defining eligibility for a condition that lacks an 'event', widely agreed diagnostic criteria and demonstrated pathology
- Collection of individual-level data may compromise an individuals' privacy and lead to stigmatization
- Individuals often have more than one diagnosis (e.g., chronic pain and arthritis) that requires monitoring

Option 2 (create a national network of centres with a coordinating 'hub' to provide chronic pain-related decision-support)
- Resources (time and money) are required to meaningfully involve people living with chronic pain in the development and evaluation of medical device technology

Option 3 (implement a cross-payer, cross-discipline model of patient-centred primary health care-based chronic pain management)
- Individuals with chronic pain may need more specialized and urgent pain management for acute exacerbations or injuries than may be feasible in primary health care

Care provider Option 1 (create a model patient/registry/treatment-monitoring system in a single jurisdiction)
- Primary health care providers will require training and support in how to use the registry/monitoring system
- Primary health care providers may perceive decision supports as a threat to their professional authority
- Health care providers, particularly physicians, have to ensure that a patient-centred primary health care model is integrated with specialty and community-based pain services
- Chronic pain has not traditionally been considered a chronic disease to be managed by primary health care providers

Organization Option 1 (create a model patient/registry/treatment-monitoring system in a single jurisdiction)
- Organizations must be sensitive to the personal health information being collected and how it is used.
- All clinical and non-clinical members of health care teams need to be aware of the processes that need to be in place for effective use of information technologies
- Collaborative work arrangements need to be established and maintained between primary healthcare organizations, secondary and tertiary care organizations that can support these primary healthcare organizations (e.g., chronic pain management programs in academic health science centres), and payers.

System Option 1 (create a model patient/registry/treatment-monitoring system in a single jurisdiction)
- Resources must be in place to ensure sustainability of information systems in the longer term.
- The need for public health surveillance must be balanced with the need for individual privacy.

Possible strategies to address the barriers
- Initiate a national stakeholder-engagement process to raise awareness of health system issues within the chronic pain community and to raise awareness of chronic pain issues within the health policy and systems community
- A similar approach in Canada might mobilize evidence from sources including (but not limited to) academic pain centres, interest groups representing people living with chronic pain and the public, health provider associations and other sources.

*The information in this table is based on what was available at the time of publication of the evidence brief (December 2009)*

<table>
<thead>
<tr>
<th>Levels</th>
<th>Potential barriers</th>
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<tbody>
<tr>
<td>Individual</td>
<td>Option 1 (create a model patient/registry/treatment-monitoring system in a single jurisdiction)</td>
</tr>
<tr>
<td></td>
<td>- Operational challenge in defining eligibility for a condition that lacks an 'event', widely agreed diagnostic criteria and demonstrated pathology</td>
</tr>
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<tr>
<td>Care provider</td>
<td>Option 1 (create a model patient/registry/treatment-monitoring system in a single jurisdiction)</td>
</tr>
<tr>
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<td>System</td>
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<tr>
<td></td>
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</table>

options. Several participants also voiced strong support for the research and support functions that a national network of centres with a coordinating ‘hub’ (option 2) could achieve for people living with chronic pain and their providers. Many also supported the idea of a cross-discipline model of patient-centred, primary health care-based chronic pain management. This model could include: primary health care practices/clinics assumed the primary responsibility for chronic pain management; facilitated access to mentoring for these practices/clinics to support and enable them to fulfil this role; and opportunities for these practices/clinics to periodically engage multidisciplinary and multimodal secondary- and tertiary-level supports for those people living with chronic pain requiring more complex care and support.

As part of the deliberation about implementation considerations, several dialogue participants noted that prospects for success for the registry/system (option 1) would be much greater if implemented in the form of a program of research rather than as a government or regional health authority initiative. Some argued that successful implementation of a national network of centres (option 2) hinged on getting the right champions (clinicians, leaders in teaching institutions and people living with chronic pain) around the table from the beginning. Several dialogue participants suggested that brokering and supporting the implementation of a cross-discipline model (option 3) could be facilitated in the short term through demonstration projects, coupled with rigorous monitoring and evaluation, and in the long term through a systematic effort to “move beyond the early adopters.”

In the last deliberation about next steps that different constituencies could take, one dialogue participant argued that governments seem unprepared to take action in the short term, so “stakeholders have to be the ones who make it happen.” Several dialogue participants argued that the critical next step should be to engage those who could take action, including key opinion leaders (both those leading the push for strengthened chronic pain management and those in primary health care practices), regional health authorities and government. A number of dialogue participants argued that success stories need to be identified, their cost-effectiveness relative to the status quo studied, and the findings from this effort popularized in a systematic way.

### Topic 2: Engaging health system decision makers in supporting comprehensive chronic pain management in provincial and territorial health care systems in Canada

The second dialogue was designed to build on the key finding from the first dialogue that long-term sustainable action is constrained by the lack of attention paid to chronic pain by health system decision makers. Given this, the second dialogue was focused on how to more systematically engage health system decision makers in supporting comprehensive chronic pain management in provincial and territorial health care systems in Canada. For the purposes of this brief and dialogue, decision makers included policymakers (and those who support policymakers), and regional health authority staff. We present below a summary of the key findings from the issue brief and the key themes of the deliberations. For those interested in additional information, the issue brief (11) and dialogue summary (25) are available on the McMaster Health Forum website (www.mcmasterhealthforum.org).

#### Key findings from the issue brief:

The lack of health system decision maker engagement in supporting comprehensive chronic pain management in provincial and territorial health care systems in Canada can be understood by considering four sets of inter-related issues, outlined along with a summary of contributing factors in Table 4. Similar to the first brief, to promote discussion about the pros and cons of potentially viable options, we selected three, which we outline in Table 5 along with a summary of the key findings. Finally, we identified implementation barriers at the level of individuals, care providers and systems, as well as possible strategies to address the barriers, which we outline in Table 6.

#### Summary of dialogue 2:

The dialogue brought together 16 participants from across Canada, which included six health system decision makers, four individuals from groups representing people living with chronic pain and professionals, five researchers and one representative from another stakeholder group.

During the deliberation about the problem, most participants noted that the lack of health system decision maker engagement is largely the result of a lack of awareness of the problem. Themes related to this lack of awareness that emerged during the dialogue include: a lack of understanding of chronic pain (which in turn relates to a lack of consensus about what chronic pain actually is, and a lack

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**TABLE 4**

Features of the problem of the lack of health system decision maker engagement in supporting comprehensive chronic pain management (and its causes)*

<table>
<thead>
<tr>
<th>Issue</th>
<th>Factors contributing to the issue</th>
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<tbody>
<tr>
<td>Lack of awareness of chronic pain</td>
<td>• Chronic pain may not garner sufficient attention because it is often associated with or the result of one or more physical or psychological comorbidities and, as a result, it is often seen as a symptom rather than a disease or condition in its own right (1)</td>
</tr>
<tr>
<td>Lack of awareness of limitations in existing programs and services</td>
<td>• General limitations in the availability of and access to primary health care services are likely felt particularly intensely by those living with chronic pain, given they may be seen as more complex (and hence be less likely to be taken on as patients), they may have greater needs for care (and hence be more likely to suffer the consequences of a lack of care), and they may be more likely to seek out care in suboptimal settings such as emergency rooms (and hence be more likely to suffer the consequences of inappropriate care)</td>
</tr>
<tr>
<td>Gaps in health system arrangements that limit the attention given to chronic pain</td>
<td>• A variety of health system arrangements contribute to the problem, including:</td>
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<td></td>
<td>o gaps in existing delivery arrangements (eg, lack of well-established packages of care/guidelines for the management of chronic pain, lack of a comprehensive continuum of care, and lack of integration with other models of proactive and coordinated care for chronic conditions);</td>
</tr>
<tr>
<td></td>
<td>o financial arrangements (eg, lack of visibility of the public and private costs of chronic pain management and lack of financial incentives for effective chronic pain management at the primary health care level); and</td>
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<tr>
<td></td>
<td>o governance arrangements (eg, lack of clear policy authority and lack of training and accreditation for health care providers and clinics to deliver care to people living with chronic pain)</td>
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<tr>
<td>Limited reach of existing efforts to engage health system decision makers in supporting chronic pain</td>
<td>• Several efforts have been made with the goal of developing and implementing a national pain strategy including briefs from the Canadian Pain Coalition and Canadian Pain Society that contained recommendations to government about the creation of a strategy (49), and broad-based endorsement for a national pain strategy at the Canadian Pain Summit, which garnered significant attention (26);</td>
</tr>
<tr>
<td></td>
<td>• While initiatives exist in several provinces (eg, British Columbia, Alberta, Ontario, Quebec and the Atlantic provinces), calls for a national pain strategy persist (28)</td>
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</tbody>
</table>

*The information in this table is based on what was available at the time of publication of the issue brief (April 2011)
Supporting chronic pain management across Canadian health systems

TABLE 5
Three options for engaging health system decision makers in supporting comprehensive chronic pain management*

<table>
<thead>
<tr>
<th>Option</th>
<th>Option focus and elements</th>
<th>Summary of key findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Launch an advocacy campaign</td>
<td>• Launching an advocacy campaign may involve using a variety of advocacy approaches to: o increase attention to the issue in general; and o elevate the visibility of chronic pain on provincial and territorial governments’ agendas in particular (with this goal being ideally positioned within a more general effort to encourage better informed health system decision making) • Approaches to advocacy might include: o traditional media for public engagement, such as print, radio and television; o ‘new media’ for public engagement, such as mass short messages (MSMs) and other mobile phone-based strategies, as well as online petitions and other Internet-based approaches; and o efforts to directly engage government officials</td>
<td>• Traditional media has been found by two high-quality but older reviews to positively influence individual health-related behaviours (50,51) • We did not identify systematic reviews that address whether and how the three types of advocacy initiatives outlined in the adjacent column (traditional media, ‘new media’ and direct engagement) increase the attention paid to an issue by decision makers</td>
</tr>
<tr>
<td>2. Create a multistakeholder provincial or national working group</td>
<td>• This option involves raising awareness and support among policymakers who could or should be paying attention to chronic pain, ideally in the context of a broader effort to engage all relevant stakeholders in supporting improvements to chronic pain management (with the stakeholders including the full range of health system decision makers, health care providers, researchers and provincial/national coalitions or nongovernmental organizations) • Elements of a multistakeholder provincial or national working group could include: o establishing a national network of stakeholder groups with a coordinating ‘hub’; o engaging key opinion leaders who can take action, both those leading the push for strengthened chronic pain management and those in primary health care practices, regional health authorities and government; o equipping these key opinion leaders (and the stakeholder groups from which they are drawn) with the necessary tools to take action, which may in turn include: • compelling data and stories about the current burden of chronic pain and the implications (eg, costs) of not addressing it; • regularly updated, evidence-based packages of care/guidelines for the management of chronic pain at the primary health care level and in related fields of practice; • mechanisms to coordinate across fields of practice and across relevant disease groups that are often linked to chronic pain; and • periodically identified priorities for new primary and secondary research and the communication of these priorities to relevant funders</td>
<td>• One medium-quality systematic review found a lack of evidence about the effects of multistakeholder networks (specifically public health partnerships) on health outcomes but qualitative studies included in the review suggested that some partnerships increased the profile of health inequalities on local policy agendas (52) • Four older systematic reviews related to engaging opinion leaders (in the clinical context), three of which were high quality (53-55) and one low quality (56) ○ Each review focuses on the clinical context, but still offer helpful insights about the potential effects of using local opinion leaders who can lead the push for engaging health system decision-makers in strengthening chronic pain management ○ Both of the high-quality reviews found minimal evidence about local opinion leaders, but one concluded that opinion leaders with or without another intervention were generally effective for improving appropriate care (53), and the other concluded that there was insufficient evidence to determine the effectiveness of local opinion leaders in the field of physiotherapy (54) ○ The remaining two reviews similarly concluded that there is insufficient evidence to determine whether local opinion leaders are effective for supporting clinical practice (55,56)</td>
</tr>
<tr>
<td>3. Develop chronic pain policy portfolios and strategic foci</td>
<td>• Developing policy portfolios or strategic foci would provide the opportunity to coordinate responses to chronic pain within and across governments, regional health authorities, and the stakeholder community and could include: o engaging and liaising with other relevant policy areas within the government and regional health authorities to coordinate the development and implementation of relevant policies and programs; and o engaging and liaising with relevant stakeholders to inform the development and implementation of programs and services in the community • Elements of this option might include: o mapping what existing policy portfolios are relevant to supporting chronic pain management, particularly at the primary health care level; and o establishing an integrated portfolio to support chronic pain management or a coordinating role that would work across other relevant portfolios and departments.</td>
<td>• No systematic reviews addressing any of the elements of this option were identified • Key messages that emerged from previous efforts in Canada to reallocate resources and decision making to support a shift in perspective may provide helpful insight: ○ In the early 1990s, the province of Prince Edward Island initiated a process of ‘cross-sectoral reallocation’ that emphasized “broad determinants of health, client focus in service delivery, pooling of human services, integration and coordination of services, and the establishment of regional governance” (28,57-61) ○ An analysis of instruments put in place to facilitate the shift toward the broad determinants of health revealed that regional governance can help ensure integration and coordination within regions but that there is a need for a central authority to ensure equity between regions (28) ○ Additional elements cited as important facilitators of the process include: fostering an organizational culture that is supportive of change; and starting with low-profile changes that can demonstrate how it can work (28)</td>
</tr>
</tbody>
</table>

*The findings in this table are based on what was available at the time of publication of the issue brief (April 2011)
Wilson et al

TABLE 6
Potential barriers to implementing the options in the issue brief*

<table>
<thead>
<tr>
<th>Levels</th>
<th>Potential barriers</th>
</tr>
</thead>
<tbody>
<tr>
<td>Individual</td>
<td>Option 1 (launch an advocacy campaign)</td>
</tr>
<tr>
<td></td>
<td>• People living with chronic pain may be unwilling (eg, due to the stigma associated with chronic pain) or unable (eg, due to the limitations related to having chronic pain) to be meaningfully engaged in advocacy campaigns</td>
</tr>
<tr>
<td></td>
<td>• There is a wide array of advocacy campaigns for many different diseases with each competing for the attention of the broader public</td>
</tr>
<tr>
<td></td>
<td>Option 2 (create a multistakeholder provincial or national working group)</td>
</tr>
<tr>
<td></td>
<td>• People living with chronic pain may be unwilling (eg, due to the stigma associated with chronic pain) or unable (eg, due to the limitations related to having chronic pain) to be meaningfully engaged in the activities of a working group</td>
</tr>
<tr>
<td></td>
<td>Option 3 (develop policy portfolios in ministries and strategic foci in regional health authorities)</td>
</tr>
<tr>
<td></td>
<td>• Not applicable – such a change would likely not be visible to individuals</td>
</tr>
<tr>
<td>Care provider</td>
<td>Option 1 (launch an advocacy campaign)</td>
</tr>
<tr>
<td></td>
<td>• Providers or associations of providers attempting to directly engage health system decision makers may not have the time or skills required to make the case for better supporting chronic pain management</td>
</tr>
<tr>
<td></td>
<td>Option 2 (create a multistakeholder provincial or national working group)</td>
</tr>
<tr>
<td></td>
<td>• Providers may be skeptical about or unwilling to implement the recommendations from a working group</td>
</tr>
<tr>
<td></td>
<td>Option 3 (develop policy portfolios in ministries and strategic foci in regional health authorities)</td>
</tr>
<tr>
<td></td>
<td>• Not applicable – such a change would likely not be visible to care providers</td>
</tr>
<tr>
<td>Organization</td>
<td>Option 1 (launch an advocacy campaign)</td>
</tr>
<tr>
<td></td>
<td>• Organizations attempting to directly engage health system decision makers may not have the time or skills required to make the case for better supporting chronic pain management</td>
</tr>
<tr>
<td></td>
<td>• Key health system advocacy organizations (eg, medical associations) may not be willing to devote resources to advocacy for chronic pain compared to other conditions that have bigger impacts on their members’ interests</td>
</tr>
<tr>
<td></td>
<td>Option 2 (create a multistakeholder provincial or national working group)</td>
</tr>
<tr>
<td></td>
<td>• Health care delivery organizations may not be willing to participate in a working group for chronic pain compared with other conditions that have bigger impacts on their organization</td>
</tr>
<tr>
<td></td>
<td>• Organizations with existing chronic pain strategies may not be willing to participate in a working group that is covering ground that they have already covered</td>
</tr>
<tr>
<td></td>
<td>Option 3 (develop policy portfolios in ministries and strategic foci in regional health authorities)</td>
</tr>
<tr>
<td></td>
<td>• Regional health authorities are increasingly focused on broad issue domains (eg, chronic disease) that encompass many different diseases rather than on specific issues</td>
</tr>
<tr>
<td>System</td>
<td>Option 1 (launch an advocacy campaign)</td>
</tr>
<tr>
<td></td>
<td>• There is a wide array of advocacy campaigns for many different diseases with each competing for the attention of health system decision makers</td>
</tr>
<tr>
<td></td>
<td>Option 2 (create a multistateholder provincial or national working group)</td>
</tr>
<tr>
<td></td>
<td>• There are limited resources available to support the development and ongoing activities of a working group</td>
</tr>
<tr>
<td></td>
<td>Option 3 (develop policy portfolios in ministries and strategic foci in regional health authorities)</td>
</tr>
<tr>
<td></td>
<td>• Ministries of health, and those seeking to influence them, are increasingly focused on broad issue domains (eg, chronic disease) that encompass many different diseases rather than on specific issues</td>
</tr>
</tbody>
</table>

Possible strategies to address the barriers

• Given that several options may be pursued simultaneously and that option elements may be combined in different and creative ways, identifying ‘cross-cutting’ implementation strategies may be an important first step

• One possible cross-cutting implementation strategy may be the development, pilot testing and iterative redevelopment of a package of communication materials that highlight the ways in which chronic pain affects people’s lives, the costs associated with the status quo, and success stories (and how their cost-effectiveness compares with the status quo)

*The information in this table is based on what was available at the time of publication of the issue brief (April 2011)

of those living with it, a lack of documentation of the broader economic impact of chronic pain, and a lack of documentation of success stories in chronic pain management.

In deliberating about the three options to address the problem, most dialogue participants strongly endorsed the creation of a national multi-stakeholder network. Participants indicated that such a network should be comprised of organizations and committed individuals who would work collaboratively to raise awareness about chronic pain, and increase support for and coordination in comprehensive chronic pain management. Several dialogue participants called for including in the network those involved in chronic disease management, primary health care, and other domains that have already been prioritized (particularly those that are highly relevant to people living with chronic pain). Most participants believed that an advocacy campaign would be an important function for this network. A number of dialogue participants also endorsed the idea of developing chronic pain policy portfolios within government, and strategic foci within regional health authorities. It was suggested that these would optimally be nested within broader portfolios/foci such as chronic diseases to ensure there is a clear ‘anchor’ for chronic pain within health systems.

Five significant challenges related to implementation were identified by dialogue participants: identifying a leadership model; ensuring the capacity and willingness of existing organizations and individuals to engage in creating and sustaining the network; weighing the advantages of working within a small group of provinces and territories to achieve some early wins, versus working across all provinces and territories simultaneously; securing the resources to design, launch and operate the network and any advocacy campaign it develops; and scaling up the efforts to identify and harness data and to produce and synthesize research evidence that supports the work of the network and the content of any advocacy campaign.

The deliberation about next steps focused on the need to mobilize toward building a national network that would bring together existing organizations and committed individuals. Some early wins for the network may include the endorsement of a national pain strategy, which would include agreed-upon definitions. Several dialogue participants emphasized the primacy being given to an ‘evidence-based’ and ‘grass-roots’ approach, and being certain not to lose these features in a rush to a ‘big bang’ solution.

Progress following the dialogues
Progress has been made in several areas that were deliberated about during the stakeholder dialogues (26), including: launching an advocacy campaign (eg, a delegation went to Parliament Hill in the lead up to the Canadian Pain Summit in April 2012); the initiation of a national stakeholder-engagement process (National Pain Summit) to raise awareness of health system issues within the chronic pain community and to raise awareness of chronic pain issues within the health policy and systems community (this was highlighted as a possible implementation strategy in the first dialogue); and developing chronic pain policy portfolios and strategic foci (eg, Ontario’s efforts to develop a chronic pain plan for the province). In Australia, such advocacy and stakeholder engagement efforts (eg, the National Pain Summit held there in 2010) were the main catalysts that led to them being the first country in the world to develop a national strategy for the treatment and management of pain (27).

Similar momentum and interest among policymakers and stakeholders seems to be building in Canada toward developing a similar strategy for the country.

**DISCUSSION**

Principal findings
To inform the first dialogue, we found that systematic reviews supported the use of evidence-based tools for strengthening chronic pain management, including patient education, self-management supports, interventions to implement clinical practice guidelines, and multidisciplinary approaches to pain management. While research evidence about patient registries/treatment-monitoring systems is limited, many dialogue participants argued strongly that such a system is needed. In addition, many saw a registry as a precondition for moving forward with other options, including creating a national network of chronic pain centres with a coordinating ‘hub’ to provide chronic pain-related decision support, and a cross-payer, cross-discipline model of patient-centred primary health care-based chronic pain management. For the second dialogue, we found systematic reviews indicating that traditional media can be used to positively influence individual health-related behaviours, and that multistakeholder partnerships can contribute to increasing the attention paid to issues regarding policy agendas. We also found evidence from previous efforts in Canada for ‘cross-sectoral reallocation’, which highlighted that regional governance can help ensure integration and coordination within regions (28). The evidence also indicated that fostering an organizational culture that is supportive of change, and starting with low-profile changes that can demonstrate how it can work, were important facilitators of cross-sectoral reallocation processes (28).

Dialogue participants emphasized: the need to mobilize behind an effort to build a national network that would bring together existing organizations and committed individuals; an early win for the network may include the endorsement of a national pain strategy; and the need to not lose the primacy given to an ‘evidence-based’ and ‘grass-roots’ approach in a rush to a ‘big bang’ solution.

Strengths and limitations
Our process had two notable strengths and two important limitations. First, because we were not actively engaged in work related to chronic pain, we were able to act as neutral convenors of the stakeholder dialogues, which allowed us to better achieve our overriding objective of supporting evidence-informed policymaking. The second strength is that we paired the best available research evidence with a robust deliberative process that gives voice to the tacit knowledge and real world views and experiences of those involved in and/or affected by the issue. The main limitation of the present paper is that the findings reported from the evidence and issue briefs are based on the research evidence that we identified at the time of finalizing each (December 2009 for the evidence brief and April 2011 for the issue brief). We determined that presenting the evidence from the original documents was the optimal approach as it provides a picture of what dialogue participants had reviewed before each of the stakeholder dialogues. However, we provided a monthly evidence service to stakeholders for one year following each dialogue, which kept them updated about new systematic reviews that had been published about each of the options in the briefs. The second limitation is that we were unable to engage our target size of 18 to 22 participants in each dialogue, with only 13 participants in the first dialogue and 16 in the second. In addition, we only engaged two policymakers in the first dialogue, which was the rationale to convene the second where we were able to engage six policymakers.

Implications for policy
The actionable messages coming out of these dialogues are clear: chronic pain management needs to be properly addressed in Canada and it is critical to engage leaders who can take action. Developing a national network and thereafter a national pain strategy are important initiatives that garnered broad-based support in the dialogues. Progress has been made towards these goals, although sustained efforts are required to build on this progress. In both dialogues, the idea of success stories or quick wins, which highlight the advantage of interventions, were identified as important next steps toward developing such a strategy.

Implications for research
What was apparent from the evidence and issue brief is that numerous research gaps exist. Whether the focus was the incidence and distribution of chronic pain in Canada, possible options to address problems with its management or key implementation considerations, there were few systematic reviews, and many of those that did exist were dated and of mixed quality. These gaps likely explain the assertion by dialogue participants that there is a lack of well-documented cases for why chronic pain is an important issue that health system decision makers should focus on. While there have been some efforts to this end, such as a book that provides a health policy perspective about chronic pain (29), the dialogue participants were speaking directly to the lack of awareness of such efforts, the need to supplement them with stories about how chronic pain affects the lives of those living with it, and documentation of the broader economic impact of chronic pain. In addition to this, future research may also be focused at the level of people living with chronic pain by examining the implications of classifying chronic pain as a disease in Canada; the provider level by analyzing their perceptions to different initiatives or treatment of chronic pain; the organizational level by evaluating the effectiveness of approaches to providing comprehensive chronic pain management; and at the health system level by examining the feasibility of creating a national network or studying the impact of chronic pain policy portfolios.

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REFERENCES


31. Statistics Canada. CANSIM Table 105-5001: Health indicator profile, annual estimates, by age group and sex, Canada, provinces, territories, health regions (2007 boundaries) and peer groups, occasional. Canadian Community Health Survey 2009.


