The need for knowledge translation in chronic pain

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One in five Canadians suffers from some form of persistent or chronic pain. The impact on individual lives, families and friends, the health services sector and the economy is huge. Reliable evidence is available that the burden of persistent pain can be markedly reduced when available knowledge is applied. Bridging the quality chasm between chronic pain and the care process will require a unique confluence of opinion from all stakeholders committed within a focused community of practice to address the impact of pain. Various levels of success in this regard have been demonstrated when there is exchange, synthesis and ethically sound application of research findings within a complex set of interactions among researchers and knowledge users. It is now critical to accelerate the capture of the benefits of research for Canadians through improved health, more effective and responsive services and products, and a strengthened health care system to bring about health reform and health care reform across Canada as it pertains to the one in five Canadians living with chronic, disabling pain.

The overarching outcome of such an initiative needs to be promoted to sustain a balanced portfolio of curiosity- and needs-based research, which along with existing knowledge, can be mobilized and applied for the benefit of Canadians, the health care system and the economy.

Key Words: Chronic disease management; Chronic pain; Knowledge exchange; Organizational change; Research into practice; Research utilization

Clause 4 of Bill C-13, the Canadian Institutes of Health Research (CIHR) Act of the Canadian Parliament (1), states:

The objective of the CIHR is to excel, according to internationally accepted standards of scientific excellence, in the creation of new knowledge and its translation into improved health for Canadians, more effective health services and products and a strengthened Canadian health care system...

Research generating knowledge in pain is attempting to understand mechanisms, complexities, incidence and prevalence, and the personal, social and economic costs, etc. While much remains to be learned, at the same time we must recognize that we are much further ahead than we were 20 years ago, or even 10 years ago. An important question, though, is how much this mounting knowledge is being applied to those who suffer debilitating pain, which is the presumed end-purpose of knowledge generation. This is the entry point for ‘knowledge translation’, the term widely used to refer to the process of research use or the application of knowledge to receptor communities.

More formally, knowledge translation at CIHR is defined as the exchange, synthesis and ethically sound application of research findings within a complex set of interactions among researchers and knowledge users – to accelerate the capture of the benefits of research for Canadians through improved health, more effective services and products, and a strengthened health care system.

How important is knowledge translation to the area of pain chronic or persistent pain in particular? The present overview is designed to argue that Canada, the Canadian health care system, Canadian society and the Canadian economy are in dire need of application of knowledge translation and exchange to debilitating pain. In fact, a national strategy is needed to promote implementation of research findings, application and appraisal of clinical practice guidelines, and uptake of decision support tools by health care practitioners and decision-makers.

KEEPING IN STEP WITH SHIFTING NEEDS

Effective knowledge translation initiatives must be directed at ensuring that Canadians live well, as free from undue pain and suffering as possible, engaging consumer patients and their local caregivers, along with academic, health service delivery, voluntary sectors and government organizations as a community of practice directed at implementing and sustaining capacity-building initiatives throughout Canada in support of health...
care renewal, navigating the complexities of ensuring that systems of care are available when patients and families need them. Terms used in this document are adapted from references 2 and 3, and <www.ewenger.com>.

The World Health Organization has recognized the shift in needs of health care, from the acute model to a chronic disease model. Irrespective of the level of prosperity, health care systems worldwide face disparities in access to care, and limitations in assets and resources. Policy and service decision-makers face the challenge of matching these limited assets and resources to the changing needs, to make significant improvements in caring for chronic conditions. Leadership combined with a willingness to embrace change and innovation will have far more impact than simply adding capital to health care systems. In the World Health Organization’s Innovative Care for Chronic Conditions (4) it is stated that:

Health care is organized around an acute, episodic model of care that no longer meets the needs of many patients, especially those with chronic conditions. Decreases in communicable diseases and the rapid aging of the population have produced this mismatch between health problems and health care, and chronic conditions are on the rise. Patients, health care workers, and most importantly, decision-makers, must recognize that effective chronic condition care requires a different kind of health care system. The most prevalent health problems such as diabetes, asthma, heart disease, and depression require extended care policies. Funding mechanisms and priorities, and by established health care providers, but the Institute of Medicine report shifts that focus from the providers to the systems in which they work – “Current care systems cannot do the job. Trying harder will not work. Changing systems of care will.”

In the paragraphs below, it will be pointed out that chronic pain inflicts a heavy burden on individual Canadians, the Canadian health care system and the Canadian economy. The means to bring about improvement in all these domains exist today. The chronic disease management model in Figure 1 is modified from the original by Glasgow et al (8).

**CHRONIC PAIN AS A DISEASE**

Not all pain is a disease. Pain is a protector to us most of the time, at least for those of us who do not live with constant pain. Acute pain, such as that following trauma or surgery, constitutes a signal to a conscious brain about the presence of noxious stimuli and/or ongoing tissue damage. This acute pain signal is useful and adaptive, warning the individual of danger and the need to escape or seek help. Acute pain is a direct outcome of the noxious event, and is reasonably classified as a symptom of underlying tissue damage or disease. When the original disorder resolves, so do its symptoms.

For some, though, pain is a constant companion, persisting long after its usefulness as an alarm signal has passed, and indeed, often long after the tissue damage has healed. Chronic pain in these patients may not be directly related to their initial injury or disease condition, but rather to secondary changes including some that occur in the pain detection system itself. Thus, the mechanisms underlying chronic or persistent pain may be quite different from acute pain. It is becoming increasingly clear that mechanisms are triggered that sustain the pain even when the cause is no longer active. Take phantom limb pain, for example – the painful limb may be removed, yet the pain persists (9,10). Postherpetic neuralgia is too often a lasting outcome of the cutaneous eruptions of herpes zoster infection (11,12). The pain does not resolve when the original cause resolves. The same may be said of the neuropathic pain that persists after cutting certain intercostal nerves during open-chest surgery (13).

Chronic pain may also be considered as a disease somewhat akin to considering myocardial infarction as a disease. Myocardial infarction may be caused by smoking, hypertension or diabetes. Multiple factors contribute to the cause of myocardial infarction, but myocardial infarction is the disease.
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Similarly, it can be argued that persistent pain is a disease, and can result from multiple conditions.

The European Federation of International Association for the Study of Pain Chapters views chronic pain as a disease (14), and they have lobbied the European Parliament to declare chronic pain as a disease.

Pain is a major healthcare problem in Europe. Although acute pain may reasonably be considered a symptom of disease or injury, chronic and recurrent pain is a specific healthcare problem, a disease in its own right (15).

Chronic pain as a disease is also not without precedent. It can be argued that trigeminal neuralgia is painful and is a disease, as are migraines. Central poststroke pain is not a symptom, but a lasting disease.

Chronic pain differs from acute pain not only with regard to pathophysiology, but also with regard to associated features. Chronic pain often sets the stage for the emergence of a complex set of physical and psychosocial changes that are an integral part of the chronic pain problem and add greatly to the burden on the pain patient. These include immobility and consequent wasting of muscle, joints, etc; depression of the immune system and increased susceptibility to disease, fatigue, disturbed sleep; poor appetite and nutrition; dependence on medication; overdependence on family and other caregivers; overuse and often inappropriate use of professional health care systems; poor performance on the job or inability to work; disability; isolation from society and family; and turning inwards, anxiety, fear, bitterness, frustration, depression and suicide (16).

Let us consider the overall impact of chronic pain and its summation of symptoms and comorbidities. Is it such a
problem that a system-wide paradigm shift in management and treatment will have a significant effect?

THE BARE FACTS ABOUT CHRONIC PAIN

Types of chronic pain

Chronic pain can arise from sundry causes. Some of the more common types of chronic pain include those of osteoarthritis, rheumatoid arthritis, low back, shoulder and neck, headache (including migraine), cancer pain, myofascial pain syndromes, post-thoracotomy pain, chronic regional pain syndromes, stump and phantom limb pain, neuropathic pain, herpes zoster (shingles) and postherpetic neuralgia, trigeminal neuralgia, diabetic neuropathy, temporomandibular joint disorder, post-mastectomy pain, angina pectoris, chronic visceral pain syndromes, and others.

To date, there are no comprehensive figures outlining the incidence of the various chronic pain syndromes and their costs to Canadian society. However, investigators in various countries have begun to compile information of this nature, which illustrates the magnitude of suffering due to chronic pain.

Prevalence of chronic pain

In a random survey of 2012 Canadians, chronic pain was reported by 29% of respondents, with increased frequency in women and older age groups. The average duration of pain was 10.7 years and the average intensity was 6.3 on a 10-point scale. Almost 70% of those reporting pain were worried about addiction (17). Estimates of the prevalence of activity limitations related to neck pain alone are in the range of 11% to 14% in both Ontario and Quebec (18,19). However, this number is not universally acknowledged – according to workers’ compensation statistics, work-related neck pain represents a minor problem that a system-wide paradigm shift in management and treatment will have a significant effect?

Impact of chronic pain

Multifactorial issues surround chronic pain, where direct and indirect costs must be factored into assessing impact, as well as factors such as the high occurrence of comorbidity, attitude toward chronic pain and chronic pain patients among health care providers, fears and beliefs of chronic pain sufferers, fears of taking painkiller drugs, psychosocial context, etc. Thus, while the following paragraphs will include an attempt to address specific issues, these issues must be considered in the context that there is a close inter-relationship between all the factors.

Chronic pain as a disability

In a Canadian survey (17), almost one-half of those reporting chronic pain were unable to attend social and family events and the mean number of days absent from work per person in the past year due to chronic pain was 9.3. A Toronto study (30) indicated that among medical illnesses, chronic pain was the second major cause of suicide, only after bipolar disorder, and ahead of depression and psychotic disorder. Richard Holden, a former Member of the National Assembly in the Quebec Legislature who represented Westmount, left a suicide note indicating that his excruciating pain led him to suicide. Of 19 adults with disabling hip and knee osteoarthritis in a Toronto cohort (31), most had comorbidities such as heart disease and diabetes. Sadly, adherence to pain medication differed from that of other prescribed medications because participants were reluctant to take painkillers and when they did, they generally took them at a lower dose or frequency than prescribed.

Chronic pain is typically accompanied by a constellation of other disorders. Older persons, particularly women and the less educated, are most severely impacted and disabled by chronic pain (23,29,32). In a US study of 55,686 patients with painful neuropathic disorders (33), older persons, women and the less educated were more likely to have other pain-related conditions, including fibromyalgia (6.0%), osteoarthritis (13.6%) and other chronic comorbidities such as coronary heart disease (13.6%) and depression (6.4%). In a European study of 18,980 randomly selected subjects (34), 16.5% reported a depressive symptom and of these, 27.6% had at least one type of chronic pain. Among the 4.0% diagnosed with major depressive disorder, 43.4% had at least one type of chronic pain. Neuropathic pain patients reported significantly more sleep disturbance and daytime somnolence as well as less quality and adequacy of sleep than patients in the general US population (35). Of 749 children and adolescents in a German study (36), 83% had experienced pain during the preceding three months and reported that pain led to sleep problems (54%), inability to pursue hobbies (33%), eating problems (51%), school absence (49%) and inability to meet friends (47%). A Scottish random sample of 4611 adults (37) indicated that the prevalence of ‘any’, ‘significant’ and ‘severe’ chronic pain had progressively more adverse associations with employment, and interfered with daily activities and all measured dimensions of general health. Individuals with spinal cord injuries and pain were more likely than those without pain to experience frequent interference with a variety of daily activities, including sleep (38). The pain had more negative effects on quality of life than the extent of the spinal injury (39). Of 255 patients with painful diabetic...
Peripheral neuropathy in a US study (40), during the previous three months, 60% had two or more health professional consults, 59% reported decreased home productivity, 8.5% reported activity limitations and 64.4% of those who worked reported missing work or decreased productivity due to pain; 62.7% had other neuropathic and musculoskeletal pain conditions. Whether pain associated with multiple sclerosis (41), fibromyalgia (42,43), herpes zoster (44,45) or other disorders that carry chronic pain, the data reflect a huge impact of pain and increased likelihood of comorbid conditions.

In a recent review of patients referred to a Danish pain centre (46), mean pain severity was 7 on a 10-point scale, quality of life measures were severely reduced, 58% had depression or anxiety disorder, 63% had neuropathic pain and 73% of these were taking opiates on referral even though this did not provide adequate pain relief. The study showed that health-related quality of life of chronic pain patients is among the lowest observed for any medical condition (46). Data from a study in Sweden (47) indicate that spinal pain is very common among men and women 35 to 45 years of age and that it is related to marked limitations in lifestyle for approximately one-quarter of those who experience pain.

Clearly, then, chronic pain must be recognized directly and indirectly through comorbidities as a major factor having an impact on the quality of life and the life habits of a large number of Canadians.

Health services usage
The 1994 to 1995 National Population Health Survey (48) indicated that 3.9 million Canadians, or 17% of the population over 15 years of age, suffer from chronic pain. Of those, 70% rated the pain as moderate to severe and as interfering with daily activities. Those with severe chronic pain made more physician visits (mean of 12.9 versus 3.8 visits) and stayed in hospital longer (mean of 3.9 versus 0.7 days) compared with those without any chronic pain in the previous year.

In a European study of 5803 people with musculoskeletal pain (49), up to 27% did not seek medical help. Of those who did see a doctor, several waited months or years, 55% of patients who saw a doctor are currently receiving drug treatment for their pain. Patients typically reported that communication with doctors was poor. Within the context that pain is the most common complaint of those suffering musculoskeletal conditions, it was reported (50) that in the US between 1990 and 1992, patients with musculoskeletal conditions made 315 million physician visits, had more than eight million hospital admissions and experienced approximately 1.5 billion days of restricted activity. A study in Finland (51) found that, from a pool of 5646 patient visits to primary health care services, 40% identified pain as the reason for their visit. One-fifth of patients reported having experienced pain for over six months. One-quarter of the pain patients of active working age were receiving paid sick leave. From an Australian stratified sample of 17,543 telephone interviews (32) with a 70.8% response rate, disability from chronic pain was associated with a greater number of primary care visits, hospital admissions and emergency department visits.

These data appear to implicate pain as a leading cause of health service usage by chronic pain patients, but dedicated surveys implicate factors beyond just the pain. While it is generally acknowledged that patients with chronic pain use health care services significantly more than persons without chronic pain, what is perhaps surprising is that less than one-third of the care is for the primary pain condition across different types of pain (51-54). Increased usage is often due to ill-defined conditions, lower priority chronic disease, acute disease and mental health care (55).

Economic costs
In view of efforts to align care along more economical avenues, it is interesting to note that in a study of the economic impact of chronic pain and disease in Canada (56), resource use measures were found to be driven more by disease prevalence than intensity of resource use. Costs associated with multiple sclerosis in Canada, for example, are reported to be positively correlated with levels of self-reported pain, with an estimated six-month overall burden of pain in multiple sclerosis patients to be over $79 million (41).

In an attempt to quantify the total cost of chronic noncancer pain to the Irish economy in 1995, a study from a sample of 95 patients estimated that chronic pain had already cost the economy £1.9 million by the time of their referral to a multidisciplinary pain clinic (57). Total health care expenditures incurred by individuals with back pain in the US reached $90.7 billion in 1998 (51), those for neck pain reached US$686 million in the Netherlands in 1996 (58). In a study of 25 randomly selected Finnish health centres over a four-week period (59), 1123 patients visited general practitioners because of musculoskeletal pain. Laboratory tests were ordered for 12%, imaging for 24%, analgesics prescribed for 61% and sick leave prescribed for 25%; the mean cost per patient was as high as €530 per visit. A study of the socioeconomic costs of pain syndromes in the UK (60) estimated the direct health care cost of back pain in 1998 to be £1.6 billion. However, this direct cost was considered to be insignificant compared with the cost of informal care and the related production losses, which totalled £10.7 billion. A US study of 16,567 patients in a retrospective administrative database analysis for the period 1996 to 2001 (61) reported that direct medical costs amounted to $357 per month for each patient; the authors suggested a tailored approach to care for high utilisers. Specific numbers are also available for other types of chronic pain, including postherpetic neuralgia (44), neuropathic pain (62), multiple sclerosis (41), back and neck pain (63), rheumatoid arthritis (64) and others. It is important to note that overall costs for chronic pain care have been increasing despite the emergence of evidence-based guidelines (63,65).

A German study of the usage of specialist care for low back pain found that 573 of 1342 patients were seeking specialist care. This high use of specialist care was attributed to the absence of a functioning primary care gate-keeping system for patient selection (66) and it was suggested that overall costs could be reduced with patient education, encouragement of physical activity and simple pain medications.

This section has examined direct costs. Let us now examine data on some of the indirect costs.

Chronic pain and the workplace
The Nuprin Pain Report (67), commissioned to study the impact of chronic pain in the US, revealed that patients with chronic pain had reduced capacity to work and enjoy life. The
The annual number of lost days at work was estimated at four billion, or 23 days for every adult. In a Swedish study on low back pain (68), indirect costs for patients were largely due to absenteeism and reduced productivity, and constituted 85% of overall costs.

Data from the American Productivity Audit (69) indicated that, during a two-week period, 13% of the total work force experienced a loss in productive time due to a pain condition. Surprisingly, 76% of the lost productive time was through reduced performance while at work rather than from absenteeism (69,70). Similar results were reported from an Australian study (71), in which chronic pain sufferers who worked with pain were more common (mean 83.8 days in six months) than chronic pain sufferers who lost work days due to pain (mean 4.5 days). A more recent Australian study (72) estimated that chronic pain sufferers lost 9.9 million work days due to chronic pain, equating a cost of AU$1.4 million. However, reduced effectiveness at work translated to 36.5 million lost work day equivalents, equalling AU$51 billion per year. A survey of 4839 chronic pain sufferers in Europe (73) showed that 61% were less able or unable to work outside the home, 19% had lost their jobs and 13% had changed jobs because of their pain, 60% had visited their doctor about their pain two to nine times in the past six months, only 2% were being treated by a pain management specialist and 48% had inadequate management of their pain. A study in the Netherlands (74) found that musculoskeletal diseases are the fifth most expensive disease category regarding hospital care, and the most expensive regarding work absenteeism and disability (1.7% of the gross national product). A recent systematic review of the costs of low back pain (75) concluded that economic costs, taking into account direct and indirect costs, must be considered a substantial burden on society. Interestingly, the review did not identify any studies estimating the costs from a societal perspective.

Data reported from the Alberta Workers’ Compensation Board (76) indicated that there was significant improvement in return-to-work outcomes when a continuum of care model was implemented to guide rehabilitation service delivery, with a cost-savings of $21.5 million over a two-year study period.

### Figure 2

System change must be designed to address components of chronic disease management principles applied across the care continuum for chronic pain for a system-wide strategic action that draws on existing success stories in Canada and a care continuum perspective on factors impacting on health measures and quality of life. The framework is based on the wide range of health-related disciplines that must join forces if opportunities to reduce disability associated with chronic pain are to be realized. In this respect, it is essential that system-level change is accompanied by and supportive of the empowerment and active participation of individuals, their families and communities. The present project is a first step toward this change as a reality by moving evidence and applying it to the care continuum.
Adherence as a target for a knowledge translation strategy

An extensive list of guidelines and systematic reviews is available from reliable sources for the management and treatment of chronic pain (77), and methods for assessment of evidence-based pain management are clearly laid out (78,79). In fact, the plethora of guidelines and systematic reviews may be an obstacle to use; there is a systematic review of systematic reviews of low back pain (80) and a systematic review of low back pain guidelines (81).

There are too many guidelines to begin to list here, but the important issue is whether these guidelines are used and make a difference to the patient (65); limited research has addressed the impact of adherence to guideline recommendations on clinical outcomes and costs of care (82) and adherence to guidelines has been advocated to decrease the use of ineffective therapies and to improve patient outcomes and more cost-effective care (83). On the one hand, a Montreal clinical trial confirmed that adherence to guidelines improves pain and functional status at six-month follow-up (84) and adherence to clinical guidelines for low back pain has been shown to have clinical and economic benefits (85,86). However, management of low back pain in a cohort of general practitioners in Ireland was not consistent with European clinical guideline recommendations and it was found that most of the costs incurred were attributable to nonadherence (87). Barriers to using evidence-based guidelines include lack of time, insufficient skills, uncertainty and resistance to findings that challenge traditional practices (88,89). Recognizing this gap between knowledge and practice, a call has been made to promote research into development of effective professional behaviour change interventions (90-92).

Attitude as a barrier to effective pain management

In a telephone survey of 2400 adults in Alberta and Saskatchewan (94), it was found that back pain beliefs among patients are not in harmony with current scientific evidence. Having access to appropriate information influences pain management outcomes. Attitude profiles of patients with chronic pain impact clinical status and quality of life (95,96), as well as fear-avoidance beliefs, which occur early in low back pain patients (97), for example, and have been reported to negatively impact on patient outcome, patient disability and health care contacts (98-100).

Health care providers can play an important role in providing appropriate patient information and thus may influence patient fear-avoidance beliefs and reduce costs (101). We have come a long way from the concept of accepting mild pain as...
Chronic pain is considered a chronic disease that imposes a heavy burden on patients and caregivers, on the health care system and on the economy. Chronic pain is generally undertreated and under-managed in the current health care environment in Canada. This burden translates to need related to functional impairment, due to the disease of chronic pain coupled with the disability that it creates.

Based on the evidence above, it is clear that there are measures that can be established to address the burden of chronic pain from the perspective of international standards of life habits. These can include nutrition, fitness, personal care, communication, housing, mobility, responsibilities, interpersonal relations, community life, education, work, and recreation.

Evidence is available that specific approaches to health care, deriving from the chronic disease management model outlined above, if applied, can improve these outcomes measures. This evidence translates to opportunity and calls for system change.

System change can come when, within a research environment, multistakeholder communities of practice bring together patients and their home and community caregivers, with motivated health care providers and informed community partners within a positive policy environment. These constituents may be brought together through the logic model outlined below. Chronic pain crosses all health care sectors, and shows a marked comorbidity with other chronic diseases. Thus, system change in chronic pain care should serve to improve outcomes in other chronic diseases in addition to pain.
The chronic care model
It is generally understood that the chronic pain patient is often a complex patient and, as several different factors have been shown to correlate with or be predictive of chronic pain, there are increasing calls to incorporate management approaches that are effective across illnesses (55,61). We shall examine one of these models developed to address optimal approaches to management of chronic illness. The Chronic Care Model is an evidence-based, conceptual framework developed first by Wagner et al (108). In the ACT Report of May 2002 (6), Wagner outlines a road map for a set of system changes that can increase optimal care, and includes the following list, rearranged from the original. Through a managed approach to research, it may be possible to incorporate these six areas to improve outcomes for people suffering chronic pain in Canada.

Self-management support: There is now considerable evidence that individual and group interventions that promote patient empowerment and the acquisition of self-management skills are effective in diabetes, asthma and other chronic conditions. They emphasize the crucial role that patients play in setting goals, establishing action plans, identifying barriers to effective self-management and problem-solving to overcome the barriers.

Delivery system design: Planned visits and active follow-up are central to assuring high-quality chronic illness care. Planned visits use relevant patient information, evidence-based guidelines and organized approaches to assure that all patients receive recommended services. Such visits can be individual or in groups, and include attention to self-management and preventive interventions, as well as acute problems. High-quality chronic illness care also requires close follow-up of the patient’s condition and treatments. This can be done by telephone or e-mail, as well as in person. Many of the critical follow-up tasks do not require clinical training and could be most efficiently and consistently performed by nonprofessional team members.

Provider decision support: Data indicate that guidelines become effective behaviour-change agents for health care providers only when they are woven into the fabric of patient care through effective professional education, reminders, and ongoing feedback and reinforcement. Guideline adherence can be increased by incorporating guidelines into a registry, flow sheets and patient assessment tools coupled with educational support from clinical opinion leaders such as medical specialists.

Community resources: Organizations large and small, in urban and rural areas, in solo practices and in integrated health systems have found that community resources supplement and support their efforts to improve the care of patients with chronic illnesses. Chronically ill patients can benefit from a variety of services and resources that are not available from their health care provider, such as educational offerings, peer support groups and exercise programs. Increasing access to effective community resources through linkages with relevant organizations and agencies is a cost-effective way to optimize care.

Clinical information systems: A registry or database of key information on all patients with a chronic condition is the glue that holds an effective chronic care system together. Registries facilitate monitoring and planning care for individual patients and for the practice as a whole. An effective registry reminds caregivers of needed services and can generate materials for providers and patients that support planned visits. To enhance the care for the practice as a whole an effective registry provides feedback on performance and can identify which patients are in particular need of attention.

A supportive health care organization: Given the challenges of packed appointment schedules and tighter budgets, changes at the level of practice are unlikely to happen unless it is a priority for the organization and its leaders. Strong senior leadership plays an instrumental role in providing motivation, securing resources and removing the barriers that may stall quality improvement activities.

EXECUTION PLAN

Knowing is not enough; we must apply.
Willing is not enough; we must do.
– Goethe

To focus on these areas, system change must address the chronic pain gap through a structured framework such as the one illustrated in Figure 2.

Logic model for the research themes as communities of practice
A logic model, such as that illustrated in Figure 3, identifies linkages between the activities of a programme and the achievement of its outcomes. It succinctly clarifies the set of activities that make up the programme and the sequence of outcomes that are expected to flow from these activities. The logic model is an illustration of how the activities of the programme are expected to lead to the achievement of the final outcomes.

In Nonaka’s reformulation of Polanyi’s definitions (109), extrinsic knowledge is knowledge that the individual holds explicitly and consciously in mental focus, and may communicate to others. Tacit knowledge is often subconscious, internalized, and the individual may or may not be aware of what he or she knows and how he or she accomplishes particular results. Some also use the term ‘embedded knowledge’, which is knowledge that is embedded in a process but not in an explicit way; that is, it requires other knowledge for it to be extracted (110).

This logic model can be applied in planning, implementation, evaluation and communication.

Organizational framework: The next step is to fit this logic model into an organizational framework designed to create a favourable environment for execution, to assess and evaluate impact and success, and promote clearly defined stakeholder engagement. An example is illustrated in Figure 4.

The following considerations are an attempt to outline research pathways that require adequate funding support to address the needs outlined above. These considerations are directed toward synergizing activities to make a compelling case for further large scale funding obtained from a variety of sources. The case is made for infrastructure support that will enable comprehensive research programs that focus on:

1. Pain cause and cure research;
2. Research on maximizing effectiveness and efficiency of health and social services support for the clientele of interest with a focus on disability that addresses enhanced life habits participation, prevention of deterioration, and optimization of health services usage to address the huge burden of chronic pain; and

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3. Knowledge mobilization that puts evidence-informed innovations, as per steps 1 and 2, into practice and evaluates the effectiveness of these innovations.

A coordinated approach to managed research is necessary to promote exchange and sharing of specific goals, methods, tools and assessments between active participants, to ensure cross-discipline learning, direction and strategy.

Partnering strategy

Partnership development will be an ongoing program. As research initiatives evolve, new opportunities for partnership will announce themselves, and a mobilizing plan for action will be developed and pursued based on shared vision and common values, which will build commitment, engage participants and use participants’ time well. The purpose will be to recruit and engage active and sustained involvement of additional participants in the action plan, with clearly defined benefits to this coordinated effort and to the partners, as well as to raise additional funds to enable new initiatives to be undertaken beyond those currently available. Broad representation must be sought from local public health system partners, other community organizations, and community residents. Participants will need to identify resource needs and determine sources for meeting those needs. For each partner, tools such as a project proposal, a master calendar and action-linked agendas must be developed. A process for clarifying assignments and managing the work must also be outlined.

The case for action in context

Bridging the quality chasm between chronic pain and the care process will come from a unique confluence of opinion from all stakeholders committed within a focused community of practice to address the impact of chronic pain on the one in five Canadians who suffer chronic, debilitating pain. Need for improvement in care is announced abundantly in the literature. Operational solutions with traction must be applied to this national burden through an organizational logic model to support mainstream structures at the local level. At the outset, specific action items that must be addressed will include a survey and inventory of needs, best practices, and an inventory of services and gaps coupled with stakeholder validation of gaps, preferences and priorities. It is anticipated that through activities involving all stakeholder groups, a national program will develop a defensible case for a paradigm shift in health, based on the principle that the generation, translation and diffusion of evidence is the purview and product of research-community engagement if reliable knowledge is to be socially robust.

The overarching impact anticipated from the outputs of a managed approach to research is to promote and sustain a balanced portfolio of curiosity-based and needs-based research which, along with existing knowledge, can be mobilized and applied for the benefit of Canadians, the Canadian health system and the Canadian economy. This would have to be accomplished through the key outcomes of improving effectiveness and efficiency of health and social services support for those who suffer chronic, debilitating pain with a focus on disability that addresses enhanced life habits participation, prevention of deterioration, and optimization of health service usage to address the huge burden of chronic pain disorders and disabilities.

The added value of such a national program is as follows. Creation of a unified stakeholder voice to make recommendations for public and voluntary sector policy development to address the huge burden of chronic pain. Creation of a defensible case for research-community interactions in this domain, to ensure that reliable knowledge addresses burden by being socially robust. Enhancement of the funding base supporting operations and, on this basis, potential for leverage from the public purse, for incremental funding.

CONCLUSIONS

Pain has traditionally been seen as secondary to something else – the result is undertreated or untreated pain, unnecessary suffering, heavier reliance on the health care system, loss of productivity in the work force, absenteeism, increased comorbidity, and a cycle of illness that overall exacts a heavy toll on Canadians and the Canadian economy. Reliable evidence is available that these burdens can be markedly reduced when available knowledge is applied. Various levels of success in this regard have been demonstrated when there is exchange, synthesis and ethically sound application of research findings within a complex set of interactions among researchers and knowledge users. A nationally based managed approach to pain research is needed to accelerate the capture of the benefits of research for Canadians through improved health, more effective and responsive services and products, and a strengthened health care system through promoting and integrating success stories in health reform and health care reform across Canada.

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