Physician-to-physician telephone consultations for chronic pain patients: A pragmatic randomized trial

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BACKGROUND: The impact of telephone consultations between pain specialists and primary care physicians regarding the care of patients with chronic pain is unknown.

OBJECTIVES: To evaluate the impact of telephone consultations between pain specialists and primary care physicians regarding the care of patients with chronic pain.

METHODS: Patients referred to an interdisciplinary chronic pain service were randomly assigned to either receive usual care by the primary care physician, or to have their case discussed in a telephone consultation between a pain specialist and the referring primary care physician. Patients completed a numerical rating scale for pain, the Pain Disability Index and the Short Form-36 on referral, as well as three and six months later. Primary care physicians completed a brief survey to assess their impressions of the telephone consultation.

RESULTS: Eighty patients were randomly assigned to either the usual care group or the standard telephone consultation group, and 67 completed the study protocol. Patients were comparable on baseline pain and demographic characteristics. No differences were found between the groups at six months after referral in regard to pain, disability or quality of life measures. Eighty percent of primary care physicians indicated that they learned new patient care strategies from the telephone consultation, and 97% reported that the consultation answered their questions and helped in the care of their patient.

DISCUSSION: Most primary care physicians reported that a telephone consultation with a pain specialist answered their questions, improved their patients’ care and resulted in new learning. Differences in patient status compared with a usual care control group were not detectable at six-month follow-up.

CONCLUSIONS: While telephone consultations are clearly an acceptable strategy for knowledge translation, additional strategies may be required to actually impact patient outcomes.

Key Words: Chronic pain; Consultation; Knowledge translation; Primary care; Specialist; Telephone

Chronic pain is a common health condition (1,2), whose prevalence and impact will continue to increase as the population ages (3). Pain is the subject of one in three primary care visits (4); however, primary care physicians (PCPs) report lacking confidence and expertise in managing it (5). This gap between patient need and providers’ ability has been attributed to a lack of training, access to guidelines and tools supporting best practice (6,7).

While there is a growing consensus that the most complex chronic pain cases are best cared for by tertiary care multidisciplinary teams (8,9), access to these is a barrier. Most teams are located in urban centres and have an average wait time of six months (7). Wait times of this duration are associated with adverse quality of life and mental health outcomes (10).

Most chronic pain care will continue to be provided in primary care settings, hence, the importance of developing and implementing strategies for assisting PCPs with pain-related best practices. One such strategy is direct consultation between pain specialists and PCPs without the patient being present; a ‘curbside consultation’ (11,12).

Les consultations téléphoniques entre médecins pour les patients atteints de douleur chronique : un essai aléatoire pragmatique

HISTORIQUE: On ne connaît pas les effets des consultations téléphoniques entre spécialistes de la douleur et médecins traitants au sujet des soins des patients atteints de douleur chronique.

OBJECTIFS: Évaluer l’effet de consultations téléphoniques entre spécialistes de la douleur et médecins traitants au sujet des patients atteints de douleur chronique.

MÉTHODOLOGIE: Les patients dirigés vers un service interdisciplinaire de gestion de la douleur chronique étaient attribués au hasard entre les soins habituels du médecin traitant ou une consultation téléphonique de leur cas entre un spécialiste de la douleur et le médecin traitant. Les patients ont rempli une échelle d’évaluation numérique de la douleur, l’indice d’incapacité causé par la douleur et le formulaire court en 36 questions lors de l’aiguillage, puis trois et six mois plus tard. Les médecins traitants ont répondu à un bref sondage pour évaluer leurs impressions de la consultation téléphonique.

RÉSULTATS: Quatre-vingts patients ont été répartis au hasard entre le groupe de soins habituels ou le groupe de consultation téléphonique standardisé, et 67 ont terminé le protocole d’étude. Les patients partageaient une douleur et des caractéristiques démographiques comparables en début d’étude. Six mois après l’aiguillage, les chercheurs n’ont constaté aucune différence entre les groupes quant aux mesures de la douleur, de l’incapacité ou de la qualité de vie. De plus, 80 % des médecins traitants présentaient des résultats qui semblaient acceptables et pourraient être adoptés d’autres stratégies pour parvenir à de réels résultats cliniques chez les patients.
The present study used a pragmatic, randomized experimental design to explore whether telephone consultations with pain specialists were acceptable to PCPs and led to benefits for patients compared with wait list controls. The trial was registered at www.clinicaltrials.gov (NCT01923324).

METHODS

Research questions

For the comparison of ‘usual care’ (UC), with care that included a ‘standard telephone consultation’ (STC) between patients’ PCPs and a pain specialist, the following questions were addressed:

1. Were there changes in patient outcomes?
   a. Pain
   b. Perceived disability
   c. Quality of life

2. Is it acceptable?
   a. Is STC perceived as a knowledge transfer strategy by the PCP
   b. Is the PCP satisfied

3. Do resource requirements of the Chronic Pain Centre (CPC) change?

Eligibility

Patients were eligible for the present study if they were referred to the Alberta Health Services Calgary CPC with neurological or musculoskeletal chronic pain, they met the CPC’s referral criteria (chronic pain of >3 months’ duration, not a Workers Compensation client) and were competent in English to complete study questionnaires. Patients were excluded if: they were at increased risk for suicide; they had a very complex case; the referring PCP could not reliably identify the patient’s pain problem; or their condition warranted expedited CPC referral. Very complex cases (eg, widespread pain associated with very significant psychiatric or psychological disease) were identified using the preassessment questionnaire patients were asked to complete after referral to the CPC.

All PCPs and patients included in the study provided consent to participate.

Sample size

A sample of 100 patients (50 per group), assuming 20% attrition to follow-up, was calculated as being able to detect a minimally clinically relevant difference between the groups for a mean pain score on the numerical rating scale (NRS) of >2, with 80% power at the 5% significance level.

Outcome measures

Outcome measures included an NRS for pain intensity (13), the Pain Disability Index (PDI) (14), the Short Form (SF)-36 Health Survey (15), a seven-point Patient Global Impression of Change scale, the Pain Treatment Satisfaction Scale (16), a PCP satisfaction questionnaire and a knowledge transfer questionnaire. Resource use at the CPC comparing UC with STC was estimated using the CPC’s resource use tracking system.

Intervention

For patients randomly assigned to the STC group, the telephone consultation was scheduled at a time mutually convenient for both the PCP and the pain specialist, shortly after referral was received at the CPC. The consultation followed a semistructured interview format designed to: clarify the PCP’s goals and expectations of treatment; review the pain problem; consider relevant comorbidity and previous pain treatment; and make suggestions for future pain management by the PCP. Immediately following the consultation, the pain specialist dictated a summary of proceedings and faxed it, along with appropriate treatment protocols and patient information materials, to the PCP.

Physician telephone consultations

Research questions

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The PCP could use this information as he/she saw fit for the best care of the patient. A routine follow-up telephone call, in a semi-structured interview format designed to clarify the patient’s progress, as well as to problem solve and refine the management recommendations between the two physicians, was scheduled three months after the structured telephone consultation. At the six-month follow-up telephone call, the PCP was asked specific questions regarding the process, satisfaction with the process and perceived knowledge transfer.

Patients randomly allocated to the UC group received an appointment for assessment by a CPC pain specialist approximately six months after randomization, which was representative of normal referral wait times. The UC group did not receive any additional information from the CPC.

Outcome assessments were collected from both patient groups at three and six months. Patients were blinded as to group assignment.

Study design

Patients were allocated to either the UC or STC group using a random permuted block design, with assignments contained in opaque, numbered envelopes held by the study coordinator. Cases assigned to the STC group were the subject of a telephone discussion between the pain specialist and the referring PCP. Cases assigned to the UC group were entered into normal CPC wait list processes. Baseline assessment of outcome was derived from the routine preassessment questionnaire completed by all CPC patients. End point data collection was collected using self-completed mail surveys for all patients and physicians consenting to participate in the study.

Clinical outcomes were measured three and six months after the STC for the STC group, and at three months after referral and at the time of CPC assessment for the UC group (approximately six months after referral). PCP satisfaction was recorded after each STC contact. The present study was approved by the Conjoint Health Research Ethics Board at the University of Calgary (Calgary, Alberta).

RESULTS

Eighty subjects were recruited for the present study, and two withdrew (one from each group). The research database contained 78 subjects (41 in the STC group and 37 in the UC group). Characteristics of
patient participants are provided in Table 1. A full complement of 100 subjects was unable to be collected due to funding and time restraints.

Were there changes in patient outcomes?

NRS average pain: Both groups demonstrated a modest, but insignificant, decrease in pain (according to the NRS) over the study period (Figure 1). The mean reduction in average pain score at three months was 0.91 for the STC group and 0.79 for the UC group (P=0.708 [t test]). At six months the reduction was 0.96 and 1.28, respectively (P=0.552 [t test]). The proportion of subjects with a difference of >2 units (considered to be a clinically meaningful decrease in pain score [13]) was considered across the treatment groups for both the three-month and the six-month data. Four of 34 (11.8%) patients in the STC group and three of 29 (10.3%) patients in the UC group had at least a 2-unit reduction in pain after three months (P=0.858 [Z test]). At six months, five of 24 (20.8%) patients in the STC group and five of 25 (20.0%) patients in the UC group had a reduction of at least 2 units (P=0.942 [Z test]).

NRS worst pain: The largest reduction in the least amount of pain experienced by patients occurred between baseline and six months for both groups, and was similar for the STC and UC groups; the mean reduction at three months was 0.96 for the STC group and 0.74 for the UC group (P=0.775). At six months, the mean reduction in score for the least amount of pain was 0.92 and 0.78, respectively (P=0.796).

NRS current pain: Reduction in current pain was greater, but insignificant, at three months for the STC group at 1.18 and 0.55 for the UC group (P=0.180). At six months, the mean reduction in current pain scores were 0.83 and 0.84, respectively (P=0.991).

Patient global perception of change: Only 47 patients responded to this question (23 from the STC group and 24 from the UC group), and no statistically significant differences were observed according to a one-sided Mann-Whitney U test (P=0.566). Additionally, the proportion of patients indicating ‘much improved’ or ‘very much improved’ was considered according to intervention group. In the STC group, eight of 23 patients indicated substantial improvement, compared with three of 24 patients in the UC group (P=0.072). To further investigate this, a logistic regression model was considered, again, yielding a treatment effect that was insignificant (P=0.082).

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The mean pain scores for the worst pain experienced by patients were slightly lower, but insignificant, for the STC group at three months after exposure to the intervention (0.61 mean score reduction in the STC group and 0.10 in the UC group [P=0.211]). At six months, the mean reduction was 0.38 and 0.93, respectively (P=0.362).

SF-36 physical dimension: For the SF-36 physical dimension, the increase at three months was 2.21 in the STC group and 1.66 in the UC group (P=0.693). At six months the increase was 2.07 and 2.64, respectively (P=0.747). Neither increase was statistically significant.

SF-36 mental health dimension: For the SF-36 mental dimension scores, uniform decreases were observed indicating worsening mental health status in both groups. The reduction at three months was 2.34 in the STC group and 3.18 in the UC group (P=0.719). At six months the reduction was 0.81 and 3.52, respectively (P=0.399).

Is it acceptable?

Knowledge transfer related to evidence-based practice: Based on the exact binomial distribution, it can be reported with 95% confidence that the proportion of PCPs in the STC group affirming their experience of evidence-based learning (either agree or strongly agree) was between 63.1% and 91.6%.

Primary care physician satisfaction with STC process and outcomes: Table 2 presents the responses by PCPs in the STC group to questions about the process and outcomes of the intervention, along with the corresponding binomial exact CIs. These indicate substantial support for STCs in clinical practice.

Do the resource requirements of the CPC change?

Resource utilization: To estimate whether the availability of STCs reduced the consumption of CPC resources compared with resources used by patients receiving UC, the time spent in direct and indirect CPC staff contact was calculated (Figure 2). CPC resource utilization with reference to pain specialists and other CPC team members (nursing, psychology, occupational therapy, physiotherapy, social work, kinesthesiology etc) for direct care, indirect care and for total care was similar for both the STC and UC groups. No reduction in resource utilization in association with STC use was evident.

DISCUSSION

Structured telephone consultations between pain specialists and PCPs were very well received by recipients as a knowledge transfer strategy, and were perceived as contributing to improved quality of care. However, no statistically significant differences in patient outcomes were observed between the STC and UC groups.

There are a number of possible explanations for this discrepancy. The failure to fully recruit the intended sample caused the study to be...
underpowered. This is supported by the tendency of the STC group to have more positive average outcome scores than the control (ie, UC) group. The wide variability in individual scores diminishes the likelihood of detecting group differences in an underpowered study and highlights the inconsistency of response to the treatment plan. The lack of significant benefit for STC patients may be related to inconsistent exposure to the recommended treatments. It is possible that the recommendations were not carried out, either because the PCP or the patient objected, or because the patient did not return to see the PCP in time for the possible benefits of treatment to be observed when follow-up occurred. It is also possible that when the treatment plan called for sequential titrated medication trials or patient participation in community-based self-management education, the three and six month follow-up intervals were not synchronized to capture the treatment effects that may have occurred. It is also feasible that the pain management introduced by PCPs ahead of any specialist consultation was, in practice, as effective as that recommended by the pain specialists.

However, the strong response from PCPs in favour of telephone consultation indicates that the intervention may have merit. Chronic pain is notoriously difficult to treat. Duration of pain has a significant effect on the likelihood of therapeutic response, and more timely advice regarding treatment may be helpful to PCPs. The ability to obtain quick reassurance that one is doing all one reasonably can to assuage the problem is understandably beneficial, and may lead to service efficiencies if it reduces unnecessary referrals and empowers PCPs to manage pain without pain specialist intervention. More strategically, telephone consultations involve a specialist and a generalist engaging in a learning process that may be transferred to similar cases and circumstances. In addition, telephone consultations potentially require less investment in scheduling, travel, office resources and compensation-related issues than conventional referral, and have potential to be a widely available and easily accessible resource.

Future studies should consider a detailed health economic analysis of different types of consultation processes, and should consider other options for communicating with PCPs such as eConsult (17) and faxing recommendations (18) to PCPs.

SUMMARY
The present trial demonstrated that structured telephone consultations between pain specialists and PCPs are a well accepted strategy for communicating evidence-based approaches regarding chronic pain management and leads to conditions of equipoise in patient clinical health status when compared with patients undergoing usual care. Standard telephone consultations for PCPs continue to be offered by the Calgary CPC and have become a standard of care.

FUNDING: Alberta Heritage Foundation for Medical Research (currently Alberta Innovates – Health Solutions)

DISCLOSURES: The authors have no financial relationships or conflicts of interest to declare.
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