Can Medical Cannabis Therapies be Cost-Effective in the Non-Surgical Management of Chronic Knee Pain?

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ABSTRACT

INTRODUCTION: Chronic knee pain is a common musculoskeletal condition, which usually leads to decreased quality of life and a substantial financial burden. Various non-surgical treatments have been developed to relieve pain, restore function and delay surgical intervention. Research on the benefits of medical cannabis (MC) is emerging supporting its use for chronic pain conditions. The purpose of this study was to evaluate the cost-effectiveness of MC compared to current non-surgical therapies for chronic knee pain conditions.

METHODS: We conducted a cost-utility analysis from a Canadian, single payer perspective and compared various MC therapies (oils, soft gels and dried flowers at different daily doses) to bracing, glucosamine, pharmaceutical-grade chondroitin oral non-steroidal anti-inflammatory drugs (NSAIDs), and opioids. We estimated the quality-adjusted life years (QALYs) gained with each treatment over 1 year and calculated incremental cost-utility ratios (ICURs) using both the mean and median estimates for costs and utilities gained across the range of reported values. The final ICURs were compared to willingness-to-pay (WTP) thresholds of $66,714, $133,428 and $200,141 Canadian dollars (CAD) per QALY gained.

RESULTS: Regardless of the estimates used (mean or median), both MC oils and soft gels at both the minimal and maximal recommended daily doses were cost-effective compared to all current knee pain therapies at the lowest WTP threshold. Dried flowers were only cost-effective up to a certain dosage (0.75 and 1 g/day based on mean and median estimates, respectively), but all dosages were cost-effective when the WTP was increased to $133,428/QALY gained.

CONCLUSION: Our study showed that MC may be a cost-effective strategy in the management of chronic knee pain; however, the evidence on the medical use of cannabis is limited and predominantly low-quality. Additional trials on MC are definitely needed, specifically in patients with chronic knee pain.

KEYWORDS: Cost-effectiveness, cost-utility, chronic knee pain, medical cannabis

Introduction

Chronic knee pain is a common musculoskeletal condition, which usually leads to disability, decreased quality of life and a substantial financial burden.¹ Knee osteoarthritis (OA) is one of the leading causes of chronic knee pain.² The Global Burden of Disease 2010 study estimated that the prevalence of radiographically confirmed knee OA was about 4% in the global population, and knee OA was ranked as the 11th highest contributor to disability worldwide.³

On the basis of different hypotheses surrounding the pathophysiology of knee OA, various treatments have been developed and researched, all of which intend to relieve pain, restore function and delay the necessity for a surgical joint replacement. Current non-surgical options for knee OA pain-relief include treatments such as opioids, non-steroidal anti-inflammatory drugs (NSAIDs), acetaminophen, viscosupplementation, glucosamine, chondroitin and injection therapies, as well as more conservative means such as exercise, weight control and bracing. Given the number of available interventions for these patients, there is an ongoing emphasis on establishing how they compare to each other in terms of treatment effects, patient satisfaction and tolerability and healthcare costs.⁴

Research on the potential analgesic benefits of medical cannabis (MC) is emerging. Cannabis works on the endocannabinoid system – one of the body’s natural analgesic systems and a viable target for reducing pain.⁵ Delta-9-tetrahydrocannabinol (THC) and cannabidiol (CBD) are 2 major components of MC. They both treat pain, but only THC has psychoactive properties which can cause a sense of euphoria and heightened sensory perception.⁶ THC and its analogues have been synthesized for pharmaceutical use for years, whereas the pace of synthesising CBD was slower. Though guidance and suggestions exist, no universally accepted standards have been established on the proper dosing of MC products in chronic pain populations.³ Currently, on the global market, dronabinol and nabilone are synthetic forms of THC, and Nabiximol and Sativex are a...
combination of THC and CBD; however, these cannabinoid-based products are manufactured by pharmaceutical companies whereas MC can potentially be a natural form of therapy.

There is also a risk of adverse effects with MC, which primarily occur with products containing higher levels of THC, as it has been shown to be associated with psychoactive effects.\(^8,9\) It is important to consider the specific side-effects that are associated with MC, which include fatigue/drowsiness, dizziness, dry mouth, cough/phlegm/bronchitis (when smoked), anxiety, euphoria, nausea and some effects on cognitive ability, as their potential impact on a person’s day-to-day life can vary between patients.\(^8,10\)

There has been a growing body of evidence supporting the use of MC for chronic pain conditions, such as neuropathic pain,\(^11,12\) pain associated with multiple sclerosis\(^13\) and fibromyalgia.\(^14\) In terms of treating arthritis-related pain, published clinical trials are very few; however, a number of trials are ongoing, such as the CBD Treatment in Hand Osteoarthritis and Psoriatic Arthritis (NordCAN, NCT03693833) and the Cannabinoid Profile Investigation of Vapourized Cannabis in Patients With Osteoarthritis of the Knee (CAPRI, NCT02324777) studies. Other than its analgesic effect, MC also shows some effects that may be highly desirable to patients with chronic pain. For example, adding MC to a patient’s treatment protocol could decrease the amount of opioids needed for pain relief and, consequently, reduce the likelihood of opioid-related adverse effects and addiction.\(^15\)

Cost-effectiveness analysis (CEA) compares the relative costs and health benefits of different interventions.\(^16\) An intervention is considered cost-effective, relative to another intervention, when the incremental cost-effectiveness ratio (ICER), or cost-utility ratio (ICUR), is less than one’s willingness to pay for the added health benefit. One commonly used measure of health benefit is the quality-adjusted life years (QALYs), which incorporates both the quality and quantity of life in a given health state.\(^17\)

Past cost-effectiveness studies on the use of MC products have been conducted in the context of multiple sclerosis,\(^18\) but, as there is limited evidence on MC for chronic knee pain or knee arthritis, there is currently no cost-effectiveness study on these topics. Little is known about whether the health benefits provided by MC could be a cost-effective method for treating chronic knee pain. The purpose of this study was to evaluate the cost and health benefits of MC compared to non-surgical treatments currently used to treat chronic knee pain conditions from a Canadian, single payer perspective.

Methods

Literature search

We conducted a systematic literature search in the MEDLINE database from inception to November 19th, 2019 to collect utility and cost data from published clinical trials or economic evaluations. We developed the structured search strategies using indexed terms and free-text terms related to the patient population (ie, chronic knee pain or knee arthritis) and interventions (ie, cannabis, opioids, nonsteroidal anti-inflammatory drugs (NSAIDs), bracing, glucosamine and chondroitin). We selected these interventions for comparison due to their similarities with MC in terms of their frequencies and methods of administration (ie, daily use/intake and oral ingestion [for pharmacological or dietary supplements]); injection therapies are generally considered a non-surgical, pharmacological intervention as well, but only require a single injection procedure or a small number of injections over just a few weeks, so we deemed such treatments unsuitable for comparison with MC. We included studies published in English, but did not restrict our search by the publication date. We also searched the references lists of included studies and previously performed related reviews for additional eligible articles. In terms of the inclusion of clinical trial data, we prioritized randomized controlled trials (RCTs). We only included trials that provided sufficient baseline and follow-up data so that we could calculate the utilities gained in the study.

After completing the search, we identified no eligible clinical trial data on MC in a chronic knee pain population; therefore, we included outcome data from studies investigating MC for other chronic, non-cancer pain conditions and used this information to quantify utility scores for this treatment. We did not restrict the inclusion of such trials to RCTs due to the limited evidence base on this topic.

For cost data, we prioritized studies conducted in Canada to ensure the values were most representative of costs that would be incurred in Canada. If we did not identify any cost studies conducted in Canada for a given treatment, we referred to studies performed in the United States (US) and included their cost data in our analysis. Then, if we did not locate any cost data for a given treatment in a US study, we referred to economic literature outside of North America.

Data extraction

Three reviewers independently extracted the relevant data. We created the data extraction forms Google sheets and pilot-tested it across the reviewers. We examined all extracted data in duplicate and resolved any discrepancies through discussion. We extracted study characteristics, outcome data expressed as a utility score or that could be converted to a utility score via an established mapping algorithm,\(^19-23\) and relevant cost data related to the treatment medication, additional prescription time (if applicable), and required concomitant therapies (if applicable).

Treatment utility scores

We used quality of life (QoL) outcomes to estimate utility scores or converted outcome measure scores to utilities using previously published mapping algorithms. These included the following outcome measures: (1) EuroQoL5D (EQ-5D), (2) Western Ontario and McMaster Universities Osteoarthritis Index
(WOMAC), (3) 36-item or 12-item Short Form Survey (SF-36 or SF-12), (4) Knee Injury and Osteoarthritis Outcome Score (KOOS), (5) Fibromyalgia Impact Questionnaire (FIQ), (6) 4-item Patient Health Questionnaire, (7) Health Utilities Index Mark 3 (HUI) and (8) Health Assessment Questionnaire (HAQ). We ensured all utility scores were on a 0 to 1 scale, where 1 represented perfect health. When multiple utility scores were available for a given treatment, we considered the entire range of utility scores and calculated a mean and median of this range.

Cost data

We used a Canadian, single payer perspective with respect to costs. For current knee pain therapies (ie, opioids, nonsteroidal anti-inflammatory drugs (NSAIDs), bracing, glucosamine and chondroitin), we acquired cost data for a given treatment from the economic literature, prioritising studies conducted in Canada; otherwise, we referred to studies conducted in the US and, then, outside of North America for treatment costs. For the cost of MC products, we referred to the Spectrum Therapeutics website. We converted all cost data to 2019 Canadian dollars (CAD), using the Bank of Canada Inflation Calculator when needed.

When multiple cost estimates were available for a given treatment, we considered the entire range of costs and calculated an arithmetic mean and median of this range. For MC, celecoxib and opioids, only we considered medication costs. For bracing, we included both the cost of the orthotic device and time spent on brace fitting. For non-selective NSAIDs (diclofenac, naproxen and ibuprofen), we also added the cost of a proton-pump inhibitor for gastrointestinal protection. For the cost of diclofenac, we applied the same cost estimates retrieved for naproxen and ibuprofen, as we did not identify any cost data specific to diclofenac. We considered any other costs related to the treatment of chronic knee pain to be equivalent between therapies.

Cost-utility analysis

We calculated cost-utility ratios as the cost per QALY gained. For each included study, we subtracted the average utility score at the study’s latest follow-up (up to 1 year) by the average baseline utility score. We then calculated incremental cost-utility ratios (ICURs) between treatments under 2 different scenarios:

1. Mean values from the range of costs and utilities gained
2. Median values from the range of costs and utilities gained

Based on the available cost data provided on the Spectrum Therapeutics website, we classified MC products as:

- Oils
- Soft gels
- Dried flowers

The minimal and maximal recommended doses of MC oils and soft gels were provided by a representative from Spectrum Therapeutics (Table 1), as was the price per gram of dried flowers. Oils and soft gels are priced differently depending on the THC:CBD content, so we calculated the mean and median price across the range of options, according to both minimal and maximal dosing recommendations. As there is no standard dosing recommendation for the inhalation of dried flowers, we examined outcomes across a range of different doses between 0.5 to 1.25 g per day, based on average per month consumption estimates in 2018 published by Statistics Canada. On the Spectrum Therapeutics website, dried flowers can be purchased in either a 2- or 15-g jar, which results in a different price per gram (cheaper with a 15-g jar at $6.53 vs $8 CAD per gram with a 2-g jar); for our analysis, we assumed the purchase of a 15-g jar. Due to the lack of evidence in this area, we could not differentiate treatment effects between the different types of MC (ie, oils, soft gels, dried flowers and the variable THC:CBD ratios); therefore, we assumed similar treatment effects across all MC products. We conducted data extraction and analysis using Microsoft Excel (Version 2010, Microsoft, Redmond, WA, USA) over a 1-year time horizon.

For comparisons against a willingness-to-pay (WTP) threshold, we used values of $50000, $100000 and $150000 USD, which, currently, are approximately equal to $66714, $133428 and $200141 CAD, respectively.

Table 1. Dosing guidance for medical cannabis.

| PRODUCT | THC:CBD | STARTING DAILY DOSE | MAXIMAL DAILY DOSE (IF NEEDED) |
|---------|---------|---------------------|-------------------------------|
| Oils    | 26.3:<1 mg/mL | 0.1 mL | 0.4 mL (↑ by 0.1 mL/day from day 1) |
|         | 10:10-20 mg/mL | 0.25 mL | 1 mL (↑ by 0.25 mL/day from day 1) |
|         | <1:20 mg/mL | 0.25 mL | 1 mL (↑ by 0.25 mL/day from day 1) |
| Soft gels | 2.5 mg THC capsules | 2.5 mg | 10 mg (↑ by 2.5 mg/day from day 1) |
|         | 10 mg THC capsules | 10 mg | 17.5 mg (↑ by 2.5 mg/day from day 1) |
|         | 5 & 20 mg CBD capsules | 5 mg | 30 mg (↑ by 5 mg/day from day 1) |
Results

Utility scores

The estimated utilities gained for each treatment are presented in Table 2, showing the mean, median and range of QALYs gained over 1 year across the included trials. Regardless of the estimate chosen, MC appears to provide the greatest QALYs gained among all treatments, while ibuprofen results in the least. Generally speaking, pharmaceutical-grade chondroitin and opioids also provide more favourable gains in QALYs over 1 year.

Costs

Treatment costs are summarized in Table 3. Based on the average estimate of cost data, MC oils can range from $164 to...

Table 2. Utility scores (QALYs gained over 1 year).

| TREATMENT                              | ESTIMATE BASED ON MEAN VALUE | ESTIMATE BASED ON MEDIAN VALUE | RANGE OF VALUES |
|----------------------------------------|------------------------------|--------------------------------|-----------------|
| Medical cannabis                       | 0.052                        | 0.068                          | 0.004–0.071     |
| Bracing                                | 0.015                        | 0.014                          | 0.008–0.025     |
| Glucosamine                            | 0.022                        | 0.019                          | 0.003–0.057     |
| Chondroitin (pharmaceutical-grade)     | 0.034                        | 0.041                          | 0.008–0.045     |
| Celecoxib                              | 0.026                        | 0.021                          | 0.003–0.089     |
| Diclofenac                             | 0.023                        | 0.014                          | 0.006–0.052     |
| Naproxen                               | 0.022                        | 0.018                          | 0.009–0.043     |
| Ibuprofen                              | 0.011                        | 0.011                          | 0.007–0.014     |
| Opioids (Tramadol, oxycodone)          | 0.041                        | 0.042                          | 0.021–0.06      |

Table 3. Costs for 1 year of treatment (2019 CAD, rounded to nearest dollar).

| TREATMENT                              | DELIVERY METHOD | ESTIMATE BASED ON MEAN VALUE | ESTIMATE BASED ON MEDIAN VALUE |
|----------------------------------------|-----------------|------------------------------|--------------------------------|
| Medical cannabis                       | Oral (for oils and soft gels) | Oils (minimal dose): $164 | Oils (minimal dose): $205 |
|                                        |                 | Oils (maximal dose): $657 | Oils (maximal dose): $821 |
|                                        |                 | Soft gels (minimal dose): $616 | Soft gels (minimal dose): $389 |
|                                        |                 | Soft gels (maximal dose): $1296 | Soft gels (maximal dose): $1314 |
|                                        | Smoked (for dried flowers) | Dried flowers (0.5 g/day): $1192 | Dried flowers (0.5 g/day): $1192* |
|                                        |                 | Dried flowers (0.75 g/day): $1788 | Dried flowers (0.75 g/day): $1788* |
|                                        |                 | Dried flowers (1 g/day): $2383 | Dried flowers (1 g/day): $2383* |
|                                        |                 | Dried flowers (1.25 g/day): $2979 | Dried flowers (1.25 g/day): $2979* |
| Bracing                                | Orthotic        | Device + time on brace fitting: $551 | Device + time on brace fitting: $341 |
| Glucosamine                            | Oral            | $401                         | $401*             |
| Chondroitin (pharmaceutical-grade)     | Oral            | $811                         | $811*             |
| Celecoxib                              | Oral            | $2760                        | $1616             |
| Diclofenac                             | Oral            | Medication + PPI: $1376      | Medication + PPI: $1536 |
| Naproxen                               | Oral            | Medication + PPI: $1490      | Medication + PPI: $1536 |
| Ibuprofen                              | Oral            | Medication + PPI: $1205      | Medication + PPI: $1410 |
| Opioids (Tramadol, oxycodone)          | Oral            | $1834                        | $1318             |

*Mean estimate = median estimate.
$657 per year, and soft gels can range from $616 to $1296 per year; the yearly cost of smoking dried flowers can vary dramatically depending on how many grams are smoked per day. For current knee pain therapies, orthotic intervention with bracing and dietary supplementation with glucosamine are the cheapest therapies, pharmaceutical-grade chondroitin is priced in the mid-range, and oral NSAIDs and opioids are, generally, priced highest.

Cost-utility

Based on mean estimates (Table 4) and a WTP threshold of $66,714 CAD, all forms of MC consumption were cost-effective compared to any knee pain therapy, except dried flowers at 1 and 1.25 g/day; dried flowers at 1 g/day compared to pharmaceutical-grade chondroitin and dried flowers at 1.25 g/day versus glucosamine, pharmaceutical-grade chondroitin and opioids were only cost-effective when the WTP was increased to $133,428 CAD/QALY gained.

Based on median estimates (Table 5) and a WTP threshold of $66,714 CAD, all forms of MC consumption were cost-effective compared to any knee pain therapy, except dried flowers at 1.25 g/day; dried flowers at 1.25 g/day versus pharmaceutical-grade chondroitin were only cost-effective when the WTP was increased to $133,428 CAD/QALY gained.

Regardless of the estimates used (ie, mean or median estimates), both MC oils (Figure 1) and soft gels (Figure 2) at both the minimal and maximal recommended daily doses were cost-effective compared to all current knee pain therapies at the lowest WTP threshold (ie, $66,714/QALY gained). Medical cannabis dried flowers were only cost-effective compared to all

### Table 4. Incremental cost-utility ratios (ICURs) based on mean utility and mean cost estimates (cost/QALY gained).

| MEDICAL CANNABIS VS. | BRACING | GLUCOS. | CHON. | CELE. | DICLO. | NAPROX. | IBU. | OPIOIDS |
|---------------------|---------|---------|-------|-------|-------|---------|------|---------|
| Oils (min)          | Dom.    | Dom.    | Dom.  | Dom.  | Dom.  | Dom.    | Dom. | Dom.    |
| Oils (max)          | $2886*  | $8518*  | Dom.  | Dom.  | Dom.  | Dom.    | Dom. | Dom.    |
| Soft gels (min)     | $1766*  | $7159*  | Dom.  | Dom.  | Dom.  | Dom.    | Dom. | Dom.    |
| Soft gels (max)     | $20148* | $29830* | $26938* | Dom. | Dom. | Dom.    | Dom. | Dom.    |
| Dried flowers (0.5 g/day) | $17320* | $26342* | $21126* | Dom. | Dom. | Dom.    | Dom. | Dom.    |
| Dried flowers (0.75 g/day) | $33424* | $46204* | $54229* | Dom. | $14189* | $9913* | $14211* | Dom. |
| Dried flowers (1 g/day) | $49529* | $66066* | $87332* | Dom. | $34736* | $29775* | $28744* | $49967* |
| Dried flowers (1.25 g/day) | $65633* | $85928* | $120436* | $8421* | $55283* | $49637* | $43277* | $104136* |

Abbreviations: Cele., celecoxib; Chon., chondroitin; Diclo., diclofenac; Dom., dominated (ie, MC product is more effective and cheaper than the comparator treatment); Glucos., glucosamine; Ibup., ibuprofen; Naprox., naproxen.

Positive ICUR indicates that the MC product is more effective but also more costly than the comparator treatment.

*Cost-effective at a WTP of $66,714 CAD/QALY gained.

#Cost-effective when WTP increased to $133,428 CAD/QALY gained only.

### Table 5. Incremental cost-utility ratios (ICURs) based on median utility and median cost estimates (cost/QALY gained).

| MEDICAL CANNABIS VS. | BRACING | GLUCOS. | CHON. | CELE. | DICLO. | NAPROX. | IBU. | OPIOIDS |
|---------------------|---------|---------|-------|-------|-------|---------|------|---------|
| Oils (min)          | Dom.    | Dom.    | Dom.  | Dom.  | Dom.  | Dom.    | Dom. | Dom.    |
| Oils (max)          | $8896*  | $8567*  | $362* | Dom.  | Dom.  | Dom.    | Dom. | Dom.    |
| Soft gels (min)     | $886*   | Dom.    | Dom.  | Dom.  | Dom.  | Dom.    | Dom. | Dom.    |
| Soft gels (max)     | $18021* | $18623* | $18612* | Dom. | Dom. | Dom.    | Dom. | Dom.    |
| Dried flowers (0.5 g/day) | $15757* | $16128* | $14084* | Dom. | Dom. | Dom.    | Dom. | Dom.    |
| Dried flowers (0.75 g/day) | $26791* | $28288* | $36153* | $3650* | $4668* | $5041* | $6618* | $18080* |
| Dried flowers (1 g/day) | $37826* | $40449* | $58221* | $16328* | $15702* | $16958* | $17071* | $40998* |
| Dried flowers (1.25 g/day) | $48860* | $52609* | $80290* | $29006* | $26736* | $28875* | $27525* | $63916* |

Abbreviations: Cele., celecoxib; Chon., chondroitin; Diclo., diclofenac; Dom., dominated (ie, MC product is more effective and cheaper than the comparator treatment); Glucos., glucosamine; Ibup., ibuprofen; Naprox., naproxen.

Positive ICUR indicates that the MC product is more effective but also more costly than the comparator treatment.

*Cost-effective at a WTP of $66,714 CAD/QALY gained.

#Cost-effective when WTP increased to $133,428 CAD/QALY gained only.

$657 per year, and soft gels can range from $616 to $1296 per year; the yearly cost of smoking dried flowers can vary dramatically depending on how many grams are smoked per day. For current knee pain therapies, orthotic intervention with bracing and dietary supplementation with glucosamine are the cheapest therapies, pharmaceutical-grade chondroitin is priced in the mid-range, and oral NSAIDs and opioids are, generally, priced highest.
knee pain therapies at the lowest threshold up to a certain dosage (0.75 and 1 g/day based on mean and median estimates, respectively).

**Discussion**

The purpose of this study was to estimate the cost-effectiveness of MC in patients with chronic knee pain. Prior cost-effective studies have looked at cannabinoid-based therapies in multiple sclerosis with Sativex and chronic neuropathic pain with smoked cannabis.\(^9^7\)-\(^1^0^1\) This analysis showed that, depending on the chosen WTP threshold, MC therapies could be a cost-effective strategy relative to current non-surgical knee pain therapies, namely, bracing, glucosamine, pharmaceutical-grade chondroitin, various oral NSAIDs and opioids. More specifically, MC oils and soft gels at both the minimal and maximal recommended daily doses were all cost-effective at the most conservative (ie, lowest) WTP value of $66,714 CAD (or $50,000 USD) per QALY gained compared to current knee therapies; MC consumption via dried flowers may only be cost-effective up to a certain daily dose, unless the WTP is increased to $133,428 CAD (or $100,000 USD) per QALY gained.

In terms of AEs, the extent to how related and how serious these AEs are must also be considered. The reported estimate for bracing appears high (ie, 67%), but this was just from 1 study and the nature of adverse effects related to the device are likely not severe, largely being associated with superficial skin irritation and patient discomfort while wearing the device.\(^3^5\) Glucosamine and chondroitin have been shown to be generally well-tolerated\(^1^0^2,\(^1^0^3\), however, certain patients may be allergic, some may experience minor gastrointestinal effects and they are usually administered with a salt, which would increase daily sodium intake.\(^1^0^3-\(^1^0^5\) In addition, there is
some uncertainty regarding glucosamine’s effect on glucose metabolism.\textsuperscript{103,104} Toxicities that have been considered related to chronic oral NSAID use, especially non-selective types, are gastrointestinal bleeding, erosion or ulceration, dyspepsia, chronic diarrhea, cardiovascular risks, renal failure and colonic perforation,\textsuperscript{106–109} while those related to opioids include constipation, respiratory depression, nausea, urinary retention, hyperalgesia, behavioural side-effects and, more importantly, overdose-related mortality due to addiction and increased tolerance.\textsuperscript{8,107,110} In order to limit the gastrointestinal effects of non-selective NSAIDs, gastrointestinal protective agents, such as proton pump inhibitors, and COX-2 inhibitors were developed, but those come at increased costs and there is still a risk.\textsuperscript{107,109} Also, COX-2 inhibitors may lessen gastrointestinal toxicity, but they may still increase the risk of adverse cardiovascular effects.\textsuperscript{107,110,111}

Adverse events related to MC are primarily seen with products containing higher levels of THC as it has been shown to be associated with the psychoactive effects of MC consumption.\textsuperscript{8,9} The current recommendation for THC consumption is no more than 30 mg/day and, preferably, in conjunction with CBD.\textsuperscript{8} In the 2 included trials that reported the risk of AEs following MC therapy, the 1 associated with a higher incidence of AEs (88% over 12 months) prescribed herbal MC with 12.5% THC,\textsuperscript{10} whereas the 1 reporting the lower incidence (10% over 3 months) initially provided patients with an oral MC capsule containing a 1:1 ratio of THC to CBD, though a vapor pen containing 2 mg of THC per 0.1 mg of CBD (20:1 ratio) was also prescribed to any patient experiencing breakthrough pain.\textsuperscript{30} The most common side-effects that have been seen with MC include fatigue/drowsiness, dizziness, dry mouth, cough/phlegm/bronchitis (when smoked), anxiety,
Strength and limitations

A strength of our study was that we retrieved our estimates for utility scores and cost data, when possible, from a systematic review of the literature and, except with MC, we extracted utility scores from RCT. We considered all eligible studies in order to limit bias in our final estimates. As our study was from a Canadian, single payer perspective, prioritized cost data from studies conducted in Canada, when feasible, so that our final cost estimates were most representative of what patients would have to pay in Canada. We also performed 2 types of analyses (ie, mean and median estimates) in order to assess if our final conclusions could change depending on the estimates included in our analysis.

Our study was limited by the fact that we had to use evidence from MC studies that were on patient populations not exactly representative of our target patient population (ie, chronic knee pain). The patients included in these MC studies predominantly had chronic, noncancer pain, though it is unclear if such patients benefit from MC differently than those who have chronic knee pain. Also, due to the limited evidence in this area, we had to include observational data to acquire utility estimates for MC. Another potential limitation was that, when utility scores were not provided in a study, we mapped scores from other outcome scoring measures in order to estimate utility scores. This included a heterogeneous set of different outcome measures and it is unclear how accurate they are at actually capturing utility. Lastly, again due to the limited evidence, we assumed that the effects of MC were the same regardless of the route of administration (ie, oils, soft gels or dried flowers), THC:CBD ratio or daily dosing, which may not truly be the case.

Conclusion

Our study showed that MC therapies may be a cost–effective strategy in the non-surgical management of chronic knee pain relative to current knee pain therapies; however, the evidence on the medical use of cannabis is limited and predominantly low-quality. Additional trials on MC are definitely needed, specifically in patients with chronic knee pain. Future investigations should also focus on establishing if there are any differences in therapeutic effects between the different methods of MC consumption, THC:CBD ratio and daily dosing. The relatedness and severity of adverse events must also be further evaluated as this should be an important part of the decision-making process due to their potential impact on a patient's quality of life and healthcare costs.

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REFERENCES

1. Kim W, Jin YS, Lee CS, Bin SI, Lee SY, Choi KH. Influence of knee pain and low back pain on the quality of life in adults older than 50 years of age. PM R. 2015;7:955-961.
2. Hadler NM. Knee pain is the malady–not osteoarthritis. Ann Intern Med. 1992;116:598-599.
3. Cross M, Smith E, Hoy D, et al. The global burden of hip and knee osteoarthri-ritic estimates from the global burden of disease 2010 study. Ann Rheum Dis. 2014;73:1323-1330.
4. Roosemann T, Wensing M, Joest K, Backenstraß M, Mahler C, Szczesny J. Problems and needs for improving primary care of osteoarthritic patients: the views of patients, general practitioners and practice nurses. BMC Musculoskelet Disord. 2006;7:48.
5. Kruse E, Riaux D, McDougall JJ. Mechanisms and mediators that drive arthritis pain. Curr Osteoporos Rep. 2015;13:216-224.
6. Philpott HT, O’Brien M, McDougall JJ. Articularization of early phase inflammatio-n by cannabidiol prevents pain and nerve damage in rat osteoarthritis. Pain. 2017;158:2442-2451.
7. Savage SR, Romero-Sandoval A, Schatman M, et al. Cannabis in pain treat-ment: clinical and research considerations. J Pain. 2016;17:654-668.
8. MacCallum CA, Russo EB. Practical considerations in medical cannabis admin-istration and dosing. Eur J Intern Med. 2018;49:12-19.
9. Levinsonn EA, Hill KP. Clinical uses of cannabis and cannabinoids in the United States. J Neurol Sci. 2020;411:116717.
10. Ware MA, Wang T, Shapiro S, Collet JP; COMPASS study team. Cannabis for the management of pain: assessment of safety study (COMPASS). J Pain. 2015;16:1233-1242.
11. Lee G, Grovey B, Furnish T, Wallace M. Medical cannabis for neuropathic pain. Curr Pain Headache Rep. 2018;22:8.
12. Lynch ME, Campbell F. Cannabinoids for treatment of chronic non-cancer pain: a systematic review of randomized trials. Br J Clin Pharmacol. 2011;72:735-744.
13. Nielsen S, Germansos R, Weier M, et al. The use of cannabis and cannabinoids in treating symptoms of multiple sclerosis: a systematic review of reviews. Curr Neurol Neurosci Rep. 2018;18:3.
14. Habib G, Arval S. Medical cannabis for the treatment of fibromyalgia. J Clin Rheumatol. 2018;24:255-258.
15. Narang S, Gibson D, Wasan AD, et al. Efficacy of dronabinol as an adjuvant treat-ment for chronic pain patients on opioid therapy. J Pain. 2008;9: 254-264.
16. Sanders GD, Neumann PJ, Basu A, et al. Recommendations for conduct, meth-odological practices, and reporting of cost-effectiveness analyses: second panel on cost-effectiveness in health and medicine. JAMA. 2016;316:1093-1103.
17. Berg J, Lindgren P, Fredrikson S, Kobelt G. Costs and quality of life of multiple sclerosis in Sweden. Eur J Health Econ. 2006;7(Suppl 2):S75-585.
18. Herzog S, Shanahan M, Grimison P, et al. Systematic review of the costs and benefits of prescribed cannabis-based medicines for the management of chronic illness: lessons from multiple sclerosis. PharmacoEconomics. 2018;36:67-78.
19. Ara R, Brazier J. Deriving an algorithm to convert the eight mean SF-36 dimension scores into a mean EQ-5D preference-based score from published studies (where patient level data are not available). Value Health. 2008;11:1331-1141.
20. Carreño A, Fernández I, Badía X, Varela C, Roset M. Using HAQ-DI to esti-mate HUI-3 and EQ-5D utility values for patients with rheumatoid arthritis in Spain. Value Health. 2011;14:192-200.
21. Collado-Mateo D, Chen G, Garcia-Gordillo MA, et al. Fibromyalgia and quality of life: mapping the revised fibromyalgia impact questionnaire to the preference-based instruments. Health Qual Life Outcomes. 2017;15:114.

22. Frank P, Luberkin EI, Gold MR, Tancredi DJ, Jia H. Mapping the SF-12 to the EuroQol EQ-5D index in a national US sample. Med Decis Making. 2004;24:247-254.

23. Grootendorst P, Marshall D, Pericak D, Bellamy N, Feeny D, Torrance GW. A model to estimate health utilities index mark 3 utility scores from WOMAC index scores in patients with osteoarthritis of the knee. J Rheumatol. 2007;34:534-542.

24. Spectrum Therapeutics. Accessed February 2020. https://shop.spectrumtherapeutics.com/collections/available

25. Bank of Canada. Inflation calculator. Accessed February 2020. https://www.bankofcanada.ca/rates/related/inflation-calculator/

26. Statistics Canada. StatsCannabis data availability: crowdsourced cannabis prices, third quarter 2018. Accessed February 2020. https://www150.statcan.gc.ca/n1/daily-qotidien/180104/dq180104-eng.htm

27. Cameron D, Uebes J, Norström F. On what basis are medical cost-effectiveness thresholds set? Clashing opinions and an absence of data: a systematic review. Glob Health Action. 2018;11:147828/1447828.

28. Marcotte E, Tarenex B, Kat D, Sato KG, Rosen S. Thresholds for the cost-effectiveness of interventions: alternative approaches. Bull World Health Organ. 2019;97:118-124.

29. Neumann PJ, Cohen JT, Weinstein MC. Updating cost-effectiveness—the case of the $50,000-per-QALY threshold. N Engl J Med. 2014;371:796-797.

30. Bellnier T, Brown GW, Ortega TR. Preliminary evaluation of the efficacy, safety, and costs associated with the treatment of chronic pain with medical cannabis. Ment. 2018;8:110-115.

31. Capano A, Weaver B, Burkman E. Evaluation of the effects of CBD hemp extract on opioid use and quality of life indicators in chronic pain patients: a prospective cohort study. Postgrad Med. 2019;132:56-61.

32. Yassin M, Orton A, Robinson D. Effect of adding medical cannabis to analgesic treatment in patients with low back pain related to fibromyalgia: an observational cross-over single centre study. Clin Exp Rheumatol. 2019;37(Suppl 116):13-20.

33. Cherian JJ, Bhave A, Kapadia BH, Starr R, McElroy MJ, Mont MA. Strength and functional improvement using pneumatic brace with extension assist for end-stage knee osteoarthritis: a prospective, randomized trial. J Arthroplasty. 2015;30:747-753.

34. Draganeich L, Reider B, Schlich S, Neupert A, Hilgmann M, Tournais A, Regnier JY. Impact of chondroitin sulphate on health utility in patients with knee osteoarthritis: towards economic analysis. J Med Econ. 2009;12:356-360.

35. Petersen W, Ellermann A, Henning J, et al. Non-operative treatment of unicompartmental knee osteoarthritis: a prospective, randomised, double-blind, placebo-controlled pilot study. Arthritis Rheumatol. 2019;27:85-91.

36. Yamamoto GJ, Ocampos GP, Luzo MCM, da Silva CAC, de Farias FES, de Andrade JOS, et al. The effect of glucosamine supplementation provides relief from knee osteoarthritis symptoms: a randomized controlled pilot study. Acta Ortop Bras. 2019;27:85-91.

37. Braham R, Dawson B, Goodman C. The effect of glucosamine supplementation on clinical outcomes between glucosamine sulfate-potassium chloride and glucosamine sulfate for painful knee osteoarthritis. ScientificWorldJournal. 2012;2012:902676.

38. Chopra A, Saluja M, Tillu G, et al. Ayurvedic medicine offers a good alternative to glucosamine and celecoxib in the treatment of symptomatic knee osteoarthritis: a randomized, double-blind, placebo-controlled pilot study. J Rheumatol. 2013;40:71-77.

39. Frestedt JL, Walsh M, Kneer W, Lehnhardt K, Seidel EJ, Minz D, et al. Efficacy of peanut paste 5% versus celecoxib for osteoarthritis-related knee pain: post hoc analysis of a 12 week, prospective, randomized, active-controlled, open-label, parallel-group trial in adults. Clin Rheumatol. 2008;30:2366-2377.

40. Giordano N, Fioravanti A, Papakostas P, Montella A, Giorgi G, Nuti R. The effect of oral celecoxib on health-related quality of life in patients with knee osteoarthritis: a randomized, double-blind, placebo-controlled, parallel-group trial. Br J Sports Med. 2013;47:79-90.

41. Kivitz A, Fairfax M, Sheldon EA, et al. Comparison of the effectiveness and tolerability of liposomal 5% versus celecoxib for osteoarthritis-related knee pain: a randomised controlled 13-week study versus placebo and celecoxib. Clin Rheumatol. 2016;35:110-117.

42. Lehmkan R, Brzosko M, Kopsa P, et al. Efficacy and tolerability of lumiracoxib and celecoxib versus diclofenac in the management of osteoarthritis of the knee. Open Rheumatol J. 2011;5:195-202.

43. Rother M, Lavins BJ, Kneer W, Lehnhardt K, Seidel EJ, Mazgareanu S. Efficacy and safety of epicutaneous ketoprofen in Transfersome (IDEA-033) versus ketoprofen-free vehicle (TDT 064) and oral celecoxib for knee pain associated with osteoarthritis. Rheumatology (Oxford). 2013;52:1303-1312.

44. Selvan T, Rajah K, Nairn MS, Mathew EM. A clinical study on glucosamine sulfate versus combination of glucosamine sulfate and NSAIDs in mild to moderate knee osteoarthritis. ScientificWorldJournal. 2012;2012:902676.

45. Trc T, Bohova J. Efficacy and tolerance of enzymatic hydrolysed collagen (EHC) vs. glucosamine sulphate (GS) in the treatment of knee osteoarthritis (KOA). J Orthop Surg. 2011;19:141-148.

46. Wangroongsub Y, Tanavalee A, Wilairatana V, Ngarmukos S. Comparable clinical outcomes between glucosamine sulfate-potassium chloride and glucosamine sulfate sodium chloride in patients with mild and moderate knee osteoarthritis: a randomized, double-blind study. J Med Assoc Thai. 2003;96:805-811.

47. Beuyrer O, Scholissien S, Neupert A, Hilgmann M, Tournais A, Regnier JY. Impact of chondroitin sulphate on health utility in patients with knee osteoarthritis: towards economic analysis. J Med Econ. 2009;12:356-360.

48. Fardellone P, Zaim M, Saurel C, Mahu E. Comparative efficacy and safety study of two chondroitin sulphate preparations from different origin (avian and bovine) in symptomatic knee osteoarthritis of the knee. Open Rheumatol J. 2013;7:1-12.

49. Rondanelli M, Braschi V, Gasparri C, et al. Effectiveness of non-animal chondroitin sulphate supplementation in the treatment of moderate knee osteoarthritis in a group of overweight subjects; a randomized, double-blind, placebo-controlled study. Arthritis Rheum. 2003;48:3102-3111.

50. Conaghan PG, Dickson J, Bolten W, Cerv C, Rother M. A multicentre, randomized, placebo- and active-controlled trial comparing the efficacy and safety of topical ketoprofen in a transdermal gel (IDEA-033) with ketoprofen-free vehicle (TDT 064) and oral celecoxib for knee pain associated with osteoarthritis. Rheumatology (Oxford). 2013;52:1303-1312.

51. Conaghan PG, Dickson J, Bolten W, Cerv C, Rother M. A multicentre, randomized, double-blind, non-inferiority trial versus celecoxib. Ann Rheum Dis. 2016;75:37-44.

52. Kivitz A, Fairfax M, Seidel EJ, et al. Comparison of the effectiveness and tolerability of liposomal 5% versus celecoxib for osteoarthritis-related knee pain: post hoc analysis of a 12 week, prospective, randomized, active-controlled, open-label, parallel-group trial in adults. Clin Rheumatol. 2008;30:2366-2377.

53. Lehmann R, Brazoko M, Kopsa P, et al. Efficacy and tolerability of lumiracoxib 100 mg once daily in knee osteoarthritis: a 12-week, randomized, double-blind study versus placebo and celecoxib. Curr Med Res Opin. 2005;21:517-526.

54. Hochberg MC, Marriot-Pellerier J, Monfort J, et al. Combined chondroitin sulphate and glucosamine for painful knee osteoarthritis: a multicentre, randomised, double-blind, placebo-controlled trial. Curr Med Res Opin. 2001;19:313-324.

55. Lehmann R, Brazoko M, Kopsa P, et al. Efficacy and tolerability of lumiracoxib versus combination of glucosamine sulfate and NSAIDs in mild to moderate knee osteoarthritis. Arthritis Rheum. 2006;54:3441-3445.

56. Lehmann R, Brazoko M, Kopsa P, et al. Efficacy and tolerability of lumiracoxib versus combination of glucosamine sulfate and NSAIDs in mild to moderate knee osteoarthritis. Arthritis Rheum. 2006;54:3441-3445.
66. Case JP, Balouzas AJ, Block JA. Lack of efficacy of acetaminophen in treating symptomatic knee osteoarthritis: a randomized, double-blind, placebo-controlled comparison trial with diclofenac sodium. *Arch Intern Med.* 2003;163:169-178.

67. Kasemsuk T, Saengpetch N, Sibmooh N, Unchern S. Improved WOMAC score following 16-week treatment with bromelain for knee osteoarthritis. *Clin Rheum.* 2016;35:2531-2540.

68. Pareek A, Chandurkar N. Comparison of gastrointestinal safety and tolerability of alocfenac: a multicenter, randomized, double-blind study in patients with knee osteoarthritis. *Curr Med Res Opin.* 2013;29:849-859.

69. Petrella RJ, DiSilvestro MD, Hildebrand C. Effects of hyaluronate sodium on pain and physical functioning in osteoarthritis of the knee: a randomized, double-blind, placebo-controlled clinical trial. *Arch Intern Med.* 2002;162:292-298.

70. Pinosorsk P, Kanokkangsudal P, Itharat A. The clinical efficacy and safety of the sahastara remedy versus diclofenac in the treatment of osteoarthritis of the knee: a double-blind, randomized, and controlled trial. *Evid Based Complement Alternat Med.* 2015;2015:103376.

71. Sangedee T, Teekachanathae S, Sanapanachon K, et al. Electrocupuncture versus diclofenac in symptomatic treatment of osteoarthritis of the knee: a randomized controlled trial. *BMC Complement Altern Med.* 2002;2:3.

72. Simons LS, Grieser LM, Naeve Z, Bookman AAM, Shaninhouse ZJ. Efficacy and safety of topical diclofenac containing dimethyl sulfoxide (DMSO) compared with those of topical placebo, DMSO vehicle and oral diclofenac for knee osteoarthritis. *Pain.* 2009;143:238-245.

73. Tugwell PS, Wells GA, Shainhouse JZ. Equivalence study of a topical diclofenac solution (pennsaid) compared with oral diclofenac in symptomatic treatment of osteoarthritis of the knee: a randomized controlled trial. *J Rheumatol.* 2004;31:2002-2012.

74. Kvitva A, Eisen G, Zhao WW, Bevitt T, Recker DP. Randomized placebo-controlled trial comparing efficacy and safety of valdecoxib with naproxen in patients with osteoarthritis. *J Pain Pract.* 2002;51:530-537.

75. Kuptniratsakul V, Pinthong T, Bunjob M, Thanakhumtorn S, Chinswang-watanauk P, Thamlikitkul V. Efficacy and safety of Derris scandens Benth. extracts in patients with knee osteoarthritis. *J Altern Complement Med.* 2011;17:147-153.

76. Loo RM, Kholov A, Koppenkin S, et al. Efficacy and safety of flavocoxid, a novel therapeutic, compared with naproxen: a randomized multicenter controlled trial in subjects with osteoarthritis of the knee. *Adv Ther.* 2010;27:731-742.

77. Lohmander LS, McKeith D, Svensson O, et al. A randomised, placebo controlled clinical trial. *Cartilage.* 2006;27:1590-1598.

78. Schnitzer TJ, Kvitva AJ, Lipata RS, Sanders N, Hee A. Comparison of the COX-inhibiting nitric oxide donor AZD3582 and rofecoxib in treating the signs and symptoms of Osteoarthritis of the knee. *Arthritis Rheum.* 2005;53:827-837.

79. Svensson O, Malmenas M, Fajutrao L, Roos EM, Lohmander LS. Greater reduction of knee than hip pain in osteoarthritis treated with naproxen, as evaluated by WOMAC and SF-36. *Arch Rheum Dis.* 2006;65:781-784.

80. Watt FE, Blauwert MB, Falkhoury A, Jacobs H, Smullers R, Lane NE. Tropo-myosin-related kinase A (TrkA) inhibition for the treatment of painful knee osteoarthritis: results from a randomized controlled phase 2a trial. *Osteoarthritis Cartilage.* 2019;27:1590-1598.

81. Kuptniratsakul V, Dajpratham P, Taechaarpornkul W, et al. Efficacy and safety of Derris scandens Benth. extracts compared with ibuprofen in patients with osteoarthritis. *Arthritis Rheum.* 2002;45:449-456.

82. Schnitzer TJ, Kvitva AJ, Lipata RS, Sanders N, Hee A. Comparison of the COX-inhibiting nitric oxide donor AZD3582 and rofecoxib in treating the signs and symptoms of osteoarthritis of the knee. *Arthritis Rheum.* 2005;53:827-837.

83. Simons O, Malmenas M, Fajutrao L, Roos EM, Lohmander LS. Greater reduction of knee than hip pain in osteoarthritis treated with naproxen, as evaluated by WOMAC and SF-36. *Arch Rheum Dis.* 2006;65:781-784.

84. Underwood M, Ashby D, Carnes D, et al. Topical or oral ibuprofen for chronic osteoarthritis of the knee: a multicenter study. *Health Technol Assess.* 2009;13:1-148.

85. Babul N, Noveck R, Chipman H, Roth SH, Gana T, Albert K. Efficacy and safety of aceclofenac with diclofenac: a multicenter, randomized, double-blind study in osteoarthritis. *Arch Intern Med.* 2009;169:234-242.

86. Mukhopadhyay K, Ghosh P, Ghorai P, Hazra A, Das AK. Oxaceprol versus tramadol for treatment of knee osteoarthritis: a randomized controlled study. *Expert Rev Pharmacoecon Outcomes Res.* 2010;10:35-44.

87. Slof J, Gras A. Sativex® in multiple sclerosis spasticity: a cost-effectiveness model. *Expert Rev Pharmacoecon Outcomes Res.* 2012;12:439-441.

88. Katz JN, Smith SR, Collins JE, et al. Cost-effectiveness of nonsteroidal anti-inflammatory drugs and opioids in the treatment of knee osteoarthritis in older patients with multiple comorbidities. *Osteoarthritis Cartilage.* 2016;24:409-418.

89. Lu L, Pearce H, Roome C, Shearer J, Lang IA, Stein K. Cost effectiveness of oromucosal cannabis-based medicine (Sativex®) for spasticity in multiple sclerosis. *PharmaEconomics.* 2012;30:1157-1171.

90. Slof J, Ruiz I, Vila C. Cost-effectiveness of Sativex in multiple sclerosis spasticity: new data and application to Italy. *Expert Rev Pharmacoecon Outcomes Res.* 2015;15:379-391.

91. Black C, Clar C, Henderson R, et al. The clinical effectiveness of glucosamine, NSAIDs, and selective COX-2 inhibitors in the treatment of symptomatic knee osteoarthritis. *Value Health.* 2003;6:144-157.

92. Katz JN, Smith SR, Collins JE, et al. Cost-effectiveness of nonsteroidal anti-inflammatory drugs and opioids in the treatment of knee osteoarthritis in older patients with multiple comorbidities. *Osteoarthritis Cartilage.* 2016;24:409-418.

93. Lohmander LS, McKeith D, Svensson O, et al. A randomised, placebo controlled comparative trial of the gastrointestinal safety and efficacy of AZD3582 and rofecoxib in treating the signs and symptoms of Osteoarthritis of the knee. *Arthritis Rheum.* 2005;53:827-837.

94. Sherman AL, Ojeda-Correal G, Mena J. Use of glucosamine and chondroitin in the management of human osteoarthritis? *Acta Med Indones.* 2011;42:35-44.

95. Anderson JW, Nicolosi RJ, Borzelleca JF. Glucosamine effects in humans: a systematic review and economic evaluation. *Arthritis Care Res.* 2004;55:47-61.

96. Tyree GA, Sarkar T, Bellows BK, et al. A cost-effectiveness model for adjunctive smoked cannabis in the treatment of chronic neuropathic pain. *Can J Anesth.* 2019;66:62-72.

97. Korpina S, Zuyere O, Neuprez A, et al. Glucosamine sulphate in the treatment of knee osteoarthritis: cost-effectiveness comparison with paracetamol. *Int J Clin Pract.* 2010;64:756-762.

98. Hermans J, Koopmanschap MA, Bierma-Zeinstra SM, et al. Productivity costs and medical costs among working patients with knee osteoarthritis. *Arthritis Care Res.* 2012;64:853-861.

99. Lee PY, Winfield TG, Harris SR, Storey E, Chandratry A. Unloading knee brace is a cost-effective method to bridge and delay surgery in unicompartamental knee arthritis. *BMJ Open Sport Exerc Med.* 2016;2:e000195.

100. Woods B, Manca A, Weatherly H, et al. Cost-effectiveness of adjunct non-pharmacological interventions for osteoarthritis of the knee. *Health Technol Assess.* 2009;13:i-iv, i-xvi.

101. Sherman AL, Ojeda-Correal G, Mena J. Use of glucosamine and chondroitin in the management of human osteoarthritis? *Acta Med Indones.* 2011;42:35-44.

102. Anderson JW, Nicolosi RJ, Borzelleca JF. Glucosamine effects in humans: a systematic review and economic evaluation. *Arthritis Care Res.* 2004;55:47-61.

103. Sherman AL, Ojeda-Correal G, Mena J. Use of glucosamine and chondroitin in the management of human osteoarthritis? *Acta Med Indones.* 2011;42:35-44.