Medical Marijuana: Clearing Away the Smoke

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Abstract: Recent advances in understanding of the mode of action of tetrahydrocannabinol and related cannabinoid ingredients of marijuana, plus the accumulating anecdotal reports on potential medical benefits have spurred increasing research into possible medicinal uses of cannabis. Recent clinical trials with smoked and vaporized marijuana, as well as other botanical extracts indicate the likelihood that the cannabinoids can be useful in the management of neuropathic pain, spasticity due to multiple sclerosis, and possibly other indications. As with all medications, benefits and risks need to be weighed in recommending cannabis to patients. We present an algorithm that may be useful to physicians in determining whether cannabis might be recommended as a treatment in jurisdictions where such use is permitted.

Keywords: Cannabis, chronic pain, pain.

INTRODUCTION

In this article we review evidence that cannabis may be useful as medicine. We discuss potential indications for its use and provide an algorithm to guide medicinal cannabis recommendations.

The reasons for a revival of interest in medicinal cannabis are multiple, and beyond the scope of this review, but include increasing anecdotal and clinical study reports of potential benefit, advances in understanding of the endocannabinoid signaling system upon which cannabis acts, as well as growing public acceptance that cannabis should be available as a medicine if a physician recommends it.

BRIEF REVIEW OF PAST CLINICAL STUDIES ON MEDICINAL CANNABINOIDS

As recently as a decade ago a review of the world literature on the status of the efficacy and safety of cannabinoids for pain and spasticity revealed that only nine randomized studies of acceptable quality had been conducted [1]. All of these were single dose studies comparing oral synthetic THC (or cannabinoid analogs or congeners) to codeine or placebo. Two were “N of 1” randomized trials and two were of very small samples of acute post-operative pain. The remaining trials primarily addressed chronic cancer-related pain. Taken as a group it appeared that oral cannabinoids (e.g., THC 10mg) outperformed placebo and were analgesically equivalent to codeine 60mg; higher doses (THC 20mg) were comparable to codeine 120mg, but had a much higher incidence of adverse effects, particularly sedation [2]. Authoritative reviews judged cannabinoids as being unlikely to have a role in acute pain management, but suggested there was enough evidence for efficacy in chronic neuropathic pain and muscle spasticity to warrant further research [1].

RECENT STUDIES ON MEDICINAL CANNABIS

In the past decade, the scope and rigor of research has increased dramatically. This research has employed cannabis, cannabis-based extracts, and synthetic cannabinoids delivered by smoking, vaporization, oral, and sublingual or mucosal routes.

Studies on Smoked Cannabis

Smoking cannabis provides rapid and efficient delivery of THC to brain. THC can be detected immediately in plasma after the first puff of a cigarette; peak concentrations occur within 10 minutes, then decrease to approximately 60% of peak by 15 minutes and 20% of peak by 30 minutes, but there can be wide inter-individual variation in concentrations achieved [3]. Rapid onset and predictable decay means that self-titration of dosing is attainable.

Chronic Pain

A series of randomized clinical trials at the University of California Center for Medicinal Cannabis Research (CMCR) investigated the short-term efficacy of smoked cannabis for neuropathic pain. Sponsored by the State of California Medical Marijuana Research Act of 1999, and conducted under the auspices of the Department of Health and Human Services, the National Institute on Drug Abuse, and the Food and Drug Administration, this research allocated participants to smoke cannabis cigarettes containing from 1% to 8% THC by weight (4 to 32 mg THC) or to placebo cannabis cigarettes from which THC had been extracted. The total daily dose of THC ranged from 4 mg to 128 mg. Two trials enrolled patients with painful HIV peripheral neuropathy [4, 5]; one consisted of mixed neuropathic pain due to peripheral or central dysfunction of the nervous system (i.e., com-
Chronic Pain

Most research using oral preparations has targeted neuropathic pain and spasticity associated with multiple sclerosis (MS). These randomized trials suggest that dronabinol (up to 25 mg daily) significantly reduces pain compared to placebo (50% “improved” on dronabinol compared to 30% on placebo, p < 0.05) [10], with a number-needed-to-treat for 50% pain reduction of 3.5, which is in the range of efficacy observed for standard non-opioids [11]. Effects on spasticity are mixed: there may be no observable change in examiner-rated muscle tone, but patients report significant relief [10].

There is less research with nabnilone, although one three-week randomized crossover trial reported that nabnilone 2mg provided modest analgesia, comparable to dihydrocodeine 240mg daily in neuropathic pain [12].

Nausea-Emesis and Appetite Stimulation

Although serotonin receptor (5 HT3) antagonists (e.g., ondansetron, Zofran®) and Substance P/neurokinin-1 (NK-1) receptor antagonists (e.g., aprepitant, Emend®) are the mainstay for treatment, dronabinol and nabnilone are also FDA-approved for control of acute and delayed nausea and emesis due to cancer chemotherapy. Meta-analyses indicate these cannabinoids are equivalent to or more effective than metoclopramide and neuroleptics, but their side effect profile is less favorable in terms of sedation, dizziness, dysphoria, hypotension, and anxiety [13, 14]. There are no head-to-head comparisons of cannabinoids with serotonin 5 HT3 receptor or Substance P/NK-1 receptor antagonists.

Anorexia, early satiety, weight loss and cachexia are prevalent in late stage cancer and advanced HIV disease, but there are few effective treatments. Trials in AIDS patients with clinically significant weight loss indicated that dronabinol 5mg daily significantly outperformed placebo in terms of short term appetite enhancement (38% vs. 8% at 6 weeks), and that these effects persisted for up to 12 months [15, 16], but were not accompanied by significant differences in weight gain, perhaps because of disease-associated energy wasting. The major practical limitations are the accompanying psychoactive side effects, and the problems of oral administration (eg, delayed onset of action, variable absorption, extended duration of effects).

Studies on Cannabis-based Extracts

Outside the US, extracts of whole plant are licensed and available in capsules (Cannador®, with the main constituents being THC and the non-psychoactive plant cannabinoid, cannabidiol, in a ratio of 2:1. Rectal suppositories are also used to deliver THC hemisuccinate. Several small to medium-sized, randomized, controlled trials in MS suggest improvements in pain and perceived spasticity at daily doses of THC ranging from 7.5mg to 27.5mg [10, 17, 18]. In some trials [19] but not others [10, 20] observer-assessed spasticity also improved.

Studies with Alternative Delivery Systems

The hazards of smoking and the pharmacokinetic limitations of ingestion of cannabinoids has led to a search for alternative systems of administration. One alternative is devices which vaporize cannabis leaves by heating the plant product to below the temperature of combustion (175-225 degrees C), permitting inhalation of volatilized gases minus hazardous pyroles produced by burning. Preliminary work using plant material with a range of THC content (e.g., 1-7% THC) suggests that there is rapid onset, with peak concentrations and six-hour area under the plasma concentration curves comparable to those achieved by smoking [21]. Vaporization is not a perfect solution since carbon monoxide is formed, but levels are significantly lower than with smoking [21]. Clinical trials are currently in progress at the CMCR assessing the efficacy of vaporized cannabis as an analgesic in chronic neuropathic pain.

Sublingual delivery systems of whole cannabis plant extract, which employ metered spray devices to deliver measured doses of THC (2.7mg) and cannabidiol (2.5mg), are undergoing Phase IIb/III trials in the US, and are licensed elsewhere for cancer pain and multiple sclerosis-associated neuropathic pain and spasticity (nabiximols, Sativex®). The apparent advantages of such systems are known cannabinoid
concentrations, predetermined dosing aliquots, and time-out systems which may help prevent overuse. Some placebo-controlled trials suggest significant analgesia in neuropathic pain due to multiple sclerosis [22] and mixed neuropathy (e.g., post-herpetic, traumatic, vascular neuropathies, [23] but others do not [20]. Other controlled trials suggest efficacy for cancer-related pain inadequately responsive to opioid analgesia [24]. Responders participating in the open label extension phases of controlled trials appear to maintain analgesia on one-year follow-up [23].

In regard to spasticity in multiple sclerosis, a recent meta-analysis combining three trials with nabiximols in over 600 patients noted that mean intensity of patient rated spasticity was significantly reduced compared to placebo [20, 25, 26], and that the proportion of “responders” (30% reduction) was also significantly greater, with about 37% on the cannabinoid compared to 26% on placebo experiencing relief. Those reporting relief of spasticity seemed to maintain their gains over one year follow-up [27]. As with other studies noted above, observer-rated spasticity is often not reduced [20, 25, 28]; however, a recent CMCR study did find a significant reduction in observed spasticity among those administered active smoked marijuana vs. placebo marijuana [58].

PRESERVATION OF MASKING IN CLINICAL TRIALS

Because of the acute psychoactive effects of the experimental agent there is understandable concern that blinding cannot be preserved in placebo-controlled clinical trials of cannabinoids, particularly with cross-over designs. Few studies assess masking, but two cross-over trials tested maintenance of the blind by asking participants to “guess” assignment at different points of the study. Results suggest that participants, whether they are naïve or experienced cannabis users, are in the first week of a trial no more likely than by chance to guess assignment [5, 9]. With continued exposures rates of correct guesses exceed 75%, but exceed chance only in a high potency arm (9%) [9]. In another study correct guessing was related to two factors: whether the subject received placebo or cannabis first; and when during the study the participant guessed assignment [5]. Among individuals randomized to receive placebo first, guessing was no better than chance through the end of the first treatment week, whereas the majority of those randomized to receive cannabis first correctly guessed their treatment assignment at all time points. Furthermore, by the conclusion of the study, when all subjects had been given the opportunity to compare the cannabis placebo and treatments, even those randomized to receive placebo first correctly guessed their treatment assignment [5]. This raised the possibility that some of the pain reduction was placebo driven. Secondary analyses to assess whether correct treatment guessing influenced treatment responses showed that in the placebo group during the first treatment week, when guessing was no better than chance, cannabis still provided pain relief superior to that of placebo. This finding suggests that although placebo effects were present, treatment effects were independent [5].

RISKS AND MANAGEMENT OF MEDICINAL USE OF CANNABINOIDS

Acutely and over the longer term cannabis may have unwanted systemic and psychoactive adverse effects that must be taken into consideration in chronic pain populations, who have high rates of co-occurring medical illness (e.g., cardiovascular disease) and co-morbid psychiatric and substance use disorders. In general these effects are dose-related, are of mild to moderate severity, appear to decline over time, and are reported less frequently in experienced than in naïve users. Reviews suggest the most frequent side effects are dizziness or light-headedness (30%-60%), dry mouth (10%-25%), fatigue (5%-40%), muscle weakness (10%-25%), myalgia (25%), and palpitations (20%) [17]. Cough and throat irritation are reported in trials of smoked cannabis [9]. Tachycardia and postural hypotension are infrequent but caution is warranted in patients with cardiovascular disease, and possibly younger adults who intend to embark on very vigorous physical activity. At higher doses, sedation and ataxia with loss of balance are frequent. Participants in some but not all studies report euphoria: the relative absence of psychoactive effect has been attributed to the observation that plasma concentrations obtained in clinical trials are often <25% of those achieved by “recreational” users (eg, 25ng/ml vs >100ng/ml) [9]. After repeated smoked or oral marijuana doses, tolerance is rapidly acquired (in two to 12 days) to many of its adverse effects, e.g., cardiovascular, autonomic, and many subjective and cognitive effects [29]. After exposure is stopped, tolerance is lost with similar rapidity.

There is little systematic data on timeline to tolerance of either adverse or therapeutic effects, like analgesia. Concerns have long been voiced that rapid tolerance to adverse effects might portend tolerance to beneficial effects [29]. Data from studies using oral sprays of cannabinoids or dronabinol in multiple sclerosis report that individuals can reduce the incidence and severity of adverse effects by downward self-titration without loss of analgesia [17]. Other studies in this population note that overall the incidence and severity of adverse effects diminishes over time without evidence of tolerance to analgesic effects [20, 22]. Yet it is rare that clinical trials of cannabinoids extend follow-up beyond 12 weeks, leaving questions on maintenance of gains or need for dose escalation unanswered [10, 26]. One study with 12-month follow-up concluded there may be sustained analgesia for pain associated with multiple sclerosis, where about 30% of cannabinoid-treated participants report continued “improvement” at 12 months compared to about 15% on placebo [30] on doses conservatively limited to a maximum of 25mg THC daily. This suggests that pain relief may be sustained without dose increases. But the study design was not intended to determine the proportion of patients who experienced diminution of effect, or whether dose escalation, even within the set boundary, was needed for maintenance of efficacy.

There are risks to be considered in assessing the potential of cannabinoid therapeutics. Cannabis, like other analgesics, can be associated with dependence and a withdrawal syndrome, occurring in a dose-dependent fashion [29]. Under controlled conditions in healthy, experienced users of marijuana, withdrawal from a “low” daily dose (ie, oral THC 10 mg every 3-4 hours for 5-21 days) commences within 12 hours, is diminished by 24 hours, and is complete in 48 to 72 hours [29]. Other short term experiments with oral THC (20 to 30mg four times daily) and smoked cannabis (1% and 3% THC cigarettes four times daily) reveal an abstinence syn-
drome characterized by anxiety, irritability-restlessness, insomnia, stomach pain and decreased appetite [31, 32], with mood effects more prominent at the higher dosages. In research specifically designed to establish the time line of abstinence among regular heavy users (4 cigarettes daily), symptoms peak at 2 to 3 days, and persist for up to 2 weeks, although sleep disturbance may continue for up to 6 weeks [33]. In light of abstinence effects, standard practice in clinical trials administering a maximum of 25 mg THC daily is to use a tapering scheme to conclude therapy, with a 20% per day dose reduction [30]. Patients discontinuing higher dose cannabinoids for analgesia might warrant a longer tapering regimen, but this has not been studied.

Fatal overdose with cannabis alone has not been reported. In terms of acute drug interactions, additive effects of cannabinoids, anticholinergics, and CNS depressants should expected (e.g., increased sedation, dizziness, dry mouth, confusion). Cannabinoids are metabolized by several enzyme systems, including Cytochrome P450 (CYP 2C9, CYP 3A4) and can induce or inhibit CYP 3A4, but there is little evidence of important drug-drug interactions based on CYP 450 systems. Smoking itself (e.g., cannabis or tobacco) induces CYP 1A2, and may increase clearance of some antipsychotics (e.g., olanzapine, clozapine) and antidepressants (e.g., some tricyclics, mirtazapine) [34, 35]. Overall then, the acute medical risks of THC as used in clinical trials are rather low.

There can be adverse psychiatric side effects. THC intoxication and euphoria can be disturbing, particularly to elderly patients. Anxiety and panic attacks occur, as do frank psychotic reactions (principally paranoia), as well as so-called “paradoxical” effects of dysphoria, dejection, and depressed mood [36, 37]. Although unlikely to be a factor in the application of cannabinoids for pain, there is concern that early adolescent use of cannabis may heighten later risk of psychosis [36, 38], and evidence that genetic variation (single nucleotide polymorphisms) heightens vulnerability [39].

Acute cannabinoid intoxication adversely impacts processing speed, attention, learning and recall, perception of time and velocity, reaction time and psychomotor abilities in a dose-dependent fashion [40]. Formal neuropsychological testing in clinical trials reveals mild impairment at usual analgesic doses [6, 23]. While cannabis can acutely impair skills required to drive motor vehicles in a dose-related fashion, epidemiological data are inconclusive with regard to the association of traffic accidents and use of cannabis [41]. There is speculation that cannabis use is associated with increased awareness of impairment (e.g., altered perception of time and speed), which results in compensatory behavioral strategies. What is clearer from experimental and epidemiologic data is that driving under the influence of both alcohol and cannabis in combination confers greater risk of accidents than the risk of either drug alone [41].

The longer-term health risks of cannabis are unclear, and the evidence is based on non-medical use [42]. Long-term use of inhaled cannabis may be associated with dependence and increased respiratory symptoms; but some epidemiological studies have not found more lung disease in long-term users, once the effects of tobacco are accounted [43]. Long-term use of inhaled cannabis has not been associated with increased risk of lung or gastrointestinal cancers [44], although a meta-analysis found evidence of premalignant changes in the respiratory tract [45]. There is some evidence that among individuals with pre-existing cardiac disease, cannabis users have an increased risk of myocardial infarction in the hour after smoking cannabis compared to non-users [46]. A recent meta-analysis showed no major residual effects on neurocognitive functioning in long term daily-users of cannabis [47]. THC rapidly crosses the placenta and accumulates in breast milk of nursing mothers [3], but there is no systematic evidence of direct or behavioral teratogenicity.

In reviewing the possible acute and long term adverse effects of cannabinoids as therapeutic agents one needs also to be mindful that other agents that are used for treatment of pain or spasticity also have adverse effects. Opioids produce sedation, nausea, constipation and dependence, withdrawal from which results in serious abstinence syndrome with much more severe effects – e.g. severe autonomic, gastrointestinal, and psychotic – than the rather mild cannabis withdrawal phenomena. Tricyclic antidepressants and antiepileptic drugs commonly prescribed for chronic pain have psychotropic (e.g. sedation), anticholinergic (e.g. constipation, dizziness, palpitations, visual disturbance, urinary retention), and neuromuscular effects. Drugs for spasticity produce sedation (e.g. baclofen), hypotension (e.g. tizanidine) and serious interactions with antibiotics (e.g. tizanidine and ciprofloxacin). Benzodiazepines that are sometimes prescribed for spasticity can produce sedation, psychomotor incoordination, memory lapses, and paradoxical reactions, as well as dependence and withdrawal syndromes. Opioids and sedative-hypnotics are also drugs of abuse, and their ability to induce physiological dependence and serious withdrawal states exceed those of cannabis. Therefore, judgements on relative benefits and risks of cannabinoids as medicines need to be viewed within the broader context of risk-benefit of other agents as well [48].

**PATIENT SELECTION FOR CANNABINOID THERAPY**

Oral THC (eg, dronabinol) is FDA-approved as a second line agent for chemotherapy-associated nausea and emesis, and appetite stimulation. Dronabinol (and nabilone) have some evidence of efficacy for chronic neuropathic pain; whole plant cannabis extracts delivered by capsule or oral mucosal spray has been approved in Europe for analgesia in neuropathic pain and control of painful muscle spasticity. Patient selection for these agents would seem to be rather straightforward, and focus on therapeutic response to conventional treatments, consideration of possible psychotropic (eg. sedation effects if combined with alcohol) and cardiovascular effects, risk of dependence and an abstinence syndrome, and acknowledgement that there is narrow empirical basis for efficacy compared to standard treatments. Patients should be educated about expected adverse effects. The pharmacokinetics of orally administered cannabinoids would seem to decrease likelihood of diversion or abuse.

On the other hand, prescription of inhaled cannabis for medical purposes is legal in some US jurisdictions, and neuropsychologists, who are likely to be asked about the advisability of prescribing or recommending “medical mari-
**Table 1. Medical Board/Office of the Attorney General of California Guidelines for Medical Marijuana**

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<th>Physicians Recommending Medical Marijuana Need to:</th>
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<tr>
<td>1. Take a history and conduct a good faith examination of the patient;</td>
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<td>2. Develop a treatment plan with objectives;</td>
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<td>3. Provide informed consent, including discussion of side effects;</td>
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<td>4. Periodically review the treatment’s efficacy;</td>
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<td>5. Obtain consultations, as necessary; and</td>
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<tr>
<td>6. Keep proper records supporting the decision to recommend the use of medical marijuana.</td>
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http://www.mbc.ca.gov/board/media/releases_2004_05-13_marijuana.html

establish a similar routine of urine toxicology testing, compliance checklists, and co-enrollment in a formal substance abuse treatment facility, just as is recommended for a trial of opioid analgesics [52]. Finally, chronic pain may be associated with major depression, which complicates treatment, and which must be diagnosed and independently treated for successful pain management. All of these factors being considered, if the decision is made to proceed, the clinician must formulate and document a treatment plan and the patient’s agreement to abide by whatever guidelines are established. The clinical “trial” would entail establishing a therapeutic “dose,” appropriately monitoring for adverse effects and misuse, and assessing outcome in terms of pain, mood, and function. Based on the literature of efficacy in neuropathic pain, there could be evidence of an effect within a minimum of two weeks. Response rates have been noted to increase, however, between two and four weeks in previous neuropathic pain registration trials for gabapentin and duloxetine [57]; one might select a longer duration in difficult-to-treat cases. Considering all these factors, one would then decide with the patient whether continued treatment is warranted. A possible algorithm to guide physician decision-making is presented in Fig. (1).

CONCLUSION

Evidence is accumulating that cannabinoids may be useful medicine for certain indications. Control of nausea and vomiting and the promotion of weight gain in chronic inanition are already licensed uses of oral THC (dronabinol cap-

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**Fig. (1).** A decision tree approach for physicians who may be considering recommending medicinal cannabis to a patient. This decision tree suggests some key points that a physician may need to consider in making his/her determination. In this case, a patient is assumed to present with persistent neuropathic pain. Initially, a determination needs to be made that the patient’s signs and symptoms are indeed consistent with this diagnosis. Assuming a patient does not respond favorably to more standard treatments (e.g., antidepressants, anticonvulsants, etc), or cannot tolerate those, and the patient is willing to consider medicinal cannabis, the physician needs to determine risk versus benefit. Among these considerations is whether there is a history of substance abuse or serious psychiatric disorder that might be exacerbated by medicinal cannabis. Even if such risks exists, this does not necessarily preclude the use of medicinal cannabis; rather coordination with appropriate substance abuse and psychiatric resources is necessary, and based on such consultation a risk benefit ratio can be determined. In patients in whom the ratio appears favorable, the physician needs to discuss alternative modes of cannabis administration which may include oral, smoked, or vaporized systems. Once risks and benefits are evaluated and discussed with the patient, cannabis treatment may commence as with other psychotropic medications, with attention being paid to side effects as well as efficacy. In addition, there needs to be attentiveness to potential for misuse and diversion, which can then trigger a decision to discontinue.

**Key**

1. Daily or almost daily pain with typical neuropathic characteristics for at least 3 months; affects life quality.
2. Standard Rx = e.g., antidepressants, anticonvulsants; opioids; nonsteroidal anti-inflammatory drugs.
3. For example, at least 30% reduction in pain intensity.
4. Consider past experience, possible past history of side effects; willingness to smoke.
5. Determine history of substance abuse. If yes, or at “high risk” aberrant for drug behavior; proceed with close observation; possibly coordinate with substance abuse treatment program.
6. Efficacy = at least 30% reduction in pain intensity.
ACKNOWLEDGEMENT

Supported by the Center for Medicinal Cannabis Research at the University of California, San Diego.

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