EXPLORING THE EFFICACY AND SAFETY OF CANNABIS IN THE MANAGEMENT OF FIBROMYALGIA

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ABSTRACT

Fibromyalgia is a chronic health condition characterized by chronic pain fatigue, sleep disturbances and many other symptoms affecting a patient’s quality of life. Patients with fibromyalgia often visit rheumatology outpatients with a long list of symptoms and often receive multiple medications. Many have seen multiple specialists and have done a lot of reading about alternative modalities of treatment. The limited effectiveness of conventional therapy coupled with widespread media attention raises the question of cannabis use.

This review examines the literature on cannabis use in fibromyalgia against the context of the international variation in legal frameworks, the available products and the outcomes reported. A detailed review was performed using the EMBASE and PUBMED databases.

It was concluded that despite the interest in the use of cannabinoids in the management of fibromyalgia, there is insufficient evidence to prescribe the currently available licensed medicines or to recommend the complementary health products available for legal purchase.

There is a need for more global clinical randomised trials to accurately determine medicinal cannabis short and long-term efficacy and safety for its acute and chronic use.

Keywords: Fibromyalgia, Medicinal cannabis, Chronic pain, Fatigue, Licensed medicines

INTRODUCTION

Fibromyalgia (FM) is the label applied to a chronic pain syndrome. After around 30 y of intensive study, it remains poorly understood [1]. The condition is associated with chronic pain, fatigue and disturbed sleep patterns that all negatively impact a patient’s quality of life (Qol). The global incidence is around 2.7% of the population are affected [2] overall but within Europe it is reported as around of which 4.7% [3]. It is the third most common rheumatic condition, with only lumbar pain and osteoarthritis being more common [4]. Conventional treatments for pain components of FM include paracetamol, non-steroidal anti-inflammatory drugs and opioids, which are frequently used with antidepressants. However, while the conventional therapy achieves chronic pain reduction of approximately 50%, this is only 10-20% better than placebo [5-8]. These poor outcomes prompted many patients to look for alternatives, such as cannabinoids which is widely discussed in on social media, leading many patients to express interest in trying their use and some to report self-medication with medical, commercial and/or illicitly obtained cannabinoids [9, 10]. Medical cannabis treatment is not currently approved for FM in the UK, but has been extensively investigated for chronic pain management internationally [10-16]. According to the UK national health services (NHS), the traditional management for FM ranges from simple analgics, exercise and CBT or psychotherapy to the use of opioids, antidepressants, antiepileptics and muscle relaxants [17]. The listed symptoms are all or some of: increased sensitivity to pain, extreme tiredness (fatigue), muscle stiffness, difficulty sleeping, problems with mental processes (known as "fibro-fog"), such as problems with memory and concentration, headaches and irritable bowel syndrome (IBS), a digestive condition that causes stomach pain and bloating [17].

Cannabinoids and chronic pain pathophysiology

Cannabis and cannabinoids are reported to modulate pain and reduce inflammation through the endocannabinoid system. Cannabinoid receptors are distributed throughout central and peripheral nerves, with lower numbers found in the brain stem. They also occur in peripheral, non-nervous tissue [18]. The endogenous molecules that bind to these receptors are derived from fatty acid metabolism which is ubiquitous within the body. Endocannabinoids are believed to modulate nociceptive sensory pathways and alter pain perception in chronic pain states [19, 20]. It has further been suggested that a deficiency of endocannabinoids may be part of the pathophysiology in FM [21]. An alternative theory is that stress may be a major component in FM [22]. Modulation of stress and inflammation to improve emotional and cognitive functioning appears an attractive option in FM management [23-25]. Failed or inadequate response to conventional analgesics, and media reporting, leads many patients to consider cannabis to help control their FM symptoms, as it has been reported to be effective in other chronic pain conditions, without serious side effects [26, 27]. The identification and characterization of the endocannabinoid system has triggered an increase in research, but to date, there are few and limited clinical trials of cannabinoids in FM. Small studies with short-term follow-up, and a wide variation in products and approaches used, arising from the wide variations in international regulations and controls, limit their utility as a basis for public health policy [28-31].

Cannabinoid receptors

Cannabinoid receptor-1 (CB1) is a G-protein coupled receptor linked to the psychotropic effects of cannabis. CB1 is widely distributed in the brain, demonstrating high concentrations in the frontal cortex, basal ganglia and cerebellum, on axons and presynaptic terminals [32]. CB2, is similarly a G-protein coupled receptor, more commonly found on peripheral cells, particularly within the immune system. The CB2 variant Q63R has been linked with autoimmune conditions and inflammatory disorders [33]. It has been suggested that the Q63R polymorphism can increase the risk of autoimmune disease and the CB2 variant Q63R has been linked with autoimmune conditions and inflammatory disorders [33].

Many cannabinoids have been identified, and the number present, and their relative proportions, vary as discussed below [34]. Two of the principal components of therapeutic interest are tetrahydrocannabinol (THC) and cannabidiol (CBD) [34, 35]. THC is
the major psychoactive substance reported to affect mood, orientation, appetite and pain perception. CBD is not psychoactive, but is reported to have anxiolytic, antidepressive and anti-inflammatory effects [36]. At the receptor level, for CBL, THC is a partial agonist whilst CBD is a negative allosteric modulator [36]. Due to their varying properties and molecular interactions, the relative proportion of THC to CBD in cannabis products are believed to be the primary determinants of the effects and adverse effects observed; however, many other cannabinoids also possess activity giving rise to unique patterns seen with particular products and delivery methods [35]. This is also important in terms of ongoing consistency and quality control in products not subject to the legal controls required for licensed medicinal products [38].

Trials of medical cannabis in fibromyalgia

A Cochrane review in 2016 did not support the use of cannabinoids in FM [38]. Only two RCTs were included. Both studies focussed on nabilone, and showed promising results, with one trial demonstrating that nabilone effectively reduced anxiety, pain and Fibromyalgia Impact Questionnaire (FIQ) scores [30], and that it was superior to amitriptyline in resolving sleep disturbance, but without improvement in mood, pain or QoL[31]. The studies were limited by small sample size and short duration of follow-up. The authors noted that, neither herbal, plant-based nor were synthetic cannabis products currently licensed for FM in any jurisdiction. A trial of a pharmacological cannabinoid preparation in FM for sleep disturbance [40] (n=2705), used an online questionnaire for self-reporting. Only 383 (1.4%) responses were received. Only 44% of those who used cannabis (84%) used medicinal cannabis (MC), and 55% of them used it in combination with commercial cannabis (CC). Only those who used CC alone or in combination with MC, showed improvement in pain (94%), sleep (93%), depression (87%), anxiety (62%), ability to work (64%) and ability to drive (74%) [41]. In another study by Habib and Artul [42] MC was used (n=26, 18-34g/month for up to 2.2 mo). Using the FIQ, patients self-reported improvements in all parameters. Additionally, 50% stopped taking any other medications for fibromyalgia [42]. The review of Banerjee et al. supported these findings [43]. The review by Goen and Amital [44] discussed that chronic pain including FM is common component in all rheumatoid diseases, and does not respond well to traditional medications. The authors acknowledged the use of MC in the treatment of FM and that the evidence on long-term efficacy and safety are still controversial. Animal studies have shown accumulated immune-modulatory effect, however, this has not yet studied in human to the same extent [44].

Sagy et al. [45] found that in their sample (n=367, 82% females), the use of MC for 6-months reduced pain significantly (<p<0.001) and 82% of patients experienced remission. This suggests the standardisation of MC compounds and regimens, which is hindering the compilation of evidence from the current literature. Boehnke et al. [46] used a cross-sectional, anonymous survey to examine CBD efficacy in FM (n=2701). Participants self-reported reduction in pain on a scale provided, and only 50% reported minor side effects.

Van de Donk [47] conducted a placebo-controlled 4-way crossover trial comparing 4-varieties of inhaled MC in the treatment of FM compared to a placebo. None of the tested varieties was superior to placebo in managing spontaneous or ‘electrical’ (neuropathic) pain. In another study, Yasin and Robinson [48] concluded that conventional analgesia provided slight pain improvement in FM compared to baseline, whilst a greater improvement was achieved when MC was added and was maintained at 6 mo follow-up. Chaves et al. [49] explored the benefit of a THC-rich cannabis oil on symptoms and QoL of FM in 17 women (8 in intervention group and 9 in placebo group) with a low socioeconomic profile in Brazil. The dose was titrated up according to symptoms during an 8-weeks period and FIQ was used before and after dosing. The authors concluded that the cannabis group had a significant improvement in their FIQ on the "feel good," "pain," “do work,” and “fatigue” items compared to the placebo group at the end of the intervention, however long-term benefit and safety were not assessed. The systematic review by Fitzcharles et al. [28] included 4 RCTs covering 160 patients with FM, RA or musculoskeletal pain, examining the efficacy of nabilone. They found no significant difference between nabilone and placebo with respect to pain relief in FM patients and in patients with musculoskeletal pain. However Cameron et al. [50] noted serious methodological limitations in published studies, that prevent researchers and policymakers from being able to make a definitive conclusion on cannabinoid use for pain management in FM.

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The committee noted that some studies concluded chronic pain improvement but that the improvement was statistically and clinically insignificant (mean of 0.4 on a scale ranging from 0 to 10) and patients continued to use other pharmacological pain management products at the same level. They concluded the cost outweighed the benefit, making it not sustainable. They found there was no evidence for the use of CBD alone (either as a pure product or containing traces of THC). Therefore, the committee recommended that CBD should not be offered unless as part of a clinical trial. People who have FM or persistent treatment-resistant neuropathic pain are often taking high doses of medicines for pain relief over long periods. These can cause nausea, drowsiness, mood disturbance and fatigue. The committee noted that this is a significant population of people with chronic pain (around 15%). The committee recommended the research centre CBD in adults with FM and/or treatment-resistant neuropathic pain [51].

Possible reasons for the current poor evidence

It is clear that even optimal therapy with conventional analgesics leaves many patients with residual pain and seeking further solutions, and the internet has empowered discussion across international boundaries. Patients are communicating, and as shown in the study by Li [10] many are self-medicating or at least experimenting with self-medication using a wide variety of forms and products. As stated in the EU report [38] this need, in the context of the diverse international legislative framework and low levels of control of ‘natural products’ has led to a very wide diversity in products. Medical products are strictly controlled, containing accurately defined amounts of one or more stated ingredients or extracts. The controls for complementary therapies are less rigid. A licensed medicine will normally contain between 99% and 101% of the stated ingredient. Even when a plant extract, it will normally be accurately standardised on one or two stated ingredients, but herein lies an important issue. A typical plant extract will have several hundred identifiable compounds. The total amount, and relative proportions, of these, are affected by: the variety of the plant, the conditions in which it was grown (soil and climatic conditions), the season, time of day and specific weather conditions under which the plant was harvested, storage conditions and processes applied. Common extraction procedures include aqueous (neutral, acid or alkaline) or alcoholic extraction; all of these will produce different proportions of individual compounds. To minimise these variables, clinical trials of complementary products commonly use the product from a single batch.

The study by Li [10] was novel. A smartphone application (App) (Rekap+) was used to report the use of the cannabinoids for pain relief, the form used, the relief gained and negative aspects experienced in real-time. A total of 2,987 people reported 20,513 cannabinoid administrations. Their results reflected the factors mentioned above. Almost all patients reported pain relief. On a typical pain scale from 0-10 average reported pain relief was 3 points on a par with a mid-level pharmaceutical analgesic. None of the derived products reported was found as effective as the whole cannabis flower. Reported relief varied with different types of pain, although encouragingly for use in FM, it appeared most effective in musculoskeletal pain, but standard deviations were wide, showing large individual variation in response. Use of concentrates appeared to increase reports of negative side effects without increased pain relief compared to whole cannabis.
chronic pain [n=468], showed 73% of participants decreased [and some ceased completely] to use their prescription opioids and 31% discontinued benzodiazepines, however, again, nearly one third of patients had little or no benefit, and almost 50 % of patients noted feelings of intoxication.

There remains therefore clear evidence that Cannabis can be of considerable benefit to certain patients, with some types of pain, but is very far from being a panacea. The current licenced medical products derived from cannabis show a very different pattern of effects compared to whole plant derived administration methods. A great deal more research is required to clearly establish its role in FM.

Cannabis has well-known drawbacks as a medicine, including the potential for dependence and addiction; risk of motor vehicle accidents, psychotic and psychopathic experiences and cognitive impairment [15], but have been claimed to be relatively less severe than misuse of either prescription or non-prescription opioids [10]. Although studies have linked cannabis use to an increased risk of opioid misuse [15, 52], the US experience suggests that extending the legal use of cannabis has reduced opioid misuse amongst patients with chronic pain [53, 54].

CONCLUSION

Recent studies suggest that, in time, the use of cannabinoids will add a useful contribution to pain management in this challenging condition, but the evidence to date does not support the use of the products currently available in the UK. If governments could be persuaded to formalise the legal framework around cannabis products internationally, this would greatly facilitate the design of larger, multi-centre trials, and encourage the development of products manufactured to the standards for licensed medicines, whilst better controlling the poorly regulated products found in many countries.

Should cannabis-based products be used to manage FM? For the present, not unless as part of controlled and regulated clinical trials.

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AUTHORS CONTRIBUTIONS

Authors 1and2 conducted the initial review and first draft. Authors 3 and 4 reviewed the evidence and completed the final draft of the manuscript.

CONFLICT OF INTERESTS

No conflict of interest known

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