Cannabidiol in Humans—The Quest for Therapeutic Targets

Simon Zhornitsky and Stéphane Potvin

Abstract: Cannabidiol (CBD), a major phytocannabinoid constituent of cannabis, is attracting growing attention in medicine for its anxiolytic, antipsychotic, antiemetic and anti-inflammatory properties. However, up to this point, a comprehensive literature review of the effects of CBD in humans is lacking. The aim of the present systematic review is to examine the randomized and crossover studies that administered CBD to healthy controls and to clinical patients. A systematic search was performed in the electronic databases PubMed and EMBASE using the key word “cannabidiol”. Both monotherapy and combination studies (e.g., CBD + ∆9-THC) were included. A total of 34 studies were identified: 16 of these were experimental studies, conducted in healthy subjects, and 18 were conducted in clinical populations, including multiple sclerosis (six studies), schizophrenia and bipolar mania (four studies), social anxiety disorder (two studies), neuropathic and cancer pain (two studies), cancer anorexia (one study), Huntington’s disease (one study), insomnia (one study), and epilepsy (one study). Experimental studies indicate that a high-dose of inhaled/intravenous CBD is required to inhibit the effects of a lower dose of ∆9-THC. Moreover, some experimental and clinical studies suggest that oral/oromucosal CBD may prolong and/or intensify ∆9-THC-induced effects, whereas others suggest that it may inhibit ∆9-THC-induced effects. Finally, preliminary clinical trials suggest that high-dose oral CBD (150–600 mg/d) may exert a therapeutic effect for social anxiety disorder, insomnia and epilepsy, but also that it may cause mental sedation. Potential pharmacokinetic and pharmacodynamic explanations for these results are discussed.
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Key words: cannabidiol; THC; cannabis; multiple sclerosis; pain; social anxiety disorder; epilepsy; insomnia; schizophrenia

1. Introduction

The cannabis plant has been used by humans for thousands of years in medicine for its sedative/hypnotic, antidepressant, analgesic, anticonvulsant, antiemetic, anti-inflammatory, anti-spasmodic and appetite-stimulating effects [1]. The plant is composed of a complex chemical mixture that includes phytocannabinoids, terpenoids, flavanoids, steroids and enzymes [2]. Phytocannabinoids—the most cannabis-specific of these constituents—bind to receptor sites normally activated by endogenous cannabinoids such as anadamide and 2-arachidonylglycerol (2-AG). It is widely believed the most psychoactive phytocannabinoid is delta-9-tetrahydrocannabinol (\(\Delta^9\)-THC), which acts as a partial agonist at cannabinoid CB1 receptors—found primarily in the central nervous system (CNS), and CB2 receptors—found primarily on cells of the immune system [3,4]. However, apart from \(\Delta^9\)-THC, a number of other phytocannabinoids are present in significant quantities in cannabis (e.g., cannabidiol, cannabinol, cannabichromene), and they may be responsible for some of the plant’s many putative medicinal properties. In animal studies, cannabidiol (CBD) has been receiving growing attention for its antiemetic, anticonvulsant, anti-inflammatory, and antipsychotic properties [5–8]. This broad range of therapeutic effects may be a result of CBD’s complex pharmacological mechanisms [9]. Apart from \(\Delta^9\)-THC, CBD is the sole cannabinoid that has been thoroughly tested in humans in numerous controlled experimental studies as well as clinical trials for multiple sclerosis, neuropathic pain, schizophrenia, bipolar mania, social anxiety disorder, insomnia, Huntington’s disease and epilepsy. Surprisingly, however—up to this point—reviews and meta-analyses on the topic of CBD in humans have not considered a large number of experimental and clinical studies that administered CBD-alone and/or in combination with \(\Delta^9\)-THC, versus \(\Delta^9\)-THC-alone [10–12]. The inclusion of these studies is essential to understanding the therapeutic potential of CBD and its mediation by pharmacokinetic and pharmacodynamic factors.

The present review is aimed to comprehensively examine the effects of CBD in humans. We will begin with a brief overview of the pharmacokinetic and pharmacodynamic properties of CBD. Next, we will systematically examine the controlled experimental and clinical trials of CBD in order to elucidate its potential therapeutic role in human central nervous system (CNS) disorders.

2. Pharmacokinetics

CBD undergoes a significant first-pass effect leading to the formation of a number of metabolites, most notably, 7-hydroxy-CBD and CBD-7-oic acid [13,14]. The half-life of CBD in humans was found to be between 18–33 h following intravenous administration, 27–35 h following smoking, and 2–5 days following oral administration. Bioavailability of oral and smoked CBD in humans was found to be around 6% and 31%, respectively, providing further support for a substantial first-pass effect [13,15–17]. Oral administration of CBD (~700 mg) over six weeks to 14 Huntington’s disease patients resulted in a low, narrow plasma range of 5.9–11.2 ng/mL [15]. Oral cannabis extract (10 mg
Δ9-THC; 10 mg CBD) produced markedly lower levels of CBD (range = 0–2.6 ng/mL) at 30–120 min after administration and absorption was increased with food [18,19].

Recent in vitro studies have shown that CBD is a potent inhibitor of multiple cytochrome P450 enzymes including CYP1A2, CYP2B6, CYP2C9, CYP2D6 and CYP3A4 [20–23]. Consequently, CBD may be expected to exhibit significant pharmacokinetic interaction with other pharmacological agents. In some studies, CBD has been shown to slightly augment levels of Δ9-THC (metabolized by CYP2C9, CYP2C19, and CYP3A4) by decreasing its conversion to 11-hydroxy-THC [19,24]. Moreover, animal studies found that CBD reduced the potency of some anticonvulsants and enhanced the potency of others; however, it is uncertain whether this effect resulted from a pharmacokinetic mechanism [25,26]. Pharmacokinetic interactions with other medications are probable, but studies are lacking.

3. Pharmacodynamics

CBD possesses affinity for CB1 and CB2 receptors in the micromolar range; however, despite this very low affinity, CBD seems to antagonize CB1/CB2 agonists with $K_B$ values in the nanomolar range [9]. Some have suggested that the reason for these conflicting findings may be that CBD acts as a non-competitive inverse agonist, thereby blocking the ability of agonists to activate CB1/CB2 receptors [9]. Moreover, CBD has been found to antagonize the putative novel cannabinoid receptor GPR55, and the abnormal-CBD receptor at nanomolar concentrations [27,28]. In addition, there is evidence that CBD activates 5-HT1A serotonergic and TRPV1–2 vanilloid receptors, antagonizes alpha-1 adrenergic and µ-opioid receptors, and inhibits synaptosomal uptake of noradrenaline, dopamine, serotonin and gaminobutyric acid and cellular uptake of anandamide at micromolar concentrations [29–32]. Studies also suggest that CBD may act on mitochondria Ca$^2+$ stores, block low-voltage-activated (T-type) Ca$^2+$ channels, and stimulate activity of the inhibitory glycine-receptor [33,34]. Finally, CBD has been shown to both stimulate and to inhibit activity of fatty amide hydrolase (FAAH; responsible for the degradation of anandamide) [35–37].

4. Methods

A systematic search was performed in the electronic databases PubMed and EMBASE using the key word “cannabidiol”. This search looked for human randomized and crossover studies published up to 1 April 2012. Both monotherapy and combination studies (e.g., CBD + Δ9-THC) were included. Studies that administered CBD in the form of cannabis cigarettes were included if the percentage of CBD was provided (studies which compared cannabis cigarettes with negligible amounts of CBD (<1%) were excluded). Pharmacokinetic studies and studies that only compared the combination of CBD/Δ9-THC with placebo were excluded. Finally, studies that compared different routes of administration (e.g., oral versus oromucosal) were excluded.

5. Results

A total of 34 studies were identified. Sixteen of these were experimental studies, conducted in healthy subjects (Table 1) and 18 were conducted in clinical populations (Table 2). Of the clinical trials
included patients with multiple sclerosis (six studies), schizophrenia and bipolar mania (four studies), social anxiety disorder (two studies), neuropathic and cancer pain (two studies), cancer anorexia (one study), Huntington’s disease (one study), insomnia (one study), and epilepsy (one study).

5.1. Experimental Studies in Healthy Controls

5.1.1. Oral or Intravenous CBD-Alone

Six studies administered oral CBD-alone to healthy volunteers. An early study by Hollister [38] did not find any subjective or physiological effects with oral or intravenous CBD (100 mg PO and 30 mg IV) among 10 healthy volunteers. Additionally, a crossover study of oral CBD (200 mg) with, and without alcohol revealed no effect of the former on time production, finger tapping, cancellation test, and differential aptitude test [39]. There was also no difference in performance on these tests when CBD was added to alcohol, versus alcohol-alone; however, plasma alcohol levels in the CBD group were significantly lower compared to the alcohol-alone group. Another crossover study among 11 healthy volunteers revealed that plasma cortisol levels decreased during placebo treatment (in agreement with its normal circadian rhythm) and this decrease was attenuated by oral CBD (300 or 600 mg) [40]. Here, subjects reported CBD to have a sedative effect. A parallel-group study by the same authors compared the effects of diazepam, CBD (300 mg) and ipsapirone (a 5-HT1a agonist) among 40 individuals on anxiety before, during, and after a speech test [41]. Their results revealed that diazepam decreased anxiety before and after the speech test, whereas ipsapirone decreased it during, and CBD decreased it only after the speech test. More recently, a crossover study by Crippa et al. [42] showed that CBD (400 mg) decreased subjective anxiety and increased mental sedation among 10 healthy subjects, relative to placebo. Another crossover study found that treatment with 10 mg oral Δ9-THC increased levels of anxiety, intoxication, sedation, and psychotic symptoms among 15 participants, whereas CBD (600 mg) was inactive [43,44]. The authors also found that Δ9-THC increased the number of skin conductance response fluctuations during processing of intensely fearful faces, whereas CBD decreased it and there was a trend for reduced anxiety [45,46].

5.1.2. Oral CBD/Ketamine

One crossover study examined the effects of oral CBD (600 mg) or placebo pretreatment on ketamine-induced psychiatric symptoms among 10 healthy volunteers [47]. Results revealed that significantly CBD increased ketamine-induced activation (as measured by the Brief Psychiatric Rating Scale, but failed to reduce ketamine-induced positive and negative symptoms, relative to placebo.

5.1.3. Oral CBD/Nabilone

One crossover study examined the effects of oral CBD (200 mg) alone, and combined with nabilone (1 mg), relative to nabilone alone, in nine male subjects [48]. Here, CBD and nabilone caused mild sedation when administered alone. Moreover, CBD marginally reduced nabilone-induced intoxication and impairment in binocular depth perception—a model of impaired perception during psychotic states.
<table>
<thead>
<tr>
<th>Study</th>
<th>N (CBD)</th>
<th>Dosing</th>
<th>Outcome (≥ greater; ≤ less)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hollister [38]</td>
<td>9 (5)</td>
<td>Fixed-dose; CBD 100 mg, PO; CBD 30 mg, IV</td>
<td>CBD = no subjective or physiological effects</td>
</tr>
</tbody>
</table>
| Consroe et al. [39]   | 10 (10) | Fixed-dose; CBD 200 mg, PO                  | CBD = PBO (time production)  
CBD = PBO (finger tapping)  
CBD = PBO (cancellation test)  
CBD = PBO (differential aptitude test)                                                                                                                                 |
| Zuardi et al. [40]    | 11 (11) | Fixed-dose; CBD 300 mg or 600 mg, PO       | CBD > PBO (sedation)  
CBD < PBO (normal circadian decrease in cortisol level)                                                                                                                                                                 |
| Zuardi et al. [41]    | 40 (10) | Fixed-dose; CBD 300 mg; DZP 10 mg; IPS 5 mg, PO | DZP < IPS < CBD < PBO (speech test-induced anxiety)                                                                                                                                                                       |
| Crippa et al. [42]    | 10 (10) | Fixed-dose; CBD 400 mg, PO                  | CBD > PBO (mental sedation)  
CBD < PBO (anxiety)                                                                                                                                                                                                          |
| Borgwardt et al. [43] | 15 (15) | Fixed-dose; CBD 600 mg; Δ9-THC 10 mg, PO    | CBD < PBO < Δ9-THC (skin conductance response to fearful faces)  
CBD < PBO (anxiety p = 0.06)  
CBD = PBO (sedation, intoxication)                                                                                                                                                                                   |
| Winton-Brown et al. [44] | 15 (15) | Fixed-dose; CBD 600 mg, Δ9-THC 10 mg, PO   | CBD > PBO (ketamine-induced activation [BPRS])  
CBD = PBO (ketamine-induced positive and negative symptoms)                                                                                                                                                             |
| Fusar-Poli et al. [45,46]| 15 (15) | Fixed-dose; CBD 600 mg, Δ9-THC 10 mg, PO   | CBD > PBO (ketamine-induced activation [BPRS])  
CBD = PBO (ketamine-induced positive and negative symptoms)                                                                                                                                                             |
| Hallak et al. [47]    | 10 (10) | Fixed-dose; CBD 600 mg, PO; ketamine 0.25 mg/kg, IV | CBD > PBO (ketamine-induced activation [BPRS])  
CBD = PBO (ketamine-induced positive and negative symptoms)                                                                                                                                                             |
| Leweke et al. [48]    | 9 (9)   | Fixed-dose; CBD 200 mg; NAB 1 mg, PO        | NAB > CBD + NAB > CBD (binocular depth perception deficit)  
CBD & NAB > PBO (sedation)  
NAB > CBD + NAB > CBD & PBO (intoxication)                                                                                                                                                                             |
| Karniol et al. [49]   | 40 (5)  | Fixed-dose; CBD 15 mg, 30 mg, 60 mg; Δ9-THC 30 mg, PO | CBD [15 mg] + Δ9-THC > Δ9-THC (pulse rate)  
CBD [30 & 60 mg] + Δ9-THC < Δ9-THC (pulse rate)  
Δ9-THC > CBD [all doses] + Δ9-THC (time production impairment)                                                                                                                                                        |
| Hollister and Gillespie [50]| 15 (15) | Fixed-dose; CBD 40 mg; Δ9-THC 20 mg, PO    | CBD + Δ9-THC > Δ9-THC (duration and intensity of intoxication)  
CBD + Δ9-THC > Δ9-THC (time to onset of intoxication)  
CBD + Δ9-THC = Δ9-THC (pulse rate)                                                                                                                                                                               |
### Table 1. Cont.

<table>
<thead>
<tr>
<th>Study</th>
<th>N (CBD)</th>
<th>Dosing</th>
<th>Outcome (≥ greater; ≤ less)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zuardi et al. [51]</td>
<td>8 (8)</td>
<td>Fixed-dose; CBD 1 mg/kg; Δ9-THC 0.5 mg/kg, PO</td>
<td>Δ9-THC &gt; CBD + Δ9-THC (anxiety, intoxication)</td>
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<tr>
<td></td>
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<td>CBD + Δ9-THC = Δ9-THC (pulse rate)</td>
</tr>
<tr>
<td>Juckel et al. [52]</td>
<td>24 (24)</td>
<td>Fixed-dose; CBD 5.4 mg; Δ9-THC 10 mg, PO</td>
<td>CBD + Δ9-THC &gt; Δ9-THC (MMN amplitude)</td>
</tr>
<tr>
<td>Roser et al. [53,54]</td>
<td></td>
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<td>CBD + Δ9-THC &lt; PBO (right-hand tapping frequency)</td>
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<td></td>
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<td>CBD + Δ9-THC = Δ9-THC (P300 amplitude)</td>
</tr>
<tr>
<td>Nicholson et al. [55]</td>
<td>8 (8)</td>
<td>Fixed-dose; CBD 15 mg; Δ9-THC 15 mg, OMC</td>
<td>CBD + Δ9-THC &lt; Δ9-THC (impairment of immediate and delayed word recall)</td>
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<tr>
<td></td>
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<td></td>
<td>CBD + Δ9-THC = Δ9-THC (digit symbol substitution, choice reaction time, sustained attention, six-letter memory recall)</td>
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<td></td>
<td></td>
<td></td>
<td>CBD + Δ9-THC &gt; Δ9-THC (awake time before sleep, sleepiness and fatigue upon awakening)</td>
</tr>
<tr>
<td>Dalton et al. [56]</td>
<td>15 (15)</td>
<td>Fixed-dose; CBD 150 µg/kg; Δ9-THC 25 µg/kg, INH</td>
<td>Δ9-THC &gt; CBD + Δ9-THC (intoxication)</td>
</tr>
<tr>
<td>Ilan et al. [57]</td>
<td>23 (23)</td>
<td>Fixed-dose; CBD (1% versus 0.2%) Δ9-THC (3.6% versus 1.8%), INH</td>
<td>CBD + Δ9-THC = Δ9-THC (heart rate, intoxication)</td>
</tr>
<tr>
<td>Bhattacharyya et al. [58]</td>
<td>6 (6)</td>
<td>Fixed-dose; CBD 5 mg; Δ9-THC 1.25 mg, IV</td>
<td>Δ9-THC &gt; CBD + Δ9-THC (positive symptoms)</td>
</tr>
</tbody>
</table>

IPS = ipsapirone; DZP = diazepam; NAB = nabilone; OMC = oromucosal administration; PO = oral administration; CBD = cannabidiol; Δ9-THC = delta-9-tetrahydrocannabinol; IV = intravenous; BPRS = Brief Psychiatric Rating Scale; MMN = mismatch negativity; PBO = placebo.

### Table 2. Clinical trials.

<table>
<thead>
<tr>
<th>Study</th>
<th>N (CBD)</th>
<th>Subjects</th>
<th>Time</th>
<th>Dosing</th>
<th>Outcome(s) (≥ greater; ≤ less)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Consroe et al. [15]</td>
<td>15 (15)</td>
<td>Huntington’s</td>
<td>6 weeks</td>
<td>Flexible-dose; CBD 700 mg, PO</td>
<td>CBD = PBO (chorea severity)</td>
</tr>
<tr>
<td>Carlini and Cunha [59]</td>
<td>15 (15)</td>
<td>Insomnia</td>
<td>Acute</td>
<td>Fixed-dose; CBD 40 mg, 80 mg, 160 mg, NTZ 5 mg PO</td>
<td>CBD [160 mg] &gt; PBO (sleep duration)</td>
</tr>
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<td></td>
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<td></td>
<td>CBD [all doses] &lt; PBO (dream recall)</td>
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<td>CBD [all doses] = NTZ = PBO (sleep induction)</td>
</tr>
<tr>
<td>Study</td>
<td>N (CBD)</td>
<td>Subjects</td>
<td>Time</td>
<td>Dosing</td>
<td>Outcome(s) (≥ greater; ≤ less)</td>
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<tr>
<td>Cunha et al. [61]</td>
<td>15 (8)</td>
<td>Epilepsy</td>
<td>2–18 weeks</td>
<td>Flexible-dose; CBD 200–300 mg, PO</td>
<td>CBD &lt; PBO (seizures)</td>
</tr>
<tr>
<td>Crippa et al. [62]</td>
<td>10 (10)</td>
<td>Social anxiety disorder</td>
<td>Acute</td>
<td>Fixed-dose; CBD 400 mg, PO</td>
<td>CBD &lt; PBO (anxiety)</td>
</tr>
<tr>
<td>Bergamaschi et al. [63]</td>
<td>24 (12)</td>
<td>Social anxiety disorder</td>
<td>Acute</td>
<td>Fixed-dose; CBD 600 mg, PO</td>
<td>CBD &lt; PBO (anxiety)</td>
</tr>
<tr>
<td>Leweke et al. [64]</td>
<td>42 (21)</td>
<td>Schizophrenia</td>
<td>4 weeks</td>
<td>Fixed-dose; CBD 600 mg; AMI 600 mg, PO</td>
<td>CBD = AMI (positive symptoms)</td>
</tr>
<tr>
<td>Zuardi et al. [65]</td>
<td>3 (3)</td>
<td>Schizophrenia</td>
<td>4 weeks</td>
<td>Fixed-dose; CBD—up to 1,280 mg, PO</td>
<td>CBD = PBO (positive and negative symptoms)</td>
</tr>
<tr>
<td>Zuardi et al. [66]</td>
<td>2 (2)</td>
<td>Bipolar I disorder</td>
<td>4 weeks</td>
<td>Fixed-dose; CBD—up to 1,280 mg, PO</td>
<td>CBD = PBO (mania)</td>
</tr>
<tr>
<td>Hallak et al. [67]</td>
<td>28 (9)</td>
<td>Schizophrenia</td>
<td>Acute</td>
<td>Fixed-dose; CBD 300 mg or 600 mg, PO</td>
<td>CBD [600 mg] &gt; CBD [300 mg] &amp; PBO (Stroop Color Word Test errors)</td>
</tr>
<tr>
<td>Killestein et al. [68,69]</td>
<td>16 (16)</td>
<td>Multiple sclerosis</td>
<td>4 weeks</td>
<td>Flexible-dose; Δ9-THC 5–10 mg; Cannabis extract 5–10 mg (20–30% CBD), PO</td>
<td>CBD + Δ9-THC &gt; Δ9-THC &gt; PBO (side-effects); CBD + Δ9-THC &amp; Δ9-THC = PBO (spasticity); CBD + Δ9-THC &gt; PBO (TNF-alpha)</td>
</tr>
<tr>
<td>Zajicek et al. [70] (CAMS)</td>
<td>630 (211)</td>
<td>Multiple sclerosis</td>
<td>15 weeks</td>
<td>Flexible-dose; CBD (to 12.5 mg/d); Δ9-THC (to 25 mg/d), PO</td>
<td>CBD + Δ9-THC &amp; Δ9-THC = PBO (pain); CBD + Δ9-THC &amp; Δ9-THC = PBO (spasticity); CBD + Δ9-THC &amp; Δ9-THC = PBO (spasms); CBD + Δ9-THC &amp; Δ9-THC = PBO (sleep quality)</td>
</tr>
<tr>
<td>Freeman et al. [71] (CAMS-LUTS)</td>
<td>255 (88)</td>
<td>Multiple sclerosis</td>
<td>13 weeks</td>
<td>Flexible-dose; CBD (to 12.5 mg/d); Δ9-THC (to 25 mg/d), PO</td>
<td>CBD + Δ9-THC &amp; Δ9-THC &lt; PBO (urinary incontinence)</td>
</tr>
<tr>
<td>Strasser et al. [72]</td>
<td>243 (95)</td>
<td>Cancer anorexia</td>
<td>6 weeks</td>
<td>Fixed-dose; CBD 2 mg; Δ9-THC 5 mg, PO</td>
<td>CBD + Δ9-THC &amp; Δ9-THC = PBO (appetite, nausea, mood)</td>
</tr>
<tr>
<td>Study</td>
<td>N (CBD)</td>
<td>Subjects</td>
<td>Time</td>
<td>Dosing</td>
<td>Outcome(s) (≥ greater; ≤ less)</td>
</tr>
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</tbody>
</table>
| Johnson et al. [73] | 177 (60)| Cancer pain                             | 2 weeks| Flexible-dose; CBD 20–30 mg; Δ9-THC 22–32 mg, OMC | CBD + Δ9-THC < PBO (pain; NRS)  
Δ9-THC < PBO (pain; BPI-SF)  
CBD + Δ9-THC > PBO (nausea)  
CBD + Δ9-THC & Δ9-THC > PBO (cognitive deficits) |
| Brady et al. [74]   | 15 (15) | Multiple sclerosis                      | 8 weeks| Flexible-dose; CBD & Δ9-THC 34 mg #, OMC     | Δ9-THC < BAS (spasticity)  
Δ9-THC > BAS (sleep quality)  
CBD + Δ9-THC & Δ9-THC < BAS (pain)  
CBD + Δ9-THC & Δ9-THC < BAS (incontinence) |
| Wade et al. [75]    | 20 (20) | Multiple sclerosis (14/20) + neuropathic pain | 2 weeks| Flexible-dose; CBD & Δ9-THC 45 mg #, OMC     | Δ9-THC & CBD < PBO (pain; VAS)  
CBD + Δ9-THC & Δ9-THC & CBD = PBO (pain; NRS)  
CBD + Δ9-THC & Δ9-THC < PBO (spasms; VAS)  
Δ9-THC > PBO (appetite; VAS)  
CBD + Δ9-THC > PBO (sleep quality; VAS)  
Δ9-THC > PBO (memory impairment) |
| Notcutt et al. [76] | 34 (34) | Multiple sclerosis (16/34) + neuropathic pain | 5 weeks| Flexible-dose; CBD & Δ9-THC 2.5 mg per spray, OMC | CBD + Δ9-THC & Δ9-THC < CBD & PBO (pain)  
CBD + Δ9-THC & Δ9-THC > CBD > PBO (sleep quality) |
| Berman et al. [77]  | 48 (48) | Neuropathic pain                        | 2 weeks| Flexible-dose; CBD & Δ9-THC 20 mg or 8–10 sprays per day #, OMC | CBD + Δ9-THC & Δ9-THC < PBO (pain; BS-11)  
Δ9-THC < PBO (pain; SF-MPQ)  
CBD + Δ9-THC & Δ9-THC = PBO (pain disability)  
CBD + Δ9-THC & Δ9-THC > PBO (sleep quality) |

CBD = cannabidiol; Δ9-THC = delta-9-tetrahyrdocannabinol; AMI = amisulpride; MS = multiple sclerosis; SAD = social anxiety disorder; OMC = oromucosal; PO = oral; PBO = placebo; IV = intravenous; INH = inhalation; # = mean dose; BPI-SF = Brief Pain Inventory Short Form; SF-MPQ = short-form McGill Pain Questionnaire; VAS = visual analogue scale; NRS = numerical rating scale; NTZ = nitrazepam; BAS = versus baseline value.
5.1.4. Oral CBD/Δ9-THC

Four studies administered oral CBD alone and/or together with oral Δ9-THC. A parallel-group study tested different doses of oral CBD (15, 30, 60 mg) alone, and combined with oral Δ9-THC (30 mg), relative to Δ9-THC-alone (30 mg) and placebo in 40 male subjects [49]. The authors found that, when given alone, CBD had little effect on pulse rate and psychological outcomes. Interestingly, there was a non-significant increase of 53% in pulse rate following the combination of CBD (15 mg) and Δ9-THC (30 mg); however, there was a significant decrease in pulse rate when the higher doses of CBD (30 and 60 mg) were combined with Δ9-THC. Furthermore, the 30 mg and 60 mg doses of CBD significantly attenuated Δ9-THC-induced increases in pulse rate as well as the number of “psychological reactions” (anxiety and panic), and all doses of CBD reversed Δ9-THC-induced impairment on a time estimation task. Moreover, a crossover study compared the addition of oral CBD (40 mg) to oral Δ9-THC (20 mg), relative to Δ9-THC-alone in 15 male subjects [50]. Results revealed that that CBD slightly increased time to onset, overall intensity, and duration of the subjective intoxication produced by oral Δ9-THC (20 mg), without affecting pulse rate. Another crossover study administered oral CBD (1 mg/kg) alone, and in combination with oral Δ9-THC (0.5 mg/kg) to eight male and female subjects [51]. The authors found that CBD had little effect on its own; however, it reduced Δ9-THC associated subjective intoxication and anxiety, without affecting pulse rate.

More recently, a crossover trial randomized 27 male and female subjects to treatment with oral CBD (5.4 mg) combined with oral Δ9-THC (10 mg), compared to Δ9-THC-alone and placebo [52–54]. First, the authors evidenced that the CBD/Δ9-THC combination significantly increased auditory evoked mismatch negativity (MMN) amplitude, relative to placebo, whereas Δ9-THC-alone exerted no effect [52]. Second, they found that Δ9-THC treatment led to a reduction of P300 amplitude, but this effect was not reversed by CBD [53]. Third, they found that the CBD/Δ9-THC combination, but not Δ9-THC-alone, reduced right-hand tapping frequencies versus placebo [54]. Finally, no significant differences were found for subjective intoxication or plasma levels of Δ9-THC or its metabolites in the subjects as a whole.

5.1.5. Oromucosal CBD/Δ9-THC

One crossover study investigated the effects of a combination of oromucosal CBD (15 mg) and Δ9-THC (15 mg), relative to Δ9-THC-alone (15 mg) on sleep and cognition in eight male and female subjects [55]. Measures were taken before sleep, during sleep and upon awakening. Results demonstrated that Δ9-THC-alone increased sleepiness 30 min after rising, and decreased latencies to early morning sleep, relative to placebo. The CBD/Δ9-THC combination increased awake time before sleep, but also increased sleepiness and fatigue, compared to placebo. Lastly, no significant differences between the treatments were noticed on digit symbol substitution, choice reaction time, sustained attention, and six-letter memory recall. However, Δ9-THC-alone attenuated immediate word recall and delayed word recall, whereas the CBD/Δ9-THC combination did not [55].
5.1.6. Smoked or Intravenous CBD/Δ9-THC

Three studies administered CBD/Δ9-THC through smoking/intravenously. One early crossover study investigated the effects of smoked CBD (150 µg/kg) alone, and in conjunction with smoked Δ9-THC (25 µg/kg), relative to Δ9-THC-alone in 15 male subjects [56]. The authors found that CBD failed to exert any effects on its own and did not change Δ9-THC-induced increase in heart rate as well as impairment of stability of stance, motor performance, manual coordination and working memory. On the other hand, CBD decreased the “psychological high” associated with Δ9-THC ($p < 0.05$) [56]. Another crossover study administered cannabis cigarettes with various concentrations of Δ9-THC (3.6% versus 1.8%), CBD (1% versus 0.2%) and other minor cannabinoids to 23 male and female subjects [57]. They found CBD did not affect increases in heart rate and subjective intoxication produced by Δ9-THC. However, participants who received the lower dose of Δ9-THC tended to report more anxiety when paired with the higher dose CBD, relative to when paired with the lower dose CBD. By contrast, participants who received the higher Δ9-THC dose reported less anxiety when CBD content was high and more anxiety when CBD content was low [57]. Lastly, a crossover study administered CBD (5 mg) and Δ9-THC (1.25 mg) intravenously, relative to Δ9-THC-alone, in six male and female subjects [58]. The authors found that the addition of CBD blocked Δ9-THC-induced increases in Positive and Negative Syndrome Scale (PANSS) total scores.

5.2. Clinical Trials in Patient Populations

5.2.1. Oral CBD-Alone

Nine clinical trials administered CBD-alone via the oral route. One early crossover trial comparing CBD (mean dose = 700 mg/d) with placebo among 15 Huntington’s disease patients did not find any significant differences in chorea severity, side-effects, clinical lab tests and other safety outcomes after 6-weeks of treatment [15]. On the other hand, a crossover trial comparing CBD (40, 80, and 160 mg) with placebo and nitrazepam (5 mg) among 15 insomniac volunteers revealed that duration of sleep significantly increased following administration of the high-dose CBD (160 mg); however, dream recall was reduced, relative to placebo [59,60]. The same authors subsequently conducted a 6-week, placebo-controlled, parallel-group study of CBD (200–300 mg/d), added to antiepileptic drugs, among 15 treatment-refractory epileptic patients [61]. Here, they found that four out of eight CBD-treated patients evidenced significant improvement in their condition, whereas only one patient improved in the placebo group.

More recently, a crossover study demonstrated that CBD (400 mg) decreased subjective anxiety among 10 treatment-naïve patients with social anxiety disorder, relative to placebo, and this was accompanied with significant changes in regional cerebral blood flow [62]. Similarly, a placebo-controlled, parallel-group study among 24 treatment-naïve social anxiety disorder patients showed that CBD (600 mg) significantly reduced anxiety, cognitive impairment, and discomfort in speech performance, in response to a simulation public speaking test [63].

Preliminary data from a double-blind, randomized trial of 42 patients with acute schizophrenia revealed that CBD (600 mg) and amisulpride equally reduced psychotic symptoms after four weeks of treatment [64]. However, a placebo-controlled case-series did not find CBD to be effective among
three treatment-resistant schizophrenia patients over four weeks [65]. Another placebo-controlled case-series by the same authors did not find a significant benefit of CBD for two bipolar mania patients after about four weeks of treatment [66]. Finally, a parallel-group among 28 schizophrenia patients that individuals who were treated with the low dose of CBD (300 mg) and placebo improved significantly more on the Stroop Color Word Test over two experimental sessions, relative to those treated with the high dose of CBD (600 mg) [67].

5.2.2. Oral CBD/Δ9-THC

Four clinical trials administered CBD/Δ9-THC via the oral route. One crossover study treated 16 multiple sclerosis (MS) patients with flexible doses of cannabis extract (5–10 mg [20–30% CBD]), relative to Δ9-THC-alone (5–10 mg) over the course of four weeks. Results demonstrated that the CBD/Δ9-THC combination resulted in significantly more adverse events (e.g., dizziness, somnolence, ataxia), compared to Δ9-THC-alone. No positive trends in efficacy (e.g., pain, tremor, spasticity, cognition) were noted for either of the treatments and they equally worsened participants’ global impressions, relative to placebo [68]. An immunological analysis revealed that patients who were treated with the CBD/THC combination evidenced a modest increase in TNF-alpha in LPS-stimulated whole blood and patients with high adverse event scores had an increase in plasma IL-12p40 [69]. Both of these immunomodulators have been linked with disease progression in MS. In addition, a parallel-group study treated 630 MS patients (Cannabinoids in Multiple Sclerosis [CAMS] study) with flexible doses of oral CBD (up to 12.5 mg/d) combined with Δ9-THC (up to 25 mg/d), versus Δ9-THC-alone and placebo [70]. The authors noted no evidence for a distinction between the treatments in efficacy (e.g., pain, tremor, spasticity, sleep) or adverse events, except for a tendency for CBD/Δ9-THC to increase gastrointestinal side-effects, relative to Δ9-THC-alone. A sub-analysis showed that CBD/Δ9-THC and Δ9-THC-alone significantly reduced urinary incontinence versus placebo [71]. On the other hand, a parallel-group study that treated 243 cancer-related anorexia patients with fixed doses of oral CBD (2 mg/d) and Δ9-THC (5 mg/d), relative to Δ9-THC-alone and placebo was terminated at interim analysis due to lack of difference between study arms [72].

5.2.3. Oromucosal CBD/Δ9-THC

Five trials administered CBD/Δ9-THC via oromucosal sublingual drops. A parallel-group study treated 177 patients with cancer-related intractable pain with flexible doses of oromucosal CBD (20–30 mg/d) and Δ9-THC (22–32 mg/d), relative to Δ9-THC-alone and placebo [73]. Results showed that the combination of CBD and Δ9-THC was significantly better than placebo at decreasing pain on the neurological rating scale (NRS), whereas Δ9-THC-alone showed a non-significant reduction. By contrast, Δ9-THC-alone was more efficient than placebo at decreasing mean total pain on the Brief Pain Inventory—Short Form (BPI-SF; last 24 h) and no significant differences were found in EORTC Quality Of Life questionnaire (QLQ-C30) pain subscore or the amount of breakthrough opiate medication that was required. Additionally, CBD/Δ9-THC increased nausea and vomiting on the QLQ-C30 subscore, but not on the NRS, relative to placebo. Finally, no significant differences were noted on the NRS memory or concentration subscores, or on the QLQ-C30 cognitive functioning subscore [73]. In addition, a crossover trial that treated 15 MS patients with a 1:1 ratio of CBD/Δ9-THC
relative to ∆9-THC-alone revealed that patients preferred ∆9-THC-alone because they found it more effective for controlling urinary symptoms and needed less of it to achieve a therapeutic effect [74]. Analysis of secondary outcome measures revealed that ∆9-THC-alone was significantly better than CBD/∆9-THC for spasticity and sleep but both treatments equally improved VAS pain scores [74]. Another crossover trial treated 24 patients with MS and neuropathic pain with CBD/∆9-THC (1:1 ratio; 2.5 mg per spray), relative to CBD-alone, ∆9-THC-alone and placebo [75]. Analysis of data showed that CBD-alone and ∆9-THC-alone significantly improved VAS pain scores versus placebo; however, none of the treatments improved pain as measured by the NRS. Moreover, both CBD/∆9-THC and ∆9-THC-alone improved spasm, but only the latter improved spasticity and appetite, whereas only the former improved sleep quality. Lastly, administration of ∆9-THC-alone produced the greatest subjective intoxication and reduction in Short Orientation-Memory-Concentration test score, relative to placebo [75].

Furthermore, a crossover trial treated 34 patients with MS and neuropathic pain CBD/∆9-THC (1:1 ratio; 2.5 mg per spray), compared to CBD-alone, ∆9-THC-alone and placebo [76]. Results revealed that CBD/∆9-THC and ∆9-THC-alone were equally beneficial for pain and all three treatments (including CBD-alone, but less so) improved sleep quality, relative to placebo. Of the 28 patients that obtained benefit, 14 found CBD/∆9-THC and ∆9-THC equally satisfactory, 11 preferred CBD/∆9-THC, two preferred ∆9-THC-alone, and one found ∆9-THC-alone and CBD-alone equally satisfactory [76]. An additional crossover trial treated 48 neuropathic pain patients (brachial plexus avulsion) with CBD/∆9-THC (1:1 ratio; 2.5 mg per spray), compared to ∆9-THC-alone and placebo [77]. The authors found that the combination of ∆9-THC-alone (but not CBD/∆9-THC) significantly decreased pain on the short form McGill Questionnaire, relative to placebo. However, both treatments significantly decreased pain ratings on an 11-point Box Scale and neither treatment significantly decreased pain on the Pain Disability Index. Finally, both treatments equally improved sleep quality.

6. Discussion

Experimental studies suggest that high-dose CBD may decrease anxiety and increase mental sedation in healthy individuals. Clinical trials suggest the high-dose CBD may be useful for the treatment of social anxiety disorder, and possibly, insomnia and epilepsy. The anxiolytic effect associated with CBD may be the result of its 5-HT1A agonism, which has been evidenced in a number of behavioral studies [78–81]. Paradoxically, some animal studies have found that the dose-response of CBD follows an inverted U shape, leading to an anxiogenic effect through its agonism of TRPV1–2 receptors (which are believed to be responsible for detection and regulation of body temperature, and thermal nociception) [82]. Alternatively, it is possible that the anxiolytic properties of CBD are mediated by its action at CB1 receptors, because CB1 antagonists were found to attenuate amphetamine and/or nicotine-induced anxiety in mice [83]. Indeed, there is evidence that both CBD and AM404 (an anandamide transporter/FAAH inhibitor and TRPV1 agonist) facilitated extinction of contextual fear memory in rats and this was reversed by the CB1-receptor antagonist SR141716A, but not by the TRPV1-selective antagonist, capsazepine [84]. It is equally possible that the anxiolytic effect of CBD is explained by inhibition of FAAH. Animal studies have demonstrated that FAAH inhibitors possess anxiolytic properties in a number of paradigms including marble burying, light/dark box, elevated zero maze, and isolation-induced ultrasonic emission test [85–87]. However, CBD has been shown to both
inhibit and stimulate activity FAAH [35–37]. Consequently, it is yet unclear whether there is a role for inhibition of endocannabinoid catabolic enzymes in the anxiolytic effects of CBD.

A possible analgesic effect of CBD-alone, and CBD added to ∆9-THC was observed in two studies among mixed neurogenic (MS and neuropathic pain) and cancer pain patients [73,75]. However, both studies administered low doses of CBD (2.5 mg CBD and ∆9-THC per spray), used more than one scale to measure pain outcomes and their results were not consistent across scales. Also confounding interpretation is the fact that some of the clinical trials allowed the use of “rescue medication”, which may have led overestimation of the effects of CBD-alone [75–76]. In support, other studies did not find an analgesic effect of CBD-alone, or in combination with ∆9-THC [70,74,77]. In animals, there is evidence that ∆9-THC and cannabiol (a weak partial CB1 agonist) suppressed the abdominal constriction response to formic acid in mice, whereas CBD was inactive at doses of up to 200 mg/kg [88]. In that study, the analgesic effects of ∆9-THC and cannabiol were additive and CBD antagonised these effects in a dose-dependent manner. Likewise, ∆9-THC, cannabiol and cannabis extract produced an analgesic effect in the hot-plate test in mice; however, CBD was without effect at doses of up to 30 mg/kg [89]. By contrast, CBD (5 mg/kg IP or 25 mg/kg PO) was shown to block disease progression in murine collagen-induced arthritis—an animal model of rheumatoid arthritis—and this was accompanied by reductions in type-II collagen-specific proliferation, interferon—gamma production, and release of tumour necrosis factor-alpha by synovial cells [90]. The same group later demonstrated that the synthetic CBD derivative—HU-320—exerted more potent effects in the same direction [91]. More recently, there is evidence that cannabidiol derivative, O-1602, reduced movement-evoked firing of nociceptive C fibres in a rat model of acute inflammatory joint pain [92]. Interestingly, this effect was blocked by the GPR55-receptor antagonist, O-1918, but not by the CB1 and CB2 antagonists, AM281 and AM630, respectively. As a whole, these data indicate that CBD and its analogues may be beneficial for pain resulting from inflammation, however, human studies on this topic are lacking.

The strength of the antiepileptic effects of CBD may be difficult to judge clinically because of its potent antagonism of multiple CYP isoenzymes, potentially reducing plasma levels of concomitant anticonvulsants. Preclinical data has indicated that CBD displays antiepileptiform and antiseizure properties in vitro and in vivo CBD may possess antiepileptic properties via different mechanisms. For instance, there is evidence that CBD can block low-voltage-activated (T-type) Ca2+ channels, and increase the activity of inhibitory glycine receptors [33,34]. More recently, Jones et al. [5] used extracellular multi-electrode array recordings to show that CBD decreased epileptiform activity in the Mg2+-free and 4-aminopyridine in vitro models of hippocampal epilepsy in the mammalian hippocampus—a key epileptogenic brain region. Additionally, the authors examined the effects of CBD (1, 10, and 100 mg/kg) in vivo using the pentylenetetrazole model of generalized seizures. Their results revealed that the incidence of severe seizures and mortality was significantly attenuated in rats treated with the high dose of CBD (100 mg/kg), relative to vehicle-treated rats. The antiepileptic effects associated with CBD were suggested to be due a potentially CB1 independent mechanism because CBD acted with only low affinity at CB1 receptor and displayed no agonist activity in [35S]guanosine 5'-O-(3-thio)-triphosphate assays in cortical membranes. In support of this interpretation, there is evidence that the anticonvulsant properties of CBD in the maximal electroshock
model were not affected by the CB1-receptor antagonist, SR141716A, whereas those of ∆9-THC and the CB1-receptor agonist, WIN55,212-2, were blocked [93].

Despite its putative benefits for social anxiety disorder, insomnia and epilepsy studies suggest that high-dose CBD (400–700 mg) may increase mental sedation in normal individuals and aggravate cognitive deficits in schizophrenia—without altering physical sedation [42,62,67]. While the mechanism that is responsible for these effects is not clear, the fact that they exist is not surprising because most anxiolytics/sedatives/anticonvulsants produce their therapeutic action by decreasing CNS activation, and consequently, alertness. Some research does suggest, however, that CBD may improve cognition when used in combination with ∆9-THC. For instance, there is evidence that mixed neurogenic patients given oromucosal CBD/∆9-THC performed as well as patients given placebo on the Short Orientation-Memory-Concentration test, whereas patients given ∆9-THC-alone performed significantly worse than placebo-treated patients [75]. Similarly, there is evidence that healthy subjects given oromucosal CBD/∆9-THC (15 and 15 mg) performed equally well as placebo-treated individuals on tests of delayed and immediate word recall, whereas subjects treated with ∆9-THC-alone performed significantly worse than placebo-treated subjects on those tasks and they exhibited less wakefulness [55]. However, the treatments were not significantly different from placebo on digit symbol substitution, choice reaction, sustained attention, six-letter memory recall, digit memory recall. Moreover, Karniol et al. [49] showed that oral CBD (15, 30, and 60 mg) inhibited the time production impairment associated with ∆9-THC (30 mg) [49]. On the other hand, Dalton et al. [56] found that a high dose of smoked CBD (150 µg/kg) failed to block perturbations of stability of stance, motor performance, mental performance induced by a much lower dose of smoked ∆9-THC (25 µg/kg).

Current evidence is equivocal regarding a potential antipsychotic effect of CBD. For example, Zuardi et al. [65,66] did not find CBD monotherapy (up to 1,280 mg) to be effective relative to placebo in a case-series of bipolar mania and treatment-resistant schizophrenia patients. Likewise, the CB1 antagonist, SR141716, was ineffective for the treatment of positive or negative symptoms in schizophrenia [94]. In another study, oral CBD (600 mg) enhanced the psychomotor activating effects of intravenous ketamine, without significantly altering ketamine-induced psychiatric symptoms [47]. On the other hand, preliminary data from a four-week, randomized-controlled trial of CBD (600 mg) versus amisulpride (600 mg) for schizophrenia did not reveal any significant differences between the groups—suggesting that the former exerted an antipsychotic effect [64].

Intriguingly, some studies show that CBD can potentiate and some that it can attenuate the psychotomimetic effects associated with ∆9-THC, depending on the measure, route of administration, and dose-ratio between the cannabinoids. For example, Karniol et al. [49] found that a low dose of CBD potentiated ∆9-THC-induced increases in pulse rate, whereas an equal or higher dose of CBD attenuated these increases. A dose-dependent interaction was also evidenced in the study by Ilan et al. [57] wherein high doses of CBD potentiated anxiety induced by low doses of ∆9-THC, but they reduced anxiety induced by high doses of ∆9-THC. In addition, Hollister and Gallespie [50] found that oral CBD (40 mg) caused a slight delay and prolongation/intensification of the psychotomimetic effects of ∆9-THC (20 mg). By contrast, Dalton et al. [56] showed that a high dose of smoked CBD (150 µg/kg) minimally ($p < 0.05$) inhibited the psychotomimetic effects associated with smoked ∆9-THC (25 µg/kg). However, there is also evidence that large doses of intravenous CBD (5 mg) completely blocked elevations in PANSS positive symptoms induced by intravenous ∆9-THC (1.25 mg) [58].
Another group found significantly greater MMN amplitude values at central electrodes following treatment with combined CBD (5.4 mg) and Δ9-THC (10 mg; but not Δ9-THC-alone), indicating that the former may have exerted an antipsychotic effect [52]. Finally, a study using nabilone (1 mg) showed that the molecule significantly impaired binocular depth inversion (an illusion of visual perception that provides a model of impaired perception during psychotic states) and this effect was partially reversed by CBD (200 mg) [48].

Overall, the human data regarding CBD’s potential to reverse the cognitive perturbations and psychotomimetic symptoms induced by Δ9-THC are difficult to interpret due to the possibility of a pharmacokinetic interaction between CBD and Δ9-THC (or other molecules) following oral/oromucosal administration. A review of 1970s studies found that the ratio of CBD/Δ9-THC was 8.1 when the CBD displayed antagonistic effects and 1.8 when it enhanced the effects of Δ9-THC [95,96]. Moreover, there is evidence that combination of CBD (1–10 mg/kg IP over 21 days) with equivalent doses of Δ9-THC increased blood and brain levels of the latter, decreased levels of 11-OH-THC and THC-COOH, and augmented the anxiogenic and locomotor suppressant effects and social interaction deficits seen with Δ9-THC [97]. Interestingly, CBD did not change the THC-induced decrease in CB1 receptor binding and none of the treatments altered 5-HT1A binding, suggesting that pharmacokinetic factors may have played a role.

The presence of a pharmacokinetic interaction between CBD and Δ9-THC is supported by results of two early phase studies of nabiximols—an oromucosal spray that contains an equivalent dose of the cannabinoids [98,99]. For instance, the addition of CBD (20 mg) to Δ9-THC (20 mg) significantly increased the area under-the-curve (AUC) of 11-hydroxy-THC [98]. The same group also demonstrated that concomitant CBD (10 mg) and Δ9-THC (10 mg) lead to a significantly later Tmax for Δ9-THC, relative to treatment with Δ9-THC-alone [99]. These pharmacokinetic data roughly correspond to mean intoxication ratings across time, indicating that CBD slightly delayed and prolonged the subjective effects associated with Δ9-THC. By contrast, Nadulski et al. [18,19] found that oral CBD (5.4 mg), combined with Δ9-THC (10 mg) (non-significantly) increased the AUC and maximum concentration of Δ9-THC by approximately 20%, suggesting that CBD inhibited the conversion of Δ9-THC into 11-hydroxy-THC. Nevertheless, the pharmacokinetic impact of CBD was small compared to other factors such as gender and body mass index.

On the other hand, it remains true that a few human studies showed that oral/oromucosal CBD attenuated the psychoactive and therapeutic effects associated with Δ9-THC at low doses and dose-ratios between the cannabinoids [49,55,74,75]. Indeed, Karniol et al. [49] found that oral CBD (15 mg) was sufficient to reverse Δ9-THC-induced (30 mg) impairment on a time estimation task. In a similar fashion, there is evidence that oromucosal CBD (15 mg) attenuated verbal memory deficits induced by Δ9-THC (15 mg) [55]. Clinical studies using equal ratios of CBD to Δ9-THC show that the former may alter both the benefits and side-effects associated with the latter. For instance, CBD attenuated the antiemetic effects of Δ9-THC in cancer patients [73]. Moreover, CBD attenuated the antispastic, memory-impairing, and intoxicating effects associated with Δ9-THC and increased the prevalence of adverse events among MS patients (e.g., dizziness, ataxia, gastrointestinal) [68,71,73–75]. Nonetheless, clinical trials did not consistently show significant benefit/drawbacks of combining CBD with Δ9-THC. Some of the variability in results may be attributed to the fact that studies contained small sample sizes and measured multiple variables, leading to the possibility of a Type-I error(s).
Likewise, some variability may be explained by the fact that a number of clinical trials treated patients with CBD/Δ9-THC during a “run-in” period, which may have biased results towards CBD/Δ9-THC because individuals who did not respond or could not tolerate the medication would have withdrawn early [74–76]. Alternatively, CBD may exhibit a flat dose-response curve, whereby all doses are able to partially reverse the effects of Δ9-THC because of its non-competitive antagonist action at CB₁ receptors [9]. A final explanation for the disparate results is that oral CBD has the ability to attenuate/potentiate some and (but not other) effects associated with Δ9-THC due to activity at receptors other than CB₁, even at low doses and small ratios of CBD/Δ9-THC. Such an effect may be one reason why some studies found contradictory results using similar dose-ratios between the cannabinoids [51,55].

7. Conclusions

Experimental studies indicate that a high-dose of inhaled/intravenous CBD is required to inhibit the effects of a lower dose of Δ9-THC. Some experimental and clinical studies also suggest that oral/oromucosal CBD may prolong and/or intensify Δ9-THC-induced effects, whereas others suggest that it may inhibit Δ9-THC-induced effects. A balance between pharmacokinetic and pharmacodynamic factors may be responsible for the disparate findings, depending on the measure, route of administration and dose-ratio between the cannabinoids. Moreover, preliminary clinical trials suggest that high-dose oral CBD (150–600 mg/d) may exert a therapeutic effect for social anxiety disorder, insomnia and epilepsy, but also that it may cause mental sedation. On the other hand, trials did not consistently observe any benefits/drawbacks of adding low-dose CBD to Δ9-THC for clinical conditions such as MS, neuropathic and cancer pain, and cancer-associated anorexia. Likewise, studies did not consistently observe benefits of CBD monotherapy in bipolar mania or schizophrenia patients.

Future studies should investigate clinical applications of high-dose oral CBD for disorders such as anxiety, neuropathic pain, inflammatory pain, multiple sclerosis, insomnia and epilepsy. Future trials should also administer CBD to clinical patients for prolonged periods of time in order to simulate the “real world” setting. If CBD is not found to be beneficial in these trials, new more selective and more bioavailable molecules need to be developed in order to harness the full therapeutic potential of cannabinoid molecules. Currently, the most promising candidates are inhibitors of endocannabinoid catabolic enzymes (e.g., anandamide, FAAH) for the treatment of anxiety and depressive disorders [100].

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