Effectiveness of Cannabidiol Oil for Pediatric Anxiety and Insomnia as Part of Posttraumatic Stress Disorder: A Case Report

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ABSTRACT

Introduction: Anxiety and sleep disorders are often the result of posttraumatic stress disorder and can contribute to an impaired ability to focus and to demonstration of oppositional behaviors.

Case Presentation: These symptoms were present in our patient, a ten-year-old girl who was sexually abused and had minimal parental supervision as a young child under the age of five. Pharmaceutical medications provided partial relief, but results were not long-lasting, and there were major side effects. A trial of cannabidiol oil resulted in a maintained decrease in anxiety and a steady improvement in the quality and quantity of the patient’s sleep.

Discussion: Cannabidiol oil, an increasingly popular treatment of anxiety and sleep issues, has been documented as being an effective alternative to pharmaceutical medications. This case study provides clinical data that support the use of cannabidiol oil as a safe treatment for reducing anxiety and improving sleep in a young girl with posttraumatic stress disorder.

INTRODUCTION

Cannabidiol (CBD) oil is a naturally occurring constituent of industrial hemp and marijuana, which are collectively called cannabis. CBD oil is 1 of at least 85 cannabinoi d compounds found in cannabis and is popular for its medicinal benefits. After tetrahydrocannabinol (THC), CBD oil is the second-most-abundant component of cannabis. Other names for CBD oil include CBD-rich hemp oil, hemp-derived CBD oil, or CBD-rich cannabis oil. Considered to be generally safe, CBD has been used medicinally for decades. However, CBD is not medical marijuana and should be distinguished from high-CBD strains of medical marijuana, which do contain THC, such as “Charlotte’s Web.”

The most abundant compound in cannabis, THC is also a cannabinoi d. The THC component induces the psychoactive effect, “high.” A cannabis plant has different amounts of CBD and THC depending on the strain and thus provides different recreational or medicinal effects. The cannabinoi d profile of industrial hemp or medical marijuana is ideal for people looking for the medical benefits of CBD without the “high” of the THC.

The mechanism of action of CBD is multifold. Two cannabinoi d receptors are known to exist in the human body: CB1 and CB2 receptors. The CB1 receptors are located mainly in the brain and modulate neurotransmitter release in a manner that prevents excessive neuronal activity (thus calming and decreasing anxiety), as well as reduces pain, reduces inflammation, regulates movement and posture control, and regulates sensory perception, memory, and cognitive function.

An endogenous ligand, anandamide, which occurs naturally in our bodies, binds to the CB1 receptors throughout the G-protein coupling system. CBD has an indirect effect on the CB1 receptors by stopping the enzymatic breakdown of anandamide, allowing it to stay in the system longer and provide medical benefits.

CBD has a mild effect on the CB2 receptors, which are located in the periphery in lymphoid tissue. CBD helps to mediate the release of cytokines from the immune cells in a manner that helps to reduce inflammation and pain.

Other mechanisms of action of CBD include stimulation of vanilloid pain receptors (TRPV-1 receptor), which are known to mediate pain perception, inflammation, and body temperature. In addition, CBD may exert its anti-anxiety effect by activating adenosine receptors which play a significant role in cardiovascular function and cause a broad anti-inflammatory effect throughout the body. At high concentrations, CBD directly activates the 5-HT1A serotonin receptor, thereby conferring an antidepressant effect. Cannabidiol has been found to be an antagonist at the potentially new third cannabinoi d receptor, GPR55, in the caudate nucleus and putamen, which if stimulated may contribute to osteoporosis.

Since the 1940s, a considerable number of published articles have dealt with the chemistry, biochemistry, pharmacology, and clinical effects of CBD. The last decade has shown a notable increase in the scientific literature on CBD, owing to its identification for reducing nausea and vomiting, combating psychotic disorders, reducing inflammation, decreasing anxiety and depression, improving sleep, and increasing a sense of well-being. Findings presented at the 2015 International Cannabinoi d Research Society at its 25th Annual Symposium reported the use of CBD as beneficial for kidney fibrosis and inflammation, metabolic syndrome, overweight and obesity, anorexia-cachexia syndrome, and modification of osteoarthritic and other musculoskeletal conditions.

Although studies have demonstrated the calming, anti-inflammatory, and relaxing effects of CBD, clinical data from actual cases is minimal. This case study offers evidence that CBD is effective as a safe alternative treatment to traditional psychiatric medications for reducing anxiety and insomnia.

CASE PRESENTATION

A ten-year-old girl presented in January 2015 for a reevaluation of behaviors related to her diagnosis of posttraumatic stress disorder (PTSD) secondary to sexual
abuse. Her chief issues included anxiety, insomnia, outbursts at school, suicidal ideation, and self-destructive behaviors. Her grandmother, who has permanent custody of the patient and her younger brother, accompanied her.

Our patient had been seen for an initial evaluation in January 2012 and received a diagnosis of PTSD secondary to sexual abuse on the basis of her history, clinical observations, and behaviors (Table 1).

Her father had died 6 months earlier in a motor vehicle accident, and our patient’s maternal grandparents became her permanent guardians. Before her father’s death, our patient had no supervision from her father and very little supervision from her mother. An 11-year-old boy had molested her when she was 3 years old. Her medical history included her mother having methadone addiction, alcoholism, bipolar disorder, and depression. Her mother used marijuana her entire pregnancy with the girl. The patient presented in January 2012 as displaying aggressive, disobedient, impulsive, and sexually inappropriate behaviors. She also demonstrated low self-esteem and anxiety and had poor sleep (restless, interrupted, and unable to sleep alone).

Workup during 2012 included laboratory studies, which ruled out a thyroid dysfunction and an iron or vitamin D deficiency. The patient was started on a

<table>
<thead>
<tr>
<th>Date</th>
<th>Presentation</th>
<th>Medications</th>
<th>Supplements</th>
<th>Other</th>
</tr>
</thead>
<tbody>
<tr>
<td>January 31, 2012</td>
<td>New evaluation: 7.5-year-old girl. History of sexual abuse and neglect. Issues: Insomnia, sexual behaviors. Diagnosis: PTSD secondary to sexual abuse.</td>
<td>None</td>
<td>Melatonin, 1 mg/night</td>
<td>February 14, 2012, laboratory values: TSH, 2.46 mIU/L (reference range, 0.47-4.68 mIU/L); ferritin: 21 ng/mL (reference range, 10-150 ng/mL). February 16, 2012, laboratory values: Vitamin D₃: 39 ng/mL (reference range, 20-50 ng/mL).</td>
</tr>
<tr>
<td>February 20, 2012</td>
<td>Sleeping 2-3 hours/night. Started counseling; Cooperative and good behavior at counseling session. Anxious, traumatized.</td>
<td>Clonidine, 0.05 mg (half tablet) at bedtime</td>
<td>Inositol, 3 g 3 times/d; EPA fish oil, 500 mg/d</td>
<td>Eye movement desensitization and reprocessing therapy recommended</td>
</tr>
<tr>
<td>February 22, 2012</td>
<td>Did not do well with clonidine because of hallucinations, so she discontinued that treatment. Behavior still very rough; sleep poor.</td>
<td>Started imipramine therapy, 25 mg at bedtime</td>
<td></td>
<td>March 7, 2012: ECG was normal</td>
</tr>
<tr>
<td>August 8, 2012*</td>
<td>Good summer. In play therapy. Overall better sleep and energy with imipramine therapy. Patient’s 6-year-old brother also now in therapy.</td>
<td>Imipramine, 25 mg at bedtime</td>
<td></td>
<td></td>
</tr>
<tr>
<td>January 21, 2015</td>
<td>Returned for evaluation and treatment after 3 years. Suicidal ideation; cut self on leg; defiant and stubborn. Had psychotherapy 3 years straight twice a month. Sleeps with brother; can’t sleep alone.</td>
<td>Off all medications for past 18 months</td>
<td>Melatonin, 5 mg; St John’s wort, 450 mg twice/d; magnesium, 300 mg/d; diphenhydramine, 25 mg/night</td>
<td></td>
</tr>
<tr>
<td>February 16, 2015</td>
<td>Hard to manage. Has outbursts at school.</td>
<td>Magnesium and St John’s wort: stopped treatment; EPA fish oil, 750 mg/d; diphenhydramine, 25 mg/night</td>
<td></td>
<td>February 11, 2015: Normal cortisol and DHEA levels</td>
</tr>
<tr>
<td>March 16, 2015</td>
<td>Better overall. Started animal-assisted therapy.</td>
<td>EPA fish oil, 750 mg/d; diphenhydramine, 25 mg/night</td>
<td></td>
<td>Started a regimen of CBD oil, 25 mg (1 capsule/d) at 6 pm</td>
</tr>
<tr>
<td>April 14, 2015</td>
<td>Sleeping better with CBD treatment. Getting biofeedback. Has stomachaches. Mood is more at ease.</td>
<td>EPA fish oil, 750 mg/d; diphenhydramine, 25 mg/night</td>
<td>CBD oil, 25 mg (1 capsule/d) at 6 pm</td>
<td></td>
</tr>
<tr>
<td>May 26, 2015</td>
<td>“Ghosts” waking patient up at night.</td>
<td>EPA fish oil, 750 mg/d</td>
<td>CBD oil, 25 mg (1 capsule/d) at 6 pm</td>
<td></td>
</tr>
<tr>
<td>July 22, 2015</td>
<td>Sleeping better; able to sleep in own room 3-4 nights/wk.</td>
<td>EPA fish oil, 750 mg/d</td>
<td>CBD liquid, 12 mg (in 4 sublingual sprays)/night; 12 mg more (in 4 sublingual sprays) during the day as needed for anxiety, typically 3 or 4 times/wk</td>
<td></td>
</tr>
<tr>
<td>August 24, 2015</td>
<td>Sleeping well. Handling school well.</td>
<td>EPA fish oil, 750 mg/d</td>
<td>CBD oil, 25 mg (1 capsule)/night; CBD liquid, 6-12 mg (in 2-4 sublingual sprays) as needed for anxiety, typically 2 or 3 times/wk</td>
<td></td>
</tr>
</tbody>
</table>

* There were additional visits in 2012 with no substantial changes.

CBD = cannabidiol; DHEA = dehydroepiandrosterone; ECG = electrocardiogram; EPA = eicosapentaenoic acid; PTSD = posttraumatic stress disorder; TSH = thyroid stimulating hormone.
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regimen of 1 mg/night of melatonin, which helped her sleep duration. Three grams of inositol 3 times a day and 500 mg/d of eicosapentaenoic fish oil were also helpful in reducing her anxiety. A trial of clonidine was implemented, which resulted in hallucinations and thus was discontinued. The patient was switched to a regimen of 25 mg of imipramine at bedtime to decrease her anxiety, which appeared to be helpful. Counseling sessions were started. The patient continued psychotherapy for 3 years, but she was not seen again in our clinic until the return visit in January 2015, when she was not receiving any of her medications and supplements.

At the patient’s return in January 2015, she demonstrated the same prominent symptoms as at her initial presentation. At that time, the initial treatment included the following supplements and medications to assist with her sleep and anxiety: melatonin, 5 mg/night; magnesium, 300 mg/d; and diphenhydramine (Benadryl), 25 mg/night. Our patient demonstrated slight gains but was still having outbursts at school and was reportedly difficult to manage at home. In addition, her underlying anxiety continued.

Cannabidiol oil was explored as a potential additional treatment to help her insomnia and anxiety, but we deferred for two months while we waited for a response from other interventions. The grandmother preferred reducing the pharmacologic load given her granddaughter’s medication history. Consequently, the grandmother (her caregiver) reported: “My granddaughter provided full informed consent. Our patient was administered the Sleep Disturbance Scale for Children and the Screen for Anxiety Related Disorders (SCARED) before taking the CBD oil and each month afterward for the next 5 months. Test scores on the Sleep Disturbance Scale for Children and Screen for Anxiety Related Disorders demonstrated an improvement (Table 2).

A trial of CBD supplements (25 mg) was then initiated at bedtime, and 6 mg to 12 mg of CBD sublingual spray was administered during the day as needed for anxiety. A gradual increase in sleep quality and quantity and a decrease in her anxiety were noted. After 5 months, the patient was sleeping in her own room most nights and handling the new school year with no difficulties. No side effects were observed from taking the CBD oil.

DISCUSSION

Studies repeatedly recognize the prevalence of an anxiety-provoked sleep disorder after a traumatic experience. Our patient was definitely experiencing this phenomenon, which was aggravated by daily stressful activities.

The main finding from this case study is that CBD oil can be an effective compound to reduce anxiety and insomnia secondary to PTSD. A review of the literature suggests some benefits from the use of CBD because of its anxiolytic and sleep-inducing effects. Animal studies support use of this treatment and report that “CBD may block anxiety-induced [rapid eye movement] sleep alteration via its anxiolytic effect on the brain.”

The strength of this particular case is that our patient was receiving no pharmacological medications (other than non-prescription diphenhydramine) but only nutritional supplements and the CBD oil to control her symptoms. Her scores on the sleep scale and the anxiety scale consistently and steadily decreased during a period of 5 months (see Table 2). She was ultimately able to sleep through the night most nights in her own room, was less anxious at school and home, and displayed appropriate behaviors. The patient’s grandmother (her caregiver) reported: “My grandmother’s behaviors are definitely better being on the CBD. Her anxiety is not gone, but it is not as intense and she is much easier to be around. She now sleeps in her own room most of the time, which has never happened before.”

Further study will need to be conducted to determine the permanency of our patient’s positive behaviors and how long she will need to continue taking the CBD oil. We do not have a reasonable foundation to recommend dosing from the scientific literature. However, in our experience, this supplement given 12 mg to 25 mg once daily appears to provide relief of key symptoms with minimal side effects. Our patient did not voice any complaints or discomfort from the use of CBD. We routinely asked about headache, fatigue, and change in appetite or agitation in addition to conducting a routine psychiatric evaluation. Although CBD is considered generally safe, the long-term effects are yet to be studied.

The ultimate goal is to gradually taper her off the use of CBD oil and transition our patient into lifelong coping strategies such as yoga, meditation, and various other therapeutic activities.

Table 2. Patient’s clinical progress in sleep and anxiety

<table>
<thead>
<tr>
<th>Date of visit</th>
<th>Sleep scale scorea</th>
<th>SCARED scoreb</th>
</tr>
</thead>
<tbody>
<tr>
<td>March 16, 2015</td>
<td>59</td>
<td>34</td>
</tr>
<tr>
<td>May 25, 2015</td>
<td>42</td>
<td>24</td>
</tr>
<tr>
<td>July 22, 2015</td>
<td>41</td>
<td>19</td>
</tr>
<tr>
<td>August 24, 2015</td>
<td>37</td>
<td>16</td>
</tr>
<tr>
<td>September 22, 2015</td>
<td>38</td>
<td>18</td>
</tr>
</tbody>
</table>

a A score of more than 50 is considered indicative of a sleep disorder on the Sleep Disturbance Scale for Children.
b A SCARED score over 25 indicates a high probability of a childhood anxiety disorder.

SCARED = Screen for Anxiety Related Disorders.

References

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Marijuana and Medicine

Scientific data indicate the potential therapeutic value of cannabinoid drugs, primarily [tetrahydrocannabinol], for pain relief, control of nausea and vomiting, and appetite stimulation; smoked marijuana, however, is a crude [tetrahydrocannabinol] delivery system that also delivers harmful substances.