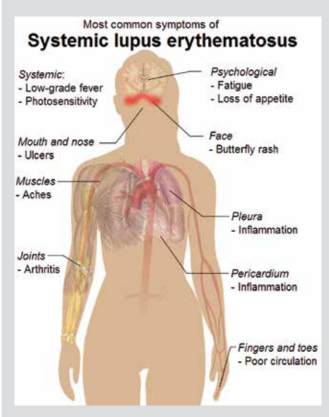
Cannabidiol in the Management of Comorbid Rheumatoid Arthritis, Lupus, and Raynaud's Disease

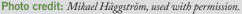
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Introduction

Rheumatoid arthritis (RA), systemic lupus erythematosus (SLE), and Raynaud's disease, are chronic inflammatory autoimmune diseases characterized by pain, inflammation, and fatigue.¹⁻³ Treatment presents a clinical challenge for several reasons, including the progressively degenerative nature of autoimmune diseases, the involvement of multiple pain mechanisms, and the adverse side effects of pain medications. Even pain treatments with low addiction profiles may pose an implicit risk, such as liver or kidney toxicity.

Presently, there are limited, if any, modern studies examining the effects of cannabidiol (CBD) products on pain and other outcomes in RA, SLE, or Raynaud's disease.⁴ This case





report describes the potential efficacy and safety of a daily, high-dose, medical grade CBD product (ie, "Hemp CBD") in the treatment of persistent pain and inflammation in a patient with multiple autoimmune disorders.

In autoimmune disorders such as RA, SLE, and Raynaud's disease, an abnormal and chronic inflammatory response occurs in various tissues that over time results in the observed degenerative features and symptoms of the conditions. For many patients with these diseases, pain and accompanying loss of mobility are the most common and debilitating daily symptoms.

Currently, use of cannabinoids in the treatment of autoimmune conditions in the United States presents both clini-

cians and patients with considerable challenges, including the lack of conformity between individual state and federal cannabis/hemp laws, minimal funding to support the clinical study of hemp- and cannabis-derived products, heterogeneity of patient symptomology (particularly in elderly patients), and quality inconsistency of cannabis/hemp-derived products.4-6 Multiple substantiated sources suggest that CBD's anti-inflammatory properties are significant.^{7,8} There also are anecdotal patient reports of symptom relief when using CBD products for inflammatory conditions. However, there currently is a lack of general knowledge about the effect of cannabinoids in autoimmune diseases and potential dosing regimens. The authors of a recent meta-analysis stated that, "There are no clinical trials of medical cannabis in rheumatology arthritis."9 A few studies have investigated the effects of cannabis obtained outside of a state program (ie, illicitly) in RA, but to our knowledge, no previously published clinical data or case reports exist on the efficacy of CBD-containing products compliant with state and federal regulations outlined in the 2018 Farm Bill in patients suffering from advanced autoimmune disorders.^{6,10} The aim of this article is to provide clinicians and patients with new insights on treatment and dosing applications of CBD for inflammatory disorders.

Medication	Dosage	Condition	Provider	Duration, Year
Gabapentin	300 mg daily	Pain	Primary Care 1	30 days, 2015
Gabapentin	600 mg daily	Pain	Primary Care 1	60 days, 2015
Gabapentin	1200 mg daily	Pain	Primary Care 1	90 days, 2015
Prednisone	20 mg daily	Inflammation	Primary Care 2	30 days, 2016
Methocarbamol	500 mg PO, 4x daily	Pain	Primary Care 2	30 days, 2016
Potassium	1500 mg (20 mEq) daily	Muscle cramping	Primary Care 2	30 days, 2016
Tramadol	50 mg as needed, but not to exceed 150 mg daily	Pain	Primary Care 2	30 days, 2016
Prednisone	20 mg daily	Inflammation	Primary Care 2	90 days, 2017
Tizanidine	4 mg PO x 8 h	Pain	Primary Care 2	30 days, 2017
Prednisone	10 mg daily	Swelling	Rheumatologist	1 year, 2018
Leflunomide	20 mg daily	Inflammation	Rheumatologist	1 year, 2018
Amlodipine	10 mg daily	Finger ulcers	Rheumatologist	1 year, 2018
Nitro paste	25 mg nightly	Finger ulcers	Rheumatologist	1 year, 2018
CBD isolate medium-chain triacylglyceride oil tincture	600 mg daily	Pain	Preventive Medicine Physician	60 days, 2019
CBD isolate medium-chain triacylglyceride oil tincture	400 mg daily	Pain	Preventive Medicine Physician	60 days, 2019
CBD isolate medium-chain triacylglyceride oil tincture	200 mg daily	Pain	Preventive Medicine Physician	Present

Table 1. Medication History

Medical History

The patient is a 50-year-old woman with pain and mobility-related symptoms of multiple autoimmune disorders. She was diagnosed with Raynaud's disease in 2015, RA in 2016, and SLE as well as scleroderma in 2017. She has been managed by conventional treatments (eg, gabapentin, prednisone, tramadol, tizanidine, and leflunomide) on and off for many years, achieving only intermittent alleviation of her pain, inflammation, and joint swelling (Table 1). Moreover, prolonged use of prednisone (at doses of 10–20 mg/d) and nonsteroidal anti-inflammatory drugs resulted in significant adverse events that now prevent the patient from safely tolerating the ongoing use of these agents.

Assessment

The patient presents with subjective complaints including pain and swelling of the hands, low back, hips, right knee, and feet, with exacerbations of low back and hip pain. The patient reports that the pain limits her ability to sit or walk. She reports enduring daily pain at work and a typical pain score of 7/8 out of 10. On an average of 2 out of every 20 work days, when the pain reached a 10 and her "feet were so swollen she couldn't wear any shoes or walk at all," she had to call in sick. Objective assessment indicated decreased range of motion in the cervical, thoracic, and lumbar spine; decreased range of motion and strength in shoulders bilaterally; and decreased strength of the right lower limb. With the exception of bilateral pedal edema, no other significant swelling was found. Laboratory evaluation revealed significantly elevated levels of the inflammatory biomarkers C-reactive protein (CRP) and erythrocyte sedimentation rate (ESR; Westergren method).

Management

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before treatment and on day 28. Confirmatory urine drug testing and blood analysis were performed on the final day of treatment by independent third-party laboratories (Quest Diagnostics and TriCore Laboratories, respectively).

Follow-Up

Significant improvement of pain and mobility-related symptoms was reported within 72 hours of treatment, reaching a maximum therapeutic effect by day 10. Symptoms related to mood (decreased anxiety, increased sense of well-being) continued to improve up to day 21 of treatment and remained increased until day 28. McGill Pain score decreased from 52 of 78 pretreatment to 25 of 78) on day 28 (Tables 2–4). SF-36 scores improved considerably across all 9 health domains (Table 4).

Pretreatment CRP and ESR values were 4.4 and

"Laboratory blood analysis demonstrated decreased inflammatory markers by day 28, further substantiating the patient's self-reported improvement from a biochemical perspective." —*Christian Shaw, MD, PhD*

Table 2. McGill Pain Questionnaire, Section 1:What Does Your Pain Feel Like?

Group #	Descriptor	Pre treatment	Day 28	Net difference
1	Temporal	4	1	3
2	Spatial	1	1	0
3	Punctate pressure	4	2	2
4	Incisive pressure	1	1	0
5	Constrictive pressure	4	2	2
6	Traction pressure	3	1	2
7	Thermal	2	1	1
8	Brightness	4	3	1
9	Dullness	4	1	3
10	Sensory, miscellaneous	4	1	3
11	Tension	3	1	2
12	Autonomic	1	1	0
13	Fear	2	1	1
14	Punishment	2	1	1
15	Affective-evaluative- sensory	2	1	1
16	Evaluative	3	1	2
17	Sensory, miscellaneous	3	1	2
18	Sensory, miscellaneous	2	2	0
19	Sensory	2	1	1
20	Affective-evaluative- sensory	2	1	1

Table 3. McGill Pain Questionnaire, Section 2: How Does Your Pain Change With Time?

Question	Pretreatment		Day 28	
	Response	Points	Response	Points
Which word or words would you use to describe the pattern of your pain?	Continuous, steady, constant	1	Brief, momentary, transient	3

Table 4. McGill Pain Questionnaire, Section 3: How Strong Is Your Pain?

Question	Pretreatment		Day 28	
	Response	Points	Response	Points
Which word describes pain right now?	Excruciating	5	Mild	1
Which word describes it at its worst?	Excruciating	5	Distressing	3
Which word describes it when it is least?	Discomforting	2	Mild	1



Figure 1. Laboratory testing result of the cannabidiol product.

% = % (w/w) = Percent (Weight of Analyte / Weight of Product)

* Total Cannabinoids results reflects the absolute sum of all cannabinoids detected.

** Total Potential THC/CBD is calculated using the following formulas to take into account the loss of a carboxyl group during decarboxylation step.

Total THC = THC + (THCa (0.877)) and Total CBD = CBD + (CBDa (0.877))

CBD, cannabidiol; **CBG**, cannabigerol; **THC**, delta-9-tetrahydrocannabinol.

For the complete laboratory report, please visit www.ajendomed.com.

Scale	Pretreatment, %	Day 28, %	
Physical functioning	15	50	
Role limitations due to physical health	0	75	
Role limitations due to emotional problems	0	67	
Energy/Fatigue	0	70	
Emotional well-being	36	76	
Social functioning	0	88	
Pain	23	90	
General health	15	15	
Health change	0	100	

Table 5. Scores on the SF-36

SF-36, 36-Item Short Form Survey 1.0.

48 mg/dL, respectively. At day 28, these values were 2.2 and 39 mg/dL, respectively. Adverse effects of treatment were mild and transient, and were limited to esophageal and stomach irritation after swallowing the CBD tincture.

Conclusion

Since completion of the 28-day CBD trial at the end of December 2018, the patient has been using nothing but CBD for her conditions with much success. Her CBD dose was titrated from 600 mg daily for 2 months, to 400 mg daily for 2 months, and 200 mg daily thereafter. The patient discontinued DMARDs 2 weeks prior to start of study and has not resumed any prescribed medications for rheumatic diseases since that time nor does she have any interest in doing so.

She no longer feels it necessary to see her

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rheumatologist. Notably, prior to participating in the CBD trial, the patient's rheumatologist intended to start her on a biologic due to her lack of response with conventional DMARDs.

This case demonstrates that a highly purified (99.9%) CBD isolate tincture of 600 mg daily was well tolerated and appeared highly effective in decreasing systemic inflammation while improving quality of life and pain scores on highly validated assessment tools. CBD did not appear to affect the kinetics of existing medications or lead to significant drug-drug interactions.

Discussion

An increasing number of reports and articles on individuals with RA using cannabis to treat their symptoms is available, although systematic studies regarding efficacy in conditions such as RA, and in patients facing multiple autoimmune conditions, are lacking.^{1,7-12} In this case study, the patient reported experiencing significant pain relief after 72 hours of high-dose CBD treatment. The patient reported greatly improved mobility and mood experienced by approximately day 10. Multidomain qualityof-life metrics reinforced the findings, indicating marked improvement between assessments taken pretreatment and on day 28 of treatment. Laboratory blood analysis demonstrated decreased inflammatory markers by day 28, further substantiating the patient's self-reported improvement from a biochemical perspective. Finally, confirmatory urine drug testing proved absent for any detectable tetrahydrocannabinol, a considerable finding within itself, as many patients suffering from inflammatory pain disorders are reluctant to use CBD products due to workplace drug testing concerns.13

Although this study is limited in its generalizability as an N=1 case report, the results are encouraging and highlight the need for future well-controlled clinical trials to investigate the efficacy of commercially available, federal and state regulatory-compliant CBD products as additional therapeutic options for inflammatory and autoimmune conditions.

Additionally, we call for the implementation of a publicly available database for cataloging clinical outcome data on commercially available and regulatory-compliant CBD products used for medical conditions. This would enable such information to be systematically mined for therapeutically relevant insights, especially in the absence of much needed evidence-based research, to guide clinical decisions on CBD and cannabinoid-based treatment options until the appropriate randomized control trials are completed.

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Dr. Shaw has no financial information to disclose.

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