Case 1

A two-year-old boy with an ependymoma was admitted for administration of chemotherapy (cyclophosphamide, methotrexate, etoposide and cisplatin). Previous cycles of the same chemotherapy had been well tolerated, with no hypotension. Independent of medical advice, the patient’s parents had been giving him Δ9-tetrahydrocannabinol and cannabidiol (THC–CBD) oil drops orally. Five days after his last chemotherapy dose, the patient was noted to be lethargic through the day, with downward-trending blood pressure readings. The patient’s blood pressure dropped to 50/30 mm Hg during deep sleep, and his skin appeared mottled, with delayed capillary refill. His heart rate was not substantially elevated, at 107 beats/min; his temperature, respiration rate and oxygen saturation levels were normal. Upon being roused, the child was upset but responsive, with normal findings on neurologic and cardiorespiratory examination. He had maintained good urine output and did not have clinical signs suggestive of dehydration.

The patient’s mother then reported that the family had independently increased the number of THC–CBD oil drops over the preceding few days from one drop three times daily to three drops three times daily. The last dose had been given 10–15 hours before the nadir in blood pressure.

Case 2

A four-year-old girl at very high risk of acute lymphoblastic leukemia was admitted for a bone marrow transplant. By day 30, she had full engraftment of transplanted marrow. During this period, she received treatment for suspected fungal lesions of her spleen but did not have fungemia. Despite frequent episodes of intermittent fever, the patient was never hypotensive.

On day 30, the patient’s mother administered a few drops of THC–CBD oil of unknown concentration to see if it would help alleviate symptoms of discomfort and pain. Nine hours later, while the patient was asleep, her diastolic blood pressure dropped from a baseline of 47 mm Hg to 33 mm Hg, and her systolic pressure dropped from 99 mm Hg to 80 mm Hg. She was febrile at 38.6°C and had a pulse of 120 beats/min. Over the preceding two days, the patient had been persistently febrile with normal blood pressure readings. She had been given empiric antibiotic and antifungal therapy, but blood culture results remained negative.

The patient was tired but alert and had a Glasgow Coma Scale score of 15. Her distal pulses were palpable but not bounding, and she did not appear clinically dehydrated.

The patient was given a rapid 20-mL/kg bolus of normal saline and maintenance fluid support, which corrected the hypotension. Empiric antibiotic therapy was started, although blood culture results were subsequently negative. The parents were insistent on continuing the THC–CBD oil but agreed to resume the lower dose. The patient’s blood pressure remained normal, even during subsequent periods of fever and sepsis.

Discussion

Patients with chronic or life-threatening illnesses have been using medical marijuana for symptom management and for possible anticancer activity.1,2 Cannabis has gained societal endorsement, with
legislation allowing for its use in a medical context, and commercial clinics and dispensaries available to the public. The medical community is becoming more accepting of medical marijuana use in outpatient and inpatient settings. However, physicians are often caught managing the medical, legal and psychosocial aspects of cannabis without the benefit of reliable safety data and dosing parameters.

More than 60 different cannabinoids have been isolated from cannabis, the most prominent of which include THC and CBD. Cannabinoids act to varying degrees on the CB1 and CB2 receptors found in the central and peripheral nervous systems. Native cannabis and cannabinoid derivatives may have effects on symptoms such as nausea, cachexia–anorexia, neurologic and cancer-related pain, chemosensory alterations and anxiety. Yet there is surprisingly little scientific information to guide cannabis use in medical practice.

The cannabinoid system has been shown to modulate cardiovascular physiology in animal models. In rats, intravenous administration of anandamide, an endogenous cannabinoid primarily acting on the CB1 receptor, has been found to have a triphasic effect on blood pressure, characterized by a transient drop followed by a brief rise and then a sustained period of hypotension. The human cardiovascular system appears to be affected by THC. A placebo-controlled trial showed that, among people who used cannabis regularly, systolic blood pressure dropped with an oral THC dose of 30 mg but increased at doses greater than 75 mg. A concomitant dose-dependent elevation in heart rate was noted. In a randomized trial involving adults with intractable cancer-related pain, hypotension was seen in 3 of 60 participants given a commercially available spray of THC–CBD extract (2.7 mg THC, 2.5 mg CBD) but in none of the participants in the THC-only and placebo groups.

The effects of cannabinoids in the pediatric population are not well described. In young children, the overarching symptom of cannabis toxicity appears to be somnolence or neurologic depression. Six cases of accidental marijuana ingestion by children seen in a children’s hospital resulted in coma, all reversible with supportive care. Of the six children, one 17-month-old child had documented hypotension. Other case reports have also described coma associated with cannabis toxicity in young children, in the absence of reported blood pressure changes. A systematic review of the effects of cannabinoids for the control of chemotherapy-induced nausea showed a relative risk for hypotension of 2.23 (95% confidence interval 1.75–2.83) in 30 randomized trials involving adults and children given cannabinoids (nabidolone, dronabinol or levonantradol).

In the cases we have described, causation cannot be assumed based on our observations, because both children were undergoing treatment for serious illness, and the illness itself, comorbidities and other medication could have contributed to the hypotension. Using the Naranjo Adverse Drug Reaction Probability Scale, we ranked the reaction in our two cases as “possible.” However, several measures of the scale, such as response to placebo and drug level measurement, were unknown in our patients and therefore could not contribute to the score. Another complicating variable is the unknown concentration and composition of the THC substance administered by the children’s parents. At present, these compounds are unregulated, often extracted and processed in facilities without consistent oversight to ensure product consistency. Non-THC ingredients may have contaminated the products, causing the adverse events.

We have reported these cases to the Canada Vigilance Program, which offers consumers, health professionals and drug producers an opportunity to report adverse effects of both regulated and unregulated health products (www.hc-sc.gc.ca/dhp-mps/medeff/databasdon/index-eng.php).

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