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Thrombotic microangiopathy associated with synthetic cannabinoid receptor agonists

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Abstract: Marijuana is one of the most commonly used recreational drugs in the United States. As marijuana is illegal in the majority of countries, the use of readily available and unregulated synthetic cannabinoids (SCBs) has increased. Little is known about the potential adverse effects of SCBs especially in regards to their nephrotoxicity. Case reports of acute kidney injury (AKI) from acute tubular injury secondary to their use have been reported. However, the exact pathology, mechanism, and extent of renal injury remain unknown. We report the first case of biopsy proven thrombotic microangiopathy (TMA) associated with SCBs resulting in AKI. The patient suffered significant morbidity with loss of renal function eventually requiring renal replacement therapy.

Keywords: Thrombotic microangiopathy (TMA); acute kidney injury (AKI); synthetic marijuana; synthetic cannabinoids (SCBs)

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Introduction

Synthetic cannabinoids (SCBs) are chemicals that produce several marijuana-like effects in humans. In recent years, they have become increasingly abused as they are cheap, easily available on the internet, and are not detected by routinely employed urine toxicology screening tests (1,2). One in nine high school seniors admitted using SCBs in 2011, making it the 2nd most prevalent illicit drug after marijuana (3). SCBs are dissolved in a volatile solvent and mixed with an assortment of plant leaves, such as Indian Warrior, Lion’s Ear, Dog Rose and/or Marshmallow leaves, which are themselves purported to have psychotropic effects upon smoking (4). Uneven distribution of SCBs to the plant mixture can result in drug “hot spots”, making dosing unpredictable and difficult, increasing the risk of overdose (5). There are commercially available, quasi-legal, unregulated “herbal incense” under names such as K2, K3, Spice, Smoke and Dream, just to name a few (6).

While usually labeled “not for human consumption” distributors and consumers understand that these products are to be used like marijuana primarily to attain a subjectively pleasant cannabimimetic psychotropic effects. This label has kept these products from being subjected to the Federal Analogue Act of 1986. This transfers all responsibility for the user’s safety from the manufacturers and distributors to the consumer, who is often unaware of the product’s potential danger (7). The most infamous of which are JWH- 018, JWH-073, JWH-200, CP-47, 497 and cannabicyclohexanol (7). Reports of acute kidney injury (AKI) attributed to it are on the rise but the underlying pathophysiology is elusive (8). Previous reports of AKI have shown diffuse patchy tubular injury favoring an ischemia related injury (8). We are reporting the first case of a young adult who presented with AKI after the use of SCBs and found to have biopsy proven thrombotic microangiopathy (TMA) as the major pathological pattern of the organ injury.
Case presentation

A 20-year-old man with no prior medical history presented to the emergency department after 2 episodes of generalized tonic-clonic seizures as witnessed by his mother. He complained of nausea and vomiting prior to the seizure. On arrival, he was conscious, then suffered from another 3-minute seizure episode. Intravenous levetiracetam 1,000 mg was given as initial treatment. Patient had no past history of seizures. His initial blood pressure was 176/88 mmHg and heart rate was 104/minute. Physical examination was otherwise unremarkable. He underwent computed tomography of his head, which was negative for bleeding or any other pathology.

Admission labs were consistent with anemia and AKI, with hemoglobin of 8.3 mg/dL, hematocrit of 27.4 mg/dL, and elevated serum creatinine of 3.2 mg/dL. He developed thrombocytopenia (platelets trended down from 162,000/mm³ on admission, to 147,000/mm³ on day 1 and 124,000/mm³ on day 2. On day 4 his serum lactate dehydrogenase levels were elevated at 309 U/L increasing to 530 U/L on day 10. Schistocytes were noted on peripheral smear and serum haptoglobin level low, less than 8 mg/dL, consistent with intravascular hemolysis. Initial urinalysis showed a specific gravity of 1.018, 3+ protein, 5–10 red blood cells per high power field, and RBC casts on urine microscopy. Urine protein to creatinine ratio was 5.06 on spot testing. A routine drug screen was positive for tetrahydrocannabinol. Serological investigation was negative for antinuclear antibody (ANA), anti-neutrophil cytoplasmic antibody (ANCA), anti-glomerular basement membrane (GBM) antibody, hepatitis screens, and HIV antibody screen.

Renal ultrasound was unremarkable. Since the patient admitted to daily use of SCBs over the prior few weeks, a sample was sent to the University of California in San Francisco for toxicology analysis and showed the presence of CP 47, CP 497 and Hebrew University (HU) 320. Using the “Criteria for Assessing Reports of Drug-Induced Thrombocytopenia and DITMA is important to diagnose as it is a preventable cause of TMA, generally managed first by withdrawing the suspected drug. The diagnosis of DITMA is made clinically with findings of MAHA and TMA associated with concurrent drug exposure. However, management of a patient with suspected DITMA may be challenging as DITMA and other primary thrombotic microangiopathies including TTP and complement-mediated TMA, have a similar clinical presentation but require different management modalities. A detailed clinical history is the key to making the correct diagnosis.

Recently published systematic review included 78 drugs known to cause TMA (9). We propose adding SCBs to this list. Individual agents identified in the reported case include CP 47, CP 497, and HU 320. Using the “Criteria for Assessing Reports of Drug-Induced Thrombocytopenia and
Levels of Evidence for a Causal Relation between the Drug and ‘Thrombocytopenia’ in our patient, a probable level of causality can be ascertained as described (10). These include that the suspected drug was taken before thrombocytopenia occurred and recovery from thrombocytopenia was complete and sustained after the drug was discontinued. Moreover, the suspected drug should be the only drug taken before the onset of thrombocytopenia, or other drugs were continued or reintroduced with a sustained normal platelet count. Additionally, to the extent possible with secondary documentation, all other potential causes of thrombocytopenia were reasonably excluded (10). This is further presented in Table 1.

The exact mechanism of drug induced TMA remains elusive but some reports suggest an idiosyncratic, acute immunologic reaction whereas others suggest a direct toxic effect, which may be either acute dose-related toxicity or chronic dose and duration-dependent toxicity (11). Some patients had severe kidney injury and were described as having HUS; others had minimal kidney injury. It is

<table>
<thead>
<tr>
<th>Mechanism</th>
<th>Pros</th>
<th>Cons</th>
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<tr>
<td>Complement dysfunction</td>
<td>Low suspicion for TTP (ADAMTS 67%); low suspicion for HUS (no diarrhea or GI symptoms)</td>
<td>No genetic testing available</td>
</tr>
<tr>
<td>Drug induced TMA</td>
<td>Low suspicion for TTP (ADAMTS 67%); low suspicion for HUS (no diarrhea or GI symptoms)</td>
<td>Definite criteria not met</td>
</tr>
</tbody>
</table>

Table 1 Pathogenesis of synthetic cannabinoid induced TMA

Figure 1 Renal and cardiac biopsy of patient. (A) Representative H&E stained image of a glomerulus and an adjacent arteriole showing thrombotic microangiopathy (3 prong arrow). Glomerulus displays focal mesangial expansion with increase in reticulated matrix material associated with partial endocapillary obliteration and focal double contouring of the capillary walls (arrow) (x100); (B) a trichrome stained section reveals marked tubular injury possibly secondary to marked vascular and microvascular thrombotic microangiopathy (x50); (C) representative electron microscopic image of a glomerular capillary showing mesangial expansion with increase in granulated mesangial matrix associated with circumferential mesangial interposition and double contouring of capillary walls (arrow) (x5,000); (D) representative cardiac biopsy showing marked edematous thickening and obliteration of 2 inter myocytic arterioles (thick arrows). A normally patent arteriole is also present (thin arrow mid bottom area).
possible that patients with severe HUS presentation may have an underlying complement dysregulation.

Inherited and acquired complement alternative pathway dysregulation, is related to various renal diseases by defective complement control, leading to the deposition of activated products, known as C3 glomerulopathies. Current modalities of treatment available for complement regulation include blockade with the anti-C5 monoclonal antibody eculizumab or by plasma substitution with success seen in several cases (12).

The agents included in SCBs may provide the second hit by unmasking the abnormality resulting in endothelial damage. Our patient failed to respond with plasmapheresis which makes it more likely that TMA was caused either by the direct toxic effect of these substances or by unmasking an underlying complement dysregulation.

There are various reports of kidney injury secondary to SCB related with acute tubular injury or allergic interstitial nephritis as the predominant clinical diagnosis, with no reported cases of biopsy proven TMA. There were four cases reported in which AKI was associated with synthetic cannabinoids with acute tubular injury as predominant pathology in renal biopsy (13). The Center for Disease Control and Prevention reported 16 cases of SCB induced AKI in multiple states with acute tubular injury demonstrated on 6 renal biopsies out of 8 performed, and acute interstitial nephritis found in 3 of 8 biopsies (14).

We are reporting for the first time SCB induced acute injury associated with biopsy proven TMA/vasculopathy in both kidney and heart. Arterioles and arteries of all sizes up to arcuate size were affected. In view of the marked vascular involvement including those of larger size, it was concluded that acute tubular injury was most likely secondary to severe vasculopathy. TMA may be more prevalent in SCBs but perhaps remain underreported due to the lack of renal biopsy in the majority of such cases.

Cannabinoid 1 and cannabinoid 2 receptors are present in almost all cell types in the kidney and play an important role in regulating normal renal physiology via the endocannabinoid system (15). Changes in these receptors are seen in pathological states like diabetic nephropathy, which causes the cannabinoid 2 receptor downregulation. Cannabinoid 1 receptor upregulation has been documented in IgA nephropathy, in acute interstitial fibrosis, and in mesangial fibrosis (16). One may postulate a cannabinoid receptor mediated injury to endothelial cells and/or podocytes, serves as an initial insult interfering with vascular endothelial growth factor receptor, leading to thrombosis and TMA.

**Conclusions**

The reported case represents the first biopsy-proven TMA secondary to SCB, not previously reported in English literature. We recommend adding SCB to the list of drugs associated with DITMAs. The implication that SCBs has deleterious effects, including DITMA necessitates further regulation on the manufacturing and selling of SCBs. Clinicians should be aware of these effects of SCBs as the drug has become more popular and these have not been reported with marijuana use. Further reporting and research of the SCBs and marijuana effects could better characterize SCBs association with renal injury.

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**Footnote**

*Conflicts of Interest:* The authors have no conflicts of interest to declare.

*Informed Consent:* Written informed consent was obtained from the patient for publication of this manuscript and any accompanying images.

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