ASRA Pain Medicine consensus guidelines on the management of the perioperative patient on cannabis and cannabinoids

Shalini Shah 1,1, Eric S Schwenk 2,2, Rakesh V Sondekoppam 2,2, Hance Clarke 4,4, Mark Zakowski 5, Rachel S Rzasa-Lynn 6, Brent Yeung 7, Kate Nicholson 8, Gary Schwartz 9,10, W Michael Hooten 11,11, Mark Wallace 12, Eugene R Viscusi 10,2, Samer Narouze 13

ABSTRACT

Background The past two decades have seen an increase in cannabis use due to both regulatory changes and an interest in potential therapeutic effects of the substance, yet many aspects of the substance and their health implications remain controversial or unclear.

Methods In November 2020, the American Society of Regional Anesthesia and Pain Medicine charged the Cannabis Working Group to develop guidelines for the perioperative use of cannabis. The Perioperative Use of Cannabis and Cannabinoids Guidelines Committee was charged with drafting responses to the nine key questions using a modified Delphi method with the overall goal of producing a document focused on the safe management of surgical patients using cannabinoids. A consensus recommendation required ≥75% agreement.

Results Nine questions were selected, with 100% consensus achieved on third-round voting. Topics addressed included perioperative screening, postponement of elective surgery, concomitant use of opioid and cannabis perioperatively, implications for parturients, adjustment in anesthetic and analgesics intraoperatively, postoperative monitoring, cannabis use disorder, and postoperative concerns. Surgical patients using cannabinoids are at potential increased risk for negative perioperative outcomes.

Conclusions Specific clinical recommendations for perioperative management of cannabis and cannabinoids were successfully created.

WHAT IS ALREADY KNOWN ON THIS TOPIC
⇒ There is a wide variability in consumption patterns among the US population (recreational, medicinal, frequent, infrequent) as well as in formulations (cannabidiol, tetrahydrocannabinol, or combination).
⇒ Although cannabis is reputed to have medicinal and psychoactive effects, its implications in the perioperative care setting are still largely unknown.

WHAT THIS STUDY ADDS
⇒ The guidelines offer critical education about terminology, pharmacology, and clinical implications associated with cannabinoid therapy in the perioperative period.

HOW THIS STUDY MIGHT AFFECT RESEARCH, PRACTICE OR POLICY
⇒ Identifying current evidence-based aspects and knowledge gaps of cannabinoid therapy should facilitate both clinicians and researchers in improving the care of patients in the perioperative period.

INTRODUCTION

This clinical practice guideline on the perioperative use of cannabis is designed to be a tool to help clinicians make evidence-based decisions regarding the perioperative management of patients who consume cannabis, who are presenting for surgery with increasing frequency. While many of the perioperative risks and challenges related to perioperative cannabis, such as how to advise patients preoperatively, the effects of cannabis on anesthetic medications, and the interaction between cannabis, opioids, and pain, have been described in the literature, there is no single document that summarizes all of these concerns and provides evidence-based recommendations. Flexibility in this clinical practice guideline is intended to enable person-centered decision-making that takes into account an individual’s expected health outcomes and well-being within the context of various regulatory environments. This document covers preoperative, intraoperative, and acute postoperative care considerations and several questions specifically focus on issues related to regional anesthesia and acute pain, which are the focus of the readers of this journal. The guidelines are not intended to limit or deny care nor affect the rights of patients or providers nor do they define standard of care. They are not intended to replace clinical judgment. In the imperfect setting of heterogeneous data, limited data, controversial topics, and bias inherent to expert opinion, compliance with the recommendations may not result in improved outcomes compared with personalized medicine. This guideline is also not intended to prompt the rapid tapering or discontinuation of cannabinoids or opioids for patients, nor is it intended to serve as a law, regulation, and/or policy
that dictates clinical practice or a substitute for US Food and Drug Administration (FDA)-approved labeling.1

BACKGROUND
Cannabis is the most commonly used psychotropic substance after alcohol and the most common recreational drug used in the USA with about 10% of the population (26 million people) reporting monthly use in 2017 according to the US Substance Abuse and Mental Health Services Administration (SAMHSA).2,3 To contextualize the risk of cannabis, SAMHSA approximates 1 in 10 people who use marijuana will become addicted; when they start before the age of 18, the rate of addiction rises to 1 in 6.4 Legalization and decriminalization of cannabis over the last decade has led to increased interest and literature on the subject, and has given rise to overt legal use and commercialization.5 With the rising prevalence of both medical and recreational cannabis use in the general population, anesthesiologists, surgeons, and perioperative physicians must have an understanding of the effects of cannabis on physiology in order to provide safe perioperative care. There is a critical need to summarize the existing cannabis literature in order to provide perioperative physicians and others with tools to address the challenges that arise from managing patients taking cannabis and cannabinoids.6

The purpose of this document is to provide background on cannabinoid terminology and relevant pharmacology as well as expert guidelines on perioperative management of patients who consume cannabis and cannabinoids in order to improve clinical care and future research and provide guidance to regulatory agencies to fully understand the impact of cannabis on anesthesia and pain care. This document covers preoperative, intraoperative, and acute post operative care considerations.

Definitions and terminology
Explanation of terminology will not only help clarify the subsequent discussions in this document but will assist physicians in their interactions with patients as well as understanding of the literature related to cannabis. Informed and accessible discussion with patients supports improved patient communication and trust and may contribute to improved outcomes.

A glossary of the most common and relevant terms for medical cannabis and related uses are presented in table 1.

Cannabinoid pharmacology
Cannabis plants include several species, the most common of which are Cannabis sativa, Cannabis indica, and Cannabis ruderalis. Cannabinoids are chemicals derived from cannabis (phytocannabinoids), such as cannabidiol (CBD) and ∆9-tetrahydrocannabinol (THC); synthetic medications, such as nabilone, dronabinol, nabilon; and endogenous cannabinoids that stimulate cannabinoid receptors, such as arachidonoyl ethanolamine (anandamide, AEA) or 2-arachidonoylglycerol (2-AG).

Table 1  Cannabis terminology*

<table>
<thead>
<tr>
<th>Term</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cannabis</td>
<td>All plant materials, components, and derivative products of the cannabis plant, including flowers, leaves, seeds, stalks, and other materials and cannabis resin, extractions, and other derivative products. Cannabis is listed in Schedule 1 of the Controlled Substances Act in the USA.</td>
</tr>
<tr>
<td>Marijuana, marihuana</td>
<td>Historical slang with Mexican roots adopted in the 1930s during the American prohibition efforts. Marijuana continues to be used interchangeably with cannabis in reference to plant strains containing high THC. Given the racial stigma, the word marijuana is becoming less used in favor of cannabis.</td>
</tr>
<tr>
<td>Hemp</td>
<td>Describes a collection of cannabis cultivars with specific properties, namely high production of fiber and seeds with minimal production of THC.</td>
</tr>
<tr>
<td>Cultivars (varieties, strains)</td>
<td>Distinct cultivars of the cannabis plant having unique genetic signature and expressing distinct chemical composition. Colloquially referred to as strains.</td>
</tr>
<tr>
<td>Cannabis extracts</td>
<td>Highly concentrated preparations of cannabis which are produced via a variety of manufacturing techniques.</td>
</tr>
<tr>
<td>Terpenes</td>
<td>Aromatic compounds that exist in unique profiles in different strains and may provide some therapeutic benefits.</td>
</tr>
<tr>
<td>Cannabinoid-based medicines</td>
<td>A general term used to describe therapeutic cannabis or cannabinoid-based products in which cannabinoids are the primary active pharmaceutical ingredient. This term is applied regardless of origin as plant-derived or synthetic cannabinoids.</td>
</tr>
<tr>
<td>Pharmaceutical or prescription cannabinoids</td>
<td>Cannabis-based treatments that have been approved as medical treatments for specific indications. Examples include nabilone (Cesamet), dronabinol (Marinol), cannabidiol (CBD; epiDiolex), and nabilonis (1:1 preparation of THC:CBD, eg. Sativex, not available in the USA).</td>
</tr>
<tr>
<td>Medical cannabis</td>
<td>Cannabis-based treatments that are not approved medical treatments but have been legalized and regulated for patient access. Medical cannabis is differentiated from non-medical cannabis by a unique access program and a required medical authorization.</td>
</tr>
<tr>
<td>Recreational cannabis use</td>
<td>Non-medical use for pleasure or leisure.</td>
</tr>
<tr>
<td>Recent cannabis use</td>
<td>Use within the past 30 days</td>
</tr>
<tr>
<td>Heavy cannabis use</td>
<td>Daily or near-daily use</td>
</tr>
<tr>
<td>Endocannabinoids</td>
<td>Endogenous cannabinoids produced by the body and active at cannabinoid receptors. The most well-known endocannabinoids are anandamide and 2-arachidonoylglycerol</td>
</tr>
<tr>
<td>Phytocannabinoids</td>
<td>Cannabinoids that are produced by the cannabis plant, primarily in the female flower. More than 100 unique cannabinoids have been identified. Common phytocannabinoids include ∆9-THC, CBD, cannabiol, and cannabigerol.</td>
</tr>
<tr>
<td>∆9-THC</td>
<td>THC is the primary cannabinoid in almost all varieties of cannabis. THC is the primary psychoactive agent and contributes the most therapeutic effects as well as adverse effects and intoxication of cannabis.</td>
</tr>
<tr>
<td>CBD</td>
<td>CBD is usually the other well-characterized cannabinoid found in cannabis. It has potential analgesic, anti-epileptic, anxiolytic, and anti-inflammatory properties, which inspired the selective breeding of cannabis strains with high concentrations of CBD and minimal THC concentration.</td>
</tr>
</tbody>
</table>

*Information extracted from references 307 308.

CBD, cannabidiol; ∆9-THC, ∆9-tetrahydrocannabinol.
**Δ9-tetrahydrocannabinol**

THC and CBD are among the most well-studied cannabinoids. THC was identified in 1940 while CBD was first identified in 1964. THC is the main psychoactive compound of the cannabis plant. THC concentration varies in different cannabis-based products ranging from 3% in marijuana to 80% in hashish oil.

**THC pharmacodynamics and mechanism of action**

THC acts as a partial agonist at the cannabinoid receptor 1 (CB1) and cannabinoid receptor 2 (CB2) and has high affinity at CB1 compared with CB2. Both CB1 and CB2 receptors are G-protein-coupled receptors. Activation of cannabinoid receptor inhibits adenylyl cyclase activity with subsequent reduction of intracellular cyclic adenosine monophosphate (cAMP) level or promotes mitogen-activated protein kinase activity. Decreased cAMP level leads to activation of voltage-gated potassium channels and inhibition of calcium channels, thus inhibiting neurotransmitters release.

Within the spinal cord, CB1 receptors have been localized to multiple areas involved in nociceptive processing including the superficial dorsal horn, the dorso-lateral funiculus, and lamina X. The activation of CB1 receptors in central nociceptive processing regions and primary afferents inhibits the release of neurotransmitters via decreased calcium conductance and increasing potassium conductance, which forms the possible anatomical basis for the analgesic action of cannabinoid agonists.

CB1 receptor activation mediates the psychoactive properties of cannabis, including changes in mood or consciousness, memory processing, and motor control. In animal studies, THC activates CB1 receptors. THC-induced dopamine release in the endocannabinoid system has been postulated as a potential mechanism of action for brain reward.

THC has many other non-CB1-receptor-mediated and CB2-receptor-mediated effects. Other receptors that THC modulates include G-protein receptor 55 and transient receptor potential cation channels (eg, transient receptor potential cation channel subfamily V members: TRPV1, TRPV2, TRPA1) and the serotonin receptors, 5-hydroxytryptamine (5-HT)

**THC pharmacokinetics**

Inhalation or smoking is the most common form of consumption. Other routes of administration include vaporization, oral spray, edibles, tinctures, other oral mucosal/sublingual routes (eg, capsules and lozenges), transdermal topical (cannabis-infused lotions and oils), and rectal routes.

Smoking cannabis produces rapid absorption, shorter duration of action and higher blood concentration of THC.

Absorption and bioavailability of THC depends on the type of smoking device, the depth of inhalation, puff duration, smoking habits (breath-holding), and the composition of cigarettes.

Vaporization offers a potential risk-reduction tool with a similar pharmacological profile as smoking.

In contrast to smoking as a method of delivery, oral absorption of cannabis is slow, variable, and highly dependent on the fat content of associated food ingestion. Based on bioavailability alone, the conversion factor between inhalation and oral absorption has been estimated at 2.5. Oral ingestion undergoes extensive hepatic first-pass metabolism with bioavailability of 10%–20%.

THC has high plasma protein binding and large volume of distribution. The plasma concentration of THC can follow two-compartment, three-compartment, or four-compartment models.

THC is metabolized in the liver via microsomal hydroxylation and oxidation by cytochrome P450 (CYP) enzymes CYP3A4, CYP2C9, and CYP2C19. THC is metabolized into an active form 11-OH-THC. Additional breakdown then results in 11-nor-9-carboxy-THC, the inactive metabolite.

**Cannabinoid**

CBD is a non-psychoactive phytocannabinoid and has been associated with analgesic, anti-inflammatory, anticonvulsant, anxiolytic, and antipsychotic effects. Although CBD has been shown to have potent anticonvulsant effects in humans, there are no studies supporting analgesic effects in humans. A recent study showed that 400 mg of CBD was not superior to placebo with respect to analgesia in patients presenting in the emergency room with acute low back pain. CBD may work synergistically with THC to produce its analgesic effect while decreasing psychoactive and cognitive side effects, such as sedation and memory impairment. CBD has proven to be well tolerated, showing low toxicity in several studies. Common side effects of CBD include somnolence, fatigue, and change in appetite and sleep pattern.

**CBD pharmacodynamics and mechanisms of action**

CBD has low affinity for both CB1 and CB2 receptors. It acts as a negative allosteric modulator of the CB1 receptor and as a weak inverse agonist of the CB2 receptor. CBD also interacts with other non-cannabinoid targets, including serotonin 1A receptors, vanilloid receptor 1 (TRPV1), and adenosine A2A receptors, all of which regulate the perception of pain. The inverse agonist activity at the CB2 receptor may explain the anti-inflammatory effects of CBD.

CBD may also act as an antagonist of the orphan receptor GPR 55. Moreover, CBD can act as an allosteric modulator of the μ- and δ-opioid receptors. The antipsychotic effects of CBD may be explained by enhancing anandamide signaling through inhibition of its reuptake and enzymatic deactivation.

**CBD pharmacokinetics**

CBD, like THC, undergoes extensive hepatic first-pass metabolism with per os (PO) consumption, with animal models demonstrating a range of 10%–13%. Only one study has reported the bioavailability of CBD in humans (31% following smoking), and half-life depends on dose and route of administration. CBD undergoes phase I and phase II metabolism. CBD is metabolized predominantly by CYP enzymes CYP3A4 and CYP2C19. Consequently, drugs that inhibit or induce these CYP enzymes would increase or decrease CBD levels, respectively. Phase II metabolism occurs through uridine 5’-diphospho-glucuronosyltransferase (UGT) 1A9 and 2B7.

Similar to that of THC, CBD plasma level decreases rapidly after smoking and follows a multiphasic pattern. The half-life of CBD has been estimated to be 27–35 hours after smoking or inhalation and 2–5 days after oral administration.

CBD is excreted both in urine and feces. Unlike THC, a large portion of CBD is excreted unchanged in the feces.

The most abundant metabolites are the inactive hydroxylated 7
(or 11)-carboxy derivatives of CBD, with the active 7 (or 11)-hydroxy CBD as a minor metabolite.\textsuperscript{31}

**Cannabinoid drug interactions**

The most clinically significant cannabinoid drug interactions are additive pharmacodynamic interactions when co-administered with other agents with similar physiological effects. In the presence of cannabinoids, sedation may be increased with administration of other central nervous system depressants, benzodiazepines, opioids, alcohol, and antihistamines, while tachycardia may increase with tricyclic stimulants, sympathomimetics, and antidepressants.\textsuperscript{32}

As noted, THC is metabolized predominantly by CYP3A4 and CYP2C9, while CBD is metabolized predominantly by CYP3A4 and CYP2C19. Therefore, drugs that inhibit or induce these CYP enzymes would increase or decrease THC and CBD levels, respectively.\textsuperscript{24, 48} Cannabinoids can competitively inhibit their own metabolizing enzymes, especially with sensitive substrates.\textsuperscript{53, 54}

Both THC and CBD may inhibit CYP3A4 and CYP2D6, with CBD having the most potent action. Moreover, CBD is a potent inhibitor of CYP1A1, CYP1A2, CYP1B1, CYP2B6, CYP2C8, CYP2C9, CYP2C19, CYP2D6, and CYP3A4.\textsuperscript{55-56} CBD is also a potent inhibitor of UGT1A9 and UGT2B7. Accordingly, plasma levels of UGT1A9 substrates, such as diflunisal, propofol, or fenofibrate and UGT2B7 substrates, such as gemfibrozil, lamotrigine, morphine, or lorazepam, may be increased when coadministered with CBD.\textsuperscript{57}

In addition to the pharmacodynamic synergistic effects that cannabis has with coadministered opioids, there are potential pharmacokinetic interactions as well.\textsuperscript{58, 59} In the perioperative setting, special attention should be paid to potential cannabinoid interactions with warfarin, direct oral anticoagulants, and clopidogrel (table 2).\textsuperscript{60-62} Common cannabinoid drug interactions are summarized in table 3.\textsuperscript{63-67}

**METHODS**

The initiation of these guidelines began with the submission of a proposal to the American Society of Regional Anesthesia and Pain Medicine (ASRA Pain Medicine) jointly by the cannabis and perioperative medicine special interest groups within the society on November 29, 2020. The proposal was reviewed and approved first by the Guidelines and Regulatory Advocacy Committee and subsequently by the Board of Directors. The members of the steering committee who created the proposal included SS, ESS, and SN. The steering committee selected potential writing committee members based on national reputation, publishing history on the topic of cannabis, and/or clinical expertise in managing patients taking cannabinoids. The Perioperative Use of Cannabis and Cannabinoids Guidelines Committee was charged with drafting responses to the key questions created by the steering committee with the overall goal of producing a document that focuses on the safe management of surgical patients taking cannabinoids. The ASRA Pain Medicine board and the steering committee provided clear directions that the committee’s objective was neither to endorse nor oppose cannabinoid use but rather to provide evidence-based guidelines that are practical for the average anesthesiologist or other perioperative physicians managing patients already taking cannabinoids.

Once the initial roster of the committee was determined, the initial meeting took place on January 5, 2021. Two members present at the initial meeting later notified the steering committee they would not be able to continue in their roles, and one was replaced. The steering committee assigned 2–3 members to each question based on expertise and availability. After initial written responses including recommendations with levels of certainty were submitted to the steering committee, they were edited, and a modified Delphi method was used in which all committee members voted on the recommendations via email with possible responses of ‘approve’, ‘approve with changes’, or ‘disapprove’. Individual responses were sent to the steering committee only and were not shared with other committee members. It was determined prior to voting that 75% agreement was needed to reach consensus for a given recommendation. After the first round of voting, it was determined that 2 of the question responses required substantial modification such that a second round of voting for those two questions was needed. Any changes suggested by committee members were considered by the steering committee and incorporated if agreed on. The results of each round of voting, including a summary of the narrative comments from committee members, are shown in online supplemental appendix table 6. The document was then approved by the ASRA Pain Medicine guidelines committee and subsequently the board of directors.

Literature searches for each question were performed using PubMed, Embase, and the Cochrane Database of Systematic Reviews. Reference lists of relevant publications were also searched. Specific search terms and strategies used were designed

| Table 2 | Cannabinoid drug Interactions with anticoagulants and antiplatelets used with permission from Samer Narouze, MD, PhD \textsuperscript{60-62} |
|---------------------------------------------------------------|
| **Cannabinoids drug interactions with anticoagulants and antiplatelets** |
| Drug | Effect | Intervention |
| Warfarin | THC and CBD can cause competitive inhibition of CYP2C9 and inhibit metabolism of the S-warfarin isomer, leading to supratherapeutic international normalized ratio levels. | ▶ Check INR within 3 days |
| DOACs (direct-acting oral anticoagulants) | CBD and possibly THC can increase DOACs level due to competitive inhibition of P-glycoproteins, and to a lesser extent CYP3A4. | ▶ Close monitoring |
| | | ▶ Consider using other anticoagulants or discontinue CBD/THC |
| Clopidogrel | CBD and possibly THC can increase clopidogrel level due to the competitive inhibition of CYP2C19. | ▶ Consider using another antiplatelet |
| Heparin/fondaparinux | No known interactions as these agents are processed by endothelial and renal cells and not metabolized by CYP enzymes, UGT, or P-glycoprotein. | |
| Platelets | Immune thrombocytopenia with synthetic cannabis. | ▶ Unlikely to have significant clinical effects |
| | | ▶ Immune thrombocytopenia is rare |
| CBD, cannabidiol; THC, tetrahydrocannabinol. | | |


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Table 3  Summary of cannabinoid-drug interactions used with permission from Samer Narouze, MD, PhD 63–67

<table>
<thead>
<tr>
<th>Enzyme</th>
<th>Drugs</th>
<th>Effects and interventions</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>CYP2C9</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CYP2C9 inducers</td>
<td>Antiarhythmics: amiodarone</td>
<td>▶ Decrease THC level</td>
</tr>
<tr>
<td></td>
<td>Anticonvulsants: valproic acid</td>
<td>▶ Unlikely to have significant effect on CBD</td>
</tr>
<tr>
<td></td>
<td>Antidepressants: fluoxetine</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Fluconazole, metronidazole, sulfamethoxazole</td>
<td></td>
</tr>
<tr>
<td>CYP2C9 inhibitors</td>
<td>Carbamazepine, rifampin</td>
<td>▶ Increase THC level</td>
</tr>
<tr>
<td></td>
<td>Unlikely to have significant effect on CBD</td>
<td></td>
</tr>
<tr>
<td>CYP2C9 substrates</td>
<td>Warfarin</td>
<td>▶ CBD and possibly THC may increase drug levels</td>
</tr>
<tr>
<td></td>
<td>Buprenorphine</td>
<td>▶ Decrease dose of substrate</td>
</tr>
<tr>
<td></td>
<td>Non-steroidal anti-inflammatory drugs (NSAIDs): celecoxib, naproxen, ibuprofen, naproxen.</td>
<td>▶ Monitor for toxicity and side effects</td>
</tr>
<tr>
<td></td>
<td>Anticonvulsants: phenobarbital, phenytoin</td>
<td>▶ Check INR within 3 days</td>
</tr>
<tr>
<td></td>
<td>Fluvastatin, rosiglitazone, rosuvastatin, sulfonylureas, losartan, valsartan</td>
<td></td>
</tr>
<tr>
<td><strong>CYP2C19</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CYP2C19 inducers</td>
<td>Anticonvulsants: carbamazepine, phenytoin, phenobarbital</td>
<td>▶ Decrease CBD and THC levels</td>
</tr>
<tr>
<td></td>
<td>Rifampin, rifampicin, ketoconazole, St. John’s wort</td>
<td></td>
</tr>
<tr>
<td>CYP2C19 inhibitors</td>
<td>Antidepressants: fluoxetine, fluvoxamine</td>
<td>▶ Increase CBD and THC levels</td>
</tr>
<tr>
<td></td>
<td>Chloramphenicol, felbamate, isoniazid</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Protease inhibitors</td>
<td></td>
</tr>
<tr>
<td>CYP2C19 substrates</td>
<td>Antidepressants: amitriptyline, citalopram, bupropion</td>
<td>▶ CBD and THC may increase drug levels</td>
</tr>
<tr>
<td></td>
<td>Anticonvulsants: clonazepam, diazepam, phenytoin, phenobarbital</td>
<td>▶ Decrease dose of substrate</td>
</tr>
<tr>
<td></td>
<td>Antipalelets: clopidogrel</td>
<td>▶ Monitor for toxicity</td>
</tr>
<tr>
<td></td>
<td>Proton pump inhibitors: omeprazole, pantoprazole</td>
<td>▶ Consider using alternative antipalelate instead of clopidogrel</td>
</tr>
<tr>
<td><strong>CYP2D6</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CYP2D6 Substrates</td>
<td>Opioids: codeine, morphine, hydrocodone, tramadol</td>
<td>▶ CBD&gt;THC may increase drug levels.</td>
</tr>
<tr>
<td></td>
<td>Anticonvulsants: valproate</td>
<td>▶ Decrease dose of substrate</td>
</tr>
<tr>
<td></td>
<td>Antidepressants: amitriptyline, citalopram, nortriptyline</td>
<td>▶ Monitor for toxicity and side effects</td>
</tr>
<tr>
<td></td>
<td>Antipsychotics: clozapine, haloperidol, risperidone</td>
<td>▶ Monitor for opioids augmentation</td>
</tr>
<tr>
<td></td>
<td>Antiarhythmics: amiodarone, dronedarone, flecainide, propafenone</td>
<td>▶ Monitor QTc for antidepressants and antiarrhythmics</td>
</tr>
<tr>
<td></td>
<td>β-blockers: carvedilol, metoprolol</td>
<td></td>
</tr>
<tr>
<td><strong>CYP3A4</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CYP3A4 Inducers</td>
<td>Anticonvulsants: carbamazepine, phenytoin, phenobarbital, topiramate</td>
<td>▶ Decrease CBD and THC levels</td>
</tr>
<tr>
<td></td>
<td>Cimetidine, pioglitazone, rifampin, St. John’s wort</td>
<td></td>
</tr>
<tr>
<td>CYP3A4 Inhibitors</td>
<td>Antiarhythmics: amiodarone, dronedarone, quinidine, diltiazem, verapamil</td>
<td>▶ Increase CBD and THC levels</td>
</tr>
<tr>
<td></td>
<td>Anticonvulsants: valproate</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Antifungals: ketoconazole, itraconazole, posaconazole</td>
<td></td>
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<tr>
<td></td>
<td>Macrolides: clarithromycin, erythromycin</td>
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<tr>
<td></td>
<td>Protease inhibitors</td>
<td></td>
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<tr>
<td></td>
<td>Tyrosine kinase inhibitors</td>
<td></td>
</tr>
<tr>
<td>CYP3A4 Substrates</td>
<td>Opioids: fentanyl, alfentanil, methadone</td>
<td>▶ Increase CBD and THC levels</td>
</tr>
<tr>
<td></td>
<td>Benzodiazepines: midazolam</td>
<td></td>
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<tr>
<td></td>
<td>Calcium channel blockers: amiodarone, felodipine</td>
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<tr>
<td></td>
<td>Calcineurin inhibitor: cyclosporine, tacrolimus</td>
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<td></td>
<td>PDE5 inhibitors: sildenafil</td>
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<tr>
<td></td>
<td>Propafenone</td>
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<tr>
<td></td>
<td>Statins</td>
<td></td>
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<tr>
<td></td>
<td>Zaleplon, zopiclone, zolpidem</td>
<td></td>
</tr>
<tr>
<td><strong>UGT1A9 (Phase II)</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>UGT1A9 Substrates</td>
<td>Analgesics/NSAIDs: acetaminophen, ibuprofen, difunlusal</td>
<td>▶ CBD increases substrate levels</td>
</tr>
<tr>
<td></td>
<td>Anesthetics: propofol</td>
<td>▶ Consider decreasing substrate dose</td>
</tr>
<tr>
<td></td>
<td>Anticonvulsants: valproate</td>
<td>▶ Monitor for side effects or toxicity</td>
</tr>
<tr>
<td></td>
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<td>DOACS: dabigatran</td>
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<td>Canagliflozin, dapagliflozin, irinotecan, mycophenolate mofetil, regorafenib, sorafenib</td>
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<td>Anticonvulsants: carbamazepine, valproate, lamotrigine</td>
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<td>Ezetimibe, losartan</td>
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p-glycoprotein (transport protein)  

Continued
by the working group assigned to each question. There were no limitations applied to the searches.

Recommendation grades assigned were based on the US Preventive Services Task Force (USPSTF) definitions of evidence strength, which specifies grade A-D or I for insufficient evidence to recommend. A level of certainty was also assigned to the statements (tables 4 and 5). This system has been used in previous ASRA guidelines documents and allows flexibility for high-grade recommendations to be made despite a lack of level 1 studies in the literature. The ASRA Pain Medicine Board of Directors reviewed and approved this document on August 8, 2022.

**Question 1: Should all surgical and procedural patients requiring anesthesia be screened for cannabinoids preoperatively and if so, what information should be obtained?**

The American Society of Anesthesiologists (ASA) standards for preanesthesia care state that before delivery of anesthetic care, the anesthesiologist is responsible for determining the medical status of the patient and developing a plan of anesthesia care, including ‘… the medical history, including previous anesthetic experiences and medical therapy.’ The ASA statement encompasses any and all medically relevant substances, including cannabinoids, regardless of whether they are medicinal, recreationally used, or illicitly obtained.

Cannabinoids are the most used addictive substances globally; therefore, physicians should screen for them during the preoperative evaluation. All patients should be questioned about cannabinoid use, dose and frequency, route of administration, and time of last use. A review of 28 studies with 65,720 participants by the USPSTF in 2020 found no evidence for benefits or harms from screening for drug use. However, this review did not specifically address perioperative screening where the implications of cannabinoid use may be more acute. Cannabinoids can produce significant physiologic changes and can potentially interact with anesthetics that can lead to complications. Drug screening by laboratory analysis of urine, saliva, blood, or hair generally detects only THC or carboxy-THC and is not recommended unless clinically indicated, such as for acute intoxication.

In addition to screening for cannabinoid use and verifying details as stated above, perioperative physicians should evaluate patients for acute intoxication. Practitioners should be aware of cannabinoid interactions with other medications, anesthetics, and physiologic changes. Literature has shown that acute cannabis intoxication could be detrimental in the perioperative period (see also sections questions 2, 3, and 8). It may be useful to use a standardized tool to screen preoperatively for cannabis use. The Cannabis Use Disorder Test has been validated and used to identify cannabis use disorder (CUD).

Recreational cannabis and cannabinoids have the potential to be mixed or laced with other substances, including pesticides, heavy metals, and carcinogens, which could have significant effects on perioperative outcomes.

A recent retrospective cohort study showed an association between chronic cannabinoid use and a 20% increase in incidence of postoperative nausea and vomiting (PONV). Because prophylaxis for PONV may be initiated preoperatively, screening for cannabinoid use can help identify patients at higher risk. Patients taking cannabinoids preoperatively may also report increased postoperative pain levels, which could affect perioperative management. This is discussed in more detail later in the document.

Unfortunately, few studies have addressed potential interactions between cannabinoids and anesthetic agents, and there is little consistency among the various preparations regarding dosage and CBD/THC ratio. A derivative of cannabis, Δ8-THC, is currently being produced and sold in the USA through a legal loophole. There is a paucity of research on this substance, but it has been known to produce psychoactive effects similar to...
the more well-known Δ9-THC.79 Until additional studies are performed, we recommend approaching this substance the same as one would manage Δ9-THC. The inconsistency and lack of regulation of the plethora of cannabinoid products being consumed makes perioperative management challenging.

Statement: Cannabinoids are the most commonly used recreational drugs in the USA and other countries, and the use of cannabinoids, both recreational and medicinal, may result in physiologic derangements. They may have interactions with other medications and treatments in the perioperative period.

Level of Certainty: Moderate

Recommendation 1: Universal screening for cannabinoids should be performed prior to surgery and should include type of cannabis or cannabinoid product, time of last consumption, route of administration, amount, and frequency of use. Grade A

Recommendation 2: Universal toxicology screening for cannabinoids is not currently indicated based on insufficient available evidence. Grade D

Question 2: What evidence exists to guide the decision to continue or stop cannabinoids perioperatively and/or postpone elective surgery?

As cannabinoids are increasingly used in the USA and the legal landscape continues to change and expand, anesthesiologists are increasingly being confronted with perioperative cannabinoid use and hence require guidance as to whether or not to continue cannabinoid products perioperatively as well as when to consider postponing elective surgery. The following sections summarize the effects of cannabinoids on the various organs systems with emphasis on the implications for perioperative management and additionally summarize the available guidance related to continuing or stopping cannabinoids perioperatively.

Cannabis use is associated with a dose-dependent impairment of cognitive function and performance. The ability of a patient to provide informed consent is one of the principal concerns when encountering an intoxicated patient perioperatively because of alterations in perception and memory function.74 80 Depending on the chronicity and quantity of use, a patient may or may not exhibit signs and symptoms of acute intoxication such as anxiety, paranoia, or frank psychosis. It is well known that cognitive and performance skills can be impaired in both recent/new users and chronic/heavy users.81 As demonstrated in a systematic review and meta-analysis involving 80 clinical trials,82 most driving-related cognitive skills recovered within approximately 5 hours (and almost all within 7 hours) of inhaling 20 mg of Δ9-THC. Oral Δ9-THC-induced impairment may take longer to resolve, and regular cannabinoid users experienced less impairment than ‘other’ (mostly occasional) cannabinoid users. Similar studies in perioperative patients are lacking. Still, the current evidence does support the return of some executive function by 5 hours after consumption. These data should be used in conjunction with clinical assessment of cognition and competence.

The cardiovascular effects of cannabinoids are biphasic. When used acutely and in low doses, there is an activation of sympathetic nervous system resulting in tachycardia and hypertension but at escalating doses and especially with chronic usage, bradycardia and hypotension can result from increased parasympathetic tone.83 In new users, the acute effects following cannabis smoking result in an increase in heart rate and blood pressure initially within the first 60 min from β-adrenergic stimulation and parasympathetic inhibition, followed by norepinephrine-mediated increases in heart rate and blood pressure for up to 120 min following cannabis use.84 These acute cannabinoid effects can impact perioperative hemodynamics and myocardial oxygen demand and may increase the risk of myocardial infarction (MI).84 85 While the evidence of increased risk of postoperative MI (OR of 1.88 (95% CI 1.31 to 2.69)) is known to exist in the surgical patient as shown in a recent cohort study,86 the period of increased risk since last use is not well characterized.

A multicenter case-crossover study of 3882 patients with MI revealed that 124 had smoked cannabis in the year preceding the MI.87 Of those 124 patients, 37 reported smoking it within 24 hours of MI onset and nine reported smoking within 1 hour of MI symptom onset. In addition to these nine patients, three patients reported smoking cannabis between 60 and 120 min before the onset of symptoms. Case-crossover analysis that controlled for differences between patients showed that within 1 hour after smoking cannabis, the risk of MI onset was elevated 4.8-fold (95% CI 2.9 to 9.5; p=0.001) compared with periods of nonuse. In the second hour after smoking cannabis, the relative risk was 1.7 (95% CI 0.6 to 5.1; p=0.34), suggesting a rapid decline in the cardiac effects after smoking cannabis.87 Cannabis smoking has also been found to reduce the anginal threshold by
48% after a single use as compared with 23% after smoking a nicotine cigarette in patients with chronic stable angina.88

A nationwide retrospective database study covering the period from 2010 to 2014 revealed that there was a 60% increase in-hospital mortality among marijuana users managed for acute MI.89 More recently, a systematic review and case descriptive analysis of cannabis use and acute MI revealed that the onset of acute MI symptoms is within 5 hours in the majority of cases.90 A systematic review of cannabis-induced cardiovascular effects, which included more than 4 million patients, showed that the most common events were tachycardia and various types of dysrhythmias, including atrial fibrillation, atrial flutter, atrioventricular block, and ventricular fibrillation. The researchers concluded that cannabis use is associated with rare but potentially life-threatening instances of cardiac dysrhythmia.91

Another systematic review, including 3695 individuals with toxicity from the use of synthetic cannabinoids, found that the most common cardiovascular effect was tachycardia in 30%. In addition, death occurred in 0.2%, stroke in 0.1%, and MI in 0.09%.92 Given the risk of prolonged cardiovascular effects, conservative recommendations suggest avoiding cannabinoids for 72 hours prior to surgery.93 94 While cannabinoid use does not appear to be a risk factor for coronary artery disease95–99 it may trigger acute coronary syndromes in those with underlying atherosclerotic heart disease.90 100–102

Smoking cannabis has been associated with a dose-dependent impairment of large airway function resulting in airflow obstruction and hyperinflation, but it has seldom been associated with frank emphysema.103–106 Of critical importance to the anesthesiologist are the effects that justify postponement of surgery. The limited available evidence suggests that the airway inflammation following marijuana smoking is similar to that following tobacco.107 It is not uncommon to encounter wheezing and productive cough suggestive of chronic bronchitis.108 Given the available evidence, it would be prudent to consider those who regularly smoke cannabis to be at similar risk for complications to those who smoke tobacco and a heavy/regular cannabis smoker to be at risk for chronic obstructive pulmonary disease (COPD).

Another airway concern in cannabis users is the possibility of airway inflammation, rhinopharyngitis, or uvular edema.109 110 All of which may contribute to airway obstruction in the perioperative period, especially with recent use.111

When considering whether to stop cannabinoids perioperatively, several factors come into play including:

► Medicinal versus recreational use.
► The dose, frequency and chronicity of use.
► The THC:CBD ratio of the product(s).
► The route of administration.

Some synthetic THCs, (eg, dronabinol), and synthetic analogs of delta-9-THC (eg, nabilone) as well as cannabis-derived (extracted directly from the plant, for example, CBD (Epidiolex) are FDA approved for chemotherapy-induced nausea/vomiting, HIV/AIDS-associated wasting, and rare forms of epilepsy (Lennox-Gastaut syndrome, Tuberous Sclerosis and Dravet’s syndrome), respectively. Abrupt cessation of these cannabinoid-based medications in the perioperative period for these patients is discouraged as such measures may adversely affect their perioperative course; hence, it is probably prudent to continue their use perioperatively until further safety data are available. On the other hand, the consensus panel believes recreational use should be discouraged similar to cigarette smoking.

CBD is generally well tolerated, and even with chronic usage and high doses (up to 1500 mg PO per day), CBD failed to produce any psychoactive, cardiovascular, cognitive, or psychomotor effects similar to Δ9-THC.112 113 Pure CBD containing product Epidiolex (100 mg/mL) is approved for certain forms of epilepsy and similar to other antiepileptic medications, should not be abruptly discontinued perioperatively. Safety of perioperatively continuing other FDA approved cannabinoid compounds such as dronabinol (synthetic THC compound), Nabiximol (Sativex 2.7 mg THC and 2.5 mg CBD per spray) and nabilone (Cesamet synthetic cannabinoid mimicking THC) is currently unknown. Similarly, safety data for other formulations containing CBD can be continued perioperatively, safety data are lacking and there is no standardization of THC:CBD ratios in cannabinoid-containing products. In a 2017 study published in JAMA, THC was detected in 21% of samples marketed as ‘pure CBD’ products, some at high concentrations.114

Despite the effects of cannabinoids on multiple organ systems, the scientific literature remains unclear as to whether to stop or continue medical cannabinoids preoperatively. A recent consensus-based guideline recommended reducing cannabinoid use 7 days prior to surgery (to less than 1.5 g/day of smoked cannabis, 300 mg/day of CBD oil, 20 mg/day of THC oil) while cautioning not to attempt any tapering strategies within 6 days of elective surgery and not to attempt tapering a day prior to surgery.115 Contrary to this recommendation, recent reviews of perioperative cannabinoid recommended cessation 72 hours prior to surgery.94 95 An even more conservative recommendation was recently provided, in which the authors recommended up to 10 days of cessation of oral cannabis consumption.116 The authors acknowledged the lack of evidence and their recommendation was based entirely on the half-lives of CBD and dronabinol and not on the physiological effects.

Given that cannabis smoking can negatively affect airway resistance, cardiovascular physiology, and cognition, anesthesiologists may consider discouraging cannabis use prior to surgery similar to tobacco smoking, unless its use is medically indicated. Any decision to cancel or postpone an elective surgery/procedure in a cannabis-consuming patient may be further directed by any clinical concerns about underlying cardiopulmonary problems or problems with mentation consequent to cannabis use. Given that this is an area of emerging evidence, further research is needed in order to provide evidence-based recommendations about the perioperative management of cannabinoids.

Statement 1: Acute effects of cannabis use can result in altered mental status and impairment of decision-making capacity. Hence, the frequency and the timing of the last dose of cannabis use are important. Level of Certainty: High

Statement 2: Smoking cannabis can cause increases in heart rate and blood pressure that is prominent within the first 1–2 hours of usage. Level of certainty: Moderate

Statement 3: Smoking cannabis may lead to a higher risk of perioperative acute MI within the first 1–2 hours. Level of certainty: Moderate

Statement 4: Smoking cannabis may have deleterious effects on airway resistance and respiratory adverse events. Level of certainty: Moderate

Statement 5: There is a lack of published data on the perioperative cardiovascular effects following other routes of cannabinoid administration. Level of certainty: Moderate

Recommendation 1: Patients should be counseled on the potential risks of continued perioperative cannabinoids. Grade B

Recommendation 2: We recommend postponing elective surgery in patients who have altered mental status or impairment of decision-making capacity due to acute cannabis intoxication. Grade A
Recommendation 3: We recommend delaying elective surgery for a minimum of 2 hours after cannabis smoking because of increased perioperative risk of acute MI. Grade C

Recommendation 4: With other cannabinoids routes (non-smoking) of administration, consider weighing the risks and benefits before proceeding with elective surgery given the temporal association of cannabis usage and adverse cardiovascular effects. There is a lack of published data to recommend a specific duration. Grade I

Question 3: For patients on concomitant cannabis and opioid use preoperatively, does existing evidence provide guidance on tapering of cannabinoids prior to surgery?

As the combination of cannabinoids and opioids is becoming more common in perioperative patients, an understanding of the effects on perioperative pain and risk of adverse events is essential. A comparison of data from 2012 to 2017 found that cannabinoid use increased more than 60% while opioid use decreased approximately 30%. The changes were not associated with changes in perioperative complications. Although some evidence suggests that cannabinoid use has improved opioid-related adverse events and reduced overall use, more substantial evidence demonstrates that cannabinoids can worsen pain and increase postoperative opioid use.

In a retrospective study of 71 patients undergoing primary unilateral total knee arthroplasty, preoperative exposure to cannabinoids did not affect short-term outcomes. However, in retrospective study involving 21,276 adults treated for traumatic injury, a higher rate of mechanical ventilation was observed. A recent prospective study on perioperative cannabis use compared 79 current cannabis users to 1,256 non-cannabis users undergoing elective surgery. A majority of current cannabis users were reportedly using cannabis medicinally, primarily for pain.

The results of this study showed higher levels of pain, poorer quality of life, and greater likelihood of using opioids or benzodiazepines in cannabis users compared with non-cannabis users prior to and 3 to 6 months following surgery.

Statement 1: Chronic use of THC may worsen postoperative pain, increase postoperative opioid use and precipitate the development of postoperative hyperalgesia. Level of certainty: Moderate

Statement 2: There is a lack of high-quality evidence describing the risks of concomitant opioids and cannabinoids in the perioperative period and in addition few studies have addressed the benefits and risks of preoperative cannabinoid tapering. We are uncertain of the overall benefit of preoperative cannabinoid tapering. Level of certainty: Low

Recommendation 1: We recommend that the frequent cannabis user be counseled on the potentially negative effects on postoperative pain control. Low-dose, medically supervised use likely has a lower risk of negative effects. Grade A

Recommendation 2: We cannot recommend for or against the routine tapering of cannabis and cannabinoids in the perioperative period. Grade I

Question 4: What are the specific concerns of chronic cannabinoid use in a parturient presenting for labor or cesarean section?

Cannabinoids are the most common addictive substances used by pregnant women and use in this patient population has been increasing over the last few decades. The 2019 SAMHSA survey reported 5.4% of all pregnant women using marijuana and 1.7% of all pregnant women using marijuana daily or almost daily in the USA, increasing to over 8% with screening. The cannabinoid system is prevalent throughout the human body, with receptors in maternal, placental and fetal tissues.

Cannabinoids affect both maternal and fetal physiology, cross the placenta and have been associated with preterm delivery, lower birth weight, and other adverse outcomes. Thus, the anesthesiologist must be familiar with the physiologic effects and interactions to understand the usage and safety concerns of cannabinoids during pregnancy and the puerperium.

Placental and fetal effects

Cannabinoids readily cross the placenta. THC crosses the placenta and fetal levels are about 10% of maternal levels and higher with chronic exposure in animal studies. Prenatal exposure to THC has been associated with decreased birth weight, decreased pancreatic islet density, and glucose intolerance after birth in rats. The endocannabinoid system is present in the myometrium and placenta, with changes reported in disease states including endometriosis and preeclampsia.

CB1 and CB2 receptors are present in the uterus and the placenta. Also present are other cannabinoid system components, including fatty acid amide hydrolase (FAAH), N-acylethanolamide-specific phospholipase D (NAPE-PLD), and endocannabinoid agonists AEA. Indeed, plasma AEA levels may decrease from first to second to third trimester but increase dramatically during labor.

The endocannabinoid system appears to be involved in placental development and trophoblast proliferation, with CB1 receptor knockout mice having smaller placentas. In addition, CB1 receptor levels are higher in the placental tissue of preeclamptic women.

Endocannabinoids affect normal fetal brain development, neuron proliferation, differentiation, and neurotransmitter levels.

Human fetal central nervous system CB1 receptors are present at 14 weeks of gestation, and the number of receptors increases with increasing gestational age. Cannabinoids are lipophilic, easily crossing the placenta, resulting in exposure to THC, CBD, or other components of marijuana. Prenatal marijuana exposure has been associated with decreased problem solving, visual-motor coordination, visual analysis, decreased attention span and behavior problems in offspring. Visual evoked potentials have been shown to be delayed at 18 months of age, while subsequent neuroimaging of young adults aged 18–22 show functional MRI scan differences with increased effort required for executive function tasks.

Prenatal cannabis exposure has been associated with adolescent vulnerabilities to psychopathology, sleep problems, lower cognition and lower gray matter volume, providing more evidence of long term effects in the offspring.

Maternal effects

The use of recreational marijuana for nausea or vomiting during pregnancy almost doubled to 11% from 2009 to 2016 in one California study. First trimester use of cannabis was associated with symptoms ranging from mild and severe nausea or vomiting during pregnancy. Cannabinoids (cannabis, Δ9-THC, nabilone, levonantradol and nonabine) activate CB1 receptors in the brainstem and enteric nervous system and may act as antiemetics.

High levels of cannabis consumption may cause an induced hyperemesis syndrome causing some to seek hot baths as heat activates vanilloid receptor type 1 and releases substance P, mediating nausea. A review found evidence that cannabis helps


Original research
Pre-eclampsia

Pre-eclampsia spectrum, a hypertensive disorder of pregnancy, occurs in 5%–8% of pregnancies. Cannabinoids may be involved in preeclampsia, as changes in the endocannabinoid system in preeclampsia have been noted, including a high number of CB1 receptors in the placenta, an increase in NAPE-PLD, a decrease of FAAH, and decrease in plasma AEA. Endocannabinoids influence nitric oxide production, which affect regulation of placental blood flow. FAAH blockade, which increases anandamide, led to a decreased response to angiotensin II contraction in normal and pre-eclamptic model mice, while monoacylglycerol lipase blockade, which increases 2-AG, reduced response in pre-eclamptic mice. CB1 receptor blockade had no effect on the angiotensin response in this model. The effect of exogenous cannabinoids on the hypertensive response or interaction with clinical preeclampsia in humans is currently unknown. An increase in blood pressure may be seen following acute cessation of cannabis in chronic, high-quantity users but not occasional users. In one retrospective multivariate analysis adjusted for use of other substances, there was no difference in the overall distribution of hypertensive disorders in women who used marijuana.

Breast feeding

While the American College of Obstetrics and Gynecologists (ACOG) and the FDA advise against consuming THC, CBD, and marijuana while breast feeding, some have suggested that the benefits of breast feeding may outweigh the effects of cannabinoids in breast milk. There are large variations in the concentration of cannabinoids in breast milk, but they may be present for a prolonged period after maternal consumption. Maternal cannabis use in the early postnatal period was positive in 5% of the population surveyed in one study. THC can be found in breast milk for up to 6 days after maternal consumption, although it may be concentrated up to sevenfold compared with maternal plasma in chronic consumption. The relative infant dose, the amount absorbed, has been estimated to be 2.5% (range 0.4%–8.7%) of the maternal dose. One study found the human milk:plasma partition coefficient of 6:1 with a median THC value of 3.2 ng/mL in breast milk. The half-life of THC in breast milk is up to 17 days. Large variations in the concentration of THC in breastmilk have been noted with a peak of up to 420 ng/mL.

Predelivery anesthetic considerations

Long-term, high-quantity cannabis users may benefit from a predelivery high-risk anesthesia clinic evaluation to assess potential interactions and improve outcomes with development of a multidisciplinary plan. Prenatal marijuana use adversely affects fetal brain development and subsequent behavioral self-regulation, a precursor to later, more serious problems in childhood. Yet, many mothers continue to use cannabis and believe in its safety. Stopping marijuana use before 10 weeks of gestational age may prevent these effects and higher maternal cholesterol levels seem to mitigate some of marijuana’s adverse effects on the fetus. A majority of pregnant women had poor knowledge about the risks of marijuana use during pregnancy, and 90% were more likely to use marijuana in pregnancy if it were legal.

Intrapartum anesthetic considerations

A history of occasional or recreational use of marijuana likely does not pose a risk with neuraxial anesthesia for labor analgesia or cesarean delivery. A parturient admitted with acute cannabis intoxication but without long-term, high-quantity use may be susceptible to interactions based on the physiology discussed in other sections of these guidelines. Within a 2-hour window from consumption, norepinephrine levels may be increased with potential for cardiovascular, anesthetic and vasopressor interactions. The potential for acute cannabis intoxication might reduce the amount of opioid medication needed but data are limited to animal studies.

Long-term, high-quantity use may be associated with increased parasympathetic tone, decreased heart rate, and postural hypotension. Short-term, high-quantity use may be associated with thermoregulatory changes, hypothermia, and shivering, which may be worsened with vasodilation of regional anesthesia, general anesthesia, or change in temperature set point by neuraxial opioids.

Strong CB1 agonism (or synthetic cannabinoid, eg, ‘K2’/‘spice’) may lead to or be associated with preeclampsia use may cause or be associated with symptoms resembling preeclampsia, cerebral ischemia, or PRES. Long-term cannabinoid exposure may cause a cross tolerance to opioids and require greater use of opioids. Acute intoxication with cannabinoids may augment the analgesia of mu and kappa opioid agonists, thus potentially reducing the dosage requirement for opioids.

Postoperative analgesic regimens should maximize non-opioid analgesics, use a multi-modal enhanced recovery protocol, and avoid cannabis as an adjunct due to concerns over cannabinoid passage via breastmilk to the neonate.

Summary of considerations of cannabinoid use in pregnancy and during the peripartum period

Anesthesiologists should be aware of the increasing use of cannabis and cannabinoids preconception, during pregnancy, and in the postpartum period. Cannabis or cannabinoids cannot be recommended during labor, cesarean delivery, or in the immediate postpartum period at this time, and the FDA and ACOG recommend avoiding cannabis/cannabinoids during pregnancy.
and breast feeding. The effects of cannabis use during pregnancy may include an increase in the odds of anemia, low birth weight, premature birth, need for neonatal intensive care unit services, and altered brain development. 137 178 THC and other cannabinoids enter human breast milk and may further impact neonatal development. Pregnant patients should be educated about the risks of maternal cannabis use on the fetus/neonate. 175 178 179

Statement 1: While cannabis use during pregnancy and in the postpartum period has the potential for adverse maternal and fetal physiological complications, there is currently no evidence to suggest that there are any specific implications with neuromuscular blockade and analgesia for labor or cesarean section. Level of certainty: Moderate

Recommendation 1: Pregnant patients should be educated and counseled about the risks of maternal cannabis use on the fetus/neonate. Grade A

Recommendation 2: Cannabis use during pregnancy and immediate postpartum period should be discouraged. Grade B

Question 5: Should the intraoperative doses of anesthetics and analgesics be adjusted in patients who have taken cannabinoids preoperatively?

A key issue for anesthesiologists caring for patients who have short-term or long-term exposure to cannabinoids is to determine what adjustments, if any, are needed with respect to doses of routine perioperative medications. Evidence on this topic is derived from a disparate group of human studies with significant limitations and data from preclinical animal studies. In an animal study of rats administered sevoflurane and various doses of CBD with or without morphine, the addition of CBD did not enhance the minimum alveolar concentration (MAC) of sevoflurane. 180 These results stand in contrast to those of other researchers, such as Müller et al., 181 who studied the effects of adding THC to sevoflurane in 38 rats using a blinded protocol and reported that MAC was reduced by 26%. Stoelting et al. 182 in a 1973 study reported that THC decreased the MAC of halothane in dogs. However, the translation of the study findings to humans was uncertain because of species differences in THC sensitivity. Another animal study from 1973 reported that THC lowered the MAC of cyclopropane, which is a volatile anesthetic no longer used clinically. 183

Human studies are limited in number and quality. A retrospective analysis of 118 patients who were undergoing tibial fracture repair reported that the MAC requirements of sevoflurane in cannabis users was greater compared with non-cannabis users. 184 Cannabis use was defined as any prior use in the month before surgery. However, there was no difference observed in desflurane and propofol MAC between groups and an accompanying editorial cautioned about making changes to practice based on a study with inconsistent and limited findings. 185 In a retrospective study that involved a random sample of 250 endoscopy patients, 25 were identified as cannabis users. 186 The authors reported that greater midazolam, fentanyl, and propofol doses were administered to individuals in the cannabis group but multiple confounders were identified, including procedure duration and the potential of individual provider differences in administered dosages. In addition, the absolute difference in drug doses between groups was small and of uncertain clinical significance. Finally, in a randomized, single-blinded study, Flisberg et al. 187 reported that regular cannabis users required greater propofol doses for successful laryngeal mask airway insertion than nonusers (314.0 mg ± 109.3 vs 263.2 ± 69.5 mg, p < 0.04). The timing of cannabis use in relation to the time of anesthesia induction was unclear and requires clarification in future prospective trials.

A final issue relates to the use of intraoperative anesthetic depth monitors, such as the bispectral index (BIS), in patients taking cannabinoids preoperatively. In a double-blind, randomized controlled trial (RCT) 27 patients undergoing elective orthopedic surgery under general anesthesia were allocated to 1 of 4 interventions: (1) high-dose cannabis, (2) low-dose cannabis, (3) active placebo, or (4) placebo. 188 The form of cannabis administered was nabiximols, which is administered as an oral spray and is not available in the USA. In this study, nabiximols was administered 20 min prior to induction of general anesthesia. The authors reported that the average BIS values were higher in the high-dose cannabis group but presented no data to suggest that higher BIS readings indicated ‘lighter’ anesthetic depth. Other human studies have found altered gamma neural oscillations in cannabis users, but it remains unclear how those changes would affect anesthetic management. 189 190

Finally, in their consensus recommendations on the perioperative management of cannabis, Ladha et al. 115 suggested giving ‘extra consideration to greater depth of anesthesia during induction and maintenance of anesthesia’ in cannabis users. They also stated that acutely intoxicated cannabis users might have lower anesthetic requirements and recommended the use of intraoperative electroencephalogram (EEG) monitoring. However, there is a lack of evidence about the reliability and clinical utility of processed EEG monitoring in cannabis users.

Statement: In light of the overall weak quality of evidence and absence of RCTs, the effect of preoperative cannabis use needs to be investigated further but the limited evidence suggests that depending on the timing of last cannabis consumption it may have an effect on lowering anesthetic requirements in the acutely intoxicated user and increasing anesthetic requirements in the long-term regular user (not acutely intoxicated). Level of certainty: Low

Recommendation 1: Consideration should be given to adjusting induction and maintenance doses of anesthetic agents based on clinical presentation and timing of the last consumption of cannabis in surgical and procedural patients. Grade C

Recommendation 2: There is insufficient evidence to recommend for or against the use of intraoperative EEG monitoring in patients who have taken cannabinoids. Grade I

Question 6: Does acute or chronic cannabis exposure require any adjustment of ventilator settings to accommodate for possible V/Q mismatch, smoke inhalation injury, or other lung pathology?

Long-term cannabis exposure

Anesthesiologists caring for patients using cannabis-containing products long-term must be aware of alterations in pulmonary function and respiratory physiology that may affect ventilation intraoperatively and postoperatively. While not as well studied as tobacco smoking, cannabis smoking has drawn comparisons to tobacco smoking. 103 For example, cannabis smoke contains many of the same chemical and particulate components found in tobacco smoke that are known to damage lung tissues. 103 Cannabis smoke contains greater levels than tobacco smoke of several toxic substances including acetaldehyde, hydrogen cyanide, and nitrogen oxides. 191 192 Among patients with chronic COPD, initiating use of prescription oral cannabinoids (nabiximols, dronabinol) was not associated with increased hospitalization, emergency department or outpatient visits, or...
diagnosis of pneumonia, but all-cause mortality was greater among new users of prescription cannabinoids. Alternatively, individuals receiving higher-dose cannabinoids compared with controls experienced increased rates of hospitalization for COPD symptom exacerbation and pneumonia. In a prospective study, 339 subjects were allocated to 1 of 4 groups: (1) cannabis smoking only, (2) tobacco smoking only, (3) combined cannabis and tobacco smoking, and (4) non-smokers. Cannabis smoking was associated with reductions in the forced expiratory volume in 1 s (FEV₁) to forced vital capacity (FVC) ratio and increased lung capacity. Furthermore, one smoked cannabis cigarette had the respiratory effect of 2.5–5 tobacco cigarettes, and overall pathology was worse when cannabis was combined with tobacco. These findings suggest that smoking cannabis is associated with development of an obstructive respiratory pathology. Longitudinal and cross-sectional studies have demonstrated that long-term cannabis smoking leads to chronic bronchitis and airflow obstruction involving large airways. Symptoms including coughing, excessive sputum production, and wheezing have been associated with cannabis use. In a population-based study involving 1037 young adults, smoking cannabis at least once per week was also associated with increased incidence of morning cough, sputum production, and wheezing. When these patients discontinued smoking cannabis, symptoms resolved. Cannabis vaping is also associated with increased likelihood of chronic bronchitis symptoms compared with never vaping cannabis, even when adjusting for nicotine vaping, cannabis smoking, tobacco smoking, and sociodemographic factors. These findings are not, however, universal, as several studies have found no or limited associations between cannabis use and pulmonary symptoms.

Study of the pulmonary effects of regular cannabis smoking is complicated by the high prevalence of concomitant tobacco use, which is inconsistently controlled for in participant recruitment and statistical analyses. A prospective observational study evaluating the effects of marijuana use on lung function found that while there was no overall association between pulmonary symptoms and cannabis smoking, the subset of patients with a 10-year history of smoking filter-less cannabis cigarettes did report some symptoms. Among young, intermittent smokers, cannabis co-use with intermittent tobacco smoking was more strongly associated with the presence of respiratory symptoms than use of either substance alone. A similar association was found in an observational study of older participants, in whom tobacco use, alone or combined with cannabis, was associated with respiratory symptoms but no significant association was found for cannabis use alone.

A retrospective database analysis of 8932 patients with CUD and an equal number of matched controls found that regular cannabis use was associated with a significantly greater risk of asthma, COPD, and pneumonia diagnoses. Patients who also used tobacco had the greatest prevalence of these diagnoses, but a higher prevalence of all three was associated with tobacco use disorder alone compared with isolated CUD.

Multiple studies have found reduced FEV₁/FVC among people who smoke cannabis. This finding may be due to increases in FVC with preserved or smaller increases in FEV₁ rather than reductions in FEV₁ due to obstruction, as several studies have found an increase in FVC among cannabis smokers. This also may reflect inadequate control for the effects of tobacco use in some studies. It is notable that several studies that found no significant reduction in FEV₁ did observe greater airway resistance among cannabis smokers compared with non-smokers. Like tobacco use, recent cannabis use was associated with reduced exhaled nitric oxide, which may contribute to impaired bronchodilation.

Cannabis smoking has been associated with the formation of lung bullae in relatively young patients in several case series and multiple case reports, but it is unclear if this is a causative association. Cannabis smoking has also been associated with pneumothorax, and combining cannabis use with tobacco use may increase this risk. However, in general, cannabis use does not appear to be associated with emphysema.

Short-term cannabis exposure
Less is known about the effects of acute cannabis exposure on lung function compared with chronic exposure. The immediate effects of oral or inhaled THC induce bronchodilation both in healthy subjects and those with asthma, but aerosolized THC or cannabis smoking may also result in an irritation of the airways and bronchoconstriction in some individuals with reactive airway disease. Evidence of harm is limited to small case series and case reports. A small clinical trial and several case reports of patients with acute shortness of breath and hemoptysis within hours of cannabis use have been published. Alveolar hemorrhage was also reported in another case report. Several cases of cannabis use-related lung injury have also been documented and associated with vaporized cannabis products attributed to additives or contaminants such as vitamin E acetate. The resulting lung injury has been defined as e-cigarette vapor associated lung injury (EVALI) which results in severe lung pathology such as acute eosinophilic pneumonia, diffuse alveolar hemorrhage, lipid pneumonia, and respiratory- bronchiolitis interstitial lung disease. Statement 1: There is low-quality evidence that patients taking only oral cannabinoids do not experience significant changes in pulmonary function. Level of certainty: Low

Statement 2: There is conflicting evidence as to whether any ventilatory changes should be made for patients with chronic or acute cannabis exposure via inhalation. Acute cannabis inhalation may result in bronchodilation but may also cause airway irritation and bronchoconstriction in susceptible individuals. Long-term use of inhaled cannabis is likely associated with the development of obstructive lung disease-like patterns such as chronic bronchitis. Level of Certainty: Low

Recommendation 1: Based on the studies reviewed, patients taking only oral cannabis do not need any adjustments in ventilatory settings. Grade C

Recommendation 2: Adjustment of ventilatory settings should be considered since obstructive lung disease-like patterns may be associated with chronic cannabis consumption by inhalation, particularly in patients with comorbid conditions that are associated with an increased risk of pulmonary pathology. Grade C

Recommendation 3: Evidence is insufficient to guide ventilation settings following acute cannabis use via inhalation. Grade I

Question 7: Do patients taking perioperative cannabinoids require any special postoperative considerations? If so, for how long?

The immediate effects of THC administration in humans results in dose-related tachycardia and increases in cardiac index. In the acute postoperative phase of care, MI, arrhythmias, stroke, cardiac arrest, and cardiomyopathy have been reported. A retrospective cohort analysis evaluating perioperative outcomes in major elective surgeries and CUD demonstrated an adjusted OR of postoperative MI of 1.88 (95% CI 1.31 to 2.69, p<0.001) for patients with a reported active CUD compared with those

without. Mittleman et al interviewed 3882 patients with acute MI and conducted a case-crossover analysis that controlled for differences among patients, and found that within 1 hour after smoking marijuana, patients’ risk of MI onset was elevated 4.8-fold (95% CI 2.9 to 9.5; p<0.001) compared with periods of nonuse. However, in the second hour after smoking, the relative risk was 1.7 (95% CI 0.6 to 5.1; p=0.34), suggesting a rapid decline in the cardiac effects of marijuana. The supply-demand mismatch in oxygen coupled with the increased cardiovascular oxygen demands as a result of hypertension and tachycardia may explain the higher perioperative risk of acute MI among patients with CUD. Cannabis may induce hypothermia which has the potential to aggravate postoperative shivering. Shivering will adversely affect the supply-demand mismatch in oxygen and may further increase the odds of postoperative MI. In another study the authors also demonstrated that active CUD may be associated with higher adjusted odds of suffering a postoperative acute cerebrovascular event.

In a retrospective cohort of 510,007 patients reviewed between 2006 and 2015, CUD was also associated with a significantly higher incidence of acute MI (p=0.001) and perioperative stroke with vascular surgery (p=0.031). Those with CUD had a higher incidence of perioperative MI (3.3% vs 2.1%; OR 1.56; 95% CI 1.09 to 2.24; p=0.016) and perioperative stroke (5.5% vs 3.5%; OR 1.59; 95% CI 1.20 to 2.12; p=0.0013) than patients without CUD. In another large retrospective cohort analysis of patients with CUD undergoing major elective surgeries, the adjusted odds of perioperative MI was 1.88 (95% CI 1.31 to 2.69, p<0.001) times higher for patients with a reported active CUD (89 of 13,603; 0.7%) compared with those without (46 of 13,603; 0.3%) an active CUD (unadjusted OR 2.88; 95% CI 2.34 to 3.55; p<0.001). Owing to limitations in administrative data, it is unclear if this represents a true effect or selection bias; however, these findings do warrant further investigation. Calapai et al demonstrated several findings that indicate that CBD can modify the deleterious effects on the blood–brain barrier (BBB) caused by inflammatory cytokines and may play a pivotal role in ameliorating BBB dysfunction consequent to ischemia. In a retrospective evaluation of the Personality and Total Health Through Life study (n=2404), Hemachandra et al found a 3.3-fold risk of shock/transient ischemic attack in cannabis users within the past year, but this elevated risk was specific only to participants who used cannabis at least weekly. Patients using perioperative cannabinoids also appear to have a 3.24-fold increased risk of developing hypotension postoperatively (95% CI 1.12 TO 9.36, p=0.03).

Gastrointestinal system

Cannabis consumption has been associated with poor oral hygiene, caries, and periodontal disease, which may have implications for airway management. Cannabis consumption is also linked to delayed gastric emptying, delayed intestinal motility, cyclical vomiting, cannabis hyperemesis syndrome and in surgical patients may also increase the risk of PONV. The endocannabinoid system is involved in hepatic homeostasis with derangements in this system being implicated in hepatic steatosis, fibrogenesis, and hepatic cirrhosis. Apart from its effect on CB1 and CB2 receptors, cannabis can exert its effect on the gut vanilloid receptors resulting in mesenteric vasodilation.

Question 8: Are there special considerations for concomitant opioid and cannabinoid use and should postoperative opioid prescriptions be adjusted prior to discharge?

Chronic pain is one of the most common indications for medical cannabis prescriptions. However, not all patients who use cannabis for pain relief do so with the guidance of a knowledgeable clinician and therefore may use doses and formulations that lead to greater adverse effects, including alterations in pain and responses to opioids. Furthermore, it may be difficult to ascertain

Postoperative analgesia and hyperalgesia

As explained elsewhere in this guideline, there is growing evidence suggesting that long-term, frequent and infrequent marijuana users may experience increased postoperative pain and may require more opioids compared with non-users. In a systematic review and meta-analysis of 4259 patients, Abdallah et al demonstrated that patients receiving cannabinoids appeared to have an increased weighted mean difference of pain at 12 hours by 0.83 cm on a 10-cm visual analog scale (VAS) (95% CI 0.04 to 1.63, p=0.04) but no differences in severity of rest pain at 24 hours. In addition, a retrospective pilot study investigating acute pain management in four trauma centers across the USA found that cannabis users reported higher pain scores and consumed larger quantities of opioids for management of acute pain (25%–37% higher) compared with non-cannabis users. In another, Jamal et al reported the effects of recreational cannabis smoking (prevalence 11.9%) on postoperative pain management within the first 24 hours after abdominal surgery. Cannabis users required a higher dose of postoperative morphine, extrapolated to 23% increased dose requirement in a model controlling for age, preoperative morphine use, and other comorbidities, and therefore increased vigilance to pain and opioid use in the postoperative phase may be required.

A recent prospective study of 1335 adults undergoing elective surgery showed that cannabis users have higher pain levels and greater opioid use before and after surgery.

Dose-dependent effects of inhaled marijuana may contribute to hyperalgesia. While the mechanism for hyperalgesia is still unclear, it is postulated that it may be a phenomenon of long-term use and TRPV1 modulation. Further discussion about the role of cannabis in pain will be discussed in subsequent questions.

Statement 1: Acute cannabis intoxication and active CUD may be associated with increased risk for acute postoperative MI and cerebrovascular morbidity. Level of certainty: Moderate

Statement 2: A cannabis-using patient may have delayed gastrointestinal motility and may also be at a higher risk for PONV. Level of certainty: Moderate

Statement 3: Cannabis users and patients with CUD may be associated with higher postoperative pain scores and opioids use. Level of certainty: Moderate

Recommendation 1: Based on the currently available evidence, we do not recommend the routine use of additional postoperative monitoring for cardiac or neurological adverse events. However, we do recommend increased vigilance given that cardiac and neurovascular events do frequently occur in the postoperative period. Grade C

Recommendation 2: Based on the currently available evidence, we recommend using multimodal analgesia incorporating regional analgesia if appropriate and using opioids as rescue medication. Patients may need additional follow-up for adequacy of analgesia and the need for adjusting postoperative pain medications accordingly. Grade C
the quantity of cannabinoid consumed as package labeling may be wildly inaccurate, even when obtained from state-licensed dispensaries.114

Multiple surveys of patients with chronic pain have found a significant number of reported reductions in opioid use through the use of cannabis, and case series report similar findings.250–256 This outcome has not been consistently observed in RCTs and reviews of cannabis for chronic pain.257–259 A propensity-matched retrospective analysis of a large national database found that non-medical use of cannabis was associated with a significantly greater risk of prescription opioid use disorder, while both medical and non-medical use purposes were associated with an increased risk of prescription opioid misuse.260 However, such findings may depend on what tool is used to define opioid use disorder as some (such as the DSM-5 (The Diagnostic and Statistical Manual of Mental Disorders, Fifth Edition) criteria) overestimate the diagnosis in chronic pain patients. There are abundant preclinical data demonstrating interactions between the endogenous cannabinoid system and opioid pathways. CB1 and mu-opioid receptors may interact and have been shown to co-localize in the dorsal horn of the spinal cord in rats and share intracellular signaling pathways.261–263 Cannabinoids may act on opioid receptors and opioid antagonists may block some of the effects of THC.263, 264 Local administration of naloxone inhibited the antinociceptive effects of an experimental CB2-receptor agonist in rats, and this drug was found to stimulate beta-endorphin release from cultured human skin cells.265 CB1 and CB2 knockout mice display reduced antinociceptive responses to spinal and peripheral morphine but not systemic morphine compared with wild-type mice, suggesting that interactions between opioid and cannabinoid receptors may not contribute to supraspinal antinociception266. Intrathecal THC produced antinociception in a rat model of neuropathic pain that was not reversed by naloxone.267 Several CB2-receptor agonists have been found to have analgesic effects that are additive or synergistic with opioids depending on the drug combinations and pain models studied.268 269 In rats, systemic THC has an additive analgesic effect when combined with morphine, while, a full cannabinoid-receptor agonist showed synergistic effects in a pain model.270 A meta-analysis of 19 preclinical studies found a synergistic effect between cannabis and opioids.271

In patients taking stable doses of opioids for chronic pain, the administration of vaporized cannabis over 5 days resulted in statistically significant reductions in chronic pain without alterations in blood levels of opioids or changes in pulse oximetry values.271 It has been postulated that cannabinoid-opioid interactions may have a role in treating opioid withdrawal.272 273 Cannabinoids may ameliorate the symptoms of opioid withdrawal in humans and animals; studies of adjunctive cannabinoid administration during opioid detoxification have not found an increased rate of serious adverse events.274 275

A single oral dose of cannabis extract containing 10–15 mg THC after surgery produced significant analgesia and significant sedation compared with a 5 mg dose suggesting a dose-dependent effect although there was no placebo group for comparison.276 Retrospective analyses have found the addition of dronabinol to the postoperative pain regimen was associated with improved pain control, significantly reduced opioid consumption, independent of patients’ preadmission cannabis use and consequently significantly shorter length of stay.114 277 However, when authors compared opioid consumption between groups per same inpatient days, there was no difference in opioid consumption. Unfortunately, these studies did not examine the frequency of adverse events between the groups.

A single dose of the cannabinoid levonantradol produced significant but not dose-dependent analgesia after surgery compared with placebo but was associated with frequent but primarily mild side effects.277 In other randomized placebo-controlled trials, oral THC, dronabinol, nabilone, and novel experimental cannabinoid-receptor agonists given perioperatively did not appear to reduce postoperative pain or opioid use compared with placebo, and perioperative high-dose nabilone was associated with higher pain scores at rest and with movement compared with placebo.128 278–282 There is little evidence to support the use of cannabinoids for acute pain.283 A 2020 meta-analysis of cannabis for the treatment of acute postoperative pain based on eight RCTs and four observational studies found no difference in rest pain at 1, 6 or 24 hours after surgery or cumulative opioid consumption at 2 or 24 hours postoperatively compared with controls.284 Rather, patients receiving cannabinoids reported significantly greater pain scores compared with controls in three studies that evaluated pain at 12 hours after surgery. Notably, patients who received cannabinoids had 3.24 times greater odds of developing postoperative hypotension compared with controls (95% CI 1.12 to 9.36; p = 0.03).284

Few clinical or experimental studies of cannabis and opioid coadministration in healthy participants have reported adverse effects on vital signs. In one study involving healthy volunteers, the co-administration of dronabinol 20 mg with an opioid failed to produce analgesia for any pain modality and antagonized morphine analgesia for pressure pain.285 Dronabinol alone significantly increased heart rate but changes in oxygen saturation were only observed when THC and morphine were combined. An intravenous dose of THC after oxymorphone increased sedation and further decreased ventilation and CO2-ventilatory response without significantly changing respiratory rate.286 The combination of oxymorphone and THC increased cardiac index and heart rate while decreasing total peripheral resistance, but blunted the cardiovascular response to increasing CO2.286 In a study specifically designed to assess safety, a bolus of fentanyl 1 μg/kg following by either 400 or 800 mg of oral CBD resulted in no respiratory depression or cardiovascular complications.286 Among patients undergoing endoscopy, those who reported daily cannabis use required more fentanyl for procedural sedation than patients who did not use cannabis, but no significant group differences in postprocedure recovery time or frequency of cardiopulmonary events were reported.287 Multiple observational studies reported a positive association between preoperative cannabis use and postoperative pain and opioid use. A retrospective cohort study of outcomes after total knee arthroplasty, which excluded patients with a history of opioid use, found no difference in postoperative inpatient opioid administration between cannabis users and non-users.288–289 Among trauma patients, a history of marijuana use was associated with higher opioid use and pain scores during hospitalization.289 A survey study of musculoskeletal trauma patients reported that patients using cannabis during recovery reported greater levels of pain relief and reduced opioid consumption.289 However, among patients who reported marijuana use during recovery from a musculoskeletal injury, there was an increase in total prescribed opioid dose and duration of opioid use compared with patients who had never used marijuana, even among patients who specifically reported that marijuana decreased their opioid use.123

In adolescent trauma patients, a history of marijuana use was positively associated with duration of opioid use after injury.290 Interestingly, a history of marijuana use was associated with significantly higher postoperative opioid use yet a trend toward lower pain scores after bariatric surgery.126
The assessment of cannabis use on postoperative pain and opioid use is complicated by frequent comorbid preoperative opioid use. Several retrospective studies of patients undergoing elective surgery reported that patients using cannabis preoperatively were significantly more likely to also use opioids and benzodiazepines before surgery compared with patients who did not use cannabis. A retrospective cohort study of early postoperative pain after major orthopedic surgery in patients who reported a history of cannabis or cannabinoid use found that patients using preoperative cannabis were not more likely to use opioids compared matched controls. The patients using cannabis had significantly greater pain scores at rest and with movement, and higher incidence of sleep disruption compared with propensity-matched control patients, with a non-significant trend toward greater opioid use among patients who used cannabis. In contrast, Jamal et al found significantly higher post-operative opioid use among cannabis users than non-users, but this finding lost significance (p=0.06) once additional variables including preoperative opioid use were included in the statistical model.

Statement 1: Cannabinoid studies in patients taking opioids for chronic pain suggest that there may be a therapeutic benefit of low-dose THC on pain and opioid use, but the opioid-sparing effect is not apparent in the setting of acute pain. Level of certainty: Low

Statement 2: None of the studies reviewed identified any increase in significant adverse events (moderate to severe respiratory depression or nausea/vomiting) with the co-administration of THC and an opioid in experimental studies with healthy volunteers, and few studies reported on these outcomes in the clinical setting. Level of certainty: Low

Statement 3: There is evidence of increased pain and opioid requirements postoperatively among patients who use cannabis. Level of certainty: Low

Recommendation 1: Opioids may be administered when indicated for the management of perioperative pain in patients who use cannabis with increased vigilance. Grade C

Recommendation 2: There is insufficient evidence to recommend for or against adjusting postoperative opioid prescriptions in surgical patients who consume cannabinoids. Grade I

**Question 9: How do cannabis withdrawal symptoms present in the postoperative period and is there evidence for specific treatment?**

Cannabis withdrawal symptoms (CWS) can occur in the postoperative period. A thorough preoperative cannabis use history aimed at understanding the quantity of cannabis routinely consumed may be difficult to obtain but is important for assessing the risk of withdrawal. Human and animal studies have suggested there is minimal risk of developing CWS with products containing CBD only. The DSM-5 criteria for the diagnosis of CWS includes the abrupt cessation of prolonged or heavy cannabis use accompanied by three or more symptoms including irritability or anger, anxiety, insomnia, decreased appetite, restlessness, altered mood and a physical symptom causing significant discomfort (i.e. abdominal pain, tremors, sweating, fever, chills, or headache). Symptoms of CWS can occur 24 to 72 hours after cannabis cessation, peak in the first week, and can last up to 2 weeks. ‘Heavy cannabis use’ is not defined in the diagnostic criteria of the DSM-5. The magnitude of cannabis withdrawal can be related to the quantity of cannabis consumed and is often seen following cessation of prolonged high-quantity cannabis use. A Canadian panel of experts suggested in a consensus document that there should be a high index of suspicion for cannabis withdrawal postoperatively in patients who consume greater than 1.5 g/1 of inhaled cannabis or 20 mg/day THC-dominant cannabis oil. In addition, patients consuming a cannabis product with an unknown phytocannabinoid content more than 2–3 times per day are at risk for cannabis withdrawal.

Cannabis users who are at risk for withdrawal (see above) should be monitored for cannabis withdrawal postoperatively. The Cannabis Withdrawal Scale can be used to monitor severity of the withdrawal symptoms. Withdrawal for a hospitalized patient can manifest as disrupted sleep, inadequate pain control, changes in opioid use, agitation, restlessness, and even early discontinuation of treatment.

Animal models suggest the cerebellum may mediate cannabis withdrawal. Another postulated mechanism is that regular cannabis use is associated with a down-regulation and desensitization of cortical and subcortical CB1 receptors, which begins to reverse after 24–48 hours of abstinence before returning to normal in approximately 4 weeks. Several pharmacological treatments for CWS exist including cannabinoid agonists and gabapentin, and N-acetylcysteine. A phase 2a pilot study examined the safety and efficacy of gabapentin for the treatment of cannabis dependence. A 12-week, randomized, double-blind, placebo-controlled clinical trial was conducted in 50 treatment-seeking outpatients for CUD. Gabapentin 1200 mg significantly decreased withdrawal symptoms as measured by the Marijuana Withdrawal Checklist compared with placebo. The FAAH inhibitor, PF-7845, reduced CWS and cannabis use in a double-blind, placebo-controlled, parallel group phase 2a trial of 48 men with CUD. A larger replication trial is currently ongoing. At the time of this document, replacement therapy with a cannabinoid agonist has the most supportive evidence as a treatment to minimize CWS. The most common cannabinoid agonists available in the USA are dronabinol, a synthetic form of THC, and nabilone, a synthetic THC derivative. Dronabinol significantly reduced CWS at a dose of 20 mg twice daily but failed to show efficacy in reducing CUD. Nabilone attenuated cannabis withdrawal and relapse in an experimental medicine study. The panel acknowledges that dronabinol and nabilone are not FDA-approved for this indication in the USA. Both are approved by the FDA for the treatment of nausea and vomiting associated with cancer chemotherapy in patients who have failed to respond adequately to conventional antiemetic treatments.

Statement: CWS may present in the postoperative period. Highest risk patients are those consuming high quantities or unknown amounts of THC containing products. The risk is considered to be less with individuals consuming CBD dominant (>10:1 CBD to THC ratio) products. Level of certainty: Moderate

Recommendation 1: Patients using cannabis should be counseled regarding the risk of CWS. Postoperatively, patients that consume cannabis routinely should be monitored for CWS using a validated and reliable scale. Grade C

Recommendation 2: The expert panel came to the consensus that initiating a cannabinoid agonist such as dronabinol at a low dose is the best choice to treat severe CWS postoperatively. Grade C

**CONCLUSION**

The medical, social, and political landscape of cannabis is fluid, changing on an almost daily basis. Cannabinoid use in the perioperative setting has significant potential negative medical implications. We hope these guidelines will help both clinicians and researchers in their pursuit of optimal patient care. In accordance with the National Academy of Medicine’s standards for

developing clinical practice guidelines the ASRA Pain Medicine task force will continue monitoring newly released relevant publications following the publication of this guideline and may revise the entire document or specific sections if new evidence warrants updated recommendations.

Author affiliations
1 Dept of Anesthesiology & Perioperative Care, UC Irvine Health, Orange, California, USA
2 Anesthesiology and Perioperative Medicine, Sidney Kimmel Medical College at Thomas Jefferson University, Philadelphia, Pennsylvania, USA
3 Anesthesia, University of Iowa Healthcare, Iowa City, Iowa, USA
4 Anesthesiology and Pain Medicine, Univ Toronto, Toronto, Ontario, Canada
5 Anesthesiology, Cedars-Sinai Medical Center, Los Angeles, California, USA
6 Anesthesiology, University of Colorado Health, Aurora, Colorado, USA
7 Anesthesiology and Perioperative Care, University of California Irvine, Irvine, California, USA
8 Anesthesiology, Mayo Clinic, Rochester, Minnesota, USA
9 Anesthesiology, Maimonides Medical Center, Brooklyn, New York, USA
10 Anesthesiology, Division of Pain Medicine, University of California San Diego, La Jolla, California, USA
11 Center for Pain Medicine, Western Reserve Hospital, Cuyahoga Falls, Ohio, USA
12 Anesthesiology, Division of Pain Medicine, University of California, San Francisco, California, USA
13 Dept of Anesthesiology & Perioperative Care, UC Irvine Health, Orange, California, USA
14 Anesthesiology, Maimonides Medical Center, Brooklyn, New York, USA
15 Anesthesiology, Mayo Clinic, Rochester, Minnesota, USA
16 Anesthesiology, University of Colorado Health, Aurora, Colorado, USA
17 Anesthesiology and Perioperative Care, University of California Irvine, Irvine, California, USA
18 National Pain Advocacy Center, Golden, Colorado, USA
19 AAFP Integrative Pain Care, Melville, New York, USA
20 Anesthesiology, Maimonides Medical Center, Brooklyn, New York, USA
21 Anesthesiology, Mayo Clinic, Rochester, Minnesota, USA
22 Anesthesiology, Division of Pain Medicine, University of California San Diego, La Jolla, California, USA
23 Center for Pain Medicine, Western Reserve Hospital, Cuyahoga Falls, Ohio, USA

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Twitter Shalini Shah @shalinishahMD, Eric S Schwenk @ESchwenkMD, Rakesh V Sondekoppam @rakesh6282, Hance Clarke @drhclarke, Gary Schwartz @garyschwarzmd and Samer Naroze @NarozeMD

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ORCID iDs Shalini Shah http://orcid.org/0000-0001-5200-225X Eric S Schwenk http://orcid.org/0000-0003-3464-4149 Rakesh V Sondekoppam http://orcid.org/0000-0003-1132-9066 Hance Clarke http://orcid.org/0000-0003-4975-3823 W Michael Hooten http://orcid.org/0000-0001-5645-6335 Eugene R Viscusi http://orcid.org/0000-0003-0260-4396 Samer Naroze http://orcid.org/0000-0003-1849-1402

Original research


