Evidence-based Synthesis Program

QUERI

Benefits and Harms of Cannabis in Chronic Pain or Post-traumatic Stress Disorder: A Systematic Review

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PREFACE

The VA Evidence-based Synthesis Program (ESP) was established in 2007 to provide timely and accurate syntheses of targeted healthcare topics of particular importance to clinicians, managers, and policymakers as they work to improve the health and healthcare of Veterans. QUERI provides funding for four ESP Centers, and each Center has an active University affiliation. Center Directors are recognized leaders in the field of evidence synthesis with close ties to the AHRQ Evidence-based Practice Centers. The ESP is governed by a Steering Committee comprised of participants from VHA Policy, Program, and Operations Offices, VISN leadership, field-based investigators, and others as designated appropriate by QUERI/HSR&D.

The ESP Centers generate evidence syntheses on important clinical practice topics. These reports help:

- Develop clinical policies informed by evidence;
- Implement effective services to improve patient outcomes and to support VA clinical practice guidelines and performance measures; and
- Set the direction for future research to address gaps in clinical knowledge.

The ESP disseminates these reports throughout VA and in the published literature; some evidence syntheses have informed the clinical guidelines of large professional organizations.

The ESP Coordinating Center (ESP CC), located in Portland, Oregon, was created in 2009 to expand the capacity of QUERI/HSR&D and is charged with oversight of national ESP program operations, program development and evaluation, and dissemination efforts. The ESP CC establishes standard operating procedures for the production of evidence synthesis reports; facilitates a national topic nomination, prioritization, and selection process; manages the research portfolio of each Center; facilitates editorial review processes; ensures methodological consistency and quality of products; produces "rapid response evidence briefs" at the request of VHA senior leadership; collaborates with HSR&D Center for Information Dissemination and Education Resources (CIDER) to develop a national dissemination strategy for all ESP products; and interfaces with stakeholders to effectively engage the program.

Comments on this evidence report are welcome and can be sent to Nicole Floyd, ESP CC Program Manager, at <u>Nicole.Floyd@va.gov</u>.

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EXECUTIVE SUMMARY

INTRODUCTION

Eight states and the District of Columbia have legalized cannabis use for recreational purposes, and 28 states plus the District of Columbia have legalized cannabis for medical purposes. Recent studies suggest that 45-80% of individuals who seek cannabis for medical purposes do so for pain management, and an estimated 6%-39% of patients prescribed opioid medication for pain are also utilizing cannabis. Over one-third of patients seeking cannabis for medical purposes list post-traumatic stress disorder (PTSD) as the primary reason for the request. Approximately 15% of Veterans who are treated in Department of Veterans Affairs (VA) outpatient PTSD clinics report recent (past 6 months) cannabis use.

Given the social, political, and legal changes surrounding cannabis use, physicians in both VA and non-VA settings will increasingly need to engage in evidence-informed discussions about the potential benefits and harms of cannabis use with their patients. Despite the rapidly moving legislative landscape, there is little comprehensive and critically appraised information available about what is known and not known about cannabis use for the treatment of chronic pain or PTSD.

The objectives of this systematic review are to: 1) assess the physical and mental health outcome effects of cannabis in patients with chronic pain; 2) assess the physical and mental health outcome effects of cannabis in patients with PTSD; 3) assess the impact of short- and long-term cannabis use on the risk of adverse effects such as pulmonary diseases, cardiovascular diseases, cancer, cannabis use disorder (CUD), and psychosis in the general adult population; and 4) provide a broad overview of more recently recognized "emerging harms" of cannabis use.

METHODS

DATA SOURCES AND SEARCHES

We developed search strategies in consultation with a research librarian. We searched multiple data sources including Ovid MEDLINE, Embase, PubMed, PsycINFO, PILOTS Database, EMB Reviews (CDSR, DARE, HTA, Cochrane CENTRAL, *etc*), and grey literature sources from database inception through February 2016.

STUDY SELECTION

We included English-language studies of plant-based cannabis preparations including wholeplant preparations (*eg*, cannabis cigarettes, hashish, oils), whole plant extracts such as nabiximols (an oromucosal spray delivering 2.7 mg tetrahydrocannabinol [THC]/2.5 mg cannabidiol [CBD], currently available by prescription only in Europe), and capsular THC/CBD preparations. We did not include synthesized, pharmaceutically-prepared cannabinoids such as dronabinol or nabilone because the efficacy of synthetic cannabinoid preparations for chronic pain was examined in 2 recent review articles. We were broadly inclusive of different types of cannabis preparations because there are many different cannabis preparations in dispensaries, and clinicians may therefore encounter patients using many different forms.

To address the efficacy of cannabis in treating chronic pain or PTSD, we examined controlled clinical trials or rigorously designed observational studies with control groups that adjusted for important confounders and used validated outcome measures. We determined our study selection criteria for pre-specified harms based on whether the likelihood of the adverse outcome might be substantially different in populations with chronic pain or PTSD. For example, we anticipated that rates of depression and anxiety in patients with chronic pain or PTSD were likely to be substantially different than the general population, so we only included studies reporting these harms in the specific populations of interest. In contrast, we thought it unlikely that rates of pulmonary effects or cancer would be particularly influenced by the presence of chronic pain or PTSD, so we included studies in general adult populations for these outcomes.

Given the broad scope of this review, we summarized data from existing good-quality systematic reviews when available to address each question and outcome of interest and then added individual studies meeting inclusion criteria that were published after the end search date of the included review, or were not included in a prior systematic review.

DATA ABSTRACTION AND QUALITY ASSESSMENT

From each study, we abstracted the following where available: study design, objectives, setting, population characteristics, subject inclusion and exclusion criteria, number of subjects, duration of follow-up, the study and comparator interventions (formulation, strength, *etc*), important co-interventions, health outcomes, healthcare utilization, and harms. We assessed study quality and graded the strength of evidence using published criteria.

DATA SYNTHESIS AND ANALYSIS

We qualitatively synthesized the evidence on the benefits and harms of cannabis. For the subgroup of neuropathic pain studies, we conducted a study-level meta-analysis of the proportion of patients experiencing clinically significant (\geq 30%) pain relief.

RESULTS

RESULTS OF LITERATURE SEARCH

We included 12 systematic reviews and 48 primary studies after reviewing 10,875 titles and abstracts.

SUMMARY OF RESULTS FOR KEY QUESTIONS

Key Question 1. What are the effects of cannabis on health outcomes and healthcare utilization for adults who have chronic pain?

We found low-strength evidence that cannabis preparations with precisely defined THC:CBD content (most in a 1:1 to 2:1 ratio) have the potential to improve neuropathic pain but insufficient evidence in other patient populations. Most studies are small, many have methodologic flaws, and the long-term effects are unclear given the brief follow-up duration of most studies. The applicability of these findings to current practice may be low in part because the formulations studied may not be reflective of what most patients are using, and because the consistency and accuracy of labeled content in dispensaries are uncertain.



Key Question 2. What are the effects of cannabis on health outcomes and healthcare utilization for adults who have PTSD?

We found insufficient evidence examining the effects of cannabis in patients with PTSD. We found 2 observational studies comparing outcomes in cannabis users to a control group that had not used cannabis; cannabis use was not associated with improved outcomes in either study. We found no evidence addressing whether effects differed according to other comorbidities in patients with PTSD.

Key Question 3. What are the harms associated with cannabis use in adults?

General Adverse Events

Data from 2 systematic reviews examining cannabis for chronic pain suggest that cannabis may be associated with a higher risk of short-term adverse effects, although rates of adverse events did not significantly differ between groups in the additional trials we reviewed. While most adverse events were mild, there were possible treatment-related serious adverse events such as suicide attempts, paranoia, and agitation.

Medical Harms

Pulmonary effects

Moderate-strength evidence from 2 well-designed cohort studies suggest that low levels of cannabis smoking do not adversely impact lung function over about 20 years in young adults, but there is some evidence suggesting that heavy (*ie*, daily) use may have the potential to cause adverse pulmonary effects over an extended period of time. There were no studies in older users, or in those with medical comorbidities such as chronic obstructive pulmonary disease (COPD) or heart disease.

Cardiovascular events

There is insufficient evidence from 2 studies about the effect of cannabis use on the risk of cardiovascular events, due to methodological limitations including lack of longitudinal exposure measurement and potential recall bias.

Cancer

A meta-analysis of 9 case-control studies provided low-strength evidence that cannabis use does not appear to be associated with an increased risk of head and neck or lung cancer. There was insufficient evidence about the effects of cannabis on testicular or transitional cell cancer. We found no studies examining the effects on other types of cancer.

Motor vehicle accidents

Moderate-strength evidence from a recent meta-analysis of 21 multi-national observational studies found that acute cannabis intoxication was associated with a moderate increase in collision risk (odds ratio [OR] 1.35; 95% confidence interval [CI], 1.15 to 1.61).

Mental Health-related Harms

Suicidal behaviors

We found no studies examining the effects of cannabis use on suicide risk in patients with chronic pain or PTSD. A review and meta-analysis of 4 epidemiological studies in general populations found significantly increased odds of suicide death (pooled OR 2.56; 95% CI, 1.25 to 5.27) with any cannabis use.

Mania

We found no studies examining the effects of cannabis on the risk of mania among persons with PTSD or chronic pain. A systematic review of 6 longitudinal studies in other populations detected an association between cannabis use and exacerbation of manic symptoms in patients with known bipolar disorder, and an increased incidence of new-onset mania symptoms among populations without a diagnosis of bipolar disorder (OR 2.97; 95% CI, 1.80 to 4.90).

Psychosis

A systematic review and 7 studies consistently found an association between cannabis use (specifically related to THC content) and the development of psychotic symptoms (low-strength evidence). There is evidence of a dose-response relationship, and there is experimental evidence documenting the risk of acute, transient psychotic symptoms within hours of use; however, no studies were specifically in PTSD or chronic pain populations.

Cognitive effects

One systematic review of studies in general populations provides moderate-strength evidence that active, long-term cannabis use is associated with small negative effects on all domains of cognitive function, but there was insufficient evidence of cognitive effects in past users.

Cannabis use disorder (CUD)

Cannabis use was associated with incident cannabis use disorder (adjusted odds ratio, 9.5 [CI, 6.4 to 14.1]) in a large (N = 34653) prospective cohort study.

We found no studies comparing rates of CUD in chronic pain or PTSD populations to other populations.

Other studies of CUD provide potentially relevant cross-sectional data examining the prevalence of CUD among patients with chronic pain. For example, one large cross-sectional study of Veterans using administrative data found that about 2% of Veterans with non-cancer pain had a diagnosis of CUD, and that this proportion increased (up to about 4%) among subgroups with higher numbers of opioid prescriptions. In a non-VA study using structured diagnostic interviews the prevalence of cannabis abuse was 2.4% and cannabis dependence was 0.9%.

Emerging Harms

Chronic cannabis use has been associated with a severe form of cyclic vomiting called the cannabinoid hyperemesis syndrome. There have also been reports of serious infectious diseases including aspergillosis and tuberculosis associated with smoked cannabis, and a severe acute illness associated with intravenous cannabis use. The recent availability of edible forms of



cannabis with high THC content has been associated with episodes of severe acute psychosis. There is mixed evidence regarding the effects of cannabis on violent behavior.

Key Question 4. What are important areas of ongoing research and current evidence gaps in research on cannabis for chronic pain or PTSD, and how could they be addressed by future research?

We identified 10 ongoing randomized controlled trials (RCTs) examining the effectiveness of cannabis for a variety of chronic pain conditions, including several populations included in this report (3 studies for cancer pain and 2 studies for neuropathic pain), as well as conditions for which there is currently very little or no evidence (osteoarthritis, sickle cell disease, low back pain, and ulcerative colitis).

There are 2 recently initiated RCTs examining the benefits and harms of cannabis for PTSD that should add to the body of evidence.

SUMMARY AND DISCUSSION

KEY FINDINGS AND STRENGTH OF EVIDENCE

We reviewed the literature examining benefits of cannabis in chronic pain and PTSD populations, as well as literature examining potential harms relevant to these populations. We found low-strength evidence that cannabis preparations with precisely defined THC-cannabidiol content (most in a 1:1 to 2:1 ratio) may alleviate neuropathic pain but insufficient evidence in populations with other types of pain. Most studies are small, many have methodological flaws, and the long-term effects are unclear given the brief follow-up of most studies. Among neuropathic pain studies, we found a discrepancy between continuous and dichotomous pain outcomes. Possible interpretations are that cannabis is simply not consistently effective or that, although cannabis may not have clinically important effects on average, subgroups of patients may experience large effects. We did not find data to clarify which subgroups of patients are more or less likely to benefit.

We found no trials that met our inclusion criteria examining the effects of cannabis in PTSD populations, and there was insufficient evidence from observational studies to draw conclusions about its effectiveness in patients with PTSD.

In younger populations, light to moderate cannabis use does not appear to be associated with adverse pulmonary effects over the long-term, but pulmonary effects have not been studied in older populations or individuals with comorbid medical conditions. There is insufficient to low-strength evidence examining the effects of cannabis use on the risk of various types of cancer. There is consistent evidence that suggests an association between cannabis use and psychotic symptoms, as well as cognitive impairment in active users in general populations, though there is limited evidence specific to patients with chronic pain or PTSD. There are a number of adverse effects that appear to be related to cannabis use and may be important for clinicians to be familiar with, but whose incidence has not been well-characterized. These include infectious disease complications, cannabis hyperemesis syndrome, and violent behavior.

The summary of findings and strength of evidence supporting these findings are detailed in the table that follows.



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Summary of Evidence for the Benefits and Harms of Cannabis in Chronic Pain or PTSD Populations

N studies (N combined participants)		Findings	Strength of Evidence ^a	Comments	
Chronic Pain					
 Chronic Pain Multiple sclerosis (MS) 	 4 Low ROB studies (combined N=1017; 24 to 424 per study): 2 of THC/CBD capsules 1 of nabiximols 1 of sublingual spray delivering THC, CBD, or THC/CBD combined 3 Unclear ROB studies of nabiximols (combined N=562; 36 to 337 per study) 7 High ROB studies (combined N=430; 13 to 160 per study): 3 of nabiximols 2 of THC/CBD capsules 1 of smoked THC 1 of oral THC 	Favorable effect on pain and spasticity: Significant relief from patient-reported muscle stiffness, pain, and spasticity occurred with 12 to 15 weeks of treatment with THC (2.5 mg)/CBD (1.25 mg) capsules in 2 studies. A 12-week study of nabiximols (2.7 mg THC/2.5 mg CBD oromucosal spray) reported significant improvement in spasticity. A sublingual spray delivering 2.5 mg of CBD, THC, or both for sequential 2-week periods reported mixed effects. THC alone significantly improved pain and spasticity, but CBD alone and THC/CBD combined had inconsistent effects.	Low	Few methodologically rigorous studies, but fair number of patients; inconsistent results; little long-term data; restrictive entry criteria in largest study which only included patients with initial response in run-in phase; applicability to formulations available in dispensaries may be low	
	4 Low ROB studies (combined N=1017;	Other outcomes:			
	 4 Low ROB studies (combined N=1017; 24 to 424 per study): 2 of THC/CBD capsules 1 of nabiximols 1 of sublingual spray delivering THC, CBD, or THC/CBD combined 	Small improvements in sleep in 4 studies: Self-reported sleep quality improved in 2 studies of THC/CBD capsules. Nabiximols were significantly superior to placebo for reducing sleep disruption in a 12-week study (N=241). Sleep improved significantly in a small study (N=24) of a sublingual spray containing 2.5 mg each of CBD:THC. Other: Nabiximols were significantly superior to	Low (sleep)	Few methodologically rigorous studies, but fair number of patients; inconsistent results; little long-term data; restrictive entry criteria in largest study which only included patients with initial response in run-in phase; applicability to current practice may be low	
		placebo for Barthel Activities of Daily Living $(P=.0067)$, Physician Global Impression of Change $(P=.005)$, Subject Global Impression of Change $(P=.023)$, and Carer Global Impression of Change $(P=.005)$ in Function in a 12-week study (N=241).	Insufficient (other outcomes)	Only one study of nabiximols – not tested otherwise	

	N studies (N combined participants)	Findings	Strength of Evidence ^a	Comments
• Neuropathic pain	 11 low ROB studies (combined N = 593) 4 of smoked THC (combined N = 150) 3 of vaporized THC (combined N = 97) 3 of nabiximols (combined N = 312) 1 of oromucosal spray delivering THC or THC+CBD (N = 34) 1 unclear ROB study of nabiximols (N = 30) 1 high ROB trial (N = 125) 	Studies did not find a clinically significant between-group difference on continuous pain scales, but a higher proportion of intervention patients had clinically significant pain relief up to several months later. In a meta-analysis of 9 studies, intervention patients were more likely to report \geq 30% improvement in pain (combined RR, 1.43 [95% CI, 1.16–1.88]; $I^2 = 38.6\%$; $P = 0.111$).	Low	Few patients enrolled in most low ROB studies; inconsistent results; marked differences among studies in dosing and delivery mechanism; brevity of study duration; low applicability to formulations available in dispensaries.
	1 Low ROB study of smoked THC (N=23)	Other outcomes reported in low ROB studies: A study of vaporized cannabis reported that 25 mg with 9.4% THC administered as a single smoked inhalation 3 times daily resulted in significant improvements in sleep quality.	Insufficient	Only one small study
General/other/mixed populations2 Low ROB studies: - 1 trial of sublingual spray delivering THC, CBD, or THC/CBD combined (N=34) - 1 observational study of cannabis containing 12.5% THC (smoked, oral, or vaporized) (N=431) 3 Unclear ROB studies of nabiximols (combined N=428; 10 to 360 per study) 3 High ROB studies (combined N=265; 18 to 177 per study): 2 of nabiximols		Small improvements in pain, but no effect on sleep, mood, quality of life.	Insufficient	Only one small low ROB study in which the bulk of the patients had MS; larger observational study had high drop-out rate
- 2 of nabiximols - 1 of THC capsules PTSD 2 observational studies in Veterans with PTSD: - 1 Medium ROB (N=2276) - 1 High ROB (N=700)		Cannabis was not associated with an improvement in mental health symptoms.	Insufficient	No trials; only 2 observational studies with methodologic flaws

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	N studies (N combined participants)	Findings	Strength of Evidence ^a	Comments
Harms				
• General AEs	2 systematic reviews of chronic pain	Cannabis-based treatments were associated with an overall higher risk of short-term, non-serious AEs.		Consistent findings except for serious AE
• Medical harms				
Ø Pulmonary function	2 Low ROB prospective cohort studies with 20-32 years follow-up (combined N=6053)	In young adults, low levels of cannabis smoking did not adversely affect lung function over about 20 years. A previous meta-analysis of 5 studies found no	Young adults: Moderate Older adults: No evidence	Two well-done prospective cohort studies, but limited information about effects of heavy use and no information ir
	1 systematic review of 5 observational studies (3 cohort, 2 cross-sectional) (combined N=851)	increased risk for pulmonary adverse effects, OR (95% CI): 0.80 (0.46-1.39).		older or multimorbid populations
Ø Cardiovascular	 2 High ROB observational studies: 1 case-crossover (N=3882) 1 cohort study (N=2097) 	Cannabis use at the time of myocardial infarction was not associated with mortality after mean 12.7 years follow-up, but longitudinal use was not assessed. Risk of myocardial infarction within an hour of cannabis use was significantly elevated compared with periods of non-use but this finding may be inflated by recall bias, OR (95% CI): 4.8 (2.9-9.5).	Insufficient	Recall bias; inadequate controlling for confounders; lack of longitudinal exposure data
Ø Cancer				
§ Lung	 patient-level meta-analysis of 6 case- control studies (2150 cases) High ROB cohort study (N=49,231) 	The meta-analysis found no association between light cannabis use and lung cancer.	Low	Recall bias; mostly light users, few heavy users; the large cohort study had no information about exposure over time
§ Head/neck/ora 1	Meta-analysis of 9 case-control studies (5732 cases)	No association between cannabis use and cancer, OR (95% CI): 1.02 (0.91-1.14); generally consistent across studies and no evidence of dose-response.	Low	Imprecise exposure measurement with potential recall bias; ever use among studies ranged from 1 to 83%
§ Testicular Meta-analysis of 3 High ROB case-cor studies (719 cases)		 An increase in cancer risk for weekly users Insufficient compared to never-users appeared with non- seminoma cancers but not seminoma cancers, OR (95% CI): 1.92 (1.35-2.72). 		Potential confounding from recall bias and tobacco use
§ Transitional cell	1 High ROB VA case-control study (52 cases)	Risk of cancer with > 40 joint-years cannabis use compared to none, OR 3.4 (unadjusted, P=.012).	Insufficient	One very small case-control study with several methodologic flaws



	N studies (N combined participants)	Findings	Strength of Evidence ^a	Comments
Motor vehicle accidents	Meta-analysis of 21 observational studies (combined N=239,739)	Increase in collision risk, OR (95% CI): 1.35 (1.15-1.61).	Moderate	The small but significant increase in risk was seen consistently across numerous sensitivity analyses and after adjustment in meta-regression analyses
 Mental health Ø Suicidal behaviors 	No studies in chronic pain or PTSD populations.		No evidence (chronic pain or PTSD)	Meta-analysis of 4 studies in the general population reported significantly increased odds of suicide with any cannabis use, OR (95% CI): 2.56 (1.25-5.27).
Ø Mania	No studies in chronic pain or PTSD populations		No evidence (chronic pain or PTSD)	A systematic review found an increased incidence of new- onset mania symptoms among populations without a diagnosis of bipolar disorder, OR (95% CI): 2.97 (1.80 to 4.90).
Ø Psychosis	 systematic review studies including patients without psychotic symptoms at baseline: 3 Low ROB studies 3 Medium ROB studies 1 High ROB study 	History of cannabis use was associated with an increase in risk of developing psychotic symptoms.	Low	Consistent evidence from large observational studies and some evidence of increased risk with higher levels of use; consistent with data from small experimental studies suggesting risk of acute psychosis in some patients; magnitude of risk unclear and not specifically studied in chronic pain or PTSD populations
	1 systematic review of 33 studies	Active long-term cannabis use associated with small negative effects on all aspects of cognition. Mixed, inconsistent findings on long-term effects in past users.	Moderate Insufficient (past use)	Consistent data from large number of studies on effects on active long-term use, but inconsistent findings from smaller number of studies regarding effects in those that were abstinent and no data available specifically in chronic pain or PTSD populations



	N studies (N combined participants)	Findings	Strength of Evidence ^a	Comments
ØCUD	One large cohort study (N=34,653; N = 1279 past year cannabis use in last year)	OR incident CUD 9.5 (95% CI 6.4-14.1) Prevalence CUD (among those using in last year) 36% Prevalence past year cannabis dependence 7.7% Prevalence past year cannabis abuse 28%	Low	In cross-sectional studies, the prevalence of CUD in chronic pain populations was about 2%

Abbreviations: AE = adverse event; CBD = cannabidiol; CI = confidence interval; CUD = cannabis use disorder; MS = multiple sclerosis; N = number; OR = odds ratio; PTSD = post-traumatic stress disorder; ROB = risk of bias; THC = tetrahydrocannabinol; VA = Department of Veterans Affairs.

^a The overall quality of evidence for each outcome is based on the consistency, coherence, and applicability of the body of evidence, as well as the internal validity of individual studies. The strength of evidence is classified as follows:

- High = Further research is very unlikely to change our confidence on the estimate of effect.
- Moderate = Further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.
- Low = Further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.
- Insufficient = Any estimate of effect is very uncertain.

APPLICABILITY

Efficacy trials often examined the use of precisely prepared THC:CBD preparations in capsular or spray form. Cannabis forms available in medical or recreational dispensaries vary widely: the content of preparations may not be known, may vary significantly from what is studied, and the actual contents may differ from what is labeled on the product. There is virtually no information to guide discussions of benefits and harms in older populations or populations with multiple comorbidities. The best observational studies we found typically included younger, healthier populations. We found relatively little information about mental health harms specifically in chronic pain or PTSD populations, but information about harms such as cognitive impairment obtained from other populations may still provide useful information for counseling patients for the time being.

RESEARCH GAPS/FUTURE RESEARCH

There is no conclusive information about the benefits of cannabis in chronic pain or PTSD populations and limited information about its harms, so methodologically strong research in almost any area of inquiry is likely to add to the strength of evidence. It appears that the United States (US) government is poised to lift restrictions on access to cannabis for research, which may speed the development of this evidence base that has lagged far behind policy changes regarding the use of cannabis for medical purposes in many states.

CONCLUSIONS

Although cannabis is increasingly available for medical and recreational use, there is very little methodologically rigorous evidence examining its effects in patients with chronic pain or PTSD. There is limited evidence suggesting that cannabis may improve pain and spasticity in patients with MS, but no consistent, high-quality data showing benefit from cannabis for the treatment of pain in other populations. Cannabis use is associated with an increased risk of short-term adverse effects, but data on its effects on long-term physical health vary. Cannabis use is associated with cognitive impairment in active users and potentially serious mental health adverse effects such as psychotic symptoms, though the absolute risk and application specifically to chronic pain and PTSD populations are uncertain.

Abbreviation	Term
AHRQ	Agency for Healthcare Research and Quality
CBD	Cannabidiol
CI	Confidence interval
COPD	Chronic Obstructive Pulmonary Disease
CUD	Cannabis use disorder
DoD	Department of Defense
DSM-5	Diagnostic and Statistical Manual of Mental Disorders, Fifth Edition
EDSP	Early Developmental Stages of Psychopathology
FEV1	Forced expiratory volume
FVC	Forced vital capacity
HIV	Human immunodeficiency virus
HR	Hazard ratio
ICD	International Statistical Classification of Diseases and Related Health Problems
IOM	Institute of Medicine
MS	Multiple sclerosis
N	Number
NRS	Numeric rating score
OR	Odds ratio
PICOTS	Patient population, intervention, comparator, outcome, timing parameters, and study designs
PTSD	Post-traumatic Stress Disorder
QOL	Quality of life
RCT	Randomized controlled trial
ROB	Risk of bias
Т	Time point
THC	Tetrahydrocannabinol
UK	United Kingdom
US	United States
VA	Department of Veterans Affairs
VAS	Visual Analogue Scale

Abbreviations Table

EVIDENCE REPORT

INTRODUCTION

Cannabis use has become more common among United States (US) adults, with the prevalence of adults reporting past-year cannabis use nearly doubling between 2001 and 2013 to one in 10 adults.¹ Young adults ages 18-29 are nearly 4 times more likely to have used cannabis in the past year than adults ages 45-64.

The use of cannabis for medicinal purposes has also become increasingly accepted. In California, which was the first state to legalize cannabis for medical purposes in 1996, about 5% of all adults reported having used cannabis for medical purposes in 2012.² In a recent poll, 76% of physicians supported the use of cannabis for medical purposes in certain circumstances.³

Eight states and the District of Columbia have legalized cannabis use for recreational purposes, and 28 states plus the District of Columbia have legalized cannabis for medical purposes. Both houses of Congress recently passed H.R. 2577, which would allow federally-employed physicians working for the Veterans Health Administration to recommend cannabis for medical purposes to Veterans if appropriate in states that have legalized its use.⁴

The conditions that would qualify a patient to use cannabis for medical purposes differ across states, but nearly all include chronic pain itself or diseases which are likely to cause chronic pain (such as multiple sclerosis [MS]-related spasticity). Several states also list post-traumatic stress disorder (PTSD) as a qualifying condition, which is of particular importance to Veterans and, indeed, was one of the rationales cited for the genesis of H.R. 2577.

Approximately 30% of Americans currently experience chronic pain,⁵ a figure that is estimated to increase as the population ages and manages more chronic medical conditions.⁶ Recent studies suggest that 45-80% of individuals who seek cannabis for medical purposes do so for pain management^{7,8} and among patients who are prescribed long-term opioid therapy for pain, an estimated 6%-39% are also utilizing cannabis.^{9,10}

Recent research suggests that over one-third of patients seeking cannabis for medical purposes in states where it is legal list PTSD as the primary reason for the request.¹¹ Approximately 15% of Veterans who are treated in Department of Veterans Affairs (VA) outpatient PTSD clinics report recent (past 6 months) cannabis use.¹²

In the past, use had been limited to inhalation or ingestion of parts of the whole plant of the genus Cannabis. More recently, many more formulations of cannabis have become available in recreational and medical cannabis dispensaries including an array of edibles, oils, tinctures, as well as plant extracts with varying ratios of the 2 active ingredients of cannabis: tetrahydrocannabinol (THC) and cannabidiol (CBD). There are also 2 purely synthetic cannabinoids available in the US by prescription only (dronabinol and nabilone).

Given the social, political, and legal changes surrounding cannabis use, physicians in both VA and non-VA settings will increasingly need to engage in evidence-informed discussions with their patients about the potential benefits and harms of cannabis use. Despite the rapidly moving legislative landscape, there is little comprehensive and critically appraised information available



about what is known and not known about cannabis use for the treatment of chronic pain or PTSD.

The objectives of this systematic review are to: 1) assess the physical and mental health outcome effects of cannabis in patients with chronic pain; 2) assess the physical and mental health outcome effects of cannabis in patients with PTSD; 3) assess the impact of short- and long-term cannabis use on the risk of adverse effects such as pulmonary diseases, cardiovascular diseases, cancer, cannabis use disorder (CUD), and psychosis in the general adult population; and 4) provide a broad overview of more recently recognized "emerging harms" of cannabis use.

METHODS

TOPIC DEVELOPMENT

The research questions for this systematic review were developed after a topic refinement process that included a preliminary review of published peer-reviewed literature, and consultation with internal partners, investigators, and stakeholders. The proposed Key Questions are as follows:

Key Question 1: What are the effects of cannabis on health outcomes and healthcare utilization for adults who have chronic pain?

Key Question 1A: Do the effects differ by patient subgroup, such as patient medical and mental health comorbidities?

Key Question 2: What are the effects of cannabis on health outcomes and healthcare utilization for adults who have PTSD?

Key Question 2A: Do the effects differ by patient subgroup, such as patient medical and mental health comorbidities?

Key Question 3: What are the harms associated with cannabis use in adults?

Key Question 3A: Do the harms differ by patient subgroup, such as patient medical and mental health comorbidities?

Key Question 4: What are important areas of ongoing research and current evidence gaps in research on cannabis for chronic pain or PTSD, and how could they be addressed by future research?

A protocol describing the review plan was posted to a publicly accessible website before the study was initiated.¹³

SEARCH STRATEGY

Search strategies were developed in consultation with a research librarian. To identify relevant articles, we searched MEDLINE, PubMed, EMBASE, PsycINFO, PILOTS Database, EMB Reviews (CDSR, DARE, HTA, Cochrane CENTRAL, *etc*), and grey literature sources from database inception through February 2016 (Appendix A). We reviewed the bibliographies of relevant articles and contacted experts to identify additional studies.

To identify in-progress or unpublished studies for Key Question 4, we searched ClinicalTrials.gov, International Clinical Trials Registry Platform (WHO ICTRP), ISRCTN Registry, NIH Reporter, AHRQ Gold, and the American Cancer Society Database of Studies. We also queried the Technical Expert Panel and used snowball sampling techniques to identify relevant ongoing research.

STUDY SELECTION

The criteria for patient population, intervention, comparator, outcome, timing parameters, and study designs (PICOTS) that apply to each key question are specified in Table 1. We included English-language studies of plant-based cannabis preparations or whole plant extracts such as



nabiximols, which is a non-synthetic pharmaceutical product with a standard composition and dose (oromucosal spray delivering 2.7 mg THC/2.5 mg CBD) available only in select European countries. We did not include synthesized, pharmaceutically-prepared cannabinoids such as dronabinol or nabilone because the efficacy of synthetic cannabinoid preparations for chronic pain was examined in 2 recent review articles.¹⁴⁻¹⁶ However, we broadly defined plant-based cannabis preparations to include any preparation of the cannabis plant itself (*eg*, cannabis cigarettes, hashish, oils), or cannabis plant extracts. We chose to be broadly inclusive of herbal preparations because US dispensaries offer a wide variety of concentrations and products, and clinicians may encounter patients who have used a variety of preparations.¹⁷

To address the efficacy of cannabis in treating chronic pain or PTSD (Key Questions 1 and 2), we examined controlled clinical trials or rigorously designed observational studies with control groups that adjusted for important confounders. Appendix B provides the study selection criteria in detail.

Our study selection criteria to examine harms (Key Question 3) depended on the outcome of interest. In initial discussions within our research group and in consultation with our technical expert panel, we categorized a prespecified list of harms of interest according to whether the likelihood of the outcome might be substantially different in populations with chronic pain or PTSD. For example, we anticipated that rates of depression and anxiety in patients with chronic pain or PTSD were likely to be substantially different than the general population. In contrast, we thought it unlikely that rates of pulmonary effects or cancer would be particularly influenced by the presence of chronic pain or PTSD. We felt that the incidence of adverse cognitive effects and psychotic symptoms in the general population was likely to provide information that was relevant to chronic pain and PTSD populations, though we recognized that, theoretically, chronic pain and PTSD populations might have a different risk. We chose, therefore, to look more broadly at these outcomes but to report population-specific data where available. In an effort to provide clinicians with at least descriptive information about important harms likely to be related to cannabis use whose incidence and relative risk has not been well-characterized, we also included case series and descriptive studies of these "emerging harms," such as cannabis hyperemesis syndrome and infectious diseases associated with various preparations.

We conducted a primary literature search, but given the broad scope of this review, we summarized data from existing systematic reviews when available to address each question and outcome of interest and then added individual studies meeting inclusion criteria that were published after the end search date of the included review, or were not included in a prior systematic review. We only included reviews that fulfilled key quality criteria: 1) clearly reported their search strategy; 2) reported inclusion and exclusion criteria; and 3) conducted an appraisal of the internal validity of the included trials.¹⁸ If there was more than one review within each category fulfilling these criteria, we prioritized the most recent review and, if there were several recent reviews meeting quality criteria, we prioritized those with the broadest scope. We discussed the ultimate choice of which reviews to include as a group and resolved any disagreements through consensus.

Table 1. PICOTS and Key Questions

Key Question (KQ)	KQ 1. What are the effects of cannabis on health outcomes and healthcare utilization for adults who have chronic pain? KQ 1A: Do the effects differ by patient subgroup, such as patient medical and mental health comorbidities?	 KQ 2. What are the effects of cannabis on health outcomes and healthcare utilization for adults who have PTSD? KQ 2A: Do the effects differ by patient subgroup, such as patient medical and mental health comorbidities? 	KQ 3. What are the harms associated with cannabis use in adults?KQ 3A: Do the harms differ by patient subgroup, such as patient medical and mental health comorbidities?			KQ 4. What are important areas of ongoing research and current evidence gaps in research on cannabis for chronic pain or PTSD, and how could they be addressed by future research?
Population	Adults with chronic pain	Adults with PTSD	Adults (not of	therwise specified)		Adults with chronic pain or PTSD
Interventio n		uding marijuana, hashish, tincture, ha maceutically prepared cannabinoids		•		
Comparato r	Any comparator			. ,		
Outcomes	 Validated measures of pain intensity and pain-related function (including spasticity) Validated measures of pain-related outcomes (mood, depression, anxiety) Validated measures of sleep quality Validated measures of quality of life Utilization of health services Reduction in opioid use or dosage Social functioning/disability/ employment 	 § Validated PTSD clinical interviews and symptom inventories, such as: Clinician Administered PTSD Scale (CAPS), PSTD Checklist (PCL), PTSD Symptom Scale (PSS), Posttraumatic Diagnostic Scale (PDS), <i>etc</i> § Validated measures of mental health symptoms commonly associated with PTSD (mood, depression, anxiety) § Validated measures of sleep quality § Validated measures of sleep quality § Validated measures of quality of life § Utilization of health services § Reduction in benzodiazepine use or dosage § Social functioning/disability/ employment 	General Population	 psychotic population) Cardiovascular events Pulmonary outcomes (eg, forced expiratory volume [FEV1]) Infectious disease complications Mortality Cognitive effects (eg, intelligence quotient [IQ], SLUMS Saint Louis University Mental Status [SLUMS]) 	infections § Cannabinoid hyperemesis syndrome § Other emerging harms	Not applicable
		employment	Chronic	§Other substance	§CUD	

Key Question (KQ)	 KQ 1. What are the effects of cannabis on health outcomes and healthcare utilization for adults who have chronic pain? KQ 1A: Do the effects differ by patient subgroup, such as patient medical and mental health comorbidities? 	KQ 2. What are the effects of cannabis on health outcomes and healthcare utilization for adults who have PTSD? KQ 2A: Do the effects differ by patient subgroup, such as patient medical and mental health comorbidities?	KQ 3. What are the harms associated with cannabis use in adults?KQ 3A: Do the harms differ by patient subgroup, such as patient medical and mental health comorbidities?		KQ 4. What are important areas of ongoing research and current evidence gaps in research on cannabis for chronic pain or PTSD, and how could they be addressed by future research?	
			Pain or PTSD patients	use/substance use disorder §Mental health symptoms (not including psychotic symptoms) including depression, anxiety, <i>etc</i> §Employment §Weight gain §Diversion §Utilization of health services §Insomnia	Withdrawal symptoms	
			Exclude: Ima	aging findings, lab/blood tes	t results.	
Timing	Short- and long-term outcon	nes				
Study design	(randomized or non-random rigorous observational studi control/cohort studies) that a	es with a comparison group (case- adjust for important confounders. narrative reviews, opinions, case	Study designs included for KQ1 and KQ2, plus case series for certain harms (see Outcomes box). <u>Exclude:</u> Non-systematic or narrative reviews, opinions, cross-sectional studies, and individual case reports.			Not applicable

One of 9 investigators examined titles and abstracts for potential relevance to the key questions using Abstrackr.¹⁹ We dual-reviewed a random 5% sample of abstracts in order to ensure reliability between reviewers. Two investigators independently reviewed the full text of all potentially relevant articles for inclusion. Disagreements were resolved through consensus using a third reviewer.

DATA ABSTRACTION

Data from published reports were abstracted into a customized database by one reviewer and confirmed by a second reviewer. From each study, we abstracted the following where available: study design, objectives, setting, population characteristics, subject inclusion and exclusion criteria, number of subjects, duration of follow-up, the study and comparator interventions (formulation, strength, *etc*), important co-interventions, health outcomes, healthcare utilization, and harms.

QUALITY ASSESSMENT

Two reviewers independently assessed the quality of each study (Appendix C). Disagreements were resolved through discussion. To assess the quality of trials we used a tool developed by the Cochrane Collaboration.²⁰ Each trial was given an overall summary assessment of low, high, or unclear risk of bias. To assess the risk of bias of observational studies we considered potential sources of bias most relevant to this evidence base and adapted existing assessment tools.^{21,22} While there are no validated criteria for ranking observational studies, we chose to assign a summary risk of bias rating to represent confidence in each study's results as follows:

- High risk of bias: studies with one or more methodologic deficiencies which would be considered "fatal flaws"; in other words, an answer of "no" to the question: "Are study results believable, taking study limitations into consideration?" For example, studies with minimal information about the exposure of interest would be considered as having a high risk of bias.
- Medium risk of bias: studies that had important methodologic deficiencies that were not fatal flaws, but should be considered when weighing the strength of evidence. For example, recall bias is an inherent limitation to case-control studies that is important to consider in this evidence base.
- Low risk of bias: studies that had no or minor methodologic deficiencies and reflect the strongest observational study designs.

DATA SYNTHESIS

We qualitatively synthesized the evidence on the benefits and harms of cannabis. For the subgroup of neuropathic pain studies, we conducted a study-level meta-analysis of the proportion of patients experiencing clinically significant (\geq 30%) pain relief (Appendix D) using the profile-likelihood random-effects model²³ to combine risk ratios (RRs). We assessed the magnitude of statistical heterogeneity among the studies using the standard Cochran's chi-square test the I^2 statistic.²⁴ All analyses were done using Stata/IC, version 13.1 (StataCorp).

RATING THE BODY OF EVIDENCE

We assessed the overall strength of evidence for outcomes using a method developed for the Agency for Healthcare Research and Quality's (AHRQ) Evidence-based Practice Centers (EPCs).²⁵ The AHRQ EPC method considers study limitations, directness, consistency,



precision, and reporting bias to classify the strength of evidence for individual outcomes independently for randomized controlled trials (RCTs) and observational studies, with supplemental domains of dose-response association, plausible confounding that would decrease the observed effect, and strength of association, as well as separate guidance for applicability.²⁶ Ratings were based on the following criteria:

- High = Very confident that the estimate of effect lies close to the true effect for this outcome. The body of evidence has few or no deficiencies, the findings are stable, and another study would not change the conclusions.
- Moderate = Moderately confident that the estimate of effect lies close to the true effect for this outcome. The body of evidence has some deficiencies and the findings are likely to be stable, but some doubt remains.
- Low = Limited confidence that the estimate of effect lies close to the true effect for this outcome. The body of evidence has major or numerous deficiencies (or both). Additional evidence is needed before concluding either that the findings are stable or that the estimate of effect is close to the true effect.
- Insufficient = No evidence, unable to estimate an effect, or no confidence in the estimate of effect for this outcome. No evidence is available or the body of evidence has unacceptable deficiencies, precluding reaching a conclusion.

PEER REVIEW

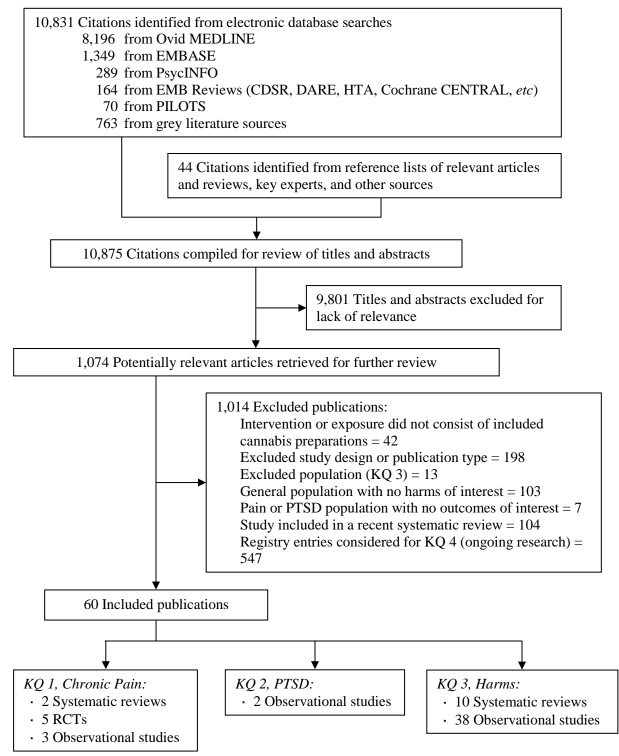
A draft version of this report was reviewed by 8 individuals with technical expertise and clinical leadership. Their comments and our responses are presented in Appendix D.

RESULTS

LITERATURE FLOW

We included 12 systematic reviews and 48 primary studies after reviewing 10,875 titles and abstracts (Figure 1).

Figure 1. Literature Flow Diagram





KEY QUESTION 1: What are the effects of cannabis on health outcomes and healthcare utilization for adults who have chronic pain?

KEY QUESTION 1A: Do the effects differ by patient subgroup, such as patient medical and mental health comorbidities?

Summary of Findings

In this systematic review of the literature, we found limited evidence on the potential benefits and harms of cannabis use in chronic pain populations. We found low-strength evidence that cannabis preparations with precisely defined THC:CBD content (most in a 1:1 to 2:1 ratio) have the potential to improve neuropathic pain but insufficient evidence in other patient populations. Most studies are small, many have methodologic flaws, and the long-term effects are unclear given the brief follow-up duration of most studies. The applicability of these findings to current practice may be low, in part because the formulations studied may not be reflective of what most patients are using, and because the consistency and accuracy of labeled content in dispensaries are uncertain.

Two recent systematic reviews examined the efficacy of cannabis and cannabinoids for the treatment of chronic pain,^{14,15} and reported mixed findings for the management of various chronic pain symptoms related to conditions such as MS, fibromyalgia, peripheral and central neuropathy, human immunodeficiency virus (HIV), rheumatoid arthritis, and cancer. Specifically, across a subset of 8 trials (N=1370) that evaluated non-synthetic cannabinoids (THC or nabiximols), cannabis treatments were associated with a non-significant trend toward benefit (proportion showing greater than 30% reduction in pain: 37% versus 31%; odds ratio [OR] 1.41; 95% confidence interval [CI], 0.99 to 2.00]) compared to placebo and no difference in quality of life among groups.¹⁴ While the authors concluded that there is low- to moderatestrength evidence supporting efficacy of cannabis in chronic pain (limited mainly to MS or neuropathic pain), a separate group reviewed and re-analyzed a similar set of published articles, and determined that there is insufficient to low-strength evidence examining the use of medical cannabis to treat chronic non-cancer pain.¹⁵ Our own interpretation of the evidence is consistent with the latter review because the vast majority of the trials cited in support of a moderatestrength evidence rating were methodologically flawed. Both reviews found insufficient evidence examining the use of medical cannabis for pain related to other conditions such as cancer, rheumatoid arthritis, and musculoskeletal pain.

While the prior reviews included the pharmaceutical, synthetic prescription medications dronabinol and nabilone, studies of these drugs did not contribute substantially to the body of evidence for chronic pain. There was only one small study with high risk of bias examining the effects of nabilone in chronic pain.

We included eligible trials identified by the prior reviews, and found an additional 8 studies²⁷⁻³⁴ that met our inclusion criteria and were not included in the prior reviews. Those additional studies included patients with pain related to MS (4 studies) and mixed pain-related conditions (4 studies). Table 2 presents the overall findings of studies that examined pain and other outcomes in patients with chronic pain. Table 3 presents the findings of RCTs that reported pain outcomes.

No studies directly compared effects according to patient comorbidity. Rather, we describe detailed findings according to patient subgroup below.



Table 2. Studies of the Overall Effects of Cannabis in Patients with Chronic Pain

Study, setting, design (N patients) Risk of bias (ROB)	Sample description Mean age (SD) % male	Intervention and comparator	Primary findings	Adverse effects	
Multiple sclerosis (N	IS)				
Notcutt 2012 ²⁹ UK, 5 sites RCT (N=36) Unclear ROB	Age 57 100% Caucasian 41.7% male MS: 16.4 years Spasticity: 12.7 years Nabiximols use: 3.6 years; Subjects experienced ongoing benefit with nabiximols. Mean daily dose of nabiximols: 8.25 sprays Mean baseline scores, Treatment vs placebo: Spasticity score on NRS: 3.6 (SD=1.7) vs 4.13 (SD=2.2), Disability scale (EDSS): 6.75 vs 6.92	T: Nabiximols (oromucosal spray delivering 2.7 mg THC/2.5 mg CBD), mean daily dose 7.7 sprays. C: Placebo, mean daily dose 9.0 sprays	 4-week treatment period Pain: NR Spasticity: no differences (P>.1) between groups on NRS score. Treatment failure, defined as cessation of nabiximols use, worsening of spasticity, or increase in anti-spasticity meds: 44% of nabiximols group vs 94% of placebo group (hazard ratio for failure in placebo group 0.335, 90% CI: 0.162-0.691, P=.013 in favor of nabiximols group). Other: No differences in sleep disruption NRS score, Modified Ashworth Scale, Timed 10-meter walk test, or Motricity Index, CGIC ease of transfer; statistically significant improvement in nabiximols vs placebo group on SGIC (OR 4.55, 90% CI: 1.59- 14.00, P=.017) and CGIC general function scores (OR 18.55, 90% CI: 3.94-118.77, P=.001). 	During treatment period, 83% (15/18) on nabiximols and 78% (14/18) on placebo had treatment-related AEs, most commonly pain (2 vs 5), spasticity (2 vs 3), muscle spasm (4 vs 4), and depressed mood (0 vs 2); 4 participants had severe AEs (2 vs 2).	

Study, setting, design (N patients) Risk of bias (ROB)	Sample description Mean age (SD) % male	Intervention and comparator	Primary findings	Adverse effects	
Novotna 2011 ²⁷ Europe, 51 sites RCT (N=241) Low ROB	Age 48.6 40% male 100% white/Caucasian, 18% with previous cannabis use in last year, MS years: 12.6 Spasticity years: 7.7 Mean baseline spasticity score on NRS 7.0 To qualify for the RCT, patients must have had at least a 20% reduction in spasticity NRS score with initial exposure to nabiximols.	T = Nabiximols (oromucosal spray delivering 2.7 mg THC/2.5 mg CBD). C = Placebo oromucosal spray. Maximum permitted dose was 12 sprays in any 24 hour period.	Pain: NR Spasticity: Change in mean NRS score at 12 weeks: -0.84 (95% Cl, - 1.29 to -0.40), <i>P</i> =.0002 % with at least 30% improvement, T vs C: 74% vs 51%; OR 2.73 (95% Cl, 1.59-4.69), <i>P</i> =.0003. Other: Nabiximols were significantly superior (<i>P</i> <.05) to placebo for sleep disruption, Barthel Activities of Daily Living, Physician Global Impression of Change, Subject Global Impression of Change, and Carer Global Impression of Change in Function.	No difference between groups; no AEs occurred in > 10% in either group. Most common AEs were vertigo, fatigue, muscle spasms, and urinary tract infections.	
Ungerleider 1987 ²⁸ US, single site Double-blind, placebo-controlled, crossover clinical trial (N=13) High ROB	Age 48.3 39% male 53% wheelchair bound 60% with prior cannabis use	T (THC) or C (placebo) for 5 days, followed by 2 day wash-out and 5 day trial with crossover drug. Patients were initiated at varying oral doses of THC (range: 2.5 to 7.5 mg in first paired trial). If patient had inadequate relief, they could be re-randomized and started at a higher dose (increased by 2.5 mg to maximum 15 mg).	Pain: NR Spasticity: self-report on scale of 1 to 5, where 5=more) was lower with T: 2.2 (SD=0.9) vs C: 3.4 (SD=0.7), P=.03; improvement started at 7.5 mg dose. No change from baseline on physician ratings on all measures (limb weakness, limb spasticity, limb coordination, gait impairment, reflexes; all P -values > 0.05).	No difference in AEs for 7.5 mg THC vs C. AEs were more frequent and less tolerable with higher doses of THC. Common AEs: weakness, dry mouth, dizziness, relaxation, mental clouding, short term memory impairment, and spatial- time distortions.	

Study, setting, design (N patients) Risk of bias (ROB)	Sample description Mean age (SD) % male	Mean age (SD)		Adverse effects	
Wade 2003 ³⁰ UK, single site outpatient clinic Pilot study: double- blind, placebo- controlled single- patient cross-over RCT (N=24) Low ROB	Age 48 50% male Types of pain: MS (n=14) Brachial plexus damage (n=1) Limb amputation due to neurofibromatosis (n=1) Target symptoms: pain (n=13), muscle spasm (n=17), spasticity (n=9), impaired bladder control (n=11), tremor (n=8)	Pump-action sublingual spray delivering 2.5 mg T1: CBD T2: THC T3: Both THC and CBD, 1:1 ratio C: Placebo Maximum permitted dose was 120 mg every 24 hours.	Mean (SD) daily VAS (0-100) over last 7 days of each 2-week period, <i>P</i> -value vs C: Pain: baseline 30.1 (17.8) to T1: 54.8 (22.6), <i>P</i> <.05 T2: 54.6 (27.4), <i>P</i> <.05 T3: 51.3 (27.0), <i>P</i> =NS C: 44.5 (22.7) Spasm: baseline 40.9 (18.5) to T1: 54.6 (19.1), <i>P</i> =NS T2: 58.4 (22.3), <i>P</i> <.05 T3: 55.8 (24.4), <i>P</i> <.05 C: 47.3 (22.6) Spasticity: baseline 29.0 (16.1) to T1: 47.8 (18.5), <i>P</i> =NS T2: 57.3 (22.2), <i>P</i> <.05 T3: 43.8 (15.6), <i>P</i> =NS C: 42.3 (18.1)	AEs reported by 33% in CBD, 55% in THC, 30% in CBD:THC, and 48% in placebo; Common AEs during periods of cannabinoid use included headache (n=5), nausea (n=3), diarrhea (n=4), sleepiness (n=3) fall (n=3); 3 patients withdrew during open-label phase due to one each of intoxication, vasovagal episodes, and sublingual burning sensation; one patient withdrew during the blinded phase due to excess sensitivity to THC; Some patients in all periods took rescue medications.	

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Study, setting, design (N patients) Risk of bias (ROB)	Sample description Mean age (SD) % male	Intervention and comparator	Primary findings	Adverse effects	
Other chronic pain					
Fiz 2011 ³⁴ Spain, single site Retrospective cohort study (N=56) High ROB	Adults with fibromyalgia, with moderate to severe symptoms, and who were resistant to pharmacological treatment. Age 50 years 5% male (users 7%, non- users 4%) Median disease duration: 5.0 years in users, 4.0 years in non-users	T: cannabis use, method of administration: smoking 11%; oral 46%; combined 43%. C: non-users (for QOL comparison) Duration of use: 40% < 1 year 32% 1 to 3 years 29% ≥ 3 years THC/CBD content NR.	Pain: 2 hours post-cannabis use, VAS (100 mm) scores showed significant mean reduction in pain (37.1 mm reduction) and stiffness (40.7 mm reduction), P <.001. Other: Patients used cannabis for almost all symptoms associated with fibromyalgia with no reported worsening of symptoms (strong relief reported by 81% for sleep disorders to 14% for headaches). 68% of patients reported reduction in pharmacological treatment (not otherwise specified) when they started using cannabis. Increased perception of well-being (40.0 mm increase); relaxation (27.6 mm increase), and somnolence scores (20.0 mm increase) were significantly increased from baseline, P <.05; QOL: (SF-36) mental health component summary score was significantly higher in users (mean=29.6, SD =8.2) compared to non-users (mean=24.9, SD=8.9), P<.05; No significant group differences found on SF-36 physical component (P =.53), PSQI (P =.73), FIQ (P =.36).	Dry mouth (61%) Sedation (43%) Dizziness (36%) High (32%) Tachycardia (29%) Conjunctival irritation (25%) Hypotension (21%) No serious AEs reported.	

Study, setting, design (N patients) Risk of bias (ROB)	Sample description Mean age (SD) % male	Intervention and comparator	Primary findings	Adverse effects
Notcutt 2004 ³³ Canada Single site "N of 1"double-blind, placebo-controlled, crossover RCT (N=34) Low ROB	Age 46.7 32% male 100% with chronic pain (mostly neuropathic) 47% with MS	Sublingual spray that delivered 2.5 mg each of: T1: THC T2: CBD T3: both CBD and THC (1:1 ratio), C: 0.1 mL matching placebo; In order to qualify for the study, patients must report benefit during a run-in period; 8-week trial where each week for first 4 weeks they randomly received a different cannabinoid or placebo; at start of each week, patients underwent supervised titration and each preparation was then given in random order over next 4 weeks so that each patient received each cannabinoid or placebo for 2 separate one-week periods; patients administered sprays daily and titrated up to a dose of their choosing depending on onset of side effects and attenuation of pain (range 1 to 8 sprays daily)	Pain: VAS 10 cm Symptom 1 score (median [IQR]) pain reduction: T1: 4.63 (1.74-6.06) T2: 5.45 (3.6-7.4) T3: 4.4 (2.6-5.8) C: 5.9 (2.8-7.3) T1 and T3 both significantly better than C (P <.05, P <.01, respectively) Symptom 2 score (median [IQR]) pain reduction: C: 4.98 (2.61-7.50) T1: 4.08 (1.33-5.43) T2: 5.03 (3.16-6.88) T3: 4.28 (2.33-5.51) T1 and T3 significantly better than placebo (P =.054, P <.001, respectively) 38% (9/24) of patients had a decrease in VAS of 50% or more for either symptom 1 or symptom 2 when using active preparations vs placebo; all 9 patients experienced this with THC and/or THC:CBD and 3 of these patients also had reduction with CBD. Other: Sleep Quality: Percentage of "good" nights during trial period, median (IQR): T1: 42.9% (57.2-35.7) T2: 36.9% (47.9-28.6) T3: 55.4% (78-34.5) C: 17.0% (35.7-3.6) T1, T2, and T3 were each significantly better than placebo (P <.001, P <.001, P <.05, respectively).	Side effects: Most commonly drowsiness, euphoria/dysphoria, and dry mouth; hallucination in one patient; vasovagal in one patient; change in neural function in 2 patients (return of absent ankle reflex, return of touch sensation to dermatome)

Study, setting, design (N patients) Risk of bias (ROB)	Sample description Mean age (SD) % male	Intervention and comparator	Primary findings	Adverse effects	
Storr 2014 ³² Canada, outpatient GI clinic Retrospective cohort study (N=313) High ROB	Adults with IBD Age 39.6 years (non-users 40.2, users 36.6 years) 31% male (27.4% non- users; 50.0% users) Note: Significant between group differences in race, household income, and education level (<i>P</i> <.05) Mean time since IBD diagnosis was 13.9 (range: 1 to 40) years in users, 13.2 (range: 1 to 43) years in nonusers; Among users vs non-users, 75.0% vs 71.9% had Crohn's Disease, 17.9% vs 20.2% had ulcerative colitis, 7.1% vs 8.0% had indeterminate colitis. Note: Significant between- group difference in type of disease (<i>P</i> =.035)	Patients self-reported cannabis use; varied between smoking (95%), oral (9%) and drinking (5%); no info provided about dose or frequency Comparator: non-users (<i>ie</i> , those who did not endorse cannabis use for treatment of IBD)	Risk of surgery for those with Crohn's Disease was significantly associated with cannabis use for at least 6 months vs never use (OR 5.03; 95% Cl, 1.45-17.46) after controlling for multiple factors; Intermittent use was not associated with higher surgery rates vs never use (OR 1.28; 95% Cl, 0.31-5.27). Risk of hospitalization for IBD was not associated with cannabis use for at least 6 months (OR 2.86; 95% Cl, 0.96-8.46) or intermittent (OR 1.99; 95% Cl, 0.41-9.73) cannabis use vs never use	Most cannabis users experienced side effects like anxiety, increased appetite, dry mouth, drowsiness, and a "high" (75% of users); generally rated as mild in severity; 19.6% reported that they needed a "high" to get symptom improvement while remainder did not	

Study, setting, design (N patients) Risk of bias (ROB)	Sample description Mean age (SD) % male	Intervention and comparator	Primary findings	Adverse effects	
Ware 2015 ³¹ Canada, 7 sites Prospective cohort (N=431) Low ROB	Age: 49.0 (cannabis vs control: 45.5 vs 52.4) 43.1% male (cannabis vs control: 51.2% vs 35.2%) Groups differed significantly for age and gender (<i>P</i> <.001) Type of pain, cannabis vs control: Nociceptive 16.3% vs 18.1% Neuropathic 38.6% vs 32.4% Both 45.1% vs 49.5% Mean pain intensity 6.6 (range: 0 to 10) vs 6.1	T: Cannabis contained 12.5 ± 1.5% THC; max of 5 g/day; median daily dosing was 2.5 g/day Patients used any delivery system that they were comfortable with (27% smoked, 61% combined smoking, oral, and vaporization, 8% consumed orally) C: Non-cannabis users	Pain: greater reduction in pain intensity noted in cannabis users: VAS (0-10 pain intensity over last 24 hours, mean (SD): T: 5.54 (2.11) C: 6.10 (2.13) Difference = 1.10 (95% Cl, 0.72- 1.56) Significant reduction in average pain intensity over 1 year with T (change=0.92; 95% Cl, 0.62-1.23) but not C (change=0.18; 95% Cl, - 0.13-0.49) Other: Mood: POMS (total mood disturbance): Cannabis = 23.92 (SD 19.04); Control = 27.09 (SD 21.29), fixed regression coefficient (-5.52, P=.0060; higher scores equal more mood disturbance) QOL: SF-36. Improvement of physical function among cannabis users at 1 year (1.62 points higher; 95% Cl, 0.10-3.14); No between or within group differences on mental component.	T vs C: Serious AEs: no sig. difference, 13% vs 19%; 40 vs 56 events Adjusted IRR (95% CI) for event = 1.08 (0.57-2.04) Most common AEs: surgical/medical procedures 25% vs 20% GI disorders 25% vs 13% Most common serious AEs in cannabis group: -abdominal pain (n=3, 12%), -intestinal obstruction (n=3, 12%) -nephrolithiasis (n=3, 12%) -2 withdrawals from treatment due to serious side effects (1 convulsion, 1 alcohol problem); Cannabis users had significantly higher number/rate of non- serious AEs (T vs C: 818 vs 574 events), adjusted IRR for event = 1.73; 95% CI, 1.42-2.14); Most common AEs, cannabis group: -nervous system: n=165 (20%) -gastrointestinal: n=109 (13.4%) -respiratory: n=103 (12.6%); Cannabis group had significantly higher rates, unadjusted IRR (95% CI): nervous system disorders 2.05 (1.46-2.86); respiratory disorders 1.77 (1.16- 2.70); infections disorder 1.51 (1.04-2.20); and psychiatric disorders 2.74 (1.45-5.18) vs control group. No significant between group differences were found in pulmonary or neurocognitive function.	

Abbreviations: AE = adverse event; C = control/comparator group; CBD = cannabidiol; CGIC = Carer Global Impression of Change; CI = confidence interval; EDSS = Expanded



Disability Status Scale; FIQ = Fibromyalgia Impact Questionnaire; GI = gastrointestinal; IBD = inflammatory bowel disease; IQR = interquartile range; IRR = incidence rate ratios; MS = multiple sclerosis; N = number; NR = not reported; NRS = Numeric Rating Scale; NS = not significant; OR = odds ratio; POMS = Profile of Mood States; PSQI = Pittsburgh Sleep Quality Index; QOL = quality of life; RCT = randomized controlled trial; ROB = risk of bias; SD = standard deviation; SF-36 = 36-Item Short Form Health Survey; SGIC = Subject Global Impression of Change ; T = treatment group; THC = tetrahydrocannabinol; UK = United Kingdom; US = United States; VAS = Visual Analogue Scale.

Footnotes on concomitant therapy:

- Fiz 2011: Participants continued their current pharmacologic regimen; at baseline (users vs non-users), analgesic/anti-inflammatory drugs used by 75% vs 64%, antidepressants used by 50% vs 61%, anxiolytics used by 36% vs 36%, opioids used by 21 vs 39%, myorelaxants used by 4% vs 21%, hypnotics used by 18% vs 29%.
- Notcutt 2004: Patients maintained their regular medications and were allowed to use non-cannabinoid medication for breakthrough pain as long as they documented it (n=7 patients used rescue THC:CBD during trial).
- Notcutt 2012: Participants maintained other medications at stable doses: 16% taking baclofen; 16% taking benzodiazepines; 16% taking analgesics and antipyretics; 12% taking quinine or derivatives; 3% taking antiepileptics; 3% taking amantadine; 3% taking herbal supplements.
- Novotna 2011: Antispasticity agents and/or disease-modifying medications were maintained at a stable dose for 30 days prior to and throughout the study. 13% taking adamantane derivatives, 22% taking benzodiazepine derivatives, < 0.5% taking dantrolenes, < 0.5% taking naltrexone, 24% taking antiepileptics, 73% taking centrally-acting medications, 58% taking baclofen, 17% taking tizanidine, 17% taking tolperisone, 1% taking "other" medications.
- Storr 2014: Patients continued all other prescribed medications; 35.7% taking aminosalicylates, 42.6% taking steroids, 41.4% taking immunomodulators, 37.9% taking analgesics, 24.8% taking narcotics, 17.2% taking loperamide, 32.0% taking biologicals, 9.7% taking IV medications, 32.0% taking complimentary and alternative medicine.
- Wade 2003: Patients continued current medication regimen and were asked not to use any other cannabis.
- · Ware 2015: Patients continued pharmacotherapy (opioids, antidepressants, and anticonvulsants).

Table 3. Characteristics and Findings^a of RCTs of the Effects of Cannabis Extracts on Pain Outcomes

Triol	Pain Type	N	Intervention	Formulation; Dosage; Duration ≥30% Pair	Patients Achieving	Mean Differei in Change Fro		Overall
Trial					≥30% Pain Reduction, - T vs C, n/N <i>(%)</i>	NRS Pain Scale, points ^b	VAS Pain Scale, mm ^c	 Risk of Bias
Abrams, 2007 ³⁵	Neuropathic sensory, HIV- associated	55	Smoked THC, 4%; 1 cigarette/d (0.9 g)	12 d	13/25 vs 6/25 (52.0 vs 24.0)	_	_	Low
Berman, 2004 ³⁶	Neuropathic brachial plexus avulsion	48	Nabiximols (THC oromucosal spray); ≤ 48 sprays/d; crossover	2 wk (no washout)	_	_	_	Low
Ellis, 2009 ³⁷	Neuropathic sensory, HIV- associated	34	Smoked THC, started at 4% and adjusted as necessary; 4 smoking sessions/d; crossover	5 d (2-wk washout)	_	_	-	Low
Lynch, 2014 ³⁸	Neuropathic chemotherapy- induced	18	Nabiximols; ≤12 sprays/d	4 wk (2-wk washout)	_	_	_	Low
Notcutt, 2004 ³³	Mostly neuropathic; 47% MS	34	Sublingual spray delivering 2.5-mg THC, 2.5-mg CBD, or 2.5 mg each; 1 to 8 sprays/d	8 wk	THC: 9/24 vs NR (37.5 vs NR) CBD: 3/24 vs NR (12.5 vs NR) THC+CBD: 9/24 vs NR (37.5 vs NR)	_	_	Low
Nurmikko, 2007 ³⁹	Neuropathic pain with allodynia	125	Nabiximols; ≤48 sprays/d	5 wk	16/63 vs 9/62 (25.4 vs 14.5)	-	-8.03 (-13.83 to -2.23)	High
Selvarajah, 2010 ⁴⁰	Neuropathic diabetic peripheral	30	Nabiximols; maximum unclear	12 wk	8/15 vs 9/14 (53.3 vs 64.3)	_	9.50 (-11.30 to 27.80)	Unclear
Serpell, 2014 ⁴¹	Neuropathic peripheral with allodynia	246	Nabiximols; ≤24 sprays/d	15 wk	34/123 vs 19/117 (27.6 vs 16.2)	-0.34 (-0.79 to 0.11)	−2.86 (−7.22 to 1.50)	Low

Evidence-based Synthesis Program

Trial Pain Type		N	Intervention Formulation; Dosage;	Duration	Patients Achieving	Mean Differe in Change Fro		Overall – Risk of
Triai	Study Design	Duration	≥30% Pain Reduction, - T vs C, n/N <i>(%)</i>	NRS Pain Scale, points ^b	VAS Pain Scale, mm ^c	Bias		
Wallace, 2015 ⁴²	Neuropathic diabetic peripheral	16	Vaporized THC, 7%, 4%, or 1%; 4 h observation at each dose; crossover	4 h (2-wk washout)	1% THC: 10/16 vs 10/16 (62.5 vs 62.5) 4% THC: 12/16 vs 10/16 (75.0 vs 62.5) 7% THC: 13/16 vs 10/16 (81.3 vs 62.5)	_	_	Low
Ware, 2010 ⁴³	Neuropathic, postsurgical or posttraumatic	23	Smoked THC, 2.5%, 6%, or 9.4%; crossover	5 d (9-d washout)	_	-	_	Low
Wilsey, 2008 ⁴⁴	Neuropathic	38	Smoked THC, 3.5% or 7%; 9 puffs; crossover	6 h (3- to 21- d washout)	3.5% THC: 4/36 vs 2/33 (11.1 vs 6.1) 7% THC: 0/34 vs 2/33 (0.0 vs 6.1)	_	_	Low
Wilsey, 2013 ⁴⁵	Neuropathic, peripheral	39	Vaporized THC, 1.29% or 3.53%; 4 puffs at 1 h after baseline, 4 to 8 puffs at 3 h; crossover	6 h (3- to 7-d washout)	1.29% THC: 21/37 vs 10/38 (56.8 vs 26.3) 3.53% THC: 22/36 vs 10/38 (61.1 vs 26.3)	_	1.29% THC: -11 3.53% THC: -10	Low
Wilsey, 2016 ⁴⁶	Neuropathic, spinal cord injury	42	Vaporized THC, 2.9% or 6.7%; 400 mg using Foltin Puff Procedure at 8 to 12 puffs over 240 min, adaptable dose design	8 h	2.9% THC: 18/26 vs 8/18 (69.2 vs 44.4) 6.7% THC: 31/35 vs 8/18 (88.6 vs 44.4)	_	_	Low
Collin, 2010 ⁴⁷	MS	337	Nabiximols; ≤ 24 sprays/d	14 wk	_	_	-	Unclear
Corey- Bloom, 2012 ⁴⁸	MS	37	Smoked THC, 4%; one 800-mg cigarette	3 d (11-d washout)	_	_	_	Unclear
Langford, 2013 ⁴⁹	MS	339	Nabiximols; ≤12 sprays/d	14 wk	84/167 vs 77/172 (50.3 vs 44.8)	0.17 (-0.62 to 0.29)	-	Unclear
					<u></u>		144	

Evidence-based Synthesis Program

Trial Pain Type		NI	Intervention	Duration	Patients Achieving	Mean Differe in Change Fro		Overall
TTA	Pain Type	Ν	Formulation; Dosage; Study Design	Duration	≥30% Pain Reduction, T vs C, n/N <i>(%)</i>	NRS Pain Scale, points ^b	VAS Pain Scale, mm ^c	 Risk of Bias
Rog, 2005 ⁵⁰	MS	66	Nabiximols; ≤ 48 sprays/d	5 wk	_	-1.25 (-2.11 to -0.39)	−6.58 (−12.97 to −0.19)	Low
Van Ameron- gen, 2017 ⁵¹	MS	24	Orally ingested THC, 99% (EPC002A, Namisol); 1.5 or 5 mg 3 times/d	2 wk	_	Week 2: -1.09 (-1.98 to -0.20) (<i>P</i> = 0.018) Week 4: -0.85 (-1.74 to -0.04) (<i>P</i> = 0.061)	_	Unclear
Wade 2003 ³⁰	MS (67%)	24	Pump-action sublingual spray delivering 2.5-mg THC, 2.5-mg CBD, or 2.5 mg each; ≤120 mg/d; crossover	2 wk (no washout)	_	_	Baseline: 30.1 (SD, 17.8) 2 nd week of each group: CBD: 54.8 (SD, 22.6; <i>P</i> < 0.05) THC: 54.6 (SD, 27.4; <i>P</i> < 0.05) THC+CBD: 51.3 (SD, 27.0; <i>P</i> = NS) Placebo: 44.5 (SD, 22.7)	Low
Wade, 2004 ⁵²	MS	160	Nabiximols; ≤ 48 sprays/d	6 wk	_	_	-	Unclear
Zajicek, 2003 ⁵³	MS	657	THC/CBD capsules; ≤ 25 mg/d	15 wk	_	_	-	High
Zajicek, 2012 ⁵⁴	MS	279	THC/CBD capsules; ≤ 25 mg/d	12 wk	_	_	_	Low
Johnson, 2010 ⁵⁵	Cancer	119	Nabiximols; ≤ 8 sprays/d	2 wk	23/53 vs 12/56 (43.4 vs 21.4)	-0.32 (-0.86 to 0.22)	-	Unclear
		117	THC oromucosal spray	2 wk	12/52 vs 12/56 (23.1 vs 21.4)	-0.67 (-1.21 to -0.14)	-	
Noyes, 1975 ⁵⁶	Cancer	10	THC capsules; 5, 10, or 15 mg; crossover	1 d (no washout)	_	_	_	High

Evidence-based Synthesis Program

		N	Intervention	Duration	Patients Achieving	Mean Differen in Change Fro		Overall
Trial	Pain Type	Ν	Formulation; Dosage; Study Design	Duration	≥30% Pain Reduction, T vs C, n/N <i>(%)</i>	NRS Pain Scale, points ^b	VAS Pain Scale, mm ^c	 Risk of Bias
Portenoy, 2012 ⁵⁷	Cancer	360	Nabiximols; 1 to 4, 6 to 10, or 11 to 16 sprays/d	9 wk	1 to 4 sprays: 30/91 vs 24/91 (33.0 vs 26.4) 6 to 10 sprays: 26/87 vs 24/91 (29.9 vs 26.4) 11 to 16 sprays: 22/90 vs 24/91 (24.4 vs 26.4)	1 to 4 sprays: −0.75 (−1.28 to −0.22) 6 to 10 sprays: −0.36 (−0.89 to 0.18) 11 to 16 sprays: −0.09 (−0.62 to 0.44)	_	Unclear
De Vries, 2016 ⁵⁸	Abdominal pain (includes chronic pancreatitis, postsurgical pain)	65	Orally ingested THC, 99% (EPC002A, Namisol); step-up phase: days 1 to 5, 3 mg 3 times/d; days 6 to 10, 5 mg 3 times/d; stable dose phase: days 11 to 52, 8 mg 3 times/d	7 wk	_	-1.6 (SD, 1.78) vs -1.9 (SD, 2.18) (<i>P</i> = 0.92)	_	High
Blake, 2006 ⁵⁹	Rheumatoid arthritis	58	Nabiximols; ≤48 sprays/d	5 wk	_	_	-3 (-18 to 9)	Unclear

Detailed Findings According to Patient Subgroup

Multiple Sclerosis (MS)

Two prior systematic reviews and 4 additional published trials examined the effects of cannabisbased preparations on pain and spasticity in patients with MS. Overall, there is low-strength evidence to support cannabis-based treatments for the potential to improve pain, spasticity, and sleep in select populations with MS, but results were inconsistent across studies. The body of evidence is limited by the paucity of methodologically rigorous studies, inconsistent findings across studies, the lack of long-term outcomes, and the small number of patients included in many trials. Moreover, the largest low risk of bias trial used restrictive entry criteria which may reduce the applicability of the evidence to broader populations.

A recent systematic review included 11 (2,653 participants) trials examining the use of cannabis preparations compared with placebo (it also included studies of synthetically produced cannabinoids which are not covered in our review).¹⁴ The authors of this review found low- to moderate-strength evidence mostly from trials of nabiximols on spasticity in MS. However, the findings were mixed with evidence of no effect on some spasticity related outcomes and small effects on others. Moreover, 9 of 11 trials had high or unclear risk of bias; only 2 of the trials were found to be at low risk of bias.

One RCT analyzed data from 414 patients from 33 outpatient neurology and rehabilitation centers in the United Kingdom (UK).⁵³ Patients were randomized to cannabis extract (containing 2.5 mg THC) and matched placebo capsules. The study had a 5-week dose titration phase and a 10-week maintenance phase; the maximum allowable dose was 25 mg daily. The study results did not identify a significant effect on mean change in spasticity between groups (mean changes in groups were 1.24 and 0.92 for cannabis extract and placebo, respectively). On secondary outcome measures, there were no differences in timed 10-minute walk test, self-reported mobility, disability score, or general health. Participants randomized to cannabis extract had a greater likelihood of self-reported improvement on 3 of 9 symptom categories (including spasticity, pain, and spasms).

In a study of 277 patients with MS, patients were randomized to cannabis extract (contained 2.5 mg THC) and matched placebo capsules.⁵⁴ The study had a 2-week dose titration phase and a 10-week maintenance phase; the maximum allowable dose was 25 mg. The proportion of patients who achieved significant relief from muscle stiffness was 29.4% in the cannabis group versus 15.7% in the placebo group (OR 2.26; 95% CI, 1.24 to 4.13; P = .004, one-sided). Secondary analyses were also in favor of the cannabis group, as patients reported improvements in body pain, muscle spasms, and sleep quality.

Another systematic review focused on non-cancer pain treatment and covered literature over the same time frame. This review differed in that it intentionally re-analyzed data excluding unpublished studies (most of which were industry-funded). They identified 4 studies (510 participants) examining the efficacy of cannabis preparations for patients with pain related to MS (2 other studies examined synthetically produced cannabinoids, which are not part of our review).¹⁵ The authors concluded that there was low-strength evidence showing no significant difference between cannabis preparations and placebo in improving pain in patients with MS.

We identified an additional 4 trials (314 participants) examining cannabinoids to treat spasticity and/or pain in patients with MS.²⁷⁻³⁰ Two studies were rated as low risk of bias,^{27,30} one was at high risk of bias,²⁸ and one was unclear.²⁹ In a large multicenter European trial with low risk of



bias (N=241), patients with MS and moderately severe spasticity were randomized to open-label nabiximols or placebo if they initially experienced at least a 20% reduction in spasticity Numeric Rating Scale (NRS) during an open-label nabiximols run-in period. Over half (52.2%) of participants failed to meet this criteria and were not enrolled. Active treatment consisted of nabiximols, containing 2.7 mg THC and 2.5 mg CBD delivered via oromucosal spray. Participants self-titrated their dose; the maximum permitted dose was 12 sprays in any 24 hour period. The intervention lasted for 12 weeks, with the final follow-up visit 2 weeks after treatment completion. The intervention group experienced a significant reduction in mean spasticity score from baseline to end of treatment compared with the placebo group (change in mean NRS score -0.84 [95% CI, -1.29 to -0.40]). The number of responders (defined as at least a 30% improvement in spasticity from baseline) was significantly higher in treatment versus placebo (74% versus 51%; OR 2.73; 95% CI, 1.59 to 4.69). The study medication was also superior to placebo for 6 of 15 secondary outcomes.

The remaining 3 trials revealed mixed findings. In a 5-day treatment study, patients with MS treated with THC 7.5 mg had no significant differences in any outcome (limb weakness, limb spasticity, limb coordination, gait impairment, reflexes) based on physician rating, though patient self-reported spasticity was lower when on THC versus placebo when doses were 7.5 mg or higher.²⁸ In a double-blind cross-over trial with 20 patients with MS or other neurological diagnosis, participants received each of THC, CBD, THC and CBD, and placebo for 2 weeks in randomized order.³⁰ Study findings were mixed: pain relief assessed with a Visual Analog Scale (VAS) was improved for both the THC and CBD groups relative to placebo, but not the group receiving THC and CBD combined; spasm VAS score improved following use of THC and combined THC and CBD; spasticity improved for THC only; and no significant improvements were seen in coordination or bladder control. Study medications, relative to placebo, were not consistently associated with significant treatment benefit on other secondary outcome measures. In a 5-site study of 36 patients who demonstrated a positive response to nabiximols during an open-label phase, participants were randomized to 4 weeks of continued nabiximols use or placebo.²⁹ Those randomized to placebo were more likely than participants randomized to nabiximols to demonstrate a treatment failure (defined as increase in spasticity, addition of antispasticity medicine, or treatment drop-out): treatment failure was observed in 44% of the nabiximols group versus 94% of the placebo group (hazard ratio [HR] 0.335; 90% CI, 0.162 to 0.691). Findings on secondary outcomes were mixed. The risk of bias from this trial is unclear, as it was underpowered and participants who withdrew from the trial may have returned to taking other medications before returning for formal study withdrawal visit.

Neuropathic Pain

Thirteen trials examined the effects of cannabis-based preparations on neuropathic pain (Table 3). Participants had central or peripheral neuropathic pain related to various health conditions. Of these studies, 11 trials were determined to be at low ROB,^{33,35-38,41-46} 1 as having unclear ROB,⁴⁰ and 1 as having high ROB.³⁹ Overall, we found low-strength evidence that cannabis may improve pain in some patients with neuropathic pain. Studies generally did not find clinically significant differences on continuous pain scales between groups, but a higher proportion of intervention patients experienced clinically significant pain relief at up to several months of follow-up. In a meta-analysis of nine studies that reported \geq 30% pain reduction, intervention patients were more likely to report improvement in pain (RR 1.43, 95% CI 1.16 to 1.88; I²=38.6%, p = 0.111; Figure 2). Most studies were small, few reported outcomes beyond 2 to 3 weeks, and none reported long-term outcomes.



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Figure 2. Odds of achieving \geq 30% pain reduction with cannabis compared to placebo in trials of patients with neuropathic pain

Study	Intervention	Duration		RR (95% CI)	Events, Treatment	Events, Control
Nurmikko 2007	Nabiximols	5 wks		1.75 (0.84, 3.66)	16/63	9/62
Selvarajah 2010	Nabiximols	12 wks —	•	0.83 (0.45, 1.53)	8/15	9/14
Langford 2013	Nabiximols	14 wks	- H	1.12 (0.90, 1.41)	84/167	77/172
Serpell 2014	Nabiximols	15 wks		1.70 (1.03, 2.81)	34/123	19/117
Wallace 2013	THC oromucosal	4 hrs -		1.20 (0.75, 1.93)	12/16	10/16
Abrams 2007	THC smoked	12 days		2.17 (0.98, 4.79)	13/25	6/25
Wilsey 2011	THC smoked	6 hrs	+	1.83 (0.36, 9.36)	4/36	2/33
Wilsey 2013	THC vaporized	6 hrs	∔_ ■	2.32 (1.28, 4.20)	22/36	10/38
Wilsey 2016	THC vaporized	8 hrs		1.72 (1.16, 2.55)	31/42	18/42
Overall (l ² = 38.6%,	p = 0.111)		\Diamond	1.43 (1.16, 1.88)	224/523	160/519
		.25	1 4			
	← F	avors placebo	Favors can	nabis →		

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In the largest RCT, 246 patients with peripheral neuropathic pain self-titrated nabiximols up to a maximum allowable dose of 24 sprays/day or received a placebo.⁴¹ Those who completed the study (79 nabiximols and 94 placebo) and responded positively to the intervention demonstrated a significant decrease in pain (OR 1.97, 95% CI 1.05 to 3.70). However, among all participants, including those who did not have an intervention response, the reduction in the NRS pain scale did not reach clinical or statistical significance. The second-largest low ROB RCT included 55 patients with HIV-associated sensory neuropathy who were randomized to smoke either 3.56% THC cigarettes or placebo 3 times daily for 5 days. Among those who completed the study, 52% (n=13) of the treatment group demonstrated a clinically significant (> than 30%) reduction in pain compared to 24% (n=6) in the placebo group.³⁵

A one-year prospective-cohort study (n=431) among patients with nociceptive and neuropathic chronic non-cancer pain provides information about long-term treatment effects.³¹ Cannabis users experienced a reduction in average pain intensity (VAS) that was stable across 4 time points over a one-year period among cannabis users, but the change was small and not clinically significant (0.92 change, 95% CI 0.62 to 1.23).

Other/Mixed Pain Conditions

Overall, there are a limited number of studies of patients with chronic pain that are not related to MS or neuropathy. Generally, the evidence is inconsistent and of low quality. As noted above in the prior systematic reviews, there were 2 studies with unclear risk of bias which both included patients with cancer-related pain (described more below); 3 other studies had a high risk of bias (and are not summarized here).^{14,15} We found only 2 additional studies, one low risk of bias RCT³³ and one observational study (N=465) (Table 2).³¹

Of the additional studies, the best evidence for the treatment of mixed pain conditions comes from a randomized, double-blind, placebo-controlled, crossover trial that was conducted in the UK among 34 patients with various pain conditions, 47% of whom were diagnosed with MS.³³ Participants were each administered 3 different medicinal cannabis extract preparations (1:1 THC/CBD, CBD only, THC only) and a placebo control group over an 8-week trial period. Participant-reported that pain symptoms decreased significantly among the THC:CBD and THC only groups compared to CBD only and placebo group (P < .001) and 38% (9/24) patients had a decrease in VAS of 50% or more when using active preparations versus placebo. No significant improvements were found on validated measures of sleep, general health, and mood among the THC:CBD and THC only groups. There were no follow-up assessments conducted to determine whether symptom improvements were maintained over time.

An observational prospective-cohort study of 431 patients provides some information about long-term treatment effects.³¹ This study assessed the efficacy of a standardized herbal cannabis product $(12.5\% \pm 1.5\%$ THC titrated up to a recommended maximum of 5g daily) among patients with chronic non-cancer pain over the course of 1 year. Participants in the cannabis group were defined as "patients using cannabis as part of their treatment" and were compared to individuals from the same clinics who denied using cannabis. Compared with baseline, there was a significant reduction in average pain intensity in cannabis group (0.92 change [95% CI, 0.62 to 1.23]), but not in control group (0.18 change [95% CI, -0.13 to 0.49]) at 1 year after adjusting for demographic variables, other substance use, and pain-related variables. Also, a greater reduction in pain intensity was observed among cannabis users versus controls (1.10 difference [95% CI, 0.72 to 1.56]). The cannabis group reported a significant reduction in mood disturbance, as well as improved physical quality of life compared to controls. All changes were stable across the 3-,



6-, and 12-month follow-ups. The limitations of this study were that the majority (66%) of the cannabis users were experienced, making the generalizability to cannabis-naïve users difficult, and this study reported a high drop-out rate (over 30%), which may be a source of selection bias. Reasons for attrition among the cannabis group included perceived lack of efficacy, experience of adverse events, and/or a dislike of the study product. However, authors noted that those who dropped out were comparable to those who completed the study.

The 2 studies of patients with cancer-related pain had an unclear risk of bias and were both included in one of the aforementioned systematic reviews.^{56,57} In a randomized, double-blind, placebo-controlled graded dose study, patients with opioid-refractory cancer pain received a placebo or one of 3 doses of nabiximols (low: 1 to 4 sprays per day; medium: 6 to 10 sprays per day; or high: 11 to 16 sprays per day) during a 5-week treatment period. A separate double-blind, placebo-controlled crossover study evaluated cancer patients who each received placebo, 10 and 20 mg of THC, and 60 and 120 mg of codeine over 5 successive days. These studies both found an improvement in cancer-related pain among medical cannabis users who ingested a 10 mg THC capsule over a 7 hour observation period⁵⁶ and among the low-dose (1 to 4 sprays per day) and medium-dose (6 to 10 sprays per day) nabiximols groups.⁵⁷ The nabiximols trial also identified a significant change in an opioid composite score that was defined as either a reduction in pain with a stable opioid consumption (morphine equivalent) or a reduction in opioid consumption with stable pain (P = .038) among those only in the low-nabiximols dose group.⁵⁷ Methodological limitations of the nabiximols trial were a high attrition rate (27%), the exclusion of patients who reported highly variable pain scores over the course of 3 days, and the use of a non-validated sleep measure. The study comparing THC to codeine did not utilize a validated measure of pain.⁵⁶

KEY QUESTION 2: What are the effects of cannabis on health outcomes and healthcare utilization for adults who have **PTSD**?

KEY QUESTION 2A: Do the effects differ by patient subgroup, such as patient medical and mental health comorbidities?

Summary of Findings

There are very few methodologically rigorous studies examining the effects of cannabis in patients with PTSD. We found only 2 observational studies which suggest that cannabis is potentially associated with neutral effects on PTSD or depression symptom severity, and employment status, and negative effects in terms of violent behavior, drug and alcohol abuse, and suicidal ideation. However, the strength of evidence is rated as insufficient due to the potential for bias in the 2 included studies in this review and the small number of controlled studies reporting data on benefits and harms of cannabis for treating PTSD symptoms. We found no evidence addressing whether effects differed according to other comorbidities in patients with PTSD.

Detailed Findings

We found one systematic review¹⁶ and only 2 primary studies^{60,61} meeting our inclusion criteria (Table 4), primarily because most of the literature on cannabis use in populations with PTSD was cross-sectional and/or did not include a comparison group.

The systematic review by Wilkinson and colleagues (2016) searched the literature through March 2015,¹⁶ and the 2 primary studies we included were not included in their review because they were both published after March 2015. The Wilkinson et al systematic review included 6 studies related to PTSD.⁶²⁻⁶⁷ Of the 6 included studies, 3 were on nabilone, a synthetic form of cannabis.⁶²⁻⁶⁴ One of these was an RCT, though it included only 10 participants, and the other 2 were retrospective chart review studies. The other 3 studies on non-synthetic forms of cannabis were 2 prospective open-label trials,^{65,66} and the last was a prospective observational study;⁶⁷ none of these 3 studies included a control group. Due to the focus on synthetic cannabis or the lack of a control group, none of the 6 primary studies included in the Wilkinson et al (2016) systematic review met our inclusion criteria. In spite of having broader inclusion criteria, the synthesized findings from the Wilkinson et al systematic review suggest that the evidence of the effectiveness of cannabis for reducing PTSD symptoms is insufficient.¹⁶

The primary study by Wilkinson et al (2015) examined data from all Veterans in VA specialized intensive PTSD programs from 1992 to 2011, with a total sample size of over 47,000.⁶⁰ They excluded participants who reported drinking more than 2 alcoholic drinks on one occasion, reported using any other drug 30 days prior to admission, or were referred from a drug or alcohol treatment program. The remaining participants were grouped into "never-users," "stoppers" who used cannabis prior to but not after admission, "continuing users," and "starters" who did not use cannabis prior to admission but started after admission. After balancing sample sizes across groups, they compared 4-month post-baseline outcomes for 2,276 Veterans. They included demographic covariates associated with cannabis use and found that continuing users and starters had significantly worse PTSD symptoms and greater drug abuse than never-users and stoppers at 4 months post-baseline. Starters also experienced significantly greater alcohol abuse than the other groups, and continuing users experienced significantly more violent behavior at 4 months post-baseline compared to the other groups. There were no significant differences among

the groups on employment status.

Johnson et al (2016) examined data at a single time point from Veterans entering a VA-based primary care and mental health integration program.⁶¹ This study included 350 Veterans who used cannabis and 350 non-user controls who were matched on age and gender; all cases and controls had PTSD. Compared to cannabis users, controls were significantly more likely to be married, White, employed, and financially stable. There were no significant differences between cannabis users versus controls on PTSD symptom severity or depression symptom severity. The cannabis users were significantly more likely to experience suicidal ideation and reported significantly more alcohol use (reporting on average approximately 6 alcoholic drinks per week compared to approximately 3 drinks per week in the control sample).

Table 4. Studies of the Effects of Cannabis on PTSD Symptoms

Study, setting, study design (N patients) Risk of bias (ROB)	Sample description Mean age % male	Description and duration of cannabis use and comparators	Primary findings	Other findings
Wilkinson 2015 ⁶⁰ VA retrospective cohort study (N=2276) Medium ROB	All Veterans referred for intensive PTSD treatment. Excluded those with prior drug or alcohol use. Mean age 51.7 96.7% male	Self-reported cannabis use during 4- month follow-up period: 850 never users 299 stoppers (use at admission but not at 4 months post-baseline) 296 continuing users (use at admission and 4 months post- baseline) 831 starters (no use at admission but use at 4 months post-baseline Concomitant medications: Usual medical care including psychotropic medications and psychotherapy provided to all participants.	Continuing users and starters had significantly worse PTSD symptoms than never users and stoppers: <i>F</i> =21.47, <i>P</i> <.0001	Violent behavior: Starters significantly more violence than continuing users, never users, and stoppers. F=21.28, $P<.0001$. Alcohol abuse: Starters significantly more alcohol abuse than continuing users, never users, and stoppers; continuing users significantly more alcohol abuse than stoppers. $F=88.51$, $P<.0001$. Drug abuse: Continuing users and starters significantly more drug abusethan never users and stoppers. F=176.26, $P<.0001$. Employment status: No significant differences among groups. $F=0.66$, $P=.58$.
Johnson 2016 ⁶¹ VA matched case- control cross- sectional study (N=700) High ROB	All Veterans with a probable PTSD diagnosis, who were referred for a primary care/mental health integration program based on clinical need following depression, PTSD, and alcohol use screening, or clinical judgment. Mean age 47.1 91.0% male	Self-reported cannabis use within 3 months of the assessment (n=350) Compared to no lifetime cannabis use reported at the time of assessment (n=350) Users were matched to non-users on age and gender.	Users had significantly worse PTSD symptoms than non-users: t (349) = 0.11, <i>P</i> =.91	Users vs non-users (%): Employed: 23 vs 40, χ^2 (1) = 21.38, <i>P</i> <.0001 Financially stable: 61 vs 71, χ^2 (1) = 8.15, <i>P</i> <.0001 Depression symptoms: No significant differences between groups. t (349) = 1.85, <i>P</i> =.07 Suicidal ideation: 33 vs 26, χ^2 (1) = 12.18, <i>P</i> =.04 Alcohol use: Users had significantly more alcoholic drinks per day than non-users: 6.3 vs 3.1, t (349) = 4.65, <i>P</i> <.0001

Abbreviations: N = number; PTSD = post-traumatic stress disorder; ROB = risk of bias; VA = Department of Veterans Affairs

KEY QUESTION 3: What are the harms associated with cannabis use in adults?

KEY QUESTION 3A: Do the harms differ by patient subgroup, such as patient medical and mental health comorbidities?

We searched broadly for harms and describe the evidence base for each harm category below. We found no evidence which directly compared risk across different patient subgroups, but we describe relevant information about patient characteristics below as applicable.

General Adverse Events

In the 2 systematic reviews examining cannabis for chronic pain, cannabis was overall associated with a higher risk of short-term adverse effects.^{14,15} Across all indications (not just chronic pain or PTSD) and treatment formulations (including synthetic cannabinoids), treatment was associated with an increased risk of: any adverse event (OR 3.03; 95% CI, 2.42 to 3.80), serious adverse event (OR 1.41; 95% CI, 1.04 to 1.92), and withdrawal due to adverse event (OR 2.94; 95% CI, 2.18 to 3.96).¹⁴ In the review focused on only chronic pain, cannabis was similarly associated with a higher risk of adverse events. While most adverse events were mild, there were possible treatment-related adverse events such as suicide attempts, paranoia, and agitation. In the additional trials that we reviewed, the rates of adverse events did not significantly differ between groups. Side effects were rated as minor and may be considered common effects of cannabis, such as dizziness, relaxation, short-term memory impairment, and mental clouding (Table 2).

One prospective cohort study of 431 patients study assessed the incidence of serious adverse events and adverse events over one year among patients using cannabis for chronic non-cancer pain and found no statistically significant group differences between the cannabis-using group and non-using group on serious adverse events. However, cannabis users were at higher risk for non-serious adverse events.³¹ The limitations of this study were that the majority (66%) of the cannabis users were experienced, making the generalizability to cannabis-naïve users difficult, and more frequent follow-up times among the exposure group may have artificially inflated the number of adverse events reported by cannabis users.

In addition, Notcutt and colleagues (2004) had 2 participants withdraw or break blinding due to the inability to tolerate cannabis.³³ The investigators also had to increase the time interval of the initial dosing titration from 15 minutes to 30 minutes between sprays due to 2 participants experiencing dysphoria and lightheadedness.

Medical Harms

Pulmonary Effects

Overview

One systematic review published in 2007,⁶⁸ and 2 more recent prospective cohort studies^{69,70} provide data relevant to the short- and long-term pulmonary effects of cannabis smoking.

Taken as a whole, the literature provides low-strength evidence that low levels of cannabis smoking do not adversely impact lung function over about 20 years in young adults, but there is some evidence suggesting that heavy (*ie*, daily) use may have the potential to cause adverse pulmonary effects over an extended period of time. There are no studies in older users, or in



those with medical comorbidities such as chronic obstructive pulmonary disease (COPD) or heart disease.

Detailed results

There were 12 studies included in the review that directly assessed the short-term effects of inhaled cannabis.⁶⁸ Most studies found that smoking cannabis was associated with bronchodilation up to about an hour after exposure. One study found that nearly daily cannabis use in a controlled environment was associated with increased airway resistance over 2 months. In general, it is difficult to draw firm conclusions from these short-term, small (N < 35) studies published over 2 decades ago, 4 of which did not control for concomitant tobacco use.

The best evidence examining the long-term effects of cannabis smoking on pulmonary function comes from 2 more recently published prospective cohort studies with low risk of bias (Table 5). In one US study, pulmonary function testing was conducted at baseline and 4 more times over a 20-year follow-up in a cohort of healthy young adults (N=5,016).⁷⁰ While a similar proportion of participants smoked cannabis or tobacco cigarettes, most cannabis users smoked infrequently (about twice monthly on average). Higher cumulative tobacco exposure was associated with a significant decline in forced expiratory volume (FEV1) and forced vital capacity (FVC), but cannabis exposure was actually associated with an increase in both measures over 20 years. Of note, the trends in lung function were non-linear: FEV1 levels were flat or downtrending among those with substantial levels of cannabis exposure (the equivalent of one joint daily for 7 years or more).

A birth cohort study (N=1,037) from New Zealand similarly found that FEV1 and FVC increased over time, though the change was small and not statistically significant. Most cannabis users had relatively low rates of cumulative exposure.⁶⁹ Of note, higher rates of cumulative exposure were associated with a small increase in measures of airway resistance.

The prior systematic review also examined long-term pulmonary effects of cannabis. There were 3 cohort studies; the rest were cross-sectional. One of the cohort studies was an earlier interim follow-up from the New Zealand birth cohort study. Another older study examined the effects of "nontobacco" cigarette smoking, but did not have detailed information about cannabis exposure specifically and did not have pulmonary function data for many participants. A third study followed a convenience sample of healthy young adults (mean age 33 years) over up to 8 years of follow-up.⁷¹ About one-third of the participants were heavy habitual cannabis smokers (3.5 joints per day on average), 28% smoked cannabis and tobacco, 17% smoked tobacco only, and 22% smoked neither. About two-thirds of participants had 2 or more FEV1 measures over time, and there was a similar mix of baseline smoking status in those lost to follow-up and those followed longitudinally. The authors found that, while there was a significant decline in FEV1 among tobacco users, cannabis smoking was not associated with a greater decline in FEV1 than nonsmoking.

Cardiovascular Events

Overall, there was insufficient evidence from 2 studies about the effect of cannabis use on the risk of cardiovascular events. Two publications reported analyses from the Myocardial Infarction Onset Study in which nearly 4,000 patients were interviewed just after suffering a myocardial infarction (Table 5). One study assessed the relationship between cannabis use at the time of this baseline interview and subsequent mortality over an average of 12.7 years of follow-up.⁷² There was no information about longitudinal exposure to either cannabis or tobacco use which makes it



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very difficult to assess the relationship between cannabis exposure and long-term mortality. The other analysis was a case-crossover study which compared the risk of myocardial infarction within one hour of cannabis use compared to periods of non-use based on one's pattern of use over the prior year.⁷³ This study had a high risk of bias because recall bias was a significant issue with this study and it was not clear how the authors accounted for tobacco use.

Study, setting, design (N patients) Risk of bias (ROB)	Sample description Mean age % male	Description and duration of cannabis use and comparators	Primary findings	Comments/other findings
Pulmonary effects				
Hancox 2010 ⁶⁹ New Zealand Community-based birth cohort (N=1037) Low ROB	Birth cohort of all individuals in Dunedin, New Zealand, enrolled 1972-1973	Cannabis use only: 25% Lifetime cannabis use, joint- years: 16% > 1, 84% ≤ 1 Comparators: Non-users: 23% Tobacco only: 6% Tobacco + cannabis users: 46%	32 years follow-up. FEV1 Change with each joint-year cannabis in non- tobacco smokers: 5.4 mL (95% CI, -7.1 to 18.0). Change with each pack-year tobacco: -3.9 mL (95% CI, -8.7 to 0.9). FVC Change with each joint-year cannabis in non- tobacco smokers: 13.4 mL (95% CI, -0.8 to 27.6). Change with each pack-year tobacco: 3.6 mL (95% CI, -2.0 to 9.1).	Each joint-year cannabis use also associated with a small but significant increase in airway resistance (0.029 cm H ² O, <i>P</i> =.042), and alveolar volume (28.5 mL, <i>P</i> =.021)
Pletcher 2012 ⁷⁰ US, 4 cities Community-based cohort (N=5016) CARDIA Low ROB	Healthy 18-30 year olds enrolled in 1985 Mean age 25 45% male	Cannabis users: 16% Lifetime use, median joint- years: 0.9 2 median episodes in last 30 days Comparators: Non-users: 46% Tobacco only: 17% Tobacco + cannabis users: 21%	20 years follow-up. FEV1 Highest (> 10 joint-years) vs lowest quartile lifetime cannabis exposure: +36 ml (95% Cl, -6.5 to 79). Highest (> 20 pack-years) vs lowest quartile tobacco exposure: -101ml (95% Cl, -136 to -65) FVC Highest (> 10 joint-years) vs lowest quartile lifetime cannabis exposure: +59 ml (95% Cl, 12 to 107). Highest (> 20 pack-years) vs lowest quartile tobacco exposure: -35 mL (95% Cl, -76 to 5.0).	Association between cannabis use and pulmonary function tests were nonlinear. Within low lifetime exposure group, increasing use was associated with an increase in FEV1 while the slope was level or downtrending in group with higher levels of exposure (> 7 joint- years)
Cardiovascular events				
Frost 2013 ⁷² US, multicenter Hospital-based cohort (N=2097) Determinants of Myocardial Infarction	Patients interviewed just after MI. Users vs non-users: Mean age: 44 vs 52 % male: 94 vs 77	Cannabis smoking within year prior to MI: 109/2097 (5%) Comparator: No cannabis use within prior year (95%)	12.7 years follow-up. Adjusted HR death, compared to no use: Any use: 1.29 (95% CI, 0.81 to 2.05) < weekly: 1.31 (95% CI, 0.74 to 2.35) ≥ once weekly: 1.27 (95 % CI 0.63 to 2.56)	

Table 5. Observational Studies of Cannabis Use and Cardiopulmonary Outcomes

Onset Study High ROB

Study, setting, design (N patients) Risk of bias (ROB)	Sample description Mean age % male	Description and duration of cannabis use and comparators	Primary findings	Comments/other findings
Mittleman 2001 ⁷³ US, multicenter Hospital-based case- crossover (N=3882) Determinants of Myocardial Infarction Onset Study High ROB	Patients interviewed just after MI. Mean age 44 years (cannabis users) 94% male 68% current tobacco smokers	Exposure: cannabis smoking within one hour prior to onset of MI: 9/124 (7%) Comparator: Self as control; expected frequency of cannabis use based on pattern over prior year	Risk of MI within one hour of cannabis use, compared to periods of non-use: OR 4.8 (95% CI, 2.9 to -9.5)	Sensitivity analysis without 3 patients with other triggers in hour prior: OR 3.2 (95% CI, 1.4 to 7.3)

Abbreviations: CARDIA = Coronary Artery Risk Development in Young Adults study; CI = confidence interval; FEV1 = forced expiratory volume; FVC = forced vital capacity; HR = hazard ratio; MI = myocardial infarction; N = number; OR = odds ratio; ROB = risk of bias; US = United States.

Cancer

There was low-strength evidence mainly from case-control studies that cannabis use does not appear to be associated with a higher risk of head and neck or lung cancer (Table 6). There was insufficient evidence from a smaller number of methodologically limited studies about the effects of cannabis on testicular or transitional cell cancer. We found no evidence examining the effects of cannabis on other types of cancer.

Head and neck cancer

A meta-analysis of 9 case-control studies (n=5,732 cases) showed that cannabis use was not associated with head and neck cancer (OR 1.02; 95% CI, 0.91 to 1.14).⁷⁴ Results were generally consistent across studies and there was no evidence of dose-response effect. The analyses are inherently limited by recall bias and there was a very wide range of ever cannabis use across studies, though results were consistent across different study populations.

Lung cancer

One international IPD meta-analysis of 6 case-control studies (n=2,159 cases) found no association between habitual cannabis use (≥ 1 joint-year) and lung cancer among middle-aged patients (OR 0.96; 95% CI, 0.66 to 1.38).⁷⁵ The results were consistent across different analyses, intensity of use, age of first use, and after excluding patients who had used cannabis within 2 years of diagnosis. Though the study was generally well-conducted, recall bias is an inherent limitation. The results apply most closely to persons with relatively light cannabis use as there were very few patients with a history of intense use. While this was a large study, there were few patients who were both habitual cannabis users and who had never smoked tobacco.

A large 40-year cohort study (N=49,231; n=189 lung cancer cases) from Sweden had a high risk of bias because of significant methodologic flaws including lack of long-term data on cannabis and tobacco exposure that make it difficult to interpret findings.⁷⁶ Cannabis and tobacco use were assessed only at the time of military conscription, and these exposures were related to subsequent risk of lung cancer over 40 years of follow-up.

Testicular cancer

A meta-analysis of 3 case-control studies (n=719 cases) found a small increase in the risk of testicular cancer among weekly cannabis users compared to those who never used (OR 1.92; 95% CI, 1.35 to 2.72).⁷⁷ In sensitivity analyses, the increased risk was only seen among those with non-seminoma cancers and not in those with seminoma cancers. While the meta-analysis itself was methodologically strong, there were substantial methodologic weaknesses in each of the 3 included studies rendering the meta-analysis at high risk of bias. The smallest study did not control for all important confounders including tobacco use. Results were consistent in the 2 larger and methodologically stronger studies, but response rates were very low which may exacerbate issues with recall bias.

Transitional cell cancer

One small case-control study (n=52 cases) from 2 VA urology clinics assessed the risk of transitional cell carcinoma.⁷⁸ While there was an increased risk of cancer seen with heavier cannabis use, the results are difficult to interpret because of significant methodologic flaws placing the study at high risk of bias.

Table 6. Observational Studies of Cannabis Use and Cancer Risk

Cancer type Study, setting, design (N patients) Risk of bias (ROB)	Sample description Mean age % male	Description and duration of cannabis use	Primary findings	Comments/other findings
Head and neck cancer Carvalho 2015 ⁷⁴ US, Africa, South America Meta-analysis of 13 case- control studies Hospital-based (6) and cancer-registry (5) studies Medium ROB	Patients with definitive diagnosis of head-neck cancer (in studies of moderate to high methodologic quality). Mean age NR % male NR	<u>% ever cannabis smokers:</u> Cases: range 2.4 to 83; overall 12.6 Controls: range 0.4 to 83; overall 14.3	9 studies contributed data to meta- analysis OR (95% CI) for head neck cancer among cannabis users: 1.02 (0.91 to 1.14); adjusted for age, gender, race, tobacco use	
Lung cancer Zhang 2015 ⁷⁵ International Lung Cancer Consortium North America, New Zealand, Europe Individual-level meta- analysis of 6 case-control studies (2,159 cases, 2,985 controls, combined) Medium ROB	Patients with histologically confirmed lung cancer. Cases vs controls: Median age: 57.3 vs 53.0 % male: 50 vs 53	Cannabis and tobacco use: Cases: ≥ 1 joint-year: 10% ≥ 1 joint-year, non-tobacco users: 3.0% Never smoked tobacco: 17% Controls: ≥ 1 joint-year: 11% ≥ 1 joint-year, non-tobacco users: 4.7% Never smoked tobacco: 46%	OR (95% Cl) for lung cancer among habitual (≥ 1 joint-year) users compared to non-habitual or never users: 0.96 (0.66 to 1.38); adjusted for age, sex, race, education, tobacco pack-years and status	OR among never tobacco smokers: 1.03 (0.51-2.08)
Lung cancer Callaghan 2013 ⁷⁶ Sweden Community-based cohort study (N=49,231) High ROB	Military conscripts born between 1949-1951 and inducted between 1969 and 1970 100% male	Lifetime cannabis use at time of conscription: Cases: Once (2.5%) 2-4 times (3.0%) 5-10 times (1.7%) 11-50 times (1.5%) > 50 times/"heavy" (1.7%) Controls: Never (82.5%) Tobacco only: 55.2% Tobacco + cannabis: 9.1% Cannabis with no tobacco use: 13.4%	40 years follow-up. 189 incident cases of lung cancer (by ICD-9 codes). HR (95% CI) for lung cancer among self-reported heavy users: 2.12 (1.08 to 4.14); adjusted for alcohol, COPD/asthma, socioeconomic status, occupation, tobacco	No significant association between other levels of cannabis use and lung cancer, no dose-response relationship.

<i>Cancer type</i> Study, setting, design (N patients) Risk of bias (ROB)	Sample description Mean age % male	Description and duration of cannabis use	Primary findings	Comments/other findings
Testicular cancer Gurney 2015 ⁷⁷ US Meta-analysis of 3 case- control studies (719 cases, 1419 controls combined) High ROB	Young adults with histologically confirmed testicular cancer Mean age NR; range 18 to 50 100% male	Overall proportion with ever, never, weekly, and current cannabis use NR	Cancer risk OR (95% CI), compared with never use: Weekly use: 1.92 (1.35 to 2.72), all studies adjusted for age and cryptorchidism; 2 largest studies adjusted for alcohol and tobacco use.	The association between cannabis use and cancer was only seen among non- seminoma cancers and not in seminoma cancers
			≥ 10 year use: 1.50 (1.08 to 2.09). Ever-use: 1.19 (0.72 to 1.95).	
<i>Transitional cell cancer</i> Chacko 2006 ⁷⁸ US, 2 VA sites Case-control (52 cases 104 controls) High ROB	Patients under age 60 with transitional call cancer presenting to urology clinic. Mean age 51 100% male	Cases: Smoked > 40 joint-years: 40.4% Ever smoked cannabis: 88.5% Smoked tobacco and cannabis: 76.9% Smoked tobacco only: 17.3% Smoked cannabis only: 11.5%	Joint-years cannabis use as continuous variable was significantly associated with transitional cell cancer: <i>P</i> -trend .01 (adjusted for tobacco use, smoked meat use, radiation, agent orange, and dye exposure)	
		Controls: Smoked > 40 joint-years: 15.1% Smoked cannabis: 69.2% Smoked tobacco and cannabis: 65.4% Smoked tobacco only: 27.9% Smoked cannabis only: 3.9%	Risk of cancer with > 40 joint-years cannabis use compared to none: OR 3.4 (unadjusted, <i>P</i> =.012)	

Abbreviations: CI = confidence interval; COPD = chronic obstructive pulmonary disease; NR = not reported; OR = odds ratio; ROB = risk of bias; US = United States; VA = Department of Veterans Affairs.

Motor Vehicle Accidents

Overall, we found evidence suggesting an increased risk of collision associated with acute cannabis intoxication, but the magnitude and precision of increased risk are unclear.

A 2016 systematic review of cannabis intoxication and motor vehicle accidents pooled the findings of 21 multi-national observational studies that were published between 1982 and 2015, with a combined sample size of 239,739. The meta-analysis determined a statistically significant, moderate increase in collision risk associated with acute cannabis intoxication (OR 1.35; 95% CI, 1.15 to 1.61).⁷⁹ In assessing study quality, the review authors examined the methods used to measure drug use (*eg*, self-report, or lab values from blood versus urine or saliva), crash severity, adjustment for alcohol use and other confounders, and whether the study evaluated a dose-response effect. Sub-analyses that grouped studies based on quality, design (case-control versus culpability studies), degree of adjustment for confounders, and crash severity (whether fatalities were involved) found pooled effects in the range of 1.07 to 1.81 using a random effects model, and 1.08 to 1.90 using meta-regression.

The review authors suggested that the pooled estimate may be complicated by factors affecting a user's decision to drive under the influence of cannabis. Experimental studies using simulated driving have reported that alcohol increases driving speed and risk-taking, while cannabis users tend to be aware of their impairment and drive slower and more cautiously in an effort to compensate.^{80,81} The pooled effect may underestimate the true risk of collision with acute cannabis intoxication, if users are more likely to drive when their level of impairment is low. Conversely, the pooled estimate may be inflated if cannabis users who choose to drive while intoxicated have a higher baseline risk independent of cannabis use, compared with cannabis users who choose not to drive after use.⁷⁹

A study that sought to determine a threshold for serum concentration of THC associated with driving impairment found that serum concentrations below 10 ng/mL were not associated with elevated accident risk, based on limited epidemiological data.⁸² The authors of the study reported that based on experimental studies, THC serum concentration in the range of 7 to 10 ng/mL is comparable to a blood alcohol concentration of 0.05% on degree of impairment.⁸²

Mental Health-Related Harms

Suicidal Behaviors

We found no evidence examining the effects of cannabis use on suicide risk in patients with chronic pain or PTSD.

A review and meta-analysis of epidemiological research from 1995 to 2015 found few studies on the effect of cannabis use and suicidality (suicide death, ideation, and attempt) among the general population including both adolescents and adults.⁸³ Data were insufficient to comment on the effect of acute cannabis use and suicidality. However, the review found limited evidence suggesting significantly increased odds of suicide death (pooled OR 2.56; 95% CI, 1.25 to 5.27, 4 studies) with any cannabis use. In 6 studies each, any cannabis use was significantly associated with increased odds of suicide ideation (pooled OR 1.43; 95% CI, 1.13 to 1.83) and suicide attempt (pooled OR 2.23; 95% CI, 1.24 to 4.00). Further, heavy cannabis use was associated with significantly increased odds of suicide attempt (pooled OR 3.20; 95% CI, 1.72 to 5.94). Suicide ideation was noted to be increased among heavy cannabis users, though this was of borderline significance (OR 2.53; 95% CI, 1.00 to 6.39). Cannabis use was slightly more



common among individuals who died from suicide who used non-overdose methods (11.6%) than among those who died from suicide related to overdose methods (9.2%) in general population studies. Limitations of this review included significant heterogeneity between studies with respect to measurement of cannabis exposure and control of risk factors, the use of observational studies (including case-series and cross-sectional), a small number of suicidality cases in studies, and research from a small number of geographical locations. An older review that included 7 studies on suicidal ideation or attempts (with 2 studies included in both reviews) found mixed results: 4 studies reported an association between cannabis use and increased risk of suicidal ideation, one study showed no association.⁸⁴

Mania

We found no evidence examining the effects of cannabis on the risk of mania among persons with PTSD or chronic pain.

One systematic review that included 6 prospective studies of other populations (mean follow-up 3.9 years) found support for an association between cannabis use and exacerbation or incidence of manic symptoms.⁸⁵ Among patients with known bipolar disorder, 3 studies demonstrated significant associations between cannabis use and fraction of time with mania or mania score/symptoms during follow-up, though meta-analysis was not undertaken. Further, a meta-analysis of 2 prospective community studies demonstrated an association between cannabis use and new-onset mania symptoms among those without a diagnosis of bipolar disorder (pooled OR 2.97; 95% CI, 1.80 to 4.90) with low heterogeneity between studies. The strength of the findings is limited by the small number of included studies in this review.

Psychosis

One systematic review⁸⁴ and 7 studies⁸⁶⁻⁹² provided evidence related to psychotic symptoms associated with cannabis use. Overall, studies consistently showed a relationship between cannabis use and the development of psychotic symptoms, though the magnitude of risk is uncertain. In addition, experimental studies have found acute, transient psychotic symptoms within hours of use. The Moore et al (2007) review also included studies that showed an increased risk of psychotic spectrum disorder among cannabis users. Given that many of the studies are observational, it is difficult to determine whether cannabis directly contributed to the development of psychotic symptoms or whether its use was simply more common among individuals with a preexisting tendency towards these symptoms. The possibility that cannabis contributes directly to symptom development is supported but not proven by biologic plausibility, evidence of a dose-response relationship, and the results of prospective cohort studies, described in the following sections.

Psychotic symptoms

Four studies included only participants with no psychotic symptoms at baseline.^{86-88,92} Time to follow-up ranged from 12 to 36 months; 2 of the 4 studies examined linear trends across frequencies, and the other 2 comparing higher to lower frequencies of use. All 4 studies found that participants who had ever used cannabis had an increased likelihood of any psychotic outcome (*eg*, symptoms, psychotic disorder) compared to participants who had never used. The studies also found that frequency of use correlated with the likelihood of a psychotic outcome.

Two articles provided data from the Early Developmental Stages of Psychopathology (EDSP) study, a prospective cohort study (medium risk of bias) of randomly selected adolescents and young adults aged 14 to 24 at baseline (N=3,021; mean age 18.3 years).^{86,87} Findings from these studies indicate that at the second (T2) and third time point (T3), using cannabis more than 5 times since the previous assessment (3.5 years between baseline and T2, and 4.9 years between T2 and T3) was associated with positive symptoms (OR 2.10; 95% CI, 1.61 to 2.75) and the cooccurrence of both positive and negative symptoms (OR 2.05; 95% CI, 1.18 to 3.59), but not negative/disorganized symptoms alone (OR 1.12; 95% CI, 0.91 to 1.39).⁸⁷ Among those reporting no cannabis use at baseline, cannabis use between baseline and T2 increased the risk for psychotic symptoms between T2 and T3 (adjusted OR 1.9; 95% CI, 1.1 to 3.1; P = .02). Among those reporting cannabis use at baseline, continued use at T2 was associated with psychotic symptoms at both T2 and T3 (adjusted OR 2.0, 1.0 to 3.8; P = .037).⁸⁶ In addition, a case-control study of 280 individuals presenting with a first episode of psychosis and 174 healthy controls found that after adjusting for confounders, there was no significant difference between groups in ever having used cannabis, or the duration of use. However, those experiencing a first episode of psychosis were more likely to use cannabis daily (adjusted OR 6.4; 95% CI, 3.2 to 28.6), and were more likely to use sinsemilla (adjusted OR 6.8: 95% CI. 2.6 to 25.4).⁹²

One cohort study (N=591) with a low risk of bias examined the relationship between frequency of use in adolescence and psychotic symptoms over a 30 year period. In the multivariate model, the frequency of use in adolescence (casual use: OR 1.80; 95% CI, 1.24 to 2.59; P = .002; regular use: OR 2.60; 95% CI, 1.59 to 4.23; P < .001) was a significant predictor of 'schizotypal signs' (*eg*, feeling lonely even when with people, never feeling close to another person). There was no significant relationship between cannabis use and schizophrenia nuclear symptoms (*eg*, thought insertion, thought broadcasting, thought control, hearing voices).⁸⁸

Acute cannabis-induced psychosis

Three studies examined the relationship between cannabis use and acute psychotic symptoms.⁸⁹⁻ ⁹¹ In one (moderate risk of bias) study, a double-blind cross-over RCT of 16 healthy cannabisnaïve women (mean age 23.56 years), comparing oral cannabis extract to placebo, one participant experienced psychotic symptoms (ie, "severe" somatic concern, anxiety, tension, depressive mood, suspiciousness, hallucinatory behavior, motor retardation, and "extremely severe" unusual thought contents) 3 hours after cannabis intake. Symptoms decreased without pharmacological intervention.⁸⁹ The second (low risk of bias) study compared THC plus CBD to THC plus placebo (N=48). Clinically significant positive symptoms (ie, an increase in Positive and Negative Syndrome Scale [PANSS] positive scores of 3 or more points), were more common with THC plus placebo (11 of 26 cases) compared to THC plus CBD (3 of 22 cases), $(\chi^2=4.74, P < .05)$, and individuals in the THC plus placebo group experienced greater paranoia (t=2.28, P < .05).⁹¹ The third was a (high risk of bias) case-control study comparing 140 cannabis users to 144 non-users on psychotic symptoms (ie, delusory thinking, perceptual distortion, cognitive disorganization, anhedonia, mania, and paranoia). Cannabis users were evaluated immediately after use, as well as 3 to 4 days later. Univariate results indicate more psychotic symptoms in the cannabis group ($F_{1,282} = 80.1$, P < .005), with greater effects immediately after use.⁹⁰

Cognitive Effects

One systematic review provides moderate-strength evidence that active, long-term cannabis use is associated with small negative effects on all domains of cognitive function, but insufficient



evidence of long-term cognitive effects in past users.⁹³ The review first synthesized the literature on non-acute (*ie*, residual and long-term combined) cognitive effects of cannabis use, reporting that the 33 included studies (with a combined total of 1,010 cannabis users compared to 839 controls) suggested that there is a small, non-acute effect of cannabis use on global cognitive functioning and on each of the 8 domains of cognitive functioning reported in the papers, which included abstraction/executive, attention, forgetting/retrieval, learning, motor, perceptual-motor, simple reaction time, and verbal/language domains. The authors then conducted a subgroup analysis of only 13 studies (with a combined total of 388 cannabis users and 387 controls) which examined cognitive functioning after at least 25 days of abstaining from cannabis use, described as long-term use. They reported that in this subgroup of studies examining long-term effects, there was not a statistically significant effect on global cognitive functioning, nor on any of the 8 reported cognitive domains.⁹³

Schreiner and colleagues' systematic review⁹³ documents consistent evidence supporting nonacute (*ie*, combined findings from both residual and long-term effects studies) cognitive effects of cannabis from the 33 studies included in their review, though these data are not specific to chronic pain or PTSD populations. Therefore, the strength of evidence for residual effects of cannabis use is rated as moderate. The magnitude of these non-acute effects is small overall, but because the studies all reported average cognitive impairment and not the percent of study participants with clinically significant cognitive impairment, it is not possible to provide an estimate for the range of severity of cognitive impairment experienced by the cannabis users in these studies.

The long-term effects of cannabis use on cognitive functioning are less clear, and the systematic review by Schreiner and colleagues suggests that cannabis use might not result in long-term cognitive impairment. This sub-analysis, however, was based on a relatively small sample from 13 studies with a very broad range of time since last cannabis use (ranging from an average of 25 days to an average of over 3 years). The amount of prior cannabis use reported in these studies also varied greatly, ranging from an average of weekly use to an average of using cannabis multiple times per day. This heterogeneity among the 13 included studies makes generalizations about amount and frequency of cannabis use associated with cognitive impairment impossible and could be at least part of the reason for the lack of consistent findings across studies. Most of the cognitive domains reported in these studies had inconsistent results within or across studies or more consistent but non-significant trends indicating the presence of at least mild long-term cognitive impairment. This suggests that, in at least some cognitive domains, a larger sample might yield findings of significant associations between cannabis use and cognitive impairment that is present after at least 25 days after abstinence. The evidence for a lack of long-term cognitive impairment associated with cannabis use reported in the Schreiner et al review, therefore, is rated as insufficient strength of evidence.

Cannabis Use Disorder (CUD)

The Diagnostic and Statistical Manual of Mental Disorders, Fifth Edition (DSM-5)⁹⁴ and the 10th Revision of the International Statistical Classification of Diseases and Related Health Problems (ICD-10)⁹⁵ both require multiple symptoms of significant psychiatric distress, social impairment, and adverse consequences associated with cannabis use for an individual to be diagnosed with CUD. While we did not find studies reporting prevalence estimates of CUD in the population of Veterans with PTSD, Bonn-Miller et al (2012) report that the prevalence of PTSD among Veterans with CUD was 29.05% in fiscal year 2012.⁹⁶



We did not find any articles comparing rates of CUD in chronic pain or PTSD populations to other populations.

A recent large national prospective cohort study found high prevalence of CUD (36%) among those reporting cannabis use in the past year (N = 1279).⁹⁷ Cannabis use was associated with incident cannabis use disorder (adjusted odds ratio, 9.5 [CI, 6.4 to 14.1]) in a large (N = 34,653) prospective cohort study. Cannabis use was also associated with increased odds of other substance use disorders (any substance use disorder: odds ratio [OR], 6.2; 95% CI, 4.1-9.4; any alcohol use disorder: OR, 2.7; 95% CI, 1.9-3.8.

Other studies of CUD provide potentially relevant cross-sectional data. For example, one non-VA study using structured diagnostic interviews found that the prevalence of cannabis misuse and dependence were 2.4% and 0.9%, respectively, in a primary care sample (though the proportion of patients who used cannabis was unknown).⁹⁸ Another cross-sectional study by Hefner and colleagues (2015) examined rates of CUD in a sample of over 1.3 million Veterans with chronic non-cancer pain, comparing rates of CUD among groups of Veterans based on the number of opioid prescriptions for non-cancer pain.⁹⁹ They found that 1.98% of Veterans with chronic non-cancer pain who were not prescribed opioids had a CUD diagnosis compared to 2.83% of those with 1 to 2 opioid prescriptions in one year, 3.44% with 3 to 10 opioid prescriptions, 3.28% with 11 to 19 opioid prescriptions, and 3.92% of Veterans with 20 or more opioid prescriptions in one year who were diagnosed with CUD.

Bonn-Miller et al (2015) studied 104 Veterans who had CUD and were attempting to stop using cannabis.¹⁰⁰ They reported that PTSD was associated with higher baseline rates of cannabis use and a slower decrease in cannabis use during the first 4 weeks following a quit attempt. Walsh and colleagues (2014) found that cannabis dependence was not associated with trauma exposure, but was associated with a greater number of PTSD symptoms in a sample of 1317 Jewish Israeli individuals.¹⁰¹ Finally, Kevorkian and colleagues (2015) examined data from the National Epidemiologic Survey on Alcohol and Related Conditions (N=34,396).¹⁰² They reported that while trauma exposure during one's lifetime was only very minimally associated with CUD (OR 0.997; 95% CI, 0.996 to 0.999), among trauma-exposed, cannabis-using individuals, PTSD was significantly associated with increased likelihood of CUD (OR 1.217; 95% CI, 1.214 to 1.220).

CUD may also impact response to PTSD treatment, though CUD has not been well-studied in general in PTSD populations. Bonn-Miller et al reported in 2013 that among Veterans who were enrolling in a VA, all-male, inpatient, intensive PTSD treatment program, those who had CUD experienced less improvement in PTSD symptoms during the course of treatment than those who did not have CUD upon enrollment.¹⁰³ This relationship was observed for overall PTSD symptoms as well as avoidance/numbing and hyperarousal symptom clusters, though group differences were non-significant for re-experiencing symptoms. These analyses included statistical adjustment for covariates including age, combat exposure, and depression symptoms as well as alcohol, amphetamine, cocaine, opioid, and sedative use disorders.

Emerging Harms

Infectious Diseases

Several case reports have suggested an association between smoking cannabis and invasive pulmonary aspergillosis in immunocompromised individuals.¹⁰⁴⁻¹⁰⁶ In an older study, investigators randomly selected 28 individuals with a history of cannabis smoking, 21 of whom were asymptomatic, 6 of whom had bronchitis symptoms after smoking, and 1 of whom was

diagnosed with pulmonary aspergillosis.¹⁰⁷ Serum precipitins against Aspergillus antigens were significantly more common among individuals with a cannabis smoking history compared to age-matched controls. Most cannabis cigarette samples provided by the participants had Aspergillus species detected in culture, and there was passage of fungal spores demonstrated through most of the samples.

Cannabis has been implicated as a possible contributing factor in tuberculosis clusters through the shared use of a cannabis water pipe,^{108 13841} or through the practice of "hotboxing."¹⁰⁹

Cannabinoid Hyperemesis Syndrome

Recently, a number of case series have described a syndrome of at times severe cyclic vomiting associated with chronic cannabis use called the cannabinoid hyperemesis syndrome.¹¹⁰⁻¹¹⁶ The largest case series included 98 patients from a single institution.¹¹⁷ The authors performed an institution-wide review of medical records of patients with recurrent vomiting, without an associated etiology, and known preceding cannabis use. All patients were younger than 50 years old and 95% had used at least once weekly; 68% of the patients had used cannabis for over 2 years. Most patients (86%) had abdominal pain as well. Information about the effect of hot water was available in 57 patients: 91% of these patients reported relief of symptoms with hot showers. Long-term follow-up was only available in 10 patients, so it is uncertain how many patients ultimately abstained from use and how often this resolved the symptoms. Earlier case series reported that most patients who discontinued use recovered.¹¹²

Complications from Intravenous Use of Cannabis

The intravenous marijuana syndrome is an acute illness following the injection of boiled cannabis preparations. The syndrome was last described in a synthesis of 25 case reports in 1986. In most cases, patients had a febrile illness with tachycardia, hypotension, gastrointestinal symptoms, and myalgias.¹¹⁸ The pathogenesis of the syndrome is unknown. A minority of patients had used cotton to strain the preparation prior to use suggesting some similarity to "cotton fever" that has been described in heroin users. Alternatively, it is possible that very high doses of cannabis itself could have contributed.

Aggression and Violence

Two studies investigated the effect of cannabis use on aggression and found mixed results. A retrospective study of clinical files from 4 public psychiatric outpatient facilities in Italy that included patients treated for 6 months continuously (N=1,582; 49% male, 41% with mood disorder and 27% with psychotic disorder) found cannabis use to be a risk factor for violent behavior, regardless of psychiatric disorder, sex, and age.¹¹⁹ The combination of a mental disorder and cannabis use was present in significantly more patients with violent behavior (3.9%) versus those with non-violent behavior (0.2%; OR 19.2; 95% CI, 4.4 to 118.6). Also, mental health patients who used cannabis were significantly more likely to engage in both violence towards others (OR 10.2; 95% CI, 3.8 to 27.5) and violence towards themselves (OR 5.7; 95% CI, 2.4 to 13.5). In particular, the probability of suicide increased more than 17 times (OR 17.6; 95% CI, 3.5 to 87.7) and the probability of attempted suicide tripled (OR 3.4; 95% CI, 1.5 to 9.4) among cannabis users versus non-users. Notably, cannabis use was significantly associated with being male, a family history of violent behavior, precarious employment, poor compliance with treatment, and undergoing psychotherapy, and there was a significant correlation between violent behavior and a positive family history for both substance misuse and violent behavior, suggesting that factors other than cannabis use are implicated in violent behavior.



A second study of 30 undergraduate males who received intense provocation following ingestion of either low (0.1 mg/kg), medium (0.25 mg/kg), or high (0.4 mg/kg) doses of THC found that the low-dose group tended to respond with more aggression than the high-dose group.¹²⁰ Participants in this study were randomly allocated to their THC dosing and asked to select a shock intensity to be administered to an opponent during a competition. In the absence of provocative stimulation, in which participants were not aware of their opponents' aggressive intentions (based on opponents' choice of shock level to be administered to the participant), there was no difference in shock intensity given by participants by THC dose. In the presence of provocative stimulation, participants in the low-dose group were significantly more likely to escalate shock intensity and use extremely high shock settings to retaliate against aggressive opponents compared with those in moderate and high THC dose groups (P < .05 for both). These findings suggest that aggression is not associated with cannabis use.

Miscellaneous

There are emerging issues related to newer methods of cannabis use that clinicians may encounter. "Dabbing" refers to vaporization and inhalation of butane hash oil which has THC concentrations that typically far exceed that seen in the cannabis flower. In a survey study, "dab" users (N=357) reported more trouble with tolerance and withdrawal than what they had experienced using flower cannabis.¹²¹ Edible cannabis use has become more common in recent years, especially in states in which cannabis has been legalized for recreational or medical purposes.¹²² A recent case series described 5 patients hospitalized with acute psychosis after ingestion of edible cannabis.¹²³ The patients described ingesting multiple portions in part because of the delay in onset of effect seen with edible cannabis, thus ingesting a much larger dose of THC than recommended.

A recently published (after our search dates ended) follow-up to a New Zealand birth cohort study found that cannabis use was associated with the development of periodontal disease by early midlife after adjusting for tobacco use.¹²⁴ They found no association with intermediate health outcome measures such as lipids, hemoglobin A1c, and measures of inflammation. However, nearly two-thirds of cannabis users also used tobacco, and there were relatively few people who used cannabis heavily.

KEY QUESTION 4: What are important areas of ongoing research and current evidence gaps in research on cannabis for chronic pain or PTSD, and how could they be addressed by future research?

Summary of Findings

Chronic Pain

We identified 10 ongoing RCTs examining the effectiveness of cannabis for a variety of chronic pain conditions (Table 7), including several populations included in this report (3 studies for cancer pain and 2 studies for neuropathic pain), as well as conditions for which there is currently very little or no evidence (osteoarthritis, sickle cell disease, low back pain, and ulcerative colitis). While there are several ongoing observational studies on the benefits and/or harms of cannabis, we found no studies looking specifically at chronic pain populations that would meet our inclusion criteria.

Most of the ongoing trials are relatively small, with 6 including fewer than 100 patients (mean 46 participants). However, 2 industry-funded placebo-controlled trials investigating nabiximols include roughly 400 patients each, and another parallel RCT compares vaporized cannabis to dronabinol (synthetic THC) and placebo in 120 adults. In addition to assessing pain, 5 trials will assess quality of life and/or functional status outcomes, 5 trials will look for mental health outcomes such as mood and depression, and 4 trials will examine cognitive outcomes, a harm on which there is very little current evidence in chronic pain populations. The follow-up duration for these trials is relatively short, ranging from 1 to 10 weeks (median 5 weeks).

Similar to the published studies included in this report, the most commonly used cannabis products in these ongoing trials are vaporized (3 studies) or smoked (3 studies) cannabis with known THC and/or CBD content, or nabiximols oromucosal spray (2 studies). One of these trials is a crossover RCT investigating 6 different vaporized cannabis products with varying THC and CBD content in 40 adults with painful osteoarthritis of the knee (NCT02324777). This trial may provide some evidence as to the most effective cannabis formulations or potencies; however, as a relatively small trial (40 patients) with only one day of exposure for each of the formulations, conclusions about their effectiveness will be limited. We found only one other study planning to compare different potencies of cannabis (NIH project number 5R01DA030424-03).

PI/Study Director (Registration); Study Design; Sponsors; Estimated Study Completion	Study Title	Purpose of Study	Participants; Intervention(s)/ Comparator	Outcomes and Timing
 Abrams, DI (<u>NCT01771731</u>) Crossover RCT Sponsored by the University of California, San Francisco; National Heart, Lung, and Blood Institute (NHLBI); University of Minnesota - Clinical and Translational Science Institute March 2016 	Vaporized Cannabis for Chronic Pain Associated With Sickle Cell Disease (Cannabis-SCD)	To assess whether inhaling vaporized cannabis ameliorates chronic pain in patients with sickle cell disease; assess the possible synergistic affect between inhaled cannabis and opioids; assess the clinical safety of the concomitant use of cannabinoids and these opioids; evaluate the short-term effects of inhaled cannabis on markers of inflammation and disease progression in patients with sickle cell disease.	 35 adults with sickle cell disease with ongoing opioid analgesic therapy for chronic sickle cell disease-associated pain. In a controlled inpatient setting, the contents of 1 cigarette is vaporized and inhaled at 12pm on day 1; 8am, 2pm, and 8pm on days 2-4; and 8am on day 5. 1. Cannabis cigarette: 4.7% THC and 5.1% CBD 2. Placebo cigarette: 0% THC and 0% CBD Participants to receive both treatments in random order for 5 days (2-week washout). 	Pain VAS evaluated during the 5-day inpatient exposure. Other outcomes: mood; QOL assessments; inflammation markers and disease progression from blood samples.
 Dayan, L (<u>NCT02560545</u>) Crossover RCT Sponsored by the Tel-Aviv Sourasky Medical Center September 2016 	Cannabinoids Effects on the Pain Modulation System	NR	 40 adults with at least moderate neuropathic pain (> 30 out of 100 on VAS) for ≥ 3 months, who have not responded to other painkillers or for whom they are contraindicated due to side effects. 1. Cannabis oil: 20% THC, 40 mg per 70 kg weight; route of administration not specified 2. Placebo oil 	Evaluation of pain using a questionnaire at 1 month. Other outcomes: testing of the pain-modulation system using TSA Neurosensory Analyzer.
 GW Pharmaceuticals Ltd. (NCT01262651) Parallel RCT Sponsored by the GW Pharmaceuticals Ltd.; Otsuka Pharmaceutical Development & Commercialization, Inc. 	A Study of Sativex® for Relieving Persistent Pain in Patients With Advanced Cancer	To determine the efficacy, safety and tolerability of nabiximols (Sativex) as an adjunctive treatment, compared with placebo, in relieving uncontrolled persistent chronic pain in patients with advanced cancer.	 397 adults with an advanced cancer for which there is no known curative therapy, and a clinical diagnosis of cancer-related pain which is not alleviated with their current optimized opioid treatment. 100 µl oromucosal spray administered twice daily up to a maximum of 10 sprays 	Percent improvement from baseline to the end of treatment in NRS average pain score (5 weeks). Other outcomes: change in NRS average pain; change in mean NRS worst pain; change in

PI/Study Director (Registration); Study Design; Sponsors; Estimated Study Completion	Study Title	Purpose of Study	Participants; Intervention(s)/ Comparator	Outcomes and Timing
· July 2015			per day: 1. Nabiximols (oromucosal spray delivering 2.7 mg THC/2.5 mg CBD) 2. Placebo	mean sleep disruption.
 GW Pharmaceuticals Ltd. (NCT01424566) Crossover RCT Sponsored by the GW Pharmaceuticals Ltd.; Otsuka Pharmaceutical Development & Commercialization, Inc. December 2015 	A Two-Part Study of Sativex® Oromucosal Spray for Relieving Uncontrolled Persistent Pain in Patients With Advanced Cancer	To determine the efficacy of nabiximols (Sativex) as an adjunctive medication in relieving persistent chronic pain (not breakthrough pain) in patients with advanced cancer, who have this pain even when they are on optimized/maximized chronic opioid therapy.	 406 adults with an advanced cancer for which there is no known curative therapy, and a clinical diagnosis of cancer-related pain which is not alleviated with their current optimized opioid treatment. 100 μl oromucosal spray administered twice daily up to a maximum of 10 sprays per day: Nabiximols (oromucosal spray delivering 2.7 mg THC/2.5 mg CBD) Placebo 	Mean 11-point NRS average pain score over the last 4 days of treatment period (7 weeks). Other outcomes: percentage improvement in NRS average pain score; mean NRS worst pain score; mean sleep disruption.
 Irving, P (<u>NCT01562314</u>) Parallel RCT Sponsored by GW Research Ltd June 2015 	A Pilot Study of GWP42003 in the Symptomatic Treatment of Ulcerative Colitis (GWID10160)	To determine the efficacy and safety of GWP42003 compared with placebo, by the percentage of participants achieving remission.	 60 adults with mild to moderate ulcerative colitis on a fixed dose of 5-aminosalicylic acid treatment and a with a Mayo assessment score 4-10. One of the following twice daily for 10 weeks: 1. GWP42003 (oral capsule that contains both CBD and THC) up to 250 mg twice daily 2. Placebo 	Percentage of participants achieving remission, quantified as a Mayo score of ≤ 2 (with no sub-score > 1). Other outcomes: NRS pain, Mayo total score, health-related QOL, Subject Global Impression of Change, Global Assessment of Illness Severity.
 Martinez, D (<u>NCT02675842</u>) Parallel RCT Sponsored by the New York State Psychiatric Institute December 2021 	Investigation of Cannabis for Pain and Inflammation in Lung Cancer	To investigate the efficacy of cannabis, compared to placebo, in participants undergoing radiation therapy for lung cancer.	30 adults with lung cancer receiving radiation therapy. Smoked cannabis (1 to 2 cigarettes over the course of 2 to 3 hours) administered 3 to 5 days/week in the research laboratory for 6 weeks: 1. High CBD/low THC: 15.76% CBD and	Change in pain ratings using the McGill Pain Questionnaire and the 9 item BPI at 6 weeks. Other outcomes: sickness- related impairment; physical and emotional wellbeing; QOL; tiredness; mood; appetite/eating;

PI/Study Director (Registration); Study Design; Sponsors; Estimated Study Completion	Study Title	Purpose of Study	Participants; Intervention(s)/ Comparator	Outcomes and Timing
			3.11% THC 2. Placebo: 0.0% CBD and 0.01% THC	subjective effects; cognitive status; physiological state; opioid use.
Martinez, D (<u>NCT02683018</u>) • Crossover RCT • Sponsored by the New York State Psychiatric Institute • March 2021	Investigation of Cannabis for Chronic Pain and Palliative Care	To investigate the effects of high CBD/low THC cannabis on symptoms such as pain, nausea/vomiting, and QOL in seriously ill participants.	 70 adults with one of the following medical diagnoses whose pain remains (score ≥ 3 on item 3 of the 9-item BPI) despite their current medical treatment: cancer, amyotrophic lateral sclerosis, Parkinson's disease, spinal cord injury, neuropathy, phantom limb pain, thalamic pain, pain related to injury of nerve plexus/plexi, and neuropathic facial pain. Smoked cannabis (1 to 2 cigarettes over the course of 2 to 3 hours) administered 3 to 5 days/week in the research laboratory for 4 weeks: 1. High CBD/low THC: 15.76% CBD and 3.11% THC 2. Placebo: 0.0% CBD and 0.01% THC 	Change in pain ratings using the McGill Pain Questionnaire and the 9 item BPI at 4 weeks. Other outcomes: sickness- related impairment; physical and emotional wellbeing; QOL; cognitive status; symptom prevalence, characteristics and degree of stress; psychological state and psychological wellbeing; mood; appetite.
 Ware, M & Lynch, M (NCT02324777) Crossover RCT Sponsored by Prairie Plant Systems Inc., McGill University Health Center, Dalhousie University, Algorithme Pharma Inc., Research Institute of the McGill University Health Center May 2016 	Cannabinoid Profile Investigation of Vapourized Cannabis in Patients With Osteoarthritis of the Knee (CAPRI)	To determine the analgesic dose-response characteristics of vaporized cannabinoids with varying degrees of THC/CBD ratios.	40 adults with painful osteoarthritis of the knee (NRS Pain intensity score ≥ 4 out of 10). 100 mg of finely ground herbal cannabis drug product formulation administered via the Volcano® Medic Vapourizer (percentages are mass fractions): 1. 21.9% THC and 0.8% CBD 2. 15.0% THC and 0.8% CBD 3. 9.0% THC and 5.0% CBD 4. 3.8% THC and 9.5% CBD 4. 3.8% THC and 10.0% CBD 5. 0.6% THC and 13.0% CBD 6. Placebo: < 0.3% THC and < 0.3% CBD Participants to be randomly assigned to	Change in VAS pain intensity at 3 hours post-dose (measured every 15 minutes). Other outcomes: Stiffness; physical, social and emotional functional outcomes; psychoactive adverse events; global rating of preference; VAS of drug effect; change in blood pressure and heart rate; hematocrit, liver, and renal function (1 week after final exposure)

PI/Study Director (Registration); Study Design; Sponsors; Estimated Study Completion	Study Title	Purpose of Study	Participants; Intervention(s)/ Comparator	Outcomes and Timing
			receive all 6 formulations in random order for one day of exposure (6-days washout)	
 Wilsey, BL (<u>NCT02460692</u>) Parallel RCT Sponsored by the University of California, San Diego & National Institute on Drug Abuse (NIDA) May 2020 	Trial of Dronabinol and Vaporized Cannabis in Neuropathic Low Back Pain	To assess whether treatment with vaporized cannabis or dronabinol reduces spontaneous and evoked pain more than placebo, whether there are any differences between the 2 active treatments in terms of interference with activities of daily living, mood, neuropsychological function, and psychomimetic side-effects (high, stoned, <i>etc</i>).	 120 adults with chronic low back pain (painDETECT questionnaire score ≥ 19, and daily NRS Pain intensity ≥ 3 out of 10). One of the following for 8 weeks: 1. Vaporized cannabis: 3.5% THC 2. Dronabinol 3. Placebo 	11-point pain intensity NRS. Other outcomes: mood; depression; psychoactive effects; withdrawal; marijuana subscale of the Addiction Research Center Inventory; Cold Pressor Test; Hopkins Verbal Learning Test; Grooved Pegboard Test; Wechsler Adult Intelligence Scale-III Digit Symbol Test; and driving simulation.
 Zhao, H (<u>5R01DA030424-03</u>) Crossover RCT Sponsored by National Institute on Drug Abuse (NIDA) May 2016 	The effect of vaporized cannabis on neuropathic pain in spinal cord injury	To evaluate the analgesic effects of vaporized cannabis in patients with neuropathic pain due to spinal cord injury, as well as evaluate other potential benefits and side effects, including the effect of different strengths of cannabis on mood, cognition, and psychomotor performance.	Patients with neuropathic pain due to spinal cord injury. 1. Vaporized cannabis: 3.5% THC 2. Vaporized cannabis: 7.0% THC 3. Placebo	Pain intensity and pain unpleasantness (timing NR). Other outcomes: neuropsychological functioning (attention, learning and memory, and psychomotor performance), emotional response/mood.

Abbreviations: BPI = Brief Pain Inventory; CBD = cannabidiol; NR = not reported; NRS = Numeric Rating Scale; QOL = quality of life; RCT = randomized controlled trial; THC = tetrahydrocannabinol; VAS = Visual Analog Scale.

^a Unpublished studies completed in June 2015 or later are included in the table in order to allow time for publication.

PTSD

There are 2 recently initiated studies on the benefits and harms of cannabis for PTSD using an RCT design that should add to the body of evidence (Table 8). The Colorado Department of Public Health and Environment has funded a "triple-blind cross-over placebo-controlled" trial to determine the effects of smoking 4 different types of cannabis with varying THC and CBD content on PTSD symptoms in Veterans (Bonn-Miller, NCT02759185). The anticipated completion date of the trial is April 2019. Second, Eades et al are conducting a study sponsored by Tilray and the University of British Columbia (NCT02517424). This study is a cross-over RCT of 42 adults with PTSD who will be administered differing amounts of THC and CBD (High/Low, High/High, and Low/Low) to compare PTSD outcomes as well as other mental and physical health outcomes.

There are also multiple ongoing studies of cannabis and PTSD that are not RCTs, or that investigate cannabis-related outcomes but do not specifically test the effectiveness of cannabis for reducing PTSD symptoms. For example, a VA-funded trial is described as investigating the impact of cognitive behavioral therapy for insomnia on cannabis cessation. Bonn-Miller and colleagues are investigating how cannabis use impacts PTSD and sleep in an unfunded observational study of 150 Veterans. Finally, another study funded by The Colorado Department of Public Health and Environment is assessing 150 individuals with PTSD to determine if recent medical or recreational cannabis use versus no cannabis use in the past 6 months is associated with differential trajectories of PTSD symptoms over the course of a year. Table 8 provides a summary of ongoing studies related to benefits and harms of cannabis for PTSD.

Table 8. Ongoing Studies^a of Cannabis for PTSD

PI/Study Director (Registration); Study Design; Sponsors; Estimated Study Completion	Study Title	Purpose of Study	Participants; Intervention(s)/ Comparator	Primary Outcome and Timing
Babson, K (NCT02102230) • Double-blind RCT • Funded by VA Clinical Science Research and Development CDA-2 • August 2019	The Impact of CBT-I on Cannabis Cessation Outcomes	To examine the role of a behavioral intervention for sleep on cannabis use frequency and insomnia symptoms among Veterans with CUD and insomnia.	 200 Veterans with CUD and insomnia. Randomly assigned to of the following conditions: 1. CBT for insomnia 2. CBT for insomnia + CBT-I coach (mobile app) 3. Placebo control (quasi- desensitization) 	Change in cannabis use frequency, point prevalence abstinence, and change is sleep quality post-treatment and 6 months post-treatment.
Bedard-Gilligan, M (NCT02874898) • Single Group Assignment • Funded by the National Institute on Drug Abuse (NIDA) • April 2019	Marijuana Use, Extinction Learning, and Exposure Therapy in Individuals with PTSD	To examine the effects of cannabis use on extinction learning using both a standard discriminative conditioning and extinction task at pre-treatment and response to an exposure treatment protocol. To also examine ability of a brief protocol to decrease PTSD and retain individuals in treatment for patients with and without cannabis use.	72 men and women (ages 18-65) with chronic PTSD (≥ 3 months); half are heavy cannabis smokers (≥ 5 days per week) and half are non- cannabis users (no use in last 3 months). Brief imaginal exposure protocol (6 daily sessions) for PTSD is provided to all participants.	PTSD severity (PSS-I severity) at post- treatment and 12-week follow-up; treatment drop-out (completion of less than 5 imaginal exposure sessions). Other Outcomes: Depression symptoms (QIDS), cannabis use and problems (MPS, Marijuana Frequency and Quantity Scale) assessed at post- treatment and 12-week follow-up.
Bonn-Miller, M (NCT02759185) • Crossover RCT	Placebo-Controlled, Triple Blind, Randomized Crossover Pilot Study	To evaluate the safety and efficacy of smoked cannabis of 4 different concentrations among	76 Veterans with service-related PTSD (≥ 6 months duration, moderate severity at baseline)	Change in CAPS Global Severity Score at 3 weeks and 8 weeks after randomization.
 Funded by The Colorado Department of Public Health and Environment April 2019 	of the Safety and Efficacy of Four Potencies of Smoked Marijuana in 76 Veterans with Chronic, Treatment-Resistant Posttraumatic Stress	participants with chronic,	 Smoked cannabis up to 1.8 g/day for 3 weeks: 1. High THC (more THC than CBD) 2. High CBD (more CBD than THC) 3. High THC/High CBD (equal amounts) 4. Placebo cannabis (low levels THC/CBD) 	Other outcomes: depression and anxiety symptoms; general and psychosocial functioning; sleep quality; suicidal ideation; responses to cannabis; withdrawal; blood and urine tests.

PI/Study Director (Registration); Study Design; Sponsors; Estimated Study Completion	Study Title	Purpose of Study	Participants; Intervention(s)/ Comparator	Primary Outcome and Timing
	Disorder (PTSD)		Participants to receive 2 of the 4 types of cannabis during 2 stages, each lasting 3 weeks (2-week washout).	
Bonn-Miller, M • Observational Study • Unfunded • June 2017	Evaluation of Veteran Cannabis Use and Impact on Sleep and PTSD	The present study aims to fill a large gap in the literature by providing an a priori test of the impact of cannabis, including variations in cannabinoids, on individual sleep, PTSD, and psychosocial functioning.	 150 Veterans currently using cannabis and are members of the Santa Cruz Veterans Alliance. Data are collected through repeated survey assessments every other week. In addition, all product provided to Veterans by the Santa Cruz Veterans Alliance is tested for cannabinoid content by an independent laboratory. 	The association between cannabinoid concentration and symptoms of PTSD, sleep, and psychosocial functioning over time among cannabis-using Veterans.
 Bonn-Miller, M Observational Study Funded by The Colorado Department of Public Health and Environment September 2018 	Treating PTSD with Marijuana: Clinical and Functional Outcomes	The proposed study aims to determine whether, among a sample of Colorado residents (Veterans and non- Veterans), individuals with PTSD who obtain and use cannabis from a medical or recreational dispensary, compared to a matched sample of individuals with PTSD who report no current cannabis use at study baseline (control), will exhibit lower PTSD symptom severity.	 150 adult Colorado residents with PTSD, half using cannabis from a medical or recreational dispensary in Colorado and half reporting no recent (past 6 month) cannabis use. Assessment at baseline and 3-, 6-, 9-, and 12-months following baseline. Measures include interview (MINI, CAPS- 5, TLFB), self-report, computerized neuro- psych assessments, and actigraphy for 1 week following each assessment point, and urine tests for objective verification of use status. Further, those using cannabis will report on the cannabis used and the dispensary from which it is obtained, and a sample will be procured and tested for cannabinoid and terpene content. 	PTSD symptom severity, as indexed by: (1) Self-reported overall symptom severity at each time point as assessed by the CAPS-5; (2) Self-reported and objective sleep quality at each time point as assessed by the PSQI and actigraphy; (3) Interview-based diagnosis at 12-month follow-up as assessed by the CAPS-5. Secondary Outcomes (assessed at each timepoint): (1) self-reported and objective psychosocial functioning; (2) suicidal ideation; (3) engagement in medical and psychological services.
Browne, K Mixed Methods 	Characterizing Cannabis Use in	The objective of this study is to build our	Veterans diagnosed with PTSD who report at least weekly cannabis use will be	Conduct an online survey in order to characterize cannabis use patterns

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PI/Study Director (Registration); Study Design; Sponsors; Estimated Study Completion	Study Title	Purpose of Study	Participants; Intervention(s)/ Comparator	Primary Outcome and Timing
Observational Study • University of Washington Alcohol and Drug Abuse Institute • VA Puget Sound Health Care System Research & Development • September 2017	Veterans with PTSD	understanding of cannabis use in Veterans with PTSD by: 1) characterizing cannabis use patterns and motives in Veterans with PTSD symptoms, 2) conducting a prospective examination of the day-to- day relations between PTSD symptoms and cannabis use, and 3) conducting the first effort to qualitatively describe the perspective of Veterans with PTSD who use cannabis.	invited to participate in: 1. Anonymous online survey (n=200) 2. Daily symptom and use monitoring (<i>ie</i> , IVR; n=48) 3. In-depth qualitative interviews (n=30) 4. Blood draw for cannabis biomarkers (n=48)	 and replicate previous findings related to PTSD symptoms, cannabis use, motives for use, and craving. Examine (via IVR) day-to-day relations between cannabis use and PTSD symptoms along with a one-time assessment of cannabis use motives. To conduct key informant interviews in order to characterize Veterans' beliefs about the relations between cannabis use and mental health symptoms and treatment, including the role of cannabis in PTSD symptom management, treatment for cannabis use, and PTSD treatment.
 Eades, J (NCT02517424) Crossover RCT Sponsored by Tilray and the University of British Columbia December 2018 	Placebo-Controlled, Triple-Blind, Crossover Study of the Safety and Efficacy of Three Different Potencies of Vaporized Cannabis in 42 Participants with Chronic, Treatment- Resistant Posttraumatic Stress Disorder (PTSD)	To evaluate the safety and efficacy of vaporized cannabis of 3 different concentrations among participants with chronic, treatment-resistant PTSD.	42 adults with PTSD (≥ 6 months duration, PCL-5 ≥ 40 at baseline). Approximately 50% police/military Veterans, 33-50% female, and 8-12% Aboriginal (First Nations, Metis, Inuit). Cannabis administered via vaporization up to 2.0 g/day as needed: 1. High THC/Low CBD cannabis 2. High THC/High CBD cannabis 3. Low THC/Low CBD cannabis	Change in CAPS Global Severity Score at 3 weeks and 8 weeks after randomization. Other outcomes: anxiety and depression symptoms; psychosocial functioning; preference; sleep quality; problems associated with cannabis use; suicidal thoughts or behaviors.

Abbreviations: CAPS = Clinician-Administered PTSD Scale; CBD = cannabidiol; CBT = cognitive behavioral therapy; CDA-2 = VA Career Development Award 2; CUD = cannabis use disorder; IVR = interactive voice response; MINI = Mini International Neuropsychiatric Interview; MPS = Marijuana Problems Scale; TLFB = Timeline Followback interview; PCL = Post-traumatic Stress Disorder Checklist; PSQI = Pittsburgh Sleep Quality Index; PSS = Posttraumatic Stress Disorder Symptom Scale-Interview Version; PTSD = post-traumatic stress disorder; QIDS = Quick Inventory of Depressive Symptomatology; RCT = randomized controlled trial; THC = tetrahydrocannabinol; VA = Department of Veterans Affairs.

^a Unpublished studies completed in June 2015 or later are included in the table in order to allow time for publication.

SUMMARY AND DISCUSSION

We reviewed the literature examining the benefits of cannabis in chronic pain and PTSD populations, as well as literature examining potential harms relevant to these populations. Table 10 summarizes the evidence on the benefits and harms of cannabis use. Overall, we found limited evidence on the potential benefits and harms of cannabis use in chronic pain populations. We found low-strength evidence that cannabis preparations with precisely defined THC-cannabidiol content (most in a 1:1 to 2:1 ratio) may alleviate neuropathic pain but insufficient evidence in populations with other types of pain. The applicability of these findings to current practice may be low, in part because the formulations studied may not be reflective of what most patients are using, and because the consistency and accuracy of labeled content in dispensaries are uncertain.¹²⁵ Furthermore, most studies are small, many have methodological flaws, and the long-term effects are unclear given the brief follow-up of most studies. There is insufficient evidence of effects on quality of life or functional status.

Among neuropathic pain studies, we found a discrepancy between continuous and dichotomous pain outcomes. Possible interpretations are that cannabis is simply not consistently effective or that, although cannabis may not have clinically important effects on average, subgroups of patients may experience large effects. We did not find data to clarify which subgroups of patients are more or less likely to benefit.

We found no trials examining the effects of cannabis in PTSD populations, and there was insufficient evidence from observational studies to draw conclusions about its effectiveness in patients with PTSD.

Even though we did not find strong evidence of benefit for most indications, clinicians will still need to counsel patients with chronic pain or PTSD who are using or requesting to use cannabis for therapeutic or recreational purposes. Therefore, understanding what is known and not known about potential harms of cannabis is also important.

There is moderate-strength evidence that at least light to moderate cannabis smoking does not adversely impact lung function over about 20 years. However, there is no evidence examining the effects in older patients, or those with multiple medical comorbidities. Moreover, the limited evidence examining the effects of heavy use (the equivalent of one joint daily for 7 years or more) suggests a possible deleterious effect on lung function over time.

There is low-strength evidence that light to moderate cannabis use is not associated with lung cancer or head and neck cancer diagnoses independent of tobacco use, but the data are limited to case-control studies and do not address heavy use. However, there is at least biologic plausibility that cannabis smoking has the potential to increase the risk of lung cancer based on data showing that cannabis use is associated with macrophage dysfunction, tar deposition, and cytologic abnormalities.¹²⁶ There is insufficient evidence about effects on other cancers.

While there is a biologically plausible link between cannabis use and cardiovascular risk given data showing adverse effects on hemodynamic parameters and anginal threshold,⁷³ we found insufficient evidence examining whether cannabis use is associated with cardiovascular events over the long-term.

There are potentially serious mental health and adverse cognitive effects of cannabis, though there is not enough data to characterize the magnitude of risk or in whom the risk is highest.



Cannabis appears to be associated with at least small, short-term deleterious effects on cognition in active users, but long-term effects in past users are uncertain. We found no data on the risk of mania or suicidality in chronic pain or PTSD populations specifically, but cannabis has been associated with these risks in other populations.

We found stronger data suggesting an association between cannabis use and the development of psychotic symptoms over the long-term and limited data suggesting a risk of acute psychosis immediately following cannabis use. There is no data to directly assess whether the risk of psychotic symptoms is related specifically to the THC content of the formulation used, but this is biologically plausible, there are case reports of severe acute psychosis after ingestion of edibles with very high THC concentrations,¹²³ and CBD may in fact have antipsychotic effects.^{127,128}

Intuitively, patients with PTSD or patients with serious mental illness, especially those already suffering with hypervigilance, agitation, and anger management issues, might be at higher risk of suffering serious consequences should they experience any adverse effects, especially psychotic symptoms. Observational studies in PTSD populations suggest a signal for harm, though the studies are inconclusive.^{60,61} While clinicians do not have adequate data to quantify risks and benefits for PTSD patients, they might consider discussing potentially serious mental health adverse effects during shared decision-making discussions. They also might consider discussing other evidence-based interventions recommended by the 2010 VA/Department of Defense (DoD) Clinical Practice Guideline for PTSD.¹²⁹ Specifically, "A" level interventions with "strong recommendations" for use include selective serotonin reuptake inhibitors (SSRIs), selective norepinephrine reuptake inhibitors (SNRIs), and "trauma-focused psychotherapy that includes components of exposure and/or cognitive restructuring, or stress inoculation training." Similar recommendations based on research synthesized through 2013 were made by the Institute of Medicine (IOM).¹³⁰ This IOM report noted, "A 2013 meta-analysis of treatment efficacy for PTSD was consistent with the VA/DoD guideline in finding that cognitive therapy including cognitive processing therapy (CPT); exposure therapy, such as prolonged exposure (PE) therapy; and eye movement desensitization and reprocessing (EMDR) were effective psychotherapies, and SSRIs were the most effective pharmacotherapies."

Finally, there are a number of adverse effects that appear to be related to cannabis use and may be important for clinicians to be familiar with, but whose incidence has not been wellcharacterized. These are reviewed above in the emerging harms section and include infectious disease complications, cannabis hyperemesis syndrome, inadvertent overingestion of THC and associated psychosis related to edible cannabis, and violent behavior.

Currently, the Centers for Disease Control and Prevention recommends the use of evidencebased non-pharmacologic therapy – such as physical therapy, exercise therapy, and psychologic interventions – and non-opioid pharmacologic therapy as the preferred modalities to treat chronic pain.¹³¹ After trying first-line options, clinicians may continue to struggle with the often difficult treatment of chronic pain in patients who have not responded. Cannabis may be perceived as a safer strategy in these patients.¹³² Indeed, the scale and severity of adverse events, including death, seen with opioids have not been described with cannabis use in the literature (though there is also simply less research available on cannabis than opioids).¹³² However, there are no studies directly comparing cannabis to opioids, and there is no good-quality data examining what impact cannabis use actually has on opioid use and opioid-related adverse effects. We found no observational studies that met inclusion criteria, but a growing body of cross-sectional literature suggests negative opioid-related correlates among individuals who use cannabis and opioids concurrently. These include opioid misuse;^{9,10,99,133} a greater number of opioid refills;⁹⁹ a longer



duration of opioid use; a higher dose of opioid medication prescribed;⁹ and endorsement of using opioids and other pain medications without a prescription.¹³⁴ By contrast, one recent open-label study found that pain scores and opioid use decreased over 6 months in a chronic pain population who initiated cannabis treatment, though confidence in the findings is limited by the lack of a control group and the large number of participants lost to follow-up.¹³⁵

LIMITATIONS

There are a number of limitations to this body of evidence beyond the paucity of well-conducted trials of treatment efficacy. The methodologic issues with each particular trial and observational study are detailed in the quality assessment tables (Appendix C). Applying available data to clinical practice is challenging for several reasons. The data on effectiveness largely comes from studies examining cannabis formulations with known THC and CBD content (most with 1:1 to 2:1 ratio). While dispensaries are increasingly labeling the content of offered products, there are often important discrepancies between labeled and measured content.¹²⁵

While trials were often able to standardize the dosing of the active ingredients in cannabis (THC and CBD), most of the observational studies were not able to characterize the amount of cannabis consumed beyond rough measures such as the average number of joints smoked per day. No observational studies were able to account for the potency of cannabis consumed. In a sense, this lack of precise dosing information reflects the reality of clinical practice and, therefore, the crude approximations of exposure in most studies may still provide useful information. Nevertheless, the evidence base is limited in providing very exact dose-response information beyond the relative distinctions between very heavy and infrequent use. Moreover, the evidence base on harms is limited because there are relatively few patients included in studies with a history of heavy and prolonged cannabis use.

There are also limitations in our approach to synthesizing this literature. Given the broad scope of our review, we relied on existing systematic reviews when available to identify the best available evidence. We believe we are unlikely to have missed important studies both because we only used systematic reviews meeting key quality criteria and because we searched the primary literature for more recent studies not captured by the reviews. As our intention was to provide an overview of evidence that would be important for clinicians to know in counseling patients, we included studies of harms in general populations when we thought it unlikely that the conditions of chronic pain or PTSD would independently contribute to risk (*eg*, pulmonary or cardiovascular harms when concurrent tobacco use was accounted for). Though we made these determinations through group discussion and in conjunction with a panel of experts, we acknowledge that the choices are inherently subjective to some degree and that there is still the possibility that there are residual confounders relevant to chronic pain or PTSD accounting for observed effects.

FUTURE RESEARCH

There is virtually no conclusive information about the benefits of cannabis in chronic pain or PTSD populations and limited information on harms, so methodologically strong research in almost any area of inquiry is likely to add to the strength of evidence. Fortunately, it appears that the US government is poised to lift restrictions on access to cannabis for research which should help speed the development of this evidence base which has lagged far behind policy changes regarding the use of cannabis for medical purposes in many states.¹³⁶ Also, there are studies currently being done which should also add to the evidence base in the near future (and are



summarized in Key Question 4). Table 9 lists opportunities for future research in each of the areas we reviewed.

Area of Inquiry	Research Suggestions
Efficacy of cannabis for treating chronic pain	 Populations other than MS or neuropathic pain Studies with longer follow-up duration Studies with cannabis-naïve patients Compare cannabis to other active treatments for pain, including opioids Use cannabis preparations that are routinely available to consumers in the US, especially given legalization in more states Examine the effects of different THC:CBD ratio preparations, and more study of CBD preparations Obtain blood levels of THC and CBD to assess actual level of drug exposure
Efficacy of cannabis for treating PTSD	 RCT of treatment Trials comparing to cognitive behavioral therapy, other standard treatments
CUD	Studies assessing risk of CUD in patients using cannabis
Pulmonary harms	Observational studies in older and multimorbidity populations
Cardiovascular harms	 Observational studies with more comprehensive information about exposure history
Cancer	 Larger scale observational studies of lung cancer reflecting patterns of use in the US More studies to investigate the insufficient evidence of a possible link with testicular and transitional cell cancers
Mental health harms	 Studies on acute psychosis in chronic pain and PTSD populations Identification of non-schizophrenic patients at high risk for psychosis Risk mitigation strategies for cannabis-induced psychosis Studies on mania and suicidality in PTSD populations Effects on sleep
Cognitive function	Studies in chronic pain and PTSD populations
Emerging harms	 Studies characterizing cannabis hyperemesis syndrome in a larger number of patients Studies examining treatment and follow-up of patients with cannabis hyperemesis syndrome

 Table 9. Suggestions for Future Research

CONCLUSIONS

Although cannabis is increasingly available for medical and recreational use, there is very little methodologically rigorous evidence examining its effects in patients with chronic pain or PTSD. Limited evidence suggests that cannabis may alleviate neuropathic pain, but there is insufficient evidence in other populations. There is insufficient evidence examining the effects of cannabis in PTSD populations. Among general populations, limited evidence suggests that cannabis is associated with an increased risk for potentially serious mental health adverse effects, such as psychosis. Data on its effects on long-term physical health vary; harms in older patients or those with multiple comorbidities have not been studied.

Table 10. Summary of Evidence for the Benefits and Harms of Cannabis in Chronic Pain or PTSD Populations

	N studies	Findings	Strength of Evidence ^a	Comments
Chronic Pain				
 Multiple sclerosis (MS) 	 4 Low ROB studies (combined N=1017; 24 to 424 per study): 2 of THC/CBD capsules 1 of nabiximols 1 of sublingual spray delivering THC, CBD, or THC/CBD combined 3 Unclear ROB studies of nabiximols (combined N=562; 36 to 337 per study) 	with THC (2.5 mg)/CBD (1.25 mg) capsules in 2 studies. A 12-week study of nabiximols (2.7 mg THC/2.5 mg CBD oromucosal spray)	Low	Few methodologically rigorous studies, but fair number of patients; inconsistent results; little long-term data; restrictive entry criteria in largest study which only included patients with initial response in run-in phase; applicability to
	7 High ROB studies (combined N=430; 13	reported significant improvement in		formulations available in dispensaries may be low
	to 160 per study):	A sublingual spray delivering 2.5 mg of		disperisaries may be low
	- 3 of nabiximols	CBD, THC, or both for sequential 2-week		
	 2 of THC/CBD capsules 1 of smoked THC 	periods reported mixed effects. THC alone significantly improved pain and		
	- 1 of oral THC	spasticity, but CBD alone and THC/CBD combined had inconsistent effects.		
	 4 Low ROB studies (combined N=1017; 24 to 424 per study): 2 of THC/CBD capsules 1 of nabiximols 1 of sublingual spray delivering THC, CBD, or THC/CBD combined 	Other outcomes: Small improvements in sleep in 4 studies: Self-reported sleep quality improved in 2 studies of THC/CBD capsules. Nabiximols were significantly superior to placebo for reducing sleep disruption in a 12-week study (N=241). Sleep improved significantly in a small study (N=24) of a sublingual spray containing 2.5 mg each of CBD:THC. Other:	Low (sleep)	Few methodologically rigorous studies, but fair number of patients; inconsistent results; little long-term data; restrictive entry criteria in largest study which only included patients with initial response in run-in phase; applicability to current practice may be low
		Nabiximols were significantly superior to placebo for Barthel Activities of Daily Living (P =.0067), Physician Global Impression of Change (P =.005), Subject Global Impression of Change (P =.023), and Carer Global Impression of Change (P =.005) in Function in a 12-week study (N=241).	Insufficient (other outcomes)	Only one study of nabiximols – not tested otherwise

	N studies	Findings	Strength of Evidence ^a	Comments
Neuropathic pain	 11 low ROB studies (combined N = 593) 4 of smoked THC (combined N = 150) 3 of vaporized THC (combined N = 97) 3 of nabiximols (combined N = 312) 1 of oromucosal spray delivering THC or THC+CBD (N = 34) 1 unclear ROB study of nabiximols (N = 30) 1 high ROB trial (N = 125) 	Studies did not find a clinically significant between-group difference on continuous pain scales, but a higher proportion of intervention patients had clinically significant pain relief up to several months later. In a meta-analysis of 9 studies, intervention patients were more likely to report \geq 30% improvement in pain (combined RR, 1.43 [95% CI, 1.16–1.88]; P = 38.6%; $P = 0.111$).	Low	Few patients enrolled in most low ROB studies; inconsistent results; marked differences among studies in dosing and delivery mechanism; brevity of study duration; low applicability to formulations available in dispensaries.
	1 Low ROB study of smoked THC (N=23)	Other outcomes reported in low ROB studies: A study of vaporized cannabis reported that 25 mg with 9.4% THC administered as a single smoked inhalation 3 times daily resulted in significant improvements in sleep quality.	Insufficient	Only one small study
 General/other/mixed populations 	 2 Low ROB studies: 1 trial of sublingual spray delivering THC, CBD, or THC/CBD combined (N=34) 1 observational study of cannabis containing 12.5% THC (smoked, oral, or vaporized) (N=431) 3 Unclear ROB studies of nabiximols (combined N=428; 10 to 360 per study) 3 High ROB studies (combined N=265; 18 to 177 per study): 	Small improvements in pain, but no effect on sleep, mood, quality of life.	Insufficient	Only one small low ROB study in which the bulk of the patients had MS; larger observational study had high drop-out rate
	 - 2 of nabiximols - 1 of THC capsules 			
PTSD	2 observational studies in Veterans with PTSD: - 1 Medium ROB (N=2276) - 1 High ROB (N=700)	Cannabis was not associated with an improvement in mental health symptoms.	Insufficient	No trials; only 2 observational studies with methodologic flaws

Benefits and Harms of Cannabis for Chronic Pain or PTSD

Evidence-based Synthesis Program

	N studies	Findings	Strength of Evidence ^a	Comments
larms General AEs	2 systematic reviews of chronic pain	Cannabis-based treatments were associated with an overall higher risk of		Consistent findings except for serious AE
		short-term, non-serious AEs.		SCHOUS AL
Medical harms				— и и
Ø Pulmonary function	2 Low ROB prospective cohort studies with 20-32 years follow-up (combined N=6053)	In young adults, low levels of cannabis smoking did not adversely affect lung function over about 20 years. A previous meta-analysis of 5 studies	Young adults: Moderate Older adults: No evidence	Two well-done prospective cohort studies, but limited information about effects of heavy use and no information
	1 systematic review of 5 observational studies (3 cohort, 2 cross-sectional) (combined N=851)	found no increased risk for pulmonary adverse effects, OR (95% CI): 0.80 (0.46-1.39).		in older or multimorbid populations
Ø Cardiovascular	2 High ROB observational studies: - 1 case-crossover (N=3882) - 1 cohort study (N=2097)	Cannabis use at the time of myocardial infarction was not associated with mortality after mean 12.7 years follow-up, but longitudinal use was not assessed. Risk of myocardial infarction within an hour of cannabis use was significantly elevated compared with periods of non- use but this finding may be inflated by recall bias, OR (95% CI): 4.8 (2.9-9.5).	Insufficient	Recall bias; inadequate controlling for confounders; lack of longitudinal exposure data
ØCancer				
§ Lung	1 patient-level meta-analysis of 6 case- control studies (2150 cases)	The meta-analysis found no association between light cannabis use and lung cancer.	Low	Recall bias; mostly light users, few heavy users; the large cohort study had no
	1 High ROB cohort study (N=49,231)			information about exposure over time
§ Head/neck/o ral	Meta-analysis of 9 case-control studies (5732 cases)	No association between cannabis use and cancer, OR (95% CI): 1.02 (0.91- 1.14); generally consistent across studies and no evidence of dose-response.	Low	Imprecise exposure measurement with potential recall bias; ever use among studies ranged from 1 to 83%
§ Testicular	Meta-analysis of 3 High ROB case-control studies (719 cases)	An increase in cancer risk for weekly users compared to never-users appeared with non-seminoma cancers but not seminoma cancers, OR (95% CI): 1.92 (1.35-2.72).	Insufficient	Potential confounding from recall bias and tobacco use
§ Transitional cell	1 High ROB VA case-control study (52 cases)	Risk of cancer with > 40 joint-years cannabis use compared to none, OR 3.4 (unadjusted, <i>P</i> =.012).	Insufficient	One very small case-control study with several methodologic flaws

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Benefits and Harms of Cannabis for Chronic Pain or PTSD

Evidence-based Synthesis Program

	N studies	Findings	Strength of Evidence ^a	Comments
Ø Motor vehicle accidents	Meta-analysis of 21 observational studies (combined N=239,739)	Increase in collision risk, OR (95% CI): 1.35 (1.15-1.61).	Moderate	The small but significant increase in risk was seen consistently across numerous sensitivity analyses and after adjustment in meta- regression analyses
 Mental health Ø Suicidal 	No studies in chronic pain or PTSD		No evidence	Meta-analysis of 4 studies in
behaviors	populations.		(chronic pain or PTSD)	the general population reported significantly increased odds of suicide with any cannabis use, OR (95% CI): 2.56 (1.25-5.27).
Ø Mania	No studies in chronic pain or PTSD populations		No evidence (chronic pain or PTSD)	A systematic review found an increased incidence of new- onset mania symptoms among populations without a diagnosis of bipolar disorder, OR (95% CI): 2.97 (1.80 to 4.90).
Ø Psychosis	1 systematic review	History of cannabis use was associated with an increase in risk of developing	Low	Consistent evidence from large observational studies
	 7 studies including patients without psychotic symptoms at baseline: 3 Low ROB studies 3 Medium ROB studies 1 High ROB study 	psychotic symptoms.		and some evidence of increased risk with higher levels of use; consistent with data from small experimental studies suggesting risk of acute psychosis in some patients; magnitude of risk unclear and not specifically studied in chronic pain or PTSD populations

	N studies	Findings	Strength of Evidence ^a	Comments
Ø Cognitive effects	1 systematic review of 33 studies	Active long-term cannabis use associated with small negative effects on all aspects of cognition. Mixed, inconsistent findings on long-term effects in past users.	Moderate Insufficient (past use)	Consistent data from large number of studies on effects on active long-term use, but inconsistent findings from smaller number of studies regarding effects in those that were abstinent and no data available specifically in chronic pain or PTSD populations
ØCUD	One large cohort study (N=34,653; N = 1279 past year cannabis use in last year)	OR incident CUD 9.5 (95% CI 6.4-14.1) Prevalence CUD (among those using in last year) 36% Prevalence past year cannabis dependence 7.7% Prevalence past year cannabis abuse 28%	Low	In cross-sectional studies, the prevalence of CUD in chronic pain populations was about 2%

Abbreviations: AE = adverse event; CBD = cannabidiol; CI = confidence interval; CUD = cannabis use disorder; MS = multiple sclerosis; N = number; OR = odds ratio; PTSD = post-traumatic stress disorder; ROB = risk of bias; THC = tetrahydrocannabinol; VA = Department of Veterans Affairs.

^a The overall quality of evidence for each outcome is based on the consistency, coherence, and applicability of the body of evidence, as well as the internal validity of individual studies. The strength of evidence is classified as follows:

- High = Further research is very unlikely to change our confidence on the estimate of effect.
- Moderate = Further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.
- Low = Further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.
- Insufficient = Any estimate of effect is very uncertain.

REFERENCES

- 1. Hasin DS, Saha TD, Kerridge BT, et al. Prevalence of Marijuana Use Disorders in the United States Between 2001-2002 and 2012-2013. *JAMA psychiatry*. 2015;72(12):1235-1242.
- 2. Ryan-Ibarra S, Induni M, Ewing D. Prevalence of medical marijuana use in California, 2012. *Drug and alcohol review*. 2015;34(2):141-146.
- 3. Adler JN, Colbert JA. Clinical decisions. Medicinal use of marijuana--polling results. *N Engl J Med.* 2013;368(22):e30.
- 4. 114th Congress (2015-2016). H.R.2577 Military Construction, Veterans Affairs, and Related Agencies Appropriations Act, 2017. <u>https://www.congress.gov/bill/114thcongress/house-bill/2577/summary/40</u> Accessed August 24, 2016.
- 5. Institute of Medicine (US). *Relieving Pain in America: A Blueprint for Transforming Prevention, Care, Education, and Research.* Washington, DC: National Academies Press;2011.
- 6. Ward BW, Schiller JS, Goodman RA. Multiple chronic conditions among US adults: a 2012 update. *Prev Chronic Dis.* 2014;11:E62.
- 7. Bonn-Miller M, Boden M, Bucossi M, Babson K. Self-reported cannabis use characteristics, patterns and helpfulness among medical cannabis users. *Am J Drug Alcohol Abuse*. 2014;40(1):23-30.
- 8. Ilgen MA, Bohnert K, Kleinberg F, et al. Characteristics of adults seeking medical marijuana certification. *Drug and alcohol dependence*. 2013;132(3):654-659.
- 9. Degenhardt L, Lintzeris N, Campbell G, et al. Experience of adjunctive cannabis use for chronic non-cancer pain: findings from the Pain and Opioids IN Treatment (POINT) study. *Drug and alcohol dependence*. 2015;147(ebs, 7513587):144-150.
- 10. Reisfield GM, Wasan AD, Jamison RN. The prevalence and significance of cannabis use in patients prescribed chronic opioid therapy: a review of the extant literature. *Pain medicine (Malden, Mass).* 2009;10(8):1434-1441.
- 11. Bowles DW. Persons registered for medical marijuana in the United States. *J Palliat Med.* 2012;15(1):9-11.
- 12. Boden MT, Babson KA, Vujanovic AA, Short NA, Bonn-Miller MO. Posttraumatic stress disorder and cannabis use characteristics among military veterans with cannabis dependence. *The American journal on addictions / American Academy of Psychiatrists in Alcoholism and Addictions.* 2013;22(3):277-284.
- Kansagara D, O'Neil ME, Morasco B, et al. Cannabis for the management of symptoms of chronic pain and/or PTSD. PROSPERO International prospective register of systematic reviews.

http://www.crd.york.ac.uk/prospero/display_record.asp?ID=CRD42016033623 Accessed August 24, 2016.

- 14. Whiting PF, Wolff RF, Deshpande S, et al. Cannabinoids for Medical Use: A Systematic Review and Meta-analysis. *JAMA*. 2015;313(24):2456-2473.
- Butler M, Krebs E, Sunderlin B, Kane R. Medical Cannabis for Non-Cancer Pain: A Systematic Review. 2015; <u>http://www.health.state.mn.us/topics/cannabis/intractable/medicalcannabisreport.pdf</u>. Accessed April 26, 2016.
- 16. Wilkinson ST, Radhakrishnan R, D'Souza DC. A Systematic Review of the Evidence for Medical Marijuana in Psychiatric Indications. *J Clin Psychiatry*. 2016;77(8):1050-1064.
- 17. Oregon Health Authority. Medical Marijuana Rules and Statutes. *Medical Marijuana Program (OMMP)*.



https://public.health.oregon.gov/DiseasesConditions/ChronicDisease/MedicalMarijuanaP rogram/Pages/legal.aspx. Accessed October 3, 2016.

- 18. Shea BJ, Grimshaw JM, Wells GA, et al. Development of AMSTAR: a measurement tool to assess the methodological quality of systematic reviews. *BMC medical research methodology*. 2007;7:10.
- 19. Wallace BC, Small K, Brodley CE, Lau J, Trikalinos TA. Deploying an interactive machine learning system in an evidence-based practice center: abstrackr. *Proceedings of the 2nd ACM SIGHIT International Health Informatics Symposium (IHI).* 2012:819-824.
- 20. Higgins J, Green S (editors). Cochrane Handbook for Systematic Reviews of Interventions Version 5.1.0 [updated March 2011]. Chapter 8: Assessing risk of bias in included studies. The Cochrane Collaboration, 2011. Accessed at www.handbook.cochrane.org on January 16, 2017. 2011.
- 21. Wells GA, Shea B, O'Connell D, et al. The Newcastle-Ottawa Scale (NOS) for assessing the quality of nonrandomized studies in meta-analyses. <u>http://www.ohri.ca/programs/clinical_epidemiology/oxford.asp</u>. Accessed January 11, 2016.
- 22. Viswanathan M, Ansari M, Berkman N, et al. *Assessing the Risk of Bias of Individual Studies in Systematic Reviews of Health Care Interventions*. Rockville, MD: Agency for Healthcare Research and Quality; Methods Guide for Comparative Effectiveness Reviews (AHRQ Publication No. 12-EHC047-EF);2012.
- 23. Hardy RJ, Thompson SG. A likelihood approach to meta-analysis with random effects. *Stat Med.* 1996;15(6):619-629.
- 24. Higgins JP, Thompson SG, Deeks JJ, Altman DG. Measuring inconsistency in metaanalyses. *Bmj.* 2003;327(7414):557-560.
- 25. Berkman N, Lohr K, Ansari M, et al. *Grading the Strength of a Body of Evidence When Assessing Health Care Interventions for the Effective Health Care Program of the Agency for Healthcare Research and Quality: An Update.* Rockville, MD: Agency for Healthcare Research and Quality; Methods Guide for Comparative Effectiveness Reviews (AHRQ Publication No. 13(14)-EHC130-EF);2013.
- 26. Atkins D, Chang S, Gartlehner G, et al. *Assessing the Applicability of Studies When Comparing Medical Interventions.* Rockville, MD: Agency for Healthcare Research and Quality; Methods Guide for Comparative Effectiveness Reviews (AHRQ Publication No. 11-EHC019-EF);2011.
- 27. Novotna A, Mares J, Ratcliffe S, et al. A randomized, double-blind, placebo-controlled, parallel-group, enriched-design study of nabiximols (Sativex()), as add-on therapy, in subjects with refractory spasticity caused by multiple sclerosis. *European Journal of Neurology*. 2011;18(9):1122-1131.
- 28. Ungerleider JT, Andyrsiak T, Fairbanks L, Ellison GW, Myers LW. Delta-9-THC in the treatment of spasticity associated with multiple sclerosis. *Advances in alcohol & substance abuse*. 1987;7(1):39-50.
- 29. Notcutt W, Langford R, Davies P, Ratcliffe S, Potts R. A placebo-controlled, parallelgroup, randomized withdrawal study of subjects with symptoms of spasticity due to multiple sclerosis who are receiving long-term Sativex (nabiximols). *Multiple sclerosis (Houndmills, Basingstoke, England).* 2012;18(2):219-228.
- 30. Wade DT, Robson P, House H, Makela P, Aram J. A preliminary controlled study to determine whether whole-plant cannabis extracts can improve intractable neurogenic symptoms. *Clinical rehabilitation*. 2003;17(1):21-29.

- 31. Ware MA, Wang T, Shapiro S, Collet J-P, team Cs. Cannabis for the Management of Pain: Assessment of Safety Study (COMPASS). *The journal of pain : official journal of the American Pain Society*. 2015;16(12):1233-1242.
- 32. Storr M, Devlin S, Kaplan GG, Panaccione R, Andrews CN. Cannabis use provides symptom relief in patients with inflammatory bowel disease but is associated with worse disease prognosis in patients with Crohn's disease. *Inflammatory bowel diseases*. 2014;20(3):472-480.
- 33. Notcutt W, Price M, Miller R, et al. Initial experiences with medicinal extracts of cannabis for chronic pain: results from 34 'N of 1' studies. *Anaesthesia*. 2004;59(5):440-452.
- 34. Fiz J, Duran M, Capella D, Carbonell J, Farre M. Cannabis use in patients with fibromyalgia: effect on symptoms relief and health-related quality of life. *PloS one*. 2011;6(4):e18440.
- 35. Abrams DI, Jay CA, Shade SB, et al. Cannabis in painful HIV-associated sensory neuropathy: a randomized placebo-controlled trial. *Neurology*. 2007;68(7):515-521.
- 36. Berman JS, Symonds C, Birch R. Efficacy of two cannabis based medicinal extracts for relief of central neuropathic pain from brachial plexus avulsion: results of a randomised controlled trial. *Pain*. 2004;112(3):299-306.
- 37. Ellis RJ, Toperoff W, Vaida F, et al. Smoked medicinal cannabis for neuropathic pain in HIV: a randomized, crossover clinical trial. *Neuropsychopharmacology : official publication of the American College of Neuropsychopharmacology*. 2009;34(3):672-680.
- 38. Lynch ME, Cesar-Rittenberg P, Hohmann AG. A double-blind, placebo-controlled, crossover pilot trial with extension using an oral mucosal cannabinoid extract for treatment of chemotherapy-induced neuropathic pain. *Journal of pain and symptom management*. 2014;47(1):166-173.
- 39. Nurmikko TJ, Serpell MG, Hoggart B, Toomey PJ, Morlion BJ, Haines D. Sativex successfully treats neuropathic pain characterised by allodynia: a randomised, double-blind, placebo-controlled clinical trial. *Pain.* 2007;133(1-3):210-220.
- 40. Selvarajah D, Gandhi R, Emery CJ, Tesfaye S. Randomized placebo-controlled doubleblind clinical trial of cannabis-based medicinal product (Sativex) in painful diabetic neuropathy: depression is a major confounding factor. *Diabetes care*. 2010;33(1):128-130.
- 41. Serpell M, Ratcliffe S, Hovorka J, et al. A double-blind, randomized, placebo-controlled, parallel group study of THC/CBD spray in peripheral neuropathic pain treatment. *European journal of pain (London, England).* 2014;18(7):999-1012.
- 42. Wallace MS, Marcotte TD, Umlauf A, Gouaux B, Atkinson JH. Efficacy of Inhaled Cannabis on Painful Diabetic Neuropathy. *The journal of pain : official journal of the American Pain Society*. 2015;16(7):616-627.
- 43. Ware MA, Wang T, Shapiro S, et al. Smoked cannabis for chronic neuropathic pain: a randomized controlled trial. *CMAJ* : *Canadian Medical Association journal* = *journal de l'Association medicale canadienne*. 2010;182(14):E694-701.
- 44. Wilsey B, Marcotte T, Tsodikov A, et al. A randomized, placebo-controlled, crossover trial of cannabis cigarettes in neuropathic pain. *The journal of pain : official journal of the American Pain Society*. 2008;9(6):506-521.
- 45. Wilsey B, Marcotte T, Deutsch R, Gouaux B, Sakai S, Donaghe H. Low-dose vaporized cannabis significantly improves neuropathic pain. *The journal of pain : official journal of the American Pain Society*. 2013;14(2):136-148.



- 46. Wilsey B, Marcotte TD, Deutsch R, Zhao H, Prasad H, Phan A. An exploratory human laboratory experiment evaluating vaporized cannabis in the treatment of neuropathic pain from spinal cord injury and disease. *The Journal of Pain.* 2016;17(9):982-1000.
- 47. Collin C, Ehler E, Waberzinek G, et al. A double-blind, randomized, placebo-controlled, parallel-group study of Sativex, in subjects with symptoms of spasticity due to multiple sclerosis. *Neurol Res.* 2010;32(5):451-459.
- 48. Corey-Bloom J, Wolfson T, Gamst A, et al. Smoked cannabis for spasticity in multiple sclerosis: a randomized, placebo-controlled trial. *CMAJ* : *Canadian Medical Association journal* = *journal de l'Association medicale canadienne*. 2012;184(10):1143-1150.
- 49. Langford RM, Mares J, Novotna A, et al. A double-blind, randomized, placebocontrolled, parallel-group study of THC/CBD oromucosal spray in combination with the existing treatment regimen, in the relief of central neuropathic pain in patients with multiple sclerosis. *Journal of neurology*. 2013;260(4):984-997.
- 50. Rog DJ, Nurmikko TJ, Friede T, Young CA. Randomized, controlled trial of cannabisbased medicine in central pain in multiple sclerosis. *Neurology*. 2005;65(6):812-819.
- 51. van Amerongen G, Kanhai K, Baakman AC, et al. Effects on Spasticity and Neuropathic Pain of an Oral Formulation of δ9-Tetrahydrocannabinol in Patients With Progressive Multiple Sclerosis. *Clinical Therapeutics.* 2017.
- 52. Wade DT, Makela P, Robson P, House H, Bateman C. Do cannabis-based medicinal extracts have general or specific effects on symptoms in multiple sclerosis? A double-blind, randomized, placebo-controlled study on 160 patients. *Multiple sclerosis (Houndmills, Basingstoke, England).* 2004;10(4):434-441.
- 53. Zajicek J, Fox P, Sanders H, et al. Cannabinoids for treatment of spasticity and other symptoms related to multiple sclerosis (CAMS study): multicentre randomised placebo-controlled trial. *Lancet (London, England)*. 2003;362(9395):1517-1526.
- 54. Zajicek JP, Hobart JC, Slade A, Barnes D, Mattison PG, Group MR. Multiple sclerosis and extract of cannabis: results of the MUSEC trial. *Journal of neurology, neurosurgery, and psychiatry*. 2012;83(11):1125-1132.
- 55. Johnson JR, Burnell-Nugent M, Lossignol D, Ganae-Motan ED, Potts R, Fallon MT. Multicenter, double-blind, randomized, placebo-controlled, parallel-group study of the efficacy, safety, and tolerability of THC:CBD extract and THC extract in patients with intractable cancer-related pain. *Journal of pain and symptom management*. 2010;39(2):167-179.
- 56. Noyes R, Jr., Brunk SF, Avery DA, Canter AC. The analgesic properties of delta-9tetrahydrocannabinol and codeine. *Clinical pharmacology and therapeutics*. 1975;18(1):84-89.
- 57. Portenoy RK, Ganae-Motan ED, Allende S, et al. Nabiximols for opioid-treated cancer patients with poorly-controlled chronic pain: a randomized, placebo-controlled, graded-dose trial. *The journal of pain : official journal of the American Pain Society*. 2012;13(5):438-449.
- 58. De Vries M, Van Rijckevorsel DCM, Vissers KCP, Wilder-Smith OHG, Van Goor H. Tetrahydrocannabinol Does Not Reduce Pain in Patients With Chronic Abdominal Pain in a Phase 2 Placebo-controlled Study. *Clinical gastroenterology and hepatology*. 2016;(no pagination).
- 59. Blake DR, Robson P, Ho M, Jubb RW, McCabe CS. Preliminary assessment of the efficacy, tolerability and safety of a cannabis-based medicine (Sativex) in the treatment of pain caused by rheumatoid arthritis. *Rheumatology (Oxford, England)*. 2006;45(1):50-52.

- 60. Wilkinson ST, Stefanovics E, Rosenheck RA. Marijuana use is associated with worse outcomes in symptom severity and violent behavior in patients with posttraumatic stress disorder. *The Journal of clinical psychiatry*. 2015;76(9):1174-1180.
- 61. Johnson MJ, Pierce JD, Mavandadi S, et al. Mental health symptom severity in cannabis using and non-using Veterans with probable PTSD. *J Affect Disord*. 2016;190:439-442.
- 62. Cameron C, Watson D, Robinson J. Use of a synthetic cannabinoid in a correctional population for posttraumatic stress disorder-related insomnia and nightmares, chronic pain, harm reduction, and other indications: a retrospective evaluation. *Journal of clinical psychopharmacology*. 2014;34(5):559-564.
- 63. Fraser GA. The use of a synthetic cannabinoid in the management of treatment-resistant nightmares in posttraumatic stress disorder (PTSD). *CNS neuroscience & therapeutics*. 2009;15(1):84-88.
- 64. Jetly R, Heber A, Fraser G, Boisvert D. The efficacy of nabilone, a synthetic cannabinoid, in the treatment of PTSD-associated nightmares: A preliminary randomized, double-blind, placebo-controlled cross-over design study. *Psychoneuroendocrinology*. 2015;51(7612148, qgc):585-588.
- 65. Roitman P, Mechoulam R, Cooper-Kazaz R, Shalev AY. Preliminary, open-label, pilot study of add-on oral Δ [superscript]9-tetrahydrocannabinol in chronic post-traumatic stress disorder. *Clinical Drug Investigation*. 2014;34(8):587-591.
- 66. Mashiah M. Medical Cannabis as treatment for chronic combat PTSD: promising results in an open pilot tudy. Patients Out of Time Conference. Tucson, AZ. 2012.
- 67. Reznik I. Medical marijuana/cannabis use in patients with post-traumatic stress disorder. The International Conference on Integrative Medicine. Jerusalem, Israel. 2011.
- 68. Tetrault JM, Crothers K, Moore BA, Mehra R, Concato J, Fiellin DA. Effects of marijuana smoking on pulmonary function and respiratory complications: a systematic review. *Archives of internal medicine*. 2007;167(3):221-228.
- 69. Hancox RJ, Poulton R, Ely M, et al. Effects of cannabis on lung function: a populationbased cohort study. *The European respiratory journal*. 2010;35(1):42-47.
- 70. Pletcher MJ, Vittinghoff E, Kalhan R, et al. Association between marijuana exposure and pulmonary function over 20 years. *JAMA*. 2012;307(2):173-181.
- 71. Tashkin DP, Simmons MS, Sherrill DL, Coulson AH. Heavy habitual marijuana smoking does not cause an accelerated decline in FEV1 with age. *American journal of respiratory and critical care medicine*. 1997;155(1):141-148.
- 72. Frost L, Mostofsky E, Rosenbloom JI, Mukamal KJ, Mittleman MA. Marijuana use and long-term mortality among survivors of acute myocardial infarction. *American heart journal*. 2013;165(2):170-175.
- 73. Mittleman MA, Lewis RA, Maclure M, Sherwood JB, Muller JE. Triggering myocardial infarction by marijuana. *Circulation*. 2001;103(23):2805-2809.
- 74. Carvalho MFFd, Dourado MR, Fernandes IB, Araujo CTP, Mesquita AT, Ramos-Jorge ML. Head and neck cancer among marijuana users: a meta-analysis of matched case-control studies. *Archives of Oral Biology*. 2015;60(12):1750-1755.
- 75. Zhang LR, Morgenstern H, Greenland S, et al. Cannabis smoking and lung cancer risk: Pooled analysis in the International Lung Cancer Consortium. *International journal of cancer Journal international du cancer*. 2015;136(4):894-903.
- 76. Callaghan RC, Allebeck P, Sidorchuk A. Marijuana use and risk of lung cancer: a 40year cohort study. *Cancer causes & control : CCC*. 2013;24(10):1811-1820.
- 77. Gurney J, Shaw C, Stanley J, Signal V, Sarfati D. Cannabis exposure and risk of testicular cancer: a systematic review and meta-analysis. *BMC Cancer*. 2015;15:897.



- 78. Chacko JA, Heiner JG, Siu W, Macy M, Terris MK. Association between marijuana use and transitional cell carcinoma. *Urology*. 2006;67(1):100-104.
- 79. Rogeberg O, Elvik R. The effects of cannabis intoxication on motor vehicle collision revisited and revised. *Addiction*. 2016.
- 80. Ronen A, Gershon P, Drobiner H, et al. Effects of THC on driving performance, physiological state and subjective feelings relative to alcohol. *Accident; analysis and prevention.* 2008;40(3):926-934.
- 81. Lenne MG, Dietze PM, Triggs TJ, Walmsley S, Murphy B, Redman JR. The effects of cannabis and alcohol on simulated arterial driving: Influences of driving experience and task demand. *Accident; analysis and prevention.* 2010;42(3):859-866.
- 82. Grotenhermen F, Leson G, Berghaus G, et al. Developing limits for driving under cannabis. *Addiction (Abingdon, England)*. 2007;102(12):1910-1917.
- 83. Borges G, Bagge CL, Orozco R. A literature review and meta-analyses of cannabis use and suicidality. *Journal of Affective Disorders*. 2016;195:63-74.
- 84. Moore THM, Zammit S, Lingford-Hughes A, et al. Cannabis use and risk of psychotic or affective mental health outcomes: a systematic review. *Lancet (London, England)*. 2007;370(9584):319-328.
- 85. Gibbs M, Winsper C, Marwaha S, Gilbert E, Broome M, Singh SP. Cannabis use and mania symptoms: A systematic review and meta-analysis. *European Psychiatry*. 2015;30((Gibbs M.; Winsper C.; Marwaha S.; Gilbert E.; Singh S.P.) Warwick Medical School, University of Warwick, Coventry, United Kingdom):1128.
- 86. Kuepper R, van Os J, Lieb R, Wittchen H-U, Hofler M, Henquet C. Continued cannabis use and risk of incidence and persistence of psychotic symptoms: 10 year follow-up cohort study. *BMJ (Clinical research ed).* 2011;342:d738.
- 87. Dominguez M, Saka MC, Lieb R, Wittchen HU, van Os J. Early expression of negative/disorganized symptoms predicting psychotic experiences and subsequent clinical psychosis: a 10-year study. *The American journal of psychiatry*. 2010;167(9):1075-1082.
- 88. Rossler W, Hengartner MP, Angst J, Ajdacic-Gross V. Linking substance use with symptoms of subclinical psychosis in a community cohort over 30 years. *Addiction (Abingdon, England).* 2012;107(6):1174-1184.
- 89. Kaufmann RM, Kraft B, Frey R, et al. Acute psychotropic effects of oral cannabis extract with a defined content of Delta9-tetrahydrocannabinol (THC) in healthy volunteers. *Pharmacopsychiatry*. 2010;43(1):24-32.
- 90. Mason OJ, Morgan CJM, Stefanovic A, Curran HV. The psychotomimetic states inventory (PSI): measuring psychotic-type experiences from ketamine and cannabis. *Schizophrenia research*. 2008;103(1-3):138-142.
- 91. Englund A, Morrison PD, Nottage J, et al. Cannabidiol inhibits THC-elicited paranoid symptoms and hippocampal-dependent memory impairment. *Journal of psychopharmacology (Oxford, England).* 2013;27(1):19-27.
- 92. Di Forti M, Morgan C, Dazzan P, et al. High-potency cannabis and the risk of psychosis. *The British journal of psychiatry : the journal of mental science*. 2009;195(6):488-491.
- 93. Schreiner AM, Dunn ME. Residual effects of cannabis use on neurocognitive performance after prolonged abstinence: a meta-analysis. *Experimental and clinical psychopharmacology*. 2012;20(5):420-429.
- 94. American Psychiatric Association. *Diagnostic and Statistical Manual of Mental Disorders--Fifth Edition (DSM-5)*. Washington, DC2013.
- 95. World Health Organization. *International Classification of Diseases--Tenth Revision* (*ICD-10*). Geneva, Switzerland1999.



- 96. Bonn-Miller MO, Harris AHS, Trafton JA. Prevalence of cannabis use disorder diagnoses among veterans in 2002, 2008, and 2009. *Psychological services*. 2012;9(4):404-416.
- 97. Blanco C, Hasin DS, Wall MM, et al. Cannabis Use and Risk of Psychiatric Disorders: Prospective Evidence From a US National Longitudinal Study. *JAMA psychiatry*. 2016;73(4):388-395.
- 98. Fleming MF, Balousek SL, Klessig CL, Mundt MP, Brown DD. Substance use disorders in a primary care sample receiving daily opioid therapy. *The journal of pain : official journal of the American Pain Society*. 2007;8(7):573-582.
- 99. Hefner K, Sofuoglu M, Rosenheck R. Concomitant cannabis abuse/dependence in patients treated with opioids for non-cancer pain. *The American journal on addictions / American Academy of Psychiatrists in Alcoholism and Addictions*. 2015;24(6):538-545.
- 100. Bonn-Miller MO, Moos RH, Boden MT, Long WR, Kimerling R, Trafton JA. The impact of posttraumatic stress disorder on cannabis quit success. *The American journal of drug and alcohol abuse*. 2015;41(4):339-344.
- 101. Walsh K, Elliott JC, Shmulewitz D, et al. Trauma exposure, posttraumatic stress disorder and risk for alcohol, nicotine, and marijuana dependence in Israel. *Comprehensive psychiatry*. 2014;55(3):621-630.
- 102. Kevorkian S, Bonn-Miller MO, Belendiuk K, Carney DM, Roberson-Nay R, Berenz EC. Associations among trauma, posttraumatic stress disorder, cannabis use, and cannabis use disorder in a nationally representative epidemiologic sample. *Psychology of addictive behaviors : journal of the Society of Psychologists in Addictive Behaviors*. 2015;29(3):633-638.
- 103. Bonn-Miller MO, Boden MT, Vujanovic AA, Drescher KD. Prospective investigation of the impact of cannabis use disorders on posttraumatic stress disorder symptoms among veterans in residential treatment. *Psychological Trauma: Theory, Research, Practice, and Policy.* 2013;5(2):193-200.
- 104. Cescon DW, Page AV, Richardson S, Moore MJ, Boerner S, Gold WL. Invasive pulmonary aspergillosis associated with marijuana use in a man with colorectal cancer. *J Clin Oncol.* 2008;26(13):2214-2215.
- Chusid MJ, Gelfand JA, Nutter C, Fauci AS. Letter: Pulmonary aspergillosis, inhalation of contaminated marijuana smoke, chronic granulomatous disease. *Ann Intern Med.* 1975;82(5):682-683.
- 106. Marks WH, Florence L, Lieberman J, et al. Successfully treated invasive pulmonary aspergillosis associated with smoking marijuana in a renal transplant recipient. *Transplantation*. 1996;61(12):1771-1774.
- 107. Kagen SL, Kurup VP, Sohnle PG, Fink JN. Marijuana smoking and fungal sensitization. *The Journal of allergy and clinical immunology*. 1983;71(4):389-393.
- Munckhof WJ, Konstantinos A, Wamsley M, Mortlock M, Gilpin C. A cluster of tuberculosis associated with use of a marijuana water pipe. *Int J Tuberc Lung Dis.* 2003;7(9):860-865.
- 109. Oeltmann JE, Oren E, Haddad MB, et al. Tuberculosis outbreak in marijuana users, Seattle, Washington, 2004. *Emerging infectious diseases*. 2006;12(7):1156-1159.
- 110. Ramos S, Rodrigues R, Almeida N, Sá JM, Fonseca L. Cannabinoid hyperemesis syndrome. *Psychotherapy and Psychosomatics*. 2013;82((Ramos S.; Rodrigues R.; Almeida N.; Fonseca L.) Department of Psychiatry, Centro Hospitalar Alto Ave, Guimarães, Portugal):90.
- 111. Sadiq M. Cannabis hyperemesis syndrome. Journal of Addiction Medicine. 2013;7(4):E3.
- 112. Soriano-Co M, Batke M, Cappell MS. The cannabis hyperemesis syndrome characterized by persistent nausea and vomiting, abdominal pain, and compulsive bathing associated



with chronic marijuana use: a report of eight cases in the United States. *Digestive diseases and sciences*. 2010;55(11):3113-3119.

- 113. Velasco A, Pentecost P. An unexpected etiology of cyclical vomiting. *Journal of Hospital Medicine*. 2012;7((Velasco A.; Pentecost P.) University of New Mexico, School of Medicine, Albuquerque, United States):S281.
- 114. Vujasinović M, Ivartnik M, Tretjak M. Cannabinoid hyperemesis syndrome Case report. *Zdravniski Vestnik.* 2012;81(2):159-162.
- 115. Welder JD. Some like it hot: A case of cannabinoid hyperemesis syndrome. *Journal of General Internal Medicine*. 2012;27((Welder J.D.) University of Iowa, Iowa City, United States):S480-S481.
- 116. Woods JA, Wright NJ, Gee J, Scobey MW. Cannabinoid Hyperemesis Syndrome: An Emerging Drug-Induced Disease. *Am J Ther.* 2016;23(2):e601-605.
- 117. Simonetto DA, Oxentenko AS, Herman ML, Szostek JH. Cannabinoid hyperemesis: a case series of 98 patients. *Mayo Clinic proceedings*. 2012;87(2):114-119.
- 118. Brandenburg D, Wernick R. Intravenous marijuana syndrome. *West J Med.* 1986;145(1):94-96.
- 119. Carabellese F, Candelli C, Martinelli D, La Tegola D, Catanesi R. Cannabis use and violent behaviour: a psychiatric patients cohort study in Southern Italy. *Rivista di psichiatria*. 2013;48(1):43-50.
- 120. Myerscough R, Taylor S. The effects of marijuana on human physical aggression. *Journal of personality and social psychology*. 1985;49(6):1541-1546.
- 121. Loflin M, Earleywine M. A new method of cannabis ingestion: the dangers of dabs? *Addictive behaviors*. 2014;39(10):1430-1433.
- 122. Lamy FR, Daniulaityte R, Sheth A, et al. "Those edibles hit hard": Exploration of Twitter data on cannabis edibles in the U.S. *Drug Alcohol Depend.* 2016;164:64-70.
- 123. Hudak M, Severn D, Nordstrom K. Edible Cannabis-Induced Psychosis: Intoxication and Beyond. *Am J Psychiatry*. 2015;172(9):911-912.
- 124. Meier MH, Caspi A, Cerda M, et al. Associations Between Cannabis Use and Physical Health Problems in Early Midlife: A Longitudinal Comparison of Persistent Cannabis vs Tobacco Users. *JAMA Psychiatry*. 2016;73(7):731-740.
- Vandrey R, Raber JC, Raber ME, Douglass B, Miller C, Bonn-Miller MO. Cannabinoid Dose and Label Accuracy in Edible Medical Cannabis Products. *Jama*. 2015;313(24):2491-2493.
- 126. Mehra R, Moore BA, Crothers K, Tetrault J, Fiellin DA. The association between marijuana smoking and lung cancer: a systematic review. *Archives of internal medicine*. 2006;166(13):1359-1367.
- Leweke FM, Piomelli D, Pahlisch F, et al. Cannabidiol enhances anandamide signaling and alleviates psychotic symptoms of schizophrenia. *Translational psychiatry*. 2012;2:e94.
- 128. Schubart CD, Sommer IE, Fusar-Poli P, de Witte L, Kahn RS, Boks MP. Cannabidiol as a potential treatment for psychosis. *Eur Neuropsychopharmacol.* 2014;24(1):51-64.
- 129. Management of Post-Traumatic Stress Working Group. VA/DoD Clinical Practice Guidelines for Management of Post-Traumatic Stress, Version 2.0. Department of Veteran Affairs and Department of Defense. 2010; <u>http://www.healthquality.va.gov/guidelines/MH/ptsd/cpg_PTSD-FULL-201011612.pdf</u>. Accessed November 4, 2016.
- 130. Treatment for Posttraumatic Stress Disorder in Military and Veteran Populations, Final Assessment. Washington, DC: Institute of Medicine; http://iom.nationalacademies.org/Reports/2014/Treatment-for-Posttraumatic-Stress-



Disorder-in-Military-and-Veteran-Populations-Final-Assessment.aspx, Accessed November 4, 2016. 2014.

- 131. Dowell D, Haegerich TM, Chou R. CDC Guideline for Prescribing Opioids for Chronic Pain United States, 2016. *MMWR Recommendations and reports : Morbidity and mortality weekly report Recommendations and reports.* 2016;65(1):1-49.
- 132. Abrams DI, Couey P, Shade SB, Kelly ME, Benowitz NL. Cannabinoid-opioid interaction in chronic pain. *Clinical pharmacology and therapeutics*. 2011;90(6):844-851.
- 133. DeGeorge M, Dawson E, Woster P, Ko M, Burke L, Bronstein K. An analysis of the association between marijuana use and potential nonadherence in patients prescribed hydrocodone. Poster session presented at the 2013 annual meeting of the American Academy of Pain Medicine. Fort Lauderdale, FL: 2013, April. Retrieved from http://www.painmed.org/2013posters/poster114.pdf. Accessed August 25, 2016.
- Ashrafioun L, Bohnert KM, Jannausch M, Ilgen MA. Characteristics of substance use disorder treatment patients using medical cannabis for pain. *Addictive behaviors*. 2015;42(2gw, 7603486):185-188.
- Haroutounian S, Ratz Y, Ginosar Y, et al. The Effect of Medicinal Cannabis on Pain and Quality of Life Outcomes in Chronic Pain: A Prospective Open-label Study. *Clin J Pain*. 2016.
- 136. Saint Louis C, Apuzzoaug M. Obama Administration Set to Remove Barrier to Marijuana Research. The New York Times. August 10, 2016. <u>http://www.nytimes.com/2016/08/11/science/obama-administration-set-to-remove-barrier-to-marijuana-research.html?emc=eta1&_r=0</u> Accessed August 25, 2016.
- 137. van Nierop M, Janssens M, Genetic Risk OoPI, et al. Evidence that transition from health to psychotic disorder can be traced to semi-ubiquitous environmental effects operating against background genetic risk. *PloS one*. 2013;8(11):e76690.

APPENDIX A. SEARCH STRATEGIES

Databases/Websites

- Ovid Medline
- PubMed (non-Medline materials)
- Elsevier EMBASE
- Ovid PsycINFO
- PILOTS Database (PTSD search only)
- EBM Reviews (CDSR, DARE, HTA, Cochrane CENTRAL, etc)
- Conference Papers Index
- Clinicaltrials.gov
- International Clinical Trials Registry Platform (WHO ICTRP)
- · ISRCTN
- NIH Reporter
- AHRQ Gold
- American Cancer Society Database of Studies

Search Strategies

Ovid MEDLINE(R) and Ovid OLDMEDLINE(R) 1946 to December Week 5 2015,

Ovid MEDLINE(R) In-Process & Other Non-Indexed Citations January 11, 2016

Date Searched: Tuesday January 12, 2016

#	Searches	Results
1	medical marijuana/ or cannabis/ or marijuana smoking/ or exp Cannabinoids/ or Cannabaceae/	18682
2	(cannabis or canabis or cannabinoid* or cannabidiol* or CBD or cannabacae or marijuana or marihuana or hashish or hash or ganja or ganjah or hemp or bhang or charas or THC or tetrahydrocannabinol* or tetra-hydrocannabinol* or 9?tetrahydrocannabinol* or DELTA?9?-tetrahydrocannabinol*).tw.	38570
3	1 or 2	41269
4	pain/ or acute pain/ or breakthrough pain/ or mastodynia/ or exp musculoskeletal pain/ or exp back pain/ or chronic pain/ or facial pain/ or headache/ or metatarsalgia/ or neck pain/ or exp neuralgia/ or exp nociceptive pain/ or pain, intractable/ or pain, referred/	205083
5	(pain or pains or painful* or migraine* or headache* or neuropath* or neuralgia* or arthriti* or fibromyalg*).tw.	770253
6	4 or 5	823437
7	3 and 6	2868
8	7 and (humans/ not animals/)	1331
9	7 not (humans/ or animals/)	312
10	8 or 9	1643
11	limit 10 to (case reports or comment or editorial or letter or news)	293



12	cross-section*.tw.	243912
_	10 not (11 or 12)	1313
	limit 13 to english language	1211
15	stress disorders, traumatic/ or combat disorders/ or stress disorders, post-traumatic/	26019
16	(PTSD or post-traumatic stress or posttraumatic stress).ti,ab.	23732
17	15 or 16	32767
18	3 and 17	210
19	18 and (humans/ not animals/)	131
20	18 not (humans/ or animals/)	31
21	19 or 20	162
22	limit 21 to (case reports or comment or editorial or letter or news)	9
23	cross-section*.tw.	243912
24	21 not (22 or 23)	140
25	limit 24 to english language	132
26	medical marijuana/ or cannabis/ or marijuana smoking/ or marijuana abuse/ or exp Cannabinoids/ or Cannabaceae/	22185
27	(cannabis* or canabis* or cannabinoid* or cannabidiol* or CBD or cannabacae or marijuana* or marihuana* or hashish* or hash or ganja or ganjah or hemp or bhang or charas or THC or tetrahydrocannabinol* or tetra-hydrocannabinol* or 9?tetrahydrocannabinol* or DELTA?9?-tetrahydrocannabinol*).tw.	38598
28	26 or 27	41948
29	(ae or co or de).fs.	5311331
30	(harm or harms or harmful or safe or safety or side effect* or undesirable effect* or treatment emergent or tolerability or toxic* or adrs or damag* or impair* or disorder* or abuse* or addict* or withdrawal* or (adverse adj2 (effect or effects or reaction or reactions or event or events or outcome or outcomes))).tw.	3065069
31	29 or 30	7263273
32	28 and 31	25510
33	limit 32 to (meta analysis or systematic reviews)	422
34	32 not 33	25088
35	34 and (humans/ not animals/)	13847
_	34 not (humans/ or animals/)	1758
37	35 or 36	15605
38	limit 37 to ("all adult (19 plus years)" or "young adult (19 to 24 years)" or "adult (19 to 44 years)" or "young adult and adult (19-24 and 19-44)" or "middle age (45 to 64 years)" or "middle aged (45 plus years)" or "all aged (65 and over)" or "aged (80 and over)")	8086
39	limit 38 to (case reports or comment or editorial or letter or news)	1030
40	cross-section*.tw.	243912
41	38 not (39 or 40)	6701
12	limit 41 to english language	6238

PubMed

Date searched: Friday May 6, 2016

#	Searches	Results
#10	Search (#7 OR #8 OR #9)	444
#9	Search ((((((cannabis*[tiab] OR canabis*[tiab] OR cannabinoid*[tiab] OR cannabidiol*[tiab] OR CBD[tiab] OR cannabacae[tiab] OR marijuana*[tiab] OR marihuana*[tiab] OR hashish*[tiab] OR hash[tiab] OR ganja[tiab] OR ganjah[tiab] OR hemp[tiab] OR bhang[tiab] OR charas[tiab] OR THC[tiab] OR tetrahydrocannabinol*[tiab] OR tetra-hydrocannabinol*[tiab] OR 9?tetrahydrocannabinol*[tiab] OR DELTA?9?-tetrahydrocannabinol*[tiab])))) AND (((harm[tiab] OR harms[tiab] OR harmful[tiab] OR safe[tiab] OR safety[tiab] OR side effect*[tiab] OR undesirable effect*[tiab] OR treatment emergent[tiab] OR disorder*[tiab] OR toxic*[tiab] OR advs[tiab] OR damag*[tiab] OR impair*[tiab] OR disorder*[tiab] OR adverse effects[tiab] OR adverse reaction[tiab] OR adverse effect[tiab] OR adverse effects[tiab] OR adverse reaction[tiab] OR adverse reactions[tiab] OR adverse outcomes[tiab] OR horizon scan*[tiab] OR adverse outcome[tiab] OR adverse outcomes[tiab] OR horizon scan*[tiab] OR systematic* review*[tiab] OR systematic effectiveness review*[tiab] OR comparative effectiveness review*[tiab] OR adverse review*[tiab] OR landscape review*[tiab] OR quantitative review*[tiab] OR qualitative review*or integrative review*or mixed-method* review*or mixed method* review*[tiab] OR updat* review*[tiab] OR cochrane review*or mixed method* review*[tiab] OR updat* review*[tiab] OR evidence aggregat*[tiab] OR evidence map*[tiab] OR evidence brief*[tiab] OR evidence summar*[tiab] OR rapid review*or mini* review*or pragmatic review*or targeted review*or focused review*or brief review*or short review*[tiab] OR (meta-analy*[tiab] OR metaanaly*[tiab] OR meta- meta-analy*[tiab] OR evidence synthes*[tiab] OR knowledge synthes*[tiab] OR quantitative synthes*[tiab] OR mixe4* comparison*[tiab] OR metaanaly*[tiab] OR meta-analy*[tiab] OR evidence synthes*[tiab] OR netaanaly*[tiab] OR meta-analy*[tiab] OR evidence synthes*[tiab] OR knowledge synthes*[tiab] OR meta-analy*[tiab] OR mixe4* comparison*[tiab] OR health technology assessment*[tiab] OR mini-HTA*[tiab] OR relat	16
#8	Search (((((cannabis*[tiab] OR canabis*[tiab] OR cannabinoid*[tiab] OR cannabidiol*[tiab] OR CBD[tiab] OR cannabacae[tiab] OR marijuana*[tiab] OR marihuana*[tiab] OR hashish*[tiab] OR hash[tiab] OR ganja[tiab] OR ganjah[tiab] OR hemp[tiab] OR bhang[tiab] OR charas[tiab] OR THC[tiab] OR tetrahydrocannabinol*[tiab] OR tetra-hydrocannabinol*[tiab] OR 9?tetrahydrocannabinol*[tiab] OR DELTA?9?-tetrahydrocannabinol*[tiab]))) AND (((PTSD[tiab] OR post-traumatic stress[tiab] OR posttraumatic stress[tiab]))) AND (((pubmednotmedline[sb] OR inprocess[sb] OR [publisher[sb]))))	39
#7	Search (((((cannabis*[tiab] OR canabis*[tiab] OR cannabinoid*[tiab] OR cannabidiol*[tiab] OR CBD[tiab] OR cannabacae[tiab] OR marijuana*[tiab] OR marihuana*[tiab] OR hashish*[tiab] OR hash[tiab] OR ganja[tiab] OR ganjah[tiab] OR hemp[tiab] OR bhang[tiab] OR charas[tiab] OR THC[tiab] OR tetrahydrocannabinol*[tiab] OR tetra-hydrocannabinol*[tiab] OR 9?tetrahydrocannabinol*[tiab] OR DELTA?9?-tetrahydrocannabinol*[tiab])))) AND (((pain[tiab] OR pains[tiab] OR painful*[tiab] OR migraine*[tiab] OR headache*[tiab] OR neuropath*[tiab] OR neuralgia*[tiab] OR arthriti*[tiab] OR fibromyalg*[tiab])))) AND (((pubmednotmedline[sb] OR inprocess[sb] OR [publisher[sb]))))	392
#6	Search ((meta-review*[tiab] OR meta-epidemiolog*[tiab] OR metaepidemiolog*[tiab] OR horizon scan*[tiab] OR systematic* review*[tiab] OR systematic effectiveness review*[tiab] OR comparative effectiveness review*[tiab] OR evidence review*[tiab] OR	78086

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	landscape review*[tiab] OR quantitative review*[tiab] OR qualitative review*or integrative review*or mixed-method* review*or mixed method* review*[tiab] OR research review*or scoping review*[tiab] OR umbrella review*or review of review*[tiab] OR updat* review*[tiab] OR cochrane review*or campbell review*[tiab])) OR (research* aggregat*[tiab] OR evidence aggregat*[tiab] OR evidence map*[tiab] OR evidence brief*[tiab] OR evidence summar*[tiab] OR rapid review*or mini* review*or pragmatic review*or targeted review*or focused review*or brief review*or short review*[tiab]) OR (meta-analy*[tiab] OR metaanaly*[tiab] OR meta-meta-analy*[tiab] OR evidence synthes*[tiab] OR knowledge synthes*[tiab] OR quantitative synthes*[tiab] OR research synthes*[tiab] OR pooled analy*[tiab] OR indirect* comparison*[tiab] OR mixed* comparison*[tiab]) OR (HTA[tiab] OR health technology assessment*[tiab] OR mini- HTA*[tiab] OR relative effectiveness assessment*[tiab]))	
#5	Search (pubmednotmedline[sb] OR inprocess[sb] OR publisher[sb])	2833028
#4	Search (harm[tiab] OR harms[tiab] OR harmful[tiab] OR safe[tiab] OR safety[tiab] OR side effect*[tiab] OR undesirable effect*[tiab] OR treatment emergent[tiab] OR tolerability[tiab] OR toxic*[tiab] OR adrs[tiab] OR damag*[tiab] OR impair*[tiab] OR disorder*[tiab] OR abuse*[tiab] OR addict*[tiab] OR withdrawal*[tiab] OR adverse effect[tiab] OR adverse effects[tiab] OR adverse reaction[tiab] OR adverse reactions[tiab] OR adverse event[tiab] OR adverse events[tiab] OR adverse outcome[tiab] OR adverse outcomes[tiab])	3137250
#3	Search (PTSD[tiab] OR post-traumatic stress[tiab] OR posttraumatic stress[tiab])	24584
#2	Search (pain[tiab] OR pains[tiab] OR painful*[tiab] OR migraine*[tiab] OR headache*[tiab] OR neuropath*[tiab] OR neuralgia*[tiab] OR arthriti*[tiab] OR fibromyalg*[tiab])	788713
#1	Search (cannabis*[tiab] OR canabis*[tiab] OR cannabinoid*[tiab] OR cannabidiol*[tiab] OR CBD[tiab] OR cannabacae[tiab] OR marijuana*[tiab] OR marihuana*[tiab] OR hashish*[tiab] OR hash[tiab] OR ganja[tiab] OR ganjah[tiab] OR hemp[tiab] OR bhang[tiab] OR charas[tiab] OR THC[tiab] OR tetrahydrocannabinol*[tiab] OR tetra- hydrocannabinol*[tiab] OR 9?tetrahydrocannabinol*[tiab] OR DELTA?9?- tetrahydrocannabinol*[tiab])	39258

EMBASE.COM

Date Searched: Tuesday May 10, 2016

#	Searches	Results
#1	'medical cannabis'/mj OR 'cannabis'/mj OR 'cannabis smoking'/mj OR 'cannabinoid'/exp/mj OR 'cannabaceae'/mj	28,447
#2	cannabis:ab,ti OR canabis:ab,ti OR cannabinoid*:ab,ti OR cannabidiol*:ab,ti OR cbd:ab,ti OR cannabacae:ab,ti OR marijuana:ab,ti OR marihuana:ab,ti OR hashish:ab,ti OR hash:ab,ti OR ganja:ab,ti OR ganjah:ab,ti OR hemp:ab,ti OR bhang:ab,ti OR charas:ab,ti OR thc:ab,ti OR tetrahydrocannabinol*:ab,ti OR 'tetra hydrocannabinol*:ab,ti OR '9 tetrahydrocannabinol*:ab,ti OR 9tetrahydrocannabinol*:ab,ti OR 'delta*9*tetrahydrocannabinol 11carboxylic acid':ab,ti	52,180
#3	#1 OR #2	57,164
#4	'pain'/mj OR 'breakthrough pain'/mj OR 'mastalgia'/mj OR 'musculoskeletal pain'/mj OR 'low back pain'/mj OR 'backache'/exp/mj OR 'chronic pain'/mj OR 'face pain'/mj OR 'headache and facial pain'/exp/mj OR 'metatarsalgia'/mj OR 'neck pain'/mj OR 'neuralgia'/exp/mj OR 'nociceptive pain'/mj OR 'intractable pain'/mj OR 'referred pain'/mj	243,955
#5	pain:ab,ti OR pains:ab,ti OR painful*:ab,ti OR migraine*:ab,ti OR headache*:ab,ti OR neuropath*:ab,ti OR neuralgia*:ab,ti OR arthriti*:ab,ti OR fibromyalg*:ab,ti	1,079,039
#6	#4 OR #5	1,130,556
#7	#3 AND #6	4,553



#8	#7 AND 'human'/de NOT 'nonhuman'/de	2,655
#9	#8 AND ('editorial'/it OR 'letter'/it OR 'note'/it)	80
#10	'cross-section*':ab,ti	297,421
#11	#8 NOT (#9 OR #10)	2,516
#12	#8 NOT (#9 OR #10) AND [english]/lim	2,308
#13	#8 NOT (#9 OR #10) AND [english]/lim AND [embase]/lim	2,088
#14	'posttraumatic stress disorder'/mj	23,335
#15	ptsd:ab,ti OR 'post-traumatic stress':ab,ti OR 'posttraumatic stress':ab,ti	29,813
#16	#14 OR #15	34,693
#17	#3 AND #16	314
#18	#17 AND 'human'/de NOT 'nonhuman'/de	227
#19	#18 AND 'editorial'/it	1
#20	'cross-section*':ab,ti	297,421
#21	#17 NOT (#19 OR #20)	295
#22	#17 NOT (#19 OR #20) AND [english]/lim	286
#23	#17 NOT (#19 OR #20) AND [english]/lim AND [embase]/lim	267
#24	'medical cannabis'/mj OR 'cannabis'/mj OR 'cannabis smoking'/mj OR	30,213
// Z -T	'cannabinoid'/exp/mj OR 'cannabaceae'/mj OR 'cannabis addiction'/mj	00,210
#25	#2 OR #24	57,324
#26	#25 AND ('adverse drug reaction'/Ink OR 'complication'/Ink OR 'drug interaction'/Ink OR 'drug toxicity'/Ink OR 'side effect'/Ink)	7,995
#27	harm:ab,ti OR harms:ab,ti OR harmful:ab,ti OR safe:ab,ti OR safety:ab,ti OR 'side effect*':ab,ti OR 'undesirable effect*':ab,ti OR 'treatment emergent':ab,ti OR tolerability:ab,ti OR toxic*:ab,ti OR adrs:ab,ti OR damag*:ab,ti OR impair*:ab,ti OR disorder*:ab,ti OR abuse*:ab,ti OR addict*:ab,ti OR withdrawal*:ab,ti OR 'adverse effect':ab,ti OR 'adverse effects':ab,ti OR 'adverse reaction':ab,ti OR 'adverse reactions':ab,ti OR 'adverse event':ab,ti OR 'adverse events':ab,ti OR 'adverse outcome':ab,ti OR 'adverse outcomes':ab,ti	4,055,060
#28	#26 OR #27	4,059,085
#29	#25 AND #28	25,939
#30	#25 AND #28 AND ([cochrane review]/lim OR [systematic review]/lim OR [meta analysis]/lim)	373
#31	#25 AND #28 AND ([cochrane review]/lim OR [systematic review]/lim OR [meta analysis]/lim) AND [embase]/lim	335
#32	#25 AND #28 ([cochrane review]/lim OR [systematic review]/lim OR [meta analysis]/lim) AND [embase]/lim AND [english]/lim	319
#33	#14 OR #24 OR #33	2,616
#34	#33 NOT [medline]/lim	1,592

PSYCINFO 1806 to May Week 1 2016

Date Searched: Tuesday May 10, 2016

#	Searches	Results
1	cannabis/ or hashish/ or marijuana/ or exp cannabinoids/ or tetrahydrocannabinol/ or cannabinoids/ or tetrahydrocannabinol/ or marijuana usage/ or marijuana/	11208
2	(cannabis or canabis or cannabinoid* or cannabidiol* or CBD or cannabacae or marijuana or marihuana or hashish or hash or ganja or ganjah or hemp or bhang or charas or THC or tetrahydrocannabinol* or tetra-hydrocannabinol* or 9?tetrahydrocannabinol* or DELTA?9?-tetrahydrocannabinol*).tw.	19269
3	1 or 2	19585
4	pain/ or aphagia/ or back pain/ or chronic pain/ or headache/ or myofascial pain/ or neuralgia/ or neuropathic pain/ or somatoform pain disorder/ or headache/ or migraine headache/ or muscle contraction headache/ or neuralgia/ or trigeminal neuralgia/ or pain management/	50895



44

5	(pain or pains or painful* or migraine* or headache* or neuropath* or neuralgia* or arthriti* or fibromyalg*).tw.	116341
6	4 or 5	117164
7	3 and 6	915
8	limit 7 to human	599
9	limit 7 to animal	346
10	7 not (8 or 9)	35
11	8 or 10	634
12	limit 11 to english language	582
13	limit 12 to ("column/opinion" or "comment/reply" or editorial or "erratum/correction" or letter)	54
14	12 not 13	528
15	cross-section*.tw.	54490
16	14 not 15	505
17	posttraumatic stress disorder/ or complex ptsd/ or desnos/	25127
18	(PTSD or post-traumatic stress or posttraumatic stress).tw.	33843
19	17 or 18	35163
20	3 and 19	209
21	limit 20 to human	178
22	limit 20 to animal	33
23	20 not (21 or 22)	12
24	21 or 23	190
25	limit 24 to english language	173
26	limit 25 to ("column/opinion" or "comment/reply" or editorial or letter)	9
27	25 not 26	164
28	cross-section*.tw.	54490
29	27 not 28	155
30	(harm or harms or harmful or safe or safety or side effect* or undesirable effect* or treatment emergent or tolerability or toxic* or adrs or damag* or impair* or disorder* or abuse* or addict* or withdrawal* or (adverse adj2 (effect or effects or reaction or reactions or event or events or outcome or outcomes))).tw.	816214
31	3 and 30	10099
32	limit 31 to ("0830 systematic review" or 1200 meta analysis)	111
33	16 or 29 or 32	750

EBM Reviews Databases

- · Cochrane Central Register of Controlled Trials April 2016,
- · Cochrane Database of Systematic Reviews 2005 to May 05, 2016,
- Database of Abstracts of Reviews of Effects 1st Quarter 2016,
- Health Technology Assessment 2nd Quarter 2016,
- NHS Economic Evaluation Database 1st Quarter 2016

Date Searched: Tuesday May 10, 2016

<u>#</u>	Searches	Results
1	(cannabis or canabis or cannabinoid* or cannabidiol* or CBD or cannabacae or marijuana or marihuana or hashish or hash or ganja or ganjah or hemp or bhang or charas or THC or tetrahydrocannabinol* or tetra-hydrocannabinol* or 9?tetrahydrocannabinol* or DELTA?9?-tetrahydrocannabinol*).tw.	2318
2	(pain or pains or painful* or migraine* or headache* or neuropath* or neuralgia* or arthriti* or fibromyalg*).tw.	100259
3	1 and 2	262
4	(PTSD or post-traumatic stress or posttraumatic stress).tw.	2401
5	1 and 4	20
6	(harm or harms or harmful or safe or safety or side effect* or undesirable effect* or treatment emergent or tolerability or toxic* or adrs or damag* or impair* or disorder* or abuse* or addict* or withdrawal* or (adverse adj2 (effect or effects or reaction or reactions or event or events or outcome or outcomes))).tw.	274193
7	1 and 6	1227
8	(meta-review* or meta-epidemiolog* or metaepidemiolog* or "horizon scan*" or ((systematic* or "systematic effectiveness" or "comparative effectiveness" or evidence or landscape or methodologic or methodological or quantitative or qualitative or integrative or mixed-method* or "mixed method*" or research or scoping or umbrella or "review* of" or updat* or cochrane or campbell) adj review*) or ((research* or evidence) adj2 aggregat*) or "evidence map*" or "evidence brief*" or "evidence summar*" or ((rapid or mini* or pragmatic or targeted or focused or brief or short*) adj2 (systematic or evidence or knowledge or review* or synthes*)) or meta-analy* or "metaanaly* or "meta-meta-analy*" or "evidence synthes*" or "knowledge synthes*" or "quantitative synthes*" or "qualitative synthes*" or "research synthes*" or "integrat* data analys*" or (integrative adj1 analys?s) or "pooled analy*" or (indirect* adj2 comparison*) or (mixed* adj2 comparison*) or ((reliability or validity) adj generali?ation*) or meta-aggregat* or metaaggregat* or meta-narrative* or metanarrative* or meta-review* or metareview* or meta-stud* or metastud* or meta-summar* or metasummar* or meta-synth* or "qualitative cross-case" or realist-synth* or "realist synth*" or "realist review*" or "thematic synth*" or "summary receiver operating characteristic*" or "comparative case study" or "comparative case studies").ti,ab.	41555
9	7 and 8	116
10	3 or 5 or 9	343
11	remove duplicates from 10	334
12	limit 11 to english language	308

PILOTS: Published International Literature On Traumatic Stress Database

(http://www.ptsd.va.gov/professional/pilots-database/)

Date Searched: Tuesday May 10, 2016

ab(cannabis OR canabis OR cannabinoid* OR cannabidiol* OR CBD OR cannabacae OR marijuana OR marihuana OR hashish OR hash OR ganja OR ganjah OR hemp OR bhang OR charas OR THC OR tetrahydrocannabinol* OR tetra-hydrocannabinol* OR 9?tetrahydrocannabinol* OR DELTA?9?tetrahydrocannabinol*) AND

(PTSD OR posttraumatic stress OR post-traumatic stress)



= 177 results

COS Conference Papers Index

Date Searched: Tuesday May 17, 2016

Set	Search	Results
S4	S1 or S2 or S3	711°
	(cannabis OR canabis OR cannabinoid* OR cannabidiol* OR CBD OR cannabacae OR marijuana OR marihuana OR hashish OR hash OR ganja OR ganjah OR hemp OR bhang OR charas OR THC OR tetrahydrocannabinol* OR tetra-hydrocannabinol* OR 9?tetrahydrocannabinol* OR DELTA?9?-tetrahydrocannabinol*) AND (harm or harms or harmful or safe or safety or side effect* or undesirable effect* or treatment emergent or tolerability or toxic* or adrs or damag* or impair* or disorder* or abuse* or addict* or withdrawal* or (adverse NEAR/2 (effect or effects or reaction or reactions or event or events or outcome or outcomes)))Limits applied	532°
	(cannabis OR canabis OR cannabinoid* OR cannabidiol* OR CBD OR cannabacae OR marijuana OR marihuana OR hashish OR hash OR ganja OR ganjah OR hemp OR bhang OR charas OR THC OR tetrahydrocannabinol* OR tetra-hydrocannabinol* OR 9?tetrahydrocannabinol* OR DELTA?9?-tetrahydrocannabinol*) AND (PTSD or post- traumatic stress or posttraumatic stress)Limits applied	4°
	(cannabis or canabis or cannabinoid* or cannabidiol* or CBD or cannabacae or marijuana or marihuana or hashish or hash or ganja or ganjah or hemp or bhang or charas or THC or tetrahydrocannabinol* or tetra-hydrocannabinol* or 9?tetrahydrocannabinol* or DELTA?9?- tetrahydrocannabinol*) AND (pain or pains or painful* or migraine* or headache* or neuropath* or neuralgia* or arthriti* or fibromyalg*)Limits applied	176°

ClinicalTrials.gov

Date Searched: Monday May 16, 2015

Chronic Pain Search

(pain OR pains OR painful* OR migraine* OR headache* OR neuropath* OR neuralgia* OR arthriti* OR fibromyalg*) [DISEASE] AND (cannabis OR canabis OR cannabinoid* OR cannabidiol* OR CBD OR cannabacae OR marijuana OR marihuana OR hashish OR hash OR ganja OR ganjah OR hemp OR bhang OR charas OR THC OR tetrahydrocannabinol* OR tetra-hydrocannabinol* OR 9tetrahydrocannabinol* OR Δ 9-THC) [TREATMENT] = 74 results

Post-traumatic Stress Disorder (PTSD) Search

(PTSD OR post-traumatic stress OR posttraumatic stress) [DISEASE] AND (cannabis OR canabis OR cannabinoid* OR cannabidiol* OR CBD OR cannabacae OR marijuana OR marihuana OR hashish OR hash OR ganja OR ganjah OR hemp OR bhang OR charas OR THC OR tetrahydrocannabinol* OR tetrahydrocannabinol*) [TREATMENT] = 6 results

Harms Search

(harm* OR safety OR "side effect*" OR "undesirable effect*" OR "treatment emergent" OR tolerability OR toxic* OR adrs OR damag* OR impair* OR abuse* OR addict* OR withdrawal* OR "adverse effect*" OR "adverse event*" OR "adverse outcome*") AND ((cannabis OR canabis OR cannabinoid* OR cannabidiol* OR CBD OR cannabacae OR marijuana OR marihuana OR hashish OR hash OR ganja OR ganjah OR hemp OR bhang OR charas OR THC OR tetrahydrocannabinol* OR tetra-hydrocannabinol* OR 9tetrahydrocannabinol*) AND NOT (sativex OR namisol OR POT-4 OR Levodopa OR Carbidopa)) [TREATMENT] = 65 results





WHO ICTRP Database

Date Searched: Wednesday May 18, 2016

*Due to the 256 character limit for searches, the following searches were edited to fit within the proscribed limits. All terms removed were searched separately and found to not change or add additional results compared to the searches below.

Chronic Pain Search

CONDITION = (pain or pains or painful* or migraine* or headache* or neuropath* or neuralgia* or arthriti* or fibromyalg*)

AND

INTERVENTION=(cannabis or canabis or cannabinoid* or cannabidiol* or CBD or cannabacae or marijuana or hashish or hash or ganja or ganjah or hemp or THC or tetrahydrocannabinol* or tetrahydrocannabinol* or 9-tetrahydrocannabinol* or DELTA-9-tetrahydrocannabinol*) = 45 results [24 results were from ClinicalTrials.gov, therefore only 21 results were downloaded]

Post-traumatic Stress Disorder (PTSD) Search

CONDITION = (PTSD or post-traumatic stress or posttraumatic stress) AND

INTERVENTION=(cannabis or canabis or cannabinoid* or cannabidiol* or CBD or cannabacae or marijuana or hashish or hash or ganja or ganjah or hemp or THC or tetrahydrocannabinol* or tetrahydrocannabinol* or DELTA-9-tetrahydrocannabinol*) = 4 results [all results were from ClinicalTrials.gov so no results were downloaded]

Harms Search

CONDITION = (harm* or safe or safety or side effect* or undesirable effect* or tolerability or toxic* or adrs or damag* or impair* or disorder* or abuse* or addict* or withdrawal* or adverse effect* or adverse reaction* or adverse event* or adverse outcome*)

AND

INTERVENTION=(cannabis or canabis or cannabinoid* or cannabidiol* or CBD or cannabacae or marijuana or hashish or hash or ganja or ganjah or hemp or THC or tetrahydrocannabinol* or tetrahydrocannabinol* or 9-tetrahydrocannabinol* or DELTA-9-tetrahydrocannabinol*) = 203 results [108 results were from ClinicalTrials.gov, therefore only 95 results were downloaded]

ISRCTN Registry

Date Searched: Tuesday May 24, 2016

Text search: cannabis or canabis or cannabinoid or cannabidiol or CBD or cannabacae or marijuana or marihuana or hashish or hash or ganja or ganjah or hemp or bhang or charas or THC or tetrahydrocannabinol or tetra-hydrocannabinol or 9-tetrahydrocannabinol or DELTA-9-tetrahydrocannabinol (each keyword searched individually and results reviewed) = 8 results

NIH RePORTER

Date Searched: Monday May 16, 2016

Chronic Pain Search

((cannabis OR canabis OR cannabinoid OR cannabidiol OR CBD OR cannabacae OR marijuana OR marihuana OR hashish OR hash OR ganja OR ganjah OR hemp OR bhang OR charas OR THC OR tetrahydrocannabinol OR tetra-hydrocannabinol OR 9tetrahydrocannabinol) AND (pain OR pains OR painful OR migraine OR migraines OR headache OR headaches OR neuropathy OR neuropathies OR neuralgia OR arthritis OR fibromyalgia)) | Search in: Projects | Limit Project Search To: Project



Title, Project Abstracts | Limit Publication Search To: 2015-2016 = 50 results

Post-traumatic Stress Disorder (PTSD) Search

((cannabis OR canabis OR cannabinoid OR cannabidiol OR CBD OR cannabacae OR marijuana OR marihuana OR hashish OR hash OR ganja OR ganjah OR hemp OR bhang OR charas OR THC OR tetrahydrocannabinol OR tetra-hydrocannabinol OR 9tetrahydrocannabinol) AND (PTSD OR post-traumatic stress OR posttraumatic stress)) Search in: Projects | Limit Project Search To: Project Title,Project Abstracts | Limit Publication Search To: 2015-2016 = 5 results

Harms Search

((cannabis OR canabis OR cannabinoid OR cannabidiol OR CBD OR cannabacae OR marijuana OR marihuana OR hashish OR hash OR ganja OR ganjah OR hemp OR bhang OR charas OR THC OR tetrahydrocannabinol OR tetra-hydrocannabinol OR 9tetrahydrocannabinol) AND (harm OR harms OR harmful OR safe OR safety OR "side effects" OR "undesirable effects" OR "treatment emergent" OR tolerability OR toxicity OR adrs OR damage OR impaired OR impairing OR abuse OR addicted OR addiction OR addictions OR withdrawal OR "adverse effects" OR "adverse events" OR "adverse outcomes")) (Advanced), Search in: Projects | Limit Project Search To: Project Title,Project Abstracts | Limit Publication Search To: 2015-2016 = 220 results

AHRQ Gold (Grants On-Line Database)

Date Searched: Monday May 16, 2016

cannabis OR canabis OR cannabinoid OR cannabidiol OR CBD OR cannabacae OR marijuana OR marihuana OR hashish OR hash OR ganja OR ganjah OR hemp OR bhang OR charas OR THC OR tetrahydrocannabinol OR tetra-hydrocannabinol OR 9 tetrahydrocannabinol = 0 results

APPENDIX B. STUDY SELECTION

Inclusion Codes, Code Definitions, and Criteria

****Please note:* Important background/discussion papers may be coded "B" followed by an exclusion code, with notes or key words. For example: **B–X2, pearl for references**

1. Does the intervention or exposure consist of cannabis preparations including marijuana, hashish, tincture, hashish oil, infusion, and plant extract (*eg*, Sativex)?

No " STOP. Code X1 (Not relevant to topic)

Yes " Proceed to 2.

- 2. Is the article any of the following study designs or publication types:
 - Non-systematic or narrative review
 - Opinion/editorial
 - Cross-sectional study
 - · Individual case report

No "Proceed to 3.

Yes "STOP. Code X2 (Excluded study design or publication type)

3. Does the population include adults with chronic pain or PTSD?

No " Go to 10. Yes: Chronic pain " Go to 20. Yes: PTSD " Go to 30.

X4 = lab/blood/imaging findings

X5 = superseded by previous high-quality systematic review

Questions 10-13 deal with KQ3 (harms) in the general population

- 10. Are the majority of the study subjects either of the following:
 - Younger than age 18
 - Adults diagnosed with a psychotic disorder (eg, schizophrenia)
 - No "Proceed to 11.
 - Yes " STOP. Code X10 (Excluded pop for KQ3)
- 11. Does the study report any of the following harms:
 - Fungal infections
 - Cannabinoid hyperemesis syndrome
 - Other emerging harms (potential example: sudden onset of violent behaviors)
 - No "Proceed to 12.

Yes " Code I-11 (Gen pop, rare harms, KQ3) <u>Proceed</u> with items 12 and 13. Add Code I-13 if applicable.

- 12. Does the study report any of the following harms:
 - Psychotic symptoms
 - Cardiovascular events
 - Pulmonary/FEV1 outcomes
 - Infectious disease complications
 - Traffic collisions
 - Mortality
 - No "STOP. Code X12 (Gen pop, no harms of interest reported)
 - Yes "Proceed to 13.
- 13. Does the study design include a control group? The control group should differ from the primary group in dose or duration of cannabis use (including no use). However, a study comparing onset of cannabis use during adolescence vs adulthood would be excluded.
 - No "STOP. Code X13 (Gen pop, no control group for specified harms)
 - Yes " STOP. Code I-13 (Gen pop, has control group for specified harms)

Questions 20-22 deal with chronic pain

- 20. Do the study outcomes include either of the following:
 - · Cannabis use disorder
 - Withdrawal symptoms
 - No " Proceed to 21.
 - Yes " Code I-20 (*Pain pop, no controls needed for specified harms*) <u>Proceed</u> with items 21 and 22. Add Code I-22 if applicable.
- 21. Does the study report any of the following outcomes? The list below includes effectiveness outcomes and specific adverse effects of interest:
 - · Validated measures of pain intensity and pain-related function (including spasticity)
 - · Validated measures of pain-related outcomes (mood, depression, anxiety)
 - Validated measures of sleep quality
 - Validated measures of quality of life
 - Utilization of health services
 - · Reduction in opioid use or dosage
 - Social functioning/disability
 - Other substance use/substance use disorder
 - Mental health symptoms including depression, anxiety, *etc* (not psychotic symptoms)
 - Cognitive effects (*eg*, IQ, SLUMS, or measures of memory, processing speed, attention, learning, executive function, *etc*)
 - Employment
 - Weight gain
 - Diversion
 - Insomnia
 - No "STOP. Code X21 (Pain pop, no outcomes of interest)
 - Yes " Proceed to 22.
- 22. Is the study design a controlled clinical trial, case-control, or cohort study with a comparison group?
 - No "STOP. Code X22 (Pain pop, excluded study design)
 - Yes "STOP. Code I-22 (Pain pop, addresses KQ1 and/or KQ3)

Questions 30-32 deal with PTSD

- 30. Do the study outcomes include either of the following:
 - · Cannabis use disorder
 - Withdrawal symptoms
 - No "Proceed to 31.
 - Yes " Code I-30 (*PTSD*, no controls needed for specified harms) Proceed with items 31 and 32. Add Code I-32 if applicable.
- 31. Does the study report any of the following outcomes? The list below includes effectiveness outcomes and specific adverse effects of interest:
 - · Validated PTSD clinical interviews and symptom inventories, such as:
 - ▼ Clinician Administered PTSD Scale (CAPS)
 - ▼ PSTD Checklist (PCL)
 - ▼ PTSD Symptom Scale (PSS)
 - ▼ Posttraumatic Diagnostic Scale (PDS).
 - Validated measures of mental health symptoms commonly associated with PTSD (mood, depression, anxiety)
 - Validated measures of sleep quality
 - · Validated measures of quality of life
 - Utilization of health services
 - · Reduction in benzodiazepine use or dosage
 - Social functioning/disability
 - Other substance use/substance use disorder
 - Mental health symptoms including depression, anxiety, etc (not psychotic symptoms)
 - Cognitive effects (*eg*, IQ, SLUMS or measures of memory, processing speed, attention, learning, executive function)
 - Employment
 - Weight gain
 - Diversion
 - Insomnia

No "STOP. Code X31 (PTSD, no outcomes of interest)

Yes " Proceed to 32.

- 32. Is the study design a controlled clinical trial, case-control, or cohort study with a comparison group?
 - No " STOP. Code X32 (PTSD, excluded study design)
 - Yes "STOP. Code I-32 (PTSD, addresses KQ2 and/or KQ3)

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APPENDIX C. QUALITY ASSESSMENT

Domain	Criteria
Sequence generation	Was the allocation sequence adequately generated?
Allocation concealment	Was allocation adequately concealed?
Blinding	Was knowledge of the allocated intervention adequately prevented during the study?
Incomplete outcome data	Were incomplete outcome data adequately addressed? Consider attrition, intention-to-treat analysis
Selective outcome reporting	Are reports of the study free of suggestion of selective outcome reporting?
Other sources of bias	Was the study apparently free of other problems that could put it at a high risk of bias (ROB)?
Overall assessment of potential for bias	Low/Unclear/High

Cochrane Risk of Bias (ROB) Assessment Criteria for Trials²⁰

Trials in Patients with Chronic Pain – Risk of Bias (ROB) Assessment

Study; Pain type	Sequence generation	Allocation concealment	Blinding	Incomplete outcome data	Selective outcome reporting	Other sources of bias	Overall: Low/Unclear/ High ROB
Abrams, 2007 ³⁵ HIV- associated sensory neuropathy	Yes: Randomization (1:1) to cannabis or placebo cigarettes was computer- generated by the study statistician	Yes: allocation managed by an independent research pharmacist	Yes: Treatment was double-blind, NOS. The National Institute on Drug Abuse provided identically appearing pre-rolled cannabis and placebo cigarettes	Yes: Low loss to follow-up.	Probably yes. Does not state protocol was reported prior to study.	Yes	Low
Berman, 2004 ³⁶ Neuropathic Pain from Brachial Plexus Avulsion	Yes - computer generated list	Unclear	Uncertain; notes that treatment sequence was blinded via sealed code break envelopes but no further details given	Yes: ITT analysis, attrition described	Unclear: No protocol mentioned, but outcomes reported in the methods are included in results	Unclear: No washout period between treatment regimens	Low
Blake, 2006 ⁵⁹ Rheumatoid arthritis	Unclear (permuted blocks of four)	Unclear (not reported)	Unclear (not reported)	Yes: low attrition, comparable across groups	Yes	Unclear (COI statement notes the study was funded by a drug company)	Unclear
Collin, 2010 ⁴⁷ MS	Unclear, method not described	Unclear, method not described	Probably yes, but not described in detail	Yes: ITT	Yes	The data on pain is limited only to spasticity responders.	Unclear
Corey-Bloom, 2012 ⁴⁸ MS	Unclear, method not described	Unclear, method not described	Yes: identical placebo cigarettes	Yes: < 80% attrition; also did worst case scenario analysis	Yes	Excluded high doses of narcotic medications for pain, but did not control for or examine concomitant use of analgesics because spasticity was primary criteria and outcome of interest. No mention of analgesic use for pain.	Unclear

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Benefits and Harms of Cannabis for Chronic Pain or PTSD

Evidence-based Synthesis Program

Study; Pain type	Sequence generation	Allocation concealment	Blinding	Incomplete outcome data	Selective outcome reporting	Other sources of bias	Overall: Low/Unclear/ High ROB
De Vries, 2016 ⁵⁸ Abdom-inal pain	Yes: computer- generated randomization list stratified for opioid and non-opioid users by using separate lists	Yes: central allocation ("Independent pharmacists dispensed either active or placebo tablets according to a computer- generated randomization list")	Yes: "Treatment allocation was strictly concealed from participants, investigators, and all other study personnel involved in the study until end of study and database lock."	No: ITT analysis not performed	Unclear: for several secondary outcomes (depression, quality of life, EEG, <i>etc</i>), researchers simply stated "did not change after THC treatment compared with placebo" but did not give any values.	No power calculation, likely inadequate power to detect differences. Also, what was originally supposed to be 2 trials were combined into one study "because of a disappointing recruitment."	High
Ellis, 2009 ³⁷ HIV- associated sensory neuropathy	Yes: Randomization was performed by a research pharmacist using a random number generator, and the key to study assignment was withheld from investigators until completion statistical analyses.	Yes: key was withheld from study investigators until completion of analysis.	Yes: double blind, cigarettes "were constructed of the same base material." Assessed effectiveness of blinding among participants	Moderate: 6/34 lost. ITT. Similar baseline characteristics.	Yes: all outcomes in methods were reported.	This is a specific population of HIV pain; Patients allowed to use own analgesia; Used validated scales for pain measures	Low
Johnson, 2010 ⁵⁵ Cancer	Not described; Table 1 shows general similarities between groups but baseline opioid use was lower in one group vs others	Not described	Not described for investigators; Yes for patients; bottles were similar between active and placebo, though patients on active were able to guess their group	Yes; ITT done and overall attrition >80%, though one group lost >80%	Yes	This is a specific population of cancer pain; Patients allowed to use own analgesia; Used validated scale and self-reported scale as co-outcomes for pain	Unclear

Benefits and Harms of Cannabis for Chronic Pain or PTSD

Evidence-based Synthesis Program

Study; Pain type	Sequence generation	Allocation concealment	Blinding	Incomplete outcome data	Selective outcome reporting	Other sources of bias	Overall: Low/Unclear/ High ROB
Langford, 2013 ⁴⁹ MS	Yes: "Randomization occurred using a pre-determined computer- generated randomization code in which treatment allocation was stratified by center, and used randomly permuted blocks of variable sizes."	Yes	Yes: double-blind	Yes - ITT, but in group A, 26/167 (cannabis) and 16/172 (placebo) withdrew.	Yes: outcome measures reported	No: Strong placebo effect; Placebo group patients who titrated to the maximum dose had disproportionate improvements in pain scores, and a number of these patients reached the maximum permitted dose as the study period was drawing to a close. Self- titration combined with a subjective endpoint seems therefore to have significantly impacted the placebo response.	Unclear
Lynch, 2014 ³⁸ Chemotherap y-Induced Neuropathic Pain	Yes	Yes	Yes	Yes (16/18 completers)	Yes	Yes	Low
Notcutt, 2004 ³³ Mostly neuro- pathic; 47% MS	Yes	Yes: randomization done externally	Yes: trial stated as being double-blind and delivery of intervention and placebo were matched	Unclear: for the trial portion, 71% (24/34) patients were included in analysis due to withdrawal/use of rescue meds	Yes	Excluded those who had to use rescue medications; also, only randomized pts who reported a positive response to medical cannabis.	Low

Study; Pain type	Sequence generation	Allocation concealment	Blinding	Incomplete outcome data	Selective outcome reporting	Other sources of bias	Overall: Low/Unclear/ High ROB
Notcutt 2012 ²⁹	Yes: independent statistician produced an allocation schedule using balanced permuted blocks of 4 with computer- based algorithm	Yes: independent statistician	Unclear: no methods described	Yes: had high attrition (~50%) and only some subjects met treatment failure but based on disposition tree, all included subjects were analyzed	Yes: appeared to report on relevant outcomes	No: underpowered (though CI adjusted to help with this), some participants restarted on their own nabiximols prior to final assessment (likely to reduce the effect of the drug)	Unclear: multiple areas of uncertainty; study was under- powered and patients could have restarted nabiximols prior to assessment
Novotna 2011 ²⁷	Unclear, methods not described	Unclear, methods not described	Unclear: states that trial was double-blind but no details on methods; comment that inclusion into trial based on investigator assessment that patient remained blinded during initial phase of study	Yes: ITT though patients without post- randomization efficacy data were excluded, all patients who had received one dose of medication included in safety analyses; attrition reported (12% for nabiximols group and 2% for placebo).	Yes: had clearly stated pre- specified primary outcome and included multiple secondary outcomes	Yes: no major issues identified aside from lack of clarity around the methods used for allocation, randomization	Low: though limited data on methodology around allocation and blinding, authors state that study was double- blind and had low attrition with ITT analyses and pre-specified outcomes
Noyes, 1975 ⁵⁶ Cancer	No	No	Yes (patients), No (providers)	Unclear, though 34/36 patients were reported to be completers.	No (only reported results for significant tests, refer to "other differences" that did not reach significance)	Pain measure not validated	High

Study; Pain type	Sequence generation	Allocation concealment	Blinding	Incomplete outcome data	Selective outcome reporting	Other sources of bias	Overall: Low/Unclear/ High ROB
Nurmikko, 2007 ³⁹ Neuropathic Pain Characterised by Allodynia	Yes: "randomly permuted blocks stratified by center and was generated using a computer based pseudo- random number algorithm"	No: a copy of randomization schedule in patient- specific sealed envelopes sent to the pharmacy in each center	Yes	Yes: ITT analysis, attrition described	Unclear: no protocol mentioned, but outcomes reported in the methods seem to be included in results	Unclear: "GW Pharma acted as the sponsor of the study, provided the medication, participated in the study design, coordinated the study between centers and carried out the first set of analyses. The analyses were verified by an independent statistician."	High
Portenoy, 2012 ⁵⁷ Cancer	Unclear	Unclear	Yes (patients), Unclear (research staff)	No (attrition 27%)	Yes	Yes	Unclear
Rog, 2005 ⁵⁰ MS	Yes: "Patients were randomized using a predetermined randomization code drawn up by a statistician who remained unknown to study personnel throughout the duration of the trial."	Yes: "statistician remained unknown to study personnel throughout the duration of the trialTreatment allocation was made using randomized permuted blocks of four (two active drug, two placebo), with treatments sequentially assigned."	Yes: double-blind, and "Placebo was designed to match the appearance, smell, and taste of the active formulation but contained no active components, in ethanol: propylene glycol (50:50) excipient. To facilitate blinding, patients completed pain and sleep	Yes: only 2/64 did not complete. Both received cannabis - one adverse event, one withdrew consent. ITT.	Yes: all outcomes reported in methods were reported.	Unclear: Required no change of concomitant meds, but no mention of controlling for meds/sensitivity analysis, or analyzing by med class.	Low

Study; Pain type	Sequence generation	Allocation concealment	Blinding	Incomplete outcome data	Selective outcome reporting	Other sources of bias	Overall: Low/Unclear/ High ROB
Selvarajah, 2010 ⁴⁰ Diabetic Neuropathy	Unclear: no information other than saying it was randomized	Unclear: no information	Unclear: no information about blinding except stating that it was a double-blind trial	Unclear: completed ITT analysis included but 1 patient with protocol violation was excluded.	Unclear: states "Tolerability and side effects were evaluated using standardized forms" but does not report these results (except saying 6 patients overall withdrew due to AEs). Unlikely to have introduced significant bias.	No power calculation reported, very likely the study had inadequate power to detect differences	Unclear
Serpell, 2014 ⁴¹ Peripheral neuropathic pain (PNP) associated with allodynia	Yes: a predetermined computer- generated randomization code	Unclear	Yes	Yes: ITT analysis, attrition described	Yes: protocol available (https://clinicaltrials .gov/ct2/show/NCT 00710554), study reports all outcomes mentioned	Recorded medications used; also did allodynia testing	Low
Van Amerongen, 2017 ⁵¹ MS	Unclear: randomization schedule prepared by independent statistician; allocated "on the basis of the date of eligibility of the individual because the identification numbers are assigned at that moment."	Unclear: "The schedule was sent to the hospital pharmacy, and sealed envelopes for code breaking were available for the investigator"; opaque envelopes not specified	Yes: matching placebo tablets, "All staff involved in the clinical execution of the study were blinded until all data were collected and the database was locked."	Yes: attrition with reasons reported by group, and ITT analysis performed.	Unclear: the McGill Pain Questionnaire was mentioned as a secondary endpoint in the online protocol and Methods section, but was not reported in the Results.	Yes	Unclear
Wade 2003 ³⁰ MS (67%)	Yes: sequence generated with Williams squares	Yes: stated that participants and staff were blinded.	Yes: identical sprays used with masking flavor; investigators were not aware of coding	Yes: attempted to analyze those who took rescue medications vs entire sample	Yes: looked at range of symptoms	Small sample size	Low

Evidence-based Synthesis Program

Study; Pain type	Sequence generation	Allocation concealment	Blinding	Incomplete outcome data	Selective outcome reporting	Other sources of bias	Overall: Low/Unclear/ High ROB
Wade, 2004 ⁵² MS	Unclear: permuted blocks of 4, stratified by primary symptom and center	Unclear: the pharmacist at each center was provided with a randomization scheme and assigned the treatments in sequential patient number order	Unclear: investigators did not assess the degree of blinding of patients and outcome assessors, though a stronger effect was found for pain outcome in placebo compared with active treatment.	Yes	Unclear	Unclear: study was underpowered for pain outcome	Unclear
Wallace, 2015 ⁴² Diabetic Neuropathy	Yes: "Randomization was performed by a research pharmacist using a random number permutations"	Yes: "Randomization was performed by a research pharmacist using a random number permutations and the key to study assignment was withheld from investigators until completion of statistical analyses"	Unclear: blinding may have been compromised due to crossover design and euphoria from the drug, but analyses didn't find this to be significant	Yes: did not appear to be any missing data; one patient only participating in some of the sessions were only analyzed for those sessions	Yes: reports the outcomes mentioned in the CT.gov protocol (https://clinicaltrials .gov/ct2/show/NCT 00781001) – although some of results are mostly in charts which may make it hard to abstract all data accurately	Study only enrolled 16 patients, rather than 20 in power calculation. Also, only very short-term	Low
Ware, 2010 ⁴³ Post-surgical or post- traumatic neuro-pathic pain	Unclear (just notes a Latin Square design)	Unclear (not reported)	Yes (notes factors to maintain blinding such as placebo comparability confirmed by objective assessment)	Yes (very low attrition)	Yes	Yes	Low
Wilsey, 2008 44 Neuro-pathic pain	Yes	Yes	Uncertain: no details given on blinding, beyond statement to the effect	Yes: Attrition <80% for all arms; all available data used in analysis	Yes	Pain scales were self-report; also used neurocognitive testing and evoked pain threshold	Low
Wilsey, 2013 ⁴⁵ Neuro-pathic pain	Yes	Yes	Uncertain: no details given on blinding, beyond statement to the effect	Yes: Attrition <80% for all arms; all available data used in analysis	Yes	Yes: pain scales were self-report; also used neurocognitive testing	Low

Study; Pain type	Sequence generation	Allocation concealment	Blinding	Incomplete outcome data	Selective outcome reporting	Other sources of bias	Overall: Low/Unclear/ High ROB
Wilsey, 2016 ⁴⁶ Spinal cord injury	Yes: used a Web- based random number-generating programto determine the sequence of administration.	Yes: The allocation schedule was maintained by a research pharmacist and concealed from other study personnel.	Yes: Placebo cannabis was derived from whole plant material with extraction of delta 9- THC.	Unclear: Attrition with reasons reported by group, method for handling missing data was not described.	Yes: Protocol available and all outcomes mentioned appear to be reported	No power calculation reported, but since a significant difference was found between interventions, this is unlikely to have introduced significant bias	Low
Zajicek, 2003 ⁵³ MS	Yes: The coordinating center allocated the patient a trial number and then forwarded relevant details to the central trial pharmacy, where randomization took place, using a dedicated stand- alone computer.	Yes: Throughout the study, the list of treatment allocation codes was kept at the central trial pharmacy, located separately from the coordinating office.	No: most in the active treatment group guessed correctly that they were on active treatment; potential bias despite that placebo group was effectively blinded.	Yes	Yes	Patients selected for spasticity, not pain. Power calculation based on projected effects on spasticity. Baseline pain scores not reported, only whether improved, unchanged, or deteriorated. Unclear whether pain levels were high or low to begin with.	High
Zajicek, 2012 ⁵⁴ MS	Yes: Computer generated permuted block randomization	Yes	Yes: Matched placebo capsules.	Yes	Yes	Yes	Low

Criteria	Englund 2013 ⁹¹	Kaufmann 2010 ⁸⁹
Sequence generation	Unclear - methods not described	Unclear - methods not described
Allocation concealment	Unclear - methods not described	Unclear - methods not described
Blinding	Yes - double blind, randomly allocated	Unclear - double blind. No details provided.
Incomplete outcome data	NA - all participants completed study	No - One participant developed acute psychotic symptoms and was not included in the statistical analysis, but was qualitatively described.
Selective outcome reporting	Yes - All relevant outcomes appear to be reported	Yes - appear to report all outcomes.
Other sources of bias	Yes - no major issues identified aside from lack of clarity around the methods used for sequence generation and allocation.	No - small sample/under powered.
Overall assessment of potential for bias	Low - despite lack of clarity about sequence generation and allocation concealment.	Moderate - due to lack of clarity about sequence generation and allocation concealment and small sample size.

Trials Assessing the Risk of Psychotic Symptoms with Cannabis Use – Risk of Bias (ROB) Assessment

Quality Assessment Criteria for Observational Studies, Based on the Newcastle-Ottawa Scale²¹

Representativeness of the exposed cohort Enter 0 or 1:
1 = truly representative of the average patient in the community
1 = somewhat representative of the average patient in the community
0 = selected group of users (<i>eg</i> , nurses, volunteers)
0 = no description of the derivation of the cohort
Selection of the non-exposed cohort
Enter 0 or 1:
1 = drawn from the same community as the exposed cohort
0 = drawn from a different source
0 = no description of the derivation of the non-exposed cohort
Ascertainment of exposure
Enter 0 or 1:
1 = biological test (<i>eg</i> , blood/urine)
1 = structured interview
1 = written self-report that characterizes dose (current or cumulative)
0 = written self-report without quantification of exposure
0 = no description
Precision of Exposure Dose Ascertainment
Enter 0 or 1:
1 = amount and time
0 = no information about amount and time
Ascertainment of exposure done prospectively or retrospectively Enter 0 or 1:
1 = Prospectively
0 = Retrospectively
Demonstration that outcome of interest was not present at start of study, OR baseline
assessment
Enter 0 or 1:
1= yes
0 = no
Adjustment for confounding (rendering comparability of cohorts on the basis of the design or
analysis)
Add points: Minimum 0, Maximum 2
1 = study accounts/controls for other substance use
1 = study controls for any additional factor (mental health comorbidity; medication use; severity of PTSD;
mental health comorbidity and treatment; socioeconomic status)
0 = no adjustment for potential confounders
Assessment of outcome
Enter 0 or 1:
1 = objective measure
1 = validated self-report measures
0 = no information or non-validated measures
Was follow-up long enough for outcomes to occur? Enter 0 or 1:
1 = yes (need to define adequate follow-up period for outcome of interest)
0 = no
Adequacy of follow-up of cohorts
Enter 0 or 1:
1 = complete follow-up; all subjects accounted for.
1 = subjects lost to follow-up unlikely to introduce bias; small number (less than 20 %) lost, or
description was provided of those lost.
0 = follow-up rate < 80% and no description of those lost.
0 = no statement

Criteria	Ware 2015 ³¹	Storr 2014 ³²	Fiz 2011 ³⁴
Representativeness of the exposed cohort	1 - included patients with non- cancer pain but had to be moderate/severe and refractory	1 - exposed cohort was equal for males and females although IBS impacts females at a slightly higher base rate	1 - somewhat; these are treatment resistant patients in particular
Selection of the non- exposed cohort	1 - all drawn from same clinical centers	1 - drawn from same source	0 - 2 of the recruitment sites were the same (FM associations and outpatient rheumatology) but cannabis group also recruited from cannabis association.
Ascertainment of exposure	1 - pharmacy dispensed and recorded use	0 - self-report; only method of administration (<i>ie</i> , smoking) recorded	0 - information reported about duration of cannabis use (<i>ie</i> , 1 year) and administration modality, but no info provided about dose or cannabinoid concentration.
Precision of Exposure Dose Ascertainment	1 - dosing described	0 - no dosing information provided	0 - method of administration varied among users (smoking 54%; oral 46%; combined 43%), duration and frequency of use varied among users. Dosage varied among users ("1-2 cigarettes each time when smoked or 1 spoonful each time when eating"). No info on THC/CBD content given. 39% used daily, 18% used 2-5 days per week.
Ascertainment of exposure done prospectively or retrospectively	1 - prospectively	0 - cross-sectional so ascertainment based on one timepoint	0 - exposure groups established by use status
Demonstration that outcome of interest was not present at start of study, OR baseline assessment	1 - all results compared to baseline	0 - no baseline	1 - baseline data gathered 2 hours prior to exposure
Adjustment for confounding	2 - cohort significantly different on age, gender, disability status, tobacco status, past cannabis use, drug abuse screening, average pain intensity (cannabis users higher) and medications – however, these group differences were controlled for in the inferential statistics.	2 - study adjusts for demographic variables, tobacco smoking status, time since diagnosis, and biological use	0 - no adjustments made

Observational Studies in Patients with Chronic Pain – Risk of Bias (ROB) Assessment



Criteria	Ware 2015 ³¹	Storr 2014 ³²	Fiz 2011 ³⁴
Assessment of outcome	1 - objective/validated measures used	0 - surgical history gleaned from medical chart (only measure of utilization provided) no other validated measures reported for our PICOTS. Side effects and perceived utility of cannabis for treatment of IBD symptoms all subjective and only descriptive data is provided for users.	1 - validated self-report measures for outcomes (<i>eg</i> , VAS, SF-36)
Was follow-up long enough for outcomes to occur?	1 - (12 months follow-up)	0 - no follow-up	0 - difficult to ascertain sustainability of outcomes, only 2 hours of follow-up
Adequacy of follow-up of cohorts	1 - > 20% loss to follow-up in the cannabis group but all subjects are accounted for and all subjects included in the primary safety analysis	0	1 - appears to be complete follow-up
Comments on study quality	Low ROB - there are some concerns as noted below but what is measurable by scale appears to be properly done - Study's primary outcomes were adverse events, other outcomes were secondary; Study notes that protocol changes were made but no details provided; Study did not recruit pre-specified sample size for power; Multiple adjustments and subgroup analyses were undertaken; Also, strange that inclusion into cannabis group relied on use of cannabis but there are persons included there who are cannabis naïve and who were ex-users; baseline demographics/ population details differed by group, though adjustments made in analysesthe majority (66%) of the cannabis users were experienced, making the generalizability to cannabis- naïve users difficult, and differences in the follow-up times between the control and exposure group may have artificially inflated the number of AEs reported by cannabis users.	High ROB - dosing information was not provided or consistent for users, data collection only at one time point so no f/u data provided. Minimal outcomes of interest.	High ROB - dosing information was not provided or consistent for users, participants gathered from different sources introducing selection bias; groups were established by exposure status and those using cannabis are likely to differ from others not using cannabis (although baseline characteristics are not different per study authors and this is the only way to conduct a cohort study), also concern that there were no adjustments made for other medications used, small sample size, use of self-reported measures, very limited follow-up with a pre-, post-design rather than between group comparison for primary outcome
Notes on Applicability	Patients had treatment moderate/severe, refractory chronic pain but otherwise applicable, especially since drawn from clinical centers		Patients had treatment resistant FM



Criteria	Wilkinson 2015 ⁶⁰	Johnson 2016 ⁶¹
Representativeness of the exposed cohort	1	1
Selection of the non-exposed cohort	1	1
Ascertainment of exposure	0 (self-report)	0 (self-report)
Precision of Exposure Dose Ascertainment	0 (not specific)	0 (not specific)
Ascertainment of exposure done prospectively or retrospectively	1	0
Demonstration that outcome of interest was not present at start of study, OR baseline assessment	1	n/a
Adjustment for confounding	1 (included all assessed confounders related to cannabis use)	0
Assessment of outcome	1 (validated self-report measures)	1 (validated self-report measures)
Was follow-up long enough for outcomes to occur?	1 (4 months)	n/a
Adequacy of follow-up of cohorts	1	n/a
Comments on study quality	Medium ROB	High ROB
Notes on Applicability	VA population with PTSD	VA population with PTSD

Observational Studies in Patients with PTSD – Risk of Bias (ROB) Assessment

Frost 201372 Hancox 2010⁶⁹ Criteria Pletcher 2012⁷⁰ Mittleman 2001⁷³ Carvalho 201574 Representative-1 - trulv 1 - somewhat 1 - somewhat 1 - in half the 1 - somewhat ness of the representative representative of studies, these representative representative of MI MI patients - not patients - not exposed cohort Community (birth cohort, but were hospital based study in 4 community, but community, but most patients, half the for that reason most MI patients MI patients would get studies used cities does not would get cared cared for in hospital representing represent older cancer registry different parts of for in hospital and and this was multisite patients in the data this was multisite hospital study country, community) ethnically diverse hospital study group. 1 - drawn from 1 - self-control Selection of the 1 - same 1 - same community 1 - most studies non-exposed same community community found general cohort population controls (eq, electoral rolls, random digit dialing) Ascertainment of 1 - structured 1 - interview 0 - risk of recall 1 - interview 1 - most studies interview bias, not clear how used structured exposure accurate recalled interview pattern of use over prior year was since this formed basis for control (expected frequency of hourly use) there is some potential for bias. 0 - not enough 1 - amount and 1 - amount and 0 - time only, and only 1 - most gathered Precision of information about Exposure Dose time time at baseline information about Ascertainment amount and time amount and time Ascertainment of 1 - prospectively 1 - prospectively 0 - retrospectively 0 - retrospectively 0 - retrospectively exposure done prospectively or retrospectively 1 - yes - PFTs Demonstration 1 - yes, serial PFT n/a 1 - yes (inception 0 - no that outcome of were measures, and cohort) longitudinally they adjusted for interest was not present at start of collected spirometry at age baseline PFT study, OR 15 data were baseline available and assessment outcomes were reported as change from baseline

Observational Studies of Medical Harms Associated with Cannabis Use – Risk of Bias (ROB) Assessment



Criteria	Pletcher 2012 ⁷⁰	Hancox 2010 ⁶⁹	Mittleman 2001 ⁷³	Frost 2013 ⁷²	Carvalho 2015 ⁷⁴
Adjustment for confounding	1 - for PFT outcomes, most important covariate is tobacco exposure along with gender, age, race all of which were well accounted for.	1 - accounts for tobacco exposure, age, gender which are probably most relevant for the PFT outcomes - did not account for race, SES, second hand smoke exposure, <i>etc</i>	0 - not clear that they account for tobacco use in hour prior to MI	1 - propensity score matching - adjusted for tobacco, other substance use, SES other factors	1 - most studies adjusted for tobacco use and alcohol use
Assessment of outcome	1 - PFTs, objective measure	1 - PFTs	1- objective assessment of MI outcome	1 - national death index	1 - only included studies of patients with definitive HNC
Was follow-up long enough for outcomes to occur?	1 - yes	1 - yes	n/a	1 - yes, partly - the exposed group was younger and the number of mortality events therefore relatively small, but 18 year f/u	n/a
Adequacy of follow-up of cohorts	1 - data from 98% of participants, 95% of all visits had complete data	1 - data from 96% of original cohort at 32 years	n/a	1 - national death index	n/a
Comments on study quality	Low ROB. Well- conducted, prospective cohort study. Should be one of the better sources of data for this outcome.	Low ROB. Well- conducted, prospective cohort study. Similar to Pletcher study, but did not have data on linear trends.	High ROB. Case- crossover study with several potential sources of bias including recall bias, small # patients with exposure of interest, and lack of clarity re: accounting for tobacco use.	High ROB. Information on exposure (both cannabis and tobacco) only available at baseline interview. Assess long-term mortality, but no information on total use over the period of follow-up, making it difficult to assess relationship between exposure and outcome. Moreover, cannabis users were different than non- users - confounders were adjusted for, but strong possibility of residual confounding.	Medium ROB. Ascertainment of exposure is necessarily limited by retrospective nature and issues of recall bias.

Criteria	Pletcher 2012 ⁷⁰	Hancox 2010 ⁶⁹	Mittleman 2001 ⁷³	Frost 2013 ⁷²	Carvalho 2015 ⁷⁴
Notes on Applicability	Applicable to younger populations (< 30)	Applicable to younger populations	Most cannabis users were male	Most cannabis users were male, younger than nonusers	Very wide range of ever cannabis use - some of the studies with very low rates of use may not be applicable, but the consistency of results across different study populations is reassuring.



Criteria	Zhang 2015 ⁷⁵	Callaghan 2013 ⁷⁶	Gurney 2015 ⁷⁷	Chacko 2006 ⁷⁸
Representative- ness of the exposed cohort	1 - international, mix of hospital-based and community studies	1 - nearly all (98%) 18- 20 year old males	1 - cancer registry cases with community-based controls	1 - representative of transitional cell ca population, at least in VA
Selection of the non-exposed cohort	1 - all drew controls either from same hospital/clinic, or the community	1 - drawn from same population	1 - drawn from same population (random general population in 2 studies and friends of cases in one study which is a potential source of selection bias)	0 - drawn from urology clinic, presenting for different reason - not representative of community
Ascertainment of exposure	1 - written self-report with information on duration and frequency	0 - self-report without adequate quantification	0 - interview in 2 studies and written self-report with quantification in other, but not clear that interviewers were blinded to case/control status of participant	1 - written self-report with information on duration and frequency
Precision of Exposure Dose Ascertainment	1 - amount and time	0 - minimal information about exposure over time	1 - amount and time	1 - amount and time
Ascertainment of exposure done prospectively or retrospectively	0 - retrospectively	0 - retrospectively, and only at time of conscription	0 - retrospectively	0 - retrospectively
Demonstration that outcome of interest was not present at start of study, OR baseline assessment	1 - performed additional analyses excluding patients who had used within 2 years of cancer diagnosis (to evaluate possibility of reverse causality)	0 - no, but very unlikely that outcome was present in young age group	1 - (case-control)	n/a
Adjustment for confounding	1 - adjusted for tobacco use and some other sociodemographic factors	0 - adjusted for multiple factors, but did not have a way of quantifying tobacco exposure after conscription which is likely to have been heaviest amongst those with heavier cannabis use	1 - adjusted for major confounders relevant to disease (including cryptorchidism), but one study did not adjust for alcohol or tobacco use (but was also the smallest of the studies)	0 - important confounders considered, but they did not report adequately the adjusted analyses
Assessment of outcome	1 - only histologically confirmed lung cancer	1 - based on national medical records, claims - fair validation	1 - histologically confirmed cancers	1 - confirmed cancers
Was follow-up long enough for outcomes to occur?	n/a	1 - yes	n/a	n/a
Adequacy of follow-up of cohorts	n/a	1 - 1.9% lost to f/u due to emigration	n/a	n/a

Medical Harms Observational Studies – Risk of Bias (ROB), Continued



Criteria	Zhang 2015 ⁷⁵	Callaghan 2013 ⁷⁶	Gurney 2015 ⁷⁷	Chacko 2006 ⁷⁸
Comments on study quality	Medium ROB - ascertainment of exposure is necessarily limited by retrospective nature and issues of recall bias.	High ROB - biggest issue was that the main exposure and main confounder (tobacco use) were only determined at time of conscription. High risk of residual confounding due to ongoing tobacco exposure for finding of heavy cannabis use association with lung cancer.	High ROB - the meta- analysis itself was well done, but there were methodologic deficiencies in all 3 included studies. The smallest study did not control for important confounders such as tobacco. Low response rates among controls or cases in the 2 bigger studies. There was a potential for ascertainment bias, and recall bias is also an issue. Use of friends as controls in one study is a potential source of bias. The largest and methodologically strongest study showed results consistent with overall findings, direction of effect was consistent across studies, there was a dose- response relationship, and the authors do highlight some biologic plausibility to findings.	High ROB - small study, 2 VA sites, very little information on adjusted analyses, control group were symptomatic patients in urology clinic so not representative of community, reverse causality a real concern (<i>ie</i> , cancer patients may have been using cannabis to palliate symptoms - no information on timing of use and diagnosis), recall bias
Notes on Applicability	Variety of settings, included squamous cell and adenocarcinoma patients but few patients with other types of lung cancer.			VA only - 2 sites. One of the sites located in a town with prominent textile industry (and, thus, dye exposure). Small number of patients.



Observational Studies of Adverse Mental Health Effects Associated with Cannabis Use – Risk of Bias (ROB) Assessment

Criteria	Di Forti 2009 ⁹²	Dominguez 2010 ⁸⁷	Kuepper 2011 ⁸⁶	Mason 2008 ⁹⁰	Rossler 2011 ⁸⁸	van Nierop 2013 ¹³⁷
Representative- ness of the exposed cohort	1 - first episode psychosis (presenting to the hospital)	1 - representative population study.	1 - representative population study.	0 - No information about the population from which the sample was recruited. Recreational cannabis smokers who used cannabis at least once a month. No personal history of diagnosed mental illness. Lifetime drug usage of other illicit drugs in the cannabis group commonly included amphetamines, benzodiazepines, cocaine, ketamine, LSD, and heroin.		1 - somewhat representative (siblings of individuals with psychotic disorders and healthy controls in the same geographical areas)
Selection of the non-exposed cohort	0 - No description of source. Control group was individuals with no psychotic episodes.	1 - same population	1 - same population	0 - No description of source	1 - same population	1 - same community
Ascertainment of exposure	1 - Cannabis Experience Questionnaire	1 - Munich composite international diagnostic interview (DIA-X/M- CIDI)	1 - Munich composite international diagnostic interview (DIA-X/M-CIDI)	1 - self-report and urinalysis	1 - Structured Psychopathological Interview and Rating of the Social Consequences of Psychological Disturbances for Epidemiology (SPIKE)	1 - urinalysis and CIDI

Criteria	Di Forti 2009 ⁹²	Dominguez 2010 ⁸⁷	Kuepper 2011 ⁸⁶	Mason 2008 ⁹⁰	Rossler 2011 ⁸⁸	van Nierop 2013 ¹³⁷
Precision of Exposure Dose Ascertainment	1 - Assessed type and frequency, as well as potency	0 - > or < 5 times since last exposure	0 - > or < 5 times since last exposure	0.5 - Participants contacted researchers when they were using cannabis recreationally. The study team went to meet them for testing. Dose not ascertained.	1 - frequency of use	0.5 - used interviews to determine lifetime use and urinalysis to determine current use. No information re: dose, frequency, <i>etc</i>
Ascertainment of exposure done prospectively or retrospectively	0 - retrospectively	0 - retrospectively	0 - retrospectively	1 - prospectively	0 - retrospectively	0 - retrospectively
Demonstration that outcome of interest was not present at start of study, OR baseline assessment	1 - first psychotic episode	0 - not excluded	0 - not excluded	0 - not excluded, and no baseline assessment.	1 - although clinical diagnoses of psychotic disorders were not assessed with the SPIKE at baseline through 1999, two-thirds of the sample were at "high risk" for subclinical psychosis symptoms based on Symptom Checklist 90— Revised (SCL-90-R) scores.	1 - healthy siblings of individuals with a psychotic disorder (high risk) and healthy controls.
Adjustment for confounding	2 - adjusted for age, gender, ethnicity, other stimulant use, education, and employment status.	1 - controls for depression but not other substance use	2 - Adjusted for age at baseline, sex, baseline SES, use of other drugs at baseline and T2, trauma before the age of 14 as assessed at baseline, and urban/rural environment.	1 - performed sensitivity analysis for other drug/alcohol use	2 - adjusted for sex, familial background, socio- economic status, family history of mental disorders, other family problems, and school problems, and used step wise multivariate analysis with each substance entered individually.	0 - adjusts only for age, sex, high-risk sibling status

Criteria	Di Forti 2009 ⁹²	Dominguez 2010 ⁸⁷	Kuepper 2011 ⁸⁶	Mason 2008 ⁹⁰	Rossler 2011 ⁸⁸	van Nierop 2013 ¹³⁷
Assessment of outcome	1 - hospital admission	1 - Munich composite international diagnostic interview (DIA-X/M- CIDI)	1 - Munich composite international diagnostic interview (DIA-X/M-CIDI)	0 - Psychotomimetic States Inventory (PSI) - the study is a validation study.	1 - SPIKE and SCL-90-R	1 - Community Assessment of Psychic Experience (CAPE)
Was follow-up long enough for outcomes to occur?	NA	1 - Mean T1 1.6, T2 3.5, and T3 8.4 years (range=7.3-10.5)	1 - Mean T1 1.6, T2 3.5, and T3 8.4 years (range=7.3-10.5)	1 - interested in acute symptoms. Assessed at time of exposure, then 3 to 4 days later.	1 - 30 years	0 - Mean 3.3 years
Adequacy of follow-up of cohorts	NA	0 - 84% at T2 and 73% at T3. No description provided.	0 - 84% at T2 and 73% at T3. No description provided.	NA - no follow-up other than 3-4 days post use.	1 - 57% assessed at 30 year follow-up. Description of lost provided.	1 - 78% assessed at follow-up. Description of participants lost provided.
Comments on study quality	Low ROB study despite lack of detail on ascertainment of control group. Nicely conducted retrospective study.	Moderate ROB study. Included participants with negative/disorganized symptoms at baseline.	Moderate ROB study. Included participants with negative/disorganized symptoms at baseline.	High ROB study. No information about the source of the exposed or non-exposed sample. Exposed sample used drugs in addition to cannabis, and there was no baseline assessment. No information about dose ascertained.	Low ROB study. Well- conducted, good description of follow-up and loss to follow-up, description of methods, <i>etc</i>	High ROB due to lack of controlling for important confounders, short follow-up

APPENDIX D. PEER REVIEWER COMMENTS AND AUTHOR RESPONSES

Rev #	Comment	Response					
Are t	Are the objectives, scope, and methods for this review clearly described?						
1-7	Yes	Noted.					
9	No - page 4, line 35: please add risk of cannabis use disorder to the list of adverse events in this phrase- "assess the impact of short- and long-term marijuana use on the risk of adverse effects such as pulmonary diseases, cardiovascular diseases, cancer, and psychosis in the general adult population"	This change has been made.					
Are t	here any <u>published</u> or <u>unpublished</u> studies that we	may have overlooked?					
1, 2, 4, 7	No	Noted.					
3	Yes - There is a recently published systematic review of medical marijuana in psychiatric indications (Wilkinson et al., 2016) that wasn't included. This may have been a timing issue. But now that it is published, it should be included - especially since it informs the PTSD literature.	We have added information from this recent systematic review to our report.					
5	 Yes - A couple of studies regarding harms have come out since your February 2016 deadline. Considering that the review is not likely to be formally published much before 2017. I uploaded the pdfs of these papers. One is a new analysis of the Dunedin study showing that cannabis users are more likely to develop periodontal disease. The second one is an epidemiologic study from Sweden that shows an association between early, heavy cannabis use and mortality. 	We added the new Dunedin analysis to the emerging harms section. We had assessed another analysis from the Swedish military conscript study – there was no data on ongoing tobacco or cannabis use after conscription and, since the outcomes were many decades later, the lack of exposure information made the study results very difficult to interpret.					
6	Yes - See Review.	We have reviewed the studies you suggested and included them in our report if they met our inclusion criteria or if they were relevant for background and discussion sections.					
9	Yes - In assessing risk of harm, it would be more appropriate to include studies assessing harm among daily marijuana users (whether or not they have pain or PTSD) than to assess risk of harm amongst pain or PTSD patients who do not use or who occasionally use marijuana.	We broadly included studies with varying levels of use (including heavy use) and in broad patient populations. We have clarified throughout the summary table and manuscript whether the results apply to light or heavy use and we have clearly noted when there is a lack of data on heavy (daily) use.					
Is the	ere any indication of bias in our synthesis of the evi	dence?					
1-5, 7	No	Noted.					
6	Yes - The choice of only including plant-based and not synthetic cannabinoid studies seems biased, given that they have the same molecular structure.	We have rewritten the methods and the KQ1 results section to better clarify the rationale for this decision and we note how the exclusion of synthetic cannabinoid studies would likely not have affected our overall findings (since there were no large, good quality studies of synthetics in the populations of interest for this report).					
9	Yes - There appears to be a bias in favor of state-	We agree. We have added language to the results					



	, ,
 approved retail marijuana products for treatment of pain and PTSD. The executive summary introduction states that the purpose of the paper is to examine health effects of marijuana use because of increased state legalization of marijuana plant products for the indications of pain and PTSD, but the review of the literature conflates studies of plant-based pharmaceutical grade products (i.e. Sativex) with those of retail smoked marijuana and other marijuana products. As written, the two types of cannabis products are conflated in the summaries of the evidence and in the recommendations. The differences between the two types of products need to be clearly explained and then considered separately in all of the analyses. While Sativex is not currently FDA-approved, it is approved as a pharmaceutical in other countries, is manufactured to known standards of purity and potency and is therefore distinct from retail marijuana products. It cannabidiol concentrations vary widely in retail marijuana. The trend toward higher THC and lower cannabidiol in retail marijuana renders studies of lower THC/higher cannabidiol pharmaceuticals and plant products irrelevant or only indirectly relevant to many currently marketed marijuana products. Given these differences, the level of evidence should be appropriately downgraded for "indirectness" when citing studies of cannabinoid pharmaceuticals, as these do not directly address the benefits and harms of smoked marijuana or other retail marijuana products. 	
9 In assessing potential risks, studies of "low to moderate use" are not appropriate for inclusion. When used for medical purposes, the usual pattern is daily consumption. Therefore, in order to evaluate potential risk, only studies that systematically assess for risk among daily users would be relevant to the question of potential harm from medical use. At least one cited study includes cannabis non-users in the denominator when reporting rates of cannabis use disorder among patients with pain, and is therefore implies a much lower risk of cannabis use disorder than would be expected among daily "medical marijuana" users.	With regard to the cannabis use disorder studies, we agree that we did not clearly describe the cited study and the limitations in the overall evidence base. We revised this section to clearly state that there were no studies in cannabis users. We also de-emphasized the cross-sectional data in chronic pain users in the summary of evidence section since these were not studies in a cannabis-using population. With regard to the other harms, we were broadly inclusive in part because clinicians may encounter a broad range of use among patients. We were careful to describe the evidence as being applicable to low levels of use (as with effects on pulmonary function) when appropriate, and added clarification on the lack of data (or even potential for harm in case of pulmonary function) with heavy use.
Additional suggestions or comments.	
1 Excellent review.	See above.
Clarify on page 4 and in methods the reasons for choice in key exposure (e.g., what is typically found at dispensaries, and not synthetic forms that have been	



	systematically reviewed already)	
2	My comments are all fairly minor.	We agree and have made this change.
_		
	1. A brand name, "Sativex," is used many times in	
	tables and intermittently throughout the text. I believe	
	the generic name (nabiximols) should be used instead	
	in all text and tables.	
2	2. Page 6, line 18 (also page 18, line 39): "and an	This language was clarified.
	estimated 6.2%-39% of chronic pain patients are	
	utilizing cannabis in addition to opioid medication for	
	pain management." The denominator is unclear in this	
	sentence. Should it be "among patients on opioid	
	medication for chronic pain, 6-39% also use cannabis"?	
2		We agree and have changed it to "connehic"
2	3. Page 6: The introduction alternates between "marijuana" and "cannabis." Is there any distinction? If	We agree and have changed it to "cannabis" throughout.
	not, I suggest selecting a preferred term and using it	
	consistently for clarity.	
2	4. Page 6, Methods: A brief rationale for the decision	This has been added.
	to exclude synthetic cannabinoids would be helpful.	
2	5. Page 66, last paragraph of discussion: When	We added language from the 1 st recommendation in
	considering implications for pain management, it	the CDC guidelines. We also added references and
	seems appropriate to mention that multiple	language about other evidence based pharmacologic
	pharmacological and nonpharmacological therapies	and non-pharmacologic therapies.
	have stronger evidence for chronic pain than either	
	cannabis or opioids. Given the state of the science on	
	cannabis and the existence of many efficacious	
	medical and complementary therapies for pain, I am	
	aware of no scientific rationale for singling out cannabis as an important "opioid sparing" therapeutic	
	option. (This is a common line of argument for	
	increasing cannabis availability, so I don't fault the	
	authors for mentioning it.) The first recommendation	
	from CDC guidelines on opioid prescribing, as well as	
	treatment guidelines for common conditions such as	
	back pain and arthritis, could be cited here.	
3	Overall this is a very thorough review.	We generally agree, though we have to stick to the
		strength of evidence grading approach we have used
	The risks of psychosis are underestimated and	throughout the report – we did include mention of
	understated. There is a body of evidence that	experimental studies, though they were small and
	exposure to cannabis is associated with a risk for a	had some methodologic flaws. However, we had not
	psychotic disorder. There is an entire special issue of Biological Psychiatry (April 2016) dedicated to	incorporated these into the summary statement – we
	Biological Psychiatry (April, 2016) dedicated to cannabinoids and psychosis. The authors are strongly	have changed this and clarified the extent of evidence. The SOE rating is low because much of the
	urged to review this special issue.	evidence is observational (though not entirely), it is
		difficult to know the magnitude of effect, and there is
	There is robust evidence (unlike what the review	little data specific to chronic pain and PTSD
	states) of direct experimental evidence that	populations – we have clarified this rationale
	cannabinoids at certain doses can induce psychosis-	throughout.
	like effects in healthy individuals and that	
	cannabinoids can exacerbate psychosis in individuals	
	at risk for or with an established, psychotic disorder.	
	Restating the risk of psychosis is important because	
	of the numbers of veterans with SMI who seek out	
	certification for medical marijuana. I see a number of	
	veterans diagnosed with chronic psychotic disorders	



5	who have asked for medical marijuana certification from VA doctors. They go to non-VA providers get a card, start using marijuana and end up in the hospital. While this is anecdotal, stating that the link to psychosis is "low" or "entirely observational" is not without risk. Obviously, compiling all the papers need to generate this review took a lot of effort. Overall, the review	Thank you for the suggestions – we have detailed our responses below and additionally went through the
	seems comprehensive and generally accurate. When fully refined, it will make an important contribution to our knowledge base.	entire report and did an additional round of copyediting.
	There is some sloppiness in the preparation as though the draft did not undergo careful and extensive proofreading before being sent out for review. In certain presentations of various studies there is a lack of needed detail and occasionally lack of rigor in interpretation. Most of the examples of these concerns that I could find are detailed below, but I cannot confirm that this list is exhaustive of all miscues.	
5	Page 5, lines 12-13: The assumption that rates of pulmonary effects or cancer would not be influenced by presence of PTSD or pain seems flawed at least on the basis that individuals with these disorders use tobacco at higher rates than the general population, and tobacco and cannabis might have additive or synergistic effects. In addition, it seems likely that both PTSD and pain might have subtle hormonal or immune system effects that could interact negatively with cannabis use.	We agree that there is some risk in considering studies in broader populations. We did so after considering likely important confounding factors as related to chronic pain or PTSD. We agree tobacco use is an important confounder and levels might be higher in chronic pain or PTSD populations, but the studies that contributed findings all accounted for tobacco use (and usually conducted analyses among never smokers <i>etc</i>) – studies that did not adequately control for tobacco use were downgraded in quality and did not contribute to findings. There are certainly other factors that might theoretically confound findings – we have added to the limitations section this issue (and, in general, this is one of the reasons why bodies of evidence based only on observational data typically start with a lower strength of evidence rating).
5	Page 5, line 45: Change "size" to "sizes."	We have made this change.
5	Page 6, lines 4-13: Given the nature of the uncontrolled studies reviewed, it would probably be better to say that "cannabis is potentially associated with either harmful or neutral effects" rather than is potentially harmful.	We have made this change.
5	Page 8, line 36: Change "is" to "are."	Done
5	Table (Page 9): Calls medication Sativex when text calls it nabiximols. Should use generic name throughout document to be consistent. Acronym ROB should be footnoted to explain it to anyone perusing the table.	Done
5	PTSD: It seems incorrect to say that marijuana is potentially harmful since these studies were observational. Is it likely the marijuana is causing more violence and use of other substances? Possibly, but it seems more intuitively probable that patients who are more violent and certainly who use other	We changed the executive summary paragraph accordingly. There is more detail in the main body of the report, but the strength of evidence related to bias and small number of studies is clearly indicated.



	substances are more likely to use marijuana. It is	
	more credible to say that there is no evidence that it is helpful.	
5	Page 21, lines 54-57: This sentence does not make sense. If they inhaled a 25 mg dose, the per cent THC is irrelevant because the dose would be the same. What is the preparation here? It does not sound like herbal marijuana.	Language regarding the preparation was clarified; it was indeed an herbal preparation obtained from Prairie Plant Systems Inc. (Saskatoon, Sask.). Regarding dose and potency, this is the language that the authors use to describe the potency and dose. The 0% THC was prepared using "ethanolic extraction of cannabinoids" (see Ware 2010 pg. E695). Concentrations/potencies (percent THC) were varied, but were delivered in the same dose (25 mg).
5	Page 22, lines 42-59: This study is very poorly described. The reader needs to know more about the cannabis product used. If the study was observational, how was assignment to condition determined? The word "native" should be "naïve."	Cannabis product described in more detail. Assignment to conditions described in more detail. Native changed to naïve.
5	Page 23, lines 7-21: These studies are also exceedingly poorly described. What were the basic study methodologies?	Designs for both studies were described in more detail.
5	Table 3: How can the Wilkinson study be medium risk of bias? Shouldn't it be high risk of bias? Obviously, the participants self-selected into their groups. We know that people who use marijuana are more likely to use alcohol and vice-versa. Most likely individuals with PTSD and a propensity to violence are more likely impulsive and more likely to use marijuana. The marijuana may not be causing the violence. The p- value given for primary outcome of Johnson study is inconsistent with what the text says.	We used a standard risk of bias tool to evaluate the observational studies (Newcastle-Ottawa Scale), and using this tool classified the study as medium risk of bias (individual item scores are included in the Appendix C PTSD risk of bias table). This particular study adjusted for confounders which contributed to the medium rating. We agree that causation is very difficult to assume here and this is part of what contributes to rating the body of evidence as insufficient.
		Regarding the Johnson et al. paper, we have checked the values and confirmed that those reported in our table correspond to those reported in the paper.
5	Page 26, lines 21-23: Serious adverse events mentioned twice with different ORs.	Thanks – this was a typo and was corrected (last should have been withdrawal due to AE).
5	Page 26, lines 25-26: Information on specific serious adverse events should be provided in more detail. It is hard to see how paranoia or agitation by themselves would meet the FDA definition of serious adverse event unless they resulted in hospital admission.	We believe the section provides the detail we have available, while remaining circumspect about the seriousness of most of the short-term adverse events reported. The definition of serious adverse event is not provided in the Whiting review or its review protocol. We do clarify that many of the side effects were minor and common effects of cannabis. We have rewritten the sentence and taken out the modifier "serious". The definition of serious adverse events includes medical events for which an intervention might be necessary to prevent something like hospitalization – this is obviously somewhat at the discretion of the monitoring board and investigators and we simply report what the review authors reported.
5	Page 26, line 34: add "and" between "pain" and "found."	This change has been made.
5	Page 34, line 13: Change "was" to "were."	This change has been made.



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Page 36, line 8: Describe dose and route of administration of cannabis in this study.	This change has been made.
Page 37, line 42: Delete "who were diagnosed with CUD."	We have left this statement in the text for clarification.
Page 38, lines 4-15: An apparent issue with the Bonn- Miller study described here which may warrant mention is that the Veterans who had CUD and checked into an inpatient unit presumably had to undergo cannabis withdrawal absent any treatment for it. Was it their CUD per se or the withdrawal symptoms (or both) that interfered with their treatment improvement? It would be good to know also if their PTSD severity at treatment entry was equivalent to that of the non-CUD group.	Our summary describes the results and adjustments for confounders, but we have not included a discussion about whether or not withdrawal symptoms vs CUD was responsible for the findings because it is not possible to determine based on the methods.
Page 40, line 7: Change "abuse" to "misuse."	This change has been made.
Page 41, line 4: Remove "is."	This change has been made.
Page 42, line 47: Need route of administration of cannabis oil.	This was not specified in the ClinicalTrials.gov entry; we have clarified this in table.
Page 43, line 42: 0 mg does not make sense.	This is what was reported in the ClinicalTrials.gov entry, but we have clarified (it was a titration up to 250 mg).
Page 52, Table 8: Additional suggestions: All clinical trials of cannabis should obtain blood levels of THC and CBD so that there is some objective measure of how much drug exposure has occurred. Almost all studies done thus far have been quite low dose. Thus, higher doses must be tested. CBD should be much better studied acutely and longitudinally to determine whether it is reinforcing and whether tolerance and withdrawal occur with chronic use.	Thanks, this has been added.
highlighted below Major Issues: 1. One of the larger issues with the report, as written, is the choice to exclude "synthesized, pharmaceutically prepared cannabinoids (e.g., dronabinol, nabilone)." The authors chose to include studies of whole-plant or plant-derived cannabinoid preparations, but synthetic preparations with the same exact molecular structure and delivery method were excluded. There are very few organizations that produce plant-derived cannabinoids (e.g., NIDA, GW Pharmaceuticals), whereas synthetic cannabinoids (e.g., dronabinol, nabilone) are not only more widely available to researchers, but have been produced and used in research for quite some time. Without a clear rationale, which I think would be difficult to make, the choice of excluding synthetics appears to introduce bias particularly as a number of studies on pain and PTSD have utilized synthetic preparations. For example, Jetly et al., 2015	We added rationale in methods section. We also added information to both the chronic pain and PTSD section regarding the findings from recent systematic reviews on synthetics as they relate to our populations of interest. There was an SR published that included PTSD data very recently – while it was published after our search dates ended, we did include a description of the review and the studies relevant to PTSD. We added discussion of the applicability of the synthetic studies to our questions of interest – there was only one trial of nabilone with very few patients – the other studies would not have met inclusion criteria. Regardless, even after considering all the additional studies, the authors of the recent SR came to the same conclusion re: insufficient evidence.
	administration of cannabis in this study. Page 37, line 42: Delete "who were diagnosed with CUD." Page 38, lines 4-15: An apparent issue with the Bonn- Miller study described here which may warrant mention is that the Veterans who had CUD and checked into an inpatient unit presumably had to undergo cannabis withdrawal absent any treatment for it. Was it their CUD per se or the withdrawal symptoms (or both) that interfered with their treatment improvement? It would be good to know also if their PTSD severity at treatment entry was equivalent to that of the non-CUD group. Page 40, line 7: Change "abuse" to "misuse." Page 42, line 47: Need route of administration of cannabis oil. Page 43, line 42: 0 mg does not make sense. Page 52, Table 8: Additional suggestions: All clinical trials of cannabis should obtain blood levels of THC and CBD so that there is some objective measure of how much drug exposure has occurred. Almost all studies done thus far have been quite low dose. Thus, higher doses must be tested. CBD should be much better studied acutely and longitudinally to determine whether it is reinforcing and whether tolerance and withdrawal occur with chronic use. Excellent work! Remaining points to consider are highlighted below Major Issues: 1. One of the larger issues with the report, as written, is the choice to exclude "synthesized, pharmaceutically prepared cannabinoids (e.g., dronabinol, nabilone)." The authors chose to include studies of whole-plant or plant-derived cannabinoid preparations, but synthetic preparations with the same exact molecular structure and delivery method were excluded. There are very few organizations that produce plant-derived cannabinoids (e.g., NIDA, GW Pharmaceuticals), whereas synthetic cannabinoids (e.g., dronabinol, nabilone) are not only more widely available to researchers, but have been produced and used in research for quite some time. Without a clear rationale, which I think would be difficult to make, the choice of exclud

		Evidence-based Synthesis i Togram
	of 47 patients diagnosed with PTSD who received nabilone, and Roitman et al., 2014 conducted an open-label trial of oral THC for PTSD symptoms. While there is currently debate regarding the necessity of using plant-derived versus synthetic cannabinoids in research and treatment, the heart of this debate lies in the importance of secondary cannabinoids and terpenes, which are present in plant-derived products and not in synthetic ones. As it is unlikely that the role of these secondary compounds informed the selection criteria, given that secondary cannabinoids and terpenes are not even reported in the studies discussed in this review, it seems as though it would be difficult to provide a compelling case for this choice.	
6	2. An additional consideration for the section entitled "Emerging Harms" could be the recent proliferation of new methods of cannabinoid delivery and the resulting risks of adverse events. For example, the use of "dabs" appears to be associated with particularly heightened risk of tolerance and withdrawal (e.g., Loflin & Earleywine, 2014), and the use of edibles with a number of more acute consequences (e.g., Hudak et al., 2015; Lamy et al., 2016).	We added this information to the emerging harms section.
6	3. While the authors are correct in stating that the majority of the literature describes the effects of "cannabis" or "marijuana" without a clear definition of the cannabinoid profile of the product tested or used, the authors similarly make broad comments about consequences of "cannabis," where a more nuanced understanding is emerging. For example, the authors discuss a negative consequence of cannabis use as being psychosis. While this is indeed a finding that has been described in-depth within the literature, and even tied to a genetic vulnerability (i.e., catechol-O-methyltransferase), emerging evidence suggests that the association between cannabis and psychosis is specific to THC and that CBD can actually provide anti-psychotic effects (e.g., Leweke et al., 2012). This level of nuance is not currently provided in the review.	We agree. We have added clarification that it is the THC component that is most likely to be associated with psychotic symptoms and we added a statement to the discussion that CBD has actually been studied as an antipsychotic agent.
6	4. Somewhat related to the inclusion and selection of studies for the review, it is puzzling that the Bonn-Miller, Boden, Vujanovic, & Drescher, 2013 study was not included in the list of studies of the effects of cannabis on PTSD symptoms. That study appears to meet inclusion criteria as it was prospective, involved validated measures of PTSD (i.e., PCL), and included a comparison group (CUD diagnosis was compared to those without CUD diagnosis). The sample was adults and there is no indication that they used synthetics. While the study did use data from medical records, so did the administrative study by Wilkinson et al., 2015. This is just confusing.	Although this study included a control group, the controls didn't have CUD, but might have used cannabis; therefore, it did not meet our criteria because we were comparing studies with a non- cannabis using control group.
6	5. On page 7, the authors note that they "did not find any literature comparing rates of CUD among individuals with chronic pain or PTSD to rates in other	Although these studies don't meet inclusion criteria, we have added the 2012 data on prevalence to the background paragraph of our CUD section.



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	populations" While this may be true, a study by Bonn-Miller, Harris, & Trafton (2012) documented the prevalence of PTSD among Veterans with CUD (29.05% in FY12), and a VA fact-sheet by Bonn-Miller & Rousseau utilized VA PERC data to document the percentage of Veterans with PTSD-SUD who had a CUD diagnosis (22.7% in FY14). These data seem to provide information close to what the authors note as being missing from the literature.	
6	Minor Issues: 1. The authors switch between using the terms "cannabis" and "marijuana." The manuscript may flow more nicely if consistent terminology was used throughout. Indeed, the term "cannabis" is generally preferable over "marijuana."	We agree and have made this change.
6	2. p. 7: "found that about 2% if Veterans with non- cancer" should be "found that about 2% of Veterans with non-cancer"	This change has been made.
6	3. p. 46: The description of the study by Eades et al. within the text is not consistent with the table. The table is correct and the text is inaccurate. The text should note that the three groups are "High/Low, High/High, and Low/Low".	This change has been made.
6	4. p. 46: "marijuana use versus no marijuana use in the past 6 months is associated with PTSD symptoms and sleep" should be "marijuana use versus no marijuana use in the past 6 months is associated with differential trajectories of PTSD symptoms over the course of a year."	This change has been made.
6	5. p. 48: Replace "In addition, we obtain lab analysis results of the cannabis donated through the Santa Cruz Veterans Alliance to the Veterans. This includes lab analysis results of percent cannabinoids within each product." with "In addition, all product provided to Veterans by the Santa Cruz Veterans Alliance is tested for cannabinoid content by an independent laboratory."	This change has been made.
6	6. The authors cite one of the two epidemiological studies of cannabis and PTSD (i.e., Kevorkian et al., 2015), but not the earlier study conducted among the NCS-R (i.e., Cougle et al., 2011).	The Cougle et al. study only reports data on cannabis use, not CUD, and therefore is not included in this section.
7	I was primarily interested/knowledgeable of the evidence for its use in PTSD and think that you did an excellent job reviewing that sparse literature and mentioning the fact that there are two current RCTs in progress that will add to the literature. Overall, very nice job and I have no further suggestions.	Noted.
9	Page 6-line 20. "There is low strength evidence that low levels of marijuana smoking do not adversely impact lung function over about 20 years in young adults." Low levels of marijuana smoking are irrelevant to the question of possible harm associated with "medical" that is, frequent/daily use.	We included any data regarding harms from studies that met inclusion criteria. We clearly state that these data apply to low level users and not daily users. We feel that the breadth of evidence will be useful to clinicians who can assess patients' frequency of use and decide whether or not the available data apply to an individual patient. While it is likely that many patients using medical marijuana do so daily, we do not know this to be universally true and there may be



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		substantial proportion of patients who use less frequently. In any case, the lack of information in older or multimorbid populations (which we clearly state) is perhaps an even bigger issue in applying the data in VA clinical settings – again, we attempted to present our broadest look at harms with clarification on generalizability issues.
9	Page 7-line 38. Recommend deleting this sentence: "One large cross-sectional study of Veterans found that about 2% if (sic) Veterans with non-cancer pain had a diagnosis of CUD, and that this increased to 4%" This is irrelevant to the question of the risk of cannabis use disorder among patients using marijuana for chronic pain treatment who would more likely use it multiple times daily. If it is possible to discern from the paper the prevalence of CUD among those with pain who used marijuana to treat pain, that would be worth mentioning. THC concentration would also be important to note, as more potent varieties (10 - 20+%) currently marketed would pose a greater risk for CUD than the more common low potency (3%) of a decade ago.	We have corrected this sentence and provided this information (as well as some additional, new information) on prevalence as part of our background.
9	Page 8-line 30. Ibid. "Light to moderate use" is irrelevant to the question of harm among daily users.	We have clarified that the data does not apply to heavy (daily) users.
9	Page 8 line 34- also needs to include cannabis use disorder among the serious mental health adverse events. Including indirect evidence about the risk of cannabis use disorder among daily users would better inform decision-making than the indirect study of pain patients who have not used marijuana.	We rewrote the sections on CUD to clarify that there was no evidence with which to assess rates of CUD, and we mention cross-sectional data.