BUNDED BY THE LIGHT

MARIJANA

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a knedication by any other hame

AKA - AUNT MARY, BUD, DOOBIE, DOPE, FATTY, GRASS, HASHISH, HERB, HOMEGROWN, LOCO WEED, LOVE WEED, MAGIC SMOKE, MARIHUANA, MARIJUANA, MARY JANE, MAUI-WOWIE, POT, REEFER, ROACH, WHACKY TABACKY, WEED

It runs the spectrum ...

CBD—snake oil or panacea?

By NIKOLAS HARTER • SEP 11, 2018

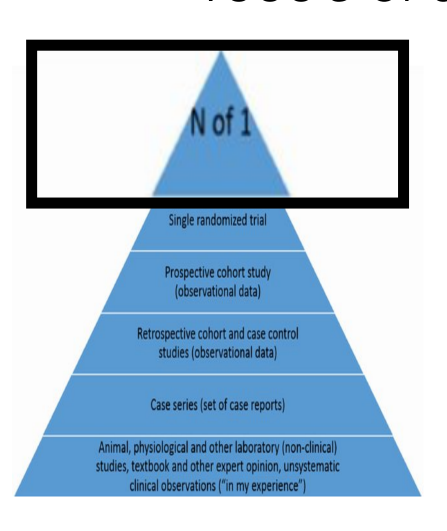








1000's of anecdotal claims



8 Miraculous Medical Marijuana Survival Stories

Meet the eight inspiring souls who say cannabis saved their lives.

Two Anecdotes About My Parents and Marijuana

DISEASES / MEDICAL CONDITIONS

7 Marvelous Medical Marijuana Miracles

Simplified guideline for prescribing medical cannabinoids in primary care

G. Michael Allan MD CCFP Jamil Ramji Danielle Perry Joey Ton Pharmd Nathan P. Beahm Pharmd Nicole Crisp RN MN NP-Adult Beverly Dockrill RN Ruth E. Dubin MD PhD FCFP DCAPM Ted Findlay DO CCFP FCFP Jessica Kirkwood MD CCFP Michael Fleming MD CCFP FCFP Ken Makus MD FRCPC Xiaofu Zhu MD FRCPC Christina Korownyk MD CCFP Michael R. Kolber MD CCFP MSc James McCormack Pharmd Sharon Nickel Guillermina Noël MDes PhD Adrienne J. Lindblad ACPR PharmD

Can Fam Phy 2018;64:111-120

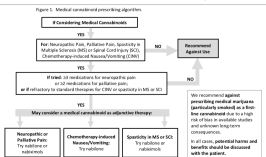
Systematic review of systematic reviews for medical cannabinoids

Pain, nausea and vomiting, spasticity, and harms

G. Michael Allan MD CCFP Caitlin R. Finley MSc Joey Ton PharmD Danielle Perry Jamil Ramji Karyn Crawford MLIS Adrienne J. Lindblad ACPR PharmD Christina Korownyk MD CCFP Michael R. Kolber MD CCFP MSc

Can Fam Phy 2018;64:e78-94

Medical Cannabinoids: Guideline Summary



Percentage of people experiencing harms

	Cannabinoids	Placebo
Sedation	50%	30%
"Feeling high"	35%	3%
Dizziness	32%	11%
Speech disorders	32%	7%
Ataxia/Muscle twitching	30%	11%
Hypotension	25%	11%
Numbness	21%	4%
Psychiatric	17%	5%
Euphoria	15%	2%
Dysphoria	13%	0.3%
Impaired memory	11%	2%
Withdraw due to harms	11%	~3%
Dissociation/Acute	5%	0%

Percentage of people experiencing benefits

Benefits	Cannabinoids	Placebo		
Chronic Pain (≥30% reduction after 4 weeks)				
Neuropathic pain	38%	30%		
Palliative pain	30%	23%		
Chemotherapy-induce	d nausea/vomitir	g (in 1 day		
Control of nausea & vomiting	47%	13%		
Spasticity (≥30% imp	provement after	6 weeks)		
Spasticity	35%	25%		

,			
	Daily Dose ²	Approximate cost/month	
Nabilone*1	2 to 6 mg	\$94 to \$305	
Nabiximols*	4 to 12 sprays	\$226 to \$903	
Medical Marijuana Dried	1 to 3 g typical use	\$250 to \$750 Based on \$8.37/g	

*Manufacturer inst price, ones not reflect pnarmacy depensing rees. "Only generic nablione covered by most provincial drug plans. "Studied doses: Nabilione 0.5mg to 8mg/day, nabisimols 4 to 48 sprays/day, smoked marijuana had THC concentrations ranging 1 to 8% up to three times a day as tolerated. Daily doses from drug monographs and Health Carada.









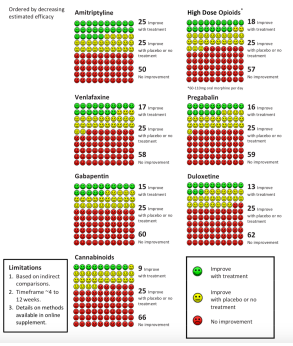






Neuropathic Pain: Pharmacotherapy Treatment

Outcome: Meaningful (~30%) Pain Improvement













Medical



Medical

Cannabinoids

This is an Evidence Synopsis

not a legal discussion

not a moral discussion

not an ethical discussion

simply a discussion of the evidence

Cannabinoids come in many forms

- liquid tinctures
- vaporizing/smoking dried buds
- \chi eating
- capsules
- lozenges
- dermal patches
- oral sprays

Medical marijuana

Manufactured cannabinoids

Nabilone,
Nabiximols (Sativex®)

Medical cannabinoids

Labelling accuracy of CBD

84 products

90%-110% labeled value ~30%

underlabeled >110% labeled value ~45%

overlabeled <90% labeled value) ~25%

THC found in ~20% of samples

JAMA 2017;318:1708-9

Some of the promoted medical uses for Cannabinoids

Tourette Syndrome

Amyotrophic Lateral Sclerosis

Huntington's Disease Parkinson's Disease

Dystonia

Glaucoma

Traumatic Brain Injury/ Intracranial Hemorrhage

Addiction

Anxiety

Depression

Sleep Disorders

Posttraumatic Stress Disorder

Schizophrenia and Other

Psychosis

Osteoarthritis

Fibromyalgia

Neuropathic Pain

HIV Pain

Dementia

Cancer

Chemotherapy-Induced Nausea and Vomiting

Anorexia and Weight Loss

Irritable Bowel Syndrome

Epilepsy

Spasticity Associated with

Multiple Sclerosis or Spinal

Cord Injury

Stolen from Mike Allan

Examples of Poor Research

Glaucoma: 1 RCTs with 6 people (no effect)

Anxiety: 1 RCT of 24 patients tested for simulated public speaking found more improvement on mood visual analogue scale

IBS: 1 RCT of 36 pts given dronabinol for 2.5, 5mg or placebo BID x 2 days: Focused on transit times

JAMA. 2015;313(24):2456-2473. National Academies of Sciences, Engineering, and Medicine. 2017. The health effects of cannabis and cannabinoids: The current state of evidence and recommendations for research. Washington, DC: The National Academies Press. doi: 10.17226/24625.

Stolen from Mike Allan

Two Primary Problems

Blinding: Attempted but rarely tested

In 2 Inhaled cannabis cross-over RCTs

1st: 57% identified all 6 phases

2nd: 90% identified active vs cannabis cigs without THC/CBD

Dronabinol, 95% of patients identified active (as did 85% of nurses) (nabilone study similar)

Inclusion: Previous users often focused on.

Of 6 inhaled RCTs: 3 required past use, 2 no limitation and 1 did not report.

In nausea/vomiting, previous use led to greater response

Naive users (no previous report of psychosis).

Together, these introduce profound bias

Can Fam Physician 2018

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The Evidence base

Lack of high level research

BIAS - many studies enrolled people with a history of cannabinoid use, unblinding was common (90%), small sample size, short duration, inconsistent reporting

Reasonable evidence for

Chronic pain - neuropathic pain and refractory pain in palliative care

Nausea and vomiting - chemotherapy

Spasticity - MS and special cord injury

The Evidence for Cannabinoids

BENEFIT

PAIN

15 RCTs - at least a 30% reduction

RR = 1.37 (95% CI 1.14 to 1.64) - NNT = 11 - 39% vs 30% - primarily nabiximol and some smoked - but larger (>150) and longer (9-15 weeks) trials show no effect

SPASTICITY

4 RCTs - a + global impression of change - 30% reduction

RR = 1.45 (1.08 to 1.95) - NNT = 7 - 50% vs 35%

N and V AFTER CHEMO

7 RCTs - absence of N and V after chemo

RR = 3.60 (2.55 to 5.09) - NNT = 3 - 47% vs 13% - primarily nabilione and delisted dronabinol - vs neuroleptics - NNT = 7 - patient preference higher than effectiveness

Can Fam Phy 2018;64:e78-94

Evidence in Mental Health

The evidence is at best, very poor - overwhelmingly case reports

depression - no RCTs - case reports

anxiety - 1 RCT - CBD powder - 24 patients who performed a simulated public speaking activity and then reported improvement on the mood visual analogue scale Neuropsychopharmacology 2011;36,1219–26

PTSD - 1RCT - nabilone (THC) - 10 patients - found benefit in some outcomes, but these results disagree with other research findings of marijuana use worsening post-traumatic stress disorder Psychoneuroendocrinology 2015;51,585–8

ADHD - 1 RCT - Sativex oralmucosal spray (THC/CBD) - 30 adults - 6 weeks primary outcome (cognitive performance) no difference - some secondary outcomes showed benefit but not after adjustment for multiple testing European Neuropsychopharmacology 2017; 27,795–808

insomnia - THC may have a short-term sleep benefit - but THC is associated with habituation to the sleep-enhancing qualities Curr Psychiatry Rep 2017 19: 23

schizophrenia - 1 RCT - CBD oral solution - 88 patients - an improvement >20% in PANSS total score - no difference - rated by their clinician as "improved" on the CGI-I scale compared with those in the placebo group (79% and 55%) - treatment difference = -0.5, 95% CI = -0.8, -0.1 AJP in Advance doi: 10.1176/appi.ajp.2017.17030325

the present evidence for medical cannabinoids is insufficient to support use in mental health conditions - and it may worsen some conditions (bipolar disorder)

 TABLE 1
 Levels of evidence for cannabis and cannabinoids as treatment

Condition	Compound	Levels of evidence	Highest level
Social anxiety disorder	Cannabis CBD THC Synthetic cannabinoids	3	3
Generalized anxiety disorder	Cannabis CBD THC Synthetic cannabinoids	3 (nabilone)	3
Major depressive disorder	Cannabis CBD THC Synthetic cannabinoids	4 -3 4	4
Bipolar disorder	Cannabis CBD THC Synthetic cannabinoids	4 -4	4
Posttraumatic stress disorder	Cannabis CBD THC Synthetic cannabinoids	3 3 (nabilone)	3
Obsessive-compulsive disorder	Cannabis CBD THC Synthetic cannabinoids	4 (Adj. dronabinol)	4
Trichotillomania	Cannabis CBD THC Synthetic cannabinoids	3	3
Tourette's disorder	Cannabis CBD THC Synthetic cannabinoids	4 3	3

^{1,} Meta-analysis with narrow confidence intervals and/or two or more RCTs with adequate sample size, preferably placebo controlled.

Depress Anxiety 2017;34:1006-17

^{2,} Meta-analysis with wide confidence intervals and/or one or more RCTs with adequate sample size.

³, Small-sample RCTs or nonrandomized, controlled prospective studies or case series, or high-quality retrospective studies.

^{-3,} Negative, small-sample RCTs or nonrandomized, controlled prospective studies or case series, or high-quality retrospective studies.

^{4,} Expert opinion/consensus.

^{-4,} Negative, case report.

"the actual science appears to have been outpaced by the development of applicable legislation and public opinion"

"the documented mental health effects associated with cannabis use also create cause for concern as existing symptomatology typically worsens with use and incidence of certain psychiatric conditions increases"

"there is little observational evidence demonstrating improvement in mental health conditions with cannabis use"

"Clinicians are consequently ill equipped to offer evidence-based advice regarding cannabis' utility as a mental health treatment"

Depress Anxiety 2017;34:1006–17

Critical Review

Cannabidiol as a Novel Candidate Alcohol Use Disorder Pharmacotherapy: A Systematic Review

Jasmine Turna, Sabrina K. Syan, Benicio N. Frey, Brian Rush, Mary Jean Costello ... See all a

First published: 30 January 2019 | https://doi.org/10.1111/acer.13964

"clear limitation to the literature is the paucity of human investigations. Human preclinical and clinical studies are needed to determine whether these positive effects in model systems substantively translate into clinically relevant outcomes."

Intractable seizures

Dravet and Lennox-Gastaut syndrome

CANNABIDIOL

open label - 214 children - 30 seizures a month down to 16 during the 12 week treatment period - severe adverse events 30% - Lancet Neurology 2015

RCT - 120 children ~20-25% greater reduction in seizure frequency - NEJM 2017

RCT - 171 children ~ 20-25% greater reduction in seizure frequency - Lancet 2018

RCT - 225 patients (avg age 15) - 14 weeks - median reduction from baseline in drop-seizure frequency - 41.9% in the 20-mg group, 37.2% in the 10-mg group, and 17.2% in the placebo group - increase of 15-25% somnolence, 10-20% decreased appetite - more in the higher dose - overall 9% in cannabidiol groups had elevated liver aminotransferases - NEJM 2018

Adverse effects - above placebo

~20%-30%

sedation, feeling high, dizziness, speech disorders

~10-20%

ataxia/muscle twitching, low blood pressure, euphoria, dysphoria

~5-10%

impaired memory, withdrawing due to side effects

~5%

dissociation/acute psychosis

Can Fam Phy 2018;64:e78-94

Cannabis-based medicines for chronic neuropathic pain in adults (Review)



Mücke M, Phillips T, Radbruch L, Petzke F, Häuser W

more with 50% or greater pain relief - 21% VS 17%; risk difference (RD) 0.05 (95% confidence interval (CI) 0.00 to 0.09); NNTB 20 (95% CI 11 to 100); 1001 participants, eight studies low quality evidence

more with 30% or greater pain relief - 39% VS 33%; RD 0.09 (95% CI 0.03 to 0.15); NNTB 11 (95% CI 7 to 33); 1586 participants, 10 studies, moderate quality evidence

more withdrew due to adverse events - 10% VS 5%; RD 0.04 (95% CI 0.02 to 0.07); NNTH 25 (95% CI 16 to 50); 1848 participants, 13 studies, moderate-quality evidence

more nervous system adverse events - 61% VS 29%; RD 0.38 (95% CI 0.18 to 0.58); NNTH 3 (95% CI 2 to 6); 1304 participants, nine studies, low-quality evidence

more psychiatric disorders - 17% VS 5%; RD 0.10 (95% CI 0.06 to 0.15); NNTH 10 (95% CI 7 to 16); 1314 participants, nine studies, low-quality evidence

CD012182-2018





Cannabis and cannabinoids for the treatment of people with chronic noncancer pain conditions: a systematic review and meta-analysis of controlled and observational studies

104 studies; 48 - neuropathic pain, 7 - fibromyalgia, 1 - rheumatoid arthritis, 13 - multiple sclerosis—related pain, 6 - visceral pain, 29 - mixed or undefined CNCP

30% reduction in pain - cannabinoids 29.0% placebo 25.9%

all-cause adverse events were cannabinoids 81.2% vs placebo 66.2%

Pain 2018;159:1932-54



Systematic reviews with meta-analysis on cannabis-based medicines for chronic pain: a methodological and political minefield

Winfried Häuser^a, Nanna B. Finnerup^{b,c}, R. Andrew Moore^d

"for any chronic pain condition, the evidence fails to meet EMA and FDA standards, and there are severe risks of addiction, long-term cognitive effects, and risk of psychosis"

"Some patients with chronic pain report substantial symptom improvement with CBM, and they, and politicians driven by the desire of public affirmation, really do not care about EMA and FDA standards."

Pain 2018;159:1906-7



Systematic reviews with meta-analysis on cannabis-based medicines for chronic pain: a methodological and political minefield

Winfried Häuser^a, Nanna B. Finnerup^{b,c}, R. Andrew Moore^d

"There is an uncomfortable parallel here with the situation with opioids, where the short-term demonstration of efficacy in chronic pain led to promotion and broad scale prescribing in the absence of good quality evidence."

"The lessons of history are not to be mocked: we do not want to see a cannabis epidemic replacing that of opioids"

Pain 2018;159:1906-7

Smoked marijuana

The Literature

the literature around smoked medical marijuana demonstrates a considerable risk of bias possibly exaggerated benefits and underreported harms

Long-term harms

(including smoking) and serious adverse effects would not be adequately captured in RCTs and so are largely unknown

Dosing

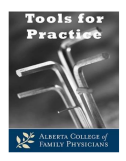
poses an issue, as THC and cannabidiol concentrations vary considerably with differing medical marijuana products

many dried medical marijuana products have THC concentrations of 15% or greater, while the highest concentration studied is only 9.4%

mode of delivery and volume per use can substantially change total intake

there is no evidence that the different formulations of medical marijuana, such as cannabis oil, are more effective or safer than dried medical marijuana

cases of pulmonary aspergillosis have been reported in immunocompromised patients



Blazing Through the Evidence on THC Versus CBD Combinations in Medical Cannabinoids.

Clinical Question: Do tetrahydrocannabinol (THC), cannabidiol (CBD), or the THC/CBD combination yield differing benefits or harms? April 23,2018

4 RCTs

1 - THC/CBD superior to THC for pain control but this was inconsistent within study and with other studies

Adverse events are prevalent in THC/CBD and individual components. While some early poor-quality research in healthy users suggest CBD may attenuate some psychiatric effects of THC, better research in real patients is needed to verify any benefits of specific components.

Is CBD non-psychoactive?

NO - it is psychoactive

Maybe better to say it is non-intoxicating

Depending on doses it does cause sedation, decreased appetite

Epilepsy studies - started at doses of 2.5 mg/kg and increased up to 10-20 mg/kg

THESE ARE NOT DOSING RECOMMENDATIONS!!!

"General recommendation" - 25 mg twice daily and then go up

Also seen

General Health: 2.5-15 mg

Chronic pain: 2.5-20 mg

Sleep disorders: 40-160 mg CBD



Cannabis for Opioid Use Disorder

The push to use cannabis in Opioid Use Disorder is being fueled by the :

Clear problem of the opioid crisis

Ineffectiveness of current treatments safety

Perceived safety of cannabis

It is NOT being fuelled by overwhelming or even marginal evidence

"There is very weak evidence to support the claim that expanding access to medical cannabis will reduce opioid overdose deaths in the United States."

Addiction 2018;113,987-98

Emerging Evidence for Cannabis' Role in Opioid Use Disorder

Beth Wiese^{1,2} and Adrianne R. Wilson-Poe^{2,*}

"The compelling nature of these data and the relative safety profile of cannabis warrant further exploration of cannabis as an adjunct or alternative treatment for OUD"

"Undoubtedly, more high-quality clinical evidence is needed to further support the use of cannabis to combat OUD"

Cannabis and Cannabinoid Research 2018;3:DOI: 10.1089/can.2018.0022

EDITORIAL doi:10.1111/add.14139

It is premature to expand access to medicinal cannabis in hopes of solving the US opioid crisis

"There is very weak evidence to support the claim that expanding access to medical cannabis will reduce opioid overdose deaths"

Issue with cohort studies

Addiction 2018;113:987-8. doi: 10.1111/add.14139.



"the evidence regarding safety, efficacy, and comparative effectiveness is at best equivocal for [chronic pain] and strongly suggests substituting cannabis for opioid addiction treatments is potentially harmful [impaired driving, psychosis, addiction]. Neither recommendation meets the standards of rigor desirable for medical treatment decisions"

JAMA Psychiatry | Original Investigation

Association of Cannabis Use in Adolescence and Risk of Depression, Anxiety, and Suicidality in Young Adulthood A Systematic Review and Meta-analysis

Users vs nonusers - 11 studies

```
depression - 1.37 (95% CI, 1.16-1.62; I_2 = 0\%) anxiety - 1.18 (95% CI, 0.84-1.67; I_2 = 42\%) suicidal ideation - 1.50 (95% CI, 1.11-2.03; I_2 = 0\%) suicidal attempt - 3.46 (95% CI, 1.53-7.84, I_2 = 61.3\%)
```

For all this type of evidence there is the issue of confounders - not all studies adjusted for these

JAMA Psychiatry. doi:10.1001/jamapsychiatry.2018.4500

Neuropathic Pain

Benefit Comparison

Can Fam Physician 2018 Feb.

Figure 2: Neuropathic Pain: Pharmacotherapy Treatment

Outcome: Meaningful (~30%) Pain Improvement



Figure 1. Medical cannabinoid prescribing algorithm. **If Considering Medical Cannabinoids** YES For: Neuropathic Pain, Palliative Pain, Spasticity in Recommend NO Multiple Sclerosis (MS) or Spinal Cord Injury (SCI), **Against Use** Chemotherapy-induced Nausea/Vomiting (CINV) YES If tried: ≥3 medications for neuropathic pain NO or ≥2 medications for palliative pain; or if refractory to standard therapies for CINV or spasticity in MS or SCI We recommend against prescribing medical marijuana YES (particularly smoked) as a firstline cannabinoid due to a high May consider a medical cannabinoid as adjunctive therapy: risk of bias in available studies and unknown long-term consequences. Chemotherapy-induced Neuropathic or Spasticity in MS or SCI: In all cases, potential harms and **Palliative Pain:** Nausea/Vomiting: Try nabilone or benefits should be discussed Try nabilone or Try nabilone nabiximols with the patient. nabiximols

Can Fam Physician 2018 Feb.







Best Available Evidence

ORAL nabilone (THC) for nausea and vomiting - NNT ~3

OROMUCOSAL SPRAY nabiximols (THC/CBD) for spasticity - NNT ~7 and neuropathic pain - NNT ~11

CBD for intractable seizures

The rest - weak evidence at best - certainly lots of anecdotal evidence that shouldn't be ignored - worth a try if other things haven't worked?

5-25% absolute increase in side effects over placebo

Long term benefit and harm - pretty much unknown

SMOKED - there is no evidence that any one formulation is more effective or safer than dried marijuana

Original Article

Epidemiological characteristics, safety and efficacy of medical cannabis in the elderly

2736 patients surveyed - mean age 75 pain - 67%, cancer - 61% 6 months - 94% reported improvement in their condition pain level was reduced from a median of 8 on a scale of 0–10 to a median of 4 adverse events were: dizziness (10%) and dry mouth (7%) 18% stopped using opioid analgesics or reduced their dose

European Journal of Internal Medicine 2018;49:44-50

Drugs & Aging (2019) 36:39–51 https://doi.org/10.1007/s40266-018-0616-5

REVIEW ARTICLE

Medical Cannabis for Older Patients

"the evidence to date for the efficacy of cannabinoids in general and medical cannabis in particular, for many medical conditions and symptoms is scanty. In contrast, there is considerable mounting evidence for harms, many of which are applicable to older individuals"

"Irrespective of the current level of evidence for medical cannabis, buoyed by media and advocacy, medical cannabis is a current reality, and clinicians must take an active role in ensuring competent patient care."

Symptoms

You primarily need to know IF it works

Safety, cost and convenience

Older medications first - safety

Head-to-head studies are uncommon

Doses in the CPS are "wrong"

N-of-1 studies

Let the patient tell you

Symptom NNTs

General anesthesia/local anesthesia - NNT ~1

PPIS, sildenafil - heartburn/"successful" intercourse NNT ~2

NSAIDs, opioids - pain NNT ~3-5

Steroids - sore throat - NNT ~3, Bell's palsy - NNT ~10

Antibiotics - acute COPD exacerbation - NNT ~5

Topical antibiotics - bacterial conjunctivitis - NNT ~7

Antidepressants - severe depression - NNT ~10

Ipratropium - asthma attack - NNT ~11

Cholinesterase inhibitors - ADAS-Cog >4 - NNT ~10

Sleeping pills - improvement in sleep quality - NNT ~13

NNTs taken from Cochrane reviews - if a Cochrane review as N/A then from a published MA

But you need to know what goes on in the placebo group

	If person "responds", what is the % chance it was the medication	
Response in the placebo group	If Benefit 10% - NNT 10	If Benefit 20% - NNT 5
0%	~100%	~100%
20%	~33%	~50%
40%	~20%	~33%

NNTs taken from Cochrane reviews - if a Cochrane review was N/A then from a published MA

The Placebo Group Effect

not the placebo effect and these are ballpark numbers

- ~0% general anesthesia
- ~5% psychosis
- ~10% sildenafil, OCD
- ~20% Alzheimer's meds, acetaminophen for headaches, side effects
- ~25% menopausal symptoms, migraine (frequency/severity)
- ~30% blood pressure goal, depression, anxiety, PTSD, PPIs/H2RA, sore throat, NSAIDs of OA, inhalers for COPD
- ~40% panic disorders

Placebo effect taken from Cochrane reviews - if a Cochrane review was N/A then from a published MA



EDITORIALS

Effectiveness of antidepressants

Lots of useful data but many important questions remain

James McCormack professor¹, Christina Korownyk associate professor²

¹Faculty of Pharmaceutical Sciences, UBC, Vancouver, British Columbia, Canada; ²Department of Family Medicine, University of Alberta, Edmonton, Alberta, Canada

Absolute response rates

placebo - roughly 40%

treatment - roughly 50%

gives clinicians and patients valuable information for decision making

10 patients with moderate to severe depression take an antidepressant for two months

five (50%) will report being "better"

in four of them the response will not be because of the drug

When a medication has "worked", if you were a betting person you would bet that it probably wasn't because the medication worked.