

Unhealthy Alcohol and Drug Use

Adult, Primary Care Clinical Practice Guideline Brief

UNHEALTHY ALCOHOL AND DRUG USE GUIDELINE USE

Full Dartmouth-Hitchcock Unhealthy Alcohol and Drug Use Adult, Primary Care Clinical Practice Guideline http://one.hitchcock.org/intranet/docs/default-source/d-h-knowledge-map-documents/uadu-guideline-final 2017.pdf

Unhealthy Alcohol and Drug Use Adult, Primary Care Clinical Practice Guideline Pocket Guide http://one.hitchcock.org/intranet/docs/default-source/d-h-knowledge-map-documents/sud-pocketguide2 6 17.pdf

Guideline Adoption Statement Source Documents:

MA-SBIRT (Massachusetts Screening, Brief Intervention and Referral to Treatment) Clinician's Toolkit² Helping Patients Who Drink Too Much: A Clinician's Guide¹

Facing Addiction in America: The Surgeon General's Report on Alcohol, Drugs and Health³

D-H Depression Management Guideline

http://one.hitchcock.org/intranet/docs/default-source/d-h-knowledge-map-documents/depression_cpg_final.pdf

D-H Behavioral Health Integration into Primary Care Model Guideline

http://one.hitchcock.org/intranet/docs/default-source/d-h-knowledge-map-documents/behavioral-health-integration-guideline-final.pdf

Definitions:

Substance Use Disorder (SUD): a condition is which use of a substance (including alcohol) leads to clinically significant impairment or distress (see DSM-5 below)

Risky Use: consumption of amounts that increase the likelihood of health_consequences.

Risky Drinking¹:

For healthy men up to age 65:

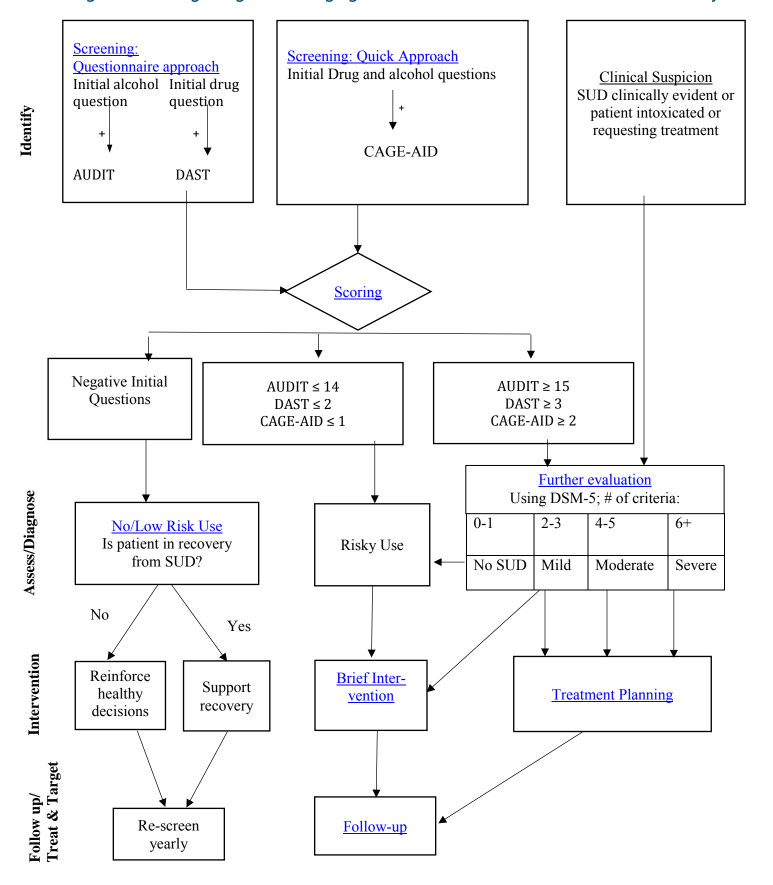
- more than 4 drinks in a day or
- more than 14 drinks in a week

For all healthy women and healthy men over age 65:

- more than 3 drinks in a day or
- more than 7 drinks in a week



Algorithm: Recognizing and Managing Alcohol and Substance Use Disorders in Primary Care



SUMMARY RECOMMENDATIONS

Patient Identification

A SUD may be suspected based on patient presentation (signs of intoxication or medical, social, or occupational consequences of substance use) or a patient requesting treatment. A potential SUD should also be investigated in patients prescribed controlled substances if there are:

- Multiple controlled substances or multiple prescribers, which may be identified thought the Prescription Drug Monitoring System (PDMP)4
- Discordant results on urine toxicology testing⁴
- Taking or craving more drug than prescribed or difficulty controlling use (including requests for early refills or reporting lost medications)4
- High opioid dosage (>50 mg morphine/day or equivalent (33mg oxycodone))⁴
- Depression, Anxiety, PTSD or history of SUD⁴

Because SUDs often go unrecognized, screening is recommended. We recommend 2 strategies (questionnaire and screening approach, below):

Screening: Ouestionnaire Approach (preferred)

Electronic health record based electronic questionnaires are administered yearly, and generate a report that includes patient's score and clinical interpretation. Positive responses to initial questions (see below in Quick Approach) trigger more in-depth (10 question) surveys: AUDIT for alcohol and DAST for other substance use. AUDIT score≥6 or DAST≥1 will trigger best practice advisories in eDH with decision support and resources.

Screening: Quick Approach- an easy to memorize approach when questionnaires aren't used

Ask about alcohol and drug use:

- Do you sometimes drink beer, wine or other alcoholic beverages? If yes, how many times in the past year have you had 5 or more drinks (4 or more for all women or men over 65) in a day? (one drink=12oz beer, 5oz wine or 1.5 oz spirits).
- In the past year have you used an illegal drug or used a prescription medication for non-medical reasons?

If answer is one or more to the first question or "yes" to the second question, use the CAGE-AID (CAGE-Adapted to Include Drugs) to assess for risk of SUD:

$\boldsymbol{\mathcal{O}}$	
	Have you ever felt that you ought to Cut Down on your drinking or drug use?
	Have people Annoyed you by criticizing your drinking or drug use?
	Have you ever felt bad or Guilty about your drinking or drug use?
	Have you ever had a drink or used drugs first thing in the morning (Eye-Opener) to steady your nerves or get rid of a
	hangover?
Sco	ore one point for each positive response.

Scoring of Screening Instruments: 3 outcomes determine next steps (below)

- Negative Initial Questions →No/Low Risk→Reinforce healthy decisions/support recovery
- AUDIT ≤ 14 , DAST ≤ 2 , CAGE-AID $\leq 1 \rightarrow \text{Risky Use} \rightarrow \text{Brief Intervention}$
- AUDIT 15-19, DAST 3-5, CAGE-AID ≥ 2→Further Evaluation
- AUDIT ≥ 20 or DAST $\ge 6 \rightarrow$ moderate to severe SUD probable, further evaluation and treatment

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Further Evaluation: Assessment for a Substance Use Disorder Using DSM-5							
<u>DSM-5</u> defines SUD as meeting 2 or more of the adapted criteria below.							
	Taking substance more or longer than intended						
	Inability to cut down or stop						
	Spending a lot of time getting/using/recovering						
	Cravings and urges						
	Not meeting responsibilities at home, work, school						
	Continued use despite causing problems in relationships						
	Giving up important social, occupational, recreational activities						
	Recurrent use leading to danger						
	Continued use when causing or worsening a physical or psychological problem						
	Tolerance (needing more to get same effect)						

□ Withdrawal symptoms relieved by taking more

Number of criteria: 0-1 2-3 4-5 6+

Interpretation: No SUD Mild SUD Moderate SUD Severe SUD

Interventions

No/Low Risk

For alcohol, advise staying within the healthy drinking limits. Recommend lower limits or abstinence if the patient has a health condition exacerbated by alcohol or takes medications that interact with alcohol. Advise against drinking during pregnancy. *If the patient is in recovery from past SUD:* congratulate them, ask how long they have been in recovery, ask whether they attend support groups or need counseling/support.

Brief Intervention (BI) for Risky Substance Use

A BI is a conversation with a patient using education and motivational interviewing techniques (see p 22-25 MA-SBIRT) to enhance a patient's motivation to change their substance use. For risky use or mild SUD, the goal is to reduce (to healthy drinking limits) or eliminate use (for illicit drugs, or consequences from or inability to cut down on alcohol). The Brief Negotiated Interview approach used in MA-SBIRT is summarized here. The NIAAA Clinician's Guide uses a more directive approach.

BI Steps (see p 9-10 and 20-21 MA-SBIRT and these videos for more detail)

- 1. Understand the patient's views of use
 - Develop discrepancy between patient's goals and values and actual behavior
- 2. Give information/feedback
 - Ask permission to give feedback. Share AUDIT/DAST/CAGE-AID scores and responses.
 - Explore possible connections to health problems.
 - Review healthy drinking limits (if focus is on risky alcohol use and abstinence is not indicated)
 - Use reflective listening and other motivational interviewing techniques
- 3. Enhance motivation
 - Ask about pros and cons of alcohol/drug use
 - Ask readiness and confidence for making changes, using 0-10 scale
- 4. Give advice and negotiate goal
 - Review concerns, summarize pros/cons
 - Ask what the patient is ready to do now
 - Offer clear advice to cut back or eliminate use

Close: Thank patient, offer written or web educational resources, negotiate follow-up

Treatment Planning for Substance Use Disorder

For patients with a SUD, the communication techniques of the BI are used to motivate patients to both eliminate use and seek further help. The treatment plan depends on severity of SUD, comorbidities, social context, resources, clinician judgment and patient preference. Many patients can be managed in primary care with medication assisted treatment (MAT- see below), self-management tools, mutual help groups, counseling, and frequent follow-up with the PCP and other primary care personnel. Consider referral for additional evaluation and treatment by an addiction specialist or program for more complex patients and those not responding to initial approaches. A minority of patients may require medically managed withdrawal (detoxification) from alcohol or opiates.

Treatment Options

- Medication Assisted Treatment for Alcohol
 - o Naltrexone (po or monthly injection)
 - o Acamprosate
 - o Disulfiram
 - o Non FDA approved: gabapentin, baclofen, topirimate
- Medication Assisted Treatment for Opiates

- o Buprenorphine: can be prescribed by PCP with a <u>waiver</u>, or through a program.
- o Methadone: requires a licensed program
- Naltrexone: for highly motivated patients and/or supervised settings, after achieving abstinence. Unlike buprenorphine and methadone, it does not help with withdrawal and craving.
- Patient Self-Management Resources and Mutual Help Groups (AA, NA, Smart Recovery)
- Specialty Treatment
 - o Outpatient: individual and group counseling, family therapy
 - o Intensive Outpatient Programs
 - o Inpatient/Residential

Follow-up/Relapse Prevention

Regular follow-up and a supportive patient-physician relationship can be therapeutic, and brief interventions are more effective when done iteratively over time. Check in about substance use or recovery at each visit. Was the patient able to meet and sustain their goal?

Yes:

- Reinforce and support continued success
- Encourage patient to return if they lapse from their goal
- Address coexisting medical and psychiatric disorders
- For Risky Use: renegotiate drinking goals as indicated, rescreen annually
- For **SUD**: Facilitate involvement in mutual help groups or other counseling, help patients to recognize and cope with relapse precipitants and cravings, coordinate care with specialist if applicable, maintain MAT for a minimum of 3 months of abstinence.

No:

- Acknowledge that change is difficult
- Support positive change and address barriers
- Relate substance use to medical or other problems as appropriate
- Consider engaging significant others
- Renegotiate the goal and plan:
 - o **Risky Use**: consider a 2-week trial of abstinence, reassess the diagnosis (inability to cut down or stop is one of the DSM-5 criteria)
 - o **SUD**: See relapse as a "learning experience" (what was the relapse trigger? How can it be avoided in the future?). Consider more intensive interventions and address coexisting medical and psychiatric disorders

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Pathways & Guidelines: Clinical Practice Guideline and pathways are designed to assist clinicians by providing a framework for the evaluation and treatment of patients. This Clinical Practice Guideline outlines the preferred approach for most patients. It is not intended to replace a clinician's judgment or to establish a protocol for all patients. It is understood that some patients will not fit the clinical condition contemplated by a guideline and that a guideline will rarely establish the only appropriate approach to a problem

Appendix 1: Medication Assisted Treatment for Alcohol (FDA approved)

(see NIAAA Guide p13-23 for details)

	Naltrexone (Depade®, ReVia®)	Extended Release Injectable Nal- trexone (Vivitrol®)	Acamprosate (Campral®)	Disulfiram (Antabuse®)
Action	Blocks opioid receptors, resulting in reduced craving and reduced reward in response to drinking.	Same as oral naltrexone, 30–day duration.	Affects glutamate and GABA neurotransmitter systems, but its alcohol-related action is unclear.	Inhibits intermediate metabolism of alcohol causing a buildup of acetaldehyde and reaction of flushing, sweating, nausea, and tachycardia if a patient drinks alcohol.
Contraindications	Currently using opioids or in acute opioid withdrawal; anticipated need for opioid analgesics; acute hepatitis or liver failure.	Same as oral naltrexone, plus inade- quate muscle mass for deep intramuscu- lar injection; rash or infection at the inspection site.	Severe renal impairment (CrCl ≤ 30 mL/min).	Concomitant use of alcohol or alcohol-containing preparations or metronidazole; coronary artery disease; severe myocardial disease; hypersensitivity to rubber (thiuram) derivatives.
Precautions	Other hepatic disease; renal impairment; history of suicide attempts or depression. If opioid analgesia is needed, larger doses may be required and respiratory depression may be deeper and more prolonged. Pregnancy Category C. Advise patients to carry a wallet card to alert medical personnel in the event of an emergency. For wallet card information, see www.niaaa.nih.gov/guide .	Same as oral naltrexone, plus hemophilia or other bleeding problems.	Moderate renal impairment (dose adjustment for CrCl between 30 and 50mL/min); depression or suicidal idea- tion and behavior. Pregnancy Category C.	Hepatic cirrhosis or insufficiency; cerebrovascular disease or cerebral damage; psychoses (current or history); diabetes mellitus; epilepsy; hyperthyroidism; renal impairment. Pregnancy Category C. Advise patients to carry a wallet card to alert medical personnel in the event of an emergency. For wallet card information, see www.niaaa.nih.gov/guide .
Serious adverse reactions	Will precipitate severe withdrawal if the patient is dependent on opioids; hepatotoxicity (although does not appear to be a hepatotoxin at the recommended doses).	Same as oral naltrexone, plus infection at the injection site; depression; and rare events including allergic pneumonia and suicidal ideation and behavior.	Rare events include suicidal ideation and behavior.	Disulfiram-alcohol reaction, hepatotoxicity, optic neuritis, peripheral neuropathy, psychotic reactions.
Common side effects	Nausea, vomiting, decreased appetite, headache, dizziness, fatigue, somnolence, anxiety.	Same as oral naltrexone, plus a reaction at the injection site; joint pain; muscle aches or cramps.	Diarrhea, somnolence.	Metallic after-taste, dermatitis, transient mild drowsiness.
Examples of drug interactions	Opioid medications (blocks action).	Same as oral naltrexone.	No clinically relevant interactions known.	Anticoagulants such as warfarin; isoniazid; metronidazole; phenytoin; any nonprescription drug containing alcohol.
Usual adult dosage	Oral dose: 50 mg daily. Before prescribing: Patients must be opioid-free for a minimum of 7 to 10 days before starting. If you feel that there's a risk of precipitating an opioid withdrawal reaction, administer a naloxone challenge test. Evaluation liver function. Laboratory followup: Monitor liver function.	IM dose: 380 mg given as a deep intramuscular gluteal injection, once monthly. Before prescribing: Same as oral naltrexone, plus examine the injection site for adequate muscle mass and skin condition. Laboratory followup: Monitor liver function.	Oral dose: 666 mg (two 333-mg tablets) three times daily; or for patients with moderate renal impairment (CrCl 30 to 50 mL/min), reduce to 333 mg (one tablet) three times daily. Before prescribing: Evaluate renal function. Establish abstinence.	Oral dose: 250 mg daily (range 125 mg to 500 mg). Before prescribing: Evaluate liver function. Warn the patient (1) not to take disulfiram for at least 12 hours after drinking and that a disulfiram—alcohol reaction can occur up to 2 weeks after the last dose and (2) to avoid alcohol in the diet (e.g., sauces and vinegars), over the counter medications (e.g., cough syrups), and toiletries (e.g., cologne, mouthwash). Laboratory followup: Monitor liver function.

Appendix 2: Medication Assisted Treatment for Alcohol (off label/not FDA approved)

- Baclofen: GABA agonist, decreases withdrawal symptoms, decreases alcohol consumption and craving, and may help anxietv.6
 - o 10-20mg tid, potentially higher (up to 300mg/day is being studied)
 - Side effects: sedation, respiratory depression, psychiatric disturbance
 - Safe in liver disease
 - o Felt not to have any abuse potential
 - o Least expensive of the options (Walmart \$4 list)
- Gabapentin: structurally similar to GABA, but doesn't bind to receptor (mechanism unclear). Decreases withdrawal symptoms⁷, improves insomnia, dysphoria and craving, and reduces alcohol consumption.⁸
 - 300 tid \rightarrow 600 tid (600 more effective)
 - o Side effects: sedation and dizziness, but dangerous side effects unlikely
 - Safe in liver disease
 - o Some concern for abuse potential
- Topiramate: modulates GABA and glutamatergic activity, decreases craving and alcohol consumption. 9,10 Better than naltrexone in one RCT.11
 - o 25 gd, increase by 25mg/week to 100-150mg BID (or 200-300 once daily ER)
 - Side effects: metabolic acidosis, renal stones, psychomotor slowing, somnolence

Appendix 3: Patient Self-Management Resources and Mutual Support Groups

Note: these are being vetted by a patient group, and will be revised

- Alcoholics Anonymous: www.aa.org
- Narcotics Anonymous: http://na.org/
- Self-Management Addiction Recovery Program: www.smartrecoverv.org
- Digital device Applications (Apps)
 - o A-CHESS: developed by researchers at the University of Wisconsin as a continuing relapse prevention support for those in recovery from alcoholism and alcohol use disorders, after they leave treatment
 - o recoveryBox- for a range of addictions
 - Sober Grid
 - o 12 Steps AA Companion

Appendix 4: Treatment Options

- http://www.dartmouth-hitchcock.org/alcohol drug.html DH guide to options in NH and VT, arranged by treatment intensity
- http://nhtreatment.org/ Recent comprehensive guide to options in NH
- https://findtreatment.samhsa.gov/ National directory
- www.addictionrecoveryguide.org National directory, also with other information and resources for the patient and their family
- www.uvmentalhealth.org Upper Valley resources for all behavioral health issues
- www.uvalltogether.org , includes a "Consumer's Guide to Substance Use Treatment" Upper Valley resources, including pdf booklet (print copies available to stock in clinic
- Substance related crises 1-844-711-4357 (HELP)
- 211 http://www.vermont211.org/ http://www.211nh.org/
- Decisions in Recovery: Treatment for Opioid Use Disorder. A patient handbook discussing MAT options http://store.samhsa.gov/shin/content/SMA16-4993/SMA16-4993.pdf