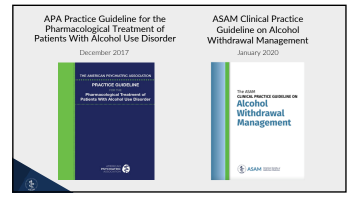
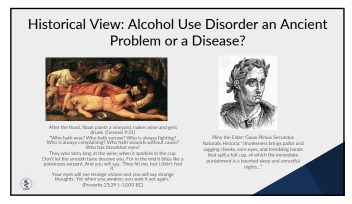




Outline

- 1. Historical View
- Neurobiology
 Epidemiology
- 4. SBIRT and Clinical Screening Test
- 5. Diagnosis
- 6. Biomarkers
- 7. Phases of Alcohol Treatment and Related Syndromes
- 8. CIWA-Ar and Management
- 9. Relapse Prevention Pharmacotherpy and Psychotherapy
- 10. New Directions
- 11. Conclusion





5

Case: RR Mr. RR is a 58 -year-old, Latino, married, male owner of a music theater in Los Angeles. He is being referred for evaluation to assess his drinking and depression after his older brother, who in the past had problems with alcohol, recommended him.



Case: RR

He presents for his evaluation thinking alcohol helps him to manage:

- Depression
- Insomnia
- Irritability and anxiety



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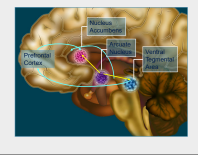
Case: RR

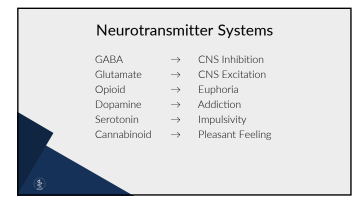
SA history: He reports that he grew up drinking. His first drink was at age four when he tasted the left-over alcohol from a party in his family home. He describes falling in love with the taste of wine and waited every weekend for his family to throw another party.

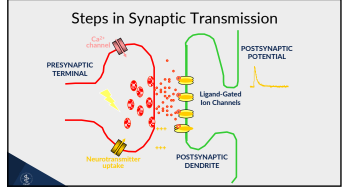


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Alcohol Use Disorder a Disease?



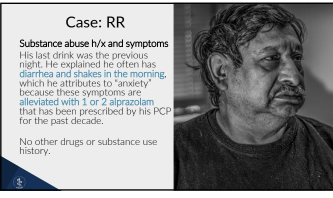


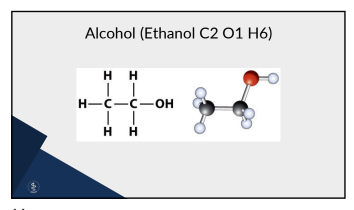


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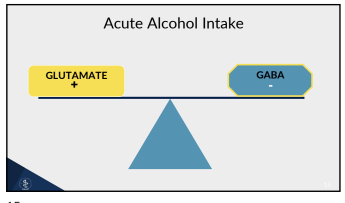


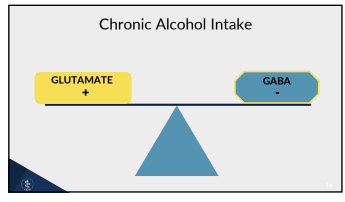


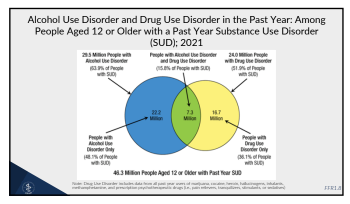




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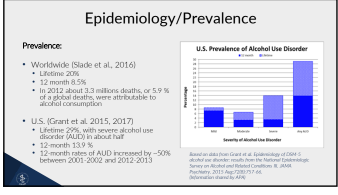


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Epidemiology Scope of Alcohol-Related Problems - ~140,000 people die (380 per day) annually from alcoholrelated causes in the U.S from 2015-2019 Nearly 29.5 million people ages 12 and older had AUD in 2021 894,000 adolescents ages 12 to 17 with AUD in 2021 4th leading preventable cause of death in U.S. is AUD

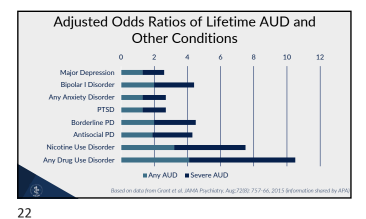
Cost and Scope of Alcohol-Related Problems - 50% of U.S. liver disease deaths attributable to alcohol misuse (2021) - 100 200 300 400 - 100 200 300 - 100 2

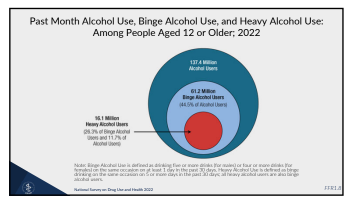
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Epidemiology/Demographics AUD affects individuals of all demographic groups Onset: 18-29 years Ethnicity (12-month prevalence): American Indian/Alaska Native 19.2% African American 14.4% White 14% Hispanic 13.6% Asian-American/Pacific Islander 10.6% Gender (12-month prevalence): Men 17.6% Women 10.4%





How Much is "too much"? **Emerging Trend-**Heavy Drinking High Intensity Binge Drinking **Drinking** A pattern of drinking that brings blood alcohol concentration (BAC) levels to Consuming ETOH at levels that are two or WOMEN: 4 or more standard drinks in a sitting. (8 or more per week.) more times the gender-0.08g/dl specific binge drinking thresholds • WOMEN: 5 or more standard drinks in a sitting. 4 or more drinks on same occasion in about 2 hours 10 or more standard drinks (or alcoholic drink (15 or more per week.) equivalents) for males and 8 or more for 5 or more drinks in same occasion in about 2 hours

COVID and Alcohol Use Disorder

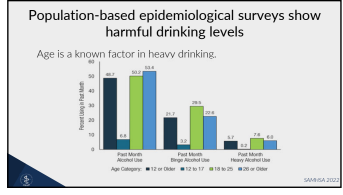
- Data from a national survey of U.S. adults on their drinking habits found that excessive drinking (such as binge drinking) increased by 21% during the COVID-19 pandemic.
- More than a dozen studies have found that 20% to 40% of individuals surveyed reported consuming more alcohol than usual during the pandemic, based on National Institute on Alcohol Abuse and Alcoholism (NIAAA) information

NIAAA National Institute on Alcohol Abuse and Alcoholism 202

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Alcohol use is increasing more in women than men in USA Nonthly Alcohol Use Percentage of U.S men and women who reported drinking alcohol in the past month Monthly Alcohol Use Percentage of U.S. men and women who reported drinking alcohol in the past month Monthly Alcohol Use Percentage of U.S. men and women who reported drinking, having alcohol use disorder, drunk driving and self reported consequences In the last decade differences narrowed further. Rates of alcohol use disorder (AUD) have increased in women by 84% over the past ten years relative to a 35% increase in men (Grant et al., 2017), Women are more likely to experience blackouts, liver inflammation, brain atrophy cognitive deficits and some cancers. (Slade T et al. BMJ 2016)

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DSM-5 : Criteria for Alcohol Use Disorders 1. Use In Larger Amounts / Longer Periods Than Intended 2. Unsuccessful Efforts To Cut Down 3. Excessive Time Spent Taking Drug 4. Failure To Fulfill Major Obligations 5. Continued Use Despite Knowledge Of Problems 6. Important Activities Given Up 7. Recurrent Use In Physically Hazardous Situations 8. Continued Use Despite Social Or Interpersonal Problems 9. Tolerance 10. Withdrawal 11. Craving

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Underdiagnoses and Unmet Treatment Needs

- Only 1 in 6 US adults report ever having asked by a clinician about their drinking behavior
- Despite high prevalence, societal cost, and available treatments, AUD remains undertreated
- <1 in 10 with a 12-month AUD diagnosis receive any treatment:
- Self-help groups
- Psychotherapy
- Pharmacological treatments
- Treatment received by patients varies based on geography, insurance coverage, and formulary restrictions

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Intoxication Features 1 drink \rightarrow BAC = ~15 mg% (0.015 g/dl) 0-100 mg/dl 100-200 mg/dl Incoordination 200-300 mg/dl 300-400 mg/dl Stage 1 Anesthesia, amnesia, hypothermia 400-600 mg/dl 600-800 mg/dl Death

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The Rules of Twenties

Going Up

- MEN: Each drink adds 20 mg/dL to one's BAL.
- WOMEN: Each drink adds 40 mg/dL to one's BAL.

Coming Down

• We metabolize 20 mg/dL every 60-90 minutes (zero order kinetics).

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Women and Pregnancy

- There are three general reasons that females show higher BACs (and greater intoxication) than males if they drink the same amount of alcohol.
- Body composition: In females a greater percentage of body mass is fat compared to males
 Result The concentration of alcohol is increased in the female bloodstream compared to the male body
- Stomach alcohol dehydrogenase (ADH): Females have very little of this enzyme compared to males
- Result Females do not metabolize alcohol before it gets out of the stomach. Therefore, the blood alcohol concentration (BAC) is higher for females versus males
- Liver ADH: Females have a less active form of this enzyme than males.
 Result Females do not metabolize alcohol as efficiently as males, thereby increasing the BA

Women and Pregnancy

Fetal Alcohol Spectrum disorders (FASD): Growth retardation, Facial malformations, Small head, Greatly reduce intelligence.

- FASD is the most common known preventable cause of mental impairment.
- The prevalence of FASD: 50 per 1,000 (May et al., 2009 and CDC
- 40,000 infants per year in US

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Case: RR

Past Medical h/x: HTN for 10 years, GERD and H/x of pancreatitis.

Medications:

- Lisinopril 40 mg qam,
- Omeprazole 20 mg daily
- Zolpidem XR 6.25 mg qhs prn for insomnia
- Alprazolam 1-2 mg tid a day for anxiety.



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Case: RR

Vital Signs: BP:150/95 Pulse: 90x'

CBC normal with the exception of Increased MCV equal 102 (80-96) Electrolytes and renal function: normal

- Hepatic function:
 GGT 141 (10-42),
 AST 60 (15-40)
 ALT 40 (10-40)

 - AST/ALT ratio 1.5
- CDT score exceeded the cutoff and so you performed a diagnostic evaluation





Preventing and Treating AUD

There are evidence-based interventions for preventing and treating AUD:

- Screening, Brief Intervention, and Referral to Treatment (SBIRT)
- Professionally-led behavioral interventions
- FDA-approved medications
- Mutual support groups, such as Alcoholics Anonymous

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SBIRT

- Screening quickly assesses the severity of substance use and identifies the appropriate level of treatment.
- Brief intervention focuses on increasing insight and awareness regarding substance use and motivation toward behavioral change.
- Referral to Treatment provides those identified as needing more extensive treatment with access to specialty care.

www.niaaa.nih.gov/guid http://www.sbirtcolorado.org/healthcare_videosandwebcasts.ph

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Screening Tools

Alcohol Screening is an Effective Prevention Strategy

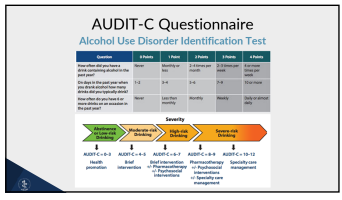
The CAGE Questionnaire

- Cut Down
- Annoyed
- Guilty
- Eye-Opener

2 or more positive responses are strongly associated with alcohol dependence.

National Institu

National Institute on Alcohol Abuse and Alcoholism (NIAAA): "Helping Patients Who Drink Too Much



The Role of Biomarkers in The Treatment of

- Provide objective outcome measures in alcohol research or evaluating an alcohol treatment program.
- Screen for individuals unable/unwilling to accurately report drinking behavior (e.g., fear, embarrassment, or adverse consequences).
- Evidence of abstinence in individuals prohibited from drinking.
- Enhance patient motivation to stop/reduce drinking.
- Diagnosis tool by assessing contribution of alcohol to the disease.
- Identify relapse early.
- · Fear of detection by biomarkers may dissuade drinking.

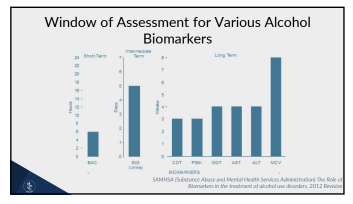
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Types of ETOH Biomarkers

- Manifestations of organ damage often due to drinking
- gamma glutamyltransferase (GGT)
 aspartate amino transferase (AST, SGOT)
- alanine amino transferase (ALT, SGPT)
 macrocytic volume (MCV)
- · Reflections of alcohol's effects on other metabolic processes -
- carbohydrate-deficient transferrin (CDT) Only FDA Approved alcohol biomarker

Direct Tests

- Reflections of alcohol use
 - ethyl glucuronide (EtG) and ethyl Sulfate (EtS)
 Phosphatidylethanol (PEth)



Characteristics of Assessment for Various Alcohol Biomarkers							
Marker	Time to Return to Normal with Abstinence	Level of Drinking	Comments	Blood test normal range			
GGT	2-4 weeks of abstinence	~ 5 drinks (>60g/day) for several weeks	Many sources of false positives—liver disease, diabetes, smoking, obesity, age, anticonvulsants, etc.	W: 0-45 U/L M: 0-53 U/L			
SGOT/AST	2-4 weeks of abstinence	Unknown but heavy	Many sources of false positives (see GGT) in addition to excessive coffee consumption	10 - 34 U/L			
SGPT/ALT	2-4 weeks of abstinence	Unknown but heavy	Many sources of false positives (see GGT) Less sensitive than AST	8-37 U/L			
MCV	Up to several months	Unknown but heavy	Slow return to normal limits even with abstinence renders it a poor independent indicator of relapse. More specific than GGT. Unlike other markers, no strong gender effect	80-100fL			
CDT	2-4 weeks	~ 5 drinks(>60g/day) for 2 weeks	Few sources of false positives. Good marker of relapse	<60 mg/L			

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Diagnostic S	ensitivity and Biomarkers	Specificity of	
	Sensitivity (%)	Specificity (%)	
CDT	69	92	
CDT/transferrin	65	93	
GGT	73	75	
AST	50	82	
ALT	35	86	
MCV	52	85	
(4)	Bell, et al	. Alcoholism: Clinical and Experiment	al Research 1994

Case: RR

His last drink was the previous night. He explained he often has insomnia, diarrhea, palpitations, and shakes in the morning, which he attributes to "anxiety" because these symptoms are alleviated with 1 or 2 alprazolam that has been prescribed by his PCP for the past decade.



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Phases of Alcoholism Treatment

Detoxification

- Primary goal is to achieve an alcohol-free state
- Wide spectrum of severity
- Drug-specific syndromes: opiates, cocaine, alcohol, benzodiazepines

Relapse Prevention

- Primary goal is to maintain an alcohol-free state
- Chronic Treatment

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Introduction Alcohol Withdrawal

Epidemiology

Neurobiology

- Neurotoxicity
 Kindling

Management of Alcohol Withdrawal

- Benzodiazepines
- Anticonvulsants

Real World Implications

- Outpatient vs. InpatientEvaluation and Management

Epidemiology of Alcohol Withdrawal

- Not well studied
- Significant symptoms occur in 13% to 71% of individuals presenting for detoxification
- Up to 10% of individuals undergoing alcohol withdrawal require inpatient medical treatment
- Estimated mortality up to 2%

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Saitz R, Mayo-Smith MF, Roberts MS, Redmond HA, Bernard, DR, Calkins DR. JAM 1994:272:519-52:

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Alcohol Withdrawal and Kindling

- Repeated episodes of alcohol withdrawal likely to worsen
- Exacerbation of symptoms may be due to a kindling process
- Positive relationship of alcohol withdrawal seizures to repeated detoxification

B

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Managing Alcohol Withdrawal

Principles of treatment

- Alleviate symptoms
- Prevent progression of symptoms
- Treat underlying comorbidities



Alcohol Withdrawal Treatment

- Substitute cross-dependent drug (benzodiazepine)
- · Gradually withdraw substitute drug
- Supplement vitamins and minerals
- Thiamine
- Multivitamin
- An array of acid-base disorders and electrolyte disorders can occur in patients with chronic alcohol-use disorder, irrespective of their social circumstances.
- Supportive treatment
 Decrease stimulation, increase fluid and caloric intake

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Alcohol Withdrawal Treatment

Thiamine Deficiency

Thiamine

- Important cofactor for several enzymatic reactions
 Cerebral glucose utilization
 Glutamate elimination

- Wernicke's Encephalopathy
 Partial to complete paralysis of extra ocular muscles
 Nystagmus

 - Ataxia Mental disturbances
- Mortality: 10-20% if untreated
 Treatment: Thiamine replacement PRIOR dextrose administration

Korsakoff's Psychosis

- Antegrade amnesia
 Confabulations

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States of AWS

- 1. Autonomic Hyperactivity
- 2. Hallucinations
- 3. Neuronal excitation
- 4. Delirium Tremens

There is not necessarily a linear progression.



States of AWS

Autonomic Hyperactivity

- Clear Sensorium
- Tremulous
- Diaphoresis
- Anxiety
- Nausea/Vomiting
- Increase cathecolamines in urine, serum and CSF
- Start 6 hrs after last drink Peak 24-48 hrs

Hallucinations

• Most common= VISUAL

Neuronal excitation

- Seizures (Generalized Tonic Clonic)
- Up to 10%
- Most common in first 24 48 hours after last drink

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States of AWS

Delirium Tremens (DTs)

- Most often occur within 72 hours after the last drink
- Delirium with Tremor
- · Autonomic hyperactivity
- Hallucinations
- Electrolyte abnormalities
- Dehydration
- Hemodynamic instability
- Mortality up to 15%
- Cardiovascular/respiratory collapse

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CIWA-Ar

Clinical Institute Withdrawal Assessment of Alcohol, Revised

- It requires under two minutes to administer
- It requires no medical knowledge
- It provides you with a quantitative score that predicts the severity of withdrawal from alcohol

Symptoms	Range of Scores
Nausea and Vomiting	0 (no nausea, no vomiting) -7 (constant nausea and/or vomiting
Tremor	0 (no tremor) - 7 (severe tremors, even with arms not extended
Paroxysmal sweats	0 (no sweat visible) - 7 (drenching sweats)
Anxiety	0 (no anxiety, at ease) - 7 (acute panic states)
Agitation	0 (normal activity) - 7 (constantly trashes about and pacing)
Tactile disturbances	0 (none) - 7 (continuous hallucinations)
Auditory disturbances	0 (not present) - 7 (continuous hallucinations)
Visual disturbances	0 (not present) - 7 (continuous hallucinations)
Headache	0 (not present) - 7 (extremely severe)
Orientation/clouding of sensorium	O (orientated, can do serial additions) - 4 (Disorientated for plac and/or person)

CIWA-Ar Determining Need of Pharmacotherapy • <8: Minimal - Mild AW, Drug therapy not necessarily indicated • 8-15: Moderate AW, Drug therapy indicated. • >15: Severe, Drug therapy absolutely indicated, consider inpatient treatment

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Mechanisms Underlying Alcohol Withdrawal • Multiple neuroadaptive changes in CNS • Decreased GABA activity • Increased glutamate activity • Upregulated calcium channel activity

- Increased noradrenergic activity
- Alcohol withdrawal is associated with increased CNS activity CNS=central nervous system; GABA=gamma-aminobutyric acid.

Anton RF, Becker HC, eds. Pharmacotherapy and pathophysiology of alcol withdrawal. (Handbook of Experimental Pharmacology.) 199

Case: RR

You apply your knowledge and training through Motivational Interviewing. Your open-ended questions and affirmations reviewed with patient's possibilities set the bases for a good rapport with Mr. RR. As part of the treatment dialogue, you showed Mr. RR. his BP elevation 150/90, CIWA:8, and his scores on the CDT, GGT and AST/ALT. You noted that the values were outside the reference ranges for the tests.



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Case: RR

You then explained, in a direct, yet empathetic manner, the significance of the scores and noted that GGT and AST/ALT levels this high can reflect liver damage and that CDT levels this high usually reflect heavy drinking. Mr. RR then agrees to start an outpatient alcohol treatment program.



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Treatment Plan

There are several evidence-based options for nonpharmacological treatment that have minimal harms:

- Motivational Enhancement Therapy (MET): manualized psychotherapy based on the principles of motivational interviewing; shown to have a small to medium effect size on achieving abstinence
- Cognitive Behavioral Therapy (CBT): focusing on the relationships between thoughts, feelings, and behaviors; help manage urges and triggers

Treatment Plan

There are several evidence-based options for nonpharmacological treatment that have minimal harms:

- Medical Management (MM): manualized treatment that provides education and strategies to support abstinence and promote medication adherence
- Community based peer support groups such as Alcoholics Anonymous (AA) and other 12-step programs: helpful in achieving long-term remission but not for replacing formal medical treatment

3

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Alcohol Detoxification Use of Benzodiazepines

- First line agent (gold standard)
- Loss of inhibition/sedation due to lack of ETOH
- Treatment: Replace the GABA activation (inhibition)
- Benzodiazepines:
 - If hepatic impairment: oxazepam or lorazepam
 - Provide dosing for 24 hour intervals patient must be re-evaluated before more is provided
 - Vital Signs
 - CIWA-Ar

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Chlordiazepoxide • Only available in oral form (PO) • Longer half life than most benzos (5-30 hrs) Diazepam • Lipophilic rapid onset of action Lorazapem • Available in oral form (PO) and IV • Half life (12-18 hrs) • Simple metabolism of hepatic glucuronidation (no active metabolite) • Ideal for patients with cirrhosis/liver damage and elderly population

Indications for Outpatient withdrawal treatment

- CIWA <8 or some with CIWA 8 -15
- No hx. of AW seizures/delirium
- No serious medical/surgical problems
- No serious psychiatric/drug hx
- Social support
- Supervision/housing available

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Indications for inpatient withdrawal treatment

- History of DTs or withdrawal seizures
- Alcohol withdrawal severity (CIWA>10) + other criteria (e.g Abnormal lab results, Utox + for other substances)
- Pregnancy
- Major medical/surgical problems
- Inability to tolerate oral medication
- Imminent risk to harm himself and/or others
- Active psychosis or cognitive impairment
- Recurrent unsuccessful attempts at ambulatory detoxification

Muncie HL Jr, Yasinian Y, Oge' L. Am Fam Physician. 2013 Nov 1;88(9):589-9

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Treatment of Mild-Moderate Alcohol Withdrawal CIWA-Ar- 8 to 14

Long-acting Benzodiazepines:

- \bullet Chlordiazepoxide (Librium) 50-100 Mg Po Q 6-8 Hrs.
- Diazepam (Valium) 10-20 Mg Po Q 6-8 Hrs.

Short-acting Benzodiazepines:

 \bullet Lorazepam (Ativan) 2-4 Mg Po Q 1-4 Hrs.

Treatment of Severe Alcohol Withdrawal CIWA-Ar > 15

Diazepam 10 mg IV

• Repeat 5 mg IV q 5 Min Until Calm

Lorazepam 4 mg po q 1 hr, PRN

- Moderate To Severe Liver Disease
- Elderly Or Confused Patients
- Very III Or Debilitated Patients
- Can Be Given PO, IV Or M

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Alcohol Detoxification

Use of Anticonvulsants

Anticonvulsants Reduce Gaba Activity

- CBZ: Reduced rebound withdrawal & post-detox drinking (Malcolm, 2002)
- Gabapentin normalizes alcohol-induced effects on GABA and glutamate; has no hepatic metabolism
- Gabapentin more effective than lorazepam in reducing post-detox drinking (Myrick, 2009)
- Gabapentin, divalproex & vigabatrin may prove useful
- Caution: CBZ & divalproex have limited use in patients with severe hepatic or hematologic disease

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Alcohol Detoxification
Anticonvulsants Effectiveness and Limitations

Advantages

I No abuse liability
Cognition
Neuroprotective
Protracted Withdrawal

Disadvantages
Limited clinical experience
Hematological side effects
Liver toxicity

When to Consider Pharmacotherapy

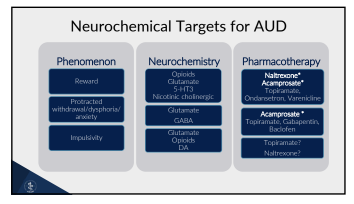
- Anti-craving Medication as the new standard of care
- Consider immediately post-detoxification for ALL patients with alcohol use disorder
- Efficacy requires counseling and/or frequent physician monitoring
- Most FDA approved medications for SUDs can be used in outpatient settings
- Exception: Methadone maintenance therapy: can only be used for treatment of opioid addiction in licensed opioid treatment programs

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	Pharmacogenetics in AUD treatment							
Medication	Genetic Variant	Outcome Moderated	Notable Studies					
Topiramate	GRIK1 (182832407)	Heavy drinking days (%); side effects	Kranzler et al., 2014 (2); Ray et al., 2009 (4)					
Naltrexone	<i>OPRM1</i> (Asn40Asp), (rs1799971), DRD4 VNTR	Heavy drinking days (%); abstinence rates; relapse to heavy drinking	Anton et al., 2008 (12); Kim et al., 2009 (13); Oslin et al., 2003 (14); Tidey et al., 2008 (15) Note: OPRM 1 predictive					
Ondansetron	LL/LS/SS (5-HTTLPR) (rs1042173), SLC6A4 (5-HTTLPR)	Drinks per drinking day; days abstinent (%)	Johnson et al., 2011 (9) value for NTX response has not been supported (Schacht, J., Randall, P., Latham, P. et al 2017)					
Sertraline	5-HTTLPR triallelic SLC6A4	Heavy drinking days (%); drinking days (%)	Kranzler et al., 2011 (8)					
Acamprosate	GATA4 (181327367)	Relapse	Kiefer et al., 2011 (10)					
Disulfiram	DBH (rs161115)	Adverse events	Mutschler et al., 2012 (11)					
(4)		Hartwell and Kranzler (2)	Batki & Pennington (2014) Am J Psychiatry 019)Expert Opinion on Drug Metabolism & Toxicology					

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Alcohol Use Disorder (Relapse Prevention) FDA Approved Naltrexone (Revia): 1994 Long Acting Naltrexone IM (Vivitrol): 2006 Acamprosate (Campral): 2004 Disulfiram (Antabuse): 1949 Nalmefene (2016) *European Medicines Agency (EMA)*



Pharmacotherapy of Alcohol Use Disorder:

Naltrexone-oral/Mechanism of Action

- · Reduces positive reinforcement (reward craving)
 - · Potent inhibitor at mu opioid receptors
- Modulates the mesolimbic dopamine system in the VTA & projections to the nucleus accumbens
- There is mixed evidence around markers that predict a favorable response to naltrexone treatment, such as:
 - Male sex
 - A positive family history of alcoholism
 - High levels of craving,
 - Polymorphism (asp variant) of the opioid receptor gene OPRM1?

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Pharmacotherapy of Alcohol Use Disorder:

Naltrexone-oral/Mechanism of Action

- The patient does not experience the full euphorogenic/reinforcing effect of alcohol.
 - suppresses/reduces endogenous opioids (beta-endorphin) involved in the reinforcing (pleasurable) and subsequent reduces DA in NAc effects of alcohol and possibly craving
- Prevents a slip from becoming a full-blown relapse

Pharmacotherapy of Alcohol Use Disorder:

Naltrexone-oral / Effectiveness

- · Effective in reducing relapse to heavy drinking.
- A meta-analysis of (N:16 studies and 2347 patients) found a:
- risk decrease (RD) for a return to any drinking (risk decrease = -0.05; 95% CI, -0.10 to -0.002; number needed to treat = 20)
- (19 studies N: 2875) found also a:
- risk decrease (RD) of binge drinking (risk decrease = -0.09; 95% CI, -0.13 to -0.04; number needed to treat = 12)
- · Medication compliance may be a limiting factor in oral treatment.

Kranzler Hr et al JAMA 2018 ; Srisurapanont M, Jarusuraisin N. Cochrane Databa Syst Rey 2005;(1):CD0018

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Pharmacotherapy of Alcohol Use Disorder:

Naltrexone-oral / Dosing and Safety

Oral Naltrexone Hydrochloride

- FDA approved dose: 50 mg per day
- · Antagonist of mu, delta and kappa opioid receptors.
- Antagonizes opioid-containing agents, but no other significant drug-drug interactions.
- Some have used 100 mg daily with rationale that naltrexone has been effective for heroin addiction at doses of 100mg-100mg-150 mg q Monday, Wednesday, and Friday; an effective plasma concentration can be obtained even if some doses are missed

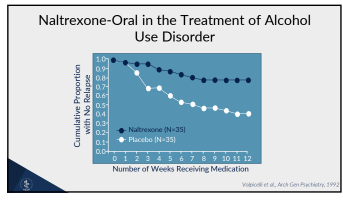
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Pharmacotherapy of Alcohol Use Disorder:

Naltrexone-oral /Dosing and Safety

- Side effects
 - · Gl: abdominal pain, diarrhea, decreased appetite, nausea
- Sedation: daytime sleepiness, fatigue, insomnia, headache
- Reversible hepatoxicity
 - · LFT's should be monitored closely (check LFT's prio starting medication)
- · Works best with complaint patients
- Requires counseling (CBT) or frequent MD monitoring visits (Project Combine, 2006)
- Efficacy questioned in women (O'Malley, 2007)

Physician's Desk Reference (www.PDR.net) and Epocrates. Accessed on September 1, 201:



Pharmacotherapy of Alcohol Use Disorder:

Long-Acting Naltrexone (IM)

Extended-Release - Injectable Naltrexone

- 1 injection per month/ 380 mg
- $\bullet\,$ 100 μm diameter microspheres of naltrexone and polymeric matrix.
- Advantages: once a month injection can be done in clinician's office.
- Better adherence with once monthly dosing
- More stable plasma concentrations compared to the oral formulation

Garbutt et al. JAMA. 2005;293:1617-1625. Physician's Desk Refere (www.PDR.net) and Epocrates. Accessed on September 1, 20

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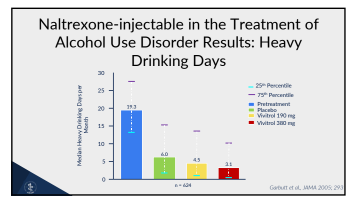
Pharmacotherapy of Alcohol Use Disorder:

Long-Acting Naltrexone (IM) Dosing and Safety

Extended-Release Injectable Naltrexone

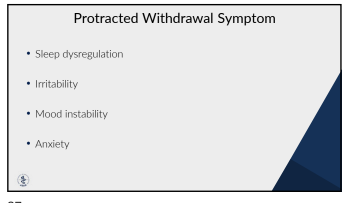
- Side effects: nausea & headaches; more sedation than with the oral formulation
- LFT's should be monitored closely
- Injection site reactions possible
- \bullet Best results in patients sober 1 week prior to starting the medication
- Efficacy shown in more severe alcoholics
- Reduction in heavy-drinking days (48.9% vs 30.9% on placebo)
- Pregnancy Category C , acceptable for use when breastfeeding

Pettinati HM Alcohol Clin Evn Res May 20



Return to A Placebo	ny D	rinking	g, Nait	rexone	· VS	Return to Hea	avy I	Jrinkin	g, Nait	rexone	vs Placeb
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National National Conference (National Conference (- "	Amount	PARIS	(meri	reactions become	Nationality, 50-mark and	-	-	100.000	00110	martine pount
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Online at at 14 1007	12	601098	80204B	0.82 (0.34 0.80)		Daller et al. ⁽⁴⁾ 1967	13	3/20 (14.3)	8/22 (24.10	0.40 (0.10 1.30)	
Anton # 4,18 1999	12	36/68 (32 N)	43,95 (96.7)	0.79 (0.60 1.80)	-+-	Return et al., ¹⁸ 2399	13	26/68 (38.2)	38/63 (60.3)	043 (0.44-0.81)	-
Oxo.er.4,™ 3000	12	79/95 (92.4)	6479301.0	142 (189-119)		Disk et al, ³⁷ 2000 Erwind et al, ³⁸ 2000	24	52/85 (87.10 183/018-01.80	\$3/79 (87.1) 105/309 (90.2)	100(0.80-1.24)	2
See 84.5 3001	12	18/31/34/3	11/18/83 D	48900140		Reynold on al, ** 2001 Marin on al, ** 2001	13	18/9410-01.0	13/34/07/3	0.87 (0.79 L.00) 0.04 (0.49 L.40)	
Krystal et al. ** 2001	15	255/418-01-01	145/299 (67.4)	9.50 (0.61 1.63)		Marris et al. 47 2001	12	29/55 (28:50	43/56/05/80	0561049-080	
Workers, 17, 2003	12	40/91 (79.2)	49/56 (87.5)	0.89 (0.75-0.80)	*	Lamin at, *1 2000	13	19/54 (30.9)	23/51 (52.9)	0.64(0.40 1.00)	
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Films et al. 7 1994	12	30/21/28 81	1100 OR D	1.00 (0.00 1.00		Ethan et al., ** 2000 Ethan et al., ** 2004	12	23/54 (94.0)	30/25/00 E	1.040.79-2.10	
Personal Page	12	20/20/20/40	2224040	1.01 (0.01 (.00)		Ribban et al., ** 2004 Auton et al., ** 2005	12	23/91 (40.3)	46/90/07/31	071030-030	
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07866m rt. 65™ 2000	12	49/57 (96-2)	3850 (N.O.	1.13 (554 1.30)		Markey vt. al. ** 2000	12	29/51 (73.6)	ONLOSS	104(03)-130	
0 Walley et 4,12 2005	16	20/34 (34.7)	30/34 (98.2)	9.79 (0.96-9.NY)		O'Mullay vt.ut, ⁵² 2007	13	35/57 (58.4)	33/90 (84.0)	1.07 (0.80-1.40)	
Baltieri-8 at, 77 2000	12	15/49 (71.4)	29/54 (10.20	0.99 (0.79-1.29)	-	D'Mallay et al, ³¹ 2006	16	22/94 (94.7)	28/34 (82.4)	6.79 (0.59 1.00)	-
Neterogeneity n2+0.01, 12+10.075, 162+1 Sec 414, 14, 10231 + 29.25, P+.06	34			0.93 (0.87-0.99)	1	Brown et al, ²¹ 2000 Mann et al, ²⁵ 2013	12	4/29 (39.0) 86/369 (38.5)	10/23 (43.5) 43/95 (48.5)	0.46(0.17.1.24) 1.05(0.80-1.34)	
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Amme.4,173006	16	345366-0.416	254,509 (62.2)	9.95 (1.88-1.83)		Test of 4, + 6, 10(22) + 53,29, P + 603 Nationalist, 500 regit cost					
Ordinal Al, 17 2008	24	95/130/79-25	94(129 (95.0)	0.99 (0.67-0.12)		Automotic (17 200)	16	263/309-063.00	235/309-(73.30	0901039-150	- 4
Personal et al, ¹⁶ 2010	14	36(49 (79 5)	30/39 (76.5)	1.80 (0.40 1.2%)	-	Dalmet A, 17 2008	24	73/320-98-83	76/130/013.29	0261079-1370	-
Microgenety of +0.00, if +0.00%, of +1.0 microfile, + 8/102.0 +0.002.0 ** . 79	10			0.87(0.91-0.00)	1	Television of a 200, C+0.000, C+1.00 Sec. of a, 4 a, 400 + 0.17, P+ 68				03919-94 1-80	F
Konder et al. ³⁷ 2004	12	139/116/02/30	243.557 IIIS.ID	0.82 (0.04 0.00)	4	Nathrouse, irgestion					110
Sarbutt et al. 77 2005	26	MERIT-00-9	295,009 (94.T)	0.00 (0.00 0.00)	-	Erander et al. 75 2004	12	123/190-077.29	132/157-094.10	6921032-1.60	P.
Materiageneity of =0.00; pf =54.40%, lef = 3 Section 4, = 4,-0(1) = 2.20; P = 14				0.96 (0.90-1.03)	Ŧ	ALX21-014. ¹⁷ 2001 Materiagnisety: v ² = 8.01, v ² = 68, 31%, v ² = 2.51 Text of a, v a, 1001 = 2.91, P = 68	12	90/152 (18.2)	78/148 (53.7)	1124030-1379	

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Pharmacotherapy of Alcohol Use Disorder:

Acamprosate/ Mechanism of Action

- Stabilizes glutamatergic neurotransmission altered during withdrawal (Littleton 1995).
- Chronic ETOH exposure alters GABA & NMDA systems
- Restores balance between inhibitory & excitatory neurotransmission
- Anticraving, reduced protracted withdrawal
- · Reduce negative reinforcement (abstinence craving)
- · No abuse liability, hypnotic, muscle relaxant, or anxiolytic properties

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Pharmacotherapy of Alcohol Use Disorder:

Acamprosate/ Effectiveness

- · Effective in improving abstinence.
- A meta-analysis (16 studies; N = 4847) concluded that acamprosate treatment was associated with a greater reduction than placebo in the risk of drinking among abstinent patients but no reduction in the likelihood of binge drinking.
- (risk decrease = -0.09; 95% CI, -0.14 to -0.04; number needed to treat = 12)
- The US trial showed efficacy only in patients motivated for abstinence.

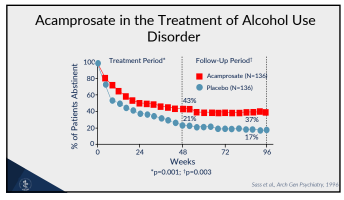
Jonas et al Jama 2014; Kranzler HR, Gage A. Am J Addict. 2008;17:70-76. M BJ et al. J Psychiatr Res. 2006;40:383

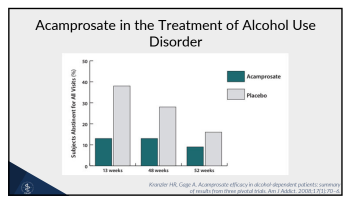
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Pharmacotherapy of Alcohol Use Disorder:

Acamprosate/Dosing and Safety

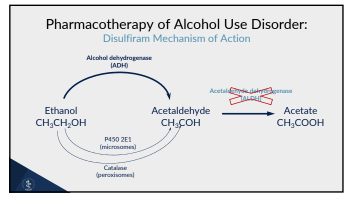
- 666 mg three times a day (2000 mg daily)
- Excreted by the kidneys; no liver metabolism
- Contraindicated: significant renal disease with creat cl <30ml/min or those who are pregnant
- Mild diarrhea (16% acamprosate vs. 10% placebo)
- Recommendation: patients with hepatic disease or those treated with opioids. Advantage
 when a patient is taking multiple medications
- No drug-drug interactions.Pregnancy category C





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Return to A	ny D	rinking	g, Acar	nprosate	e vs		Return to Heav	/y Dri	nking, A	Acampr	osate vs	
	Duration,	No./betal No. (10)		Riskratie	favors fav			Duction.	No./total No. (%)		Risk ratio	Favors Favo
Source University (Chief	sk II	Acamprisate 22(42/2014)	Raode 31(43,072.1)	(995.0)	acamposate pla	scebe	Source	vit.	Acamerosate	Placebo	(95% CI)	acampresate place
Underward of W1990	12	200/275/75/0	265(291(84.2)	0.89(0.81-0.90)	-		Disk at all 2000	24	246/289 (BS.1)	242/292 (\$2.5)	1.03/0.96-1.10)	southerny her
Pricetal ^{SE} 1992	26	42/55/06/0	45/47/95.70	0.80/0.68-0.90	- 4							
Palific et al, ¹⁶ 1995	53	294061(01-0	157/177 (86.7)	0.52 (0.65 8.93)			Kiefer et al. 37 2003	12	25/40 (62.5)	30(40 (75.0)	0.83 (0.62-1.12)	-++
Nhbroth-rt.st, ⁵¹ 25W	12	183(234(81.7)	208(224(92.5)	0.88 (0.82-0.95)			Kiefer et al ³⁷ 2003	12	20(40 (50.0)	25/40 (62.5)	0.80 (0.54-1.38)	
Sau et al, ⁵² 1996	48	75(136 (35.1)	102(136(75.0)	0.74 (0.61-0.88)	-		Worley et al. 42 2006	12	40(55(72.7)	43/61/70.9	1.08 (0.82-1.30)	1
Podrugs, ¹⁰ (He)	25	60(122(04.6)	84(124(67.2)	0.76(0.61-0.94)								
Pelc et al, ⁵⁷ 1987	13	74(126(587)	51/42/85.5)	0.69 (0.57-0.62)			Morley et al, 45 2006	12	40(55 (72.7)	39/53 (73.6)	0.99 (1.79-1.24)	+
Georgings et al. ¹² 1997	26	96(128(75.0)	116(134(86.6)	0.87(0.77-0.98)			Anton et al. ¹⁹ 2006	16	211/903 (69.6)	226/309 (73.1)	0.95 (0.86-1.05)	
Besonet al., ³⁴ 1996	11	41/55 (74.5)	47(55-(85.5)	0.87 (0.73-1.05)			Anton et al. ¹⁹ 2006.	16			1.04(0.93-1.16)	- 1
Pemperturit al, ¹⁷ 2000 Children al, ²⁶ 2000	26	87(354(53.0) 254(295(97.9)	115(166(60.1) 260/292(89.0)	0.77 (0.64-0.91)	-			16	211/903 (69.6)	207/309 (67.0)	1.04 (1.93-1.16)	
Clock et al. (** 2000 Guilland Lehert, ¹⁰ 2001)	26	254(299 (97.3) 92(34) (65.2)	290(252(3510) 109(147(74.1)	0.99(0.93-1.05)			Mason et al. 46 2006	24	143/341 (41.5)	119/260 (45.8)	0.92 (3.76-1.30)	-
Edward F 2003	12	30(40 (75.0)	37/40/52.50	0.81 (0.66-0.90)	- 2		William et al 65 2011	74	65/124 (52.4)	65/125 (52.0)	1.01 (0.79-1.38)	-
Baltieri and De Androdo, 71 2004	12	15(40)(17.5)	20/35/60/00	0.61(0.39-1.00)	7		Manuel of 6 2013					
Arton et al, ¹⁷ 2005	16	244/303 (30.5)	254(909(92.2)	0.98(0.91-1.06)				12	85(172 (51.7)	41/85 (48.2)	1.07 (1.82-1.40)	-
Morley-et.al, ⁴⁶ 2006	12	44(55 (80.0)	50/61 (62:0)	0.98 (0.82-1.16)			Mannet al,45 2013	12	85(172 (51.7)	86/159 (50.9)	1.02 (1.83-1.25)	+
Mason et al. ⁶⁵ 3006	24	128/941 (96.2)	240(260(90.8)	1.84 (1.80-1.00)			Heterogeneity: 12-0.00, 12-0.001; 12-1.00				1.00 (3.94-1.04)	1
Berger et al, ²⁰ 2013	12	48(52 (54.1)	40(49 (81.6)	1.15 (0.59-1.14)			Tex of a, = a; Q(10) = 5.90, P = 82					
Higuchiet al, ²⁵ 2015	24	86(163(52.8)	105364(64.0)	0.82 (0.68-0.99)			Oral				1.00 (0.96-1.04)	1
Percongrecky: r*=6.01, r*=77.641, r*=4.47 Text of a, = a; Q(25)=64.97, P < 001				0.86(0.83-0.95)	1		Heterogeneity: 12=0.00, 12=0.00%, 142=1.00				1.00 (1.96-1.04)	1
Overall Heterogeneitz: 1 ² + E.O.E. F = 77.64%, IF = 4.47 Test of group differences: Q,(0) = 0.00				0.88 (0.83-0.93)			Test of group differences: Q _p (3) + 0.00				02	



Pharmacotherapy of Alcohol Use Disorder:

Disulfiram/ Mechanism of Action

- Alcohol → Acetaldehyd → Acetate
- Disulfiram irreversibly binds to acetaldehyde dehydrogenase inhibiting the metabolism of acetaldehyde to acetate.
- Acetaldehyde accumulates resulting in a very unpleasant reaction Disulfiram – Ethanol Reaction (tachycardia, headache, nausea/vomiting, hypotension, sweating, warmness and flushing of the skin, dizziness, blurred vision and confusion).

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Pharmacotherapy of Alcohol Use Disorder:

Disulfiram Effectiveness

- Second Line Treatment
- In a meta-analysis of 22 studies was associated with:
- Sustained abstinent compared to control conditions only in open-label studies
- Double-blind, placebo-control study design is not helpful as both the medication and the placebo pills may (or may not) result in fear of drinking.
- Most studies are negative, but disulfiram may be helpful for a better response than control conditions when medication adherence was supervised

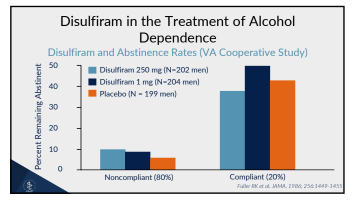
Diehl et al. Alcohol Alcohol. 2010;45:271-277. Fuller RK et al. JAMA. 1986;256:1449-5 Kranzler HR, Sayka M. Diagnosis and Pharmacotherapy of Alcohol Use Disorder. Review. JAMA. 2018;320(8):815-8.

Pharmacotherapy of Alcohol Use Disorder:

Disulfiram Dosing and Safety

- 250-500 mg daily.
 - First dose 12 hours after the last drink;
 - 500mg PO each morning for 1-2 weeks, then 250mg PO each morning
- Some liver toxicity; monitor LFTs at the beginning, 2 weeks, 3 months and then every 6 months. Caution with CAD. Contraindicated: psychosis, significant liver disease, esophageal varices, pregnancy, impulsivity, severe pulmonary disease, seizures, CRF (Barth et al., 2010)
- Inhibits hepatic microsomal enzymes and increases drug levels (phenytoin, warfarin, isoniazid, metronidazole, TCA and benzodiazepines among others)
- Pregnancy category C
- SIDE EFFECTS: skin/acneiform eruptions, drowsiness, headache, metallic taste, decreased libido/potency
 Physician's Desk Reference (www.PDR.net) and Epocrates. Accessed on March 1, 2018

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MAT + FDA Approved								
Medication (typical dose)	Mechanism of action	Adverse effects	Cautions	Lab monitoring	Other			
*Nattrexone (50-100mg PO daily or 380mg (M monthly)	Blocks opioid receptors May reduce rewarding effects of alcohol	Nausea Headache, dizziness, insomnia Anxiety *Injection site reaction	Need 7-10 days "opioid free" if patient previously receiving chronic opioids Do not use if: Current opioid use LFTs > 5x upper limit of normal	LFTs prior and during treatment	Number needed to treat to reduce heavy drinking days is 12			
*Acamprosate (666mg PO three times daily)	Levels out GABA + glutamate activity	Diarrhea	CrCl 30-50 mL/min: 333mg PO three times daily Do not use if: CrCl s 30 mL/min	Renal function (basic metabolic panel) prior and during treatment	Prolongs periods of abstinence			
*Disulfiram (250-500mg PO daily)	Blocks acetaldehyde dehydrogenase Blocks enzyme involved in dopamine metabolism	Disulfiram-alcohol reaction if combined Rare but notable: acute liver failure	Need ≥ 12h alcohol abstinence Many medication interactions Do not use if: Severe cardiac disease or coronary occlusion Primary psychotic disorder	LFTs prior and during treatment	Daily observed disulfiram Targeted disulfiram (e.g., weddings, reunions, holidays)			
(\$)			Primary psychotic disorder					

Combinations

- Naltrexone and acamprosate have different mechanisms of action and may work synergistically on cravings:
 - Naltrexone on positive reinforcement
 - Acamprosate on negative reinforcement
- Medications and psychotherapy.

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Rosner S et al. J Psychopharmacol. 2008;22:11-2:

Adapted from Kiefer F et al. Arch Gen Psychiatry. 2003;60:

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Naltrexone/Acamprosate Abstinence rates during a 12week trial with: Naltrexone 50 mg QD, Acamprosate 666 mg TID. The combination of the two medications helped alcoholics stay abstinent (P=0.002) better than each drug alone.

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Project MATCH

- Compared outcome efficacy for patients matched to treatments based on a prior hypotheses about 11 client attributes
- Treatment was for 12 weeks; follow-ups continued for years
- 12-Step programs, CBT and MET were compared
- Each of the three methods helped in the treatment of alcoholism
 - However outpatients who received TSF were more likely to remain abstinent after 1 year following treatment
- There were a few matching effects, and they were weak

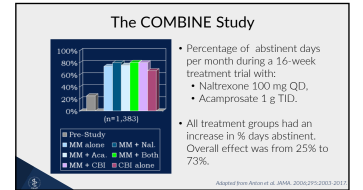
The COMBINE Study

- 1383 patients with alcohol dependence randomized to varying combinations of oral Naltrexone, Acamprosate, combined behavioral intervention (CBI) and medical management (MM)
- Patients received naltrexone, acamprosate, both, or neither
- Half of patients received psychotherapy in addition to medical management
- One patient cohort received psychotherapy alone, no pills

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JAMA. 2006;295:2003-201

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The NIAAA COMBINE Study Results

- For patients receiving MM, naltrexone, or CBI therapy, improved outcomes over placebo plus MM
 Naltrexone + MM had the best outcome
- Acamprosate did not add benefit to naltrexone or CBI, and was no more effective than placebo plus MM
- Taking tablets and seeing a health care professional was more effective than receiving CBI alone (possible placebo effect)
- One-year outcome: no significant differences among the groups

N=1383 (16 weeks trial)	Good Clinical outcome
MM and Placebo	58 %
MM and Placebo and CBI	71%
MM and Naltrexone	74%

CBI: Combined Behavioral Intervention Good Clinical Outcome: Abstinence or drinking moderate amounts without problems P<0.025 (interaction p-value 0.02)

Adapted from Anton et al. JAMA. 2006;295:2003-2017

Other Pharmacological Agents Alpha 2 agonists • Clonidine Anticonvulsants · Topiramate Gabapentin Serotonin (5-HT3) antagonists Ondansetron Mirtazapine Carbamazepine Valproic Acid Selective Serotonin Reuptake Inhibitors GABA agonist Baclofen Partial agonist for the $\alpha 4\beta 2$ nicotinic acetylcholine receptor subtype (nACH) • Varenicline Alpha1 adrenergic blocker Mu and delta opioid antagonist and partial kappa agonist Nalmefene • Prazosin

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• Identify the need of your patients to get treatment • Substance use disorders are chronic, be ready for relapses • Prevention is based on screening and early Intervention • CIWA-Ar is your best ally for AWS • AWS=BZD most effective, safest and cheapest treatment • Medications for Alcohol Use Disorder are relatively safe but modestly effective

- Acamprosate is best for preventing "the first drink."
- Naltrexone is best for "cutting down."

Conclusions

- Pharmacotherapy and psychotherapy modalities can be offered by you
- · Pharmacotherapy and psychotherapy modalities are effective and scientifically based approaches

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