

The Selection and Use of Adjuvant Therapies

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## **Disclosure**

- Courtney Kominek has nothing to disclose.
- This presentation was not a part of the presenter's official duties at the VA and does not represent the opinion of the VA
- The presentation will include "off-label" uses of some medications.

Painweek.

## **Learning Objectives**

- Describe where adjuvant analgesics act in the pain pathway and the differences in mechanism of action (MOA)
- Compare risks and benefits of different adjuvant analgesics for a given patient
- Choose an adjuvant analgesic based on current guidelines and/or evidencebased medicine as well as individual patient factors

Painweek.

Case	#1
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- MT is a 39-year-old male Iraqi War veteran presenting to clinic with complaints of low back pain that's limiting his ability to do his normal daily activities. He's tried acetaminophen, ibuprofen, and naproxen without benefit. He asks for a muscle colorest ted. relaxant today.

- relaxant today.

  He also reports flashbacks and nightmares of war. For the past few months he's been avoiding his military friends and isolating himself at home.

  Additionally, he reports hypertension controlled with hydrochlorothiazide/lisinopril 25mg/20 mg by mouth daily with a blood pressure today of 128/79 mmHg.

  Further work-up is completed and the patient is diagnosed with degenerative disc disease and post-traumatic stress disorder (PTSD).

  The patient is open to physical therapy and working with a psychologist but is also requesting medication options because the symptoms are so distressing. What might you consider prescribing?

Painweek.



Skeletal Muscle Relaxants

"WELL, FIRST, I THINK WE'LL CUT BACK ON THOSE MUSCLE RELAXANTS!"

Painweek.

Intro	du	ctio	n

Heterogeneous group

Structurally not related

2 million people per year report use of SMR

300,000 elderly patients use SMR

Associated with sedation and weakness as well as other adverse effects  $% \left( 1\right) =\left( 1\right) \left( 1\right) \left($ 

Fudin J, Raouf M. Pract Pain Manage. 2016; 16(5):1-18. See S, Ginzburg R. Skeletal muscle relaxants. Pharmacother. 2008;28(2):207-213.

	Description	Spasticity	Snaeme
	Definition	Velocity-dependent increase in muscle tone because of increased	Spasms Involuntary muscle contraction
	Etiology	Central     Upper motor neuron	Peripheral     Muscle sprain or injury
	Symptoms	• Stiffness	Nerve compression     Jerks
	*Toble adea	Hypertonicity     Hyperreflexia  ted from below reference. Used	Twitches     Cramps
NWEEK.	Fudin J, Raouf M	I. Pract Pain Manage. 2016;	16(5):1-18.
O	<b></b> On.		
Spastici	Description	Spasticity	Spasms
	Cause	Multiple sclerosis     Cerebral palsy     Spinal cord injury	Musculoskeletal pain     Fibromyalgia     Sciatica
		Traumatic brain injury     Motor neuron disease     Post-stroke syndromes	Mechanical low back pa     Herniated disk     Spinal stenosis     Myofascial pain
	FDA-approved medications	Botulinum toxin     Baclofen     Dantrolene	Carisoprodol     Chlorzoxasone     Cyclobenzaprine
		Diazepam	Metaxalone     Methocarbamol
		Riluzole	Ornhenadrine
		Riluzole     Tizanidine  rom below reference. Used with	Orphenadrine permission.
NEEK.		Riluzole     Tizanidine	Orphenadrine     permission.
<b>N</b> WEEK, 1		Riluzole     Tizanidine  rom below reference. Used with	Orphenadrine     permission.
<b>in</b> week. 1		Riluzole     Tizanidine  rom below reference. Used with	Orphenadrine     permission.
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<b>lin</b> week <u></u>		Riluzole     Tizanidine  rom below reference. Used with	Orphenadrine permission.
	Fudin J, Raouf M.	Riluzole     Tizanidine rrom below reference. Used with Pract Pain Manage. 2016; 1	• Orphenadrine opermission.  (6(5):1-18.
Neurotra	Fudin J, Raouf M.	Riluzole     Tizanidine rrom below reference. Used with Pract Pain Manage. 2016; 1	Orphenadrine permission.
Neurotra Spasm	ansmitter  Gamma-ami	Ruiuzole     Tizanidine rom below reference. Used with Pract Pain Manage. 2016; 1  S Involved in Manobutyric Glu  Obstantial Section 1.	• Orphenadrine o permission. (6(5):1-18.
Neurotra Spasm	Fudin J, Raouf M.	Ruluzole     Trandidne  rom below reference. Used with  Pract Pain Manage. 2016; 1  S Involved in Manage.  Glumobutyric  Printery  Printery	orphenadrine opermission.  (6(5):1-18.

Glycine

Inhibitory and excitatory roles

\*Adapted from below reference. Used with permission.

Painweek Fudin J, Raouf M. Pract Pain Manage. 2016; 16(5):1-18.

Acetylcholine (Ach)

• Primary neurotransmitter for sending signals from neurons to muscles

• Changes Na+ and Ca2+

# Carisoprodol

## ■MOA

- -Centrally acting
- -Changes interneuronal activity in spinal cord and descending reticular formation of brain
- -Decreases pain perception

## Dosing

- -350 mg PO QID -Max 1400 mg/day
- Avoid in children < 12 years (or EVERYONE)

Fudin J, Raouf M. Pract Pain Manage. 2016; 16(5):1-18.

See S, Ginzburg R, Skeletal muscle relaxants. Pharmacother. 2008;28(2):207-213.

Skeletal Muscle Relaxants. http://paindr.com/wp-content/uploads/2012/05/FINAL\_selayants\_2011;11(6):06.04.6-caps.ed.2 (lum.2016).

# Carisoprodol

## ADE

- -Abuse potential
- -Drowsiness, headache, vertigo, insomnia
- -Respiratory depression particularly in combo
- -Seizures with overdose, excessive use, withdrawal
- -Idiosyncratic allergic type reactions

## Kinetics

-Metabolized by CYP2C19 to meprobamate among others
• Subject to pharmacogenetic differences

Fudin J, Raouf M. Pract Pain Manage. 2016; 16(5):1-18.

See S, Ginzburg R. Skelstal muscle relaxants. Pharmacother. 2008;28(2):207-213.

Skelstal Muscle Relaxants. http://mainfacom/bracontent/fubloads/2017.05/E/IAAL s

## Chlorzoxazone

## MOA

- Acts at spinal cord and subcortical areas of brain
- Inhibition of multisynaptic reflex arcs

## Dosing

## • 500-75 mg PO TID-QID

- Dizziness, drowsiness,
- Rare hepatoxicity (monitor LFTs periodically)

ADE

- GI irritation or ulcer • Urine discoloration

Fudin J, Raouf M. Pract Pain Manage. 2016; 16(5):1-18.

See S, Ginzburg R. Skeletal muscle relaxants. Pharmacother. 2008;28(2):207-213.

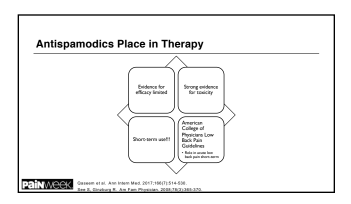
Skeletal Muscle Relaxants. http://paindr.com/wp-content/uploads/2012/05/F/NAL\_st

		_
Cyclobenzaprine		
■ MOA		
<ul><li>Structurally related to tricyclic</li><li>Not clear likely sedation</li></ul>		
<ul><li>No direct activity on skeletal n</li><li>Dosing</li></ul>	nuscle	
-5 mg PO TID -Increase up to 10 mg PO TID		
<ul><li>Avoid longer than 3 weeks</li></ul>		
<ul> <li>Kinetics         <ul> <li>Metabolized by CYP3A4, CYF</li> </ul> </li> </ul>	P1A2, and CYP2D6	
<ul> <li>ADE</li> <li>Anticholinergic side effects</li> </ul>		
-Avoid in patients with cardiac	conduction abnormalities or arrhythmias	
PainVeek See S, Graburg R. Skeletal muscle Skeletal Muscle Relaxants. http://ex	gg. 2016; 14(5):1-18. retalaarts. Planmaother. 2008;28(2):207-213. aindr.com/ter-content/unloads/2012(05F)NAL. skeletal-muscle-relaxants. 2011-11-06.odf.	
		_
	*All benzodiazepines have muscle relaxant properties	
Diazepam		
Approvals	Spasticity	
/ Applovais	• Muscle spasms	
MOA	• GABA receptor agonist → increases chloride conductance → presynaptic inhibition of spinal cord	
Dosing	• 2 mg PO BID-TID or 5 mg PO QHS	
Dosilig	Target 40 mg/day divided	
Painweek, Fudin J, Raouf M. Pract Pain Mana See S, Ginzburg R. Skeletal muscl. Skeletal Muscle Relaxoants. http://los	ga. 2016; 19(5):1-18. re relaxants. Pharmacother. 2008;29(2):207-213. ainth combine conferibulicosis/2012/05FINAL_skeletal-muscle-relaxants. 2011-11-06.pdf.	
Accessed 2 June 2017		J
		]
Diazepam		
Diazepaili		
Kinetics:	Elimination half-life 20-50 h     Active metabolites with half-life up to 100 h	
ixilletics.	Metabolized by CYP3A4 and CYP2C19	
Avoid	Elderly     Renal or hepatic impairment	
	)	
<b>AD</b> E	Abuse potential	
ADE	Dizziness, drowsiness, confusion, amnesia     Withdrawal with abrupt cessation	

Metaxalone	
• MOA	
Not established  No direct action on skeletal muscles or nerve fibers	
■ Dosing	
- 800 mg PO TID-QID ■ Kinetics	
- Bioavailability increased with high fat meal - Metabolized by CYP1A2, CYP2D6, CYP2E1, and CYP3A4	
• ADE	
<ul> <li>Dizziness, drowsiness (less compared to others), headache,</li> <li>Respiratory depression in combination</li> </ul>	
<ul> <li>Rare leukopenia and hemolytic anemia</li> <li>Avoid &lt; 12 yrs of age</li> </ul>	
<ul> <li>Avoid in patients with renal or hepatic failures</li> <li>Avoid in anemia</li> </ul>	
Fudin J, Raouf M. Pract Pain Manage. 2016; 18(5):1-18.	
Painweek See S, Ginzburg R. Skeletal muscle relaxants. Pharmacother. 2008;28(2):207-213. Skeletal Muscle Relaxants. http://apiandr.com/mys-content/uploads/2012/05/FINAL_skeletal-muscle-relaxants. 2011;13(6):6(1):6(1):6(1):6(1):6(1):6(1):6(1):6	
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Methocarbamol	
MOA Dosing ADE	
Centrally acting 1500 mg PO QID Discoloration of	
Carbamate x 2-3 days, then urine (brown-	
derivative of 750 mg PO QID black or green)	
guaifenesin Altered mental status	
Unknown status mechanism of Worsen	
muscle myasthenia gravis	
relaxation, likely sedation	
Fudin J, Raouf M. Pract Pain Manage. 2016; 16(5):1-18.	
See S, Ginzburg R. Skeletal muscle relaxants. Pharmacother. 2008;28(2):207-213.  Skeletal Muscle Relaxants. http://aaindr.com/wocontent/unloads/2012/05/FINAL_skeletal-muscle.	
relayants 2011-11-lik odt. Accessed 2 June 2017	
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Orphenadrine	
■MOA  -Anticholinergic agent	
-Structurally related to diphenhydramine	
-Inhibits antimuscarinic acetylcholine and N-methyl-D-aspartate receptors	
■ Dosing  —100 mg PO BID	
■ADE	
-Anticholinergic	
-GI disturbances     -Avoid elderly, glaucoma, cardiospasms, myasthenia gravis	
Contraindicated	
-Duodenal or pyloric obstruction or stenosing peptic ulcers Fudin J, Raouf M, Pract Pain Manage. 2016; 16(5):1-18.	
Fudin J. Raouf M. Pract Pain Manage. 2016; 18(5):1-18.  Bes S, Ginzburg R. Skeletal muscle relaxants. Pharmacother. 2008;28(2):207-213.  Skeletal Muscle Rekazants. http://gaindr.com/wp-content/uploads/2012/05/FINAL_skeletal-muscle-relaxants. http://gaindr.com/wp-content/uploads/2012/05/FINAL_skeletal-muscle-relaxants. 2018;27(2):2017.	
relayante 2011-11-106 ndf Accessed 2 lune 2017	

# Tizanidine MOA • Structurally related to clonidine • Centrally acting • Inhibits presynaptic and postsynaptic α-2 motor neurons • Potentiate glycine Fada J. Rand M. Pret Pain Manage 2016, 16(5) 1-18. Fada S. Gradys B. Salekid music entourist. Pharmacother. 2008 28(2) 297-213. Fada S. Gradys B. Salekid music relations of College 2017-213. Fada S. Gradys B. Salekid music relations of College 2017-213. Fada S. Gradys B. Salekid music relations of College 2017-213. Fada S. Gradys B. Salekid music relations of College 2017-213. Fada S. Gradys B. Salekid music relations of Why 2017. Fada S. Gradys B. Salekid music relations of College 2017-213. Fada S. Gradys B. Salekid music relations of Why 2017. Fada S. Gradys B. Salekid music relations of Why 2017. Fada S. Gradys B. Salekid music relations of Why 2017. Fada S. Gradys B. Salekid music relations of Why 2017. Fada S. Gradys B. Salekid music relations of Why 2017.

# Tizanidine ADE - Hypotension, sedation, asthenia, dry mouth - Elevated liver function tests, hepatoxicity - Monitor baseline, I, 3, and 6 months - Withdrawal syndrome with abrupt discontinuation - Avoid CrCl < 25 mL/min Fudn J, Raouf M. Pract Pain Manage. 2016; 18(5):1-18. Fig. J. Raouf M. Pract Pain Manage. 2016; 18(5):1-18. PRINTMONE Fig. S, Ginzburg R, Steelest muscle releasents. Pharmacother. 2008; 28(2):207-213. Res. S, Ginzburg R, Steelest muscle releasents. 2016; 18(5):1-18. PRINTMONE Fig. S, Ginzburg R, Steelest muscle releasents. 2016; 18(5):1-18.



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Tricyclic Antidepressants (TCA)	
	-
Painweek,	
Pain Week	
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Role in Pain Management	
First-line for neuropathic pain Second-line for neuropathic pain	
NICE     American Academy of Neurology	
Canadian Pain Society Guidelines     American Diabetes Association	
Neuropathic Pain Special Interest  Group of the International	
Group of the International Association for the Study of Pain	
European Federation of Neurological Societies	
Integrate B. Holyali, Exemine A. et al. A distribution for the first the Court Exemine Clear Grant Genthins, 2013.  Many D. G. Chin, Chine L. et al. Pan Exemine and Management Gentle (1971) 22-1.  P. M. S. Chin, C. Chin, Chin, C. Chin, Chin	
Ref III N V Cogland J. Frankin CM, et al. Neurology, 2011;78(20):1784-1785. Almal N, Crucco G, Baron R, et al. Eur J Neurol. 2010;17(6):1113-41186.	
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Role in Pain Management	
Effects	
independent of Low back pain	
BH disorder	
Effects Lower doses	
Fibromyalgia independent of compared to	
BH disorder MDD	-
<b>Point</b>	

noline muscarinic receptor, αι	=alpha-1 ad	Irenergic rec	eptor, H₁=his	tamine-1 recepto	r,
Medication	Ach M	αι	Hı	5-HT3	NE
Secondary amines					
Desipramine	+	+	+	+	++++
Nortriptyline	+	+	+	++	+++
Tertiary amines					
Amitriptyline	+++	+++	++	++++	++
Clomipramine	+	++	+	+++	++
Doxepin	++	+++	+++	++	++
Imipramine	++	+	+	+++	+++

## Tricyclic Antidepressants (TCAs)

- May initiate as follows:
  •Nortriptyline 10mg PO at bedtime
- Desipramine 25mg PO at bedtime
- Amitriptyline 10-25mg PO at bedtime
- -Increase by 10-25mg PO every 3-5 days
- -Use doses <100mg/day when possible
  -Do not exceed 50mg/day in patients on SSRI or SNRI
- -Use with caution in BPH, glaucoma, cardiac disease, and those at risk for suicide

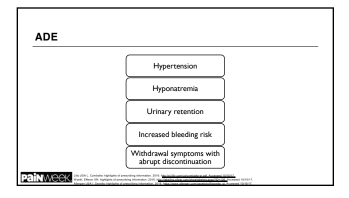
PainWeek, Lancet Neurol 2015; 162–73.

Cardi	$ac \rightarrow A$	void ir	n CV d	isease
	QTc			
Sudden cardiac death with doses > 100 mg/day	prolongation  Baseline ECG recommended by some in those >40-50 years of age Routine ECG monitoring not recommended	Arrhythmias	Tachycardia	Orthostatic hypotension

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ADE	
Anticholinergic $\rightarrow$ Elderly	
Dry mouth	
Constipation	
Urinary retention → BPH	
Tachycardia  Confusion	
Blurred vision → Glaucoma	
PainWeek Treeman MP, Markowitz JC, et al. Treatment of Patients with Major Depressive Disorder, 2010.  Insulantin completions session and insulanting and administration of the patients of th	
	_
ADE	
	-
Withdrawal symptoms	
Control of the control	
Suicide risk	
Seizure risk	
Histamine receptor antagonism $\rightarrow$ Sedation	
PainWeek Selectory AJ. Freemen MP. Markowitz JC, et al. Treatment of Patients with Major Depressive Discorder 2010.  https://www.documents.com/documents/doc	
	_
Serotonin Norepinephrine Reuptake Inhibitors (SNRIs)	
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<b>Pain</b> week.	
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Role in Pain Man	-9
	FDA-approved for
Duloxetine -	Diabetic peripheral neuropathy     FMS
Duloxetine -	Chronic musculoskeletal pain (LBP, OA)
	Second-line in American College of Physician guideline on LBP
Milnacipran	FDA-approved for FMS
Ē	• LBP
J	Diabetic peripheral neuropathy     FMS
Venlafaxine <	Chemotherapy-induced neuropathy
	Painful polyneuropathy
(	Headaches

# Common Nausea Somnolence Dry mouth Hyperhidrosis Erectile dysfunction Constipation



SNRI—Suicidality	
Warnings Fifected populations	
Timing of risk Monitoring and follow-up	
•	
Polinwood H. et al. Am   Psychony, 2008; 145(1):42-50.  Morrato BH. et al. Am   Psychony, 2008; 145(1):42-50.  Accessed by 18, 2012.  Accessed by 18, 2012.	
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SNRI Bleeding Risk	
Block serotonin uptake into platelet De-amplification of platelet aggregation Controversial data suggests: —Minimal risk of upper GI bleed as monotherapy Increased risk of upper GI bleed in combination with NSAIDs —Acid suppression therapy decreases risk	
Diluis 50, et al. Arb. Insen. May 2003;14(1):59.64. Licia Y, et al. Albamer Removal Bres. 2002;17(1):1-40.  Pali NACCHAR (A. Land Control Protection Bres. 2008;15(1):164-172.  de Alban S R et al. Arbon de Protection 2006;15(1):164-172.  de Alban S R et al. Arbon de Protection 2006;17(1):45-203.	
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Serotonin Syndrome	
Mental status changes     -Anxiety, agitated delirium, restlessness, disorientation	
<ul> <li>Autonomic hyperactivity</li> <li>Diaphoresis, tachycardia, hyperthermia, HTN, vomiting, and diarrhea</li> </ul>	-
Neuromuscular changes  Tremor, muscle rigidity, myoclonus, hyperreflexia, and clonus	
<ul> <li>Severity may range from benign to lethal</li> <li>Solely a clinical diagnosis</li> </ul>	
Patient and caregiver education paramount Consider serotonin active herbal/OTC products!!!	
- Consider Constant adults in order of a production	

<b>Duloxetine Dosing and Considerations</b>	
■ Dosing  —Initiate at 30mg PO daily x1 week, then increase to target dose of 60mg PO daily	
<ul> <li>Continue for 2 weeks at 30 mg daily in elderly</li> <li>In fibromyalgia and chronic MSK pain, no evidence that doses &gt;60mg/day provide additional benefit</li> </ul>	
■ADE  -Hyperglycemia	
-Avoid in chronic hepatic disease or cirrhosis -Avoid < 30 mL/min Control disease disease of control disease	
-Contraindicated uncontrolled closed-angle glaucoma	
PainWeck, Jily USA L Cynthille hapfright of prescribing information 2016. ** ** ** ** ** ** ** ** ** ** ** ** **	
	1
Venlafaxine Dosing and Considerations	
• Dosing	
<ul> <li>Initiate at venlafaxine SA 37.5 mg PO daily</li> <li>Titrate dose q2 weeks to 75 mg daily, 150 mg daily, 225 mg daily</li> </ul>	
OTc prolongation     Consider baseline ECG in those with cardiac disease history     Caution with renal disease – reduce doses	
- Mild to moderate: reduce total daily dose by 25-50% - Severe: reduce total daily dose by 50% or more  Caution with hepatic disease - reduce doses	
- Mild to moderate: reduce total daily dose by 50% - Severe: reduce total daily dose by at least 50% or more  • Caution uncontrolled closed-angle glaucoma	
Pain Weck  When the State of the Management 2010, Spanish and a find the state of t	
	_
Milnacipran Dosing  •FDA-approved indication for fibromyalgia	
Initial dose: 12.5mg PO once daily on Day 1     Initiation schedule;	
- Thration screenie; -12.5mg PO BID on Days 2-3 -25mg PO BID daily on Days 4-7	
-50mg PO BID thereafter ■ Target dose: 50mg PO BID (100mg/day)	
<ul> <li>Maximum: 100mg PO BID (200mg/day)</li> <li>Dose adjustment required in renal impairment</li> </ul>	

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Milnosianon Considerations	
Milnacipran Considerations	
<ul> <li>Hepatoxicity – no dose adjustment recommendations</li> <li>Use with caution in moderate renal impairment</li> </ul>	
Severe renal impairment (CrCl 5-29 mL/min), the maintenance dose should be	
reduced by 50% to 50 mg/day (25 mg twice daily). May increase to 50 mg BID  Not recommended in ESRD	
PainWeek. Allergan USA I. Savella: highlights of prescribing information. 2016. https://www.allergan.com/issosta/ord/fisnella_ci. Accessed 10/10/17,	
Post-traumatic Stress Disorder Treatment	
■ First-line ■ Suggest against	
-Sertraline -Amitriptyline	
<ul> <li>Paroxetine</li> <li>Inadequate evidence for or against</li> <li>Duloxetine</li> </ul>	
-Venlafaxine -Other TCAs  • Second-line	
- Imipramine	
VA/DoD clinical practice guideline for the management of posttraumatic stress disorder and acute	
PRINVECK Treating patients with acute stress disorder and post-traumatic stress disorder a quick reference quick. A maximum Republishin-description description.	
	•
Return to Patient Case	
Initiate venlafaxine SA 37.5 mg PO daily	
■Titrate dose q2 weeks as tolerated to at least 150 mg PO daily	
<ul> <li>Monitor BP</li> <li>Psychotherapy for PTSD</li> </ul>	
TO THE WOOL	

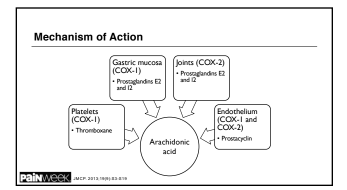
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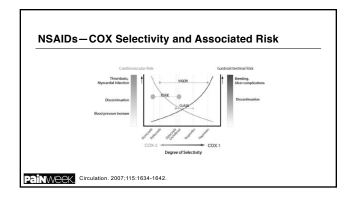
- MP is a 68-year-old male with diabetic peripheral neuropathy. His past medication history is significant for type 2 diabetes, uncontrolled hypertension, chronic kidney disease with CrCl = 43 mL/min, and benign prostatic hypertrophy (BPH).
- The patient has a history of alcohol use disorder that was in remission until recently. He reports that he has been drinking 10 beers a day lately for his pain but wants to return to sobriety.
- •The patient was started on amitriptyline 25 mg daily a week prior for pain control, and his wife is reporting that he has been confused and is having problems urinating. He presents to you for help with his pain.
- During medication reconciliation, you learn he is taking ibuprofen 400 mg q6h that he takes for knee arthritis.

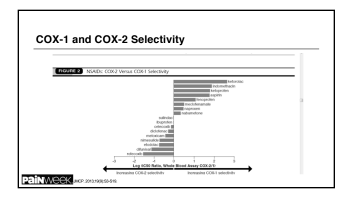
Painweek.

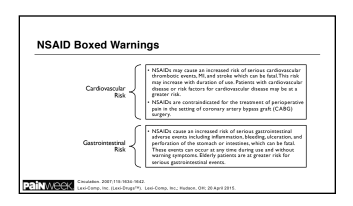
Nonsteroidal Anti-Inflammatory Drugs (NSAIDs)

Painweek.









## Celecoxib & Cardiovascular (CV) Safety

- Clinical question: How does the CV safety of celecoxib, a COX-2 selective NSAID, compare to that of a nonselective NSAID, such as ibuprofen or
- Primary composite outcome of CV death (including hemorrhagic death), nonfatal MI, or nonfatal stroke
- Mean treatment duration of 20.3±16.0 months and a mean follow-up period of 34.1±13.4 months
- In regards to the primary outcome, celecoxib was found to be *noninferior* to both ibuprofen and naproxen
- Risk of GI events was *significantly lower* with celecoxib compared to both ibuprofen and naproxen
- Study funded by Pfizer

PaiNWECK. N Engl J Med 2016; :2519-2529.

## **GI Adverse Events: Risk Factors** Moderate risk (1-2) Low risk High risk patients S years of age High dose NSAID therapy History of prior uncomplicated ulcer Concurrent use of low-dose aspirin, anticoagulants, or corticosteroids History of previously complicated ulcer Multiple risk factors (>2) No risk factors Pain Med. 2013;14:S18-S22. Am J Gastroenterol. 2008;104:728-738

## **GI Adverse Events: Prevention**

	Low GI Risk	Moderate GI Risk	High GI Risk
Low CV Risk	NSAID alone	NSAID + PPI or misoprostol	Alternative therapy or COX-2 + PPI or misoprostol
High CV Risk	Naproxen + PPI or misoprostol	Naproxen + PPI or misoprostol	Avoid NSAIDs or COX-2 inhibitors. Use alternative therapy

PaiNWEEK, Am J Gastroenterol. 2009;104:728-738

			٦
NSAIDs	and Renal Dysfunc	tion	-
	Avoid in people with GFR < 30 ml/min	Long-term therapy is not recommended	
	GFK < 30 mi/min	in people with GFR < 60 ml/min	
	Avoid with lithium	Avoid in people taking RAAS blocking agents	
Painweek.	er et al. Am J Kidney Dis. 2014;64(5):713-735.		
			_
			7
NSAIDs	and Renal Dysfunc	tion	
	Sulindac and salsalate		-
	may have less renal hemodynamic	Limit dose and frequency	
	changes	,	
	Topicals	Short-acting NSAIDs	
	.,	preferred	
Painweek.	um J Kidney Dis. 2014;64(5):713-735. Clin Kidney J. 2017;10(5):688-697.		
			_
			7
NSAIDs	and Liver Dysfunct	ion	
	moderate		
dysfuncti		Avoid in patients with	
etodolac,	doses: ibuprofen, , diclofenac	Avoid in patients with cirrhosis	
	duction: naproxen, o, sulindac		
	Proof Pain Manage 2010-12/03		
	Pract Pain Manage. 2013;13(9). J Gastroenterol Hepatol. 2014;29:13: Drugs. 2012;72(12):1646-1669.	56-1360	

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Drug-Drug	Interactions			
Anticoagulants, anti-				
Selective serotonin	reuptake inhibitors (SSRIs), serotonin	norepinephrine reuptake inhibitors (SNRIs)		
Cyclosporine, tacro	limus	_		-
Methotrexate		_		
Angiotensin convert	ing enzyme inhibitors (ACE-Is), angio	otensin receptor blockers (ARBs)		
Lithium				
Diuretics				
Ibuprofen + aspirin				
ainweek Grounds	<ul> <li>System. MICROMEDIEX 2.0, Greenwood Village, Colorad:</li> <li>n. 2014;129:907-916.</li> <li>Res Ther. 2013;15(Suppl 3):1-10.</li> </ul>	a. 21 March 2014.		
Penna!	and that, 2010, to cappe 0), 1-10.		_	
			7	
Topical P	roducts			
ainweek.			_	
			_	
Topical NS	SAIDs: Agents			
Medication	Indication	Dosing		
Diclofenac gel	Joint amenable to topical application (knee and hands)	2 g for each elbow, wrist or hand 4 g for each knee, ankle, or foot		
		Max 32 mg/day		
Diclofenac sodium topical	OA of knee	10 drops at a time on each of 4 sides of knee		
solution		40 drops QID		
Diclofenac	Topical treatment of acute	I patch to painful area BID		
	pain due to miner eteria -			
epolamine patch 1.3%	pain due to minor strains, sprains, and contusion			

## Topical NSAIDs: Place in Therapy

- American College of Rheumatology

  —Initial management of hand or knee OA may include topical NSAID
- American Geriatric Society
- -May consider topical NSAID for localized, non-neuropathic persistent pain
- European League Against Rheumatism (EULAR)
  - -Hand OA: topical NSAIDs over systemic
- -Hand or Knee OA: topical NSAIDs with clinical efficacy and safety
- National Institute for Health and Clinical Excellence (NICE)
  - -Topical NSAIDS considered in addition to nonpharmacological
  - -Consider topical NSAIDs or acetaminophen prior to PO NSAIDs

PainWeek.3MC. 2014;15:1-5

## **Topical NSAIDs: Pharmacokinetics** Gel: Solution: Absorption 2-3% 6-10% Patch Half-life ~12h Gel ~ Solution Patch 10-Time to peak 5-17h 10-14h PaiNWEEK. Packaging inform

## Lidocaine

- Topical anesthetic and Class 1b anti-arrhythmic
- Sodium channel blockade Na(v) 1.7
- Inhibition of acid sensing ion channels (ASIC)
- $\blacksquare$  Available via OTC (0.5%-4%) and prescription (5%)
- Lidocaine 5% patch applied directly to area of PHN
- -No more than 3 patches concurrently
- -12 hours on, 12 hours off ■OTC lidocaine 4% patch
- IV infusion is a potential treatment option

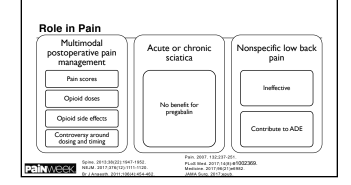
Lin J. et al. Inhibition of acid sensing ion channel currents by lidocaine in cultured mouse corotocl neurons. Amenth Analy 2011;11:2977-81.

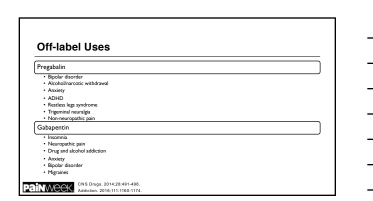
Kalle W. et al. Topical allocaine for the treatment of posterbepetic neuralgis. Cochrane Distabase Syst. Rev 2007;18:CD004946.

Contractions of the contraction of the contraction of the contraction of posterbepetic neuralgis.

Capsaicin 8% Patch	
Dose is a single, 60-minute application of up to 4 patches	
Consider	
Only physicians or healthcare professionals under after patch	
as warranted by the supervision of a application.	
return of pain   Patients may require   administer capsaicin   short-term pain	
8% patch medication postapplication	
Painweek. Capsaicin 8% patch [package insert]. Ardsley, NY: Acorda Therapeutics, Inc.; 2013.	
Capsaicin 0% patch (package insert). Alosiey, NT. Acorda Therapeutics, Inc., 2010.	
	1
Gabapentinoids	
Painweek.	
Mechanism of Action	
Structurally related to GABA and has GABA-mimetic properties	
Do not	
Alter uptake or breakdown     Convert into GABA	
Bind to GABA₃ or GABA₃	
Binds to the $\alpha 2$ - $\delta$ subunit of the voltage-gated calcium channel	
Reduces the Ca <sup>2+</sup> -dependent release of pro-nociceptive neurotransmitters	
Decreases release of glutamate, NE, and substance P	
Debreis Rei et al. Pairs 2007. 1319-227-251.  Centre F 1 & Debreis 2014 (Sent Sent Sent Sent Sent Sent Sent Sent	
I Clin Psychiatry. 2007 Mar;68(3):483-4	

# Role in Pain NICE Gabapentin - 1st line treatment for neuropathic pain ADA Diabetic Peripheral Neuropathy Consider pregabalin or duloxetine as initial approach ANN Diabetic Peripheral Neuropathy Offer pregabalin Consider gabapentin Neuropathic Pain Special Interest Group of International Association for the Study of Pain Gabapentin, pregabalin first line Addiscin. 2016;11:1160-174. Pain Neuropathic Pain Special Stripes Consider Spe





Gabapentinoi d Medication		Dosing	Renal Dose Adjustments
Gabapentin (Neurontin®)	PHN     Adjunctive treatment of partial onset seizures	Initiate at 100-300 mg PO QHS or TID. Doses can be increased by 100-300 mg/day every 1-7 days / Maximum dose 3600 mg/day Exceeding 1800 mg/day may not provide further benefit owing to saturable nonlinear kinetics	<ul> <li>30-59 mL/min – 400-1400 mg/day in 2 divided doses</li> <li>15-29 mL/min 200-700 mg in 1 daily dose</li> <li>15 100-300 mg in 1 daily</li> </ul>

Gabapentinoid Medication	FDA- Approved Indications	Dosing	Renal Dose Adjustments
Gabapentin (Gralise®)	• PHN	Take once daily with evening meal. Day 1:300 mg Day 2:600 mg Day 3-6:900 mg Day 3-6:900 mg Days 7-10:1200 mg Days 11-14;1500 mg Days 15;1800 mg Maximum dose 1800 mg/day	> 60 mL/min – none     30-60 mL/min – 600-1800 mg     < 30 mL/min do not use     Hemodialysis: do not use

abapentinoid ledication	FDA- Approved Indications	Dosing	Renal Dose Adjustments
abapentin nacarbil Horizant®)	Moderate to severe RLS     PHN	600 mg in AM x 3 days     Then increase to 600 mg PO BID.     Maximum dose 1200 mg/day	Selo mL/min no change 30.59 mL/min – initiate at 300 mg QAM × 3 days, may increase up to 600 mg BID 15-29 mL/min – 300 mg in QAM × 3 days then increase to 300 mg BID   Is 15-19 mL/min – 300 mg every other day, may increase to 300 mg QAM Hemodialysis – 300 mg after dialysis may increase to 600 mg after dialysis  Amount of the magnetic field of the selection of the select

Gabapentinoid	FDA-Approved	Dosing	Renal Dose Adjustments
Medication	DPN	Initiate at 150 mg/day in 2	1 12 11 1
Pregabalin (Lyrica)	PHN     Adjunctive therapy for partial onset seizures     Fibromyalgia     Neuropathic pain associated with SCI	or 3 divided doses. Increase dose to 300 mg/day within I week. Maximum doses vary depending on indication	> 60 mL/min – no change needed 30-60 mL/min – 75-300 mg divided BID or TIS – 150 mg divided daily or BID < 15 mL/min – 25-75 mg daily Hemodialysis – provide supplemental doses after dialysis based on daily dose

Gabapentinoid Medication	FDA- Approved Indications	 Renal Dose Adjustments
Pregabalin CR	PHN DPN	 Renal dosage adjustments needed

Medication	F	Tmax	okineti Half-life	Notes
Gabapentin IR (Neurontin)	900 mg 60% 1200 mg 47% 2400 34% 3600 33		5-7h	Bioavailability is not dose proportional
Gabapentin ER (Gralise)*		8h		Bioavailability is not dose proportional. Cmax increased 33-8% and AUC 33-118% with food depending on fat content. Absorbed from proximal small bowel by a saturable L-amino transport system.
Gabapentin enacarbil*	75%	7.3 h with food	5.1-6	Prodrug. Dose-proportional and extended exposure to gabapentin. Nonsaturable absorption
Pregabalin	90%		6.3 hours	Linear Cmax and AUC, independent of dose

Population-based nested case-control study Cases (1,256 cases) were opioid users who died of an opioid-related cause	
matched with up to 4 controls (4,619 controls)  Primary exposure was gabapentin use 120 days preceding index date	
■ 12.3% of cases and 6.8% of control were prescribed gabapentin	
•Odds increased 49% if prescribed gabapentin + opioid	
<ul> <li>High dose gabapentin (1800 mg/day) about 60% increased odds compared to moderate dose</li> </ul>	
■ Very high dose (2,200 mg/day) associated with 2-fold increased odds	
INVECK. PLoS Med. 2017;14(10:e1002396.	
, , ,	
Pregabalin Increases Overdose Odds	
Population-based, nested, case-control study in patients received opioid	
Population-based, nested, case-control study in patients received opioid 1417 cases: died of an opioid-related cause, excluding suicide or homicide	
Population-based, nested, case-control study in patients received opioid 1417 cases: died of an opioid-related cause, excluding suicide or homicide 5097 controls: matched on several characteristics	
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## **Gabapentinoid Abuse**

- Prevalence
- General population 1.1%
   Opioid use disorder
   15-22% gabapentin misuse
   40-65% abuse of gabapentin with prescription

  Dosing variety
- -Therapeutic range no prescription
- -Supratherapeutic range
  -3-20 times clinically used amounts
- -Taken as one large dose

Painweek.
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Addiction. 2016;111:1160-1174. CNS Drugs. 2014;28:491-496. Drugs. 2017;77:403-426.

Gabapentinoid Abuse	
Typically ingested with other substances	
Often used to increase high or treat withdrawal	
<ul> <li>90% of fatalities involve opioids</li> <li>Withdrawal treatment involves tapering gabapentinoid</li> </ul>	
Drugs. 2017;77:403-428.	
Brain Seg. 2016 Apr 22:80(2) Psyllottian Psylphotosism, 2011;89(2):118-22 ONS Drugs. 2014;28:891-98. Addisclin. 2016;11:1180-1174.	
PainWeek, Radiot be management of glostoperfinoid misuse. Available at:  https://doi.org/10.1001/10.10	
Role in Addiction Treatment	
■ Pregabalin	-
-Alcohol withdrawal	
<ul><li>Alcohol relapse prevention (abstinence similar to naltrexone)</li><li>Benzodiazepine/opioid withdrawal</li></ul>	
-Some evidence to prevent cocaine relapse	
<ul> <li>Gabapentin</li> <li>Evidence in opioid, THC, alcohol addictions</li> </ul>	
-Gabapentin suggested in APA AUD Guidelines	
<ul> <li>Goal of reducing or abstaining from alcohol</li> <li>Prefer topiramate or gabapentin or intolerant or did not respond to naltrexone or acomprosate</li> </ul>	
No contraindications	
CNS Drugs. 2014/28-491-498. Pacific Quideline III Agrica Quideline III A	
	1
Other Anticonvulsants	
Outer Attriconvulsaries	
<b>Pain</b> week	

# Carbamazepine (CBZ) and Oxcarbazepine (OXCBZ) • MOA: inhibit voltage-gated sodium channels and potentiate GABA • Role - CBZ drug of choice for trigeminal neuralgia - OXCBZ • Trigeminal neuralgia • Specialist setting/4th line NICE neuropathic pain guidelines Tegeted package insert. East Hanover. NJ: Novartis: 2018 March. Trigeded package insert. East Hanover. NJ: Novartis: 2018 March. Trigeded package insert. East Hanover. NJ: Novartis: 2018 March. Trigeded package insert. East Hanover. NJ: Novartis: 2018 March. Pair J Neurol. 2010;17(9):113-e1188. Neurosciences: 2015/2002/107/114. Eccellence: Clinical Guidelines. 2011

CBZ IR and XR	OXCBZ IR	OXBZ XR
Initial: 100 mg PO BID     Titrate by 100 mg PO BID     Target dose 300-900 mg/day     Max dose: 1200 mg/day	Initial: 150 mg PO BID     Titrate by 300 mg q3 days     Target: 300-600 mg PO BID     Max dose: 1800 mg/day	Initial dose: 600 mg PC daily     Tirrate by 600 mg/day weekly     Max dose: 2400 mg/day

CBZ	
-Metabolized by CYP3A4	
-Active metabolite carbamazepine 10,11, epoxide autoindu -Induces CYP3A4, CYP1A3, CYP2B6, CYP2C9, CYP2C1	
OXCBZ	
-Keto-derivative of CBZ     -Metabolized to active metabolite 10-monohydroxy oxcarb metabolism	azepine which avoids CYP
-May be better tolerated	
-20-30% have cross-reactivity with OXCBZ if allergic to CE	3Z

Carbamazepine and Oxcarbazepine	
■Common ADE  -Diplopia, abnormal vision  -Fatigue  -Allergic reactions	
-Fatigue -Allergic reactions -Dizziness -Pancytopenia -Somnolence -Agranulocytosis	
-N/V -Leukopenia -Ataxia -Serious dermatological reactions -Headache •HLA-8-1502 testing for those with Asian	
-Nystagmus ancestry -Tremor -Cardiac (BP, CHF, arrhythmias, AV block) (CBZ)	
-Abnormal gait -Elevation LFTS (CBZ)  Painweek Hd padage Peet East Honoer, NJ Novatic 2018 Merch. Hd padage Peet East Honoer, NJ Novatic 2018 Merch.	
The American Production of the Control of the Contr	<u> </u>
Lamotrigine	
<ul> <li>MOA: voltage-gated sodium channels</li> <li>Role</li> </ul>	
<ul> <li>-4<sup>th</sup> line/specialist setting for neuropathic pain</li> <li>-Data supports use in refractory trigeminal neuralgia, central poststroke pain, SCI pain</li> </ul>	
with incomplete cord lesion and brush-induced allodynia, HIV-associated neuropathy in patients on anti-retroviral therapy, and diabetic neuropathy  Dosing	
-TITRATE DOSE SLOWLY -Initiate at 25 mg PO daily x 2 weeks then increase to 50 mg/day for 2 weeks	
–Then titrate by 50 mg/day q1-2 weeks –May need to titrate to 200-400 mg/day	
Pein Research and Management 2014;18(5):230-335.  Bur J Menur 2016;17(5):1113-1188.  Neurosalité le Pinn The Pharmacologial Research of Neurosalité.  Lamida javalage insert. Research Triengle Park. Glassofimith/Gine; 201  Care Excellence Circles Guidelières. 2013.  Neurol So (2006) 27:5185-5189.	
Lamotrigine	
Common ADE     —Blood dyscrasias     —Dizziness     —Aseptic meningitis	
Nausea Insomnia Somnolence	
−Fatigue −Diplopia	
-Ataxia Severe ADE	
-FATAL OR LIFE-THREATENING HYPERSENSITIVYT	

■MOA	
-Inhibit	s voltage-gated sodium channels
-AMPA	kainate subtype of glutamate receptor
-Carbo	nic anhydrase inhibitor
-Increa	ses activity at GABA-A receptor
■ Role	
-Alcoho	ol use disorder
-Migrai	ne prophylaxis
-Neuro	pathic pain
	The American Psychiatric Association Procisio Guideline for the Pharmacological Treatment of Patients With Alcohol Use Clisoder. https://psychiatryonline.org/puidelines   Path Research and Management. 2014;19(5):2233-335.
NWEEK	Pair International Conference (Including Conference Con

## **Topiramate**

## Dosing

- -Topiramate IR
  - Initial dose: 25 mg po daily x 1 week Titrate by 25-50 mg/day

- Titrate by 25-50 mg/day

  Target dose

  Migraine: 50 mg PO BID

  Neuropathic pain 200-400 mg/day

  Topiramate XR

  Initial dose: 25 mg PO daily

  Titrate by 25mg/day q week

  Target dose: 100 mg/day

Trokendi XR package insert. Rockville, MD: Supernus Pharmaceuticals; 2018 Jan. Topamax package insert. Titusville, NJ: Janssen Pharmaceuticals, Inc; 2018 June.

## **Topiramate**

- Dose-related ADE
- -Fatigue
- -Nausea -Anorexia
- -Dizziness
- -Difficulty with memory
- -Diarrhea
- -Weight loss -Concentration/attention
- -Somnolence

- Caution
  - -Secondary angle glaucoma
  - -Metabolic acidosis
  - -Hyperammonemia
  - -Kidney stones
  - -Oligohidrosis -Hypo/hyperthermia

  - -Cognitive dysfunction
    -Renal adjustments CrCl < 70 mL/min



PatinWeek Trokendi XR package insert. Rockville, MD: Supernus Pharmaceuticals; 2018 Jan. Topamax package insert. Titusville, NJ: Janssen Pharmaceuticals, Inc; 2018 June.

	1
Return to Patient Case #2	
Discontinue amitriptyline	
After resolution of side effects	
-Initiate gabapentin 300 mg PO QHS	
-Titrate by 300 mg/day q3-5 days as tolerated	
-Max dose 1400 mg/day in 2 divided doses	
<ul> <li>Discontinue ibuprofen</li> <li>Consider topical diclofenac 1% TID</li> </ul>	
- Consider topical dictoreriac 1/6 FID	
<b>Pain</b> Week.	
	•
	_
Conclusions	
Adjuvant and coanalgesics require judicious monitoring for safe use	
<ul> <li>Extensive patient education regarding potential</li> </ul>	
adverse effects is paramount	
<ul> <li>Comorbid disease processes and concurrent medications may obscure</li> </ul>	
adverse effects	
<b>Pain</b> week.	
<del></del>	1
Learning Assessment Questions	
<b>Pain</b> week.	