

Nonopioid Analgesics:

The Selection and Use of Adjuvant Therapies

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Disclosures

Nothing to disclose

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2

Objectives

- Describe where adjuvant analgesics act in the pain pathway and their differences in mechanism of action
- •Compare risks and benefits for different adjuvant analgesics
- Choose an adjuvant analgesic based on current guidelines and/or evidence-based medicine as well as individual patient factors

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Are opioids still a concern?

- Drug overdose fatalities involving opioids in the US from 1999 to 2017¹
- ■399,230 (56.8% of all cases)
- Drug overdose fatalities involving opioids in the US in 2017 ¹
- ■47,600 (67.8% of all cases)
- Rate of overdose fatalities involving opioids in women age 30-64 from 1999 to 2017 increased by²
- **492%**

Schold L, Seith P, Keinss M, Willen N, Beldein C, Drus and Opinid-Involved Overdose Deaths — United States. 2013-2017. MMRN Mee Moot Mark Mee 2018(67:1461-146).

2017. MMRN Mee Moot Mark Mee 2018(67:1461-146).

2018. Mark M. Drus Operation Deaths Among Women Aged 30-64 Years — United States. 1999-2017. MMRN Mee Moot Mark Mey 2019(881-56.

2018. Mark M. Drus Mee 2019(881-56.

2019. Mark M. Drus Mee 2019(881-56.

Risk Factors for Opioid Overdose or Addiction

Risk factors for overdose

- Daily dose > 100 MEDD
- Long-acting (LA) or extended-release (ER) formulation
- Combination with benzodiazepines
- Long-term use (> 3 months)
 Period shortly after initiation of LA/ER formulation

Risk factors for addiction

- ■Age > 65 years
- Sleep disordered breathing
- Renal/hepatic impairment
- Depression
- Substance use disorder
- History of overdose

Volkow NJ et al. NEJM.2016;374:1253-1263. MEDD = morphine equivalent daily dose

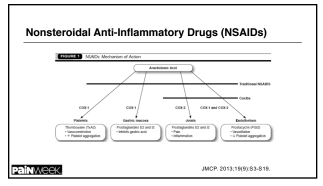
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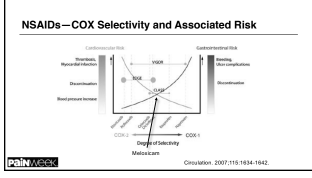
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Where Do Adjuvants Work? Painweek.

NSAID Ibuproten Naproxen Ketorolac (IV form) Meloxicam Celecoxib Corticosteroids	Normal Joint Ostecarthritis Rheumatoid Arthritis bore ercalcy frametaure grands of the formal Joint Capitule Laurillage bone enchance to logister symbolism are produced emericans and the formal Joint Capitule Cartiflage bone enchance to logister and the formal Joint Capitule Cartiflage bone enchance to logister and the formal Joint Capitule Cartiflage bone enchance to logister and the formal Joint Capitule Cartiflage bone enchance to logister and the formal Joint Capitule Cap
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Nonsteroidal Anti-Inflammatory Drugs (NSAIDs)





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 naproxen?
- периодит:
 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

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N Engl J Med 2016; :2519-2529.

11

NSAIDs and GI Adverse Effects

- •Strategies to prevent gastric mucosal damage in chronic NSAID users:
- -Proton pump inhibitor (PPI)
- -Histamine-2 receptor antagonist (H2RA)
- -Use of COX-2 selective NSAID
- ■Risk factors for NSAID-related GI toxicity:
- -History of peptic ulcer disease or upper GI bleed
- -≥65 years old
- -Presence of comorbidities such as rheumatoid arthritis
- -Concomitant use of anticoagulants, aspirin or corticosteroids

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Am J Gastroenterol. 2009;104:728-738.
 JMCP. 2013;19(9):S3-S19.
 Circulation. 2007;115:1634-1642.

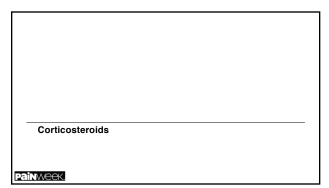
Topical NSAIDs

- Diclofenac sodium 1% gel
 Dosing:
 Upper extremity (hands, elbows, wrists): 2g applied QID up to 8g on any one joint
 Lower extremity (knees, ankles, and feet): 4g applied QID up to 16g on any one joint
- Diclofenac epolamine 1.3% patch -1 patch applied BID to the most painful area
- Both products carry the same boxed warnings but are proposed to have a more favorable safety profile than oral NSAIDs
- Most common adverse effect: application site reactions

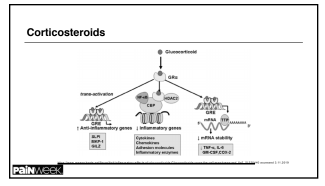
Painweek.

1.Pain Medicine 2013; 14: S35–S39. 2.Cochrane Database of Systematic Reviews 2012, Issue 9. Art. No.: CD007400.

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- Mechanism of action leads to a decrease in production of heat shock proteins intracellularly leading to a decrease inflammation
- Multiple routes of administration
- -Parenteral
- ٠IV
- •IM depot
- •Intraarticular

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16

Glucocorticoids (cont'd)

- Caution should be exercised in patients with the following conditions
- -Diabetes
- -Psychiatric history
- -Heart failure
- -Adrenal suppression
- •Taper needed when therapy exceeds 10 to 14 days
- -Immunocompromised

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17

Neuropathic Pain

- Anticonvulsants
- -Gabentin
 -Pregabalin
 -Carbamazepine/oxcarbazepine
 -Lamotrigine (off-label indication)
 -Topiramate (off-label indication)
- Antidepressants
- -TCAs (off-label indication)
 -SNRIs
- •Local anesthetics



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Anticonvulsants	
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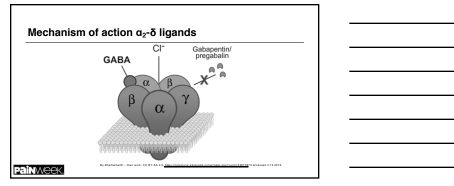
Anticonvulsants Gabapentin & Pregabalin

- Structurally related to GABA but it does not bind to GABA_A or GABA_B receptors or influence the degradation or uptake of GABA
- Binds to the $\alpha_2\text{-}\delta$ subunit of voltage-gated Ca²+ channels in CNS and peripheral nerves
- Reduces the Ca²⁺-dependent release of pro-nociceptive neurotransmitters, possibly by modulation of Ca²⁺ channel function
- Pregabalin may also interact with descending noradrenergic and serotonergic pathways in the brainstem

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J Clin Psychiatry. 2007 Mar;68(3):483-4.

20



Anticonvulsants Initial dose: 100 mg to 300 mg by mouth up to 3 times daily Increase dose based on response and tolerability to a maximum total daily dose of 3600 mg 3600 mg Renal dose adjustment required NO hepatic adjustment needed Gabapentin is not metabolized by hepatic enzymes Most common adverse effects: Dizziness and drowsiness (approx. 20%) Ataxia Fatigue Painweek. 22

Anticonvulsants (cont'd)

Pregabalin

- Initial dose: 25 mg to 150 mg by mouth once or twice a day
- Increase dose in 1 week based on tolerability to a maximum daily dose of 450 mg
 Doses up to 600 mg have been evaluated with no significant additional benefit
- Renal dose adjustment required
- NO hepatic adjustment needed

 —Pregabalin is minimally metabolized by hepatic enzymes
- Most common adverse effects:
 Dizziness and somnolence
 Peripheral edema

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23

Anticonvuls	sants: Alternative Options
Carbamazepine	
-Drug of choice f	or trigeminal neuralgia
 May require titra 	ation of dose to maximum of 1200 mg/day
-Consider obtain	ing baseline CBC and LFTs
 Consider per 	iodic monitoring of CBC and LFTs thereafter
 Oxcarbazepine 	
-Better tolerabilit	y compared to carbamazepine
-Titration begins	at 150 mg twice daily to a maximum dose of 1800 mg/day
-Patients allergic 25% allergic cro	to carbamazepine should also avoid oxcarbazepine, ss-reactivity
	Hooten M, et al. Institute for Clinical Systems Improvement. Pain: Assessment, Non-
in week	Opioid Treatment Approaches and Opioid Management. Updated September 2016. 2. Update on neuropathic pain treatment for trigeminal neuralgia. Neuroscience, 20.2.107-14 2015.

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Anticonvulsants: Alternative Options (cont'd)	
Lamotrigine (off-label indication) Data supports use in refractory trigeminal neuralgia, central poststroke pain, SCI pain with incomplex cord legions and hysthyndriged allowings. HIV-associated neuronaphy in patients on	
incomplete cord lesion and brush-induced allodynia, HIV-associated neuropathy in patients on antiretroviral therapy, and diabetic neuropathy - Most effective at doses between 200-400 mg/day - Note: follow strict titration schedule to reduce the risk of serious skin reactions	
- Immune response?	
Topiramate (off-label indication) Data supports use in diabetic neuropathy, refractory trigeminal neuralgia, and for migraine prophylaxis Dosing generally ranges from 50-100 mg/day	-
- Dosing over 200 mg is generally side-effect limiting	
Neurol Sci (2008) 27:5183-5188 R.H. Dworkin et al. / Pain 132 (2007) 237-251.	
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Anti-consularata Nessasanitisa	
Anticonvulsants – Neurocognitive	
Psychomotor reaction time Learning, memory, and executive function	
Word finding Considerable variance based on:	
–Age	
Multiple anticonvulsants Serum drug concentrations	
 All anticonvulsants appear to have some effect on neuropsych batteries 	
Meader KJ. Epileopy Res. 2006;86(1):83-87. Pendres Cd. ed. Pedart Neural 2010;42(3):187-196. North MN. Pendres SC. Obstantiantiev evens carbannazapine monotherapy for partial creet seizures. Cochrane Database of Systematic Reviews 2000; Issue 4.4. No.: C0006453, DOI: 10.1002/14851658 C0006453, pub2. Hessen, Ed. d. And Neuro Sacra (1005):11(9):134-134-136.	
4. Hessen E, et al. Acts Neurot Scand 2009;119(3):194-196.	J
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Antidepressants	
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Tricyclic Antidepressants (TCAs)

Initial dosing of TCAs

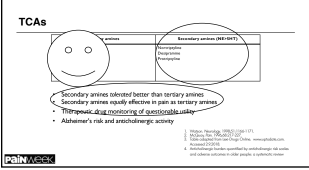
- Nortriptyline 10 mg at bedtime (off-label indication)
- Desipramine 25 mg at bedtime (off-label indication)
- Amitriptyline 10-25 mg at bedtime (off-label indication)

 -Use with caution in BPH, glaucoma, cardiac disease, and those at risk for suicide

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Lancet Neurol 2015; 162-73.

28



29

TCAs-Anticholinergic & Sedation

- Muscarinic receptor antagonists
- -Blurred vision, constipation, dry mouth, urine retention, constipation, tachycardia, confusion, delirium, increased ocular pressure
 -Secondary amines < tertiary amines
- Antihistaminergic effects (sedation, delirium)
 Maprotiline, amitriptyline, doxepin, and trimipramine

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TCAs-Cardiovascular Risk

- Orthostatic/postural hypotension
- -Alpha adrenergic blockade (even at low doses)
- Slowed cardiac conduction, tachycardia, ventricular fibrillation, heart block, and ventricular premature complexes (similar to Class Ia AA)
 Sudden cardiac death (unclear association with QTc prolongation)
- -Avoid doses > 100 mg/day amitriptyline equivalents
- Avoid in those with cardiovascular disease or established conduction abnormalities
- Screen for known heart disease, syncope, palpitations, dyspnea, or chest pain
- Baseline ECG recommended by some in those > 40 years of age (> 50 years of age based on APA Depression Guidelines)
- Routine ECG monitoring not recommended unless CV symptoms arise

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Ray WA, et al. Clin Phormocol Ther. 2004;75:234-241.
 Gelenberg AJ, et al. Practice guideline for the treatment of patients with Major Depressive Disorder, 3rd Edition. www.psychiatrycelline.org. Accessed 29.2018

31

TCAs-Behavioral Health Risks

- Abrupt discontinuation
- -Withdrawal symptoms (GI, malaise, chills, rhinitis, and myalgias)
- -Rebound depression
- Increased suicidality vs overdose toxicity
 - -Boxed warning for children, adolescents, young adults (18-24 years of age)
 - -Cardiac (QTc) and anticholinergic toxicity at doses as little as 10 x prescribed
 - Labbate, LA, Fave, M, Rosenbaum, JF, et al. Drugs for the treatment of depression. In: Handbook of Psychiatric Drug Therapy, 6th act Uppincox Williams & William, Psiliadelphia 2010.
 Dallai A et al. J Can Psychopharmocology. 1998;18:343-344.
 Frys MA, et al. Am J Psychophary 2009;165:164-172.
 Van Schweg Dr. et al. Am Graphyston; 1979;26:560-565.

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32

SNRI

Venlafaxine (off label)

- ■Initial dose: 37.5 mg to 75 mg ER by mouth
- once a day

 Increase dose by 37.5 mg to 75 mg ER daily every week -Target dose of 225 mg ER once daily
- larget cose of 225 mg EH once daily
 Renal and hepatic dosing adjustments necessary
- Discontinuing therapy should be done over 2 to 4 weeks

 Most common adverse effects

Duloxetine

- Initial dose: 30 mg by mouth once a day ■Increase dose to 60 mg ER every week

 —Maximum daily dose 120 mg
- Avoid use with severe renal or hepatic
- Discontinuing therapy should be done over 2 to 4 weeks
- Most common adverse effects
 Suicidal ideations [Black box warning]
 Children and up to 24 years of age
 Cognitive impairment

 Suicidal ideations [Black box warning]
 Children and up to 24 years of age
 Anxiety, insomnia Painweek.

SNRI (cont'd) Milnacipran for fibromyalgia —Initial dose: 12.5 mg PO once daily on Day 1 -Titration schedule: -12.5 mg PO BID on Days 2-3 -25 mg PO BID on Days 2-7 -50 mg PO BID thereafter -Target dose: 50 mg PO BID (100 mg/day) -Maximum: 100 mg PO BID (200 mg/day) -Dose adjustment required in renal impairment Painweek. 34

Serotonin Syndrome

- Mental status changes
- -Anxiety, agitated delirium, restlessness, disorientation
- Autonomic hyperactivity
 Diaphoresis, tachycardia, hyperthermia, HTN, vomiting, and diarrhea

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- Neuromuscular changes
 - $-\mbox{Tremor},$ muscle rigidity, myoclonus, hyperreflexia, and clonus
- Severity may range from benign to lethal
- Solely a clinical diagnosis
- Patient and caregiver education paramount

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Boyer EW, et al. N Engl J Med. 2005;352(11):1112-1120.
 Mackay FJ, et al. Br J Gen Proct. 1999;49(448):871-874.

35

Diagnosis of SS-Hunter Criteria

- Serotonergic agent PLUS one of the following:
- -Spontaneous clonus
- -Inducible clonus and agitation or diaphoresis
- -Ocular clonus and agitation or diaphoresis
 -Tremor and hyperreflexia
- -Hypertonia
- -Temp above 38°C (100.4°F)
- reinplatous of Clious Pro-Although Clinical dx, consider CBC, BMP, INR, CPK, LFTs, UA, chest X-ray, head CT, to rule out differentials

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Dunkley EJ, et al. QJM. 2003;96(9):635-642.

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SNRI Bleeding Risk		
■Blocked serotonin uptake into platelet		
De-amplification of platelet aggregControversial data suggests:	gation	
-Minimal risk of upper GI bleed as monotherapy -Increased risk of upper GI bleed in combination with NSAIDs		
 Acid suppression therapy decreases 	s risk	
	Dalron SO er al Arch Intern Med 2003: [AV]]: \$9,64	
Painweek.	 Dalcon SO, et al. Arch Intern Med. 2003;163(1):59-64. Lobe TK, et al. Aliment Pharmacol Ther. 2008;27(1):31-40. McClosber QJ, et al. Timat Res. 2008;15(1):161-81-72. de Abajo FJ, et al. Arch Gen Psychiotry. 2008;65(7):3795-803. 	
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Local Anesthetics		
Local Allesthetics		
Pain week.		
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Lidocaine		
■ May be used topically or by injection		
 Topical patch available in 0.5% to 5% 5% patch applied directly to area of postherpetic neuralgia¹ 		
No more than 3 patches concurrent12 hours on, 12 hours off		
■Trigger point injections ² -Lidocaine or procaine		
	nts and local anesthetic allergy history	
pain week	Kaliq W. et al. Topical lidocaine for the treatment of postherpetic neuralgia. Cochrane Database Syst Rev 2007;18:CD004846. Alvarez DJ, et al. Trigger Points: Diagnosis and management. American Family Direction 2010;16:14:14-14.	

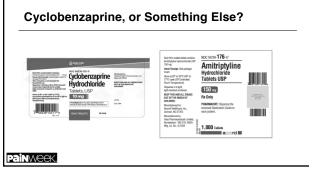
Antispasticity and Antispasmodic Agents	-
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Muscle Spasms	
	-
■Baclofen	
Tizanidine Other agents	
-Cyclobenzaprine, the TCA ?	-
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41	
Muscle Relaxants	
 Antispasticity agents Spasticity: upper motor neuron disorder characterized by muscle hypertonicity and 	
involuntary jerks - Multiple sclerosis, cerebral palsy, spinal cord injury	
Tizanidine Baclofen	
• Diazepam	
I. Cana K. et al. Pair Spayane Hasings 2304/281-0475. 2. Via Talue Herry, et al. Contrast Entirestay III A. 2. Via Talue Herry, et al. Contrast Entirestay III A. 2. Paramatistary 2004/2012/2013-11. 4. Ani Isaam Herr 2007 2012 1/10/17/481. 5. Ani Isaam Herr 2007 2012 1/10/17/481. 5. Nation Heart 2007 2012 1/10/17/481. 6. Lana Cana, Inc. (and Ough P) 6. Lana Cana, Inc. (and Ough P) 6. Lana Cana, Inc. (and Ough P) 6. Manne, (No. 117 20) 2015. 6. M	
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III. Centrally-acting agents (spasmolytic drugs) Muscle Relaxants (cont'd) Tizanidine ■ GABA analogue Agonist of α2 receptors (presynaptic) Selective GABA-B receptor agonist (↑
K+ conductance, ↓ Ca++ conductance)
 Muscle relaxant and analgesic (reduced) Reduces adrenergic input to alpha motor neurons ■ No effect on spinal cord reflex substance P) Less antihypertensive effect than ■ 5 mg PO TID, may titrate every 3 days clonidine ■2 to 8 mg PO TID ■ Max dose: 80 mg/day Max dose: 36 mg /day Adverse effects: somnolence, increased Side effects: hypotension, asthenia, elevated LFTs, hepatotoxicity seizure activity Painweek.

43

Muscle Relaxants (cont'd) • Antispasmodics - Primarily used for treatment of musculoskeletal conditions, such as back pain, sciatica, herniated discs, spinal stenosis, myofascial pain - Cyclobenzaprine - Metaxalone - Methocarbamol - Orphenadrine citrate - Carisoprodol | Carisoprodol

44



Conclusions

- Adjuvant and coanalgesics require judicious monitoring for safe use

 Extensive patient education regarding potential adverse effects is paramount

 Comorbid disease processes and concurrent medications may obscure adverse effects

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