

Rational Polypharmacy: An Update for Specific Conditions

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Disclosures	
■ Nothing to disclose	
Pain /Week	

In the news now...

Feds halt 2 Tennessee pharmacies' opioid dispensing for now

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https://apnews.com/fcae3106c7954369bf509 05b6639ab6b accessed 3.6.2019 https://www.deadiversion.usdoi.gov/21cfr/cfr/ 1306/1306 04.htm accessed 3.6.2019

Learning Objectives	
Define rational polypharmacy as it pertains to the patient in pain	
 Recognize the various pharmacological classes used in rational polypharmacy of migraine, neuropathic pain, and musculoskeletal pain conditions 	
Distinguish between rational and irrational polypharmacy in managing pain	
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How does rational polypharmacy apply to my	
practice?	
 Synergistic combinations decreasing the amount of opioid needed for pain control 	
Control	
 Using nonopioids as first line therapy can minimize or even prevent the need for opioid medications on a chronic basis 	
for opiola medications of a chronic basis	
 Shortages and regulatory constraints on the manufacture of opioids have lead to shortages and the inability of pharmacies to stock opioids and other 	
medications used in pain management	
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Definitions	
■ Polypharmacy:	
The use of two or more drugs together, usually to treat a single condition or disease	
Synergy:	
The cooperative action of two or more stimuli or drugs • Rational:	
Proceeding or derived from reason or based in reason	
Irrational: Not endowed with the faculty of reason	
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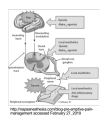
Goals of Rational Polypharmacy

- Minimize adverse effects
- -Lower doses of individual medications
- -Opioid sparing effects
- Increase adherence to the prescribed regimen
 Using synergistic combinations of medications to achieve improved outcomes compared to the individual medications
- Increase efficacy by utilizing long acting and short acting preparations

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Hitting the Target(s)

- Stimulation of nociceptors causes signal transduction to the dorsal horn —Transduction
- The spinothalamic tract transmits the signals to the brain where pain is first experienced
- -Transmission and perception
- Descending pathways from the brain attempt to block the signal from the periphery
 - -Modulation



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Medications Used in Pain Management

- Acetaminophen
- ■NSAIDs
- ■5HT_{3-1B/D} antagonists (Triptans)
- Calcitonin gene-related peptide antagonists
- Antidepressants
- Anticonvulsants
- Local anesthetics
- Skeletal muscle relaxants
- Opioids



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Acetaminophen

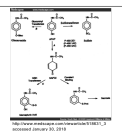
- Mechanism of action is still not entirely known
 Thought to be a partial COX inhibitor
- March 2014 FDA mandates all prescription drug combination products containing acetaminophen cap the dose at 325 mg
- Maximum daily dose limits vary based on comorbidities and who you ask
 —FDA vs Johnson and Johnson

http://www.fda.gov/drugs/drugsafety/informationbydrugclass/ucm165107.htm accessed January 30, 2018 https://www.tylenol.com/safety-dosing/usage/dosage-for-adults accessed January 30, 2018

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Acetaminophen (cont'd)

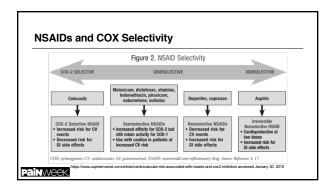
- Largest concern is unintentional overdoses
- Metabolism of acetaminophen by the liver is a saturable process
- Over the counter products and cumulative acetaminophen dosing

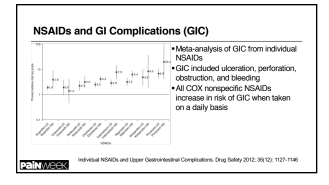


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Nonsteroidal Anti-Inflammatory Agents

- COX 1 more specific to the GI tract and renal homeostasis
- COX 2 more specific to inflammation and platelet aggregation
- Certain comorbidities limit the dosing on most NSAIDs
 - -Patients on anticoagulants
 - -Patients with renal dysfunction
 - -Pregnancy





Nonsteroidal Anti-Inflammatory Drugs

- Topical vs systemic NSAIDs
 - -Patch, cream, lotion, etc
- Range in application frequency from twice to four times daily

 Topical can provide NSAID relief at the site of inflammation without the systemic side effects
- -Cost can be a limiting factor
- -Still carry a black box warning on the labeling for cardiovascular complications

5HT_{3-1B/D} Antagonists (Triptans)

- Serotonin receptor antagonists leading to
- -Extra-cerebral vasoconstriction (5-HT_{1B})
- -Decreased inflammatory neuropeptide release (5-HT_{1D})
- Indicated for migraine treatment
 - -Abortive therapy, not prophylactic
- Dosing in general involves administration of a second dose in 1 to 2 hours if the first dose was unsuccessful in aborting the migraine

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Triptans (cont'd)							
Drug	Almotriptan	Eletriptan	Fravatriptan	Naratriotan	Ricatriptan	Sumatriotan'	Zolmitriotan
Brand Name (Manufacturer)	Axert (Janssen)	Relpax (Pfizer)	Frava (Endo)	Amerge (GSK)	Maxalt Maxalt MCT (Merck)	Imitrex (GSX) Crizetra Xsail (Avanir) Sumavel DosePro (Endo) Zembrace SymTouch (Promius)	Zomig Zomig ZMT (Impax)
Generic Available	Yes	No	Yes	Yes	Yes	Yes - for Imitrex products only	Yes - for oral tabs and ODTs only
Route of Adminstration	Oral	Oral	Oral	Oral	Oral	Oral: Nasal: SC	Oral; Nasal
Formulations	6.25, 12.5 mg tabs	20, 40 mg tabs	2.5 mg tabs	1, 2.5 mg tabs	5,10 mg tabs and 5,10 mg COTs	Imitive and genetics — Chal 2.5 6.100 mg tabs SC-4.6 mg/S FmL surfo-injector pen and refil carridag, visit? Nasal-5, 20 mg/G 1.m. c. nasal spray Onzeto Xasil 1 mg nasal powder caps Sumarel DosePto-6 mg/U 5 mL 5C needle -fee delivery system Zembrace SymTouch: 3 mg/U 5 mL, SC sudo-injector	Oral 2.5, 5 mg tabs and 2.5, 5 mg COTs Nasal 2.5, 5 mg/0.1 mi, nasal spray
Onset of Action	30-60 min	30-60 min	~ 2 hrs	1-3 hrs	30-60 min	Tabs: 30-60 min SC: ~10 min Nasal: 10-15 min	Tabs: 30-60 min Nasal: 10-15 min
Elimination Half-life	3-4 hrs	~4 hrs	~25 hrs	~6 hrs	2-3 hrs	~2 hrs	2-3 hrs
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Triptans (cont'd)

- Patients that are NOT candidates for triptan agents
 - -Ischemic heart disease
- -Uncontrolled hypertension
- -Peripheral vascular disease
 -History of cerebrovascular syndromes (stroke or transient ischemic attack)
- Multiple formulations exist for
- -Sumatriptan (nasal, SQ, oral)
- -Zolmatriptan (nasal and oral)

Calcitonin Gene-Related Peptide (CGRP) Antagonists

- Monoclonal antibodies that bind to CGRP
- -Preventing intracranial artery vasodilatation
- -Prevention of dural mast cell degranulation
- Indicated for the prevention of migraine
- Not indicated for the management of acute migraine symptoms
- Administration of the currently approved agents monthly subcutaneous injection

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AnnRevPharmacolTox.55.533-52 2015

CGRP Antagonists Currently Available

- Erenumab-aooe [Aimovig®]
- -Subcutaneous injection 70 mg once monthly
 -May increase to 70 mg twice a month in some patients
- Fremanezumab-vfrm [Ajovy®]

 —Subcutaneous injection 225 mg once monthly or 675 mg every three months
- Galcanezumab-gnlm [Emgality®]
 - -Subcutaneous injection 240 mg once then 120 mg monthly

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Lexicomp accessed 3.1.2019

CGRP Antagonists (cont'd)

- Questions that remain unanswered regarding their long term safety include
- -Hypertension
- -Nitric oxide synthase
- Platelet aggregation
 Negative impact on microvasculature
- ·Heart failure
- Diabetes



https://www.practicalpainmanagement.com/pain/h side-effects-cgrp-antagonists accessed 3.1.2019

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Tricyclic Antidepressants (TCA)	-
mcyclic Antidepressants (TCA)	
 Mechanism of action is through inhibition of norepinephrine and serotonin 	
reuptake and inhibition of sodium channel action potentials	
The antidepressant effects and the neuropathic pain analgesia are	
independent	
 Higher dosing and longer treatment time needed for antidepressant effects Caution should be exercised in patients 	
-With cardiac arrhythmias	
-Over the age of 65	
-Over the age of oo	
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Serotonin Norepinephrine Reuptake Inhibitors (SNRI)	
 Mechanism of action is through inhibition of norepinephrine and serotonin 	
reuptake	
 Dosing is generally higher for treating neuropathic pain compared to treating depression 	
Withdrawal syndromes can occur if patients are taken off SNRI therapy	
abruptly	
-Anxiety, irritability, headache, paresthesia, nervousness	
Caution should be exercised in patients with liver dysfunction, uncontrolled Caution should be exercised in patients with liver dysfunction, uncontrolled	
hypertension, or moderate cardiovascular disease	
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Antionilantias	
Antiepileptics	
■ The primary antiepileptics used in pain management work on calcium	
channels	
-Gabapentin	

-Pregabalin

-Phenytoin

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Carbamazepine for trigeminal neuralgia

Other antiepileptics have had mixed results regarding neuropathic pain
 –Valproic acid

Local Anesthetics	
Mechanism of action is through membrane stabilization of sodium channels	
preventing depolarization and signal transduction • Acute uses for local anesthesia (procedures, etc)	
-Topical application	
Cream, ointment, patch, etc Intradermal injections	
-Nerve blocks	
 Patches are indicated for the management of postherpetic neuralgia 	
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Skeletal Muscle Relaxants	
Multiple medications are included in this general taxonomy	
-Certain agents approved for spasticity	
Baclofen and tizanidine	
Others stand out for reasons other than their indication	
–Cyclobenzaprine and orphenadrine regarding their anticholinergic effects –Chlorzoxazone and potential for hepatotoxicity	
-Carisopradol and meprobamate and potential for abuse	
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Opioids	
Opioids work on multiple receptors within the CNS	
-Analgesia and adverse effects are derived from mostly mu receptors	
 There is no ceiling dose for analgesia; however, as doses increase the incidence of adverse effects increases 	
 CDC (2016) and VA/DoD (2017) guidelines outlining the use of opioids in chronic pain have been published 	
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Opioids (cont'd)

- Agonists vs partial agonists vs antagonists
 Morphine, fentanyl, methadone, etc
 - –Morphine, fentanyl, methadone, etc–Buprenorphine, nalbuphine, butorphanol
 - -Naloxone and naltrexone
- Awareness of other nonpain combination products

-Naltrexone-bupropion for weight loss

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Opioid Statistics

- Medication overdose deaths in 2016: 63,632
- Opioids (illicit and prescription)
 were involved in 66.4% of those fatalities
- Patients on > 90 morphine milligram equivalents have decreased from 11.5 to 5 per 100 patients in the US

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://www.cdc.gov/drugoverdose/pdf/pubs/2018-cdc-drug-surveilland

Opioid Statistics (cont'd) Lever feeting Links on Certain Opioid Preceptiques The control of t

Patients at Risk for Opioid Adverse Events

- Patients with sleep apnea and sleep disordered breathing
- Pregnancy
- Hepatic or renal dysfunction
- Age greater than 65
- Mental health or substance use disorders
- Nonfatal overdose history

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Opioid Metabolism • Metabolic pathways can become saturated leading to metabolism by other pathways - Codeine - Oxycodone • 2D6 → noroxycodone • 3A → oxymorphone **The policy of the saturated leading to metabolism by other pathways - Codeine - Oxycodone • 3A → oxymorphone

Immediate Release (IR) vs Extended Release (ER)	
 Initial therapy should include the use of IR formulations ER preparations are appropriate for patients 	
That routinely use the IR preparation with relief of pain	
 That are not experiencing adverse effects that decrease quality of life That are on stable doses of IR preparations and have been for an 	
appropriate time frame	
 IR and ER preparation use should be re-evaluated for safety and efficacy periodically or per state guideline 	
portourour, or per dutie gardonnie	
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Nonrational Polypharmacy	
• Utilizing two medications in the same family for the same condition	
-lbuprofen and naproxen	
-Morphine immediate release and oxycodone immediate release	
 Adding a medication that may be contraindicated based on the patients other 	
comorbidities -Methadone use in a patient with a history of QTc prolongation	
-Tramadol or use in a patient with underlying seizure history	
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Petionelizing Migraine Pain Management	
Rationalizing Migraine Pain Management	
 Use of abortive medications at the beginning of a migraine NSAIDs, triptans 	
-Opioids and dopamine antagonists (severe)	
allog of groups to die they are got onto most evitorie	
 Use of prophylactic therapy once patients meet criteria More than two migraines per month 	
-Migraine lasts for more then 24 hours	
-Use of abortive therapy more than twice per week	
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ts (nortriptyline, protriptyline)
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Rationalizing Neuropathic Pain

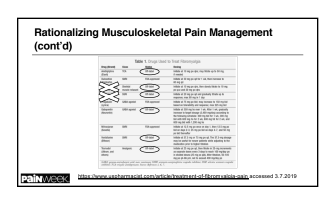
- Scheduled use of tricyclic or SNRI antidepressants at appropriate doses

 -Caution regarding the use of anticholinergic tricyclic agents
- Use of antiepileptics at appropriate doses
- -Opioids may be used in combination with the use of an antiepileptic
 -Topical local anesthetics such as patches and creams with the above

NSAIDs and acetaminophen are unlikely	Topical Agents Lidecaine Apply to mod points are of extension. Op to 5 publies Pruntes dynamical may be assisted in a single assistant and an analysis and apply and apply and apply and apply and apply apply and apply and apply apply and apply and apply and apply apply apply apply apply apply and apply apply apply and apply			
to alleviate neuropathic pain	Capusion 8% patch	remain in place for up to 12 hours in any 24 hour period. Apply to mode painful area of index sin no 68 minutes, then senous; up to 4 pathess may be applied in a single application. May repeat or snow than every 3 months upon mitum of pain. Assa should be pertiased with a topical assertities; prior to pathe, application.	erythema, depigmentation Erythema, pain, hypertemion (barrelent), pruntus, nausea, voniding, papules, edema, nacopharmyngidis	
		Tricyclic Antidepressants (TCRs)		
Anticonvulsants, local anesthetics, and ricyclic antidepressants are mainstays in	Sonippine*	10-25 mg at befilme: increase dose by 25 regides weekly it infended, usual membranice dose 1's mights as a signific befilme dose or 2 divided disses; maximum 12's regidey.	Ory mouth, condigation, anhythmic tend likely TCA to cause orthestatic hypotensius; less likely TCA to cause cognitive imporment, cause cognitive imporment, and donor, and antichologous; effects	
neuropathic pain management		10-25 reptiley increase every 3 days as necessary until decined effect achieved, usual effective door. 50-150 registey, maximum dose 150 registey.	Dry-mouth, condigation, anhythmia, cognitive impairment, orthostatic hypotension; less likely TCA to cause sedation and anticholinergic effects	
		Articonvoluerts		
 Opioids may have a place but not first or second line 	Subspentin	City 1: 300 mg, day 2: 300 mg brice daily, day 2: 300 mg 3 times/day, dose may be titoded as needed for pain relief (range: 1,800-3,800 mg/day); daily doses >1,800 mg/d o sid generally show grader benefit	Distiness, attacks, sommolence, faligue, peripheral edems, impaired cognitive function	
	Pregubalin	150 mg/bby in divided doses (75 mg twice daily or 50 mg 3 times daily; may be increased to 306 mg/bb; within 1 week to 100 mg/bb; within 1 week 2-4 weeks if follooks (Maximum dose 500 mg/bby after 2-4 weeks if follooks (Maximum dose 500 mg/bby	Dictiness, ataxia, sommolence, impaire cognitive function, peripheral edema, headache	
		Systemic Analgenics		
Muscle relaxants are controversial in terms of efficacy	Orycodone (spinit) Regular or immedial- velesse farmulation (disaper given for morphile equivalent)	2.5-15 mg every 4 hours as needed. After 1-2 medis, convert bibli duly desage to temp-acting opinid analysis and continue short-acting agent as needed	Comfigation, rauses, sonnelence, impaired cognitive function, falls	
	Tramadol Inmediate-missor formulation*	50-100 mg every 4-6 hours; maximum dasc 400 mg/day derate: 25 mg once daily, increase 25-50 mg/day in divided doses every 3-7 days as tolerabet; maximum 200 mg/day in collected over 25 vars of soe	Conditation, rauses, distriess, headache, somnolence, vomiting, pruritue, insonnia, orthostesis, tells	

Rationalizing Musculoskeletal Pain Management ■ Bone pain ■ Muscle pain ■ Tendon and ligament pain ■ Fibromyalgia ■ Joint pain ■ Nerve compression syndromes ■ More than 150 diagnoses all of which affect the locomotor system https://pmi.bmi.com/content/79/937/827





Conclusion	
Contolacion	

- Pain management typically involves more than one modality in order to manage
- Safety must take into consideration patient specific factors that will change over time
- Certain combinations can put patients at risk for adverse effects but having a complete picture of a patients medications can help prevent this

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See you at PAINWEEK

