



Rational Polypharmacy in Pain Management

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Disclosures

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Learning Objectives

- Define rational polypharmacy and when it is indicated for pain management
- List the array of medications and their MOAs that may be employed in polypharmacy
- Discuss the pharmacologic and clinical considerations of which the prescriber should be aware
- Explain painful conditions where polypharmacy might be considered



Real Patients to Consider in Our Discussion

- 60 year old male experiencing painful diabetic neuropathy and chronic LBP
- 50 year old female who experiences fibromyalgia and migraine
- 70 year old female who experiences osteoporosis, osteoarthritis, and postherpetic neuralgia
- 52 year old male who experiences post-laminectomy pain
- 28 year old female with chronic migraine



Selecting an Analgesic: Evidence and Guideline Limitations

- Paucity of trials on comparative effectiveness of different treatments
- Most treatment trials are of short duration with limited evidence for functional benefit
- Few trials evaluate strategies for choosing initial agent
- Various clinical practice guidelines may interpret evidence differently
- Clinical practice guidelines may not include latest evidence
 - Duloxetine for low back pain or osteoarthritis

Chappell AS, et al. Pain. 2009;146:253-260; Chappell AS, et al. Pain Pract. 2011;11:33-41; Kroenke K, et al. Gen Hosp Psychiatry. 2009;31:206-215; Skjarevski V, et al. Pain Med. 2010;11:648-657; Skjarevski V, et al. J Pain. 2010;11:1262-1280; Skjarevski V, et al. Spine (Phila Pa 1976). 2010;35:E579-582; Wallace M, et al. Expert Rev Neurother. 2011;11:15-27.



What Is Polypharmacy?

- Defined as
 - Intentional use of 2 or more medications to treat 1 condition
 - Example: opioids + NSAIDs to treat low back pain, OR
 - Use of 2 or more medications by 1 patient to treat multiple conditions
 - Example: antihypertensives, antidepressants, NSAIDs, and statins, OR
 - Use of 2 or more agents of the same chemical class
- Polypharmacy should be minimized whenever possible; however, it may be warranted under certain circumstances



Rational Polypharmacy

- Rational polypharmacy
 - May help reduce chronic pain
 - Reduction of chronic pain can improve patient outcomes



Rational Polypharmacy (cont'd)

- Rational polypharmacy has become an acceptable component of chronic pain management
 - More therapeutic options are being made available
- Useful to target pain that has peripheral and central mechanisms
- Consider the following when selecting combined regimens:
 - Side effects (SEs)
 - Medication interactions
 - Ease of use
 - Costs



Why Is Rational Polypharmacy Used?

- Patients with pain
 - May experience only a partial response to monotherapy
 - Cannot tolerate adverse events at analgesic doses of monotherapy
 - May have positive synergistic effects with combined agents from different medication classes



Multidrug Therapy Proposed Principles for Chronic Pain

- Purpose is to combine medications to achieve additive or synergistic analgesia
 - Potentially at lower doses (and fewer side effects) than those required with monotherapy
- Combine medications with differing mechanisms or sites of action, based on patient response, functional goals, clinical experience, and potential adverse interactions
- Consider interactions of newly added drug with current medications
- Select and use one drug at a time
- Start low, go slow for dosing and titration, particularly in older patients
- Ongoing reassessment is critical
 - Assess for clinically meaningful relief and document functional outcomes



Backonja MM, et al. Curr Pain Headache Rep. 2006;10:34-38; Gilron I, et al. Lancet. 2009;374:1252-1261.

The Chronic Pain PHARMACOLOGIC Armamentarium

- Nonopioids
 - Acetaminophen
 - NSAIDs
 - COX-2 inhibitors
 - Antidepressants
 - Anticonvulsants
 - Topical agents, other
- Opioids
 - Mu-opioid agonist
 - Partial agonists



Nonopioid Analgesics*

Chemical Class/Examples	Class Examples	Brands/Examples
Para-aminophenols	Acetaminophen	Tylenol®
Salicylates	Aspirin Choline magnesium trisalicylate (CMT) Choline salicylate Magnesium salicylate Diflunisal	Boyer® Bufferin® Trilisate® Arthropan® Doan's® Asgesic® Salgesic® Dolobid®
* Not an exhaustive list of class/examples.		



Nonopioid Analgesics (cont'd)

Chemical Class	Class Examples	Brands
Arylpropionic/propionic derivatives	Ibuprofen Naproxen Ketoprofen Flurbiprofen Fenoprofen Oxaprozin	Advil® Motrin® Aleve® Anaprox® Naprelan® Naprosyn® Orudis® Oruvail® Ansaid® Nalfon® Daypro®
Indole and indene acetic acids	Indomethacin	Indocin® Indocin® SR



Clinical Indications: Nonopioids

- Variety of acute and chronic pain types
 - Eg, trauma, post-op, cancer, arthritis
- Somatic pain
 - Muscle and joint pain, bone/dental pain, inflammatory pain, post-op pain
- APAP vs NSAIDs
 - Acetaminophen has analgesic, antipyretic effects
 - But lacks anti-inflammatory effect
 - NSAIDs have analgesic, anti-inflammatory, and antipyretic effects
 - But affect gastric mucosa, platelets

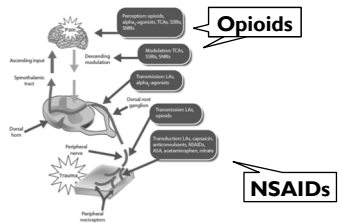


Clinical Indications: Opioids

- Moderate-severe pain unresponsive to nonopioids alone
 - Example: acute pain
 - Post-op, trauma
 - Breakthrough pain
 - Cancer pain
 - Chronic, noncancer pain



**Mechanism of Action:
Opioids + NSAIDs**



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**Adjuvant Analgesics:
Tricyclic Antidepressants**

- **Examples**
 - TCAs include amitriptyline, desipramine, doxepin, imipramine, nortriptyline
- **MOA**
 - Inhibition of reuptake of norepinephrine and serotonin
 - Analgesia is independent of antidepressant function
- **Uses**
 - Chronic pain examples: migraine, other headaches, low back pain, cancer pain, fibromyalgia
 - Neuropathic pain examples: PDN, PHN, cancer-related pain
 - Common adverse events (AEs)
 - Examples: sedation, orthostatic hypotension and anticholinergic effects (ie, dry mouth, blurred vision, constipation, urinary retention)

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**Adjuvant Analgesics:
Selective Serotonin Reuptake Inhibitors**

- **Examples**
 - SSRIs include citalopram, paroxetine, fluoxetine, sertraline
- **MOA**
 - Selectively inhibit 5-HT reuptake without affecting norepinephrine
- **Uses**
 - Examples: neuropathic pain, diabetic neuropathy
- **Common AEs**
 - Examples: anxiety, insomnia, nausea, headache, drowsiness, sexual dysfunction, withdrawal symptoms upon abrupt cessation

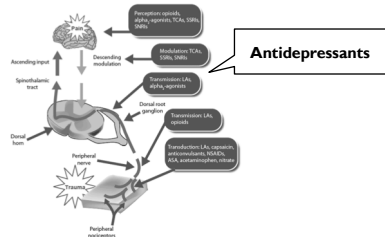
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Adjuvant Analgesics: Serotonin/Norepinephrine Reuptake Inhibitors

- Examples
 - SNRIs include duloxetine and venlafaxine
- MOA
 - Block reuptake of 5-HT and norepinephrine
- Uses
 - Example: diabetic peripheral neuropathy (DPN)
- Common AEs
 - Examples: nausea, somnolence, dizziness, constipation, dry mouth, hyperhidrosis, anorexia



Mechanism of Action: Antidepressants



Adjuvant Analgesics: Anticonvulsants

- Examples
 - AEDs include gabapentin, pregabalin, carbamazepine, phenytoin, divalproex sodium, clonazepam, levetiracetam, topiramate, lamotrigine
- MOA: exact mechanism of analgesic effect is unknown; it is thought they
 - Reduce membrane excitability
 - Suppress abnormal discharges in pathologically altered neurons
- Uses
 - Neuropathic pain (PDN, PHN, RSD, PSP, TN)
 - Cancer pain, HIV-related neuropathy, phantom limb pain
 - Migraine (prophylaxis), dysesthesia, deafferentation pain, thalamic pain
- Common AEs
 - Sedation, headache, dizziness, rash, vertigo, ataxia, nausea, diplopia



Adjuvant Analgesics: Topicals

- Examples
 - Lidocaine, Lidoderm, EMLA®, Capsaicin cream
- MOA
 - Lidocaine/prilocaine: block sodium channels and inhibit generation of abnormal impulses by damaged nerves
 - Capsaicin: depletion of substance P in sensory nerve endings
- Uses
 - Examples in acute and chronic pain
 - PHN, other neuropathic pain, mechanical allodynia
 - Pain associated with medical procedures: needle insertion, cannulation, epidural nerve blocks
- Common AEs
 - Examples: localized reaction including burning sensation



The Potential Benefits of Rational Polypharmacy

- Enhance current treatment
- Use a lower dose of a medication
- Target symptom clusters (eg, pain and depression)
- Ease the treatment of a comorbid condition (eg, control diabetes to reduce DPNP)
- Address different pain mechanisms (eg, central and peripheral mechanisms)
- Treat AEs



Considerations for Rational Polypharmacy

- Know medication toxicities
- Avoid overlapping/additive toxicities
- Know medication MOAs
- Know medication PK/PD
 - Avoid drug-drug interactions
- Have convincing evidence that the combination is more effective vs monotherapy and should not pose significantly greater safety or tolerability risks



MOA Considerations for Polypharmacy

- When using multiple medications, consider carefully the MOA of each drug:
 - Each drug should have one MOA
 - Drugs should not have broad-acting MOA
 - Drugs should not have the same MOA
 - Drugs should not have opposing MOAs



Possible Drug-Drug Interactions

- Interaction of absorption: one drug may cause an increase/decrease in the absorption of the other in the GI system
- Interaction of protein binding
- Interaction of metabolism (eg, CYPs)
- Interaction of receptor binding
- Interaction of therapeutic action



Types of Drug-Drug Interactions

- Additive
- Synergistic
- Potentiation
- Antagonism
 - Functional/physiological
 - Chemical/inactivation
 - Dispositional
 - Receptor



P450 Enzymes

- Care should be taken when coadministering drugs whose metabolism might be inhibited by other drugs in order to prevent adverse drug reactions (ADRs)
 - For example, SSRIs inhibit the metabolism of drugs mediated by certain P450 enzymes
- > 30 human CYP isoenzymes have been identified
- ≥ 90% of drug oxidation can be attributed to 6 main P450 cytochromes:
 - 1A2
 - 2C9
 - 2C19
 - 2D6
 - 2E1
 - 3A4

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Phases of Medication Metabolism

- Phase I: oxidation/reduction/hydrolysis
 - Oxidation to the parent compound or deletion of the alkyl group, reduction, and hydrolysis reactions
- Phase II: conjugation
 - Biotransformation links a parent medication molecule or product of Phase I metabolism with an endogenous substrate (eg, glucuronic acid, sulfate, or glycine)

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Induction and Inhibition

- Induction
 - Increase of enzyme metabolism by a medication
 - Increasing doses are needed to produce same effect, as the body metabolizes the drug more quickly
- Inhibition
 - Decreased enzyme activity due to direct interaction with a medication or chemical
 - Doses should be decreased due to a decrease in metabolism

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When Is Polypharmacy Indicated in Pain Management?

- To reduce medication intolerance by using a second medication that allows a lower dose of the first
 - May increase treatment compliance
- To provide analgesic efficacy at certain times of the day by giving immediate-release with long-acting medications
 - Example: control breakthrough pain in a patient taking long-acting opioids



When Is Polypharmacy Indicated in Pain Management? (cont'd)

- To use a lower dose of a medication by using a second medication
 - Example: opioid-sparing
- To address partial or nonresponse to 1 medication by adding a second medication to increase efficacy
 - Example: using 2 different antidepressants with different MOAs



When Is Polypharmacy Indicated in Pain Management? (cont'd)

- To target different symptom clusters that are a product of the disease or of the comorbid disease
 - Example: pain with associated depression, which in turn is associated with suicidal ideation
- To treat the comorbid disease with ease by aggressively treating the index disease
 - Example: treat diabetes aggressively, thereby reducing the peripheral neuropathy severity



When Is Polypharmacy Indicated in Pain Management? (cont'd)

- To address different locations of the disease process
 - Example: pain with peripheral AND central mechanisms that require medications with peripheral and central activity
 - Example: topical lidocaine patch with antidepressant
- To treat an adverse event



Checklist for Controlling Pain in a Polypharmacy Environment

- Prescribers are charged with
 - Prudent attention to the patient's past medication history, including OTC preparations
 - Vigilant surveillance of systemic function
 - Pharmacologic alternatives when medications are inappropriate



5 Principles of Polypharmacy for Pain Associated Comorbidity

- Use medications for comorbid disease with proven analgesic efficacy
- First target symptoms should ALWAYS be pain
- Target all possible pain mechanisms (eg, peripheral and central) believed to be causing the pain
- Do not aim for absolute pain relief
 - Aim for tolerable pain levels that improve QoL or function
- Use medications to address more than 1 comorbidity
 - Example: sedating antidepressant for pain, sleep, and depression



Summary of Implementing Rational Polypharmacy in the Treatment of Pain

- Polypharmacy, the use of multiple medications in a patient, should be minimized whenever possible; however, it may be warranted under certain circumstances
- Rational polypharmacy may be employed when the benefits outweigh the risks



Summary of Implementing Rational Polypharmacy in the Treatment of Pain (cont'd)

- The benefits of rational polypharmacy include:
 - Enhancing current treatment
 - Using a lower dose of a medication
 - Targeting symptom clusters
 - Easing the treatment of a comorbid condition
 - Addressing different pain mechanisms
 - Treating AEs

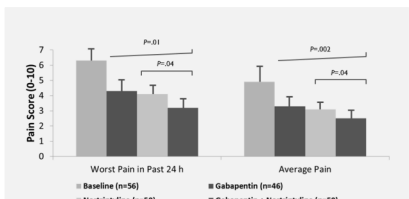


Examples of "Rational" Polypharmacy in Specific Pain Conditions

- Neuropathic pain
- Fibromyalgia
- Chronic headache
- Low back pain



Nortriptyline and Gabapentin for Neuropathic Pain



- Significant improvements were also seen with combination therapy for pain interference with mood, sleep, and enjoyment of life
- Combination therapy did not result in a markedly higher side effect burden



Gilron I, et al. Lancet. 2009;374:1252-1261.

Morphine, Gabapentin or their Combination for Neuropathic Pain

- Patients with postherpetic neuralgia or painful diabetic neuropathy
- Randomized, double-blind, active placebo-controlled, 4 period crossover trial
- Periods included active placebo (lorazepam), gabapentin, sustained release morphine, or a combination of both gabapentin and morphine each period 5 weeks
- Gabapentin and morphine combined achieved better analgesia at lower doses of each drug than either as a single agent



Postherpetic Neuralgia: 5% Lidocaine Medicated Plaster, Pregabalin, or a Combination of Both?

- Randomized, open label clinical effectiveness trial
- PHN patients with pain intensity score of greater than 4 were randomized to lidocaine plaster or pregabalin
- Patients experiencing pain intensity of 4 or less after 4 weeks remained on monotherapy
- Those who did not received both medications after 4 weeks
- Equal response between medications with monotherapy AND combining medications was well tolerated and improved response



Combination of Morphine with Nortriptyline for Neuropathic Pain

- Combination compared to monotherapy of each in patients with neuropathic pain (1:1:1)
- 3 6-week treatment periods
- Superior efficacy noted with morphine-nortriptyline combination over either monotherapy
- Constipation, dry mouth, and somnolence most frequent adverse effects



Pregabalin with Duloxetine for Fibromyalgia

- Randomized, double-blind 4 period crossover design
- 6 week periods
- Placebo, pregabalin, duloxetine, or combination
- Daily pain intensities improved most with combination
- Fibromyalgia Impact Questionnaire scores improved most with combination
- SF-36 scores improved most with combination



MIGRAINE Preventive Medications

- Anticonvulsants
 - Divalproex*
 - Gabapentin
 - Topiramate*
- Antidepressants
 - TCAs, SSRIs, MAOIs
- β-Blockers
 - Propranolol*/Timolol*
- Ca channel blockers
 - Verapamil
- NSAIDs
- 5-HT antagonists
 - Methysergide*/methergine
- Neurotoxins
 - Onabotulinum toxin A (Botox)*
- Angiotensin system
 - ACE inhibitors
 - Antagonists
- Acetylcholinesterase inhibitors?
- Other
 - Riboflavin, coenzyme Q10, Feverfew, Petasites

FDA approved.
 ACE, angiotensin converting enzyme; MAOIs, Monoamine oxidase inhibitors; SSRI, selective serotonin reuptake inhibitors; TCA, tricyclic antidepressant.
 Modi S, et al. *Am Fam Physician*. 2006;73:172-78; Nicolodi M, et al. *Headache*. 2002;42:596-602; Silberstein SD, et al. *Cephalalgia*. 2002;22:491-512.



Migraine Comorbid Conditions

- Cardiovascular
 - Heart attack/angina
 - Mitral valve prolapse
 - Hypertension or hypotension
 - Stroke
 - Raynaud's syndrome
- Mood disorders
 - Depression
 - Mania
 - Anxiety
 - Panic

Silberstein SD, et al. *Cephalalgia*. 2002;22:491-512



Silberstein SD, et al. *Cephalalgia*. 2002;22:491-512.

Migraine Comorbid Conditions (cont'd)

- CNS
 - Epilepsy
 - Essential tremor
 - Fibromyalgia
- GI disorders
 - Ulcer disease
 - Colitis
 - Irritable bowel syndrome

• Allergy/Asthma

CNS, central nervous system; GI, gastrointestinal. Silberstein SD, et al. *Cephalalgia*. 2002;22:491-512.



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Drug Treatment

- Assess coexisting conditions
- Do not treat migraine with drug contraindicated for other condition
- Do not use drug for other condition that exacerbates migraine
- Beware of drug interactions
- Pay attention to women of childbearing potential
- Select 1 drug to treat both disorders?

Silberstein SD, et al. *Headache*. 2007;47:585-599.



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Comorbid and Coexisting Disorders Monotherapy?

- Preferred, but may be exception rather than rule
 - May not be best choice for either disorder
 - β -blocker not first choice for hypertension
 - Dose for one may not be adequate for second
 - TCA migraine dose too low for depression

Silberstein SD, et al. *Headache*. 2007;47:40:585-599.
PainWeek Silberstein SD, et al. *Headache*. 2007;47:40:585-599.

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Comorbid and Coexisting Disorders Monotherapy? (cont'd)

- Therapeutic opportunities
 - Angina: β -blocker
 - Epilepsy: divalproex or topiramate
- Therapeutic limitations
 - Depression or asthma: avoid β -blockers
 - Epilepsy: caution with TCAs or neuroleptics

Silberstein SD, et al. *Headache*. 2007;47:40:585-599. Silberstein SD, et al. *Headache*. 2007;47:40:585-599. 740:585-599.

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Comorbid and Coexisting Disease Therapeutic Independence

- Treat each disorder with best drug
 - Benefits: use first-line drug for each disorder at correct dose
 - Less adverse effects?
 - Concerns
 - Drug interactions or more adverse effects
 - Contraindicated for one of the disorders

Silberstein SD, et al. *Headache*. 2007;47:40:585-599.

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Comorbid and Coexisting Disease Therapeutic Independence (cont'd)

- Examples
 - Depression: SSRI or SNRI plus AED (divalproex or topiramate)
 - Hypertension: ACE inhibitor or antagonists plus AED or TCA

AED, antiepileptic drug; SNRI, serotonin-norepinephrine reuptake inhibitor; SSRI, selective serotonin reuptake inhibitor.
Silberstein SD, et al. Headache. 2007;47(4):585-599.



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Comorbid and Coexisting Disease Depression

- Migraine and depression comorbid
- TCAs often recommended for migraine
 - However, appropriate dose for depression often higher
 - More adverse effects

Silberstein SD, et al. Headache. 2007;47(4):585-599.



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Comorbid and Coexisting Disease Depression (cont'd)

- Better approach
 - Treat depression with SSRI or SNRI and
 - Treat migraine with AED (divalproex or topiramate), β -blocker, Ca channel blocker, or even low-dose TCA

Silberstein SD, et al. Headache. 2007;47:585-599.



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Tizanidine and ibuprofen in acute low back pain

- Patients with acute low back pain randomized to receive either tizanidine 4 mg po 3 times daily with ibuprofen 400 mg 3 times daily or placebo plus ibuprofen 3 times daily
- Earlier improvement occurred in combination group, significantly better than ibuprofen alone by day 3
- More GI side effects noted with ibuprofen alone group supporting animal data that tizanidine can reduce GI side effects from NSAIDs



Conclusions

- Rational use of polypharmacy is indicated for pain management especially in specific painful conditions
- The prescriber should survey the array of medications and their MOAs that may be employed in polypharmacy
- Available studies suggest that rational approaches to polypharmacy in pain management can lead to improved analgesia and greater treatment tolerability



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