When Addiction Hurts:
Managing Acute Pain in Patients on Medications for Opioid Use Disorder (MOUD)

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Conflict of Interest Disclosure

• Author’s conflicts of interest:
  • Michelle Meyer, no conflict of interest
  • Andrea Wetshtein, no conflict of interest
Objectives

- Review the pharmacology of currently available MOUD options
- Discuss options for treating acute pain with opioids in patients on MOUD
- Describe non-opioid options and dosing for treating pain in patients on MOUD
Medications for Opioid Use Disorder (MOUD)

- The use of medications in **COMBINATION** with counseling and behavioral therapies for the treatment of substance use disorders
- Three current medications approved for use as MOUD
  1. Methadone
  2. Buprenorphine
  3. Naltrexone
Methadone

• Brand names:
  – Dolophine, Methadose

• Class:
  – Full Mu agonist and NMDA antagonist

• Use:
  – To reduce cravings and withdrawal symptoms from opioids

• Dosage Forms:
  – PO Tablets
  – PO Solution
  – IV Solution
Methadone

• Advantages:
  – Slows brain uptake and reduces euphoria in oral dosing
  – Beneficial in patients finding no response to other MOUD medications

• Disadvantages:
  – Only available through outpatient treatment programs which MOST patients must visit daily

• Clinical Pearls:
  – Start at no more than 30 mg total daily dose
  – QTc prolonging agent—watch for drug interactions
  – Use with caution in patients with liver dysfunction
Buprenorphine

• Brand Names:
  – Subutex, Suboxone, Zubsolv, Sublocade
• Class:
  – Partial Mu agonist
• Use:
  – To reduce cravings and withdrawal symptoms from opioids
• Dosage Forms:
  – Sublingual film
  – Sublingual tablet
  – Subcutaneous injection
  – Topical patch (for pain management only)
Buprenorphine

• Advantages:
  – Wider availability than methadone, patients can take at home

• Disadvantages:
  – Plain buprenorphine product has high abuse potential
  – High $$$ to patients

• Clinical Pearls:
  – Use plain buprenorphine product in pregnant women
  – Takes up to 20 min for sublingual tablet to dissolve
  – Depot injection can remain in the plasma for > 12 months after reaching steady state
Naltrexone

- **Brand Names:**
  - ReVia, Vivitrol
- **Class:**
  - Full Mu antagonist
- **Use:**
  - To diminish the reinforcing effects of opioids
- **Dosage Forms:**
  - PO tablets
  - Depot Injection
**Naltrexone**

- **Advantages:**
  - Does not result in physical dependence
  - Non-sedating

- **Disadvantages:**
  - Poor patient compliance (with PO tabs)
  - Management of acute pain crisis can pose a challenge

- **Clinical Pearls:**
  - Initiation requires prolonged abstinence – 7 days
  - Must stop PO naltrexone 48 to 72H before surgery in order for opioids to be effective
  - Within the first 14 days of depot injection, unable to overcome the mu receptor blockade for pain control
Availability of MOUD

**Methadone** = clinic
- Dosed in clinic
- Take home bottles
- Do not submit data to PMP

**Buprenorphine** = DATA2000 waiver (XDEA)
- Can be filled in multiple day prescription
- Dosing can be verified in state PMP

**Naltrexone (REMS)**
- No prescriptive restrictions
- Increasing court mandated use
- May not show up on ESI as frequently given in the office setting
Clinical Pearls for patients on MOUD

- Include the patient in the plan
- Utilize multi-modal analgesia for pain
- Require opioid tolerant doses to be effective
Patient with SUD AND Pain

Long-Acting Opioids

Short Acting Opioids

Symptomatic Medications
UNDER TREATMENT OF PAIN IN OPIOID USE DISORDER PATIENTS INCREASES INCIDENCE OF RELAPSE
Approaches to Acute Pain Crisis
Approaches to Pain Crisis in Patients on MOUD

• General Considerations
  – Maximize non-opioid therapies
  – Patients on MOUD typically require higher doses of opioids to compensate for tolerance and/or effect of their MOUD therapy
  – Try to avoid morphine as it is undistinguishable from heroin on a UDS
  – Transition to oral opioids as soon as clinically appropriate
    • IF patient has nausea consider utilizing sublingual opioids
  – Identify and avoid a patient’s previous drug of abuse and avoid as this may aggravate their addiction disorder

ALWAYS include the patient on their pain crisis plan
Pain Crisis: Sublingual/Buccal Buprenorphine

- Buprenorphine products are not required to be stopped prior to surgery
- SL films have greater bioavailability than buccal tablets
  - Monitor for over or under dosing when switching between products
- Approaches for pain management for SL buprenorphine products:
  1) Maintain outpatient dose of Buprenorphine OR Buprenorphine/Naloxone and split home regimen into divided doses given every 4-6 Hours +/- an opioid at a tolerant dose
     -- OR --
  2) Maintain outpatient dose schedule of Buprenorphine OR Buprenorphine/Naloxone +/- an opioid at a tolerant dose
     -- OR --
  3) Discontinue Buprenorphine/Naloxone and add an opioid at a tolerant dose
Pain Crisis: Subcutaneous Buprenorphine

- The depot can be “cut out” within 14 days of administration if necessary.

- Approaches for pain management for subcutaneous buprenorphine:
  - Maximize non-opioid therapy
  - Add opioids at tolerant dosing if non-opioids ineffective

- Patients will require opioid tolerant dosing for pain control if they have received an injection within the past 6 months.
Pain Crisis: Methadone

- Confirm home dose and when last dose given with methadone clinic
- Approaches to pain management with methadone:
  - Split home dose of methadone Q8H
    - Duration of analgesia of methadone is 6-8H
  --OR--
  - Continue home DAILY dose of methadone
  - May consider adding opioids at a tolerant dose for either approach

- **NOTE:** If patient is enrolled in a methadone clinic, they will have to provide evidence of continued administration of methadone during their hospitalization in able to remain enrolled in the outpatient program
Pain Crisis: ORAL Naltrexone

• Stop/Suspend use of oral naltrexone
  – Recommend to stop 72 hours prior to any PLANNED procedures
• Add opioids at an opioid tolerant dose

Other Considerations:
  – Regional anesthetic techniques
  – Ketamine
    • Typically dosed @ 0.1mg/kg/hr

• NOTE: A 7-10 day opioid free period is recommended before naltrexone can be resumed
Pain Crisis: DEPOT Naltrexone

- Determine when the last naltrexone injection was administered
- 0 to 14 days
  - Maximize adjuvant therapies
  - Regional anesthetic techniques
  - Continuous ketamine infusion at 0.1mg/kg/hr
  - Opioids are ineffective during this time frame
- 15 to 28 days:
  - All recommendations for 0-14 days +/- opioids at a tolerant dose

- **NOTE:** Coordination with naltrexone provider is key
Multimodal Therapies to the Rescue!
Rational Polypharmacy

Mechanism-Specific Treatment
Multiple targets…

Brain

Descending Inhibition
(NE, 5HT)

TCA
SNRI
Tramadol
Opioids

Central Sensitization
(Ca\(^{2+}\) channels, NMDA receptors)

Peripheral Sensitization
(Na\(^{+}\) channels)

NSAIDs
Opioids
TCA
Lidocaine

Spinal Cord

TCA
Gabapentin
Opioids
Ketamine

PNS
Multimodal Therapies

- Inpatient titration of anti-epileptic drugs can be more aggressive with balancing of sedation-caution if renal impairment
- Schedule adjunct meds to maximize effects
- Pre-operative cocktails could include APAP, anti-epileptic, NSAID, epidural or local block
- Non-pharmacological treatments-heat/ice, massage, healing touch, physical therapy
IV Acetaminophen

- Higher plasma concentrations of APAP when given IV vs oral
- Levels quickly become equivalent to oral administration
- Dose oral APAP 1 hr prior to surgery
# IV APAP Literature Recap

<table>
<thead>
<tr>
<th>Hickman et. al.</th>
<th>IV (n = 241)</th>
<th>PO (n = 245)</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Median opioid use first 24hr*</td>
<td>21.7 mg</td>
<td>21.7 mg</td>
<td>0.60</td>
</tr>
<tr>
<td>Median time to first pain med</td>
<td>41 mins</td>
<td>38 mins</td>
<td>0.90</td>
</tr>
<tr>
<td>Median time to ambulation</td>
<td>18.8 hrs</td>
<td>18.5 hrs</td>
<td>0.54</td>
</tr>
<tr>
<td>Median PACU LOS</td>
<td>2.1 hrs</td>
<td>2.2 hrs</td>
<td>0.17</td>
</tr>
<tr>
<td>Median hospital LOS</td>
<td>58.5 hrs</td>
<td>58 hrs</td>
<td>0.63</td>
</tr>
<tr>
<td>Post-op nausea</td>
<td>21.2 %</td>
<td>21.6 %</td>
<td>0.91</td>
</tr>
<tr>
<td>Post-op vomiting</td>
<td>7.9 %</td>
<td>6.9 %</td>
<td>0.73</td>
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</table>

*Opioid use in morphine mg equivalents.
PACU=post-op anesthesia care unit, LOS=length of stay

Liposomal Bupivacaine (Exparel®)

Mean Pain Score POD3

Opioid Use

LOS

Value Based Analgesia

- **IV Acetaminophen**
  - No published trials demonstrating outcome differences when compared to oral APAP
  - Cost of 975mg oral APAP=$0.06 vs IV 1000mg APAP=$48.18

- **Liposomal Bupivacaine**
  - Published data shows no significant difference in outcomes when compared to active comparators
  - Cost of bupivacaine 0.5% with epinephrine=$1.77 vs liposomal bupivacaine $301.49

Choose your patients thoughtfully with high cost/low value adding medications
Ketamine

- **Mechanism of Analgesia**: NMDA antagonism
  - Some effects on μ-opioid receptors, muscarinic receptors, monoaminergic receptors, γ-aminobutyric acid receptors, and several others

- **Analgesic, anti-inflammatory, and anti-hyperalgesic effects**

<table>
<thead>
<tr>
<th>Dosing For Pain</th>
<th>Dosing For Anesthesia</th>
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<tr>
<td><strong>Intravenous</strong></td>
<td><strong>Intravenous</strong></td>
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<tr>
<td>0.3–0.5 mg/kg bolus</td>
<td>1–4.5 mg/kg bolus</td>
</tr>
<tr>
<td>0.1–0.2 mg/kg/hour</td>
<td>0.5–4.5 mg/kg/hour</td>
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<tr>
<td><strong>Oral</strong></td>
<td></td>
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<tr>
<td>10 mg PO TID</td>
<td></td>
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<tr>
<td>1 mg/kg divided in equal doses Q8H</td>
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</tbody>
</table>
Ketamine

• Who can benefit?
  – Surgical patients with expected SEVERE post-operative pain
  – Opioid tolerant or dependent patients with exacerbation of a chronic condition or requiring surgery
  – Patients at increased risk for opioid-related respiratory depression

• Contraindications
  – Poorly controlled cardiovascular disease
  – Severe hepatic dysfunction
  – Poorly controlled psychiatric conditions with psychosis
Lidocaine Infusion

• Mechanism of Analgesia
  – Attenuates nociceptive sensitization via sodium channel blocking
  – Decreases the NMDA receptor mediated post-synaptic depolarization
  – Proposed that ischemic nerves are more affected than non-stimulated nerves

• Analgesic, anti-inflammatory, and anti-hyperalgesic effects

• Dosing regimens vary
  – 1-2mg/kg bolus followed by 0.5-3mg/kg/hr
  – 2-5mg/kg(150mg) IVPB x 1 over 2 hours
Lidocaine Infusion

• Who makes a good candidate?
  – Opioid-refractory nociceptive pain
  – Central and peripheral neuropathic pain
  – Increasing opioid requirements

• Relative contraindications
  – History of arrhythmia, ischemic disease, syncope
  – 1\textsuperscript{st} and 2\textsuperscript{nd} degree heart block
  – Use of alpha agonists or beta blockers
  – Congestive heart failure
Lidocaine Infusion

• Monitoring
  – 1st order kinetics, level proportional to infusion dose
  – Efficacy measured by patient reported and any decrease in MME
  – Side effects and early signs of toxicity via patient report

• Concern for toxicity in patients, especially renal or cardiac failure
  – CNS symptoms tongue numbness->metallic taste->light-headedness->visual changes->mental status changes->coma->respiratory arrest
  – Continuous cardiac monitoring recommended with this patient population
Putting It All Together

Apply what you’ve learned
Patient Case-Methadone

• 38 yo female presents with fever, severe L arm pain secondary to multiple abscesses. Her abscesses are going to require I&D and she is concerned with her pain control and is extremely fearful of withdrawal. She has a history of OUD, but has been in remission for the past 2 years and is being managed by the local methadone clinic. Her current MOUD is methadone 120 mg PO Qday. She usually will take acetaminophen as needed for any minor aches and pain. She requests not to be started on oxycodone, as she reports this was her drug of choice when she was using.

• What recommendations would you make to the hospitalist to control this patient’s pain?
Patient Case: Buprenorphine

• 57 y.o. male presented to your hospital for a TKA. He has a history of HTN, DM2 and substance use disorder. His SUD is currently being managed with buprenorphine-naloxone sublingual tablets 8-2mg SL twice daily. He is anxious about controlling his pain after surgery.

• What recommendations would you make to the surgery team in order to control this patient’s pain?
Patient Case: Naltrexone

• 43 yo male presents as a level 2 trauma after an MVC. He has multiple rib fractures, a sternal fracture, and a L humerus fracture. In the trauma bay he continuously rates his pain as 10/10 and has not responded to IV hydromorphone 1mg.

• A nurse finds a medication alert necklace which shows the patient is on naltrexone. He then reports that he recently received a dose of depot naltrexone 18 days ago

• What recommendations would you provide the trauma team for the management of this patient’s pain?
Patient Case: Naltrexone

• 37 yo male admitted s/p GSW to the head. He was administered one dose of depot naltrexone 7 days prior to admission. The patient’s family would like to transition to comfort measures only due to the nature of his injuries but are worried about maintaining the patient’s comfort due to his naltrexone injection.

• What are your concerns with the date of his last naltrexone injection?
• What recommendations do you have for the team to control the patient’s pain for comfort care?
Patient Case: Naltrexone

• Limited options to manage pain and end-of-life symptoms
  – Typically rely on opioids in patients not receiving naltrexone
  – Patient is still experiencing complete mu-receptor blockage as the injection was only one week prior to admission
  – Attending teams uncomfortable with the amount of opioids theoretically required to overcome blockade
  – Management of dyspnea was also difficult due to mu-receptor blockade

• Ketamine infusion was started at 0.1 mg/kg/hr
  – What are some concerns/cautions with use of ketamine?
  – What is the maximum recommended dose for pain management?
  – Can nurses push ketamine for pain management?
Patient Case: Naltrexone

• Ketamine was titrated up to 0.4 mg/kg/hr for comfort
  – Symptoms were well controlled
  – Patient respirations were easy
  – Patient passed away peacefully

• What if we didn’t know the last date of naltrexone administration? How would this have changed our therapy plan?
• What if the patient had received naltrexone 20 days prior to admission? What alternatives could be considered?
References

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PCSS Mentoring Program

- PCSS Mentor Program is designed to offer general information to clinicians about evidence-based clinical practices in prescribing medications for opioid addiction.

- PCSS Mentors are a national network of providers with expertise in addictions, pain, evidence-based treatment including medication-assisted treatment.

  - 3-tiered approach allows every mentor/mentee relationship to be unique and catered to the specific needs of the mentee.

  - No cost.

For more information visit: pcssnow.org/mentoring
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A simple and direct way to receive an answer related to medication-assisted treatment. Designed to provide a prompt response to simple practice-related questions.

Ask Now
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<td>American Medical Association</td>
<td>Southeastern Consortium for Substance Abuse Training</td>
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