

BUPRENORPHINE DOSE LIMITS: WHAT IS THE EVIDENCE?

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UNIVERSITY OF WASHINGTON PSYCHIATRY AND ADDICTION CASE CONFERENCE MAY 12, 2022





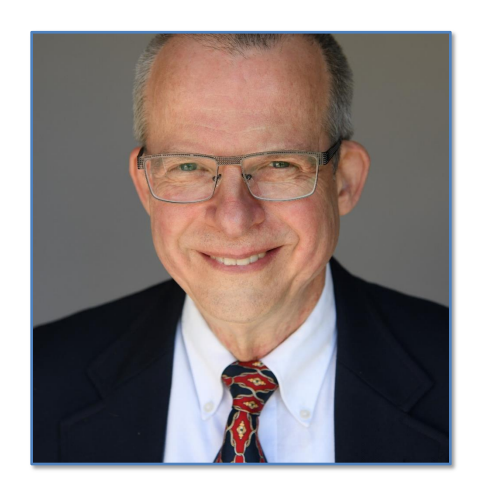


Buprenorphine Dose Limits: What is the Evidence?

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- No Financial Disclosures
- Will discuss off-label use
- May mention brand names in discussion



Buprenorphine Dose Limits: What is the Evidence?

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Buprenorphine Dose Limits: What is the Evidence?

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Buprenorphine Dose Limits: What is the Evidence?

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Professor of Obstetrics, Gynecology, and Reproductive Sciences, University of California at San Francisco

- Royalties from book: "Opioid Use
 Disorders in Pregnancy,"
 Cambridge Univ. Press, 2018
- Consulting income, McKesson, regarding neonatal withdrawal.
- Will discuss off-label use
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LEARNING OBJECTIVES

Buprenorphine Dose Limits: What is the Evidence?

Attendees will be able to:

- Discuss history and status of buprenorphine dose limits
- Discuss evidence for improved outcomes at higher buprenorphine doses
- Cite criteria for determining buprenorphine dose adequacy
- Identify special considerations for pregnant people taking buprenorphine for opioid use disorders

ACKNOWLEDGEMENTS

- Michael M. Miller, MD, DFASAM
- Richard K. Ries, MD
- Martin M. Klos, MD, MBA
- Olympia Bupe Clinic at Capital Recovery Center
- Our patients!

ISSUES WITH DOSE LIMITS

- Prior authorizations sometimes required for doses higher than 16/4 mg or 24/6 mg per day
- Pharmacy calls: Insurer will not pay for more than 2 of any film dose (may have 8x2=16 + 12 + 4 to get dose of 32 mg)
- Many Buprenorphine prescribers refuse to prescribe doses higher than 16 mg per day
- Limits referral options for patients discharging from residential treatment, especially those in rural areas!

CASE STUDY: 36-YEAR-OLD JOE

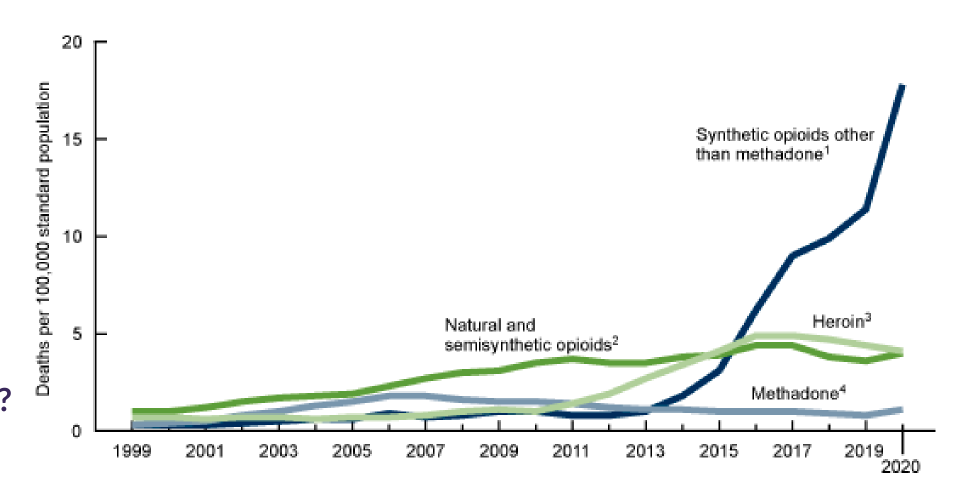
- 20 years IVDU: heroin
- Did not complete high school
- Intermittent laborer in local industry
- And then: Buprenorphine/Naloxone
 - Dose titrated upward from 8/2 Q 12 hours
 - At 32/8 able to work intermittently but always felt sick
 - At 36/9 worked full time; not sick; primary wage earner

CASE STUDY: 28-YEAR-OLD JACKIE

- SUD counselor doing intake: "You need to see her NOW!"
- Cellulitis bilateral forearms, rural ED visit yesterday
- Went to different ED on my insistence
- Airlifted to population center 2+ hours away
- 4+ weeks of IV antibiotics; osteomyelitis in knee
- Phoned me in panic: Resident MDs tell me to get off bupe!
- Outcome: stable on 16/4 Q 12 hours; gait resolved in 6 mo
- Until talked about bupe/nx at NA meeting...

WHY SHOULD WE CARE ABOUT BUPRENORPHINE DOSE?

Other medication options often not realistically available **Providing** effective treatment is a matter of life and death Fentanyl 2013 to?



MEASURES OF OUD TREATMENT SUCCESS

- Death rates
- Treatment retention
- Visit reliability ("on-time visits")
- Abstinence from non-prescribed opioids
- Abstinence from other illicit drugs
- Short-term clinical goals or "therapeutic targets"
- Long-term clinical goals or "life goals"

ARGUMENTS FOR AND AGAINST DOSE LIMITS

FOR:

- FDA package insert
- Receptor occupancy interpretations (2000-2009)
- Concerns about cost
- Concerns about diversion
- "Expert opinion"

AGAINST:

- **ASAM** guidelines
- Individual variability
- Receptor occupancy data (2010-present)
- Improved treatment retention
- Reduced illicit drug Use
- No analgesic ceiling effect
- Kappa receptor role

FDA BUPRENORPHINE-NALOXONE PACKAGE INSERT

- Dosing recommendations are based on data from trials before 2002 at doses equivalent to 6-24 mg/day.
- The recommended daily dose for maintenance is 16/4 mg.
- The maintenance dose "is generally in the range of 4/1 mg buprenorphine/naloxone to 24/6 mg buprenorphine/naloxone per day depending on the individual patient. Dosages higher than this have not been demonstrated to provide any clinical advantage."

ASAM 2020 GUIDELINES: TREATMENT GOALS

- 1. Suppress opioid withdrawal
- 2. Block the effects of illicit opioids
- 3. Reduce opioid craving and stop or reduce the use of illicit opioids
- 4. Promote and facilitate patient engagement in recoveryoriented activities including psychosocial intervention

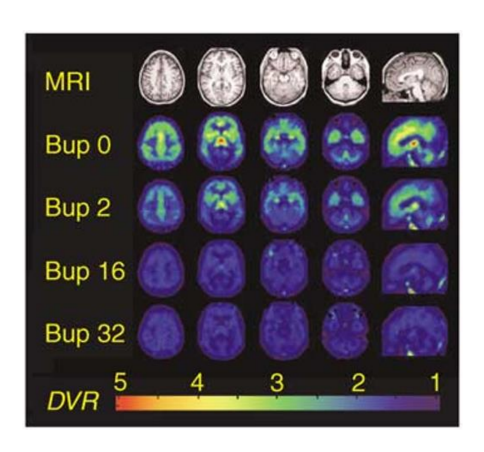
IS MY PATIENT MEETING ASAM TREATMENT GOALS?

- 1. Any cravings, any time of day or night?
- 2. Any withdrawal symptoms, any time of day or night?
- 3. Any night sweats?
- 4. Any "using dreams"?
- 5. Any use of any opioid that isn't prescribed for you and known to me?
- These questions address short-term, buprenorphine-specific issues
- Additional targets depend on relationships and mutual goals
- Test urine/saliva etc. when clinically appropriate/required

BUPRENORPHINE DOSE LIMITS: WHAT IS THE EVIDENCE?

Dose-Response Observations in Physiology & Pharmacology Studies

MU RECEPTOR OCCUPANCY (2003)



- Five heroin-dependent volunteers [no fentanyl!]
- Mu receptor occupancy (relative to placebo):

-2 mg/day: 27-47%

-16 mg/day: 80-92%

-32 mg/day: 89-98%

MU RECEPTOR OCCUPANCY CONCLUSIONS

- Buprenorphine dose-dependently increased mu receptor occupancy
- High receptor occupancy correlated with improved therapeutic effect: decreased opioid withdrawal and reward symptoms.
- Near-maximal effect occurred at 32 mg/day

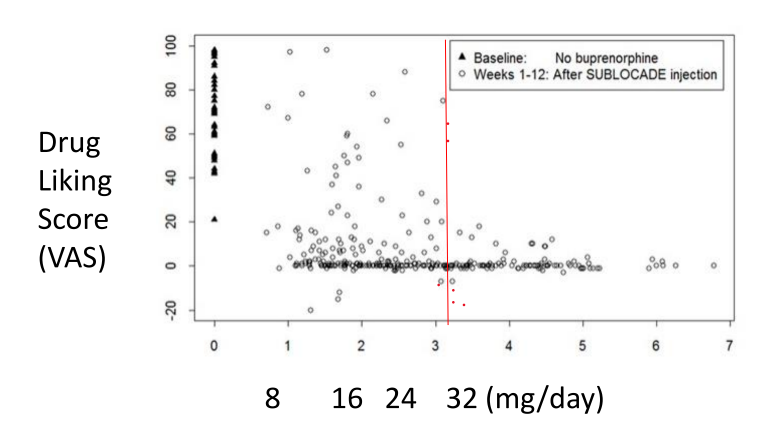
MU RECEPTOR OCCUPANCY REVIEW (2014)

- Withdrawal suppression requires >50% receptor occupancy (4 mg/day)
- Blockade of subjective opioid effects requires >80% occupancy (16 mg/day)
- Blockade of subjective effects of high opioid doses may require >90% occupancy (up to 32 mg/day)

MU RECEPTOR OCCUPANCY REVIEW (2014)

We conclude that fixed, arbitrary limits on buprenorphine doses in clinical care or limits on reimbursement for this care are unwarranted.

OPIOID REWARD SUPPRESSION > 24 MG/D



Exposure:

High-dose hydromorphone (18 mg IM)

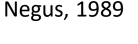
Plasma Concentration (avg ng/mL) Equivalent SL dose avg

ROLE OF KAPPA RECEPTORS

- Kappa receptor activation mediates dysphoria, anhedonia and aversion in animal models
- May contribute to negative affect in addiction
- Buprenorphine produces a dose-dependent kappa receptor antagonist effect up to 1 mg/kg in rodents

ROLE OF KAPPA RECEPTORS

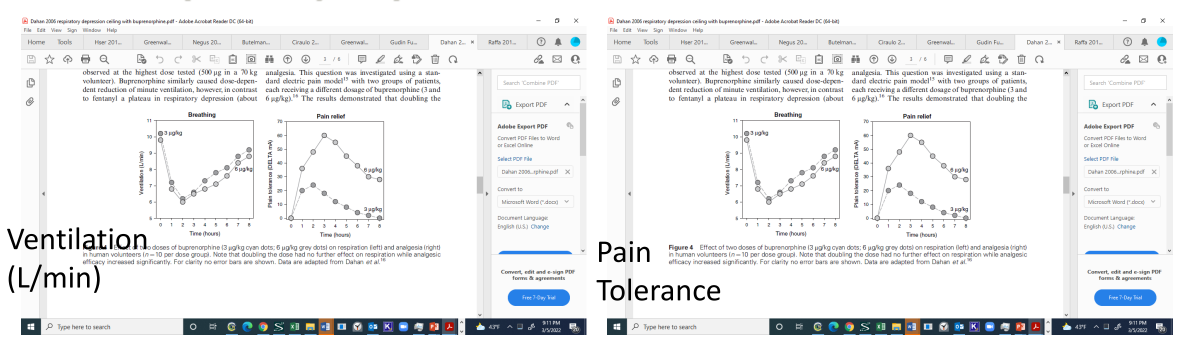
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HIGHER SERUM LEVELS: ARE THERE CEILING EFFECTS? YES ... but NOT

for respiratory depression



for analgesia

IS THERE ANY CEILING EFFECT FOR PAIN?

- Primary data above from Dahan, 2006 (0.2-0.4 mg IV per 70 kg)
- Lots of "expert opinion" says there's NO ceiling effect for pain
- How high do the experimental data go?
- Four loci of receptor activity (Gudin, 2020):
- Mu receptor partial agonism (analgesia, mood)
- Delta receptor antagonism (limits GI & respiratory depression)
- Kappa receptor antagonist (limits depression, dysphoria, suicidal tendencies)
- ORL-1 receptor agonist (inc. spinal analgesia, dec. supraspinal analgesia, dec. reward, dec. tolerance)

Higher Doses Work Better: Evidence From Clinical Research

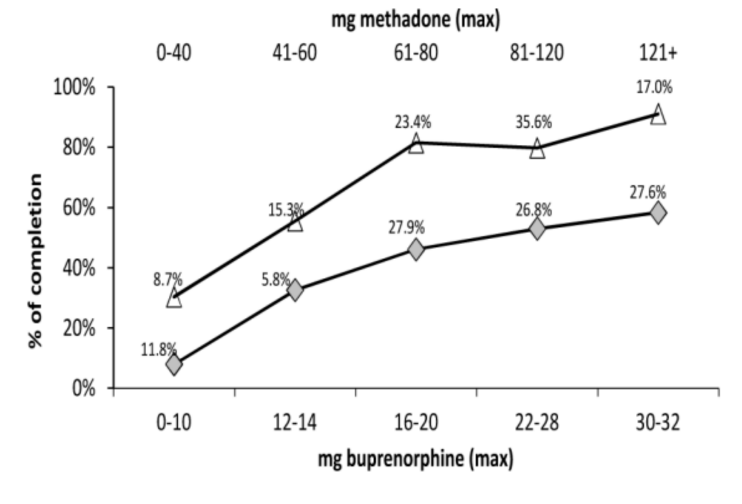
PRACTICAL MEASURES OF TREATMENT SUCCESS

- Improved treatment retention
- Reduced illicit drug use
- Patient perception of adequate dose
- Reduce complications
- Reduced pain
- Improved visit reliability

TREATMENT RETENTION

- Open-label RCT, N=1,267 at 9 OTPs in the U.S.
- Buprenorphine (n=738) vs. methadone (n=529)
- Daily observed dosing for both groups

TREATMENT RETENTION RESULTS



- → Buprenorphine (% = % of buprenorphine participants prescribed in that dose range)
- → Methadone (% = % of methadone participants prescribed in that dose range)

TREATMENT RETENTION: CONCLUSIONS

- Linear relationship between dose and treatment completion rate.
- 32 mg/day or even higher may be needed for optimal benefit for some patients.

META-ANALYSIS: RETENTION & DRUG USE

- 21 RCTs, international
- 2,703 participants

- High dose group (16-32 mg/day):
 - -Better retention in treatment than low dose group
 - -Fewer urine tests positive for opiates and cocaine

PATIENT PERCEPTION #1: PARAMETERS

- 48 outpatients, Sweden
- Heroin dependence, average duration 9.4 years
- Bupe/nx dose increased as needed up to 32 mg/day, then switched to methadone if needed

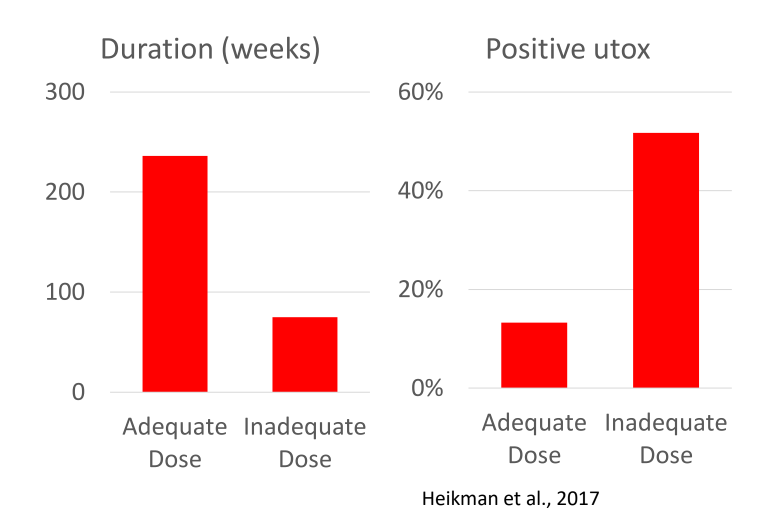
PATIENT PERCEPTION #1: RESULTS

- Out of 48 patients:
 - -17 remained on buprenorphine
 - Mean final dose 29.6 mg/day
 - -20 switched to methadone
 - No difference between groups in retention or urine test results
 - -11 dropouts

PATIENT PERCEPTION #2: PARAMETERS

- Addiction Psychiatry Outpatient Clinic, Finland
- Observational study
 - Methadone 65 mg/day (avg) (n=52)
 - ❖Bupe-nx 15 mg/day (avg) (n=8)
- Inadequate Dose (n=39) vs. Adequate Dose (n=21), by patient self-report

PATIENT PERCEPTION #2: RESULTS



Adequate Dose:

- Improved retention
- Reduced illicit drug use

PATIENT PERCEPTION: CHRONIC PAIN

- Retrospective study, 35 patients with chronic pain on high-dose opioids (>200 MME)
- Transitioned to buprenorphine SL, allowed up to 32 mg/day
- Average pain score dropped from 7.2 to 3.5
- Average buprenorphine dose: 28.1 ± 5.9 mg.

REDUCED COMPLICATIONS (HEPATITIS C)

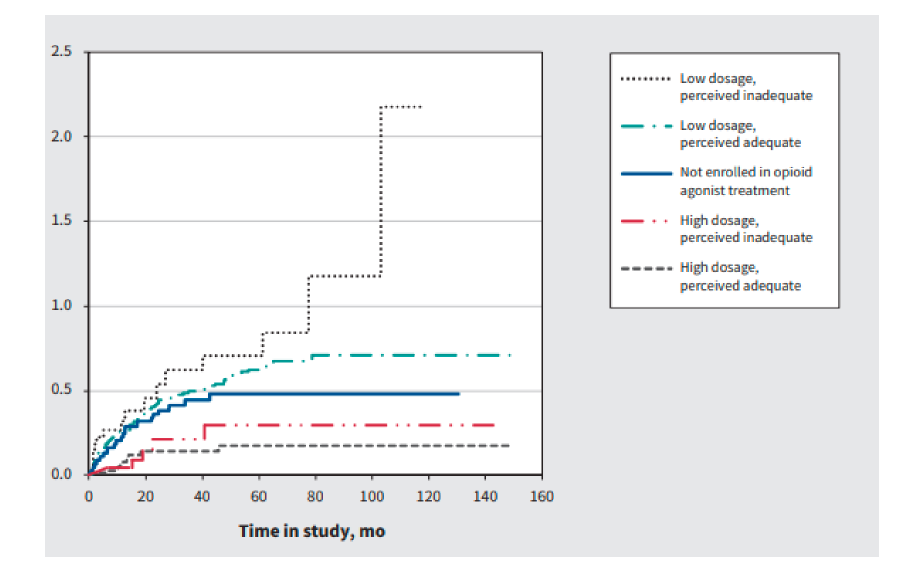
- 513 HCV-negative people who inject, Canada
- Exposure 2.8 years (average)
- Comparisons:
 - Medication Dose (high vs. low vs. none)
 - Buprenorphine high dose: >=16 mg/day
 - Methadone high dose: >=60 mg/day
 - Patient perception:
 - Adequate vs. inadequate, by self-report

REDUCED COMPLICATIONS (HEPATITIS C)

Hazard Ratio

Highest Risk: Low dose, Inadequate

Lowest risk:
High dose, Adequate



HIGH DOSES AT A LOW THRESHOLD CLINIC

- Olympia Bupe Clinic (OBC), Olympia Washington
- >1500 patients treated
- Walk-in only, no cost, medication dispensed on-site
- Team-based care, peer recovery coaches, nurse care manager
- Buprenorphine dosing to clinical effect

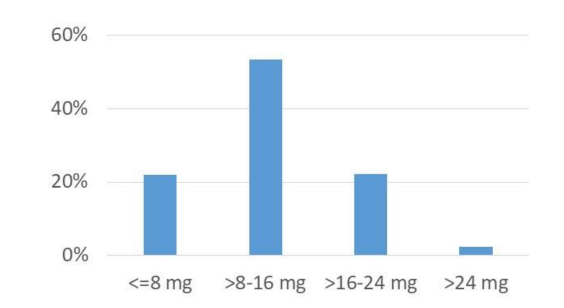
QUALITY IMPROVEMENT PROJECT

- Hypothesis: In this population, engagement consistency increases with dose up to 32 mg/day
- Data: Prescription Monitoring Program (2018-2021), Intake Database
- Measure of consistent engagement: on-time visits (+/- 1 day)
- Comparison: Prescriptions by OBC vs. non-OBC prescribers for the same patients

DISTRIBUTION OF DOSE RANGES

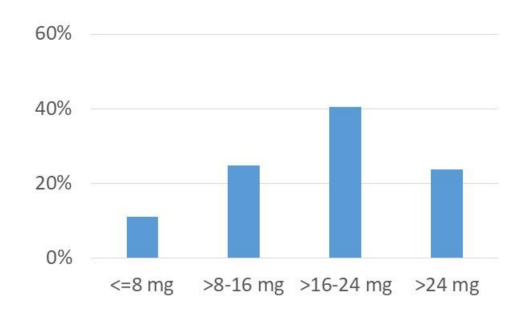
Non-OBC Prescribers

% of Rxs



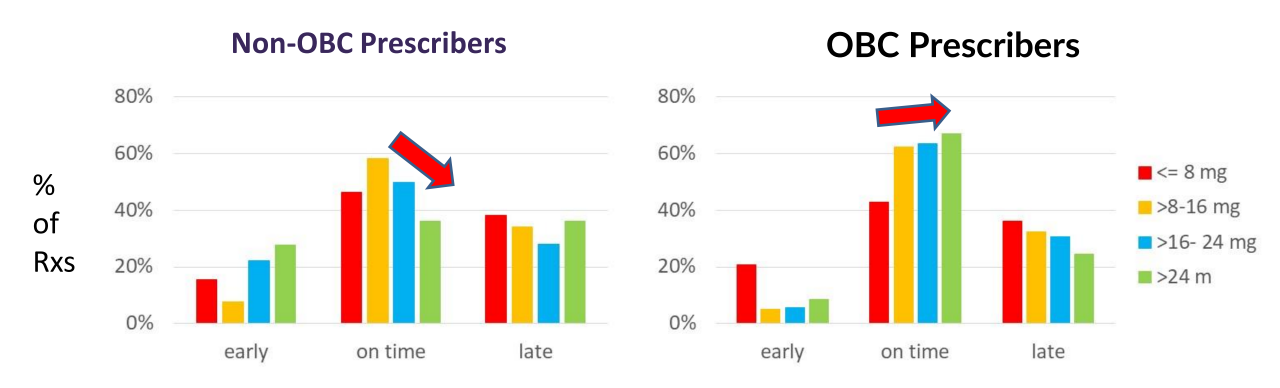
2241 Rxs, 127 patients

OBC Prescribers



3093 Rxs, 242 patients

TIMELINESS OF FOLLOW-UP VISITS

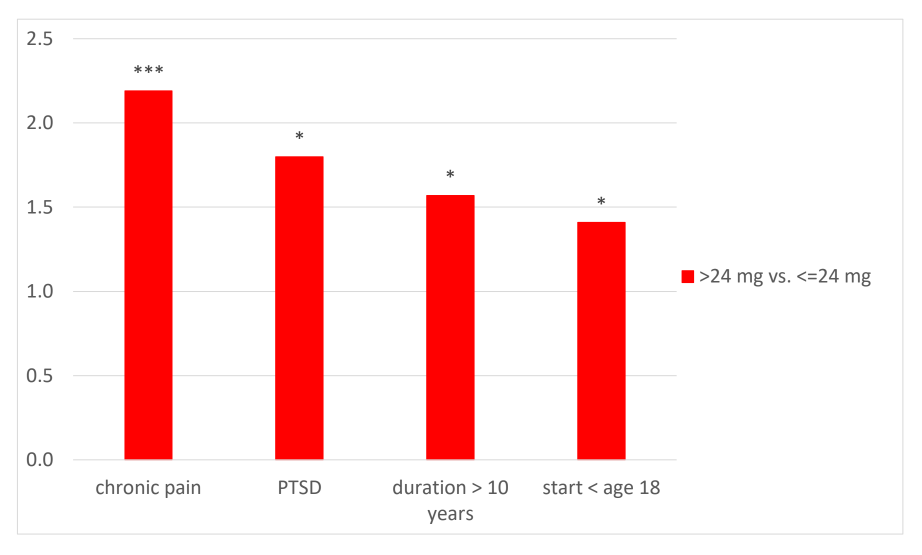


OBC on-time performance better overall OBC on-time performance improves at higher doses Why the difference?

OBC, unpublished data

Relationship of Dose With Chronic Pain, PTSD & Opioid Tolerance

Relative Risk *** p =0.0004 * p < 0.05



OBC, unpublished data

Buprenorphine dosing in pregnancy: Why 8 mg BID doesn't cut it.

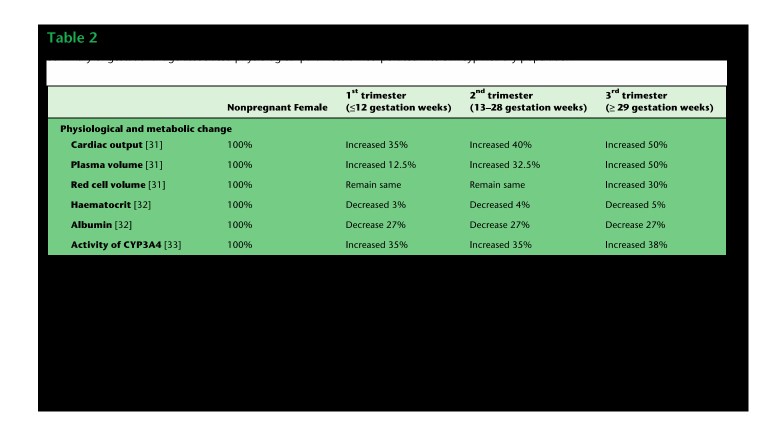
CASE EXAMPLE

- 27 y/o pregnant female at 10 weeks gestation
- On buprenorphine 16 mg/day
- Reports increased withdrawal symptoms and cravings
- At 16 weeks, dose increased to 24 mg/day
- At 24 weeks, dose increased to 32 mg/day
- At 32 weeks, requests further increase

CASE EXAMPLE

- ❖Is she diverting?
- What are the effects of pregnancy on withdrawal symptoms?
- What are the effects of buprenorphine dose on baby?

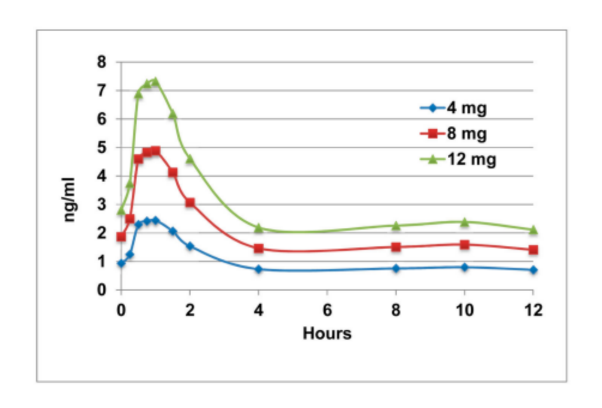
REVIEW OF PREGNANCY PHYSIOLOGY

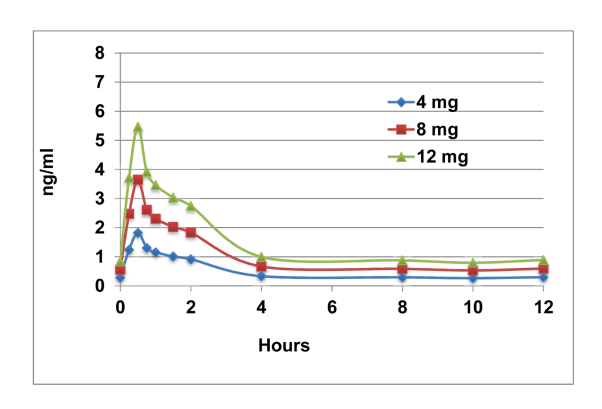


PREGNANT WOMEN NEED HIGHER DOSES

- Increased volume of distribution
- Increased clearance
- Decreased salivary pH which may decrease absorption
- Decreased time to peak concentration
- Decreased time to trough
 - Cause of nausea?
 - Relieved by split dosing?
- Plasma concentrations ~50% lower

DECREASED SERUM CONCENTRATION DECREASED TIME TO TROUGH

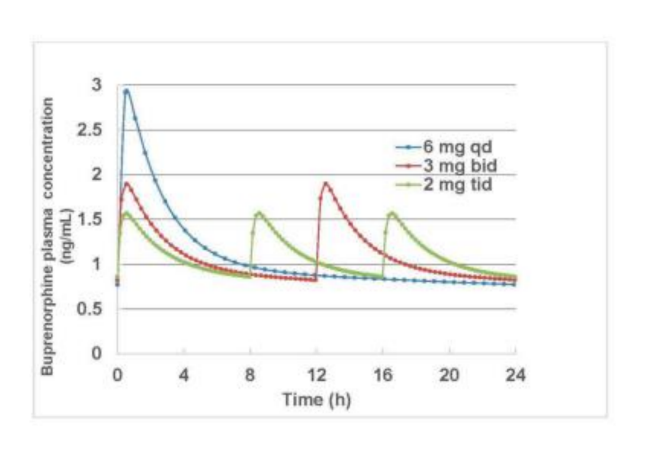




Physiologic Baseline (Postpartum)

2rd Trimester

PREGNANT WOMEN NEED MORE FREQUENT DOSING



1 ng/mL = threshold of withdrawal symptoms

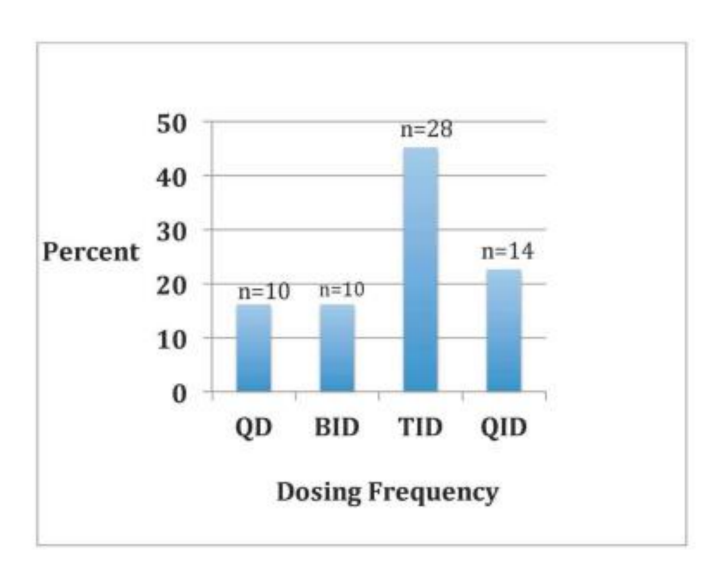
More frequent dosing results in less total time below 1 ng/mL

Once daily: 16.3h
BID: 14.4h
TID: 10.8h

Computer model of serum concentration at different dosing frequencies

Caritis et al 2017

SELF-DETERMINED DOSE FREQUENCY

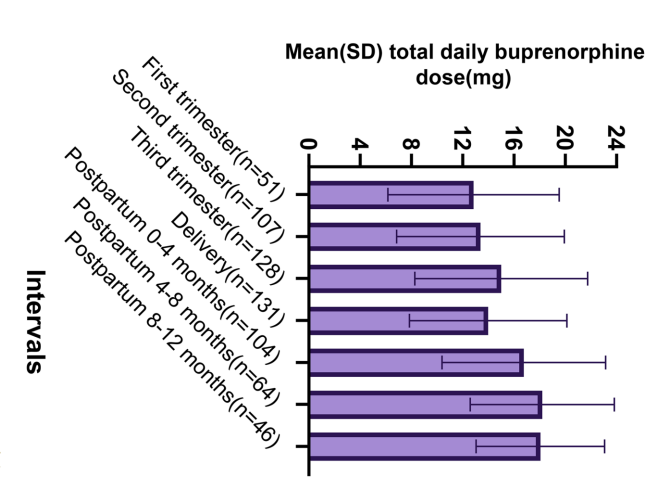


 Residential treatment facility for pregnant women.

Most choose TID dosing.

INDIVIDUAL DOSING IN PREGNANCY IS NEEDED

- Martin et al (2020)
 - Literature Review=25 studies
 - Mean dose change -12.3 to 10.5 mg/day
 - 2 studies dose retention
 - Prospective cohort study
 - Mean dose steadily increased
 - Maintained postpartum
 - Less split dosing postpartum
 - VA Medicaid limits dosing to 24 mg



BUPRENORPHINE DOSING AND NAS/NOWS

- No relationship between dose and risk of NAS/NOWS
- Positive relationship between meconium buprenorphine concentration and risk of NAS/NOWS

TAKE-AWAYS

- Buprenorphine dose is adequate if:
 - Patient perceives it as adequate, AND
 - Reduced illicit opioid use
 - Increased retention
- Clinical and preclinical evidence that 32 mg/day or higher can be helpful for some patients, particularly those with pain
- Pregnant women need:
 - Increased dose of buprenorphine
 - Increased frequency of dosing

CONTACT INFORMATION

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REPRISE: ARGUMENTS FOR AND AGAINST DOSE LIMITS

FOR:

- FDA package insert
- Receptor occupancy data

(2000-2009)

- Concerns about cost
- Concerns about diversion
- "Expert opinion"

AGAINST:

- **ASAM** guidelines
- Individual variability
- Receptor occupancy data (2010-present)
- Improved treatment retention
- Reduced illicit drug Use
- No analgesic ceiling effect
- Kappa receptor role

PATIENT INSTRUCTIONS: MAKE THE DOSE EFFECTIVE!

- Stay well hydrated so films or tablets will dissolve easily.
- No nicotine use during 20-30 minutes prior to dose.
- Place film or tablet under tongue and tuck chin to chest.
- Do not swallow excess saliva (may precipitate withdrawal symptoms). Spit out excess saliva!